Zn-Cu Alloy Nanofoams as Efficient Catalysts for the Reduction of CO₂ to Syngas Mixtures with a Potential-Independent H₂/CO Ratio
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Zn-Cu alloy nanofoams as efficient catalysts for CO\textsubscript{2} reduction to syngas mixtures with potential-independent H\textsubscript{2}:CO ratio

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Abstract: Alloying strategies are commonly used to design electrocatalysts that take on properties of their constituent elements. Herein, we explore the use of such a strategy to develop Zn-Cu alloyed electrodes with unique hierarchical porosity and tunable selectivity for CO\textsubscript{2} vs. H\textsuperscript{+} reduction. By varying the Zn:Cu ratio, tailored syngas mixtures were attained with no other gaseous products, which we assign to preferential CO and H\textsubscript{2} forming pathways on the alloys. The syngas ratios were also significantly less sensitive to the applied potential in the alloys relative to pure metal equivalents; an essential quality when coupling electrocatalysis to renewable power sources of fluctuating intensity. As such, industrially-relevant syngas ratios were achieved at large currents (\textasciitilde\textasciitilde\textasciitilde 60 mA) for extensive operating times (> 9 h), demonstrating the potential of this strategy for fossil-free fuel production.

Introduction

Electroreduction of CO\textsubscript{2} into energy-dense compounds such as carbon monoxide, formic acid, hydrocarbons and alcohols offers a promising route to store intermittent renewable energies. Multi-carbon products (ethanol, ethylene, propanol etc.) are the most valued outcome of such a process, representing a denser store of chemical energy. However, the multi-electronic and multi-protonic nature of the reactions at work commonly results in very high overpotential, low faradaic efficiency and complex product mixtures. This process is commonly carried out in aqueous media, with the anodic oxidation of water providing the source of protons and electrons. Yet, the use of aqueous electrolyte is highly challenging considering the kinetically and thermodynamically facile alternative reduction of H\textsuperscript{+} to H\textsubscript{2}. While significant efforts are currently being made to develop electrocatalysts promoting the direct CO\textsubscript{2} reduction to hydrocarbons or alcohols with limited activity for H\textsuperscript{+} reduction, an equally attractive strategy is to take advantage of the produced H\textsubscript{2} to generate valuable gas mixtures, such as syngas, a combination of CO and H\textsubscript{2}. Syngas can be used to produce hydrocarbons and alcohols through well-established industrial technologies. A two-step process that couples CO\textsubscript{2} electroreduction to syngas with its subsequent transformation to high added-value products has been proposed to be more favorable from an economic perspective since i) CO\textsubscript{2} reduction to CO occurs at moderate overpotentials and ii) a highly energy-demanding product separation step is not required.\textsuperscript{[1]} However, one of the main prerequisites for the syngas generated is control over the H\textsubscript{2}:CO ratio, that must meet different values depending on the reaction targeted: optimal H\textsubscript{2}:CO ratios of 1.5-2.2 are typically required for methanol synthesis and the Fischer-Tropsch reaction; 3 for the methanation reaction; and 1 is typically required for hydroformylation and fine chemical synthesis.\textsuperscript{[1]}

Fossil-fuel reforming reactions are currently the largest source of syngas; however these processes generate specific H\textsubscript{2}:CO ratios (Figure 1) and are based on non-renewable feedstocks. Adjusting these ratios requires an additional energy-demanding water-gas-shift reaction.\textsuperscript{[1]} In this context, electrochemical syngas generation is a more versatile and sustainable alternative, allowing a broad range of H\textsubscript{2}:CO ratios to be produced from renewable precursors (H\textsubscript{2}O and CO\textsubscript{2}).\textsuperscript{[2, 3]} The electrochemical conversion of CO\textsubscript{2} to syngas is particularly relevant in the context of renewable electricity conversion, which requires the design of electrolytic devices tolerant to the significant variations of power provided by intermittent energy sources, such as photovoltaic panels. Nevertheless, to the best of our knowledge all the electrocatalytic systems for syngas generation present a significant variation of the H\textsubscript{2}:CO ratio with applied potential, preventing an efficient coupling with such sources of electricity.

