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Cu and Zn promoted Al-fumarate metal organic frameworks for electrocatalytic CO₂ reduction†

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Metal organic frameworks (MOFs) are attractive materials to generate multifunctional catalysts for the electrocatalytic reduction of CO₂ to hydrocarbons. Here we report the synthesis of Cu and Zn modified Al-fumarate (Al-fum) MOFs, in which Zn promotes the selective reduction of CO₂ to CO and Cu promotes CO reduction to oxygenates and hydrocarbons in an electrocatalytic cascade. Cu and Zn nanoparticles (NPs) were introduced to the Al-fum MOF by a double solvent method to promote in-pore metal deposition, and the resulting reduced Cu–Zn@Al-fum drop-cast on a hydrophobic gas diffusion electrode for electrochemical study. Cu–Zn@Al-fum is active for CO₂ electroreduction, with the Cu and Zn loading influencing the product yields. The highest faradaic efficiency (FE) of 62% is achieved at –1.0 V vs. RHE for the conversion of CO₂ into CO, HCOOH, CH₄, C₂H₄ and C₂H₅OH, with a FE of 28% to CH₄, C₂H₄ and C₂H₅OH at pH 6.8. Al-fum MOF is a chemically robust matrix to disperse Cu and Zn NPs, improving electrocatalyst lifetime during CO₂ reduction by minimizing transition metal aggregation during electrode operation.

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1. Introduction

The electrochemical reduction of CO₂ into hydrocarbon fuels is a promising strategy to reduce societal reliance on fossil fuels and anthropogenic CO₂ emissions, while meeting global energy demands. To achieve this goal, the design and development of suitable functional materials which can effectively catalyse CO₂ reduction into fuels using cheap and renewable energy is required.^{1–3} Homogeneous and heterogeneous catalysts, including transition metal complexes of Pd, Re, Ru, Mn, Fe, Co, Ni and Cu, heterogenized molecular catalysts, nanostructured metals, metal chalcogenides and heteroatom-doped carbons have been explored.^{4–6} Multi-carbon products can form efficiently over Cu catalysts, being the preferred metal to promote C–C coupling reactions during CO₂ electroreduction.^{1,4} CO₂ reduction to multi-carbon products proceeds *via* a CO

intermediate, which undergoes additional multi-electron reduction, hence many studies target multifunctional catalysts in which a second active site selective for the reduction of CO₂ to CO, is incorporated alongside Cu. This enables a tandem process in which CO₂ reduction selectively produces CO for subsequent reduction and/or coupling over Cu sites to produce (oxygenated) hydrocarbons.^{5–7} Zn is selective for CO₂ electroreduction to CO,⁸ and in combination with Cu facilitates deeper reduction or coupling products (CH₃OH, CH₄, C₂H₄ or C₂H₅OH).^{9–11}

Metal-organic frameworks (MOFs) are attractive scaffolds that are readily functionalised with metal- or metal oxide-based nanoparticles for diverse applications including adsorption, membrane separation and catalysts.^{12–16} The high porosity, large surface area and chemical flexibility of MOFs renders them well-suited for fabricating multifunctional materials,^{13,14} with properties are tailored by changing the metal nodes or organic linkers, or introducing metal precursors within the pore network to create highly dispersed metal or metal oxide nanoparticles (NPs) with enhanced catalysis. Stabilization of such dispersed metal and metal oxide NPs may prevent their agglomeration and deactivation. MOFs have found application in electrocatalytic CO₂ reduction,^{13,17–20} with Jiang *et al.* reporting a Ag₂O/layered ZIF-7 catalyst, comprising Ag₂O NPs and a ZIF-7 MOF, that affords 81% faradaic efficiency (FE) for CO₂ electroreduction to CO to at –1.2 V vs. RHE in 0.25 M K₂SO₄. This value was much greater than that achieved with either ZIF-7 (25%) or Ag/C (36%) components.²¹ Hupp *et al.* embedded Cu

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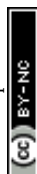
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NPs within Zr-MOF (NU-1000), obtaining a FE of 28% for CO₂ electroreduction reaction at -0.82 V vs. RHE, with formate (HCOO[−]) as the major product,²² while Beobide *et al.* reported a HKUST-1(Cu,Ru) heterometallic electrocatalyst formed by partially replacing Cu(II) nodes in the MOF with Ru(III) nodes, resulting in a combined FE of 47% for CO₂ conversion to methanol and ethanol. However, the latter electrocatalyst deactivated after 60 min of operation to a stable FE of only ~10%.²³

Metal-doped MOFs are hence promising electrocatalysts for CO₂ reduction, with multimetallic catalysts desirable to optimize product selectivity. Al-fumarate MOFs exhibit excellent thermal and chemical stability, alongside their high surface area and porosity,²⁴ but to our knowledge have not been investigated for CO₂ electroreduction. Here we explore the utility of Al-fumarate MOFs to: (i) synthesise Cu and Zn doped analogues for the cascade reduction of CO₂ to CO and subsequent multicarbon products; and (ii) improve active site dispersion and catalyst lifetime. Multimetallic Al-fumarate MOFs (Cu-Zn@Al-fum MOFs) with different metal loadings and Cu : Zn ratios were used to fabricate catalytic gas diffusion electrodes. Electroreduction of CO₂ was effective at neutral pH, with a FE of 27% to CO, 28% to hydrocarbons (CH₄, C₂H₄, C₂H₅OH) and 7% to HCOOH at -1.0 V vs. RHE.

