Journal of Materials Chemistry A



PAPER

View Article Online
View Journal | View Issue



Cite this: J. Mater. Chem. A, 2024, 12, 32821

Temperature promotes selectivity during electrochemical CO₂ reduction on NiO:SnO₂ nanofibers†

M. A. Rodriguez-Olguin, ‡^{ab} R. Lipin, ‡^c M. Suominen, d F. Ruiz-Zepeda, logo ef E. Castañeda-Morales, d A. Manzo-Robledo, d J. G. E. Gardeniers, logo d C. Flox, *dh T. Kallio, d ** M. Vandichel ** and A. Susarrey-Arce ** and ** susarrey-Arce ** and **.

Electrolyzers operate over a range of temperatures; hence, it is crucial to design electrocatalysts that do not compromise the product distribution unless temperature can promote selectivity. This work reports a synthetic approach based on electrospinning to produce NiO:SnO2 nanofibers (NFs) for selectively reducing CO₂ to formate above room temperature. The NFs comprise compact but disjoined NiO and SnO₂ nanocrystals identified with STEM. The results are attributed to the segregation of NiO and SnO₂ confirmed with XRD. The NFs are evaluated for the CO2 reduction reaction (CO2RR) over various temperatures (25, 30, 35, and 40 °C). The highest faradaic efficiencies to formate (FE_{HCOO}-) are reached by NiO:SnO₂ NFs containing 50% of NiO and 50% SnO₂ (NiOSnO50NF), and 25% of NiO and 75% SnO₂ (NiOSnO75NF), at an electroreduction temperature of 40 °C. At 40 °C, product distribution is assessed with in situ differential electrochemical mass spectrometry (DEMS), recognizing methane and other species, like formate, hydrogen, and carbon monoxide, identified in an electrochemical flow cell. XPS and EELS unveiled the FE_{HCOO}- variations due to a synergistic effect between Ni and Sn. DFT-based calculations reveal the superior thermodynamic stability of Ni-containing SnO₂ systems towards CO₂RR over the pure oxide systems. Furthermore, computational surface Pourbaix diagrams showed that the presence of Ni as a surface dopant increases the reduction of the SnO₂ surface and enables the production of formate. Our results highlight the synergy between NiO and SnO₂, which can promote the electroreduction of CO₂ at temperatures above room temperature.

Received 13th June 2024 Accepted 8th August 2024

DOI: 10.1039/d4ta04116j

rsc.li/materials-a

^eMesoscale Chemical Systems, MESA+ Institute, University of Twente, P. O. Box 217, Enschede 7500AE, The Netherlands

Department of Physics and Chemistry of Materials, Institute of Metals and Technology, Lepi pot 11, Ljubljana, Slovenia

*Instituto Politécnico Nacional, Laboratorio de Electroquímica y Corrosión, Escuela Superior de Ingeniería Química e Industrias Extractivas, Av. Instituto Politécnico Nacional S/N, Unidad Profesional Adolfo López Mateos, CP 07708 CDMX, Mexico *Department of Electrical Energy Storage, Iberian Centre for Research in Energy Storage, Campus University of Extremadura, Avda. de las Letras, s/n, 10004 Cáceres, Spain. E-mail: cristina.flox@ciiae.org

 \ddagger These authors contributed equally to this work.

1. Introduction

Anthropogenic CO2 emissions are a problem that must be addressed to decelerate greenhouse effects like global warming. According to the Intergovernmental Panel on Climate Change (IPCC) published in 2021, the planet's temperature may increase by 1.5 °C in the following decades due to the accelerated release of CO2 and other greenhouse gases, posing a threat to the global ecosystems and humankind. A call-to-action part of the energy transition goals is CO₂ recycling and utilization. One way to recycle and utilize CO2 is by producing CO2-based value-added products electrochemically. For example, electrochemical CO₂RR can enable the production of C1 (e.g., CO, $HCOO^-$, CH_4) and C2 (e.g., C_2H_4 , alcohol) products. The selectivity of C1 or C2 products is often attributed to the catalyst type or the operation conditions during CO2 electrolysis. As for the catalysts, three different groups of metals have been recognized based on the product generated: formate (e.g., Sn, 2 Pb, 3 Bi, 4,5 In, 6 Hg⁷), CO (Au,⁸ Ag,⁹ Pd,¹⁰ Zn,¹¹ Ni¹²), hydrocarbons and alcohols (Cu^{13,14}) and other multi-carbon products like amino acids, ¹⁵ which in combination with CO2 and nitrogenated sources (NH₄HCO₃) has been demonstrated to lead to the formation of

^bDepartment of Chemical Engineering, MESA+ Institute, University of Twente, P. O. Box 217, Enschede 7500AE, The Netherlands. E-mail: a.susarreyarce@utwente.nl
^cSchool of Chemical Sciences and Chemical Engineering, Bernal Institute, University of Limerick, Limerick V94 T9PX, Republic of Ireland. E-mail: matthias.vandichel@ul.ie
^cDepartment of Chemistry and Materials Science, Aalto University School of Chemical Engineering, Kemistintie 1, 02015 Espoo, Finland. E-mail: tanja.kallio@aalto.fi
^cDepartment of Materials Chemistry, National Institute of Chemistry, Hajdrihova 19, 1000, Ljubljana, Slovenia

serine.^{16,17} Among the mentioned metals, Cu is a catalyst capable of producing C1, C2, and higher multi-carbon products.¹⁸ However, copper exhibits low selectivity, resulting in a mixture of gaseous and liquid reaction compounds. Therefore, the selective production of CO₂-reduction products is a key challenge that needs attention. One way to overcome this challenge is by synthetically designing a catalyst with earthabundant elements that can cope with CO₂RR on an industrial level aimed at 90% conversion to a single product and elevated electrolyzer temperatures for practical reasons (*e.g.*, increased reaction rates or reduced overpotentials).

The reaction temperature is a crucial yet sometimes overlooked parameter in electrochemistry that can compensate for the thermal losses during CO₂ electrolysis. Mizuno et al. have delved into the electrochemical reduction of CO2 with various electrode compositions.19 In a different study, Kim et al. analyzed the effect of operating conditions, including temperature, on the electrochemical conversion of CO2 to formic acid.20 Like Mizuno et al.,19 the results underscore the intricate relationship between temperature and selectivity for CO₂RR. Recently, Löwe et al. addressed in depth how temperature variations can influence transport and, thus, formate production.21 According to the authors, the temperature notably impacts the reaction selectivity at an industrially accepted current density of 200 mA cm⁻². Specifically, the FE_{HCOO} decreases from 89% at 20 °C to 85% at 70 °C, while the production of CO increases from around 7% to 11%. Interestingly, variations of about ±10% during FE_{HCOO}- have been observed at the assessed temperatures with current densities of up to 1000 mA cm⁻². The results from Löwe et al. indicate that with an increase in temperature, the selectivity towards formate decreases while it increases for other products like CO or H₂. Other studies reported similar trends, ascribing the decreased formate production and selectivity with increasing temperature to the kinetic effects and CO2 solubility.22 Under other conditions, e.g., by varying the electrolyte solution, a fundamental understanding of the relationships between the surface coverage, pH, and kinetics has been proposed for Cu.18,23 The previously cited seminal works are therefore used as a stepping stone to understand the temperature effects over NiO:SnO₂ catalyst, whose synergistic combinations can lead to tuning the selectivity of CO₂RR products. It should be noted that other studies for NiO, SnO2, and their combinations do not deal with temperature variation conditions during CO2 electrolysis (Tables S1-S3†). NiO typically leads to CO, CH₄, and, in some cases, to $HCOO^-$ (Table S1†), while SnO_2 leads to $HCOO^-$ as the main product (Table S2†). However, their synergistic effect has only been shown HCOO⁻ products near room temperature (Table S3†).

An additional expected challenge for a single oxide composition like SnO_2 is its instability under cathodic electrochemical potentials. The instability of SnO_2 during CO_2RR can be attributed to the thermodynamic formation of various oxidized tin species. Mu *et al.* highlighted the dominance of hydroxyl radicals in the reoxidation of oxide-derived metals like Cu. Despite being thermodynamically unstable under cathodic conditions, the authors propose that the presence of $Cu^{\delta+}$

species enables the CO2RR.26 The results from Mu et al. emphasize the importance of the stability of metal oxide catalysts.26 Lately, Jiang et al. have discussed the importance of stabilizing the oxidation state of SnO₂ to achieve highly selective CO₂ electroreduction to formate.²⁷ Their study emphasizes the necessity of maintaining the oxidation state of tin during the CO₂RR, achieved by incorporating Cu single atoms into the SnO₂ lattice. Like single-atom catalysis, other strategies such as engineering the defects in oxides, have been proposed to tune the selectivity of CO₂RR²⁸ and NO_x reduction reactions.^{28,29} From the later examples, catalyst discovery should favor thermodynamically stable catalytic species that appropriately control product selectivity over various reaction conditions. This could be the case for catalysts that incorporate suitable supports or modifiers optimized for various temperature conditions to minimize the degradation of the active sites.

Aside from single Cu atoms and others like Pt and Bi,²⁷ NiO is an exciting option to stabilize SnO₂ because it can be incorporated in much higher proportions using electrospinning.^{30,31} The incorporation of NiO and SnO₂ goes hand in hand with the possibility of achieving 1D nanofibrous structures to produce non-woven architectures ideal for transport control during electrochemical reactions.^{32,33} Herein, our working hypothesis is that NiO embedding will stabilize the SnO₂ and increase the hydrogenation activity of SnO₂, further enabling the formate (HCOO⁻) production. However, it is not 'a priori' clear how NiO and SnO₂ combined in an NF can synergistically favor formate production and how temperature affects selectivity. Thus, it is essential to explore.

