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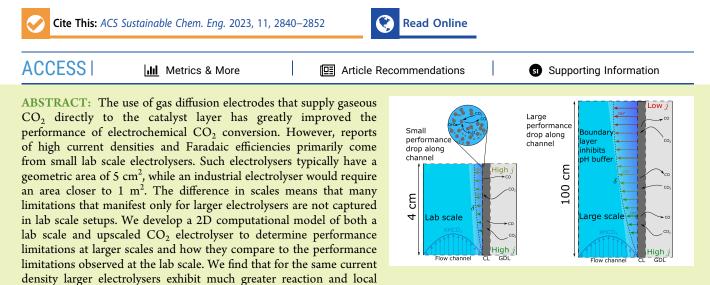
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Inhomogeneities in the Catholyte Channel Limit the Upscaling of CO₂ Flow Electrolysers

Joseph W. Blake,* Vojtěch Konderla, Lorenz M. Baumgartner, David A. Vermaas, Johan T. Padding, and J. W. Haverkort



environment inhomogeneity. Increasing catalyst layer pH and widening concentration boundary layers of the KHCO₃ buffer in the electrolyte channel lead to higher activation overpotential and increased parasitic loss of reactant CO_2 to the electrolyte solution. We show that a variable catalyst loading along the direction of the flow channel may improve the economics of a large scale CO_2 electrolyser.

KEYWORDS: CO_2 reduction, electrolysis, gas diffusion electrode, scale up, variable catalyst loading, parasitic reactions

INTRODUCTION

CO₂ from atmospheric or industrial sources can be electrochemically converted to valuable chemicals and fuels, leading to carbon neutral energy storage solutions and chemical feedstocks. The use of gas-diffusion electrodes (GDEs) allows the problems of low CO_2 solubility and diffusivity in aqueous electrolytes to be minimised, leading to high current density lab scale electrolysers.¹ However, the industrial realisation of this technology is hindered by a poor understanding of how these processes work in industrially relevant conditions. A rough benchmark of a minimum current density of 200 mA cm^{-2} has been reported for commercial feasibility,²⁻⁴ and this current density must be achieved at industrial scales. The typical lab scale cell has a geometric surface area around 5 cm² but an industrial scale cell would be significantly larger: Verma et al. proposed a surface area of 400 cm² per cell,³ while analyses that make comparisons to alkaline water electrolysis propose even surface areas from 1000 cm² to 1 m² and above.^{5,6}

However, upscaling in the flow direction leads to issues that negatively affect the performance of larger electrolysers. The high pH gradients, which have little impact on short lab scale electrolysers, lead to excessive pH increase in the extended flow direction. This high pH induces a Nernstian potential shift and depletes CO_2 through the carbonate equilibrium reactions, leading to reduced current density, Faradaic efficiency (FE), and reactant utilisation. However, the lack of experimental upscaling studies means the mechanisms behind these issues and the severity of their effects are poorly understood. In this paper we investigate the effect of scale by developing a 2D model of a typical lab scale CO_2 electrolysis cell and compare it to models that extend the geometry in the flow direction to 1 m. We identify the causes and magnitude of the performance loss and provide strategies to minimise the issues.

THEORY

Figure 1 shows a schematic diagram of an electrolyser setup^{7,8} for CO_2 electrolysis flow cell with a liquid electrolyte channel. This channel is sometimes replaced with a membrane-electrode assembly (MEA),^{9,10} and similarly, it is not unusual

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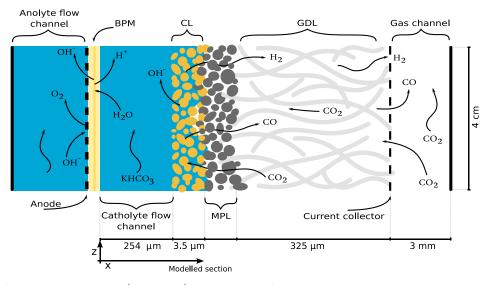


Figure 1. Diagram of a typical electrolysis cell (not to scale). From right to left: the gas channel, which supplies gas phase CO_2 ; the gas-diffusion layer (GDL), a dry macroporous structure through which CO_2 can permeate; the microporous layer (MPL), a thin hydrophobic layer that prevents electrolyte leakage into the gas-diffusion layer while allowing CO_2 dissolution into the liquid phase; the catalyst layer (CL), in which liquid electrolyte and catalyst particles embedded in the porous structure facilitate the electrochemical reaction with the dissolved CO_2 ; the catholyte flow channel, through which liquid electrolyte is supplied; a bipolar membrane (BPM), in which water dissociation supplies H⁺ and OH⁻ ions to the respective sides; the anode, on which the counter reaction is performed; and the anolyte flow channel.

for an anion-exchange membrane $(AEM)^{11-13}$ to be used in place of the bipolar membrane (BPM) shown.¹⁴⁻¹⁸ Gaseous CO₂ enters through the gas channel and passes through the macroporous gas diffusion layer (GDL) and microporous layer (MPL) to enter the liquid catholyte in the porous catalyst layer (CL), in which the heterogeneous electrochemical reaction takes place on the catalyst particles suspended in ionomer in the CL. The porous structures in the cell are modelled as macrohomogeneous domains characterised by porosity, ε_i permeability, κ , and volumetric surface area, a_v . While it is possible for the exact location of the gas-liquid interface to vary in the neighbourhood of the CL-MPL boundary when hydrophobic CLs are used,^{19,20} it is assumed in the model that the CL remains fully saturated with liquid electrolyte, while the MPL and GDL remain entirely dry at all times. This allows the combination of the GDL and MPL into one effective diffusion layer, a practice that is common in the determination of bulk properties of commercial GDLs supplied with preprinted MPLs.

Reactions. The ionomer in the CL is considered to be impregnated with evenly distributed Ag catalyst particles that facilitate heterogeneous reactions, and due to the high Faradaic efficiency (FE) of Ag catalysts towards CO, we consider only the reduction of CO_2 to CO

$$CO_2 + H_2O + 2e^- \rightarrow CO + 2OH^-$$
(1)

and the competing hydrogen evolution reaction (HER),

$$2H_2O + 2e^- \rightarrow H_2 + 2OH^-$$
(2)

While there is an argument for the HER pathways involving H⁺ and HCO₃⁻ to be included, these pathways are prevalent in near-neutral pH environments,²¹ and previous modelling studies predict that the CL environment will be very basic at current densities greater than 100 mA cm⁻²²² and that this pH will only increase in the flow-wise direction.²³ On the anode, the oxygen evolution reaction (OER),

$$4OH^- \rightarrow O_2 + 2H_2O + 4e^- \tag{3}$$

takes place.

A major complication distinguishing CO_2 electroreduction from similar technologies arises from the homogeneous reaction of CO_2 with OH^- to form bicarbonate and carbonate. These reactions can take both acidic and basic pathways, neither of which can be neglected. The basic

$$\operatorname{CO}_2 + \operatorname{OH}^- \stackrel{k_1}{\underset{k_{-1}}{\rightleftharpoons}} \operatorname{HCO}_3^-$$
 (4)

$$HCO_{3}^{-} + OH^{-} \underset{k_{-2}}{\overset{k_{2}}{\leftrightarrow}} H_{2}O + CO_{3}^{2-}$$
(5)

and acidic

$$CO_2 + H_2O \stackrel{k_3}{\underset{k_{-3}}{\rightleftharpoons}} H^+ + HCO_3^-$$
(6)

$$\mathrm{HCO}_{3}^{-} \stackrel{k_{4}}{\underset{k_{-4}}{\leftrightarrow}} \mathrm{H}^{+} + \mathrm{CO}_{3}^{2-} \tag{7}$$

reactions have the same ratios of CO_2 , HCO_3^- , and CO_3^{2-} at equilibrium for a given pH due to their coupling in the water self-ionisation reaction

$$H_2 O \rightleftharpoons_{k_{-w}}^{k_w} H^+ + O H^-$$
(8)

Furthermore, the equilibrium constants, $K_i = \frac{k_i}{k_{-i}}$, for the acidic reactions are related to the basic reactions through $K_3 = K_1 K_w$ and $K_4 = K_2 K_w$. However, the reactions have different kinetic rates and as such are each modelled individually.