2. Experimental

2.1 Preparation of Al-fum MOF

Al-fum MOF were prepared as follows.²⁴ In a 500 mL three-neck flask, 0.05 mol of Al₂(SO₄)₃·18H₂O was dissolved in 150 mL of DI water and heated to 65 °C. A 150 mL mixture containing an aqueous solution of 0.10 mol fumaric acid and 0.21 mol sodium hydroxide was injected into the reaction flask containing the aluminum precursor at 65 °C and stirred vigorously for 1 h. The obtained white suspension was filtered, then washed with DI water and warm ethanol. The washed Al-fum MOF product was dried overnight at 100 °C in air and subsequently at 130 °C in a vacuum oven.

2.2 Fabrication of Cu@Al-fum MOF

1 g of Al-fum MOF was dispersed in 50 mL of *n*-hexane and sonicated for 20 min to obtain a white suspension. Next, a certain volume of aqueous 1.45 mM Cu(NO₃)₂·3H₂O was added gradually under vigorous stirring. After stirring for 8 h, the blue solid was decanted and washed with *n*-hexane until the blue color of Cu²⁺ from the washing solution was clear. This precipitated solid was dried under vacuum at 80 °C overnight to obtain Cu²⁺@Al-fum MOF. Cu²⁺@Al-fum MOF which was then reduced by dispersing in 50 mL of CH₂Cl₂ and stirred vigorously for 30 min under N₂, prior to dropwise addition of a 25 mL NaBH₄ solution (prepared by dissolving 0.47 g of NaBH₄ in 25 mL of ethanol). The solution changed from light blue to yellow brown and then black as reduction proceeded. The product was collected by centrifugation and purified by dispersing in ethanol five times. The Cu@Al-fum MOF solid

precipitate was dried under vacuum for overnight, and hereafter is termed Cu@Al-fum.

2.3 Fabrication of Cu-Zn@Al-fum MOF

A Cu-Zn@Al-fum MOF was prepared similarly to Cu@Al-fum, except that Cu and Zn precursors were simultaneously introduced into the Al-fum MOF suspension. Briefly, a certain volume of aqueous Cu(NO₃)₂ (1.45 mM) and Zn(NO₃)₂ (3.5 mM) was mixed and added dropwise to a suspension containing 1 g of Al-fum MOF in 50 mL of anhydrous *n*-hexane under vigorous stirring. The prepared with 0.6 mL Cu²⁺ and 1.0 mL Zn²⁺ sample is termed Cu-Zn@Al-fum. After stirring overnight, the solids were collected, washed by *n*-hexane and dried under vacuum at 80 °C overnight to obtain Cu²⁺-Zn²⁺@Al-fum. The Cu²⁺-Zn²⁺@Al-fum was then dispersed in 50 mL of CH₂Cl₂ and stirred vigorously for 30 min under N₂ prior to reduction with fresh NaBH₄ solution which was added gradually into the reaction flask. The resulting Cu-Zn@Al-fum was collected, purified and dried akin to Cu@Al-fum MOF.

2.4 Fabrication of the electrode for CO₂ electroreduction

The working electrode was prepared by drop-casting aqueous suspensions of Cu@Al-fum MOFs or Cu-Zn@Al-fum MOFs onto a hydrophobic gas diffusion electrode (GDE) (dioxide, AvCarb GDS5130), on a hot plate at 80 °C. The solution for deposition was prepared with 1 mg catalyst (metal-impregnated MOFs) in 150 µL ethanol with 5 µL Nafion 5% (Sigma-Aldrich). The final 1 cm² electrode was dried in air before use.

2.5 Characterization

Morphology, structure and chemical composition of Al-fum MOFs were characterized by a transmission electron microscope (TEM, Hitachi H-7100, Japan), field emission scanning electron microscope (SEM, JEOL JSM-7600F, Japan), and powder X-ray diffraction (XRD, Shimadzu XRD-6000, Japan). Thermal stability was determined by thermogravimetric analysis (TGA, Mettler Toledo TGA/SDTA 851, USA) under a dry nitrogen flow of 30 mL min^{−1}, under heating at 5 °C min^{−1} from 32 to 700 °C. Brunauer–Emmett–Teller (BET) and Barrett–Joyner–Halenda (BJH) methods were applied to determine the specific surface area and pore size, respectively from N₂ physisorption isotherms (Quantachrome Nova 4200e porosimeter at 77 K). X-ray photoelectron spectra were acquired (Thermo Scientific K-alpha instrument) using a monochromatic Al K_α (1486.7 eV) source and charge neutralizer to investigate surface properties. Bulk Cu and Zn loadings were determined by inductively coupled plasma-optical emission spectrometry (ICP-OES, PerkinElmer, USA). Electrochemical impedance spectroscopy (EIS, Biologic SP-300) were measured with frequencies ranging from 100 kHz to 0.1 Hz at a potential of -1.0 V vs. RHE in 0.1 M KHCO₃, and the amplitude of the applied voltage was 10 mV. Gas chromatography (GC, SRI instruments, MG#5 GC) was used to analyze the gaseous products from CO₂ electroreduction and ¹H nuclear magnetic resonance (NMR, Bruker Avance III 300 MHz) was used to analyze liquid products; further details are provided in the ESI.†



