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Core-shell AgSn@SnO_x electrodes prepared via dynamic hydrogen bubbling templating and galvanic replacement demonstrated selective electrocatalytic reduction of CO_2 towards HCOOH with a faradaic efficiency of $96 \pm 4.9\%$ and partial current density of $-10.46 \pm 0.35 \text{ mA cm}^{-2}$. DFT calculations revealed that the catalyst expedites the production of formic acid via adjusting the binding energy of the HCOO^* intermediate.

Given the growing concern over climate change, the replacement of fossil fuels with greener alternatives is a must.^{1,2} Continuously rising temperature levels have left little leeway for how much carbon dioxide (CO_2) emissions the globe can tolerate.^{2,3} One attractive solution is the conversion of CO_2 into value-added chemicals. To achieve this, appropriate catalysts are needed to reduce the activation energy barrier for such reactions, which will ultimately minimize the amount of energy required for the CO_2 reduction reaction (CO_2RR). An important subject matter for the CO_2RR is determining which product is ideal for large scale production. Many techno-economic analyses argue that C_1 (CO and formate) products hold the best economic feasibility, which involve only 2-electron transfer.^{4–6} Generally, these products involve the lowest overpotential with an already demonstrated high faradaic efficiency (FE) exceeding 80% using metal electrocatalysts such as Sn,^{7–9} Ag,¹⁰ Zn,^{11–13} Bi,^{7,14–16} and Co.^{17,18}

To achieve higher FEs, however, different material design strategies must be undertaken. Among these strategies, core-shell structures offer compelling results. Pérez *et al.* prepared $\text{SnIn}@\text{InSnO}_x$ nanoparticles for the CO_2RR with an 80% FE for formate production, which was ascribed to the In-rich surface

Dynamic hydrogen bubbling templated AgSn@SnO_x electrocatalyst for selective electrochemical CO_2 reduction: adjusting the binding energy of the HCOO^* intermediate†

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of the catalyst.¹⁹ In another study, Luc *et al.* manipulated the shell thickness of AgSn/SnO_x to optimize the performance of the catalyst to 80% FE for formate.²⁰ The authors argued that the shell thickness affected the stabilization of $^*\text{CO}_2^-$, which is correlated to formate production, largely due to the electronic and lattice changes in the material. In general, core-shell structures offer wide versatility, high conductivity, and high surface area, in which all produce high performance electrocatalysts for the CO_2RR . To design these catalysts, however, many complicated multi-step synthesis techniques that rely on seed generation and growth have to be employed, which hinders their large-scale implementation.²¹ Herein, we demonstrate a facile, two-step synthesis of AgSn@SnO_x core-shell structures using electrodeposition and galvanic displacement for the selective production of formate. The synthesized catalyst adjusted the binding energy of the HCOO^* intermediate, leading to formate production with a 96% FE and -10.5 mA cm^{-2} partial current density.

The synthesis of the AgSn@SnO_x catalysts involves dynamic hydrogen bubbling template (DHBT) electrodeposition and galvanic replacement, Fig. 1a. The first step includes the galvanostatic electrodeposition of Sn from a SnCl_4 solution on copper foil, resulting in the formation of

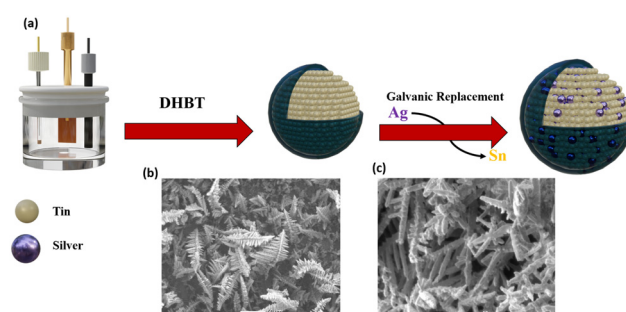


Fig. 1 (a) Synthesis steps of the AgSn@SnO_x catalyst and (b) SEM images of Sn@SnO_x and (c) AgSn@SnO_x (30 s).