The reaction rates per unit volume for species i in heterogeneous reaction r are given by

$$R_{\rm ct,i} = \sum_{r} \frac{J_r \nu_{i,r} a_{\rm v}}{n_r F} \tag{9}$$

where j_r is the local current density of reaction r, $\nu_{i,r}$ is the stoichiometric coefficient of species i in reaction r, a_y is the

volumetric surface area, n_r is the number of electrons transferred in reaction r, and F is Faraday's constant. The local current densities at the anode and cathode are determined using the anodic and cathodic branches of the Butler–Volmer equation, respectively, in the forms of

$$j_{\text{OER}} = j_{0,\text{OER}} e^{\eta_{\text{OER}} / b_{\text{OER}}}$$
(10)

$$j_{\rm HER} = -j_{0,\rm HER} e^{-\eta_{\rm HER}/b_{\rm HER}} \tag{11}$$

$$j_{\text{COER}} = -\frac{c_{\text{CO}_2}}{c_{\text{ref}}} j_{0,\text{COER}} e^{-\eta_{\text{COER}}/b_{\text{COER}}}$$
(12)

where $j_{0,r}$ are the exchange current densities, b_r are the Tafel slopes, and η_r are the activation overpotentials. The COER equation includes the concentration dependence on CO₂ with respect to a reference concentration c_{rep} but the other two reactions are assumed to have a constant reactant source of water, and as such no explicit concentration dependence is necessary. In this case, c_{ref} should refer to the equilibrium concentration, but in some derivations of kinetics a different reference concentration, such as 1 M_r^{22} is used, leading to a correspondingly altered derived value of exchange current density. The Tafel slopes are given by

$$b_r = \frac{RT}{\alpha_r F} \tag{13}$$

where α_r are the charge transfer coefficients for their respective Tafel branches and *R* is the ideal gas constant. The activation overpotentials η_r are defined as

$$\eta_r = \phi_{\rm s} - \phi_1 - E_{\rm eq,r} \tag{14}$$

where ϕ_s and ϕ_l are the electrode and electrolyte potentials, respectively, and $E_{eq,r}$ is the equilibrium potential of reaction r, corrected for the local pH through the a simplified Nernst equation assuming water and solute activities are near unity,

$$E_{\rm eq,r} = E_{0,r} - \frac{\ln(10)RT}{F} \, \rm pH$$
 (15)

Nesbitt et al. noted that this form of the Nernst equation is determined using the assumption of an acidic or near neutral electrolyte in which the reaction equation reads

$$CO_2 + 2H^+ + 2e^- \rightarrow CO + H_2O \tag{16}$$

and the rate-determining step is the second protonation by H_2O ,²⁴ but it is often assumed without verification that this remains the rate-determining step for the reaction pathway in alkaline media, eq 1. Noting that the pH dependency in eq 15 is roughly -0.0591 pH, they give simplified Nernst equations for acidic/neutral electrolytes and alkaline electrolytes,

$$E_{\rm CO_2/CO} = -0.106 - 0.00591 \text{ pH(V)}$$
 acidic/neutral (17)

$$E_{\rm CO_2/CO} = -0.933 - 0.00591 \text{ pH(V)}$$
 alkaline (18)

and note a common misnomer in literature in which eq 1 is associated with eq 17. Reaction kinetics determined under these assumptions are frequently used in literature²² and while the resulting current density-potential curves may give results similar to experiments, further experimental verification of the reaction mechanism in alkaline media and subsequent recalculation of kinetic parameters are necessary to describe the dependence of the reaction on local activities. The reaction rates from the homogeneous chemical reactions, $R_{H,i}$, are given by

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$$R_{H,i} = \varepsilon \sum_{n} \nu_{i,n} \left(k_n \prod_{\nu_{j,n}>0}^{j} c_j - k_{-n} \prod_{\nu_{j,n}<0}^{j} c_j \right)$$
(19)

where $\nu_{i,n}$ are the stoichiometric coefficients of species *i* in homogeneous reaction *n*, and $k_{\pm n}$ are the forward and backward reaction rates of reaction *n*.

Transport. The gas phase consists of 99.99% pure CO_2 at the inlet (0.01% N_2) with additional CO and H_2 components at the outlet. Transport in the gas phase from the gas channel through the GDL and MPL is modelled with Darcy's law and a mixture averaged diffusion model approximation requiring only binary diffusion coefficients.^{25,26} Darcy's law reads

$$\mathbf{v} = -\frac{\kappa}{\mu_g} \nabla p \tag{20}$$

where **v** is the gas velocity field, κ is the permeability of the porous medium, μ_g is the dynamic viscosity of the gas, and p is the pressure. The flow is assumed incompressible and density variations are neglected. The diffusive mass flux is assumed to be proportional to the mole fraction gradient through

$$\mathbf{J}_i = -\rho_i D_i \frac{\nabla x_i}{x_i} \tag{21}$$

where \mathbf{J}_i is the mass flux of species *i* relative to the average fluid velocity \mathbf{v} , ρ_i is the species density, D_i is the porosity and finite pore size corrected mixture-averaged diffusion coefficient, and x_i is the mole fraction. The mixture-averaged diffusion coefficients are determined from the binary diffusion coefficients in

$$D_i = \frac{1 - x_i}{\sum_{j \neq i} \frac{x_j}{D_{i,j}^{\text{eff}}}}$$
(22)

where D_i^{eff} are the effective binary diffusion coefficients corrected for porosity and tortuosity through the Bruggeman correlation,²⁷

$$D_{i,j}^{\text{eff}} = \varepsilon^{3/2} D_{i,j} \tag{23}$$

Gas transport is usually slightly poorer in the MPL than in the GDL because MPLs typically have lower permeabilities and porosities than GDLs, but this reduction is small compared to the effect that electrolyte intrusion would have on the GDL. Regardless, it is common for manufacturers to supply GDLs with MPLs already applied, and provide the averaged values of transport properties of the final bilayer structure rather than of the individual components. We thus elect to model the GDL and MPL as one numerical domain with uniform porosity and permeability.

In the liquid phase there is bulk flow in the electrolyte channel, but the effect of change in permeability on transition into the porous CL is sufficient that flow in the CL can be neglected.²⁸ Transport of species is modelled by assuming that concentrations remain small enough to be treated as a dilute solution and using the Nernst–Planck equation:

$$\mathbf{N}_{i} = -D_{i}^{\text{eff}} \nabla c_{i} - D_{i}^{\text{eff}} \frac{Fz_{i}}{RT} c_{i} \nabla \phi_{l} + c_{i} \mathbf{u}$$
(24)

where **u** is the liquid velocity and z_i is the species charge number. Species conservation in the steady state is ensured by

$$7 \cdot \mathbf{N}_i = R_{\mathrm{H},i} + R_{\mathrm{ct},i} \tag{25}$$

and electroneutrality is ensured by

$$\sum_{i} z_i c_i = 0 \tag{26}$$

allowing the transport equations in eqs 24 and 25 to be solved for all but one of the species, with the final concentration determined through the electroneutrality condition in eq 26. The local liquid velocity \mathbf{u} is determined assuming a Poiseuille flow profile,

$$\mathbf{u} = 6U \left(\frac{x}{W_{\rm EL}} - \left(\frac{x}{W_{\rm EL}} \right)^2 \right) \hat{\mathbf{z}}$$
(27)

where U is the average velocity. A similar method is used for the inlet gas velocity but with W_{GAS} instead of W_{EL} , and U_g prescribed based on the target single pass conversion. At the liquid-gas phase interface, the flux is described by

$$-\mathbf{n} \cdot \mathbf{J}_i = -k_i (\mathcal{H}_i p_i - c_i) \tag{28}$$

where \mathcal{H}_i is the Henry constant for species *i*, p_i is the partial pressure of that species and k_i is a characteristic cross phase mass transfer coefficient determined for CO₂ adsorption into the CL ionomer by Kas et al.²³ The Henry constant is further corrected for ionic concentrations within the electrolyte (see eq 6 in the SI) following the Sechenov equation.^{29–31} The solubilities of the remaining gas species are around 2 orders of magnitude lower than that of CO₂, so their Henry constants are negligible and they cannot enter the CL, only exit. CO₂ entering the liquid phase causes p_{CO_2} to progressively decrease along the flow channel, subsequently decreasing the flux in eq 28.