2.6 Evaluation of electrocatalytic CO₂ reduction

CO₂ electroreduction was performed by a chronopotentiostatic method at different potentials, using an H-type electrochemical cell, which consists of two compartments separated by an anion exchange membrane (Dioxide, X37-50 grade T) containing the working, counter and reference electrodes. In the anodic compartment, Pt wire was the anode for the water oxidation reaction. In the cathodic compartment, CO₂ reduction was performed using catalysts deposited on the GDE in 0.1 M KHCO₃ solution-saturated CO₂ (pH 6.8) with a Ag/AgCl reference electrode. During electrolysis, CO₂ was continuously bubbled at a flow rate of 7.5 mL min⁻¹; the headspace was sampled by online GC for quantification of H₂, CO, CH₄, C₂H₄, with formate and ethanol quantified by ion chromatography and NMR respectively. Details of FE calculations are presented in the ESI.†

3. Results and discussion

3.1 Morphological and structural characterization

Al-fum with a high specific surface area, porosity and excellent water stability was successfully synthesized by precipitating aluminum sulphate with fumaric acid at 65 °C.²⁴ Cu and Cu-Zn NPs analogues were prepared by impregnation of metal salts into the Al-fum MOFs using the double solvent method (Fig. 1).

The composition, texture and structure properties of Al-fum, Cu@Al-fum and Cu-Zn@Al-fum were first determined by ICP and N₂ porosimetry and XRD; the Cu mass and molar loading in Cu@Al-fum was similar to the combined [Cu + Zn] loading in Cu-Zn@Al-fum of ~3 wt% (Table 1). The BET surface area and

average pore diameter of Al-fum were ~1073 m² g⁻¹ and 1.7 nm respectively, in good agreement with the literature.^{24–26} Note the average pore diameter is larger than the 0.6 nm of the rhombohedral channels of the MOF framework due to mesoporous intercrystallite voids²⁷ (with a mode pore diameter of ~8 nm). The pore diameter increases after Cu and Cu/Zn doping, likely due to enlargement of intercrystallite voids. The decrease in surface areas of Cu@Al-fum and Cu-Zn@Al-fum compared to Al-fum is attributed to incorporation of dense Cu and Zn NPs within the low density MOF, and resultant partial micropore blockage. This observation concurs with a previous report of Al-fum and CuO/ZnO/AlFum MOFs wherein surface areas decreased from 910 to 416 m² g⁻¹.²⁸

The hydrophilicity of Al-fum MOF after Cu and Zn incorporation was evaluated from water adsorption isotherms (Fig. S1a†). Water adsorption decreased 3.5 times after introducing Cu and Zn into the Al-fum MOF, partly reflecting the lower surface area and pore volume, but also indicative of decreased hydrophilicity.

Fig. 2 shows the XRD patterns of Al-fum, Cu@Al-fum and Cu-Zn@Al-fum, with the former (Fig. 2a) exhibiting diffraction peaks at 2θ of ~10.6, 15.2, 21.2, 31.8 and 42.8° corresponding to the (011), (020), (022), (033) and (044) planes of monoclinic Al-fum MOF crystals.

Cu@Al-fum and Cu-Zn@Al-fum exhibit similar reflections to Al-fum, suggesting that the former retain the crystalline structure of the parent MOF. No reflections associated with copper oxides were observed (Fig. 2b), however a very weak peak at 42° in Cu@Al-fum is characteristic of copper metal. For Cu-Zn@Al-fum, small reflections at 36.4° and 43.7° are indicative of

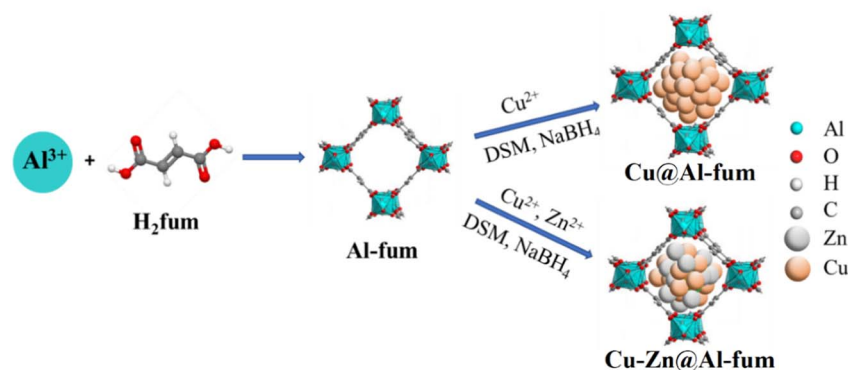


Fig. 1 Schematic of Cu@Al-fum and Cu-Zn@Al-fum MOF synthesis.

