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Techno-economic Assessment of CO$_2$ Electrolysis: How Interdependencies between Model Variables Propagate Across Different Modeling Scales

Isabell Bagemihl,*¶ Lucas Cammann,¶ Mar Pérez-Fortes, Volkert van Steijn, and J. Ruud van Ommen

ABSTRACT: The production of base chemicals by electrochemical conversion of captured CO$_2$ has the potential to close the carbon cycle, thereby contributing to a future energy transition. With the feasibility of low-temperature electrochemical CO$_2$ conversion demonstrated at lab scale, research is shifting toward optimizing electrolyser design and operation for industrial applications, with target values based on techno-economic analysis. However, current techno-economic analyses often neglect experimentally reported interdependencies of key performance variables such as the current density, the faradaic efficiency, and the conversion. Aiming to understand the impact of these interdependencies on the economic outlook, we develop a model capturing mass transfer effects over the channel length for an alkaline, membrane electrolyser. Coupling the channel scale with the higher level process scale and embedding this multiscale model in an economic framework allows us to analyze the economic trade-off between the performance variables. Our analysis shows that the derived target values for the performance variables strongly depend on the interdependencies described in the channel scale model. Our analysis also suggests that economically optimal current densities can be as low as half of the previously reported benchmarks. More generally, our work highlights the need to move toward multiscale models, especially in the field of CO$_2$ electrolysis, to effectively elucidate current bottlenecks in the quest toward economically compelling system designs.

KEYWORDS: CO$_2$ electrolysis, First-principle electrolyser model, Multiscale modeling, Optimization, Techno-economic analysis

INTRODUCTION

The current anthropogenic carbon economy does not possess the ability to reduce CO$_2$. Instead, it solely oxidizes various fossil-based carbon sources to CO$_2$, leading to increasing atmospheric concentrations. Closing the carbon cycle by converting waste CO$_2$ to bulk chemicals is a promising avenue to minimize emissions and fossil-based resource consumption. One technology offering the potential of achieving this transition is the electrochemical conversion of CO$_2$. Techno-economic studies have led researchers to identify target values for performance variables and pathways toward the profitable deployment of this emerging technology. The first techno-economic analyses studied the economic feasibility of electrochemical CO$_2$ reduction by presenting target values for the performance variables to reach a break-even point. These performance variables include the current density, the faradaic efficiency, and the cell potential, while the conversion rate is fixed. Importantly, these variables are usually assumed to be independent. Under this assumption, the threshold values for the first three variables were derived utilizing a generalized electrochemical CO$_2$ reduction plant model based on a fixed conversion rate and price indication for existing electrolyser technologies. The derived thresholds include current densities above 250–300 mA cm$^{-2}$ and cell potentials below 1.8 V to reduce capital and operational costs of the electrolyser unit, respectively. Further, faradaic efficiencies above 80–90% reduce downstream separation costs of the product. Although these thresholds provide significant guidance for experimental studies, their underlying analyses neglect the interdependencies of current density, faradaic efficiency, cell potential, and conversion on the mechanistic level. This confines the techno-economic analyses to univariate sensitivity analyses, potentially leading to overestimation of the solution space for feasible performance values and electrolyser designs. Understanding the interdependencies of the performance variables is thus crucial for...
electrolyser design, operation, and optimization. Therefore, experimental studies on electrolyser design have mostly been accompanied by modeling efforts, to capture the interdependencies at the channel scale. These models are used to resolve local effects in the electrolyser, for example, to understand concentration gradients and mass transfer limitations due to the change in pH near the catalyst layer. While these models can provide relevant insights into the interdependencies of the performance variables, they so far have not been translated into techno-economic analyses.

Channel models can additionally account for concentration gradients along the flow channel, taking into account their effect on single-pass conversion. For example, the study of Kas et al. showed an increased loss of CO$_2$ to carbonate formation at high current densities due to the limited buffer capacities of the electrolyte. This insight reveals a trade-off between current density and conversion, one of the interdependencies commonly neglected when using fixed performance variables for techno-economic analyses. This study presents a multiscale modeling approach ranging from the mechanistic channel scale over the electrolyser stack scale to the process scale (Figure 1a), assessing interdependencies on the electrolyser design level from an economic perspective.

For the multiscale model, a channel model accurate enough to capture interdependencies between the performance metrics of the electrolyser is developed, which then allows evaluation and demonstration of the influence of the interdependencies on the selected process economic indicator. The channel model is based on a first-principle model of an alkaline flow-through CO$_2$ electrolyser for the production of ethylene, capturing the

Figure 1. Overview of the connection between the process, electrolyser, and channel scale model (a) with the relevant in- and output variables and parameters to couple the scales (b). The no channel model (M1) uses fixed performance variables, while the simplistic channel model (M2) and the full channel model (M3) capture the different pathways of CO$_2$ through the electrolyser (c) with increasing level of detail.
interplay between CO₂ conversion, faradaic efficiency, and cell voltage for varying current densities. This interplay, in turn, influences the electrolyser and downstream unit investment and operating costs. Employing this multiscale framework for techno-economic assessment and optimization allows for computing the desired target performance variables based on mechanistic insights.

## MULTISCALE MODEL

The multiscale modeling framework comprises three scales: the process, the electrolyser, and the channel scale, as shown in Figure 1a. While the multiscale model is generic and independent of the desired product, we will show the model setup and results for ethylene as the main product with a target production rate of 10,000 kg d⁻¹. The choice of this gaseous throughput is motivated by the objective to investigate industrially relevant conditions while ensuring that the financial correlations used for the cost estimate remain applicable. The throughput is used to calculate investment and operating costs for the electrolyser and gas separation, herein considered to be a pressure-swing adsorption unit (PSA). The design of the electrolyser is based on a flow-through gas diffusion electrode (GDE) cell, motivated by its extensive application in experimental studies aiming for high current densities.20,28,29 The electrolyser is operated under ambient pressure and room temperature (see Table 1). The liquid catholyte flow rate is fixed to evaluate the electrolyser performance, while the liquid postprocessing is not considered in the cost estimation. It is assumed in the model that the electrolyser is continuously fed with fresh electrolyte and all formed ionic species, for example bicarbonate, leave the reactor with the liquid electrolyte stream. The gas phase is solely composed of the reactant CO₂, the target product C₂H₄, and the side product H₂, as shown in the flowchart in Figure 1a.