COMPUTATIONAL MODEL

The equations for the liquid phase are altered slightly to yield approximate expressions that prove significantly easier to resolve computationally. The homogeneous reactions are problematic due to the extreme magnitudes of the rate coefficients, so we use the assumption that the local environment will be alkaline in the vicinity of the cathodic reaction and acidic in the vicinity of the BPM to selectively neglect the alkaline pathways, K_1 and K_2 , in the BPM half of the electrolyte channel and the acidic pathways, K_3 and K_4 , in the cathodic half of the channel. Due to the coupling of their equilibrium constants we are assured that there is no significant discontinuity in net reaction rates at this halfway point so long as the concentration boundary layers do not extend further than half of the channel width. Furthermore, for numerical stability we adopt a logarithmic form of the transport equations in the liquid,

$$\mathbf{N}_{i} = -e^{C_{i}} D_{i}^{\text{eff}} \nabla C_{i} - D_{i}^{\text{eff}} \frac{Fz_{i}}{RT} e^{C_{i}} \nabla \phi_{l} + e^{C_{i}} \mathbf{u}$$
(29)

where $c_i = e^{C_i}$. C_i can now take any real value and the relation ensures that c_i is strictly positive. This entirely precludes the possibility of negative concentrations in the solution, which is otherwise a common computational issue in systems with high reaction rates and reactant depletion. Another limitation is the computational cost of the effect of local ionic strength variation on ϕ_l in eq 24. Extremely high homogeneous reaction rate coefficients lead to high species concentration sensitivities, even in regions with smooth solutions, leading to stiffness in the system. However, we can do little more to alleviate this without sacrificing significant detail in the homogeneous reactions. An approximation is made to remove the dependence of the migration term on concentration and instead use an iterative nonuniform electrolyte conductivity, given by

$$\sigma_l = \frac{F^2}{RT} \sum_i z_i^2 D_i c_i^* \tag{30}$$

where c_i^* is the average concentration in the previous iteration of the numerical solver. This effectively decouples ϕ_i from eq 29 by determining it from Ohm's law,

$$\mathbf{i} = -\sigma_l \nabla \phi_l \tag{31}$$

and the state of the previous iteration, and allowing eq 29 to solve for C_i without simultaneously solving for ϕ_l .

Despite these simplifications, the scale of the system is still too large for reasonable computation. To this end, the cell model is decomposed into the subcells shown in Figure 3. We

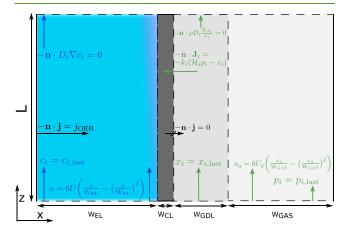


Figure 2. Schematic of the model equations and domains. The *z*-axis follows the flow along the catholyte channel of length *L*. The *x*-axis passes through the catholyte channel of width W_{EL} , the CL of width W_{GL} , the GDL of width W_{GDL} , and into the gas channel of width W_{GAS} . In black, the electrolyte current boundary conditions denote the anodic current source on the left and insulating condition on the right. In the liquid phase, blue equations denote the inlet condition on concentration, the open boundary outlet condition, and the Poiseuille flow velocity distribution in eq 27. In the gas phase, green equations denote the inlet conditions on partial pressures and mole fractions, the open boundary outlet condition, the liquid—gas mass transfer rate in eq 28, and the initial Poiseuille gas velocity distribution in the gas channel, where x_g denotes the coordinate within the gas channel and U_g is the initial average gas velocity for the simulation.

take one initial short (4 cm) basis cell and find a stationary solution, and then solve a second subcell with the downstream outlet of the previous subcell used as the inlet condition. This approximation requires that the flow be sufficiently fast that transport against the flow direction can be neglected. This condition can be written as a necessarily high bulk Péclet number, the ratio of advective to diffusive transport rates. While there is no flow at the channel walls and in the CL, these regions are fully within the concentration boundary layer and the flow-wise concentration gradient is too small for diffusion

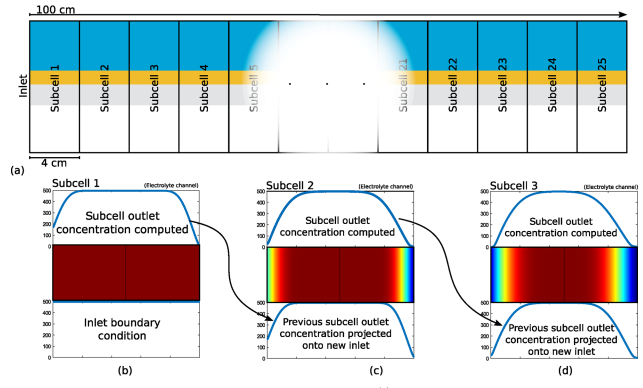


Figure 3. The simulation sequentially solves subcells divided in the flow direction. (a) Decomposition of the 100 cm cell into 25 adjacent 4 cm subcells. The initial and final subcells are coupled to inlet and outlet regions, respectively, and each internal subcell is coupled to the boundary values of the adjacent subcells. (b) The inlet HCO_3^- profile is projected onto the inlet of the first subcell, solved along the length of the subcell, then projected from the outlet onto the inlet of (c), which is subsequently solved and projected onto the inlet of (d) and so on. The other concentration and potential profiles at the top of each subcell catholyte channel and CL are projected onto the inlets of each subsequent subcell. The same method is applied between subsequent GDL and gas flow channel inlets, projecting the mole fractions and pressures in the gas phase. The numerical conditions are shown in Figure 2.

to play a non-negligible role. Instead the flow wise variation in the CL is almost entirely determined by the concentration development in the adjacent electrolyte channel and in the gas phase, in what amounts to a locally 1D model, which is to be expected of a domain whose in plane dimension is 4 orders of magnitude greater than the through plane dimension. In the 4 cm model with no subcell decomposition it was already found that in plane diffusion was negligible in the CL, and it is expected that it would become even less important when the length scale is increased.

Similarly we require that electrolyte potential varies little in the flow-wise direction. The characteristic length of a subcell far exceeds the thickness of the channel, so the flow-wise current density, j_{z} , will be negligible with respect to the flowperpendicular current density j_x . The variation of electrolyte potential in the flow-wise direction depends on the variation of electrode reaction kinetics, and while this can be large due to high pH gradients, it is still small compared to the potential drop across the channel. While the subcells allow us to reduce the size of the computed domain in favour of iteration, the above requirements still motivate us to take as long a subcell as feasible, but the size of 4 cm is chosen out of expediency. While domain decomposition and variable transformations are not uncommon in numerical modelling they are underutilised in the field, with most models of electrochemical CO2 reduction resorting to 1D,^{10,22,32,33} simplifying the homogeneous reactions,³⁴ investigating only low current densities,³⁵ or only modelling small electrolysers.²³Table 1 shows the geometric and operational parameters of the electrolyser,

with unreferenced parameters measured from an in-house experimental setup.

RESULTS

Model Validation. Figure 4 shows experimental comparisons of partial current density against cathode potential and

Table 1. Geometric and Operational Parameters for the Base $Model^a$

parameter	description	value	unit	ref
$p_{\rm abs}$	gas pressure	1	atm	
T	temperature	293.15	K	
$c_{\rm KHCO_3}$	catholyte concentration	500	mM	
Re	Reynolds number	200		
L	flow channel length	4	cm	
$W_{\rm EL}$	flow channel thickness	0.254	mm	
$W_{\rm CL}$	CL thickness	3.5	μ m	22
$W_{\rm GDL}$	GDL thickness	325	μ m	36
$W_{\rm GAS}$	gas channel thickness	3	mm	
Н	flow channel height	5	mm	
$\varepsilon_{\rm CL}$	CL porosity	0.5		
$\varepsilon_{ m GDL}$	GDL porosity	0.53		36
κ	GDL permeability	1.72×10^{-11}	m ²	36

^{*a*}For a description of all parameters, see Table S2. Flow rate and channel length in the validation cases are adjusted accordingly, with standard temperature and pressure. Channel parameters are based on an in-house experimental setup with dimensions similar to that of Wu et al.⁸.

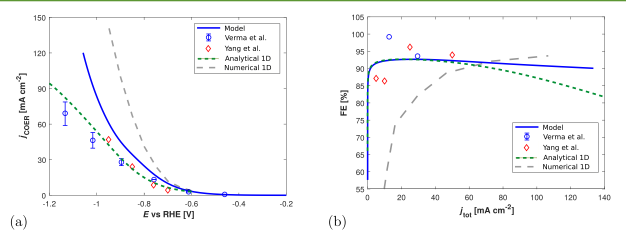


Figure 4. Model validation against experimental results from experimental results from Verma et al.⁴⁰ as well as experimental results from a similar experimental setup from Yang et al.,⁴¹ and a computational model developed by Weng et al.²² and an analytical model developed by Blake et al.³³ based on the electrolyser properties of the Verma et al. experiments. (a) Partial CO₂ current density against cathode potential and (b) the Faradaic efficiency against total current density. The line plot for the computational model is for the experimental case of a 15 mm long flow electrolyser with a 0.5 M unsaturated KHCO₃ liquid electrolyte with a liquid flow rate corresponding to Re = 2.6 and an excess gas feed.