Table 1 Elemental analysis and textural properties of Al-fum MOFs

Samples	Elemental loading ^a (wt%)		S_{BET}^b (m ² g ⁻¹)	Pore volume ^b (cm ³ g ⁻¹)	Pore diameter ^c (nm)
	Cu	Zn			
Al-fum	—	—	1073	1.4	1.7 (8.3)
Cu@Al-fum	2.92	—	335	0.8	1.9 (11)
Cu-Zn@Al-fum	0.86	2.18	560	1.2	1.9 (19)

^a ICP analysis. ^b Error in S_{BET} and pore volume $\pm 10\%$. ^c Modal value of mesopore diameter in brackets.

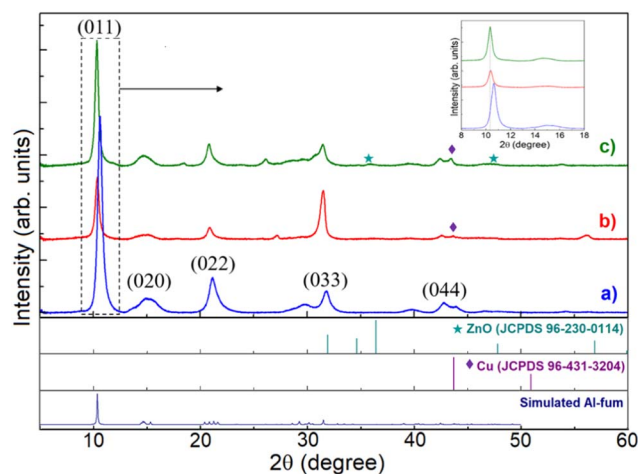


Fig. 2 XRD patterns of (a) Al-fum, (b) Cu@Al-fum and (c) Cu-Zn@Al-fum. Inset shows the magnification of the (011) peak.

ZnO (Fig. 2c), although no reflections associated with Zn metal were observed (Zn being more easily oxidized than Cu). The low intensities of Cu and ZnO reflections is consistent with their low loading in the doped MOF.²⁸ The (011) reflection of the parent Al-fum MOF slightly shifts to lower angle (by $\sim 0.3^\circ$) following Cu and Cu-Zn modification (Fig. 2 inset), indicating lattice expansion which we ascribe to lattice stress/strain or defects (dislocations or stacking faults) arising from the incorporation of Cu and Zn NPs into the MOF framework.²⁹

SEM and TEM images of Cu-Zn@Al-fum (Fig. 3) and of Cu@Al-fum (Fig. S1b†) reveal aggregates of relatively uniform sheets (average width ~ 50 – 100 nm), similar to the parent Al-fum MOF and consistent with previous reports.^{24,27} TEM images and EDX elemental maps (Fig. 3, S1b and S2†) evidence <6 nm Cu and Zn NPs uniformly distributed throughout the Al-fum MOF, akin to reports of Cu and Zn NPs dispersed in UiO-66 or Ni NPs in Zr MOFs.^{30–32}

A survey scan by XPS identified the presence of C, O, Al, Cu and Zn in Cu@Al-fum and Cu-Zn@Al-fum (Fig. S3a†).