The electrolyser performance is described based on the following five performance variables (Figure 1b): the current density \( i_{\text{tot}} \), the cell voltage \( V_{\text{cell}} \), the faradaic efficiency \( FE_{\text{C}_2\text{H}_4} \), the heterogeneous conversion \( \chi_{\text{het}} \) and the homogeneous consumption \( \chi_{\text{hom}} \). The energy efficiency is not considered separately in this work as it directly follows from the cell voltage and faradaic efficiency.15 While the first three are common terms in electrochemistry, the last two are understood as the conversion rates of (i) CO₂ due to the heterogeneous electrochemical reaction at the catalyst forming the reaction product (often referred to as single pass conversion²⁹⁻³¹) and (ii) the loss of CO₂ due to the homogeneous carbon equilibrium reactions in the liquid electrolyte, respectively (Figure 1c). The current state of the art techno-economic assessments rely on a fixed set of these performance variables, which are chosen independently of each other.⁶,¹¹,¹³ To illustrate the propagation of the interdependencies across scales, we introduce three exemplary models (see Table 2). The no channel model (M1) is based on the current state of the art and, therefore, neglects the interdependencies on the channel scale. For M1, we use a variable current density \( i_{\text{tot}} = [50–250 \text{ mA cm}^{-2}] \) in combination with fixed electrochemical variables \( V_{\text{cell}} = 3.69 \text{ V} \) and \( FE_{\text{C}_2\text{H}_4} = 0.7 \), and the fixed conversion rates \( \chi_{\text{het}} = 0.5 \) and \( \chi_{\text{hom}} = 0 \). The simplistic (M2) and full (M3) channel models are governed by the physics at the channel scale with increasing level of detail (Figure 2). They, therefore, capture interdependencies between the performance variables as further explained in the following section. All three models are embedded in the same electrolyser and process scale model.

### Channel Scale Model

The channel scale model (M2) considers two-dimensional channel scale models for a GDE-cell, which are most applicable for shallow channels in which the height is much smaller than the width \( (W = 10H \text{ in this work; Table 1}) \). The interdependencies among the performance variables \( i_{\text{tot}}, V_{\text{cell}}, FE_{\text{C}_2\text{H}_4}, \chi_{\text{het}}, \) and \( \chi_{\text{hom}} \) are captured through a mechanistic model of the GDE-cell. The GDE-cell is characterized by a gaseous and a liquid flow channel separated by a gas diffusion electrode in the cathode compartment. The anode is separated from the cathode side by a proton exchange membrane (Figure 1a), which is not explicitly modeled in this work.

The simplistic channel model (M2) solely considers the gas flow channel, with the catalyst layer modeled as an abrupt interface (see Figure 2, top). For this, the description of the homogeneous consumption \( \chi_{\text{hom}} \) is fixed and solely depends on the single channel gas flow rate \( u_g \). With a fixed additional current density \( i_{\text{f}} = 50 \text{ mA cm}^{-2} \) and no mass transport or species interactions in the gas phase, the cell model (M3) is reduced to the gas flow channel, the catalyst and the liquid boundary layer in the electrolyte compartment. Model M3 hence includes the parasitic homogeneous reactions occurring in the catalyst layer, thereby allowing to fully resolve the homogeneous consumption. The governing equations for M2 and M3 are given in the following sections, together with the relevant assumptions. The chosen channel dimensions and operating parameters are listed in Table 1. All other input parameters used in models M2 and M3 are listed in section S2. All relevant derivations, boundary conditions, the model validation, and the discussion of assumptions for model M3 are given in section S3.

### Mass Transport and Species Balance

We present a model to describe the concentration \( c_i \) of species \( k \) along the channel length and across the three layers (gas flow channel, catalyst layer, and liquid boundary layer) with coordinates \( x \) and \( y \) as defined in Figure 2. The simplistic channel model M2 and the other ⁶,¹¹,¹³

### Table 1. Overview of Channel Dimensions and Operating Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( L )</td>
<td>[m]</td>
<td>0.10</td>
<td>Channel length</td>
</tr>
<tr>
<td>( H )</td>
<td>[m]</td>
<td>1.00 × 10⁻³</td>
<td>Channel height</td>
</tr>
<tr>
<td>( W )</td>
<td>[m]</td>
<td>1.00 × 10⁻²</td>
<td>Channel width</td>
</tr>
<tr>
<td>( H_c )</td>
<td>[m]</td>
<td>3.00 × 10⁻⁸</td>
<td>Catalyst layer thickness</td>
</tr>
<tr>
<td>( \epsilon )</td>
<td>[-]</td>
<td>0.70</td>
<td>Porosity</td>
</tr>
<tr>
<td>( T )</td>
<td>[K]</td>
<td>300</td>
<td>Temperature</td>
</tr>
<tr>
<td>( P )</td>
<td>[Pa]</td>
<td>1.00 × 10⁵</td>
<td>Pressure</td>
</tr>
</tbody>
</table>

### Table 2. Values of Performance Variables for the No (M1), Simplistic (M2), and Full (M3) Channel Model, with a Variable Current Density \( i_{\text{tot}} = [50–250 \text{ mA cm}^{-2}] \)

<table>
<thead>
<tr>
<th>Model</th>
<th>( V_{\text{cell}} ) [V]</th>
<th>( FE_{\text{C}_2\text{H}_4} ) [-]</th>
<th>( \chi_{\text{het}} ) [-]</th>
<th>( \chi_{\text{hom}} ) [-]</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1</td>
<td>3.69</td>
<td>0.70²⁸</td>
<td>0.50</td>
<td>0.00²³</td>
</tr>
<tr>
<td>M2</td>
<td>( f(\chi_{\text{het}}) )</td>
<td>( f(\cdot) )</td>
<td>( f(\cdot) )</td>
<td>0.13²³</td>
</tr>
<tr>
<td>M3</td>
<td>( f(\chi_{\text{het}}) )</td>
<td>( f(\cdot) )</td>
<td>( f(\cdot) )</td>
<td>( f(\cdot) )</td>
</tr>
</tbody>
</table>

²³Performance variables which are dependent on more than one other variable are denoted as \( f( \cdot ) \). With a fixed additional current density \( (\chi_{\text{hom}} = 50 \text{ mA cm}^{-2}) \) and a single channel gas flow rate of 10 sccm, for more details see section S1.
**Simplistic channel model M2**

![Diagram of Simplistic channel model M2](image)

**Full channel model M3**

![Diagram of Full channel model M3](image)

**Figure 2.** *Simplistic* channel model (M2) with an assumed, fixed homogeneous consumption of CO$_2$, and the *full* channel model (M3) with a fully resolved homogeneous consumption of CO$_2$.