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porous dendrite structures of Sn on the surface as shown in Fig. 1b. Afterwards, the Sn film was dipped in a solution of AgNO_3 to produce a AgSn alloy (eqn (1)). An ultrathin SnO_x layer is generated spontaneously when the samples are exposed to air.

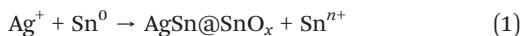


Fig. 2a shows the high resolution transmission electron microscopy (HRTEM) images of the synthesized $\text{AgSn}@\text{SnO}_x$, revealing the core-shell structure of the AgSn alloy along with an ultrathin layer of SnO_x with an average thickness of 3.55 nm. Both the Sn and Ag metals have different surface energies. This drives the spontaneous diffusion of bulk Sn atoms to the surface, where they are subsequently oxidized into a layer of SnO_x as they come into contact with air. The overall result of the Sn atoms' behaviour is that less Ag atoms are detected on the surface (compared to the bulk) despite the galvanic displacement reaction as listed in Table 1.²² Additionally, some neighbouring lattice fringes (Fig. 2b) show a diffraction spacing of 0.259 nm, verifying the formation of an intermetallic AgSn (110) phase in the bulk.²³ The X-ray diffraction (XRD) patterns (Fig. 2c) revealed no metal oxide phases, confirming the amorphous nature of the SnO_x shell. As for the X-ray photoelectron spectroscopy (XPS) measurements, the survey spectrum (Fig. S1†) shows three peaks corresponding to O1s, Ag 3d, and Sn 3d. The O1s

Table 1 Elemental analysis of $\text{AgSn}@\text{SnO}_x$ (30 s) using the XPS and EDX techniques

Elemental analysis technique	Sn	O	Ag
EDX	88%	4%	8%
XPS	58.96%	40.68%	0.35%

spectrum (Fig. S1†) shows three peaks at binding energies of 531.33, 532.51, and 533.64 eV, corresponding to (Ov), (Sn–O), and adsorbed oxygen species, respectively.²⁴ The deconvolution of the Sn 3d spectrum generates two peaks characteristic of $\text{Sn}^{4+/2+}$ (487.88 and 496.28 eV).²⁵ It is worth mentioning that the absence of Sn^0 peaks is likely due to the *ex situ* XPS measurement that results in excessive oxidation of the sample upon exposure to air for a long time; thus, the oxide species peaks overlapped with the metallic Sn^0 peaks. As for the Ag 3d spectrum (368.7 eV and 374.82 eV), it exhibits no oxide species, which indicates that the surface is predominantly covered with SnO_x . The elemental composition of the bulk and surface is further investigated *via* XPS and energy dispersive X-ray spectroscopy (EDX). Each technique measured the elemental distribution at a unique penetration depth.²⁶ XPS data (Table 1) show a lower atomic percentage of Ag compared to that from EDX analysis, leading to the conclusion that Ag diffuses into the bulk during the synthesis.

The electrochemical CO_2 reduction activity of $\text{AgSn}@\text{SnO}_x$ (30 s) was examined using linear sweep voltammetry (LSV) in CO_2 -saturated 0.1 M KHCO_3 and N_2 -saturated 0.1 M Na_2SO_4 . The pH under both conditions was confirmed to be the same (pH = 7.2). The onset potential in the case of CO_2 purging was measured to be less negative (−0.624 V vs. RHE) compared to that under N_2 purging conditions (−0.928 V vs. RHE). This likely indicates the electrochemical selectivity of $\text{AgSn}@\text{SnO}_x$ (30 s) towards CO_2 reduction instead of the hydrogen evolution reaction (HER) with a potential difference of 304 mV. Fig. S2† shows a positive shift in the onset potential of $\text{AgSn}@\text{SnO}_x$ (30 s) compared to that of bare Sn, indicating the formation of a new bulk Ag–Sn intermetallic phase rather than a simple interfacial alloy with superior catalytic activity towards CO_2 reduction.²⁷