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Supporting information for this article is given via a link at the end of the document.
An effective syngas generating system must therefore meet several key parameters:

1. The modulation of the H₂:CO ratio must not depend on the potential applied; materials that can independently fine-tune the transfer coefficients for the formation of H₂ and CO must be chosen.

2. The system must operate in a regime where the current is not limited by the diffusion of reactants; CO₂ is present in a very low concentration in aqueous solutions and therefore a system maximizing reactant mass transfer is mandatory.

3. The chosen system must be selective for syngas production only, i.e. the typical concomitant formation of formic acid must be minimized.

Previous reports have focused on monometallic molecular and heterogeneous catalytic systems, showing simultaneous CO and H₂ production, but single catalytic sites offer limited possibilities to tune catalyst selectivity. Introducing a second metal site offers an extra degree of freedom in catalyst design and thereby offers the best route to address the above criteria. By combining two metals, one with a high propensity for CO₂ reduction and another for proton reduction, ratios of syngas can easily be attained by altering the ratio of the constituent metals.

Herein, we present a series of high-surface area dendritic foams comprised of Zn alloyed with varied amounts of Cu. The foams are generated through a facile co-electrodeposition of Zn with sub-stoichiometric amounts of Cu. The Cu-doping first plays a major role in the growth of the catalysts; Cu salts in the deposition precursor solution triggers the growth of high surface area three-dimensional porous dendritic materials via a hydrogen-evolution-assisted electrodeposition approach, not possible with Zn alone. The incorporated Cu then presents sites for hydrogen evolution in the material such that H₂:CO ratios are found to correlate directly to the Cu:Zn ratio in the catalyst. The resultant porous structures also allow fast diffusion of species to the micropores, keeping the H₂:CO ratio constant in a wide potential range by preventing mass transfer limitations. Syngas production could thus be achieved at currents up to ~45 mA at ~1.2 V vs. RHE using 1 cm² flat-Zn supported catalysts. This strategy could be extended to a 1 cm² Zn-foam-supported catalyst, which exhibited syngas production over multiple hours, reaching up to ~60 mA at ~1.1 V vs. RHE.

**Results and discussion**

**Zn:Cu Alloy Growth and Characterization**

Dendritic Zn:Cu foam electrodes were generated by applying high current density (0.5 A.cm⁻²) to 1 cm² Zn plates immersed in a 0.5 M H₂SO₄ aqueous solution of metal precursors. The total metal salt concentration was kept constant and equal to 0.1 M but distributed between 5% of ‘doping’ CuSO₄ and (100−X)% ZnSO₄ with X varying between 0%, 5%, 10%, 15%, 20%, 25%, and 35%. These electrodes are labelled Zn│ZnSO₄(100−X)%CuSO₄X% in the following sections. Scanning electron microscopy (SEM) reveals that the presence of CuSO₄ in the precursor mixture, even below 5%, promotes high nano- and meso-porosity within the Zn-based ‘fern-like’ structures, which otherwise grow in a low-surface-area stacked configuration (Figure 2 and Figure S1).

![Figure 1. H₂:CO ratios reachable using fossil-fuel-based techniques versus the CO₂ electroreduction (CO₂ER) process presented in this work. Data for fossil-fuel-based processes are taken from ref [1]. Primary syngas ratio refers to the obtained products without additional gas-shift reactions to readjust the ratio.](image1)

![Figure 2. SEM images of Zn│ZnSO₄(100−X)%CuSO₄X%, where percentage of CuSO₄ (X) is 0, 5, 10, 20, 25 and 30, as indicated in the top-left corner of corresponding images.](image2)
marginally increases at higher doping levels. All alloys showed enhanced porosity, confirmed through BET measurements (Table 1); the electrodes prepared with the highest Cu doping display the highest surface area. Areas as high as 27.4 m² g⁻¹ were reached, corresponding to a roughness factor (RF) of ca. 25 once normalized by the mass of electrodeposited material. This marks a substantial improvement over the RF = 1.12 of bulk Zn foil previously recorded using atomic force microscopy. [20]