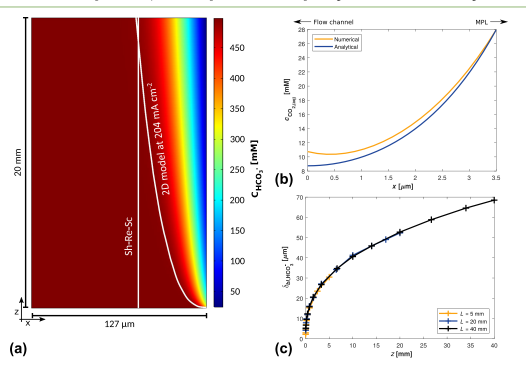


Figure 5. Analytical results for flow-wise development of concentration boundary layer thickness, compared to numerical results. To quantify the numerical boundary layer thickness, HCO $\frac{1}{3}$ is used as the characteristic species to quantify the buffering effect, rather than CO₂. In (a) the boundary layer develops from negligible thickness to a thickness greater than that of the Sherwood–Reynolds–Schmidt correlation (eq S2), which does support its validity as an average approximation, but also highlights how unreasonable it is to take such an average near the inlet. In (b) the CO₂ distribution in the CL from the model follows $\cosh\left(\frac{x}{W_{CL}}\right)$ as is expected of a 1D reaction-diffusion system.^{33,42} More information on the analytical approximations can be found in SI, section 1.2. In (c) the boundary layer thicknesses exhibit a dependence on $z^{1/3}$ as is expected of Poiseuille flow for boundary layers much thinner than the half-channel width.^{43,44}

FE against total current density, for the parameters in Table S2. The match between computational and experimental results is good, only slightly overestimating partial current density. It is expected that the deviation arises from phenomena outside of the scope of the model, such as significant bubble formation near the CL-electrolyte channel interface or liquid breakthrough in the MPL. Both of these phenomena would result in a reduction in cell performance due to reduced access to reaction sites³⁷ and increased

diffusion pathway lengths³⁸ and are both present primarily at higher current densities. As the liquid breakthrough would lead to a significant reduction in FE,³⁹ which is not observed in Figure 4b, the bubble formation explanation is favoured. Despite the short length of the validation case electrolyser, the difference between the presented 2D model and the 1D model of Weng et al. is stark. The use of one operation-independent boundary layer thickness in the Weng model leads to prohibitively low mass transfer between the electrolyte channel

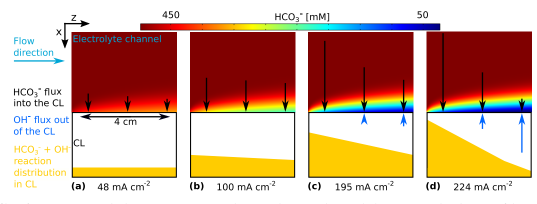


Figure 6. Profiles of HCO_3^- just outside the CL at increasing total current densities, along with the respective distributions of the reaction in eq 5 of HCO_3^- ions with OH^- in the CL. The yellow shape shows the distribution of the homogeneous reaction within the CL. In (a) and (b) relatively little carbonate buffering takes place due to low OH^- generation, and this reaction is quite uniform and almost exclusively takes place within the CL (>99%). For (c) and (d), however, the reaction rate is relatively high, occurs predominantly close to the inlet where CO_2 availability and OH^- generation are highest, and increasingly takes place in the buffer layer outside of the CL, with 10% in (c) and 30% in (d). This increasing reaction outside of the CL is due to the combined transport of buffer into the CL and CO_2 gas supply together being insufficient to buffer the generated OH^- , which is now able to diffuse out into the electrolyte channel before reacting with the bulk buffer solution.

and CL at low current densities where the boundary layer should be thinner, and unrealistically high mass transfer at high current densities where the boundary layer should be thicker. By contrast the analytical model of Blake et al. predicts the development of the concentration boundary layer in the channel and takes an average value for its thickness, but neglects the effect that the buffer of the KHCO₃ catholyte will have on this development. This leads to an overestimation of boundary layer thicknesses and a lower prediction for CO2 current density as a result. It is possible to verify against analytical mathematical approximations of scaling relations that can be extrapolated to larger length scales. Figure 5 shows that the computational results match the predicted trends well, such as the cubic root length scaling of the boundary layer thicknesses and the hyperbolic cosine shape of the CO₂ distribution in the CL. The quality of the agreement with these results is the best validation that the circumstances permit.

UPSCALING

Upscaling Study. To determine how local reaction environments change for longer electrolysers, we compare a 4 cm long electrolyser model with a 100 cm long electrolyser model. For this ratio of lengths, we chose to model the 100 cm electrolyser receiving a $25 \times$ higher CO₂ flow rate in the gas channel due to it being $25 \times$ longer. Despite this, the pressure drop in the gas phase remains small compared to the liquid pressure drop. The liquid flow rate between models is held constant. Depending on the flow geometry, this can lead to large pressure drops in the liquid phase that far exceed the pressure drops in the gas phase, up to the order of hundreds of millibars for thin single channels. Although this could lead to many problems, from mechanical stability to local liquid breakthrough when pressure variations exceed the stability window of the GDL,⁴⁵ these effects are outside of the scope of this work and are not modelled. The gas flow rate was varied to give insight into how different single pass conversions affect cell performance.

Given the focus on higher current densities (>100 mA cm^{-2}), we normalise reactant consumption by gas phase reactant supply only, as the overwhelming majority of CO₂ from the liquid flow channel is expected to be consumed in

buffer reactions before reaching the CL for current densities above 40 mA cm⁻².⁴⁶ If the yield was normalised around the total influent CO₂ from both gas and liquid phase then it would become a relatively low percentage that is dependent on the ratio of channel flow rates. This electrolyte contribution to the reaction is similar to the operational method of a bicarbonate reactor, but such reactors seldom reach high current densities and when they do, they require high electrolyte concentrations.⁴⁷ We furthermore disambiguate between the usage of consumption and conversion following the convention of Larrázabal et al.,48 defining consumption as the percentage of gas phase CO_2 that enters and is consumed in the electrolyser and conversion as the percentage of influent gas phase CO₂ converted specifically into CO. These choices are further motivated by the results of the lab-scale electrolyser model in Figure 6.

CO₂ Conversion. At the low current density in Figure 6a, the carbonate buffer reaction in eq 5 occurs almost entirely within the CL and HCO₃ concentration remains high at the edge of the CL, indicating that the pH within the CL remains well buffered and the aqueous CO2 concentration is high enough for the reaction to be performed in both bicarbonate electrolyser mode and gas supplied mode. However, at higher current densities in Figure 6c,d, HCO_3^- is depleted by the reaction in eq 5 before reaching the CL, and the buffering reaction takes place in a concentration boundary layer outside the CL, showing that the reaction can no longer be performed in bicarbonate electrolyser mode. Despite this, when tracking the total HCO₃⁻ consumed this way we still find that the majority of the reaction takes place within the CL, implying that the buffering effect is actually due to the reaction of gassupplied CO_2 to HCO_3^- through eq 4 and then CO_3^{2-} through eq 5.

The performances at different single pass conversions are shown in Figure 7. At low conversions, the supply of CO_2 is not limiting, and so both cases exhibit high FE, with the small reduction in the 100 cm electrolyser FE arising only from the increase in average boundary layer thickness, following the trend of the 4 cm case in Figure 5c. As conversion increases, the reduction in FE is far more severe in the 100 cm electrolyser than the 4 cm, as the reactant CO_2 is increasingly consumed by the parasitic reaction with OH⁻ in the regions

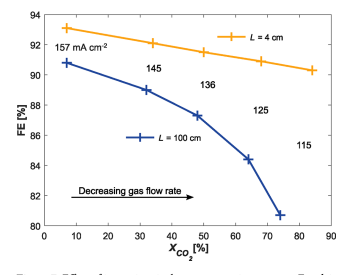


Figure 7. Effect of increasing single pass conversion, χ_{CO_2} on Faradaic efficiency in the 4 and 100 cm electrolysers. The 100 cm electrode is maintained at 3.1 V, leading to a progressive reduction in average j_{tot} as flow rate is reduced to emulate increasing single pass conversion. The 4 cm electrolyser cell potential is adjusted from 2.98 to 2.94 V to match the average j_{tot} at each corresponding gas flow rate. While FE decreases when single pass conversion is increased, the drop is more severe in the 100 cm case.

where the boundary layer is too thick to effectively buffer the OH⁻ produced in the CL. This can be seen directly in Figure 8b, in which 30% of the total consumed CO_2 is converted into HCO_3^- and CO_3^{2-} instead of CO. This high consumption can be easily erroneously interpreted as a high conversion, as in Figure 8a, in which the outlet ratio of CO_2 and CO seems much improved in the 100 cm case over the 4 cm case. It is instead the loss of reactant rather than an increase in product that skews this ratio.