The *full* channel model M3 share the same modeling domain for the gas flow channel. In the gas flow channel, pure gaseous CO$_2$ is introduced and described by plug-flow behavior. The gaseous mass transport in the porous gas diffusion layer is neglected and the catalyst layer is assumed to be fully flooded. Therefore, the phase transfer of gaseous CO$_2$ to the liquid electrolyte occurs at the interface of the gas channel and the catalyst layer. The concentration of CO$_2$ is assumed to be in equilibrium at the gas–liquid interface. The species balance for gaseous compounds (CO$_2$(g), C$_2$H$_6$(g), and H$_2$(g)) in the gas channel is described by

$$\frac{\partial z_i}{\partial x} = \frac{n_{ig}(x)}{u_x}$$

where $u_x$ is the superficial gas velocity in the gas flow channel and $H$ is the channel height (see section S.3.1 for derivation). The term $n_{ig}(x)$ denotes the molar flux (per unit area) across the gas-catalyst interface. This flux is equal to the molar production or consumption rate of the gaseous compounds over the catalyst layer height $H$ at any location $x$ in the single channel

$$n_{ig}(x) = \int_0^{Hc} (N_{k,het} + N_{k,hom})dy$$

(2)

The term $N_{k,hom}$ denotes the consumption/production rate of CO$_2$(g), C$_2$H$_6$(g), and H$_2$(g) in mol s$^{-1}$ m$^{-3}$ due to the heterogeneous electrochemical reactions, while $N_{k,hom}$ in eq 2 denotes the consumption rate of the dissolved CO$_2(aq)$ due to the homogeneous buffer reactions in the liquid electrolyte.

For the *simplistic* channel model (M2) the catalyst layer is not explicitly modeled, with the consumption rate considered as part of the heterogeneous reaction term, adding an additional current density for the homogeneous consumption rate in $n_{hom} = 50$ mA cm$^{-2}$ (see section S1). By approximating the homogeneous consumption with a fixed additional heterogeneous reaction rate, the species balance for dissolved CO$_2(aq)$ and the ionic species does not need to be solved. This approach eliminates $N_{k,hom}$ from eq 2 and therefore allows straightforward calculation of the (single channel gas flow rate dependent) consumption rate (section S1). The concentration of the gaseous compounds CO$_2$, C$_2$H$_6$, and H$_2$ along the channel length is then fully described by eqs 1 and 2.

For the *full* channel model (M3), the catalyst layer and the liquid boundary layer are fully captured by explicitly solving the species balance for all species (including OH$^-$, HCO$_3^-$, and CO$_3^{2-}$ as the ionic species considered in this work), which allows one to calculate the homogeneous consumption rate. Similar to previous modeling studies$^{19,25}$ in the following we neglect migration for all ionic species and the crossover of carbonate and bicarbonate to the anode side.$^{32–34}$ The steady state species balance of the dissolved CO$_2(aq)$ and the ionic species in the catalyst layer ($0 \leq y < H_c$) is then governed by diffusion as well as homogeneous and heterogeneous reactions$^{19}$

$$0 = N_{k,diff} + N_{k,hom} + N_{k,het}$$

(3)

with the term $N_{k,diff}$ accounting for species transport through diffusion. In the catalyst layer, this term is calculated via

$$N_{k,diff} = \epsilon^{3/2} D_k \frac{\partial^2 c_k}{\partial y^2}$$

(4)

with the diffusion coefficient $D_k$ and the prefactor arising from the porosity $\epsilon$ and tortuosity $\tau = \epsilon^{-1/2}$.$^{35}$ Outside the catalyst layer, within the boundary layer ($H_c < y < H_c + \delta(x)$), the balance equation of the dissolved CO$_2(aq)$ and the ionic species is only governed by diffusion$^{25}$

$$0 = N_{k,diff}$$

(5)

with the diffusive transport given as

$$N_{k,diff} = D_k \frac{\partial^2 c_k}{\partial y^2}$$

(6)

The formation of the diffusive boundary layer on the liquid electrolyte side hinders the transport of fresh electrolyte to the catalyst layer, resulting in an increase in local pH and homogeneous reaction rate in the catalyst layer along the channel length.$^{28}$ This effect is included by calculating the thickness of the boundary layer according to the Lévêque approximation$^{36}$

$$\delta(x) = 1.022 \left( \frac{x H D_{HCO_3}}{u_x} \right)^{1/3}$$

(7)

with the average liquid electrolyte velocity $u_x$. The Lévêque approximation entails that two assumptions need to hold for eq 7 to be a good approximation: (a) constant concentration at the catalyst–electrolyte interface and (b) a developing boundary layer with $\delta(x) \ll H_c$. Since the supply of HCO$_3^-$ is the limiting factor in retaining the electrolyte buffer capacity in the catalyst layer, the length of the boundary layer is equally calculated for all species using the diffusion coefficient $D_{HCO_3^-}$. The concentrations of the ionic species are fixed to the equilibrium concentration in the 1 M KHCO$_3$ CO$_2$ saturated electrolyte at the liquid electrolyte/boundary layer interface, and the no flux boundary condition is imposed at the catalyst/gas channel interface. Similarly, the concentration of the dissolved CO$_2(aq)$ is assumed to be in equilibrium with the gaseous CO$_2$(g) concentration at the catalyst/gas interface, and the no flux boundary condition is imposed at the catalyst/boundary layer interface (see section 10133).
S3.2. This allows the species balance in the catalyst (eq 3) and boundary layer (eq 5) to be calculated, which are coupled through eq 2 to the species balance in the gas channel (eq 1). The concentration profile in the gas channel, catalyst layer, and liquid boundary layer is thereby fully described, with calculations of the required heterogeneous production and homogeneous consumption rate given in the following sections.