Elemental composition of the electrodeposited alloys was probed by energy-dispersive X-ray spectroscopy (EDS) and inductively coupled plasma - atomic emission spectroscopy (ICP-AES) measurements (Table 1 and Figure S3-S9). The incorporation of Cu in the materials was confirmed at each loading investigated, and ICP-AES measurements revealed that the Cu:Zn ratio in the material is typically 1.5 times higher than the Cu:Zn ratio present in the metal sulfate precursor solution (Table 1), in agreement with the thermodynamically preferential reduction of Cu⁺ vs. Zn²⁺. The bulk Cu content determined for each electrode hence differs from the Cu content 'X' used for the naming Zn│ZnSO₄(100-X%)CuSO₄:X% of the electrodes which refers to the content in CuSO₄ salt initially introduced in the precursor solution.

<table>
<thead>
<tr>
<th>Electrode</th>
<th>%CuSO₄</th>
<th>% Cu (bulk)</th>
<th>% Cu (surface)</th>
<th>BET [m² g⁻¹]</th>
<th>RF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn│ZnSO₄(5%)CuSO₄(95%)</td>
<td>5</td>
<td>6.5±0.5</td>
<td>1</td>
<td>1.3</td>
<td>1.3</td>
</tr>
<tr>
<td>Zn│ZnSO₄(10%)CuSO₄(90%)</td>
<td>10</td>
<td>13±1</td>
<td>2</td>
<td>3.8</td>
<td>3.5</td>
</tr>
<tr>
<td>Zn│ZnSO₄(20%)CuSO₄(80%)</td>
<td>20</td>
<td>32±2</td>
<td>n/d</td>
<td>16.9</td>
<td>15</td>
</tr>
<tr>
<td>Zn│ZnSO₄(25%)CuSO₄(75%)</td>
<td>25</td>
<td>39.5±4</td>
<td>n/d</td>
<td>14.6</td>
<td>13</td>
</tr>
<tr>
<td>Zn│ZnSO₄(30%)CuSO₄(70%)</td>
<td>30</td>
<td>59.3±6</td>
<td>12</td>
<td>27.4</td>
<td>25</td>
</tr>
</tbody>
</table>

Table 1. Relationship between the CuSO₄ percentage in the precursor solution, the subsequent percentage of Cu incorporated in the electrode (bulk, determined by ICP-AES), at its surface (determined by XPS) and associated BET surface area and roughness factor (RF).

To gain deeper insight into the bulk and surface composition and mixing pattern of the deposits, known to have a significant influence on the catalyst selectivity[19], the electrode materials were analyzed using X-ray diffraction spectroscopy (XRD), X-ray Photoelectron Spectroscopy (XPS) and high-resolution transmission electron microscopy (HR-TEM). Diffractograms for Zn│ZnSO₄(100-X%)CuSO₄:X% electrodes confirm the strong correlation of alloy content and the relative metal stoichiometry of the soluble precursors (Figure 3). At low loading (X ≤ 10), the XRD patterns of the corresponding Zn│ZnSO₄(100-X%)CuSO₄:X% structures show hexagonal-Zn phases with slight variations of lattice parameters, suggesting a solid solution of Cu in these materials, as well as additional hexagonal phases attributed to a Cu₆Zn₉ alloy. With increasing Cu content (10 < X ≤ 30), diffraction patterns attributed to cubic Cu₂Zn₈ and Cu₃Zn₄ are observed. In some samples, further analysis at low diffraction angles revealed the presence of a low amount of namuwite (Zn₃₃Cu₆Zn₆S₉O₅(OH)₂(H₂O)₄) at the surface of the electrodes (Figure S10). This insoluble sulfate probably stays at the surface of the electrode after synthesis. It however disappears during electrolysis, as exemplified by the comparison of Zn│ZnSO₄(75%)CuSO₄(25%) XRD patterns before and after 3 hours of electrolysis (Figure S10), without affecting the morphology of the electrode, as corroborated by a drop in S content measured in SEM-EDX (Figure S3-S9).