In this work, we employ electrospinning to synthesize different NiOSnO2 NFs by varying Ni: Sn precursor molar ratios between 75:25, 50:50, and 25:75. Among the applied Ni:Sn molar ratios, 50:50 and 75:25 resulted in an NF-like morphology after annealing at 550 °C while other molar ratios or single compositions did not lead to NFs. Therefore, a detailed structural and morphological characterization (XRD, HR-SEM, and STEM-EDX) is carried out for the 50:50 (NiOSnO50NF) and 25:75 (NiOSnO75NF) NFs. Additionally, XPS and EELS are used to understand the chemical environment before and after the CO₂RR. The results show the presence of Ni³⁺ species and partially reduced SnO2 after CO2 electrolysis at 40 °C. The selectivity of the NFs is evaluated by using an electrochemical flow cell with a maximum heating capacity of 40 °C. The electrochemical experiments for NiOSnO50NF and NiOSnO75NF show HCOO $^-$ selectivities close to \sim 25% at 25 °C, while at temperatures of 30 °C and 35 °C the HCOO selectivity increased to ~30% and ~50%. An increased HCOO⁻ selectivity is observed at an electroreduction temperature of 40 °C, ca. \sim 85%, and \sim 70% for NiOSnO50NF and NiOSnO75NF, respectively. The product distribution assessed with in situ DEMS aligns with the products observed in the flow cell experiments carried out at 40 °C, where, besides methane (CH₄), HCOO⁻, hydrogen (H₂), and carbon monoxide (CO) products have also been found. DFT modeling provides insights into the reaction mechanism and the effect of temperature during the CO₂RR process. Furthermore, the computational surface Pourbaix diagram indicates that combining NiO and SnO2 increases the

hydrogenation level in the catalyst model, enabling HCOOproduction. Our results highlight the importance of catalyst discovery, as demonstrated by the synergistic effects between NiO and SnO₂, which can boost the electroreduction of CO₂ at temperatures higher than room temperature.

2. Methods

Synthesis of NiO:SnO2 nanofibers

The electrospinning technique was used to synthesize metal oxide NFs. For the NiO:SnO₂ NF, NiCl₂·6H₂O (ACS grade, Sigma Aldrich) and SnCl₂·XH₂O (ACS grade, Sigma Aldrich) were used as precursors. The stock solutions containing Ni, Sn, or their combinations were prepared by dissolving the metal salts in ethanol (100% Tech. grade, BOOM B.V., The Netherlands). Subsequently, polyvinylpyrrolidone (PVP, MW ~1 300 000 by LS, Sigma Aldrich) is added to the solution and stirred magnetically overnight. The precursor solutions were spun using a commercial electrospinning system from IME Technologies (The Netherlands) at 2.0 mL h⁻¹. NFs were obtained at 10.25 kV using a stainless-steel needle of 0.4 mm inner diameter. The collector was maintained at a separation distance of 12 cm from the needle to the aluminum collector plate. The NFs were collected at 22 °C and a relative humidity of 30%. After deposition, NFs were dried in a furnace at 80 °C for 12 h to remove the excess solvent. In a subsequent step, the NFs were annealed in two steps in air. First, the NFs were annealed at 350 °C (ramp-up rate of 1 °C min⁻¹) for 3 h to remove the organic components. Second, the NFs were annealed at 550 °C (1 °C min⁻¹) for 1 h to produce the NiO:SnO2 mixed oxides. Controls were produced using the electrospinning precursor mentioned above. The controls lead to nanoparticles (NP), hereafter labeled as NiOSnONP. In short, the control samples were prepared by directly pouring the electrospinning precursor into porcelain crucibles and calcined at 550 °C (1 °C min⁻¹) for 1 h in air.

2.2 Morphological, structural and chemical characterization

- 2.2.1 HR-SEM. High-resolution (HR)-SEM images were taken using a Zeiss MERLIN SEM microscope operated at 1.4 kV coupled with a High-Efficiency Secondary Electron Detector (HE-SE2). Samples were mounted on conductive carbon tape for analysis with no other preparation.
- 2.2.2 STEM and EELS. Annular Dark Field (ADF) Scanning transmission electron microscopy (STEM) was performed in a JEOL ARM 200 CF system operated at 80 kV. During imaging, the estimated current density was 14.5 pA while using 68-175 mrad of the annular detector's inner and outer angles. The microscope is equipped with an SDD Jeol Centurio Energy-Dispersive X-ray (EDX) spectrometer and a GIF Quantum (Gatan) Dual Electron Energy Loss Spectroscopy (EELS) spectrometer. STEM samples were prepared by dispersing 5 mg of NF sample in ethanol and sonicated for 5 min. The suspension was drop cast on Cu grids.
- 2.2.3 X-ray diffraction. X-ray powder diffraction was performed in a Bruker D2 PHASER diffractometer, using Cu Ka radiation ($\lambda = 1.5418 \text{ Å}$) operated at 30 V, 10 mA, in a range

between $2\theta = 20-85^{\circ}$, employing a step size of 0.05° and a scan speed of 0.1° s⁻¹. A low background sample holder (Bruker) was used for the powder samples.

2.2.4 Chemical surface analysis. X-ray photoelectron spectroscopy (XPS) analysis was performed in a Kratos AXIS ULTRA spectrometer using monochromated Al Kα (1486.58 eV). The electron binding energies were referenced to aliphatic carbon C 1s at 284.8 eV. The obtained peak analysis was made using the XPSPEAK41 software. Construction and peak fitting of synthetic peaks in narrow region spectra used a Shirely-type background, and the synthetic peaks were of a mixed Gaussian (30%)-Lorentzian (70%) type.

2.3 Electrochemical characterization

2.3.1 Electrochemical flow cell. The electrochemical cell for CO₂RR consisted of a custom-made microfluidic flow cell using a filter-press configuration. First, catalyst ink was made to prepare the working electrodes. The catalyst ink consisted of a mixture of 3 mg of multiwalled carbon nanotubes (MWCNTs), 7 mg of catalyst material, 228 µL of 5 wt% Nafion perfluorinated solution (Nafion and catalyst + MWCNTs in 1:1 ratio by weight) and 600 µL of 2-propanol. Then, MWCNTs are added to the ink to increase the conductivity of the samples. In all samples, the final loading of the catalysts was 1.25 mg cm⁻², covering a geometrical area of 1.8 cm² of the cathode. The working electrode consists of carbon gas diffusion electrodes (GDE) (SIGRACET 25BC) sprayed with the catalyst ink and dried at 80 $^{\circ}$ C. Commercial Ir-MMO and leak-free Ag/AgCl electrodes were used as anode and reference electrodes, respectively. The Nafion 117 membrane separates the cathode and anode chambers. The potential was referred to against the reversible hydrogen electrode (RHE).

$$U^{\text{RHE}} = U(\text{Ag/AgCl}) + U_0(\text{Ag/AgCl}) + 0.059\text{pH}$$
 (1)

The electroreduction was performed by applying a potential of -0.85 V (vs. RHE) for 2 h, using 0.5 M KHCO₃ as the electrolyte in both chambers, circulated at 23 mL min⁻¹. CO₂ gas was fed at 11 mL min⁻¹. The pH value of the CO₂-saturated electrolyte at 40 °C was 7.9. The different temperatures for electroreduction (25, 30, 35, and 40 °C) were controlled by placing the electrolyte reservoirs in a hot water bath and keeping the cell insulated. The maximum temperature capacity of our electrochemical cell was 40 °C.

An online gas chromatograph (Agilent micro-GC) was connected to the electrochemical cell to analyze the gas products (H₂ and CO). No CH₄ was observed, possibly due to the low concentration, and thus, below the detection limit of the micro-GC. After electrolysis, liquid products were analyzed with HPLC (AMinex column HPX-87X from Bio-Rad). The eluent used was 5 mM of H₂SO₄ with a 0.6 mL min⁻¹ flow rate at 65 °C. Typically, 10 mL of collected catholyte was mixed with 4 M of H₂SO₄ to decrease from pH 7.9 to pH 1-3, corresponding to formic acid formation. It should be noted that all current densities are expressed as cathodic currents. Thus, a negative value was used in the manuscript.

The faradaic efficiency (FE) for the products was calculated according to the following equation:

$$FE (\%) = \frac{(N_r)(n)(F)(V)}{O}$$
 (2)

where N_r is the number of electrons involved in obtaining the product r (2e⁻ for the formate, CO, and H₂), n corresponds to the number of r moles generated, F is the Faraday's constant (96 485 C mol⁻¹), V is the molar flow rate of CO₂ and Q is the total charge passed during electrolysis.

2.3.2 Electrochemical surface area and electrochemical impedance. The electrochemical surface area for the samples was estimated by obtaining the corresponding double-layer capacitance ($C_{\rm dl}$) from cyclic voltammetry (CV). CV for the working electrodes in the non-faradaic region was recorded in 0.5 M KHCO₃, continuously purged with N₂ at scan rates from 5 mV s⁻¹ to 80 mV s⁻¹. The $C_{\rm dl}$ was estimated for all catalysts from the slope of the linear relationship between the current density in the non-faradaic region and the scan rates in the CV. Electrochemical impedance spectroscopy (EIS) was carried out in the presence of 0.5 M KHCO₃ as a supporting electrolyte. EIS was done by applying a sinusoidal signal of 10 mV amplitude in the frequency range from 10⁵ to 0.1 Hz at an employed potential of -1.1 V ν s. SHE in the absence of CO₂ at 25 °C, while in the presence of CO₂ at 25 and 40 °C.

2.3.3 *In situ* **DEMS.** An *in situ* DEMS cell at three electrodes was used to record the ionic current and faradaic current versus potential characteristics during cathodic polarization. As for DEMS, the same WE, CE, and RE were employed. The working electrode was prepared on a glassy carbon electrode (3 mm diameter), as described in the previous section. The ionic current (mass signal) for selected mass-to-charge ratios (m/z)was recorded simultaneously with the CV profiles at a scan rate of 1 mV $\rm s^{-1}$. The CV measurements were conducted in a $\rm CO_2$ saturated 0.5 M of KHCO₃ solution within a potential window from 0.45 to -1.13 V (vs. RHE) at 40 °C. The heating increment was made with a heating resistance controlled by an Ink-bird ITC-308-WIFI controller. The ionic current (mass signal) for selected mass-to-charge ratios (m/z) was also recorded simultaneously with the faradaic current-time profile (I-t). The electrochemical cell was connected to the quadrupole mass spectrometer (Prisma QMG220) at a working pressure of ca. 2.7 \times 10⁻⁵ mbar.

2.4 Theoretical methods

2.4.1 First-principles calculations. Spin-polarized Density Functional Theory (DFT) calculations were employed to investigate the structure and thermodynamic stability of $SnO_2(110)$ models (see 2.4.1.1) using the plane wave Vienna *Ab initio* Simulation Package (VASP).^{34,35} In our samples with and without Ni doping, the $SnO_2(110)$ surface is typically the most abundantly exposed surface corresponding with $2\theta=26.5^{\circ}$.