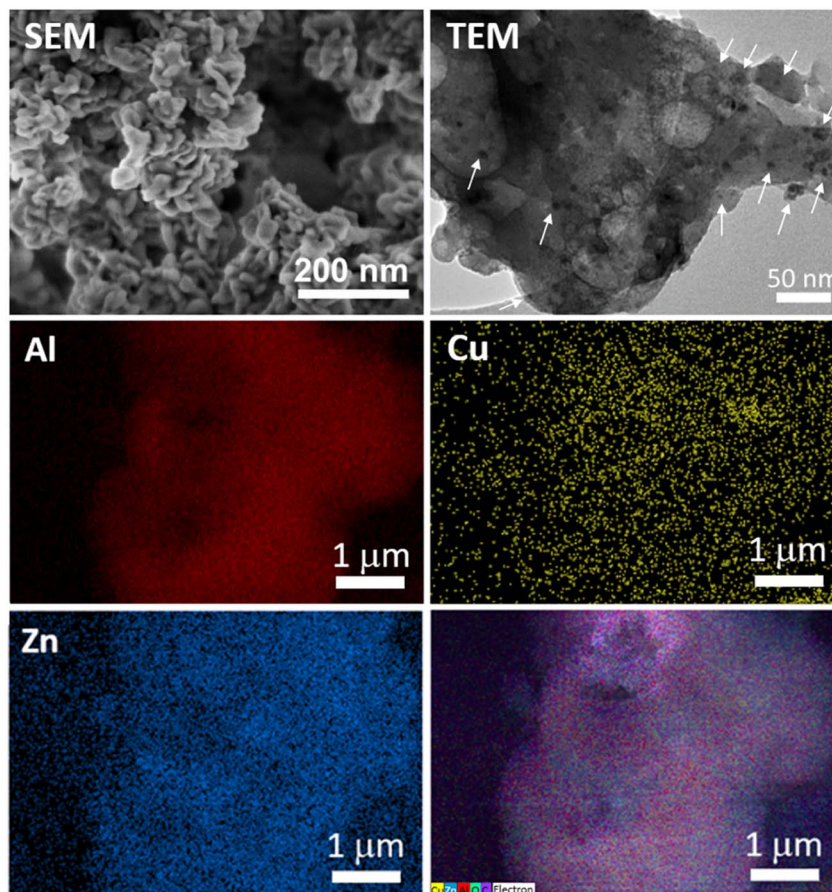


Fig. 3 (Top) SEM and TEM images, and (middle, bottom) EDX elemental maps of Cu-Zn@Al-fum. Arrows in the TEM image indicate <6 nm Cu and/or Zn NPs. FTIR spectra of Al-fum, Cu@Al-fum and Cu-Zn@Al-fum (Fig. 4a) confirmed the molecular components of the MOF framework. Bands at ~ 1610 cm^{-1} , 1430 cm^{-1} , 1158 cm^{-1} and 805 cm^{-1} are attributed to asymmetric and symmetric stretches of the carboxylate group in fumarate (which coordinates to Al^{3+} nodes).^{33,34} Bands spanning 720 – 650 cm^{-1} arise from $\text{C}=\text{C}$ and $\text{C}-\text{H}$ bending modes of the fumarate framework, while the broad band from 3400 – 3600 cm^{-1} is due to the $\text{O}-\text{H}$ stretch of in-pore or intercrystallite water. New bands are visible <650 cm^{-1} following Cu and Zn doping, attributed to $\text{Cu}-\text{O}$ stretches in Cu_2O (expected ~ 615 – 630 cm^{-1}) and CuO (expected ~ 609 – 590 cm^{-1} and ~ 530 – 508 cm^{-1}),³⁵ and copper coordinated to carboxylate groups (<450 cm^{-1})³⁶ in addition to $\text{Zn}-\text{O}$ stretches (expected <555 cm^{-1}).³⁷



Corresponding high-resolution C 1s XPS spectra (not shown) reveal three peaks with binding energies of 284.6, 286.2 and 288.4–288.8 eV, attributed to C–C/C–H/C=C, C–O and C=O groups, respectively of the fumarate framework.^{38–40} The Al-fum exhibits Al 2p spin-orbit split peaks at ~ 74.4 and 75.5 eV (Fig. S3b[†]), assigned to AlO(OH) species within the MOF framework,⁴¹ and two O 1s peaks at ~ 531.9 eV and 533.2 eV (Fig. S3c[†]) characteristic of metal and H-bonded oxygen respectively.⁴² Small shifts in the principal Al 2p and O 1s peaks to lower binding energy following metal doping (Fig. S3b and c[†]) may reflect tensile or compressive strains in the Al-fum MOF framework and a concomitant change in the Al³⁺ charge density.

Unreduced Cu@Al-fum and CuZn@Al-fum samples exhibit Cu 2p_{3/2} and 2p_{1/2} spin-orbit split peaks at 933.2 and 953.0 eV respectively, which (in conjunction with the absence of a copper satellite) indicates the presence of Cu⁺ species (Fig. S4[†]). Following NaBH₄ reduction, the Cu 2p binding energies decreased slightly to 932.6 and 952.4 eV for Cu@Al-fum and 932.8 and 952.7 eV for Cu–Zn@Al-fum (Fig. 4b and S4[†]) consistent with the formation of some metallic Cu.⁴³ In both cases, the Auger parameter calculated using the Cu LMM peak ~ 918.2 eV kinetic energy (Fig. S5[†]) was 1851.4 eV, consistent with Cu metal.⁴⁴ A similar binding energy red-shift was observed for the Zn 2p spectra following NaBH₄ reduction, from ~ 1022.3 for Zn²⁺ species in Cu²⁺Zn²⁺@Al-fum to 1021.9 eV for Cu–Zn@Al-fum (Fig. 4b and S4[†]). This shift may reflect chemical reduction of Zn²⁺ to Zn or electronic perturbation due to alloying with copper.³² It is challenging to distinguish Zn and ZnO from XPS

due to their similar binding energy values,⁴⁵ however the formation of Zn NPs in MOF pore networks is known.^{46–48}

The porosity of the Al-fum, Cu@Al-fum and Cu–Zn@Al-fum MOFs was analyzed using nitrogen porosimetry (Fig. S6a[†]), which reveals the parent and metal-impregnated MOF samples exhibit typical I isotherm behavior, indicating the existence of both microporous and mesoporous structures. The BJH pore size distribution (Fig. S6b[†]) reveals all samples show a broad range of mesopores with a mode of ~ 8 nm for Al-fum, increasing to 11 and 19 nm for Cu@Al-fum MOF and Cu–Zn@Al-fum respectively, which are attributed to intercrystallite voids, and is in good agreement with observations from SEM in Fig. 3 and S1b[†].