![Figure 3. Powder X-ray diffraction patterns of Zn│ZnSO₄(100-X%)CuSO₄:X% where percentage of CuSO₄ (X) is varied between 0 and 100.](image)

X-ray Photoelectron Spectroscopy (XPS) of Zn│ZnSO₄(100-X%)CuSO₄:X% electrodes confirms that both Cu and Zn are present at the surface of the electrode at low and high Cu loading (Table 1). The quantification of the relative Cu:Zn surface ratio indicates that the surface Cu content is around 6 times lower than the bulk Cu content determined by ICP-AES. This Cu:Zn surface ratio is unchanged before and after electrolysis (Figure S11).

![Figure 4. STEM-EDX analysis of a Zn│ZnSO₄(75%)CuSO₄(25%) fern-shaped structure before and after 3 h of electrolysis at -1.0 V vs. RHE in CO₂-saturated 0.1 M CsHCO₃. The overlay (far right) shows Cu in red, Zn in green and O in blue.](image)

High Resolution TEM (HRTEM) combined with elemental mapping with Scanning Transmission Electron Microscopy-Energy-Dispersive X-ray Spectroscopy (STEM-EDX) confirmed that homogeneous distribution of Cu and Zn was present even at the nanoscale features of the dendrite as illustrated in the typical 'fern-shaped' structure in Figures 4 and S12. Comparison of the microstructures before and after use in electrocatalytic
conditions (Figure 4, S3-S12), shows no change in the atomic distribution of the metal sites, demonstrating high structural stability of this surface. A sporadic distribution of O is seen by STEM-EDXS on the surface of the structure (more visible at lower magnification, Figure S12), which is not present after electrolysis, and is thus assigned to the aforementioned namellite phase.

The considerable influence of even small quantities of Cu on the overall porosity, surface morphology, as well as its facile intercalation within the Zn structure, suggests that Cu is not only a doping agent, but also has a strong structural role. We assign this to the 1.1 V more positive reduction potential of Cu$^{2+}$ with respect to that of Zn$^{2+}$,[21] promoting more rapid growth by providing seeding sites for Zn deposition, as confirmed by the higher Cu content in the bulk of the dendrites than at their surface. Proton reduction to H$_2$ is also exacerbated by the Cu surface sites, ensuring constant H$_2$ evolution during deposition, which promotes growth of macroporous structures. We explored this preparation protocol from an industrial application perspective, which proved scalable and versatile: deposition could be applied equally well to Zn and Cu plates, as well as Zn foam. Also, homogeneous electrodes as large as 8 cm$^2$ could be obtained with no decrease in homogeneity or change in morphology (Figure S13-S14).

Electrochemical syntheses generation
Electrocatalytic activity for syntheses production of the Zn$_2$ZnSO$_4$(100-X)CuSO$_4$X$_2$ electrodes was investigated in a two-compartment H-type electrolyzer equipped with anion exchange membrane under constant CO$_2$ flow (20 mL min$^{-1}$). In order to limit the competitive H$_2$ production and explore a broader range of H$_2$:CO ratios, electrolyses were carried out in 0.1 M CsHCO$_3$ electrolyte (pH=6.8), as previous reports demonstrated that large proportions of H$_2$ in the gas phase and accounted for substantial current densities for syngas production obtained at various potentials using Zn$_2$ZnSO$_4$(100-X)CuSO$_4$X$_2$, where percentage of CuSO$_4$ (X) is 0, 5, 10, 20, 25 and 30.