The generalized gradient approximation (GGA) with Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional was employed to compute the total energies.³⁶ The projector-augmented wave (PAW) pseudopotentials were used for the

calculations with an energy cutoff of 400 eV for the plane waves. 37,38 More in particular, 4, 6, 10, and 1 valence electrons were considered for Sn (5s²5p²), O (2s²2p⁴), Ni (3d⁸4s²), and H (1s1) atoms, respectively. The Grimme-D3(BJ) method was implemented in VASP 5.4.4 to account for the van der Waals interactions. 39,40 The atomic positions were optimized using the conjugate gradient algorithm with force and electronic convergence criteria of 0.01 eV Å^{-1} and 10^{-5} eV, a Gaussian smearing of 0.05 eV, and a $4 \times 3 \times 1$ Monkhorst-Pack k-point grid. ⁴¹ The Partial Hessian Vibrational Analysis (PHVA) was carried out only for the surface species and selected atoms at the active sites on the catalyst (involving three Sn and three bridging O atoms) surface while keeping the other atoms fixed (Fig. S1†). The numerical partial Hessian was calculated by displacing the unfixed atoms in x, y, and z-directions with ± 0.01 Å, and the corresponding vibrational modes were obtained by a singular value diagonalization procedure as implemented in the postprocessing toolkit TAMKIN.42 The zero-point corrections and free energy contributions to the reaction energies were determined from the PHVA-based partition functions to determine the pressure and temperature dependence of the Gibbs free energies.

2.4.1.1 Models. The $xNi@SnO_2(110)$ surface slabs (where x= 0, 1, 2) were constructed with 4 SnO₂ layers, the same number of layers used in previous studies. 43 We opted for the SnO₂(110) surface since it was known for its thermodynamic stability and has garnered significant interest in experimental and theoretical investigations. 44-48 The periodic slab models considered in the study were referred to as SnO2, Ni@SnO2, and 2Ni@SnO2 (Fig. 1). A vacuum of 15 Å was employed in the z-direction to avoid interactions between periodic images. The Ni@SnO₂ systems were constructed by replacing surface Sn atoms with Ni. To understand the effect of H adsorption on the studied systems during the CO2 reduction at the cathode, the 2-fold coordinated O atoms (denoted as Sn-O-Sn) on the surface were hydroxylated, forming Sn-(OH)-Sn. A subsequent H addition to Sn-(OH)-Sn results in adsorbed water bonded to two Sn-sites, i.e., Sn-(OH₂)-Sn. Additionally, larger supercell models were constructed to shed light onto the interface between NiO and SnO₂ (see section 15 in ESI for details†).

2.4.2 Surface Pourbaix diagram. Surface Pourbaix diagrams (SPDs) were used to investigate the thermodynamically stable terminations of all studied systems as a function of pH and electrode potential, $U_{\rm SHE}$ (Standard Hydrogen Electrode) at pH = 0, $P_{\rm H_2}$ = 1 bar, T = 26.85 °C, *i.e.*, 300 K.⁴⁹ Model systems up to a maximum coverage of 2H per bridging O were considered for all studied systems to construct the SPDs.

The adsorption of n hydrogens on the SnO₂ surfaces can be given by:

$$n(H^+ + e^-) + * \rightarrow nH^*$$
 (3)

where * represents the SnO₂(110) surface model onto which the hydrogen atoms can be adsorbed.

The change in Gibbs free energy upon n hydrogenation reactions with respect to the pristine $SnO_2(110)$ surface models (Fig. 1) is given by;

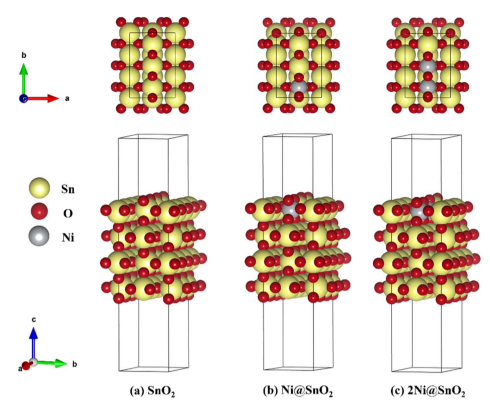


Fig. 1 Top and side view of chosen $SnO_2(110)$ model systems for (a) bare SnO_2 , (b) single Ni-doped SnO_2 (Ni@ SnO_2), and (c) double Ni-doped SnO_2 (2Ni@ SnO_2) systems.

$$\Delta G = G_{nH^*} - G^* - n \left(\frac{1}{2} G_{H_2} \right) \tag{4}$$

However, to calculate and construct the Pourbaix diagrams, the change in Gibbs free energy corresponding to a sequential hydrogenation reaction is calculated as a function of electrode potential and pH as follows;

$$\Delta G(pH, U_{SHE}) = G_{nH^*} - G_* - n \left(\frac{1}{2}G_{H_2}^0 - k_B T \ln(10)pH - eU_{SHE}\right)$$
(5)

where $G_{n\mathrm{H}^*}$ and G_* represent the free energy of the pristine and n times hydrogen-loaded model systems, respectively. The Gibbs free energy was computed for all terminations, and for a given pH and U_{SHE} conditions, the surface termination with the lowest $\Delta G(\mathrm{pH}, U_{\mathrm{SHE}})$ was considered when constructing the SPD. To compare two competing terminations, the $\Delta G(\mathrm{pH}, U_{\mathrm{SHE}})$ term is equated, *i.e.*, $\Delta G_{\mathrm{A}}(\mathrm{pH}, U_{\mathrm{SHE}}) = \Delta G_{\mathrm{B}}(\mathrm{pH}, U_{\mathrm{SHE}})$ to determine the pH $\nu s.$ U_{SHE} conditions of the equilibrium lines in the SPD. G_{nH^*} can be further described in terms of its vibrational contributions:

$$G_{nH^*} = E_{DFT} + E_{ZPVE} + \Delta E_{vib}(T) - TS_{vib} \approx E_{DFT} + F_{vib}(T)$$
 (6)

Here, E_{DFT} represents the total energy while F_{vib} is the Helmholtz vibrational energy. For the gas phase molecules, the

thermodynamic quantities such as ZPE, $\int_0^{298} C_v dT$, TS, were obtained from the ideal gas approximations from a previous report (Table S4†). The Gibbs free energy for the gas molecules at 25 °C (*i.e.*, 298 K) and 1 atm is given as:

$$G = H - TS = E_{\text{DFT}} + E_{\text{ZPE}} + \int_{0}^{298} C_{\text{v}} dT - TS + PV$$
 (7)

where $E_{\rm DFT}$, $E_{\rm ZPE}$, $\int_0^{298} C_{\rm v} {\rm d}T$ and S denote the electronic energy, zero-point vibrational energy, heat capacity, and entropy, respectively.

Results and discussions

3.1 Synthesis of NiO:SnO₂ nanofibers

Electrospinning is used to produce NiO:SnO₂ NFs. NFs are synthesized in various compositions, changing the molar ratios between Ni and Sn, *i.e.*, 75%:25% (NiOSnO25NF), 50%:50% (NiOSnO50NF), and 25%:75% (NiOSnO75NF). Similarly, NiO and SnO₂ single compositions have been electrospun. However, NiO and SnO₂ do not lead to NFs under similar electrospinning conditions. The latter includes NiO and SnO₂ formulations, which lose the NF shape after annealing. For instance, a single electrospun composition of NiO and SnO₂ NF can lead to the formation of nanoparticles after annealing (Fig. S2†). Like NiO and SnO₂, NiOSnO25NF led to nanoparticle formation after annealing (Fig. S3†). The previous results indicate that the spun NFs could not retain the fiber shape during annealing due to the unstable formation of metal oxide, leading to NF-shape

coalescence. Similar results have been observed for NiO systems, indicating that our NiCl $_2$ -PVP formulation leads to nanoparticle (NP) upon annealing. 12 This can also be the case for SnO $_2$, which could occupy additional metal oxide agents to maintain the NF shape. 50 Due to the lack of NF shape, NiOS-nO25NF has not been analyzed further. However, NiO and SnO $_2$ are still used as controls, along with NiOSnO50NP and NiOSnO75NP, which both lack NF shape.

NiOSnO50NF and NiOSnO75NF are inspected with STEM-ADF and STEM-EDX, as shown in Fig. 2. For NiOSnO50NF, Fig. 2a-d displays an NF-like shape with an NF diameter of 209 \pm 45 nm. The NiOSnO50NF comprises \sim 50 nm nanocrystals and multiple \sim 10 nm nanocrystals or smaller in diameter. The inner structure corresponds to a polycrystalline arrangement of nanoparticles with some gaps between them. Interestingly, in some cases, a row of nanocrystals forms a string along the NF with higher contrast (e.g., Fig. 2d and e, yellow arrows). From STEM-EDX mapping Ni, Sn, and O (Fig. 2e' and e'''), we have identified that these nanoparticle rows primarily comprise oxidized Sn. The contrast over the rows is attributed to densified oxidized Sn. In addition, within the NiOSnO50NF, oxidized Ni has been found to a lesser extent over the NF body; however, larger NiO particles, typically ~200 nm in diameter, decorate the NF morphology. A closer look at the interface between the NiO:SnO₂ nanocrystallites is discussed in Fig. S4.† For

NiOSnO75NF in Fig. 2f-i, the NF is more compact than in NiOSnO50NF, where nanocrystals are relatively smaller with a 35 nm diameter or less (e.g., Fig. 2i, yellow arrow). The presence of nanocrystals along the NF shape forming bright strings is less evident than in NiOSnO50NF. Moreover, NiO forms patches over the NF morphology (e.g., Fig. 2j, yellow arrow). The STEM-EDX mapping for Ni, Sn, and O in Fig. 2j' and j''' supports these results (see yellow arrows). The presence of Ni within the NF morphology is also verified by the STEM-EDX line scan (Fig. 2i and k, see dashed yellow line). The STEM-EDX line scan demonstrates that the Ni remains within the NF body but to a lower extent when compared to Sn (Fig. 2k). The NiOSnO75NF chemical species have also been investigated with EELS after CO₂ electroreduction (Fig. 2l and m). In Fig. 2l, the EELS signals for NiOSnO75NF after 22 h of CO2 electroreduction (NiOSnO75NF22h, black line) show that the oxidized tin resembles SnO₂ (blue line). However, we should not disregard the possibility of partially reduced SnO2 since the NiOSnO75NF22h has some similarities to SnO (red line). It is then suggested that NiOSnO75NF22h contains multiple oxidized Sn species (SnO_x). In Fig. 2m, the Ni L23 edge is used to obtain the L3/L2 ratio and determine the chemical environment of Ni-species in NiOSnO75NF after 2 h of CO2 electroreduction (NiOSnO75NF2h) and NiOSnO75NF22h. The EELS L3/L2 ratio shows that for NiOSnO75NF2h, more points have a wider L3/L2 ratio,12 indicating