The electrocatalytic performance of the different samples was further investigated at different overpotentials, Fig. S3.† We observed a volcano-like relationship between the galvanic replacement time and the faradaic efficiencies (FEs) and the partial current densities of HCOOH as shown in Fig. 3a and b. The maximum was observed at −0.9 V vs. RHE. The $\text{AgSn}@\text{SnO}_x$ (30 s) catalyst showed selectivity towards HCOOH with a FE of $96 \pm 4.9\%$ and partial current density of $-10.46 \pm 0.35 \text{ mA cm}^{-2}$. In contrast, the pure Sn catalyst exhibited weaker CO_2RR performance (FE = $40.41\% \pm 8.90\%$) under the same operating conditions. This enhancement likely originates from the superior surface roughness of $\text{AgSn}@\text{SnO}_x$, which can boost CO_2 gas–surface interactions, resulting in higher catalytic efficiency. Moreover, the $\text{AgSn}@\text{SnO}_x$ catalysts showed negligible HER activity, while

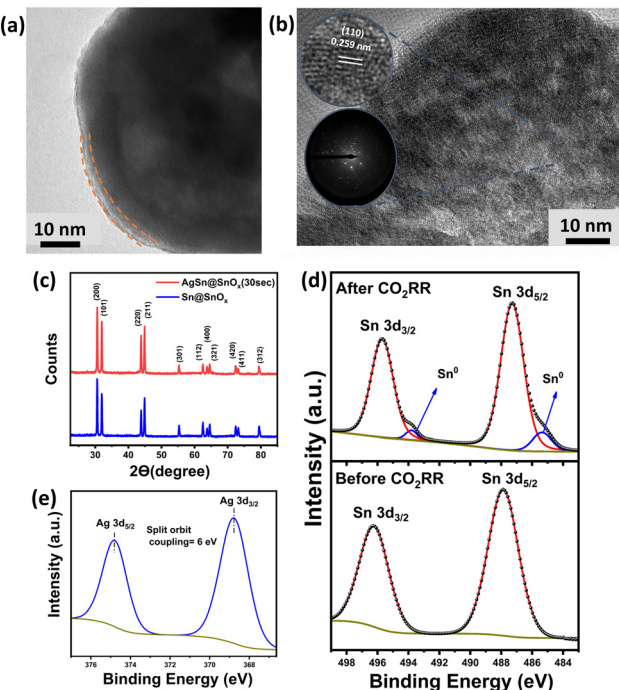


Fig. 2 (a) HRTEM image of $\text{AgSn}@\text{SnO}_x$ showing the thickness of the oxide layer, (b) d -spacing of the AgSn intermetallic phase and SAED pattern of $\text{AgSn}@\text{SnO}_x$ (30 s), (c) XRD patterns of $\text{Sn}@\text{SnO}_x$ and $\text{AgSn}@\text{SnO}_x$ (30 s), and (d) XPS patterns of $\text{AgSn}@\text{SnO}_x$ (30 s) before and after electrolysis depicting the Sn 3d spectrum, and (e) Ag 3d spectrum.

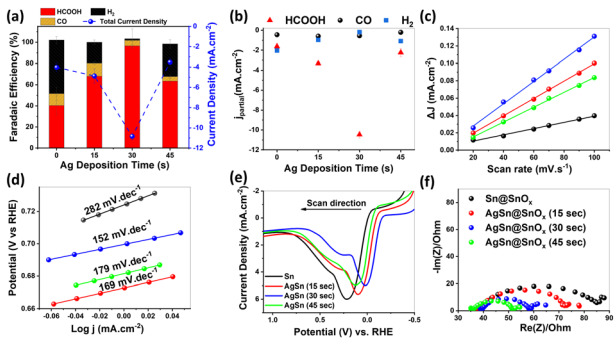


Fig. 3 (a) Faradaic efficiency and total current density as a function of the galvanic replacement time, (b) partial current densities of HCOOH, CO, and H₂ as a function of the galvanic replacement time, (c) linear regression between the cathodic and anodic current differences and scan rates, (d) Tafel analysis for the different catalysts, (e) oxidative linear sweep voltammetry conducted in Ar-saturated 0.1 M KOH and at a scan rate of 50 mV s⁻¹, and (f) EIS patterns for different samples in CO₂ saturated 0.1 M KHCO₃ within a frequency range of 100 kHz-20 mHz.

the Sn catalyst showed a modest suppression of the HER (50.53% \pm 3.41). The HER is the main competitive reaction in CO₂ electrolysis. On the other hand, the AgSn@SnO_x catalysts showed a plateau of the CO FE, despite the presence of an efficient CO promoter (*i.e.* Ag metal). This finding provides strong evidence that the surface is predominantly covered by a SnO_x shell instead of Ag.