**Heterogeneous Reactions.** The heterogeneous electrochemical reduction reactions of \( \text{CO}_2(aq) \) and \( \text{H}_2 \text{O}(l) \) occur in the catalyst layer. Copper catalysts form a wide distribution of gaseous and liquid products, which in this work are limited to \( \text{C}_2\text{H}_4 \) and \( \text{H}_2 \), by considering the following two cathodic reactions:

\[
2\text{H}_2\text{O}(l) + 2e^- \rightarrow \text{H}_2(g) + 20\text{H}^-(aq) \tag{8}
\]

\[
2\text{CO}_2(aq) + 8\text{H}_2\text{O}(l) + 12e^- \rightarrow \text{C}_2\text{H}_4(g) + 12\text{OH}^-(aq) \tag{9}
\]

Note that this is a simplification in this work, and that, to the best of the authors knowledge, no catalyst for selective ethylene production has been reported. At the anode, the oxygen evolution reaction is facilitated, i.e. 

\[
4\text{OH}^-(aq) \rightarrow \text{O}_2(g) + 2\text{H}_2\text{O}(l) + 4e^- \tag{10}
\]

The electrochemical reaction rate for the species consumed or formed at the electrodes is calculated via Faraday’s law\textsuperscript{38}.

\[
N_{k,\text{het}} = \sum_r \nu_k z_r F H \frac{c}{c_F H \text{c}} \tag{11}
\]

in which \( \nu_k \) denotes the stoichiometric coefficient for species \( k \) in reaction \( r \), \( z_r \) the amount of transferred electrons in reaction \( r \), and \( F \) the Faraday constant. The current density \( i_{\text{tot}} \) is calculated via the Tafel equation fitted to experimental data reported by Tan et al.\textsuperscript{29} (see section S3.3)

\[
i_{\text{tot}} = i_0 \exp \left( - \frac{\alpha_F}{RT} \eta \right) \tag{12}
\]

with \( i_0 \) the exchange current density, \( \alpha \) the transfer coefficient, \( R \) the universal gas constant, and \( \eta \) the applied cathode overpotential. In fitting the data for the current density \( i_{\text{tot}} \), all reported carbonaceous species are considered to be ethylene, thereby simplifying the kinetic consideration. It is further assumed that hydrogen is only produced at the onset of mass transport limitations toward \( \text{CO}_2 \),\textsuperscript{39,40} which ensures that the current density \( i_{\text{tot}} \) remains constant over the electrode length (galvanostatically controlled) leading to the following partial current densities of \( \text{C}_2\text{H}_4 \) and \( \text{H}_2 \):

\[
i_{\text{C}_2\text{H}_4}(x, y) = i_{\text{tot}} e_{\text{CO}_2}(x, y)(aq) \tag{13}
\]

\[
i_{\text{H}}(x, y) = i_{\text{tot}} - i_{\text{C}_2\text{H}_4}(x, y) \tag{14}
\]

with \( e_{\text{CO}_2}(aq) \) the \( \text{CO}_2 \) equilibrium concentration within the electrolyte at standard conditions \( (P = 1.00 \times 10^5 \text{ Pa} \) and \( T = 300 \text{ K} \)). The dissolved \( \text{CO}_2(aq) \) concentration in the liquid electrolyte \( (e_{\text{CO}_2}(x, y)(aq)) \) relates to the gaseous \( \text{CO}_2 \) concentration along the channel length \( c_{\text{CO}_2} \) through Henry’s law. The changes in the \( \text{CO}_2(aq) \) concentration over the catalyst layer height are driven by the heterogeneous consumption (eq 11) and homogeneous conversion (eq 17).

The motivation and limits of these simplified kinetics are discussed in section S3.4.

**Homogeneous Reactions.** In addition to the heterogeneous reaction, \( \text{CO}_2 \) is also consumed by homogeneous reactions within the electrolyte in the catalyst layer. These reactions are constituted by the bicarbonate-buffer reactions, balancing the pH of the solution. This reaction mechanism is only considered in model M3 and described by the following equilibrium reactions\textsuperscript{19}

\[
\text{CO}_2(aq) + \text{OH}^-(aq) \rightleftharpoons \text{HCO}_3^-(aq) + \text{H}_2\text{O}(l) \tag{15}
\]

\[
\text{HCO}_3^-(aq) + \text{OH}^-(aq) \rightleftharpoons \text{CO}_3^{2-}(aq) + \text{H}_2\text{O}(l) \tag{16}
\]

where \( k_1 \) and \( k_2 \) are the forward reaction rate constants and \( k_1 \) and \( k_2 \) the respective reverse reaction rate constants, the values to all of which are provided in Table S4. Under consideration of the above equilibrium reactions, the volumetric homogeneous reaction terms can be written as\textsuperscript{17}

\[
N_{\text{CO}_2(aq),\text{hom}} = -k_2[\text{CO}_2][\text{OH}^-] + k_1[\text{HCO}_3^-] \tag{17a}
\]

\[
N_{\text{HCO}_3^-(aq),\text{hom}} = -k_1[\text{CO}_2][\text{OH}^-] + k_2[\text{HCO}_3^-][\text{OH}^-] \tag{17b}
\]

\[
N_{\text{CO}_3^{2-}(aq),\text{hom}} = k_1[\text{CO}_2][\text{OH}^-] - k_2[\text{HCO}_3^-][\text{OH}^-] \tag{17c}
\]

with the notation \( [c] \) used for the concentration \( c \). The significance of the homogeneous reactions can be explained by the increased rate of eq 15 at higher alkalinity, which inevitably occurs at elevated heterogeneous reaction rates due to increased hydroxide production. For simplicity, it is assumed that the homogeneous reactions occur solely within the catalyst layer, where the pH and \( \text{CO}_2(aq) \) concentration are highest. The void fraction within the catalyst layer is not accounted for.