The selectivity of the alloys differs from both Cu-based catalysts (typically producing multi-carbon products) and Zn-based catalysts (typically producing CO and formic acid).[24] This disparity can be rationalized according to the analysis of Rossmeisl et al.[25] In the latter, the calculated binding energies of H* and COOH* (selected as relevant intermediates in the production of H$_2$, CO and HCOOH) were demonstrated as good descriptors for the selectivity of metallic electrodes during CO$_2$ reduction. Reasoning here on the individual metals, we notice that the introduction of Cu to Zn would aid H$_2$ evolution, as Cu possesses a lower hydrogen binding energy than Zn ($\Delta E_{\text{Cu-H}} = 0.0$ eV vs. $\Delta E_{\text{Zn-H}} = +0.5$ eV). On the other hand CO evolution should not be strongly impacted by the alloying strategy, as the adsorption energy of COOH* on Zn and Cu are comparable ($\Delta E_{\text{Cu-COOH}} = +0.5$ eV compared to $\Delta E_{\text{Zn-COOH}} = +0.7$ eV). This also explains the decrease in formic acid production with increased Cu incorporation (Figure S16), as this reaction occurs on metal sites with COOH* binding energy lower than similar to H* (i.e. a positive/near zero $\Delta E_{\text{COOH}}$), which is not the case on Cu ($\Delta E_{\text{Cu-H}} - \Delta E_{\text{Cu-COOH}} = -0.5$ eV vs. $\Delta E_{\text{Zn-H}} - \Delta E_{\text{Zn-COOH}} = -0.2$ eV).[25] Multi-carbon products are similarly not seen as their formation requires stabilized bound CO* intermediates, which are disfavored by the weaker CO* binding energy of Zn in the alloy ($\Delta E_{\text{Zn-CO}} = -0.15$ eV compared to $\Delta E_{\text{Cu-CO}} = -0.5$ eV), offering a pathway for CO release.[25] Zn and Cu would therefore synergistically increase the other metal’s selectivity, by essentially ‘turning off’ their secondary reactions. Control over product selectivity is thus afforded through facile variation of the Zn:Cu content of the alloy (Figure 5a).

These findings do not corroborate with previous studies of Cu-Zn alloys that propose the use of a Zn co-catalyst as an in situ source of CO to enhance further ethanol production on Cu sites.[26] This discrepancy may be explained by the difference in the deposition regimes used; in the latter case, Cu content is substantially higher and as such contains extended phases of multiple Cu sites that likely behave similarly to bulk copper.[27]

Figure 5a presents the H$_2$:CO ratios with respect to Zn:Cu proportions in the precursor mixture and the applied potential. The corresponding partial current densities for total syngas production are shown in Figure 5b. At each potential and with each Zn-Cu material investigated, CO and H$_2$ were the sole products in the gas phase and accounted for ~80% of the total products formed in all cases (Figure S15). Formic acid was also formed, with FE at around 20% at low Cu loadings and below 2% at high loading (Figure S16). Significantly, for all
The Zn-Cu alloy composition thus provides a way to obtain syngas with a broad range of H₂:CO ratios from 0.2 to 1.6. These ratios encompass typical values obtained using fossil-fuel-derived syngas (petrol and coal steam reforming or gasification), and can be directly used for hydroformylation, the Fischer-Tropsch reaction or methanol synthesis. Higher H₂:CO ratios (up to 3.65), in the range of those required for the methanation reaction could also be obtained when using electrodes with higher Cu loading (X> 30), yet losing the invariance of the ratio with applied potential observed at lower loadings (Figure S17).

Furthermore, the electrodes showed remarkable stability over time. Stable currents and selectivity were observed over >3 h constant potential electrolysis on Zn | ZnSO₄|CuSO₄| (Figure S18). Moreover, powder XRD patterns, XPS, SEM, STEM-XEDS and HR-TEM images after operation demonstrate that the composition and nanoscale morphology are preserved during electrolysis (Figure S3-S12).

The stability and scalability of the electrode were finally investigated on a higher surface area support to target industrially relevant currents. Depositing ZnSO₄|CuSO₄ on commercially-available Zn foam (Mesh 4, 1 cm²) generated a highly-structured surface referred to as Zn foam | ZnSO₄|CuSO₄, similar to those deposited on flat Zn (Figure S19), and afforded stable currents in the range of ~50 to ~60 mA.

As proof of the electrode’s amenability to versatile syngas production in real-world conditions, we tested its response to the potential variations commonly observed while coupling the electrolyzer to an intermittent source of energy, such as solar panels. The applied potential was varied during electrolysis with the aforementioned foam over a 300 mV range, over which time the electrode maintained a stable H₂:CO ratio (Figure 6). In total, the system was operated for more than 9 h without any decline in activity or selectivity, nor evidence of structural degradation (Figure S19). It should be noted that, while independent with respect to the applied potential in both cases, the H₂:CO ratio observed using the Zn foam differs from that obtained using flat Zn as a support. This likely is a consequence of the very different diffusion and convection regime between these two supports, affecting the local concentration of the reactants at the electrode and the final H₂:CO ratio.