Experimental studies often report a significantly higher consumption of reactant than is found in Figure 8b due to conversion to CO_3^{2-} , as well as crossover of this CO_3^{2-} through the membrane. The reports of exceptionally high crossover mainly come from electrolysers with AEMs,⁴⁹ but BPMs

inhibit the transport of the carbon carrying anions,¹⁴ so we would expect lower crossover in our case. However, our assumption that BPMs entirely prevent this ion transport may not be realistic, as unwanted crossover has still been reported in BPM electrolysers.^{50,51} There is also an argument that purely by the stoichiometry of eqs 1, 4 and 5 and balancing the local production and consumption of CO₂ and OH⁻, the minimum fraction of reactant CO_2 converted to CO_3^{2-} should be 50%.⁴⁸ This argument assumes fast equilibrium reactions though, and Weng et al. showed that the CO₂ utilisation efficiency is dependent on k_1 from eq 4,¹⁰ and we also found that the kinetic treatment is necessary as eq 4 is out of equilibrium. The 50% minimum estimate is most valid for KOH electrolytes due to their high pH and capacity for CO₂. We instead use a CO₂ saturated KHCO₃ electrolyte with a high buffer capacity, and although we do observe a large reaction between CO₂ and OH⁻ a significant portion of this comes from the catholyte supplied CO₂ while we only consider CO₂ sourced from the gas phase in Figure 8b.

The impact of the chemical consumption of CO_2 by the reaction with OH^- depends on the relative costs of the gas channel upstream and downstream processes. If the cost of separation of CO from the product stream is relatively high, then the excess CO_2 consumption actually improves the economics due to the higher CO outlet portion in the 100 cm electrolyser. Assuming that the anolyte and catholyte streams are recombined and recirculated through the cell this chemically reacted CO_2 will be converted back from CO_3^{2-} and HCO_3^{-} when the acidic anolyte stream shifts the pH back to the initial value and the CO_2 can be recovered from the solution. This means that the reaction with OH^- need not be considered as a true loss of reactant, rather as an inhibiting process that reduces j_{CO} and single pass conversion.

Reaction Inhomogeneity. The variation of j_{tot} is also greater in the 100 cm case. Figure 9 shows j_{tot} near the inlet, in the vicinity of which the boundary layers are thin and the CL is very well buffered. At the inlet, j_{tot} is far greater than j at the outlet, where both poorer buffering and reduced reactant supply. This ideal inlet region is limited only by overpotential and as such reaches a higher value of j_{tot} in the 100 cm case due

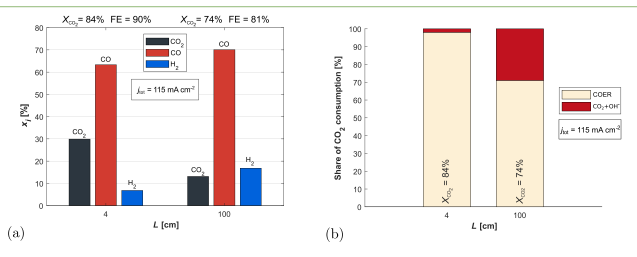


Figure 8. (a) Molar composition of the gas channel outlets in the two cases. Despite lower total CO_2 conversion and higher H_2 mole fraction, the 100 cm case exhibits a higher CO mole fraction in the gas outlet. (b) Relative magnitude of reaction sinks for CO_2 entering the CL from the gas phase. This explains the behaviour in (a), as 30% of the total CO_2 consumed is converted to HCO_3^- through the reaction with OH^- instead of being converted to CO_1 leading to a higher total consumption of CO_2 and a lower resultant CO_2 fraction in the outlet stream. It is thus the loss of reactant rather than improvement in conversion that seemingly improves the outlet composition.

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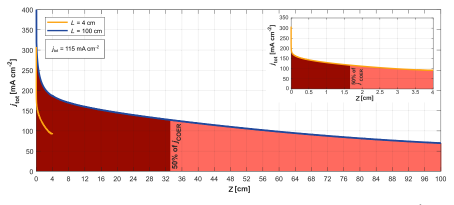


Figure 9. Local values of j_{tot} in the two electrolyser cases for an average total current density of $j_{tot} = 115$ mA cm⁻². The 100 cm reaches a higher value of j_{tot} near the inlet due to the higher cell potential (3.1 V) than the 4 cm case (2.94 V) necessary for the same average j_{tot} but falls to a lower value near the outlet due to the larger pH increase in the CL.

to the higher cell potential necessary to reach an average of 115 mA cm⁻². Despite this increased potential, j_{tot} at the outlet drops below the 4 cm case, because the pH is far higher in the 100 cm case. This inhomogeneity of the reaction distribution leads to the half of the total CO₂ reduction reaction taking place in only the first third of the electrolyser. Additional discussion of the pH gradients in the CL and catholyte channel can be found in SI, Section 2.

The inhomogeneity of the reaction distribution and failure of the buffering effect also causes a wide range of pH values to be present along the CL. The well-buffered inlet will retain a pH equal to that of the influent electrolyte, but the poorly buffered CL near the outlet will have a far higher pH unless the current density is very low or the gas phase CO_2 supply is very high. This increase in pH leads to a larger equilibrium potential correction through eq 15, a greater parasitic reaction rate and a chemical shift towards CO_3^{2-} . The electroneutrality condition requires that the high CO_3^{2-} concentration be paired with a high K⁺ concentration, which together facilitate the precipitation of K_2CO_3 on the solid phase, leading to a reduction in diffusion pathways for CO2 and potential damage to the porous structure as the K_2CO_3 crystals obstruct pores in the CL and MPL.^{39,52} These precipitates are also hydrophilic and can, along with the high pH, alter the potential of zero charge and thus electrowetting properties of the solid-liquid boundary, leading to loss of effective hydrophobicity in the MPL. This allows electrolyte breakthrough into the GDL and gas channel, leading to a negative impact on FE and stability. The penetrating electrolyte increases the diffusion pathway length for CO₂ as it blocks pores, leading to a reduction in j_{CO} while allowing HER to continue unmitigated. This pervasive issue is one of the main causes of performance loss over time, and mitigation strategies currently focus on temporal variation in reactor operation or low electrolyte strengths.^{53,54}

Discussion. By modelling the flow parallel direction as well as the through plane direction, we overcome the limitations of 1D models that require empirical or simplified analytical expressions to attempt to capture flow-wise variation. The work of Yang et al.⁵⁵ includes the flow channel in 2D but neglects variation in pH in the electrolyte. This is at odds with this work and the work of Kas et al.,²³ which both predict large variations in local pH. While this could be roughly justified by the use of high pH KOH electrolyte, it would still have a large impact on the local homogeneous reaction rates and the total loss of reactant to carbonate formation. This is not reported on, despite carbonate crossover being a serious issue,^{14,49–51}

and despite the homogeneous reaction kinetics they report being significantly faster than the heterogeneous kinetics for such a high pH. This assumption of pH homogeneity also hampers the upscaling analysis, as homogeneous properties are indifferent to scale, giving overambitious positive predictions for scalability. By contrast, the experimental scalability analysis of Jeanty et al.⁵⁶ highlights the more fundamental issues of electrolyte breakthrough and salt precipitation. Both of these support our conclusion of the importance of quantifying local pH and its effect on the carbonate equilibrium reactions as the perspired electrolyte has a pH value of 10 compared to that of the neutral K₂SO₄ catholyte and the precipitate comprised of carbonate salts for which the carbon could only have been sourced from the CO₂ feed. They also find similar trends for FE, in which the upscaled electrolyser suffered a larger reduction in FE when lowering feed gas CO₂ supply. The most interesting result is the drastic improvement when implementing the circulating pump in the scaled up cell which, despite lowering the average CO_2 concentration, leads to a much more uniform concentration profile and a break up of the boundary layer. We identify an analogous issue in the catholyte channel with nonuniform pH and thick boundary layer development as a primary concern in scalability, and the experimental success in reducing the effect in the gas channel is reassuring.