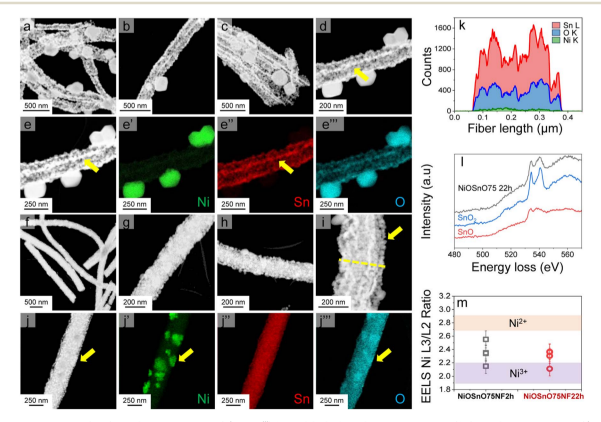


Fig. 2 STEM-ADF images (a—e) and STEM-EDX maps (e' and e''') for NiOSnO50NF. STEM-ADF images (f—j) and STEM-EDX maps (j' and j''') for NiOSnO75NF. The STEM-EDX line scan for NiOSnO75NF22h is shown in (k). EELS measurements NiOSnO75NF22h are shown in (l) and (m) for Sn and Ni, respectively. SnO_2 (blue line) and SnO_2 (red line) controls are presented in (l). In (m), the L3/L2 ratio of Ni L_{23} edge EELS for NiOSnO75NF2h and NiOSnO75NF22h are carried out over different surface areas of the crystallites.

a higher fraction of Ni²⁺ and Ni³⁺ species than NiOSnO75NF22h, which reveals a narrower L3/L2 ratio, close to Ni³⁺ species. The results denote that uncoordinated Ni³⁺ species, e.g., defects, are formed more due to longer electrolysis time attributed to the loss of NF shape after CO₂ electrolysis. 12

From the STEM-EDX results in Fig. 2, we can verify the presence of metal oxides. To confirm the oxide type, we look at the structural characteristics of NiOSnO50NF and NiOSnO75NF using XRD (Fig. 3), with NiO and SnO2 as controls. First, we describe the diffractograms of NiO and SnO2. NiO has several diffraction peaks at $2\theta = 37.2^{\circ}$, 43.3° , 62.9° , 75.4° , and 79.3° , which correspond to (111), (200), (220), (311), (222) crystallographic planes from NiO (JCPDS 65-6920), respectively.⁵¹ SnO₂ also shows diffraction peaks at $2\theta = 26.5^{\circ}$, 33.9°, 37.9°, 51.8°, 54.6°, 57.8°, 61.8°, 64.7°, 66.0°, 71.3°, and 78.7°, corresponding to (110), (101), (200), (211), (220), (002), (310), (112), (301), (220), and (321) crystallographic planes from SnO₂ respectively (ICPDS No. 41-1445).52-54 Comparing NiO and SnO2 with NiOSnO50NF and NiOSnO75NF, we observed that the crystallographic phases correspond to NiO and SnO₂. No changes in the diffraction peak positions for NiOSnO75NP used as control are observed. No evidence of a difference in crystallographic phase has been found, indicating that NiO and SnO₂ prevail in separate phases within the NF (Fig. 2 and 3).

The chemical environment is investigated with XPS to determine the type of species present over NiOSnO50NF and NiOSnO75NF. The XPS core spectra of Ni 2p, Sn 3d, O 1s, and Cl 2p for NiOSnO50NF and NiOSnO75NF are presented in Fig. 4.

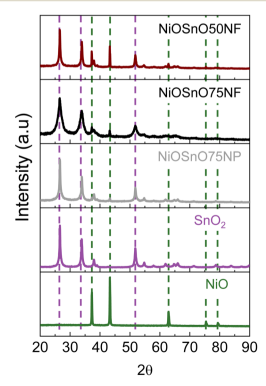


Fig. 3 XRD diffraction patterns for NiOSnO50NF and NiOSnO75NF. XRD diffraction patterns of NiOSnO75NP, SnO2, and NiO are included for controls.

NiO and SnO₂ controls are used for comparison. In Fig. 4a, Ni 2p comprises Ni $2p_{3/2}$ and Ni $2p_{1/2}$. The Ni $2p_{3/2}$ peak can be fitted into two components corresponding to Ni2+ and Ni3+ species, labeled in red. Ni2+ and Ni3+ peaks are located at 853.8 eV and 855.6 eV. 55-57 The presence of Ni³⁺ can be ascribed to the uncoordinated species, like defects. 12 The Ni³⁺/Ni²⁺ ratio for NiOSnO50NF is estimated to be close to 2.1, while NiOSnO75NF is around 8.5, indicating that Ni³⁺ is more prominent in NiOSnO75NF. Similar Ni³⁺/Ni²⁺ has been observed for NiOSnO50NP and NiOSnO75NP compared to the NF counterpart. Furthermore, a behavior opposite to NiOSnO50NF and NiOSnO75NF has been observed for NiO, in which the Ni³⁺/Ni²⁺ ratio is lower, ca. 1.3. Ni²⁺ and Ni³⁺ species have also been identified with EELS (Fig. 2), supporting our finding. The results demonstrate that the presence of Sn increases the amount of Ni³⁺ species. Next, we analyze the results for Sn 3d. In Fig. 4b, NiOSnO50NF and NiOSnO75NF show binding energy (BE) for Sn $3d_{5/2}$ around 486.3–486.5 eV, assigned to Sn⁴⁺ in SnO₂. 58-60 Compared to SnO₂ control with a BE around 486.9 eV associated with Sn⁴⁺,61 a shift to lower BE has been found for NiOSnO50NF and NiOSnO75NF. This shift can be related to reduced Sn species (e.g., SnO_r), similar to EELS, as shown in Fig. 21.

The XPS core spectra of O 1s and Cl 2p and the fitting curves have also been analyzed (Fig. 4c and d). O 1s core spectra for NiOSnO50NF and NiOSnO75NF show four different peaks resulting from NiO and SnO₂ formation within the NF body. The O(I) peak (BE 528.8-529.1 eV) is attributed to oxygen in NiO.55,56 The O(II) peak (BE 530.0-530.3 eV) is attributed to mixed oxygen species from NiO and SnO2.57,62,63 The O(III) peak (BE 531.3-531.7 eV) is attributed to surface OH groups. 56,63,64 The O(IV) peak (BE 532.9-533.5 eV) is attributed to chemisorbed water. 63,64 Similar results have been obtained for NiO. The O 1s XPS spectrum of NiO shows BE at 529.1 eV, 530.0 eV, 531.5 eV, and 533.3 eV, attributed to oxygen in NiO, 55,56 surface O²⁻ species,65,66 surface OH groups, possibly from uncoordinated Ni³⁺ species present in NiO.¹² As for SnO₂, the O 1s XPS spectrum shows BE at 530.6 eV and 531.8 eV, corresponding to oxygen in SnO₂ (ref. 67-69) and OH groups. 68,69 Cl 2p core spectra for NiOSnO50NF, NiOSnO75NF, NiO, and SnO2 show several peaks labeled as Cl(I), Cl(II), Cl(III), and Cl(IV). Cl(I) between 198.6-198.1 eV and Cl(II) between 199.7-200.2 eV are assigned to inorganic chlorine species.70-73 NiO reveals two additional peaks at Cl(III) at 200.9 eV and Cl(IV) at 202.5 eV, both corresponding to Cl- from different decomposed chemical species of chlorine salt.12,74 The concentration of Cl for all the samples remains similar, with an average atomic percentage of $1\% \pm 0.2$.

3.2 CO₂ electroreduction

The functionality of NiOSnO50NF and NiOSnO75NF for CO₂RR is assessed in Fig. 5. To elucidate the effect of NF functionality, NiOSnO50NF and NiOSnO75NF are compared to NiOSnO50NP and NiOSnO75NP, which lack the NF shape. Additionally, the results are contrasted with NiO and SnO2. The NiOSnO50NF, NiOSnO75NF, NiOSnO50NP, NiOSnO75NP, NiO, and SnO2 comparatives are conducted in a flow cell to demonstrate the

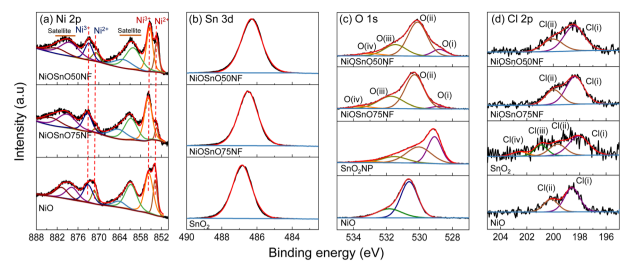


Fig. 4 (a) Ni 2p, (b) Sn 3d, (c) O 1s, and (d) Cl 2p XPS core spectra for NiOSnO50NF, NiOSnO75NF, NiO, and SnO₂.

importance of the synthesized multimetal oxide NFs. Finally, *in situ* DEMS experiments are discussed to shed light on the product pathways.

The experiments start with preparing the CO_2RR GDEs by spraying with ink containing MWCNTs, catalyst, and Nafion. The dried GDE is placed in the flow electrochemical cell using a three-electrode configuration containing a solution of 0.5 M KHCO₃ as an electrolyte. The CV in the presence of CO_2 shows the highest current density (J, mA cm⁻²) for the NiOSnO75NF, followed by NiOSnO50NF and SnO₂, with NiO showing the lowest in J (Fig. 5a). At 40 °C and -0.85 V νs . RHE, the product

distribution for NiOSnO50NP, NiOSnO75NP, NiO, and SnO $_2$ are also evaluated (Fig. 5b). It should be noted that three different potentials have been used, *i.e.*, -0.75, -0.85, and -0.95 V νs . RHE, and -0.85 V is selected since it produced the highest FE $_{\rm HCOO^-}$ and $J_{\rm HCOO^-}$. Compared to Fig. 5c and d, the results highlight the advantage of the synthesized NFs.