![Figure 6. Long-term electrolysis of a high surface area Zn foam | ZnSO₄|CuSO₄ in CO₂-saturated 0.1 M CsHCO₃. Varying the applied potential to mimic the foreseeable voltage fluctuations delivered by an intermittent renewable power source. Catalytic current is reported on the left axis (black) and corresponding H₂:CO ratio on the right axis (bulk symbols). Electrolyte was changed between each potential and roughly every hour beyond 260 minutes, causing slight signal variations.](image-url)

Conclusions

In this work, we have illustrated the use of a mixed-phased Zn-Cu alloying strategy for the one-step fabrication of a standalone material with tunable selectivity for the H₂:CO and CO₂ reduction reactions. The doping strategy not only provided fine control over catalytic selectivity, but the presence of easily-reduced Cu aided the growth of highly porous Zn-Cu alloys through seeding and hydrogen-evolution-assisted electrodeposition. H₂:CO ratios could be tuned from 0.2 to 3.65 and hence adjusted to desired ranges for syngas usage with industrially relevant currents. The catalyst activity could be related directly to the binding strength of the Cu and Zn atoms to key catalytic intermediates; a strategy that is applicable to future development of mixed-metal electrodes.

Finally, the unique ability of these materials to maintain a constant H₂:CO ratio over a broad range of applied potentials provides a new practical system to convert CO₂ to industrially relevant products using intermittent, renewable energy sources.

Experimental Section

Electrode preparation

Unless stated otherwise, electrodes were prepared on 1 cm² Zn foil (GoodFellow, 99.99%+, 1 mm) successively polished by P1200, P2400 emery paper and Al-powder followed by sonication in water before deposition. When the support employed was Cu, the same mechanical polishing procedure was applied on a 1 cm² Cu foil (GoodFellow, 99.999%, 1 mm). Each electrode was then immersed in a 0.5 M H₂SO₄ aqueous solution of 0.1 M metal salt apportioned between X% CuSO₄ and (100-X)% ZnSO₄ depending on the targeted Cu content and exposed to −0.5 A cm⁻² for 160 s using a three-electrode setup with an Ag/AgCl (KCl sat.) reference and Pt counter. In the case of the foam, 1 A was applied for 160 s. In each case the electrode was immediately rinsed with milliQ water and air-dried after deposition. CuSO₄|5H₂O (99.9%) and H₂SO₄ (99.8%), were purchased from Sigma-Aldrich and used without further purification. ZnSO₄|7H₂O (99.5%) was purchased from Roth chemicals.

Structure characterization

Imaging and EDX (Energy dispersive X-Ray spectrometry) were performed on a SU-70 Hitachi FEGSEM fitted with an X-Max 50 mm² Oxford EDX spectrometer. The imaging setup was 5 kV in order to observe surface features. Setup for quantitative analysis and mapping was 15 kV. Standards used as a reference for these voltages were purchased at Geller microanalytical laboratory (Boston, MA). Volume analyzed at this voltage is approximatively a sphere with diameter of ~700 nm. This value was calculated with Single Scattering Monte Carlo Simulation. Transmission electron microscopy images and chemical maps were acquired with a Jeol 2100F microscope operated at 200 kV. Chemical maps were acquired in STEM mode with the same microscope, equipped with Jeol system for X-ray detection and cartography. The elemental compositions of metallic electrodes
were verified with ICP-AES in a ThermoFisher iCAP 6000 device after dissolution of the metallic structures in 20% HNO₃ (Sigma-Aldrich, 65%).