The capacitive behavior of each sample was examined in order to study the intrinsic electrocatalytic activity of each sample. The double layer capacitance (C_{dl}), derived from cyclic voltammetry scanned in the non-faradaic region (Fig. S4†), showed the following trend: AgSn@SnO_x (30 s; 1280.41 μ F cm⁻²) > AgSn@SnO_x (15 s; 1000.35 μ F cm⁻²) > AgSn@SnO_x (45 s; 858.12 μ F cm⁻²) > Sn@SnO_x (359.34 μ F cm⁻²) as depicted in Fig. 3c. These results support the observed volcanic electrocatalytic performance. However, the j_{HCOOH} normalized by the electrochemical active surface area (ESCA) manifests a similar trend to the uncorrected j_{HCOOH} , indicating the minor impact of the ESCA on the catalytic performance.²⁸ Furthermore, AgSn@SnO_x (30 s) produced a plateau current density of 10 mA cm⁻² with a FE_{HCOOH} of 87.46% @ -0.9 V after 1800 min of electrolysis (Fig. S5†). The XPS spectrum of Sn 3d after the CO₂RR showed metallic Sn peaks at 485.31 and 493.79 eV, indicating the partial reduction of the SnO_x layer during the CO₂RR process.

The CO₂RR has a complicated mechanism that involves sluggish kinetic limitations related to the charge transfer and the geometrical changes of CO₂ molecules at the electrode-electrolyte interface.²⁹ Tafel analysis is the best choice to investigate the mechanistic pathways and elucidate the rate-determining step in the CO₂ reduction process. Among all samples, AgSn@SnO_x (30 s) displayed the lowest Tafel slope (152 mV dec⁻¹), verifying the faster reaction kinetics compared to its counterparts, as depicted in Fig. 3d. This value is a staunch indicator of *CO₂⁻ radical intermediate formation in the first electron transfer step.³⁰ With that in

mind, the CO₂ reduction mechanism is controlled by other factors such as the operating potential and mass transport limitations that provide a more accurate manifestation of the mechanism, so these might be discussed in the future.

Moreover, we conducted oxidative LSV to determine the adsorption affinity of HO⁻ anions as a surrogate to *CO₂⁻ radicals on different samples (Fig. 3d).^{31,32} The results demonstrated the coincidence between the kinetic findings of Tafel analysis and the adsorption affinity of HO⁻ anions on the different surfaces, indicating the better performance of AgSn@SnO_x (30 s) relative to the other samples, with the lowest adsorption energy of 0.015 V. These findings are attributed to the *in situ* formation of oxygen vacancies under the used cathodic conditions (*i.e.*, partial reduction of the oxide layer). This, in turn, stabilizes the *CO₂⁻ radicals, and results in a lower binding energy. Furthermore, EIS was used to uncover the origin of the bulk component-charge transfer relationship within the different catalysts. Nyquist plots (Fig. 3e) revealed a considerable enhancement in the charge transfer, leading to the conclusion that silver atoms diffuse to the bulk, forming a Ag-Sn intermetallic phase that immensely ameliorates the conductivity of the bulk with respect to that of pure Sn.

In fact, there are other factors that can verify the catalytic activity such as the surface energetics and the optimal binding energy of the intermediate on the surface. To this end, DFT calculations were conducted to investigate the influence of the AgSn@SnO_x core shell on the CO₂ reduction reaction (CO₂RR) activity. Fig. 4a illustrates the calculated Gibbs free energy changes for the CO₂RR and its corresponding adsorption intermediates. The formation of the HCOO* intermediate was found to be exothermic, whereas the formation process of *COOH was determined to be endothermic. Consequently, in the production of formic acid, the HCOO hydrogenation step sets the limiting potential. In contrast, in the production of CO, the rate-determining step is the formation of COOH.³³ The