Thermal analysis of Al-fum reveals mass losses at ~ 50 °C and 475 °C (Fig. S7[†]) corresponding to the removal of physisorbed water and adsorbed/coordinated solvent (~ 30 wt%) and subsequent decomposition of the fumarate organic linker (~ 35 wt%). The residual ~ 35 wt% is associated with reactively-formed alumina. Al-fum is thus thermally stable to 475 °C, in good agreement with previous reports.^{24,26,28} Impregnation with Cu and Zn NPs lowers the thermal stability, with the frameworks of Cu@Al-fum and Cu–Zn@Al-fum decomposing at 435 °C and 447 °C respectively, and new lower temperature mass losses emerging at 320 °C and 340 °C, respectively. We speculate that the presence of metal NPs promotes defect formation (missing Al³⁺ nodes or fumarate linkers) and/or lattice strain, destabilizing the parent framework, consistent with literature reports.²⁸ Nevertheless, Cu@Al-fum and Cu–Zn@Al-fum are stable to 320 °C, higher than many MOFs.^{49,50}

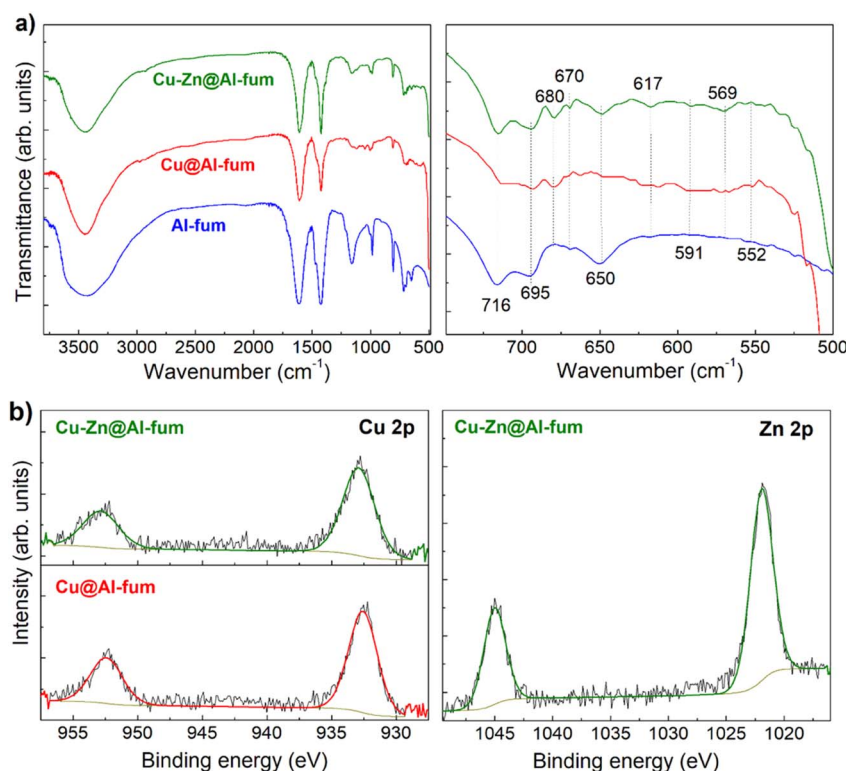


Fig. 4 (a) FTIR spectra of Al-fum MOFs, and (b) Cu and Zn 2p XP spectra of Cu@Al-fum and Cu–Zn@Al-fum.

3.2 Electrocatalytic reduction of CO₂

The electrocatalytic reduction of CO₂ over Al-fum derived catalysts was evaluated using a bespoke H-cell (Fig. S8a†), with two compartments separated by an anion exchange membrane (AEM). To maximize catalytic performance, CO₂ was injected through a glass frit at the base of cathodic compartment to produce a stream of small bubbles transported to the catalyst deposited on the hydrophobic GDE. All electrocatalysts achieved steady-state operation after 30 min time-on-stream with an example shown for Cu-Zn@Al-fum under the operating potentials studied (Fig. S8b†).

The catalytic performance of Cu-Zn@Al-fum was evaluated by linear sweep voltammetry (LSV) in the presence and absence of CO₂ (Fig. 5). Under an Ar atmosphere, liquid and gas analysis confirmed that the faradaic current density (j) was entirely due to the hydrogen evolution reaction (HER), whereas in the

presence of a CO₂-saturated solution j reflects competition between proton and CO₂ reduction. Hydrogen production was greatly suppressed under a CO₂ atmosphere, with the chemical selectivity (on a molar basis) to reduced carbon products reaching 50% for Cu-Zn@Al-fum at −1.2 V vs. RHE (Table S1†). A similar switchover from proton reduction under Ar to CO₂ reduction in a CO₂ saturated aqueous solution is reported in the literature.^{51–53} Note that the molar selectivity to reduced carbon products is always lower than the FE, as the latter accounts for the greater number of electrons required to form *e.g.* C₂H₄ (12 e[−]) than H₂ (2 e[−]). Although the electrolyte pH decreased from 8.3 under Ar to 6.8 for a CO₂ saturated solution (which could result in a positive shift in potential for H₂ generation),⁵⁴ there was no evidence for a systematic increase in H₂ production at more positive potentials (Fig. 5 and S9†). Literature reports suggest that CO₂ reduction is more pronounced at lower pH,⁵¹