In Fig. 5b, FE for NiOSnO50NP and NiOSnO75NP at 40 °C displays the product distribution, where HCOO⁻, H₂, and CO are formed during CO₂ electroreduction. The FE for HCOO⁻, H₂, and CO for NiOSnO50NP are FE_{HCOO}⁻ = 3.8%, FE_{H₂} = 80.7% and FE_{CO} = 4.7% with partial J values of J_{HCOO}^- = -0.7 mA

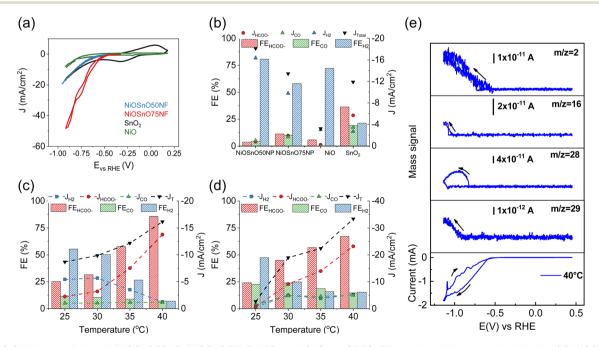


Fig. 5 (a) CV characteristics of NiOSnO50NF, NiOSnO75NF, NiO, and SnO₂ at 25 °C. FEs and partial current densities for (b) NiOSnO50NP, NiOSnO75NP, NiO, and SnO₂ at 40 °C, along with (c) NiOSnO50NF and (d) NiOSnO75NF at -0.85 V vs. RHE for 2 h over various temperatures, i.e., 25, 30, 35, 40 °C. (e) CV characteristic for NiOSnO75NF at 40 °C. In situ DEMS mass signals recorded at 40 °C are shown as a function of the applied potential (1 mV s⁻¹) for m/z = 2, m/z = 16, m/z = 28, and m/z = 29. In all cases, the pH of the bulk electrolyte remained at 7.9.

 ${\rm cm}^{-2}$, $J_{\rm H_2} = -14.7~{\rm mA~cm}^{-2}$ and $J_{\rm CO}$ of $= -0.9~{\rm mA~cm}^{-2}$. Similar product distribution is observed for NiOSnO75NP with FE_{HCOO}-= 11.2%, FE_{H_3} = 58.0%, and FE_{CO} = 10.1% have been found with $J_{\rm HCOO^-} = -1.7 \text{ mA cm}^{-2}$, $J_{\rm H_2} = -8.8 \text{ mA cm}^{-2}$ and $J_{\rm CO} =$ -1.5 mA cm⁻². For NiO, a prominent generation of H₂ is observed with $FE_{H_0} = 72.2\%$ and $J_{H_0} = -2.7$ mA cm⁻². For the same electrocatalyst, $FE_{HCOO^-} = 6\%$ and $J_{HCOO^-} = -0.2$ mA cm⁻², no CO has been detected. It could be argued that to increase HCOO⁻ formation, NiO should be selectively shaped as octahedra and not as mixed particle shapes (Fig. 2).12 SnO2 FE for HCOO⁻, H₂, and CO are 37%, 21%, and 19% with J_{HCOO^-} -5.1 mA cm^{-2} , $J_{\rm H_2} = -2.4 \text{ mA cm}^{-2}$ and $J_{\rm CO}$ of $= -3.1 \text{ mA cm}^{-2}$. The results demonstrate the importance of synergistic effects between NiO and SnO₂, particularly for the NF morphology

The effect of the temperature (25, 30, 35, and 40 °C) during CO₂R for NiOSnO50NF and NiOSnO75NF is evaluated in Fig. 5c and d. For NiOSnO50NF (Fig. 5c), the HCOO has a gradual increase in selectivity as the temperature increases, starting from FE_{HCOO}- = 25.2% at 25 °C and reaching a maximum of $FE_{HCOO^-} = 85.7\%$ at 40 °C. At 40 °C, the highest J_{HCOO^-} is observed (-13.8 mA cm⁻²). For FE_H, we observe a gradual decrease in H2 generation as a function of temperature, starting from $FE_{H_0} = 55.4\%$ at 25 °C and reaching a minimum of 7% at 40 °C. The $J_{\rm H_2}$ value for NiOSnO50NF follows a similar trend, from $J_{\rm H_2} = -5$ mA cm⁻² at 25 °C, reaching a minimum $J_{\rm H_2} =$ -1.2 mA cm^{-2} at 40 °C. FE_{CO} does not drastically decrease with increasing temperature, maintaining I_{CO} around -1 mA cm⁻² across the various conditions, with a FE $_{CO} = 7.2\%$ at 40 °C. Overall, in NiOSnO50NF, FE_{HCOO}- is favored as temperature increases while maintaining FE_{CO} constant and suppressing FEH. NiOSnO50NF and NiOSnO75NF increase the total current densities (J_T) with temperature.

Next, the electrochemical performance of NiOSnO75NF during CO₂ electroreduction is discussed (Fig. 5d). For HCOO⁻, a gradual increase in selectivity, with $FE_{HCOO^-} = 25\%$ at 25 °C and $FE_{HCOO^-} = 70\%$ at 40 °C, is observed. At 40 °C, the highest $J_{\rm HCOO^-}$ is observed with $-26.1~{\rm mA~cm^{-2}}$. The FE_H in NiOSnO75NF also follows a gradual decrease with increasing temperature, with $FE_{H_2} = 47.4\%$ at 25 °C and 15.2% at 40 °C. For H_2 , J remains at $J_{H_2} = -5$ mA cm⁻² from 30 °C to 40 °C. Lastly, FE_{CO} presents a gradual decrease with 22.6% at 25 °C and 14.5% at 40 °C with $J_{CO} = -5$ mA cm⁻² for temperatures similar to or higher than 30 °C. Although FE_{HCOO}- for NiOSnO75NF remained 15% lower than for NiOSnO50NF at 40 °C, NiOSnO75NF has a 2-fold increase in $J_{\rm HCOO}$. This 2-fold increase can be attributed to an increase in the electrochemical surface area (ECSA) as the obtained double-capacitance is higher for NiOS $nO75NF (4.68 \times 10^{-4} \text{ mF cm}^{-2}) \text{ than NiOSnO50NF } (3.78 \times 10^{-4} \text{ mF cm}^{-2})$ mF cm⁻²). Furthermore, the results at 40 °C for NiOSnO75NF are substantiated with EIS. EIS reveals less charge transfer resistance and an increased affinity in the presence of CO2 for NiOSnO75NF (Fig. S5 and Table S5†). Likewise, there is no significant effect when looking at the Tafel slopes in the presence of CO₂ (Fig. S6 and Table S6†). The Tafel slopes are somehow similar. Hence, the results indicate the existence of similar rate-determining steps in the presence of CO₂ for

temperatures close to 40 °C. Slight variations in the Tafel slopes are observed for temperatures close to 45 °C, suggesting a different rate-determining step associated with other processes, e.g., H2 competition. The results of the chronoamperometry from Fig. 5c and d are shown in Fig. S7.†

In short, a trade-off between selectivity and product yield should be found when assessing catalyst performance. However, NiOSnO50NF and NiOSnO75NF resulted in similar trends, elucidating temperature effects, which could be reasonably associated with favored reaction kinetics at high temperatures. 19-21 Such effects have not been observed during CO₂ electroreduction using synergistic catalysts shaped as NFs. Hence, the synergistic effects require an understanding of the reaction product to the fullest. Therefore, an in situ DEMS is assessed to generate insight into the reaction product pathway by detecting the formic acid (HCOOH) mass fragments for NiOSnO75NF, as it yielded the highest HCOO production at 40 °C (Fig. 5d). Mass spectrometric signals corresponding to H₂ (m/z = 2), methane (CH₄, m/z = 16), CO (m/z = 28), and HCO⁻ (m/z = 29) from HCOOH, 75 and CV are recorded simultaneously (Fig. 5e). It should be noted that mass m/z = 29 is selected as it is the most pronounced for HCOOH, and in the absence of CO₂, no CO2 reaction products are observed (Fig. S8†). Additionally, to corroborate the detection of HCOOH, formic acid is added to the electrolyte, and the mass signals associated with this organic compound are shown in Fig. S9.† Overall, the distribution of DEMS products confirmed our flow cell observations in Fig. 5d, except for CH₄, which could be expected to be below the detection limit of our gas chromatograph but captured by DEMS. It should be noted that other factors that might change reaction product selectivity to CH₄ can be related to the DEMS cell configuration as it can impact pH, generating some gradients.75

Lastly, we discuss the effect of uncoordinated Ni species found in NiOSnO NFs, which could have enabled the formation of HCOO-.12 We could expect NiO species to enhance hydrogenation over SnO2, which is more likely to be as partially reduced SnO2, i.e., SnOx, after 2 or 22 h CO2 electrolysis. Although at 2 h, the NF shape drastically changed its morphology (Fig. S10†), Ni and Sn species remained present even after 22 h of CO2 electrolysis (Fig. 2l and m). Furthermore, the Ni³⁺/Ni²⁺ ratios for NFs connect with the improved FE_{HCOO}at 40 °C. However, we should not disregard the Ni³⁺/Ni²⁺ ratio in NPs. For example, the Ni³⁺/Ni²⁺ ratio for NiOSnO75NP is 7.5, close to NiOSnO75NF (i.e., 8.5). The benefit of structuring becomes evident when comparing the Raman spectra in Fig. S11† for NiOSnO75NP and NiOSnO75NF. NiOSnO75NP contains more organic species than NiOSnO75NF, compromising the CO₂ reduction reaction activity (Fig. 2b and d).

The results highlight the advantage of the NF morphology as carbon is removed from the NiOSnO precursor due to the open fibrous structure. Similar effects have been observed for polymer-derived metal oxides, such as 3D-printed structures where carbon remnants are found.76 Hence, the carbon remnant could act as a blocking layer during CO₂RR, affecting NiOSnO75NP selectivity. This hypothesis is well aligned with NiOSnO75NF loaded with *t*-octylphenoxypolyethoxyethanol

(Triton $\times 100$) used as a surfactant, acting as a carbon-blocking agent without compromising the NF morphology after annealing (Fig. S12†). The electrochemical results of NiOSnO75NF loaded with surfactant demonstrate a change in the product distribution with low HCOO $^-$ selectivity over the explored temperature ranges (Fig. S13†). The results are substantiated further by ECSA. ECSA result for NiOSnO75NF is 4.68 \times 10 $^{-4}$ mF cm $^{-2}$, while in the presence of a surfactant or NiOSnO75NP, it decreases to 8.39 \times 10 $^{-5}$ mF cm $^{-2}$. Now that we have identified the importance of blocking agent-free catalysts, we propose a mechanism for the NiOSnONF using the most significant products (H $_2$ and HCOO $^-$), as shown in Fig. 5c and d.

