Membraneless Electrolyzers for Solar Fuels Production

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ABSTRACT

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Solar energy has the potential to meet all of society’s energy demands, but challenges remain in storing it for times when the sun is not shining. Electrolysis is a promising means of energy storage which applies solar-derived electricity to drive the production of chemical fuels. These so-called solar fuels, such as hydrogen gas produced from water electrolysis, can be fed back to the grid for electricity generation or used directly as a fuel in the transportation sector. Solar fuels can be generated by coupling a photovoltaic (PV) cell to an electrolyzer, or by directly converting light to chemical energy using a photoelectrochemical cell (PEC). Presently, both PV-electrolyzers and PECs have prohibitively high capital costs which prevent them from generating hydrogen at competitive prices. This dissertation explores the design of membraneless electrolyzers and PECs in order to simplify their design and decrease their overall capital costs.

A membraneless water electrolyzer can operate with as few as three components: A cathode for the hydrogen evolution reaction, an anode for the oxygen evolution reaction, and a chassis for managing the flows of a liquid electrolyte and the product gas streams. Absent from this device is an ionically conducting membrane, a key component in a conventional polymer electrolyte membrane (PEM) electrolyzer that typically serves as a physical barrier for separating product gases generated at the anode and cathode. These membranes can allow for compact and efficient electrolyzer designs, but are prone to degradation and failure if exposed to impurities in the electrolyte. A membraneless electrolyzer has the opportunity to reduce capital costs and operate in non-pristine environments, but little is known about the performance limitations and design rules that govern operation of membraneless electrolyzers. These design rules require a
thorough understanding of the thermodynamics, kinetics, and transport processes in electrochemical systems. In Chapter 2, these concepts are reviewed and a framework is provided to guide the continuum scale modeling of the performance of membraneless electrochemical cells. Afterwards, three different studies are presented which combine experiment and theory to demonstrate the mechanisms of product transport and efficiency loss.

Chapter 3 investigates the dynamics of hydrogen bubbles during operation of a membraneless electrolyzer, which can strongly affect the product purity of the collected hydrogen. High-speed video imaging was implemented to quantify the size and position of hydrogen gas bubbles as they detach from porous mesh electrodes. The total hydrogen detected was compared to the theoretical value predicted by Faraday’s law. This analysis confirmed that not all electrochemically generated hydrogen enters the gas phase at the cathode surface. In fact, significant quantities of hydrogen remain dissolved in solution, and can result in lower product collection efficiencies. Differences in bubble volume fraction evolved along the length of the cathode reflect differences in the local current densities, and were found to be in agreement with the primary current distribution. Overall, this study demonstrates the ability to use in-situ HSV to quantitatively evaluate key performance metrics of membraneless electrolyzers in a non-invasive manner. This technique can be of great value for future experiments, where statistical analysis of bubble sizes and positions can provide information on how to collect hydrogen at maximum purity.

Chapter 4 presents an electrode design where selective placement of the electrocatalyst is shown to enhance the purity of hydrogen collected. These “asymmetric electrodes” were prepared by coating only one planar face of a porous titanium mesh electrode with platinum electrocatalyst. For an opposing pair of electrodes, the platinum coated surface faces outwards such that the electrochemically generated bubbles nucleate and grow on the outside while ions conduct through
the void spacing in the mesh and across the inter-electrode gap. A key metric used in evaluating the performance of membraneless electrolyzers is the hydrogen cross-over percentage, which is defined as the fraction of electrochemically generated hydrogen that is collected in the headspace over the oxygen-evolving anode. When compared to the performance of symmetric electrodes – electrodes coated on both faces with platinum – the asymmetric electrodes demonstrated significantly lower rates of cross-over. With optimization, asymmetric electrodes were able to achieve hydrogen cross-over values as low as 1%. These electrodes were then incorporated into a floating photovoltaic electrolysis device for a direct demonstration of solar driven electrolysis. The assembled “solar fuels rig” was allowed to float in a reservoir of 0.5 M sulfuric acid under a light source calibrated to simulate sunlight, and a solar to hydrogen efficiency of 5.3% was observed.