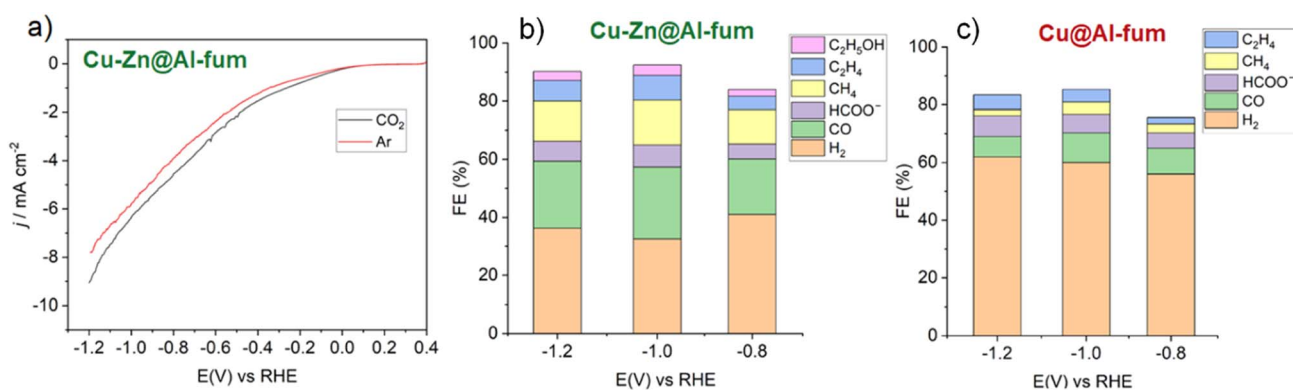


Fig. 5 (a) LSV of Cu-Zn@Al-fum/GDE in 0.1 M KHCO₃ aqueous solution saturated with CO₂ or Ar, and FE for reduction products at different cathodic potentials for (b) Cu@Al-fum, and (c) Cu-Zn@Al-fum. CO₂ was continuously bubbled at 7.5 mL min^{−1} during electrolysis.

Table 2 Comparison of electrochemical CO₂ reduction over Cu@Al-fum and Cu-Zn@Al-fum catalysts

Type of MOF	j (mA cm ^{−2})	Synthesis	Electrolyte	Maximum FE	Ref.
Ag@Al-PMOF	~6	ALD, solvothermal	0.1 M KHCO ₃	CO 56% at −1.1 V vs. RHE	55
Cu NPs embedded NU-1000 (ZIF MOF)	1.8	Solvothermal	0.1 M NaClO ₄	Formate 30% and CO 5% at −0.82 V vs. RHE	21
HKUST-1 + Cu NPs	20	Solvothermal	0.5 M NaHCO ₃	C ₂ H ₄ 12% and CH ₄ 19% at −2 V vs. SCE	57
HKUST-1 + CNT	8	Precipitation	0.5 M KHCO ₃	CH ₄ 25%, CO 5% and C ₂ H ₄ ~ 1% at −1.06 V vs. RHE	58
HKUST-1(Cu,Ru)	20	Solvent-free synthesis	0.5 M KHCO ₃	CH ₃ OH 1% and C ₂ H ₅ OH 10% at −1.84 V vs. Ag/Ag ⁺	23
Cu ₂ O-QDs@CuHHTP MOF	10.8	Solvothermal	0.1 M KCl/0.1 M KHCO ₃	CH ₄ 73% at −1.4 V vs. RHE	19
GO@Cu(BTC)	—	Hydrothermal	0.1 M KHCO ₃	HCOOH 21% at −0.1 V vs. SCE	50
MIL-53(Al)	1.2	Hydrothermal	0.5 M K ₂ CO ₃	HCOOH 19%, CO 21% at −1.1 V vs. RHE	59
CuZnDTA MOA	10	Ultrasonic	0.5 M KHCO ₃	C ₂ H ₅ OH 7%, CH ₃ OH 3% at −1.25 V vs. Ag/Ag ⁺	60
Cu@Al-fum MOF	~7	Solvothermal, double solvent	0.1 M KHCO ₃	CO 11%, CH ₄ 6%, C ₂ H ₄ 6% and formate 8% at −1.0 V vs. RHE	This work
Cu-Zn@Al-fum MOF	~5	Solvothermal, double solvent	0.1 M KHCO ₃	CO 27%, CH ₄ 16%, C ₂ H ₄ 9%, C ₂ H ₅ OH 3% and formate 7% at −1.0 V vs. RHE	This work



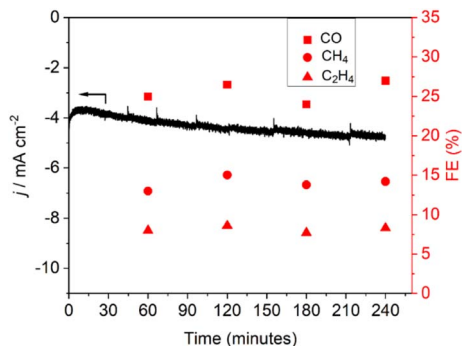


Fig. 6 Time-dependent CO₂ electroreduction over Cu-Zn@Al-fum.

which could result in the higher current density of Cu-Zn@Al-fum/GDE under CO₂ saturation (Fig. 5a). Considering that LSV curves are a convolution of catalyst activity and selectivity, differences between them can only be interpreted following analysis of evolved CO₂ reduction products, with formation of CO, HCOOH, CH₄, C₂H₄ and C₂H₅OH confirmed by GC and NMR (Table S1†). Analogous studies for Cu@Al-fum (Fig. 5c) confirmed the production of gaseous CO, CH₄, C₂H₄ and H₂ and liquid formate and ethanol (Table S1†). In contrast, Al-fum only produced H₂, CO and formate (Fig. S9†).