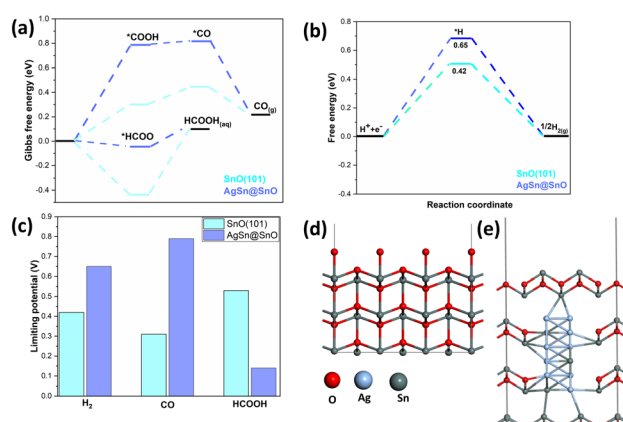


Fig. 4 Gibbs free energy diagrams for the a) CO₂RR and b) HER paths on SnO (101) and AgSn@SnO, c) limiting potentials for H₂, CO, and HCOOH formation over SnO (101) and AgSn@SnO, and d and e) optimized geometries for SnO (101) and AgSn@SnO.

implementation of the AgSn@SnO_x core shell lowers the binding strength of the HCOO^* intermediate relative to its counterpart on SnO (101), resulting in a decrease in the rate-determining step for formic acid production and an increase in the limiting potential for CO generation at the AgSn@SnO_x core shell.

Additionally, a Gibbs free energy diagram for the hydrogen evolution reaction (HER) process (Fig. 4b) was established, along with the corresponding structures of reaction intermediates, demonstrating that the AgSn@SnO_x core shell effectively suppresses hydrogen evolution by weakening the attachment of ${}^* \text{H}$ intermediates. In Fig. 4c, the limiting potential of each product in both systems is shown. In the case of the AgSn@SnO_x catalyst, the rate-determining step for producing formic acid is the lowest among the three products, referring to its prominence as the main product of the CO_2RR . Notably, the AgSn@SnO_x catalyst exhibits a further reduction in the rate-determining step for producing formic acid, while the limiting potentials for CO and H_2 generation are increased compared that for SnO (101). Therefore, based on the DFT calculations, implementing the AgSn@SnO_x core–shell structure enhances the selectivity for both formic acid and C_1 products. The optimized geometries for SnO (101) and AgSn@SnO_x are shown, for clarification, in Fig. 4d and e, respectively.

To conclude, a facile electrochemical synthesis of 3D hierarchical porous AgSn@SnO_x core–shell catalysts has been demonstrated as efficient candidates for CO_2 reduction to formate. The AgSn@SnO_x (30 s) catalyst showed excellent selectivity towards formate ($\text{FE}_{\text{HCOOH}} = 96\% \pm 4.90$; $j_{\text{HCOOH}} = -10.5 \text{ mA cm}^{-2}$ at -0.9 vs. RHE) with negligible HER activity. Tafel analysis and adsorption affinity studies suggest that AgSn@SnO_x (30 s) has faster reaction kinetics and the lowest adsorption energy, implying the formation of oxygen vacancies under cathodic conditions, which stabilize ${}^* \text{CO}_2^-$ radicals and achieve lower binding energy. DFT calculations showed that the AgSn@SnO_x core–shell structure accelerates the formation of formic acid by modifying the binding energy of the HCOO^* intermediate. Additionally, this structure improved the faradaic efficiency of C_1 production by suppressing the competitive hydrogen evolution reaction (HER), which is considered the main side reaction in the CO_2RR . The AgSn@SnO_x catalyst stands out as one of the most efficient electrocatalysts for CO_2 reduction to formate, when compared to other formate-selective electrocatalysts (Fig. S9, Table S3†). It demonstrated superior performance in terms of formate partial current density and formate faradaic efficiency. All in all, AgSn@SnO_x core–shell catalysts showed great potential for efficient CO_2 reduction to formate, which could have significant implications for sustainable energy production.

Conflicts of interest

There are no conflicts to declare.

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