Stability. The extension of the 4 cm model to the 100 cm model shows deterioration in conversion, FE, and necessary cell potential, and the model indicates that these issues will only be exacerbated with further elongation of the flow channel. While these performance losses could be in some cases surmountable, the extreme variation in local reaction environment that comes with them would lead to further issues outside of the scope of the model. While liquid breakthrough in generalised porous media is well understood, electrolyte breakthrough in the electrolytic cell is related to precipitation and electrowetting. Precipitation would require stochastic modelling and electrowetting effects are poorly understood at interfaces where charge transfer reactions take place, so both are out of the scope of the model. However, we can note that both are dependent on the local ionic concentrations and pH, the latter due to the shift in potential of zero charge, and comment that by determining the general trends in pH and ionic strength we can predict that the upscaled model will be more susceptible to breakthrough closer to the outlet. In the future, when better descriptions of these phenomena are available, it will be possible to bridge the gap between local

environment scalability and pressure stability studies^{45,57} to create a full description of breakthrough.

If the Reynolds number of the electrolyte flow channel is held constant, then the pressure drop along the channel will increase linearly with electrolyser length. The pressure variation in the liquid phase can lead to electrolyte or gas breakthrough when out of balance with the gas pressure drop, so it is ideal to scale one flow rate to ensure the differential pressures remain within MPL stability ranges. If particularly thin flow channels, such as those used in lab scale electrolysers, are required to minimise ohmic losses then the pressure drop can become excessively large. Without sufficiently pressure resistant MPLs this could lead to liquid breakthrough. Both the hydrophobicity and the mechanical stability of CLs are dependent on ionomer type and loading, which vary between experimental studies and are the subject of independent optimization. Furthermore, it becomes an issue to mechanically support a flow channel under such conditions, potentially necessitating additional structural supports in the flow channel, which could cause the Poiseuille flow to transition into plug flow. Plug flow boundary layers scale with $z^{1/2}$, rather than the $z^{1/3}$ of Poiseuille flow,⁵⁸ and as such can grow more rapidly. Specifically, we can take the ratio of the Lévêque approximation for plug flow and Poiseuille flow concentration boundary layer thicknesses,

$$\delta_{\text{plug}} = 3.643 \sqrt{\frac{Dz}{U}}$$
 and $\delta_{\text{Poiseuille}} = 1.607 \sqrt[3]{\frac{W_{\text{EL}}Dz}{U}}$
(32)

where the concentration at the edge of the boundary layer is 99% of the bulk concentration. Plug flow will lead to thicker boundary layers when

$$z > \left(\frac{1.607}{3.643}\right)^6 \frac{UW_{\rm EL}^2}{D}$$
(33)

which in our case for the diffusivity of HCO_3^- is roughly after only 11 cm. Widening the channel leads to increasing ohmic drop across the electrolyte, so a balance must be struck between electrolyte flow rate, pressure drop, and ohmic drop. For small lab scale electrolysers this balance usually favours thin channels due to the short flow channels exhibiting small pressure drops and little variation in the local pH and boundary layer, but as electrolyser length increases the benefit of lower ohmic drop will be outweighed by the cost of reduced GDL stability and increase in boundary layer thickness. This trade-off between flow velocity, flow channel length, and ohmic drop is an optimisation problem that will remain relevant even in a full-scale stack implementing short channels.

Performance Loss Minimisation. With the sources of the performance losses isolated, it is possible to propose potential solutions. The primary issue is the lack of effective buffering due to thick concentration boundary layers, and so reducing the boundary layer thickness is paramount. This could be done by introducing static mixers or altering the geometry in the electrolyte channel to induce chaotic mixing⁵⁹ and break up the developing boundary layer, and although this could worsen carbonate and product crossover,^{60,61} product crossover is only likely to be an issue when producing primarily liquid products.^{62,63} A similar issue would arise from the use of a stronger base such as KOH, in which the high pH would lead to unavoidable loss of CO₂ feed reactant to carbonate. When averaging across the entire CL for 115 mA cm⁻², we find a

dissolved CO₂ concentration of ~5 mol m⁻³, so as a rough estimate, we can replace our predicted pH of ~12.5 with 14 for KOH to find from eqs 9 and 19 that the percentage of total reacted CO₂ reacting with OH⁻ rises from ~30%, in Figure 8b, to ~80%. This chemical reaction with OH⁻ would be significantly greater than the electrochemical production of OH⁻, leading to a decreased pH boundary layer forming in the flow, in which the gas CO₂ feed is used primarily to buffer the KOH. This boundary layer would be comprised of carbonate and bicarbonate and suffer from lower conductivity, and in a recirculated electrolyte would eventually react with enough feed CO₂ to effectively become KHCO₃. In this case the share of CO₂ converted to CO would actually be improved by increasing overpotential at the cost of FE.

Circumvention of scaling issues can be preferable to prevention, so alternative parallelised geometries can be employed to retain short channels and high performance. Splitting one long channel into multiple shorter channels would require each shorter channel to maintain a comparable flow rate to and thus pressure drop to achieve greater performance, so it would be preferable to use entirely distinct shorter channels with separate inlets and outlets. While technoeconomic assessments often consider MEAs without catholyte buffers, Badgett et al. showed that the inclusion of a catholyte buffer layer improves performance⁶⁴ and only requires an increase in the capital cost of the system by around 1%.65 The supply of catholyte at sufficient flow rates to multiple channels within a cell or stack could necessitate additional equipment and, from a holistic perspective, could complicate full-scale stack designs.

Alternatively, the problem can be addressed at the source: the high production rate of OH⁻ ions. While the production of OH^- due to the reduction reaction (eq 1) is necessary and unavoidable, the low FE shows that the HER (eq 2) rate is unnecessarily high, especially in the regions with low j_{CO} . The low j_{CO} is due to reactant limitation, not lack of available reaction sites, so by reducing the catalyst loading further along the electrolyser the zeroth order HER would be reduced linearly but the COER would be impacted less as it would remain reactant limited. This would cause an overall reduction in single pass conversion, but an improvement in FE and reduction in parasitic consumption as the lowered OHproduction from HER would lower the pH in the CL. Variable catalyst loading the flow direction will be discussed in more depth in a forthcoming paper, and a brief demonstration of the efficacy of the proposed solution can be found in SI, Section 3.

CONCLUSION

We have developed a large-scale 2D computational model of a CO_2 reduction flow cell, and showed through comparison of a 4 cm lab scale model and a 100 cm upscaled model that the performance metrics of FE, conversion, and voltage efficiency decrease with electrolyser length. We found that poor electrolyte buffering due to increasing boundary layer thickness in the flow channel leads to reaction inhomogeneity in the flow direction and poor utilisation of the outlet regions of the electrolyser, resulting in a system that struggles to match performance benchmarks set in lab scale electrolysers. While some mitigation strategies have been suggested, such as electrolyte mixing, catalyst loading variation, and flow channel optimisation, it is recommended that future experimental studies test these strategies and acknowledge that not only will the necessary current densities for industrial realisation be

greater than those commonly assumed in lab scale studies, but the local reaction environments will be far less favourable than those present in typical lab scale electrolysers.

ASSOCIATED CONTENT

1 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssuschemeng.2c06129.

Description of analytical methods, additional parameters, and additional plots of local environment and variable catalyst loading (PDF)

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Notes

The authors declare no competing financial interest.

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REFERENCES

(1) Hara, K.; Kudo, A.; Sakata, T.; Watanabe, M. High Efficiency Electrochemical Reduction of Carbon Dioxide under High Pressure on a Gas Diffusion Electrode Containing Pt Catalysts. *J. Electrochem. Soc.* **1995**, *142*, L57–L59.

(2) Jouny, M.; Luc, W.; Jiao, F. General Techno-Economic Analysis of CO₂ Electrolysis Systems. *Ind. Eng. Chem. Res.* **2018**, *57*, 2165–2177.

(3) Verma, S.; Kim, B.; Jhong, H.-R.; Ma, S.; Kenis, P. J. A. A Gross-Margin Model for Defining Technoeconomic Benchmarks in the Electroreduction of CO₂. *ChemSusChem* **2016**, *9*, 1972–1979.

(4) Spurgeon, J. M.; Kumar, B. A comparative technoeconomic analysis of pathways for commercial electrochemical CO₂ reduction to liquid products. *Energy Environ. Sci.* **2018**, *11*, 1536–1551.

(5) Bahnamiri, F. K.; Khalili, M.; Pakzad, P.; Mehrpooya, M. Techno-economic assessment of a novel power-to-liquid system for synthesis of formic acid and ammonia, based on CO_2 electroreduction and alkaline water electrolysis cells. *Renewable Energy* **2022**, *187*, 1224–1240.