In Chapter 5, the design principles for membraneless electrolyzers were applied to a photoelectrochemical (PEC) cell. Whereas an electrolyzer is externally powered by electricity, a PEC cell can directly harvest light to drive an electrochemical reaction. The PEC reactor was based on a parallel plate design, where the current was demonstrated to be limited by the intensity of light and the concentration of the electrolyte. By increasing the average flow rate of the electrolyte, mass transport limitations could be alleviated. The limiting current density was compared to theoretical values based off of the solution to a convection-diffusion problem. This modeled solution was used to predict the limitations to PEC performance in scaled up designs, where solar concentration mirrors could increase the total current density. The mass transport limitations of a PEC flow cell are also highly relevant to the study of CO₂ reduction, where the solubility limit of CO₂ in aqueous electrolyte can also limit performance.
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CHAPTER 1
INTRODUCTION

1.1 Electrochemically generated fuels for energy storage

The sun is a highly abundant resource which has the potential to meet all of society’s energy demands without emitting greenhouse gases. A pitfall of solar energy is that it is intermittent, and must be stored for use during hours when the sun does not shine. An energy infrastructure which is both renewable and robust will be able to store solar electricity by transferring it into chemical energy. This can be achieved using an electrolyzer, an electrochemical reactor that uses an electrical power source to drive a thermodynamically uphill reaction. One of the simplest of these reactions is the electrolysis of water into hydrogen (H$_2$) and oxygen (O$_2$) gases. As an energy carrier, H$_2$ is storable and can be used as a fuel source for on demand electricity generation. Additionally, H$_2$ can be used as a fuel in the transportation sector. Although electric vehicles are emerging in the market for light transportation, chemical fuels will likely continue to be the dominant fuel source for commercial applications, especially in the airline and heavy freight industries.$^1$ In the chemical industry, H$_2$ will continue to be necessary for the production of ammonia for fertilizers, which is one of the leading applications for H$_2$ use today.$^2$ More broadly, electrolyzers are also of interest for the renewable production of commodity chemicals, where electrode materials have been demonstrated for the reduction of nitrogen$^3$ and carbon dioxide.$^{4,5}$ Although the design of membraneless electrolyzers is highly relevant to these processes, the focus of this dissertation is the production of H$_2$ from water electrolysis.
1.2 Water electrolysis

Hydrogen in today’s market is produced via the steam methane reforming (SMR) reaction, a process which relies on fossil fuels and releases carbon dioxide (CO₂). Water electrolysis is a more sustainable means of H₂ production, provided that there is a renewable source of electricity. An electrolyzer operates by applying a voltage across two electrodes separated by an electrolyte. In an acidic electrolyte, protons are reduced at the cathode to evolve H₂. At the anode, water is oxidized to evolve O₂. These half reactions, known as the hydrogen evolution reaction (HER) and the oxygen evolution reaction (OER) in an acidic electrolyte, are shown in Equations 1.1 and 1.2 respectively. The overall reaction, shown in Equation 1.3, is the splitting of water into H₂ and O₂ gases.

\[
\begin{align*}
2H^+ + 2e^- &\rightarrow H_2 & U^0_{H_2|H^+} = 0.0 \text{ V vs NHE} \quad 1.1 \\
H_2O &\rightarrow \frac{1}{2}O_2 + 2H^+ + 2e^- & U^0_{H_2O|O_2} = 1.23 \text{ V vs NHE} \quad 1.2 \\
H_2O &\rightarrow H_2 + \frac{1}{2}O_2 & U^0_{\text{cell}} = -1.23 \text{ V} \quad 1.3
\end{align*}
\]