**Electrolyser Scale Model.** The electrolyser scale model couples the calculated concentration profiles from the channel scale model to the process scale model (Figure 1b). First, the input variables \( \chi_{\text{het}} \), \( \chi_{\text{hom}} \), \( FE_{\text{C}_2\text{H}_4} \), and \( V_{\text{cell}} \) are calculated based on the \( \text{CO}_2(g) \) and \( \text{C}_2\text{H}_4(g) \) concentrations obtained from the channel scale models M2 and M3 for a variable \( i_{\text{tot}} \). For this, the electrolyser is assumed to be composed of a number of hydraulically, thermally, and electrically independent channels. The overall conversion achieved in the electrolyser then equals the single-channel conversion, which is calculated assuming a constant pressure as

\[
\chi_{\text{tot}} = \frac{c_{\text{CO}_2}(x = 0) - c_{\text{CO}_2}(x = L)}{c_{\text{CO}_2}(x = 0)} \tag{18}
\]

with the channel length \( L \). The heterogeneous conversion is calculated as\textsuperscript{38,39}

\[
\chi_{\text{het}} = \frac{2c_{\text{C}_2\text{H}_4}(x = L)}{c_{\text{CO}_2}(x = 0)} \tag{19}
\]

with \( 2 \) being the stoichiometric coefficient (see eq 9). The homogeneous consumption is then calculated as the difference between those figures, i.e.
\[ X_{\text{hom}} = X_{\text{tot}} - X_{\text{net}} \] (20)

The faradaic efficiency (selectivity) toward ethylene is calculated based on the product concentration and gas velocity \( u_g \) at the channel outlet \(^{32}\) (see section S4)

\[ FE_{C_2H_4} = \frac{c_{C_2H_4}(x = L)12u_gFH}{i_{\text{tot}}} \] (21)

with 12 being the amount of electrons required to reduce CO\(_2\) to C\(_2\)H\(_4\). The cell voltage is further related to the current density (see eq 12) via

\[ V_{\text{cell}} = E^0_a + \eta_a(i_{\text{tot}}) + \frac{1}{2}E^0_c + \eta_c(i_{\text{tot}}) + \eta_k(i_{\text{tot}}) \] (22)

with the constant anodic and cathodic standard potentials \( E^0_{a,c} \) and the current density dependent overpotentials \( \eta_{a,c,\Omega} \) (see section S5 for more details). The input variables for the electrolyser scale models M2 and M3 (Table 2) are then fully described by eqs 18–22.

Based on these input variables, the required electrolyser area \( A_r \), the volumetric gas flow rate \( \dot{V}_r \), the annual mass flow rate \( \dot{m}_{\text{CO}_2} \), and the power consumption \( P_r \) are calculated next (see section S4 for derivations), to estimate the investment and operating costs for the electrolyser and separation unit. The required electrolyser area is calculated taking into account the faradaic efficiency as well as the current density

\[ A_r = \frac{\dot{F}_{C_2H_4\text{target}}12F}{i_{\text{tot}}FE_{C_2H_4}} \] (23)

with the daily production target 10,000 kg d\(^{-1}\) converted to \( \dot{F}_{C_2H_4\text{target}} \approx 4.13 \) mol s\(^{-1}\). Further, the volumetric flow rate associated with the electrolyser setup is calculated from the heterogeneous conversion and target production rate as

\[ \dot{V}_r = \frac{2\dot{F}_{C_2H_4\text{target}}}{X_{\text{net}}c_{\text{CO}_2}(x = 0)} \] (24)

The annual mass flow rate of CO\(_2\) through the electrolyser then follows from the annual production target (\( \dot{m}_{C_2H_4\text{target}} = 3,500,000 \) kg yr\(^{-1}\)). The corresponding annual consumption rate of CO\(_2\) equals

\[ \dot{m}_{\text{CO}_2} = \left(1 + \frac{X_{\text{hom}}}{X_{\text{net}}}\right)^{\frac{44}{28}}\dot{m}_{C_2H_4\text{target}} \] (25)

Finally, the overall power consumption \( P_r \) follows from the product of the cell voltage and current density, i.e.

\[ P_r = V_{\text{cell}}i_{\text{tot}}A_r \] (26)

Process Scale Model. The process scale model describes the overall process and its economic performance. This model represents a simplified plant layout based on previous techno-economic analyses,\(^{6,13}\) with a CO\(_2\) feed source through direct air capture (DAC), an electrolyser unit for the electrochemical CO\(_2\)-reduction, and subsequent gas separation in a pressure swing adsorption (PSA) unit (Figure 1 a)). Liquid pre- and postprocessing steps are not taken into account. The currency used is the US dollar (multiple years).

The selected process economic indicator is the end-of-lifetime net present value (NPV) of the overall process, assuming 20 years of continuous operation.\(^5\) The NPV is calculated by taking into account the cash flow \( CF(t) \) on an annual basis as

\[ \text{NPV} = \sum_{t=0}^{20} \frac{CF(t)}{(1 + \text{IR})^t} \] (27)

in which \( t \) is the respective year of operation. The term IR denotes the interest rate and is assumed to be 10% throughout the lifetime.\(^6\) It is assumed that the plant is erected within the first year and operates at full capacity for the remaining 19 years of operation with the cash flow calculated as

\[ CF(t) = \begin{cases} \text{TCI}, & \text{for } t = 0 \\ \text{Rev} - \text{Op} - \text{M}, & \text{for } t \geq 1 \end{cases} \] (28)

where \( \text{Rev}, \text{Op} \), and \( \text{M} \) describe the annual revenue, operating costs, and maintenance costs, respectively. TCI is the total capital investment, which comprises the costs for the electrolyser, the separation unit, and all additional infrastructural facilities (balance of plant). The investment costs for the electrolyser are proportional to the required electrolyser area (\$920 m\(^{-2}\)),\(^6\) while the costs for the PSA unit scale with the overall volumetric flow rate at the electrolyser outlet with a reference cost\(^{41}\) of 1.99 MS and a reference flow rate of 1000 m\(^3\) h\(^{-1}\). The balance of plant costs are assumed to make up 35/65 of the electrolyser costs.\(^6\) The total capital investment based on the equipment costs can then be calculated as

\[ \text{TCI} = (\$920 \text{ m}^{-2}A_r)(1 + 35/65) + 1.99 \times 10^6 \left(\frac{2600 \text{ s}^{-1}}{1000 \text{ m}^3 \text{ h}^{-1}}\right)^6 \] (29)

where \( A_r \) is the required area (eq 23) and \( \dot{V}_r \) the volumetric flow rate at the electrolyser outlet (eq 24). The term \( \beta \) is a fitting factor associated with the cost correlation for the PSA unit, assumed to be 0.7 according to the regression function proposed by Paturska et al.\(^{41}\) for flow rates between 500 m\(^3\) h\(^{-1}\) and 1400 m\(^3\) h\(^{-1}\).