Surface areas were obtained from the analysis of Kr sorption isotherms measured on a BelSorp Max set-up at 77 K. Prior to the measurement, samples were treated under vacuum at 130°C during at least 7 h. Surface areas were estimated using the BET model (Kr cross-sectional area 0.210 nm²). The value derived from BET measurement, reported in m².g⁻¹, was converted to cm².cm⁻²_geometric by multiplying it by the mass of deposited electrode onto the 1 cm² flat Zn support. This provided a roughness factor (RF), as defined by the IUPAC GoldBook.[28]

Powder X-ray diffraction measurements were performed in Bragg-Brentano geometry using a BRUKER D8 Advance diffractometer with Cu Kα radiation and a Lynxeye XE detector. XPS characterization was performed using a Thermo ESCALAB 250 X-ray photoelectron spectrometer with a monochromatic Al-Kα X-ray source (hv = 1486.6 eV) operating at a pressure around 2 × 10⁻⁶ mbar. The analyzer pass energy was 50 eV for survey spectra and 20 eV for high-resolution spectra. The spectrometer was calibrated using Au 4f7/2 at 84.1 eV. Charging effects were not compensated during analysis. Spectra were recorded and analyzed using Thermo Avantage software version 5.966.

### Electrochemical performance testing

Electroanalytic and constant potential electrolysis were carried out using a Bio-logic SP300 potentiostat. A H-type cell was used with the two compartments being separated by an anion exchange membrane (AMV Selemion™, ACG Engineering) with an inter-electrode distance of 6 cm between the working and Pt counter and an Ag/AgCl reference (saturated KCl) placed at 0.5 cm from the working. 0.1 M CsHCO₃ (Sigma-Aldrich, 99.9%) aqueous solution was used as both anolyte and catholyte, the latter being CO₂-saturated preceding the experiment (CO₂, Linde, HiQ 5.2) until the catholyte pH reached 6.8. During the electrolysis, CO₂ was constantly bubbled at 20 mL min⁻¹ through a frit at the bottom of the cathodic chamber and generated gaseous products and excess CO₂ were flowed to the gaseous inlet of a gas chromatograph for online measurement.

Potentials are reported against the Reversible Hydrogen Electrode (RHE) according to the relationship E vs. RHE = E vs. Ag/AgCl + 0.197+0.059*pH.

### Products characterization

H₂ and gaseous CO₂ reduction products were analyzed by a gas chromatography set-up (GC, Multi-Gas Analyzer #5 SRI Instruments) equipped with Haysep D and MoleSieve 5A columns, thermal conductivity detector (TCD) and flame ionization detector (FID) with methanizer using Argon as a carrier gas. GC was calibrated using a standard gas mixture containing 2500 ppm of H₂, CO, CO₂, C₂H₄, C₂H₆, C₃H₆, C₃H₈, C₄H₁₀ and C₆H₁₂ in CO₂ (Messer). The liquid-phase products were quantified using ionic exchange chromatography (for oxalate – 883 Basic IC, Metrohm) and NMR spectroscopy (Bruker AVANCE III 300 spectrometer).

Faradaic efficiencies (FE) were calculated according to the following formula:

\[ FE_{\text{product}} = \frac{n_e \cdot n_{\text{product}} \cdot F}{Q} \]

Where \( n_{\text{analyzed}} \) [mol] is the sample of analyzed product, \( n_e \) [no unit] is the number of electrons involved in the formation of this product, \( F \) is the Faraday’s constant equal to 96485 C.mol⁻¹ and \( Q \) is the corresponding passed charge. Partial current density for syngas production was calculated as follows:

\[ j_{\text{syngas}} = \left( FE_{\text{H₂}} + FE_{\text{CO₂}} \right) \cdot j \]

where \( j \) refers to the total current density [mA.cm⁻²].

### Acknowledgements

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### Keywords:

CO₂ reduction • Syngas • Catalysis • Zn-Cu alloys • Electrolysis

Two is better than one: a series of bimetallic Zn-Cu electrodes were prepared and used for syngas production by CO$_2$ electroreduction, allowing a broad range of H$_2$:CO ratios to be obtained independently of the potential applied.

Zn-Cu alloy nanofoams as efficient catalysts for CO$_2$ reduction to syngas mixtures with potential-independent H$_2$:CO ratio

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