(6) Tichler, R.; Lehner, M.; Steinmüller, H.; Koppe, M.Power-to-Gas: Technology and Business Models; Springer, 2014.

(7) Whipple, D. T.; Finke, E. C.; Kenis, P. J. A. Microfluidic Reactor for the Electrochemical Reduction of Carbon Dioxide: The Effect of pH. *Electrochem. Solid-State Lett.* **2010**, *13*, B109.

(8) Wu, K.; Birgersson, E.; Kim, B.; Kenis, P. J. A.; Karimi, I. A. Modeling and Experimental Validation of Electrochemical Reduction of CO_2 to CO in a Microfluidic Cell. *J. Electrochem. Soc.* **2015**, *162*, F23–F32.

(9) Gabardo, C. M.; O'Brien, C. P.; Edwards, J. P.; McCallum, C.; Xu, Y.; Dinh, C.-T.; Li, J.; Sargent, E. H.; Sinton, D. Continuous Carbon Dioxide Electroreduction to Concentrated Multi-carbon Products Using a Membrane Electrode Assembly. *Joule* **2019**, *3*, 2777–2791.

(10) Weng, L.-C.; Bell, A. T.; Weber, A. Z. Towards membraneelectrode assembly systems for CO_2 reduction: a modeling study. *Energy Environ. Sci.* **2019**, *12*, 1950–1968.

(11) Endrődi, B.; Kecsenovity, E.; Samu, A.; Darvas, F.; Jones, R. V.; Török, V.; Danyi, A.; Janáky, C. Multilayer Electrolyzer Stack Converts Carbon Dioxide to Gas Products at High Pressure with High Efficiency. *ACS Energy Letters* **2019**, *4*, 1770–1777.

(12) Endrődi, B.; Kecsenovity, E.; Samu, A.; Halmágyi, T.; Rojas-Carbonell, S.; Wang, L.; Yan, Y.; Janáky, C. High carbonate ion conductance of a robust PiperION membrane allows industrial current density and conversion in a zero-gap carbon dioxide electrolyzer cell. *Energy Environ. Sci.* **2020**, *13*, 4098–4105.

(13) Kutz, R. B.; Chen, Q.; Yang, H.; Sajjad, S. D.; Liu, Z.; Masel, I. R. Sustainion Imidazolium-Functionalized Polymers for Carbon Dioxide Electrolysis. *Energy Technology* **2017**, *5*, 929–936.

(14) Lees, E. W.; Mowbray, B. A. W.; Parlane, F. G. L.; Berlinguette, C. P. Gas diffusion electrodes and membranes for CO2 reduction electrolysers. *Nature Reviews Materials* **2022**, *7*, 55–64.

(15) Tufa, R. A.; Chanda, D.; Ma, M.; Aili, D.; Demissie, T. B.; Vaes, J.; Li, Q.; Liu, S.; Pant, D. Towards highly efficient electrochemical CO2 reduction: Cell designs, membranes and electrocatalysts. *Applied Energy* **2020**, *277*, 115557.

(16) Salvatore, D. A.; Gabardo, C. M.; Reyes, A.; O'Brien, C. P.; Holdcroft, S.; Pintauro, P.; Bahar, B.; Hickner, M.; Bae, C.; Sinton, D.; Sargent, E. H.; Berlinguette, C. P. Designing anion exchange membranes for CO2 electrolysers. *Nature Energy* **2021**, *6*, 339–348.

(17) De Mot, B.; Hereijgers, J.; Daems, N.; Breugelmans, T. Insight in the behavior of bipolar membrane equipped carbon dioxide electrolyzers at low electrolyte flowrates. *Chemical Engineering Journal* **2022**, 428, 131170.

(18) Ma, M.; Kim, S.; Chorkendorff, I.; Seger, B. Role of ionselective membranes in the carbon balance for CO_2 electroreduction via gas diffusion electrode reactor designs. *Chem. Sci.* **2020**, *11*, 8854– 8861.

(19) Xing, Z.; Hu, L.; Ripatti, D. S.; Hu, X.; Feng, X. Enhancing carbon dioxide gas-diffusion electrolysis by creating a hydrophobic catalyst microenvironment. *Nat. Commun.* **2021**, *12*, 136.

(20) Hansen, K. U.; Jiao, F. Hydrophobicity of CO_2 gas diffusion electrodes. *Joule* **2021**, *5*, 754–757.

(21) Goyal, A.; Marcandalli, G.; Mints, V. A.; Koper, M. T. M. Competition between CO_2 Reduction and Hydrogen Evolution on a Gold Electrode under Well-Defined Mass Transport Conditions. *J. Am. Chem. Soc.* **2020**, 142, 4154–4161.

(22) Weng, L.-C.; Bell, A. T.; Weber, A. Z. Modeling gas-diffusion electrodes for CO_2 reduction. *Phys. Chem. Chem. Phys.* **2018**, 20, 16973–16984.

(23) Kas, R.; Star, A. G.; Yang, K.; Van Cleve, T.; Neyerlin, K. C.; Smith, W. A. Along the Channel Gradients Impact on the Spatioactivity of Gas Diffusion Electrodes at High Conversions during CO_2 Electroreduction. ACS Sustainable Chem. Eng. 2021, 9, 1286–1296.

(24) Nesbitt, N. T.; Smith, W. A. Water and Solute Activities Regulate CO_2 Reduction in Gas-Diffusion Electrodes. *J. Phys. Chem. C* **2021**, *125*, 13085–13095.

(25) Kee, R.; Coltrin, M.; Glarborg, P.Chemically Reacting Flow, Theory and Practice; Wiley, 2003.

(26) Millington, R. J.; Quirk, J. P. Permeability of porous solids. *Trans. Faraday Soc.* **1961**, *57*, 1200–1207.

(27) Tjaden, B.; Cooper, S. J.; Brett, D. J.; Kramer, D.; Shearing, P. R. On the origin and application of the Bruggeman correlation for analysing transport phenomena in electrochemical systems. *Current Opinion in Chemical Engineering* **2016**, *12*, 44–51.

(28) Beavers, G. S.; Joseph, D. D. Boundary conditions at a naturally permeable wall. *J. Fluid Mech.* **1967**, *30*, 197–207.

(29) Sechenov, M. On the behavior of salt solutions based on their relation to carbon dioxide. *Z. Phys. Chem.* **1889**, 117–125.

(30) Sander, R. Compilation of Henry's law constants (version 4.0) for water as solvent. *Atmospheric Chemistry and Physics* **2015**, *15*, 4399–4981.

(31) Weisenberger, S.; Schumpe, A. Estimation of gas solubilities in salt solutions at temperatures from 273 to 363 K. *Aiche Journal* **1996**, 42, 298–300.

(32) Hashiba, H.; Weng, L.-C.; Chen, Y.; Sato, H. K.; Yotsuhashi, S.; Xiang, C.; Weber, A. Z. Effects of Electrolyte Buffer Capacity on Surface Reactant Species and the Reaction Rate of CO_2 in Electrochemical CO_2 Reduction. *J. Phys. Chem. C* **2018**, *122*, 3719–3726.

(33) Blake, J.; Padding, J.; Haverkort, J. Analytical modelling of CO_2 reduction in gas-diffusion electrode catalyst layers. *Electrochim. Acta* **2021**, 393, 138987.

(34) Kotb, Y.; Fateen, S.-E. K.; Albo, J.; Ismail, I. Modeling of a Microfluidic Electrochemical Cell for the Electro-Reduction of CO_2 to CH₃OH. *J. Electrochem. Soc.* **2017**, *164*, E391–E400.

(35) El-Shafie, O. A.; El-Maghraby, R. M.; Albo, J.; Fateen, S.-E. K.; Abdelghany, A. Modeling and Numerical Investigation of the Performance of Gas Diffusion Electrodes for the Electrochemical Reduction of Carbon Dioxide to Methanol. *Ind. Eng. Chem. Res.* **2020**, *59*, 20929–20942.

(36) El-kharouf, A.; Mason, T. J.; Brett, D. J.; Pollet, B. G. Ex-situ characterisation of gas diffusion layers for proton exchange membrane fuel cells. *J. Power Sources* **2012**, *218*, 393–404.

(37) Lee, C.; Zhao, B.; Lee, J. K.; Fahy, K. F.; Krause, K.; Bazylak, A. Bubble Formation in the Electrolyte Triggers Voltage Instability in CO_2 Electrolyzers. *iScience* **2020**, *23*, 101094.