The annual revenue depends on the annual production target, and market price of ethylene (herein taken as $1.3 kg\(^{-1}\))\(^6\) as

\[ \text{Rev} = 1.3 \text{ kg}^{-1}\dot{m}_{C_2H_4\text{target}} \] (30)

The annual costs are then determined by the CO\(_2\) price (herein taken as $0.04 kg\(^{-1}\)),\(^13\) which is slightly lower than the most optimistic assumption for commercial DAC units using chemical absorption.\(^{42}\) For the electricity price, an optimistic value of $0.03 kW h\(^{-1}\) is taken based on predictions published by Haegel et al.\(^{13}\) The operating costs associated with separation are assumed to be only made up of the electricity costs (0.25 kW h m\(^{-3}\)),\(^6\) which allows calculating the overall operating costs based on the annual consumption rate of CO\(_2\) (eq 25), the overall power consumption (eq 26), and the volumetric flow rate (eq 24) as

\[ \text{Op} = 0.04 \text{ kg}^{-1}\dot{m}_{\text{CO}_2} + 0.03 \text{ kW h}^{-1} \times (8400 \text{ h yr}^{-1}\\dot{P}_r) + 0.25 \text{ kW h}^{-1} \times 30.24 \times 10^3 \text{ s yr}^{-1}\\dot{V}_r^6 \] (31)

The annual maintenance costs are taken to be 2.5% of the capital investment costs for the electrolyser, i.e.

\[ \text{M} = 0.025 \times \$920 \text{ m}^{-2}A_r \] (32)
As a comparative figure for the models (M1 to M3) the relative NPV is defined as follows

\[ NPV_{\text{rel,M}} = \begin{cases} 
\frac{NPV_{Mx}}{\max(NPV_{M1})} & \text{for } 0 < NPV_{Mx} \\
\frac{\max(NPV_{M1})}{NPV_{Mx}} & \text{for } NPV_{Mx} \leq 0 
\end{cases} \quad (33) \]

for any model \( M_x \) with \( x \in \{1, 2, 3\} \) (for further detail, see section S6.1). The highest NPV of model M1 is taken as a reference point, as at this value the investment costs for the electrolyser do not influence the overall NPV anymore and the resulting current density is commonly reported as the target value in techno-economic analyses.\(^{5,6,13}\)

**Implementation.** For the no channel model M1 the input variables to the electrolyser scale are fixed (Table 2) such that the required electrolyser area, the volumetric gas flow rate, the annual mass flow rate of CO\(_2\), and the overall power input required to achieve the target production for C\(_2\)H\(_4\) (eqs 23–26) can be straightforwardly calculated. The relative NPV (eq 33) is then calculated based on these variables as a comparison metric. For the simplistic channel model (M2) and the full channel model (M3), the input variables to the electrolyser scale depend on the output of the channel scale. Therefore, the concentrations of CO\(_2\), C\(_2\)H\(_4\), and H\(_2\) along the channel length need to be calculated first. The concentrations are then used to calculate the input variables to the electrolyser scale (eqs 18–22) and subsequently the input variables to the process scale are calculated (eqs 23–26), followed by the NPV and relative NPV.

The species balance (eq 1) in the gas channel is solved for models M2 and M3 via Heun’s method for varying current densities (\( i_{\text{cell}} \)) and gas velocities (\( u_g \)). For the simplistic model (M2), the flux across the gas-catalyst interface (eq 2) is given as a boundary condition, while for the full channel model (M3) the flux across the gas-catalyst interface is updated at every finite difference by solving the governing equations in the catalyst and boundary layer (eq 3 and S). The solution to these respective domains is found with the Matlab R2020a built-in solver bvp4c, where the extent of the boundary layer is adapted on each step according to eq 7 and the solution of the previous step is supplied as an initial guess to the current step.

The extend of the boundary layer is calculated for a fixed electrolyte flow rate of 0.54 m s\(^{-1}\) to minimize the consumption of CO\(_2\) due to the buffer reaction,\(^{25}\) while still ensuring operation in the laminar flow regime.

To gain insight into the optimal mode of operation, the model has been constructed to allow for facile use with the Matlab built-in nonlinear optimizer fmincon and fminsearch. All Matlab scripts are made available via GitHub (see Data Availability).

### RESULTS AND DISCUSSION

In this section, we first discuss the interdependencies between three performance variables on the channel scale for the no channel (M1), the simplistic channel (M2), and the full channel (M3) model. For this, we consider the heterogeneous conversion \( x_{\text{het}} \) the simplistic channel \( x_{\text{hom}} \), and the faradic efficiency \( FE_{\text{C,H}} \) and their dependence on the current density. As the cell voltage \( V_{\text{cell}} \) can be straightforwardly calculated (eq 22) it is not explicitly discussed. Second, we discuss the propagation of the interdependencies from the channel scale to the process scale model in terms of the relative NPV for varying current densities and gas velocities. The contribution of selected technical input variables to the process NPV, together with the contribution of economic parameters, is evaluated in a sensitivity analysis in Section S6.2. Lastly, the optimization results are presented and discussed in light of current developments in the literature.