Also shown in Equations 1.1 and 1.2 is the standard reduction potential \( U^0 \) of each half reaction, which is reported relative to the normal hydrogen electrode (NHE). The standard cell potential, \( U^0_{\text{cell}} = U^0_{H_2|H^+} - U^0_{H_2O|O_2} = -1.23 \text{ V} \), is the thermodynamic minimum voltage which must be applied in order for the reaction to occur. Although Equations 1.1 and 1.2 are written assuming an acidic electrolyte, an acid intermediate is not strictly necessary for the overall reaction shown in Equation 1.3. The water electrolysis reaction can also be carried out in alkaline and pH-neutral electrolytes. Provided that the pH of the electrolyte is the same at both electrodes, \( U_{\text{cell}} \) will be equal to -1.23 V across the entire pH scale. At non-standard conditions, the cell potential \( U_{\text{cell}} \) can deviate from -1.23 V, and Chapter 2 explores how to calculate these deviations using the Nernst equation.
Regardless of the composition of the electrolyte, a larger voltage must be applied to overcome barriers due to kinetics, mass transport, and ohmic resistance. These losses are shown in Equation 1.4:

$$\Delta V = |U_{cell}| + \eta_{HER} + \eta_{OER} + \eta_{MT} + IR_s$$

where $\Delta V$ is the applied voltage to the electrolyzer electrodes, $\eta_{HER}$ is the kinetic overpotential for HER, $\eta_{OER}$ is the kinetic overpotential for OER, $\eta_{MT}$ is the mass transport overpotential, $I$ is the total current passed through the electrolyzer, and $R_s$ is the ohmic resistance in the electrolyte. Each of these loss mechanisms can significantly hamper the performance of an electrolyzer, and will be explored in greater detail in Chapter 2. Minimizing these voltage penalties is necessary for electrolysis to proceed efficiently. The overall efficiency of an electrolyzer $\eta_{elec}$ is given by Equation 1.5 below:

$$\eta_{elec} = \frac{|U_{cell}|}{\Delta V} \eta_{FE}$$

where $\eta_{FE}$ is the Faradaic efficiency, or selectivity, of the electrochemical reaction. For water electrolysis, it can be generally assumed that there are no major side reactions and that $\eta_{FE}$ is 100%.

### 1.3 Economic motivations

The efficiency of an electrolyzer is an important metric for determining if it can be cost competitive with SMR for $H_2$ production. A technoeconomic analysis (TEA) by Shaner, et al. estimated that for an operating efficiency of 61%, the breakeven price for electrochemically generated $H_2$ is approximately $6.10/kg$.\textsuperscript{6} By comparison, the price of $H_2$ from SMR is approximately $1.59/kg$.\textsuperscript{7} This disparity in price can be attributed to the costs of inputs for each process. The required methane and heat input for SMR is inexpensive relative to the cost of
purified water and electricity for electrolysis. In fact, 66% of the cost of electrochemically generated H₂ can be attributed to the price of electricity consumed. By operating more efficiently, less electricity is required per kg of H₂, bringing down operating costs. However, even at 100% efficiency, the price of electrochemically generated H₂ would still not be cost competitive with SMR at today's electricity prices. A more fair comparison between the two technologies should also account for externalities such as the consequences of emitting CO₂ into the atmosphere. This could be accounted for through a carbon tax or other environmental regulations which discourage CO₂ emissions. Although H₂ from SMR can be produced at $1.59 today, an upper bound on this price would also include the cost of carbon capture and storage.

Another important consideration for the cost of H₂ from electrolysis is the capacity factor. The break-even price in the Shaner TEA assumed that the electrolyzer had a capacity factor of 97%, meaning that it was operating 97% of the time. However, an electrolyzer used for energy storage will operate at significantly lower capacity factors, particularly when coupled with solar electricity. For example, a fixed angle photovoltaic (PV) panel in Phoenix, Arizona can use an average of 6.5 hours of sunlight per day. If an electrolyzer was coupled with this PV system, its capacity factor would therefore be 27%. For this PV-electrolyzer pair to produce the same amount of H₂ at the same efficiency as the electrolyzer with a 97% capacity factor, you would need to increase the electrode area by a factor of four, and consequently the capital costs increase by a factor of four. In summary, electricity consumption accounts for 66% of the price of H₂ in the hypothetical case where an electrolyzer operates at 97% capacity factor. When the electrolyzer is used for energy storage, however, significantly higher capital investment is required to compensate for a lower capacity factor, and therefore the capital costs are expected to dominate.
Thus, efficiency improvements alone cannot make water electrolysis economically viable. There must also be reductions in the price of electricity and reductions in the capital cost of the electrolyzer itself. Fortunately, innovations in PV generation have shown steady declines in the price of renewable electricity in the past decade.\textsuperscript{10} There could also be opportunity for electrolyzers to purchase electricity at significant discount if operation is limited to hours of excess electricity generation. Decreasing the capital costs, however, requires reexamination of the reactor design of the electrolyzer itself, and is the central focus of this dissertation. In the next section, we review the current state of the art designs for water electrolyzers and the relationship between capital costs and energy efficiency.