3.3 CO₂RR mechanism

3.3.1 Surface Pourbaix diagrams. Before deriving the CO₂RR mechanism, evaluating the stability of the electrocatalyst and its surface termination under electrochemical conditions is essential. The ESI in Section 14, Table S7,† describes the stability of the electrocatalyst models in terms of their cohesive and surface formation energy. For models with an increasing Ni concentration in the SnO₂ surface, the cohesive and surface formation energies become more negative, indicating a favorable formation of Ni-doped SnO2 phases. For the pristine NiO, the cohesive energy is more positive (0.161 eV per atom) compared to the Ni-doped SnO2 systems (0.061 eV per atom and -0.040 eV per atom, respectively). SPDs have been shown to play a crucial role in elucidating thermodynamically stable terminations as a function of pH and electrode potential.12,77-79 The calculated SPDs at the experimental electrochemical conditions are shown in Fig. 6. The NiO:SnO₂ interface and stability vs. potential diagram for all models at pH = 0 is displayed in Fig. S14-S17.† In an aqueous environment, the surfaces and active sites tend to hydroxylate (Fig. 6). Fig. 6ac show the SPDs for the studied SnO₂(110) model systems with and without Ni doping, which all have 3 bridging oxygens represented by [*O, *O, *O]. In the following, hydrogen coverage refers to these three bridging oxygens, and therefore, 1 ML hydrogen coverage corresponds with the system represented as [*OH, *OH, *OH]. In the potential range of -1.5 to 1.5 V, five different terminations are found in the SPD of the studied pristine SnO₂(110) surface model (Fig. 6a). Above 1.21 V and low pH, the configuration with no adsorbed H (or H*) is stable. As the potential decreases, the adsorption of H gets pronounced, leading to a complete H* coverage of the bridging O atoms. For instance, between 1.21 V and 0.487 V at pH = 0, the termination with 0.33 ML H* is favored. The other stable terminations are 0.66 and 1.00 ML of H*. Following the hydration of the oxygens bridging two Sn atoms (Sn-O-Sn), H can further adsorb on the bridging hydroxyls, forming water molecules. The termination with 2.00 ML of H [*H₂O, *H₂O, *H₂O] is favored at potentials lower than -0.69 V. This also implies that the SnO₂ tends to reduce at cathodic electrode potentials, thus forming stable reduced surfaces which can be active for catalysis. The dashed line in red represents the standard OER and HER limits.

Next, the influence of Ni-doping of the $SnO_2(110)$ model system on the Pourbaix diagram is depicted in Fig. 6b and c.

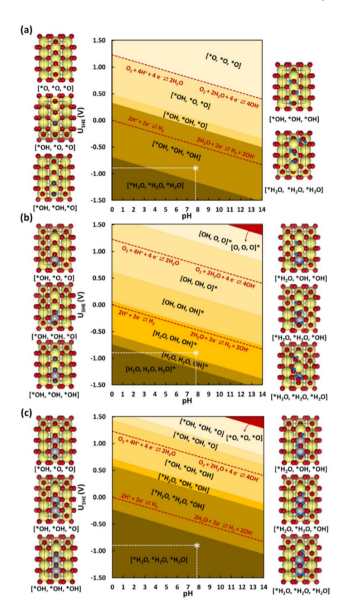


Fig. 6 Surface Pourbaix diagrams for the studied (110) surface models (a) SnO_2 , (b) $Ni@SnO_2$, and (c) $2Ni@SnO_2$. Surfaces with no H adsorbed are colored in red. Color codes: Sn (yellow), O (red), H (blue), and Ni (grey). The white star * in the diagram corresponds to typical experimental conditions used ($U_{SHE}=0.85$ V, pH = 7.9).

Unlike for SnO_2 , all H-covered terminations from 0.33 to 2.00 ML H* are present in the Pourbaix diagram for Ni-doped SnO_2 models at potentials between -1.5 and 1.5 V. In the case of a single Ni-doped $SnO_2(110)$ system, the surface with no H adsorbed is stable only at higher potentials (>2.14 V) and pH (>13). Between 1.5 and 0.06 V (at pH = 0), the two-fold bridging O* atoms (Sn-O-Sn) tend to get fully hydrogenated. At lower potentials (<0.06 V), the hydrogenated O atoms can adsorb H to form adsorbed water molecules. Below -0.74 V *versus* RHE, the surface is completely reduced with all two-fold bridging O atoms (Sn-O-Sn) forming water molecules. As a characteristic of H adsorption, on moving to a higher pH, the stable H terminations occur at lower potentials in the SPD due to the

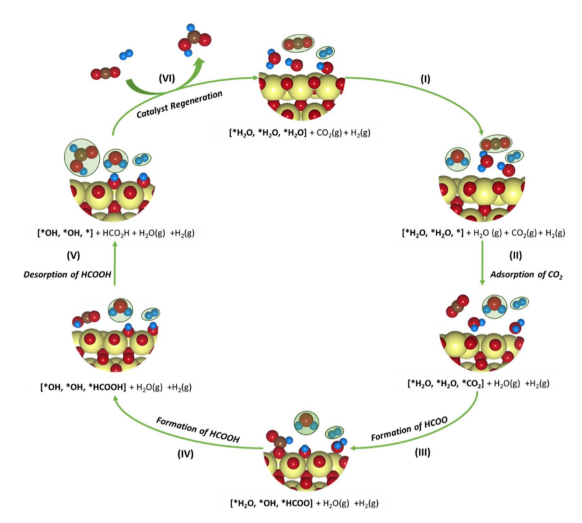


Fig. 7 Schematic representation of the plausible CO₂ reduction mechanism for the SnO₂-based model systems. The adsorbates in gaseous states are marked in green. Color codes: Sn (yellow), C (brown), O (red), H (blue).

shift of -59 meV per pH unit. Finally, another Sn atom is replaced with Ni to understand the effect of Ni concentration on the SPDs. For the 2Ni@SnO₂ surface model, all H terminations from 0.33 to 2.00 ML H* appeared in the Pourbaix diagram. The surface with no hydrogen appears only above 2.10 V and high pH. The termination with 1.00 ML coverage of H is favorable under OER conditions, while the surface with 1.66 ML H* coverage (2× H₂O*) appears at the HER limit. Interestingly, at the experimental conditions of U = -0.85 V and pH = 7.9 (Fig. 5), the surface with 2.00 ML ($3 \times H_2O^*$) is likely to be thermodynamically preferred. This also highlights that in CO₂RR conditions, the surface of the Ni-doped SnO₂ is partially reduced (e.g., SnO_r, Fig. 2 and 4), which could further tailor the electrocatalytic activity at the surface.80,81 Overall, the SPDs emphasize the reduction of the SnO₂-based catalyst surface, specifically the co-adsorption of water molecules at potentials of experimental interest.

3.3.2 CO₂RR mechanism. The SPDs allow us to determine the relevant active site model termination under operating conditions, and subsequently, CO₂ reduction pathways are determined. A typical reduction of CO₂ to HCOO⁻ (i.e., hereafter

HCOOH) over SnO2 occurs via a 2e pathway with the elementary steps described as:

$$CO_2 + * \rightarrow *CO_2 \tag{8}$$

$$*CO_2 + H^+ + e^- \rightarrow *HCOO$$
 (9)

*HCOO + H⁺ + e⁻
$$\rightarrow$$
 HCOOH + * (10)

where * denotes the active site of the catalyst. Although the proton-coupled electron transfer (PCET) mechanism assuming steps with H^+/e^- pairs (equivalent to $\frac{1}{2}H_2$) is vastly employed in the context of DFT-based studies, 43,48 it usually oversimplifies the catalytic surface in which protons can be supplied from the dissociation of water molecules at the electrode-electrolyte interface. T. Burdyny and W. A. Smith demonstrated that at current densities above 35 mA cm⁻², the proton for CO₂ reduction is supplied by the water molecules on the electrode surface, increasing the local interfacial pH.82 The resulting change in the local environment further influences the binding energies of intermediates and surface coverage on the electrode. The SPDs discussed previously highlighted this, showing that

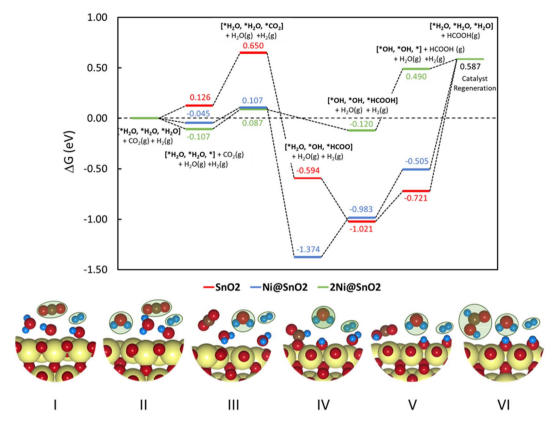


Fig. 8 Free energy diagram for CO_2 reduction to HCOOH over $SnO_2(110)$, $Ni@SnO_2(110)$, and $2Ni@SnO_2(110)$ electrocatalyst models. (Bottom panel) The binding modes of the adsorbates on the catalyst at different reaction states and the adsorbates in the gaseous state are marked in green. Color codes: Sn (yellow), C (brown), O (red), O (blue).

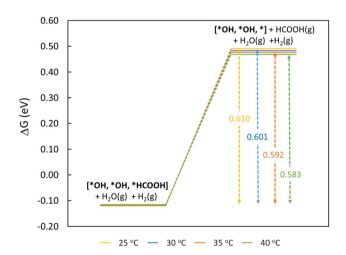


Fig. 9 Free energy diagram for rate-limiting step of HCOOH desorption as a function of temperature on the 2Ni $_{\odot}$ SnO $_{\odot}$ (110) model.

the bridged oxygen forms water molecules at lower potentials, resulting in an overall reduction of the catalytic surface. Fig. 7 presents a plausible mechanism for the CO₂RR to HCOO⁻ for the SnO₂-based models involving a PCET reaction considered in the study. It can be found that the active catalyst surface of 2Ni@SnO₂ is covered with 2.00 ML of H (*i.e.*, three H₂O per unit cell), which is represented as [*H₂O, *H₂O, *H₂O] following its

SPD. This model could approximate the local environment on the reduced surface more appropriately than the usual models that do not consider electrolyte species' co-adsorption. Although the three water molecules do not appear in the SPDs of SnO_2 and $Ni@SnO_2$ at the experimental conditions of -0.85 V and a pH of 7.9, we consider also 2.00 ML coverage models for comparison with the $2Ni@SnO_2$ system (Fig. 8).