**Interdependency of Performance Variables on the Channel Scale.** The heterogeneous conversion \( x_{\text{het}} \), homogeneous consumption \( x_{\text{hom}} \), and selectivity \( FE_{\text{C,H}} \) with varying current densities (\( i_{\text{cell}} \)) are presented for models M1 to M3 in Figure 3. Contrary to the case of fixed performance variables (M1), using a mechanistic channel scale model results in a dependency of the above-stated variables predicting the expected trend of increased conversion rates with higher current densities as shown in Figure 3a. For low conversions, this trend is linear, as the supply of CO\(_2\) to the catalyst is not limited (for more details, see section S7). For the simplistic channel model M2, the limit of the linear scaling is reached at higher conversions than that of the full channel model (M3). This can be explained by comparing the trends for the consumption rate of CO\(_2\) for models M2 and M3 in...
While the consumption rate is constant in model M2 the consumption of CO$_2$ in M3 increases with increasing current densities. Increased consumption rates of CO$_2$ limit the availability of CO$_2$ at the catalyst site and therefore limit the heterogeneous conversion, leading to a deviation from the linear scaling at lower current densities for M3 compared to M2. The steep exponential increase in homogeneous consumption for model M3 can be explained by the dependence of the formation of hydroxide ions on the current density, as seen in eqs 8 and 9, paired with the limited buffer capacity of the electrolyte. This eventually leads to higher consumption rates through the homogeneous reaction than through the heterogeneous reaction at high current densities. This, in turn, influences the selectivity, resulting in a steep decrease of $FE_{C_2H_4}$ toward the formation of ethylene with increased current density for M3 as shown in Figure 3c.

**Figure 4.** Relative NPV as a function of current density for a fixed single channel gas flow rate $u_g \times H \times W = 10$ sccm (a) and as a function of the single channel gas flow rate for a fixed current density $i_{tot} = 100$ mA cm$^{-2}$ (b). Schematic of main cost drivers for varying current densities and single channel gas flow rates with the arrows indicating the direction of decrease of the respective cost unit (c). Contours (with an equidistant spacing of 0.05) of the relative NPV for varying current densities and single channel gas flow rates for M1 (d), M2 (e), and M3 (f) with the dots indicating the optimum for model M2 and M3. For model M2 a fixed additional current density ($i_{hom} = 50$ mA cm$^{-2}$) is used to account for the homogeneous reactions, while these are fully resolved in model M3 for a fixed single channel liquid flow rate of 325 mL min$^{-1}$ ($u_l = 0.54$ m s$^{-1}$).

**Propagation of Interdependencies from the Channel Scale to the Process Scale Model.** Having established how the level of detail at the channel scale influences the input for the electrolyser scale (in the form of $\chi_{het}$, $\chi_{hom}$, $FE_{C_2H_4}$, and $V_{Cell}$ for a given $i_{tot}$, $u_g$, and $i_{hom}$ (M2) or $u_l$ (M3)), the propagation of the level of detail to the process scale is shown in terms of the relative NPV in Figure 4. The relative NPV (eq 33) compares the NPV (eq 27) for each model to the maximum NPV reached with model M1 ($NPV \approx 24$ M$\$ at $i_{tot} \approx 600$ mA cm$^{-2}$). The calculations for the maximum NPV of model M1 can be found in section S6.1 with a discussion on the negative NPV in section S8. Therefore, a decrease or increase of the NPV$_{rel,Mx}$ indicates the same relative change in the NPV, and hence, both terms will be used interchangeably in the following. In Figure 4 (a and b), the green lines depict the NPV$_{rel,M1}$ calculated with the no channel model (M1), with the current density being the only variable input parameter to the electrolyser scale (see Table 2 and Figure 1). As described in current densities is the increased heterogeneous conversion instead of the homogeneous consumption.
previous literature, the current density in M1 solely influences the electrolyser area (eq 23) and therefore the investment costs leading to the expected trend of a steady increase in the NPV with an increase in current density as shown in Figure 4a, eventually reaching the asymptotic value of 1. Contrary to model M1 an increase in current density leads to a decrease in the NPV for high current densities (see section S6.2), for both mechanistic models (M2 and M3) resulting in a clear optimum within the range of variation of the current density. A similar trend is observed when fixing the current density while varying the single channel gas flow rate, as shown in Figure 4b. While the single channel gas flow rate shows no effect on the NPV in model M1 as the input variables are fixed, both mechanistic models display again a clear optimum. The significance of the single channel gas flow rate on the NPV was also observed in the sensitivity analysis in section S2.6. This analysis reveals that the interdependencies of the input variables translate to important trade-offs on the process scale, which cannot be captured with fixed variable models such as model M1.

The maxima in the NPV for channel models M2 and M3 are explained through the interdependence of mass transfer limitation, heterogeneous conversion, and homogeneous consumption. Figure 4c shows this trade-off schematically in terms of investment and operating costs linked to the current density and single channel gas flow rate. From the previous section, we learned that low current densities retain low heterogeneous conversion rates and high selectivities. Further the high CO2 concentrations in the catalyst layer cause the mass transfer related overpotential (eq 13), and hence the overall cell potential (eq 22), to decrease. A low cell potential reduces the required power input, translating to lower operating costs for the electrolyser. However, due to the low conversion, larger electrode areas and separation units are required to maintain a specific throughput, which increases the investment costs. High current densities on the other hand increase the heterogeneous conversion rate and therefore reduce investment costs but also lead to an unwanted expense of electrons through the increased reduction of water in the hydrogen evolution reaction (eq 8), therefore increasing the operating costs. The trade-off between investment and operating costs is similarly observed for varying single channel gas flow rates. High single channel gas flow rates reduce the residence time in the channel and consequently reduce the heterogeneous conversion rate leading to an increased daily gas throughput to achieve the target production rate of ethylene. This in turn requires a larger separation unit increasing the investment costs. Low single channel gas flow rates, on the other hand, increase the heterogeneous conversion, lowering the daily gas throughput. However, this leads to a depletion of CO2 along the channel, which resulted in an increase toward the hydrogen evolution reaction. This in turn increases the operating costs of the electrolyser as electricity is now lost toward the parasitic side reaction.

The insights on the trade-offs explain the difference in the impact of current density and gas flow rate on the costs. The single channel gas flow rate mainly influences the heterogeneous conversion rates through the residence time in the channel and consequently, the investment costs of the gas separation unit and the operating costs of the electrolyser. The current density, however, influences both the investment and operating costs of the electrolyser as well as for the separation unit.

Mapping the relative NPV for all three models (M1–M3) over the space of varying current densities and single channel gas flow rates allows us to compare how the optimal operating areas vary with the level of mechanistic detail in the model. Figure 4d–f show that for all models low current densities and high single channel gas flow rates are not optimal based on the high investment costs. However, only models M2 and M3 additionally show a higher loss region for high current densities and low single channel gas flow rates. The no channel model (M1) does not display this trade-off due to a fixed conversion rate (Table 2) leading to fixed operating costs. Therefore, a clear optimum for the operating conditions is found for the models considering the interdependencies between the performance variables (indicated by the dots in Figure 4e and f).