Cu-Zn@Al-fum achieved a higher yield of CO₂ reduction products than Cu@Al-fum at all applied potentials, reaching an overall FE of 62% for CO₂ reduction products (and 32% FE for H₂) at -1.0 V vs. RHE. The total FE of Cu-Zn@Al-fum to CH₄, C₂H₄ and C₂H₅OH (8, 12 and 12 electron reductions, respectively) products alone is $\sim 34\%$ (Fig. 5b). These efficiencies (selectivities) for CO₂ reduction over Cu-Zn@Al-fum at neutral pH compare favorably to literature MOF electrocatalysts (Table 2) prepared by more complex colloidal, atomic layer deposition (ALD) or solvothermal syntheses, which predominantly yield CO.⁵⁵ Competition between CO₂ reduction and H₂ evolution will always be challenging, but tuning the solution pH could afford higher yields of multicarbon products. For Cu@Al-fum, H₂ production dominated, with a FE $> 50\%$ at cathodic potentials (Fig. 5c and Table S1†), indicating Cu was relatively poor at activating CO₂ under neutral conditions, whereas Cu-Zn@Al-fum exhibited the highest CO yield (Table S1†) consistent with the reported selectivity to this product over Zn electrocatalysts.¹² High rates of CO production over Zn are expected to promote deeper reduction and C-C coupling reactions over proximate Cu sites.^{11–13}

Further insight into the conductivity of the Al-fum MOF derived catalysts was obtained from electrochemical impedance spectroscopy (EIS) over the frequency range 100 kHz to 0.1 Hz, at a potential of -1.0 V vs. RHE in 0.1 M KHCO₃ with 10 mV amplitude for the applied voltage. Resulting Nyquist plots for Al-fum, Cu@Al-fum, and Cu-Zn@Al-fum (Fig. S10†) were fitted to a simplified Randles circuit to extract the charge transfer resistance (R_{ct} , Table S2†).⁵⁶ The first intercept on the x -axis relates to contact resistance (R_s), and includes the electrode, interfacial contact resistance between the current collector and the electroactive material, and the electrolytic solution resistance. Although R_s values were similar for all catalysts, R_{ct}

decreased significantly after Cu and Zn doping, indicating an increased electrical conductivity.

Durability of the Cu-Zn@Al-fum electrocatalyst was also assessed for CO₂ electroreduction (Fig. 6): the time-dependent current density increased by ~ 10 – 15% after 4 h time-on-stream, while FE for gaseous CO₂ reduction products remained stable.

4. Conclusions

Al-fum, Cu@Al-fum and Cu-Zn@Al-fum MOFs were synthesized and deposited on a hydrophobic gas diffusion electrode as electrocatalysts for CO₂ reduction in neutral aqueous solutions. Cu and Zn were incorporated into the parent Al-fum MOF by facile co-impregnation of Cu²⁺ and Zn²⁺ salts which were subsequently reduced to corresponding metal nanoparticles by NaBH₄. Co-doping (Cu-Zn@Al-fum) significantly improved CO₂ electroreduction compared to a singly doped Cu catalyst (Cu@Al-fum) and the Al-fum (which only catalysed proton reduction). Cu-Zn@Al-fum achieved a FE of 62% for CO₂ reduction to CO (27%), and desirable CH₄, C₂H₄ and C₂H₅OH (28%), and HCOO[−] (7%) products. This excellent selectivity under neutral pH is attributed to its lower hydrophilicity (suppressing proton reduction) and the proximity of Zn and Cu electrocatalyst sites which promote the cascade reduction of CO₂ to CO and formic acid (over Zn) and subsequent reduction of CO to CH₄ and multicarbon products (over Cu). The parent Al-fum MOF offers high thermal and chemical stability, and appears an excellent matrix to disperse and stabilize metal NPs during electrochemical operation. This approach should be amenable to diverse Earth abundant metal dopants for CO₂ electroreduction to valuable fuels and chemicals.

Author contributions

Ung Thi Dieu Thuy: conceptualization, synthesis, analysis, investigation, project administration, writing – review & editing; Tran Ngoc Huan: measure, analysis, writing – review & editing; Sandrine Zanna and Ngoc-Diep Le: measure, analysis; Jim Mensah: characterization and analysis; Venkata D. B. C. Dasireddy: characterization and analysis; Karen Wilson: writing – review & editing; Adam F. Lee: writing – review & editing; Nguyen Quang Liem: conceptualization, writing – review & editing.

Conflicts of interest

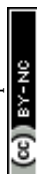
There are no conflicts of interest to declare.

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