To understand the thermodynamic feasibility of the reaction pathway proposed, we calculated the Gibbs free energy profile for each model system. The mechanism starts with the desorption of one of the H₂O molecules on 2Ni@SnO₂, leaving an empty site for CO₂ adsorption [*H₂O, *H₂O, *] (Fig. 8). From Fig. 8, the process of water desorption is exothermic and exergonic for the Ni-doped SnO₂ (110) systems compared to pure SnO₂. However, the subsequent CO₂ adsorption on the empty site is endergonic (Fig. 8), with positive reaction-free energies for SnO₂ (0.524 eV), Ni@SnO₂ (0.152 eV), and 2Ni@SnO₂ (0.194 eV) model systems which becomes more feasible if SnO₂ is Ni doped. Further, upon abstracting an H from a co-adsorbed water molecule, *CO2 can form *HCOO. For Ni@SnO2, the *HCOO intermediate is thermodynamically more stable (-1.374 eV, Fig. 8) compared to pure SnO_2 (-0.594 eV, Fig. 8). Interestingly, for 2Ni@SnO2, the CO2 molecule directly tends to form a stable HCOOH, surpassing the *HCOO intermediate state. The formation of *HCOOH from *HCOO and neighboring *OH2, is exergonic for SnO2 with a reaction-free energy of

−1.021 eV, whereas for the Ni@SnO₂ system, this process is endergonic (+0.391 eV) energy. For 2Ni@SnO2 the *HCOO intermediate is found to be protonated directly and form *HCOOH, an overall exergonic process with a reaction-free energy of -0.227 eV. From *HCOOH, the desorption of HCOOH is endergonic and requires 0.300 eV, 0.478 eV, and 0.610 eV for SnO₂, Ni@SnO₂, and 2Ni@SnO₂, respectively (Fig. 8). Therefore, it is clear that under certain reaction conditions, the desorption of HCOOH can become the ratelimiting step in the reaction. Especially, for the 2Ni@SnO2 system, with the highest adsorption free energy barrier for *HCOOH desorption, temperature facilitates the desorption process given the decreasing free energy differences from 25 °C to 40 °C (Fig. 9). These findings agree with the temperaturedependent faradaic efficiencies and partial current densities of CO₂ electroreduction (Fig. 5c and d). Finally, after the production of HCOOH, the catalyst needs to regenerate to continue with the catalytic cycle. The catalyst regeneration with two H⁺/e⁻ pairs is endergonic with the ΔG increasing in the order: 2Ni@SnO₂ (0.097 eV) < Ni@SnO₂ (1.092 eV) < SnO₂ (1.308 eV), however, this regeneration can happen electrochemically, and under cathodic potentials of -0.90 V vs. RHE this regeneration process will be spontaneous. The regeneration process is more favorable for the surfaces with more Ni species (e.g., Ni²⁺ and Ni³⁺), which indicates the influence of Ni doping on the catalytic activity of SnO₂.

Conclusions

NiOSnO NFs have been synthesized by electrospinning. NiOSnO NFs effectively function as electrocatalysts for the electrochemical CO2RR, yielding HCOO beyond the room temperature suitable to current electrolyzers. The highest faradaic efficiencies to formate are achieved with NiOSnO50NF and NiOSnO75NF at an electroreduction temperature of 40 °C. XPS and EELS analyses reveal a synergistic effect between the Ni and Sn species. Electrochemical measurements and in situ DEMS provide insights into product distribution during CO₂RR. Computational Pourbaix diagrams show that this synergistic effect arises from the dissolution of NiO under reducing conditions. DFT calculations show that embedding Ni in SnO₂ is energetically more favorable in addition to aiding the reduction of the SnO2 surface under relevant electroreduction conditions. The desorption of HCOOH is the rate-limiting step whose free energy decreases with increasing temperature from 25 °C to 40 °C, which agrees with the temperature-dependent faradaic efficiencies and partial current densities found during the experiments. Looking into the future, it is clear that catalysts like NiOSnO NFs can be further designed for other temperature conditions rather than room temperature and will, in the future, be used in CO₂ electrolyzer technologies over various temperature ranges. These findings underscore the significance of catalyst discovery and explore the potential for temperature-driven synergistic effects in metal oxide catalysts for CO₂ electroreduction.

Data availability

Data are available upon request from the authors.

Author contributions

M. A. R. O., R. L., M. V., and A. S. A. designed the experiments, analyzed the data, and wrote the first draft of the manuscript. M. A. R. O., M. S., C. F., and E. C. -M. synthesized the material and carried out electrochemical measurements. R. L. and M. V. performed the DFT calculations (1) to construct the computational Pourbaix diagrams, and (2) to unravel the relevant CO2 reduction pathways. F. R. -Z. performed STEMrelated analysis. The initial idea was coined by A. M., C. F., T. K., J. G. E., M. V., and A. S. A. All authors contributed to the final draft.

Conflicts of interest

The authors declare no competing interests.

Acknowledgements

The authors thank Mark Smithers and Gerard Kip (MESA+ Institute, University of Twente) for their support. The research leading to this report's results has received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation program (Grant Agreement No. 742004). R. L. (GOIPG/2022/442) thanks the Irish Research Council (IRC) for an IRC postgraduate fellowship. R. L. and M. V. acknowledge the Irish Centre for High-End Computing (ICHEC) for the computational facilities and support. M. S. and T. K. acknowledge funding from the Jane and Aatos Erkko foundation (USVA project). In addition, F. R.-Z. acknowledges the European Research Council (ERC) Starting Grant 123STABLE (Grant agreement ID: 852208).

References

- 1 V. Masson-Delmotte, P. Zhai, A. Pirani, S. L. Connors, C. Péan, S. Berger, N. Caud, Y. Chen, L. Goldfarb, M. I. Gomis, M. Huang, K. Leitzell, E. Lonnoy, J. B. R. Matthews, T. K. Maycock, T. Waterfield, O. Yelekçi, R. Yu and B. Zhou, IPCC, 2021: Climate Change 2021: the Physical Science Basis. Contribution of Working Group I to the Sixth Assessment Report of the Intergovernmental Panel on Climate Change, Cambridge University Press, 2021.
- 2 X. Bai, W. Chen, C. Zhao, S. Li, Y. Song, R. Ge, W. Wei and Y. Sun, Angew. Chem., 2017, 129, 12387-12391.
- 3 F. Köleli and D. Balun, Appl. Catal., A, 2004, 274, 237–242.
- 4 K. Fan, Y. Jia, Y. Ji, P. Kuang, B. Zhu, X. Liu and J. Yu, ACS Catal., 2020, 10, 358-364.
- 5 B.-Q. Miao, W.-S. Fang, B. Sun, F.-M. Li, X.-C. Wang, B.-Y. Xia and Y. Chen, Chin. J. Struct. Chem., 2023, 42, 100095.
- 6 R. Hegner, L. F. M. Rosa and F. Harnisch, Appl. Catal., B, 2018, 238, 546-556.

- 7 W. Yang, S. Chen, W. Ren, Y. Zhao, X. Chen, C. Jia, J. Liu and C. Zhao, *I. Mater. Chem. A*, 2019, 7, 15907–15912.
- 8 W. Zhu, R. Michalsky, Ö. Metin, H. Lv, S. Guo, C. J. Wright, X. Sun, A. A. Peterson and S. Sun, *J. Am. Chem. Soc.*, 2013, 135, 16833–16836.
- S. Liu, H. Tao, L. Zeng, Q. Liu, Z. Xu, Q. Liu and J. L. Luo, J. Am. Chem. Soc., 2017, 139, 2160–2163.
- 10 D. Gao, H. Zhou, J. Wang, S. Miao, F. Yang, G. Wang, J. Wang and X. Bao, *J. Am. Chem. Soc.*, 2015, **137**, 4288–4291.
- 11 J. Rosen, G. S. Hutchings, Q. Lu, R. V. Forest, A. Moore and F. Jiao, *ACS Catal.*, 2015, 5, 4586–4591.
- 12 M. A. Rodriguez-Olguin, C. Flox, R. Ponce-Pérez, R. Lipin, F. Ruiz-Zepeda, J. P. Winczewski, T. Kallio, M. Vandichel, J. Guerrero-Sánchez, J. G. E. Gardeniers, N. Takeuchi and A. Susarrey-Arce, *Appl. Mater. Today*, 2022, 28, 101528.
- 13 A. Vasileff, C. Xu, Y. Jiao, Y. Zheng and S. Z. Qiao, *Chem*, 2018, 4, 1809–1831.
- 14 S. Nitopi, E. Bertheussen, S. B. Scott, X. Liu, A. K. Engstfeld, S. Horch, B. Seger, I. E. L. Stephens, K. Chan, C. Hahn, J. K. Nørskov, T. F. Jaramillo and I. Chorkendorff, *Chem. Rev.*, 2019, 119, 7610–7672.
- 15 C. Englezos, A. Raman, D. Jonker, N. A. Ramos-Delgado, M. Altomare, H. Gardeniers and A. Susarrey, Chempluschem, 2024, 89, e202300763.
- 16 Y. Fang, X. Liu, Z. Liu, L. Han, J. Ai, G. Zhao, O. Terasaki, C. Cui, J. Yang, C. Liu, Z. Zhou, L. Chen and S. Che, *Chem*, 2023, 9, 460-471.
- 17 Y. Fang, L. Han and S. Che, *Chin. J. Struct. Chem.*, 2023, 42, 100107.
- 18 R. E. Vos, K. E. Kolmeijer, T. S. Jacobs, W. Van Der Stam, B. M. Weckhuysen and M. T. M. Koper, *ACS Catal.*, 2023, 13, 8080–8091.
- 19 T. Mizuno, K. Ohta, A. Sasaki, T. Akai, M. Hirano and A. Kawabe, *Energy Sources*, 1995, 17, 503–508.
- 20 H. Y. Kim, I. Choi, S. H. Ahn, S. J. Hwang, S. J. Yoo, J. Han, J. Kim, H. Park, J. H. Jang and S. K. Kim, *Int J Hydrogen Energy*, 2014, 39, 16506–16512.
- 21 A. Löwe, C. Rieg, T. Hierlemann, N. Salas, D. Kopljar, N. Wagner and E. Klemm, *ChemElectroChem*, 2019, 6, 4497–4506.
- 22 S. T. Ahn, S. Sen and G. T. R. Palmore, *Nanoscale*, 2022, 14, 13132–13140.
- 23 R. E. Vos and M. T. M. Koper, *ChemElectroChem*, 2022, 9, e202200239.
- 24 K. Bejtka, J. Zeng, A. Sacco, M. Castellino, S. Hernández, M. A. Farkhondehfal, U. Savino, S. Ansaloni, C. F. Pirri and A. Chiodoni, ACS Appl. Energy Mater., 2019, 2, 3081–3091.
- 25 M. He, B. Xu and Q. Lu, Chin. J. Catal., 2022, 43, 1473–1477.
- 26 S. Mu, H. Lu, Q. Wu, L. Li, R. Zhao, C. Long and C. Cui, *Nat. Commun.*, 2022, **13**, 1–8.
- 27 Y. Jiang, J. Shan, P. Wang, L. Huang, Y. Zheng and S. Z. Qiao, *ACS Catal.*, 2023, **13**, 3101–3108.
- 28 T. S. Bui, E. C. Lovell, R. Daiyan and R. Amal, *Adv. Mater.*, 2023, **35**, 2205814.
- 29 E. Castañeda-Morales, J. O. Peralta-Cruz, F. Ruiz-Zepeda, A. Susarrey-Arce, M. L. Hernández-Pichardo and A. Manzo-Robledo, *Mater. Today Energy*, 2024, 41, 101525.