**Optimization Results.** The pronounced impact of the current density and single channel gas flow rate on NPV has been discussed in the previous section. It was further shown that considering the interdependencies (model M2 and M3) between the performance variables leads to a trade-off between investment and operating costs which manifests in a clear optimum for the operating conditions. The optimization results are summarized in Table 3. Note that the optimal results for M1 are not shown as the conversion rate is fixed, and the optimal current density always lies at the upper constraint, i.e., at 250 mA cm$^{-2}$. Figure 5 compares the optimization results with literature based operating targets. It can be seen that the optimal current density for M2 lies close to the values suggested as minimal-threshold in non-mechanistic techno-economic analysis (above 200 mA cm$^{-2}$), while this value is considerably lower when modeling the consumption of CO2 as a function of the process conditions (M3). Here, the optimal current density lies at $\approx 100$ mA cm$^{-2}$, roughly half the value that is found for M2 and lower than the threshold values proposed in the literature. This is driven mainly by the prediction of a strong increase in unwanted homogeneous consumption at high current densities, rendering operation at lower current densities desirable. The observed product distribution for the formation of higher hydrocarbons and the reported increase in selectivity with

<table>
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<tr>
<th>Table 3. Overview of the Optima for Models M2 and M3 As Shown in Figure 4e and f</th>
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<td>Variable</td>
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</tr>
<tr>
<td>Input</td>
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<td>$i_{\text{tot}}$</td>
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<td>$u_g$</td>
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<td>Channel scale</td>
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<td>Electrolyser scale</td>
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<td>$P_i$</td>
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<td>$m_{\text{CO}_2}$</td>
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<td>Process scale</td>
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$^a$For model M3, the obtained optimal values show a dependency on the initial guess with a deviation of less than 5%.
increasing current density are neglected in this work. As the kinetic expressions (eqs 11–14) and the assumptions regarding the separation unit are the same in models M2 and M3, the trade-off for the heterogeneous conversion is the same in both models. Hence, the optimizer converges to a single channel gas flow rate, which facilitates a similar conversion rate for both models of slightly below 20%. This comparably low conversion is subject to how the kinetics are formulated in this work, leading to a strictly inverse relationship between the conversion and faradaic efficiency. Besides the simplification of the kinetics on the findings, other important assumptions are taken. In the remainder of the paper, we discuss their foreseen impact on our findings.

Limitations of Results. The presented channel scale model is more advanced than the majority of CO2R models reported previously, yet several phenomena are not included such as liquid product formation and their migration to and oxidation at the anode, migration of bicarbonate/carbonate ions, and hence degassing of CO2 at the anode side. While we provide a detailed discussion on the influence of these phenomena on our findings in section S9, we highlight here the influence of migration and degassing of CO2. In anion exchange membrane electrolysers for example a significant amount of CO2 crosses over to the anode side in form of carbonate ions as the main charge carrying species under steady-state conditions.32 This increases the consumption rate of CO2 compared to the predictions of model M3,34 leading to lower optimal CO2 conversions. In bipolar membrane electrolysers this effect can be minimized at the cost of increasing the ohmic resistance and hence required cell potential,33 decreasing the overall NPV (see section S8). Further, we assume the investment and operating costs of the separation unit (PSA) to depend only on the overall required single channel gas flow rate and not on the composition of the gas stream itself. Additionally, the liquid products and potential cleaning steps of the electrolytes are not considered. Conceivably, including detailed models of the required post treatment units leads to higher investment costs and optimal CO2 conversion rates.

Considering the various assumptions underlying these results, the authors stress that the herein reported optimal values shall not be understood as newly proposed target values for the performance variables. Rather, they shall showcase how combining mechanistic and techno-economic models allows for design optimizations in the field of CO2 electrolysis. More importantly, they show how the level of mechanistic detail in such models strongly influences the resulting recommendations. In this sense, the distance between the optimum points for the models M1, M2, and M3 in Figure 5 can be understood as resulting from different levels of mechanistic understanding, while the distance between the optimum points and the literature recommendations results from a discrepancy between required and currently possible electrolyser performances.

CONCLUSION AND OUTLOOK

Target values for performance variables for the low-temperature electrochemical conversion of CO2 have so far been derived from techno-economic analysis based on the assumption that the performance variables such as current density and faradaic efficiency are independent. In this work, we present a multiscale framework that incorporates mechanistic models of a GDE-based CO2 electrolyser to capture the interdependence between the performance variables required as input to the electrolyser scale: heterogeneous conversion, homogeneous consumption, faradaic efficiency, and cell voltage. This framework is used to perform a techno-economic assessment and optimization for a CO2-electrolysis-based process, revealing optimal target values for the performance variables that can strongly deviate from previously reported targets. For the herein chosen electrolyser design this manifests in an optimal current density of around half of commonly reported values. While it should be noted that the optima in this work are derived based on simplified reaction mechanisms and design considerations and therefore should not be taken as fixed optimum values for future electrolyser designs, the used approach nonetheless highlights the dependency of the mechanistic detail and interdependencies between performance variables on the economic viability of an electrolyser design. This work further presents a tool to evaluate electrolyser design choices based on an economic objective, which in its generic form can be applied to various electrolyser designs18,44–47 and CO2-reduction products, such as CO or formate19,25,48. For different electrolyser designs and products, the modeling assumptions, and hence, the economic predictions, are highly dependent on the available data; therefore, it is important to (a) move toward more holistic, multiscale modeling approaches in the field of CO2 electrolysis and to (b) communicate measured or targeted electrolyser performance with all applicable boundary conditions, including achieved conversions.

ASSOCIATED CONTENT

Data Availability Statement
The full multiscale model is available via https://github.com/IsabellBagemihl/MultiscaleModelElectrochemicalCO2Reduction.git.
Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssuschemeng.3c02226.
Detailed description including the derivation and validation of the single channel model, all material and operating parameters, derivation of electrolyser scale model, sensitivity analysis towards technical variables and economic parameters, and discussion of selected parameters and assumptions and their effect on the findings (PDF)

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Notes
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