(38) Kong, Y.; Hu, H.; Liu, M.; Hou, Y.; Kolivoska, V.; Vesztergom, S.; Broekmann, P.Visualization and quantification of flooding phenomena in gas diffusion electrodes (GDEs) used for electrochemical CO_2 reduction: A combined EDX/ICP–MS approach. J. *Catalysis;* Cambridge: Cambridge Open Engage; **2022**; This content is a preprint and has not been peer-reviewed. 40818.

(39) Leonard, M.; Clarke, L.; Forner-Cuenca, A.; Brown, S.; Brushett, F. Investigating Electrode Flooding in a Flowing Electrolyte, Gas-Fed Carbon Dioxide Electrolyzer. *ChemSusChem* **2020**, *13*, 400– 411.

(40) Verma, S.; Lu, X.; Ma, S.; Masel, R. I.; Kenis, P. J. A. The effect of electrolyte composition on the electroreduction of CO_2 to CO on Ag based gas diffusion electrodes. *Phys. Chem. Chem. Phys.* **2016**, *18*, 7075–7084.

(41) Yang, K.; Kas, R.; Smith, W. A.; Burdyny, T. Role of the Carbon-Based Gas Diffusion Layer on Flooding in a Gas Diffusion Electrode Cell for Electrochemical CO_2 Reduction. ACS Energy Letters 2021, 6, 33–40.

(42) Thiele, E. W. Relation between Catalytic Activity and Size of Particle. *Industrial & Engineering Chemistry* **1939**, *31*, 916–920.

(43) Belhocine, A.; Wan Omar, W. Z. Similarity solution and Runge Kutta method to a thermal boundary layer model at the entrance region of a circular tube. *World Journal of Engineering* **2018**, *15*, 468–477.

(44) Holzbecher, E.Weierstrass, Numerical Solutions for the Lévêque Problem of Boundary Layer Mass or Heat Flux. *COMSOL Conference 2008 Hannover*; COMSOL, 2008.

(45) Baumgartner, L. M.; Koopman, C. I.; Forner-Cuenca, A.; Vermaas, D. A. Narrow Pressure Stability Window of Gas Diffusion Electrodes Limits the Scale-Up of CO_2 Electrolyzers. *ACS Sustainable Chem. Eng.* **2022**, *10*, 4683–4693.

(46) Kas, R.; Yang, K.; Yewale, G. P.; Crow, A.; Burdyny, T.; Smith, W. A. Modeling the Local Environment within Porous Electrode during Electrochemical Reduction of Bicarbonate. *Ind. Eng. Chem. Res.* **2022**, *61*, 10461–10473.

(47) Lees, E. W.; Goldman, M.; Fink, A. G.; Dvorak, D. J.; Salvatore, D. A.; Zhang, Z.; Loo, N. W. X.; Berlinguette, C. P. Electrodes Designed for Converting Bicarbonate into CO. *ACS Energy Letters* **2020**, *5*, 2165–2173.

(48) Larrazábal, G. O.; Ma, M.; Seger, B. A Comprehensive Approach to Investigate CO_2 Reduction Electrocatalysts at High Current Densities. *Accounts of Materials Research* **2021**, *2*, 220–229. (49) Ma, M.; Clark, E. L.; Therkildsen, K. T.; Dalsgaard, S.; Chorkendorff, I.; Seger, B. Insights into the carbon balance for CO_2 electroreduction on Cu using gas diffusion electrode reactor designs. *Energy Environ. Sci.* **2020**, *13*, 977–985.

(50) Blommaert, M. A.; Sharifian, R.; Shah, N. U.; Nesbitt, N. T.; Smith, W. A.; Vermaas, D. A. Orientation of a bipolar membrane determines the dominant ion and carbonic species transport in membrane electrode assemblies for CO_2 reduction. *J. Mater. Chem. A* **2021**, *9*, 11179–11186.

(51) Blommaert, M. A.; Verdonk, J. A. H.; Blommaert, H. C.; Smith, W. A.; Vermaas, D. A. Reduced Ion Crossover in Bipolar Membrane Electrolysis via Increased Current Density, Molecular Size, and Valence. *ACS Applied Energy Materials* **2020**, *3*, 5804–5812.

(52) Cofell, E. R.; Nwabara, U. O.; Bhargava, S. S.; Henckel, D. E.; Kenis, P. J. A. Investigation of Electrolyte-Dependent Carbonate Formation on Gas Diffusion Electrodes for CO_2 Electrolysis. ACS Appl. Mater. Interfaces **2021**, 13, 15132–15142.

(53) Xu, Y.; Edwards, J. P.; Liu, S.; Miao, R. K.; Huang, J. E.; Gabardo, C. M.; O'Brien, C. P.; Li, J.; Sargent, E. H.; Sinton, D. Self-Cleaning CO_2 Reduction Systems: Unsteady Electrochemical Forcing Enables Stability. *ACS Energy Letters* **2021**, *6*, 809–815.

(54) Endrődi, B.; Samu, A.; Kecsenovity, E.; Halmágyi, T.; Sebők, D.; Janáky, C. Operando cathode activation with alkali metal cations for high current density operation of water-fed zero-gap carbon dioxide electrolysers. *Nature Energy* **2021**, *6*, 439–448.

(55) Yang, Z.; Li, D.; Xing, L.; Xiang, H.; Xuan, J.; Cheng, S.; Yu, E. H.; Yang, A. Modeling and Upscaling Analysis of Gas Diffusion Electrode-Based Electrochemical Carbon Dioxide Reduction Systems. *ACS Sustainable Chem. Eng.* **2021**, *9*, 351–361.

(56) Jeanty, P.; Scherer, C.; Magori, E.; Wiesner-Fleischer, K.; Hinrichsen, O.; Fleischer, M. Upscaling and continuous operation of electrochemical CO₂ to CO conversion in aqueous solutions on silver gas diffusion electrodes. *Journal of CO*₂ Utilization **2018**, *24*, 454–462.

(57) Baumgartner, L. M.; Koopman, C. I.; Forner-Cuenca, A.; Vermaas, D. A. When Flooding Is Not Catastrophic-Woven Gas Diffusion Electrodes Enable Stable CO_2 Electrolysis. ACS Applied Energy Materials 2022, 5, 15125–15135.

(58) Lévêque, A.Les lois de la transmission de chaleur par convection. *Ph.D. thesis*, Faculté des Sciences de Paris, 1928.

(59) Lee, C.-Y.; Chang, C.-L.; Wang, Y.-N.; Fu, L.-M. Microfluidic Mixing: A Review. *International Journal of Molecular Sciences* **2011**, *12*, 3263–3287.

(60) Erbach, S.; Epple, S.; Heinen, M.; Toth, G.; Klages, M.; Gaudreau, D.; Ages, M.; Putz, A. CO_2 Enrichment in Anode Loop and Correlation with CO Poisoning of Low Pt Anodes in PEM Fuel Cells. *Fuel Cells* **2018**, *18*, 613–618.

(61) Zhang, J.; Luo, W.; Züttel, A. Crossover of liquid products from electrochemical CO_2 reduction through gas diffusion electrode and anion exchange membrane. *J. Catal.* **2020**, *385*, 140–145.

(62) Wang, N.; Miao, R. K.; Lee, G.; Vomiero, A.; Sinton, D.; Ip, A. H.; Liang, H.; Sargent, E. H. Suppressing the liquid product crossover in electrochemical CO₂ reduction. *SmartMat* **2021**, *2*, 12–16.

(63) Li, Y. C.; Yan, Z.; Hitt, J.; Wycisk, R.; Pintauro, P. N.; Mallouk, T. E. Bipolar Membranes Inhibit Product Crossover in CO_2 Electrolysis Cells. Advanced Sustainable Systems **2018**, *2*, 1700187.

(64) Chen, Y.; Vise, A.; Klein, W. E.; Cetinbas, F. C.; Myers, D. J.; Smith, W. A.; Deutsch, T. G.; Neyerlin, K. C. A Robust, Scalable Platform for the Electrochemical Conversion of CO_2 to Formate: Identifying Pathways to Higher Energy Efficiencies. ACS Energy Letters **2020**, *5*, 1825–1833.

(65) Badgett, A.; Ruth, M.; Crow, A.; Grim, G.; Chen, Y.; Hu, L.; Tao, L.; Smith, W.; Neyerlin, K.; Cortright, R. An economic analysis of the role of materials, system engineering, and performance in electrochemical carbon dioxide conversion to formate. *Journal of Cleaner Production* **2022**, *351*, 131564.

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