- 30 M. A. Rodriguez-Olguin, R. N. Cruz-Herbert, H. Atia, M. Bosco, E. L. Fornero, R. Eckelt, D. A. De Haro Del Río, A. Aguirre, J. G. E. Gardeniers and A. Susarrey-Arce, *Catal. Sci. Technol.*, 2022, 12, 4243–4254.
- 31 M. A. Rodriguez-Olguin, H. Atia, M. Bosco, A. Aguirre, R. Eckelt, E. D. Asuquo, M. Vandichel, J. G. E. Gardeniers and A. Susarrey-Arce, *J. Catal.*, 2022, **405**, 520–533.
- 32 W. Ju, F. Jiang, H. Ma, Z. Pan, Y. B. Zhao, F. Pagani, D. Rentsch, J. Wang and C. Battaglia, *Adv. Energy Mater.*, 2019, **9**, 1901514.
- 33 X. Zong, Y. Jin, C. Liu, Y. Yao, J. Zhang, W. Luo, A. Züttel and Y. Xiong, *Electrochem. Commun.*, 2021, **124**, 106968.
- 34 G. Kresse and J. Furthmüller, *Comput. Mater. Sci.*, 1996, 6, 15–50.
- 35 G. Kresse and J. Hafner, Phys. Rev. B: Condens. Matter Mater. Phys., 1994, 49, 14251–14269.
- 36 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865–3868.
- 37 G. Kresse and D. Joubert, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **59**, 1758–1775.
- 38 P. E. Blöchl, Phys. Rev. B: Condens. Matter Mater. Phys., 1994, 50, 17953–17979.
- 39 D. G. A. Smith, L. A. Burns, K. Patkowski and C. D. Sherrill, *J. Phys. Chem. Lett.*, 2016, 7, 2197–2203.
- 40 S. Grimme, S. Ehrlich and L. Goerigk, *J. Comput. Chem.*, 2011, 32, 1456–1465.
- 41 H. J. Monkhorst and J. D. Pack, *Phys. Rev. B: Solid State*, 1976, 13, 5188–5192.
- 42 A. Ghysels, T. Verstraelen, K. Hemelsoet, M. Waroquier and V. Van Speybroeck, *J. Chem. Inf. Model.*, 2010, **50**, 1736–1750.
- 43 S. Ning, J. Wang, D. Xiang, S. Huang, W. Chen, S. Chen and X. Kang, *J. Catal.*, 2021, 399, 67–74.
- 44 D. Koziej, K. Thomas, N. Barsan, F. Thibault-Starzyk and U. Weimar, *Catal. Today*, 2007, **126**, 211–218.
- 45 I. Manassidis, J. Goniakowski, L. N. Kantorovich and M. J. Gillan, *Surf. Sci.*, 1995, **339**, 258–271.
- 46 J. Oviedo and M. J. Gillan, Surf. Sci., 2000, 463, 93-101.
- 47 L. Braglia, M. Fracchia, P. Ghigna, A. Minguzzi, D. Meroni, R. Edla, M. Vandichel, E. Ahlberg, G. Cerrato and P. Torelli, J. Phys. Chem. C, 2020, 124, 14202–14212.
- 48 C. Salvini, M. Re Fiorentin, F. Risplendi, F. Raffone and G. Cicero, *J. Phys. Chem. C*, 2022, 2022, 126–14441.
- 49 H. A. Hansen, J. Rossmeisl and J. K. Nørskov, *Phys. Chem. Chem. Phys.*, 2008, **10**, 3722–3730.
- 50 M. Safari, J. Mazloom, K. Boustani and A. Monemdjou, *Sci. Rep.*, 2022, **12**, 1–15.
- 51 Y. Luo, M. Weng, J. Zheng, Q. Zhang, B. Xu, S. Song, Y. Shen, Y. Lin, F. Pan and C. Nan, J. Alloys Compd., 2018, 750, 17–22.
- 52 X. Ye, W. Zhang, Q. Liu, S. Wang, Y. Yang and H. Wei, *New J. Chem.*, 2015, **39**, 130–135.
- 53 P. P. Dorneanu, A. Airinei, M. Grigoras, N. Fifere, L. Sacarescu, N. Lupu and L. Stoleriu, J. Alloys Compd., 2016, 668, 65–72.
- 54 N. Van Hieu, P. Thi Hong Van, L. Tien Nhan, N. Van Duy and N. Duc Hoa, *Appl. Phys. Lett.*, 2012, **101**, 1–5.
- 55 A. Kotta, E.-B. Kim, S. Ameen, H.-S. Shin and H. K. Seo, *J. Electrochem. Soc.*, 2020, **167**, 167517.

- 56 X. Xu, L. Li, J. Huang, H. Jin, X. Fang, W. Liu, N. Zhang, H. Wang and X. Wang, ACS Catal., 2018, 8, 8033-8045.
- 57 X. Xu, H. Zhang, Y. Tong, Y. Sun, X. Fang, J. Xu and X. Wang, Appl. Surf. Sci., 2021, 550, 149316.
- 58 H. Liu, F. Wang, K. Hu, B. Zhang, L. He and Q. Zhou, Nanomaterials, 2019, 9, 1250.
- 59 M. Taeño, D. Maestre, J. Ramírez-Castellanos, S. Li, P. S. Lee and A. Cremades, Nanomaterials, 2021, 11, 1-13.
- 60 Q. Ma, H. Li, J. Guo, S. Chu, Q. Zhang and Z. Lin, Mater. Sci. Semicond. Process., 2021, 128, 105762.
- 61 M. Kwoka, L. Ottaviano, M. Passacantando, S. Santucci, G. Czempik and J. Szuber, Thin Solid Films, 2005, 490, 36-42.
- 62 M. Kandasamy, A. Seetharaman, D. Sivasubramanian, Nithya, K. Jothivenkatachalam, N. Maheswari, M. Gopalan, S. Dillibabu and A. Eftekhari, ACS Appl. Nano Mater., 2018, 1, 5823-5836.
- 63 C. N. R. Rao, V. Vijayakrishnan, G. U. Kulkarni and M. K. Rajumon, Appl. Surf. Sci., 1995, 84, 285-289.
- 64 Z. Chen, T. Dedova, I. O. Acik, M. Danilson and M. Krunks, Appl. Surf. Sci., 2021, 548, 149118.
- 65 N. Weidler, J. Schuch, F. Knaus, P. Stenner, S. Hoch, A. Maljusch, R. Schäfer, B. Kaiser and W. Jaegermann, J. Phys. Chem. C, 2017, 121, 6455-6463.
- 66 B. Payne, M. Biesinger and N. McIntyre, J. Electron Spectrosc. Relat. Phenom., 2012, 185, 159-166.
- 67 J. Jeong and B. J. Lee, J. Nanosci. Nanotechnol., 2013, 13, 711-
- 68 R. Zhang, Z. Xu, T. Zhou, T. Fei, R. Wang and T. Zhang, J. Colloid Interface Sci., 2019, 557, 673-682.
- 69 P. G. Choi, N. Izu, N. Shirahata and Y. Masuda, Sens. Actuators, B, 2019, 296, 126655.

- 70 M. C. Biesinger, L. W. M. Lau, A. R. Gerson and R. S. C. Smart, Phys. Chem. Chem. Phys., 2012, 14, 2434-2442.
- 71 C. D. Wagner, W. M. Briggs, L. E. Davis, J. F. Moulder and G. E. Muilenberg, Handbook of X-Ray Photoelectron Spectroscopy, Perkin Elmer Corp, Physical Electronics Division, 1992.
- 72 C. A. Tolman, W. M. Riggs, W. J. Linn, C. M. King and R. C. Wendt, Inorg. Chem., 1973, 12, 2770-2778.
- 73 R. Félix, N. Llobera-Vila, C. Hartmann, C. Klimm, M. Hartig, R. G. Wilks and M. Bär, RSC Adv., 2018, 8, 67-73.
- 74 Y. C. Lin, Y. Y. Chen, B. Y. Yu, W. C. Lin, C. H. Kuo and J. J. Shyue, Analyst, 2009, 134, 945-951.
- 75 J. M. Mora-Hernandez, W. I. González-Suárez, A. Manzo-Robledo and M. Luna-Trujillo, J. CO2 Util., 2021, 47, 101504.
- 76 J. P. Winczewski, S. Zeiler, S. Gabel, A. Susarrev-Arce, J. G. E. Gardeniers and B. Merle, Mater. Des., 2023, 232, 112142.
- 77 A. Ngoipala, R. Lipin, R. L. Arevalo and M. Vandichel, Int. J. Hydrogen Energy, 2024, 53, 829-839.
- 78 R. Lipin, A. Ngoipala, R. L. Arevalo and M. Vandichel, *Int. J.* Hydrogen Energy, 2024, 61, 460-472.
- 79 J. W. Liao, X. Lu, B. Y. Huang, G. Q. Yu and X. B. Li, Int. J. Hydrogen Energy, 2021, 46, 9077-9086.
- 80 B. Kumar, V. Atla, J. P. Brian, S. Kumari, T. Q. Nguyen, M. Sunkara and J. M. Spurgeon, Angew. Chem., 2017, 56, 3645-3649.
- 81 H. Hu, L. Gui, W. Zhou, J. Sun, J. Xu, Q. Wang, B. He and L. Zhao, Electrochim. Acta, 2018, 285, 70-77.
- 82 T. Burdyny and W. A. Smith, Energy Environ. Sci., 2019, 12, 1442-1453.