1.4 Reactor designs for electrolyzers

An electrolyzer should be designed to operate efficiently while also ensuring that the generated gases can be collected with high purity. The efficiency of an electrolyzer is largely a product of the construction materials, but can also depend on the electrode separation distance. In general, an electrolyzer is most efficient when the electrodes are closest together. A smaller electrode separation distance will cause a decrease in the ohmic resistance and therefore increase the overall efficiency of the device. Placing electrodes too close, however, can cause the generated H\textsubscript{2} and O\textsubscript{2} to mix, which will incur downstream separation costs and possibly create an explosive mixture. An optimal electrolyzer design will balance this distance tradeoff between energy efficiency and product purity, which will largely depend on the transport mechanism of the reactants and products. Figure 1.1 shows a generalized schematic of the transport and separation processes for two established electrolyzer technologies: The polymer electrolyte membrane (PEM) electrolyzer and the alkaline electrolyzer.
A PEM electrolyzer, shown in Figure 1.1a, uses a proton exchange membrane such as Nafion to facilitate ionic transport between the electrodes. This design is a conventional approach for electrolysis, and is presently used in the chloralkali industry. Although ionically conductive, the membrane is also electronically insulating, allowing for extremely close electrode separation distances (<150 μm) without risk of electrical short circuiting. This narrow electrode separation distance is desirable because it reduces the ohmic resistance of the electrolyzer, and is sometimes referred to as a “zero gap resistance.” The membrane also serves as a rigid barrier to prevent gas permeation, ensuring that the electrochemically generated H₂ and O₂ streams have high purities that are outside the flammability range.

Figure 1.1b shows an alkaline electrolyzer, which is another conventional design for electrolyzers. Gas separation in an alkaline electrolyzer is maintained by a porous diaphragm, typically constructed out of asbestos. The ionic current between the electrodes is carried by an alkaline electrolyte which hydrates the pores of the diaphragm. The diaphragm typically has a higher ohmic resistance than the Nafion membranes in PEM electrolyzers, and thus must operate at lower current densities to achieve the same efficiency. Although the diaphragm is mostly capable of preventing product gases from crossing over and mixing, gas bubbles can enter the pores and block pathways for ionic conduction, further increasing the ohmic resistance of the cell.
In terms of energy efficiency and operating capacity, PEM electrolyzers are superior to alkaline electrolyzers. A comparison of the typical performance values for PEM and alkaline electrolyzers is given in Table 1.1. Despite the performance advantages of PEM electrolyzers, there are many costs associated with the Nafion membrane which contribute to the overall capital cost. First is the material cost of the membrane, which can account for 3% of the total cost of the electrolyzer. However, there are additional capital costs associated with the membrane beyond its material cost. In order to achieve the benefits of a zero-gap resistance, electrocatalysts must be directly impregnated onto the membrane, known as a membrane electrode assembly (MEA). The improved performance comes at the cost of durability, which is directly tied to the cost of maintenance and the overall lifetime of the electrolyzer. If any one component of the MEA fails, the entire MEA must be replaced. Studies have investigated the possibility of recycling the MEA, but the process is highly destructive and can result in significant losses in material and performance.
Table 1.1 Performance characteristics for alkaline\textsuperscript{15-17} and PEM electrolyzers\textsuperscript{18,19}.

<table>
<thead>
<tr>
<th>Electrolyte</th>
<th>Voltage Efficiency based on HHV</th>
<th>Current density</th>
<th>H\textsubscript{2} outlet pressure</th>
<th>Purity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkaline</td>
<td>25-30 % wt KOH</td>
<td>60-80%</td>
<td>0.1-0.4 A cm\textsuperscript{-2}</td>
<td>&lt; 30 bar</td>
</tr>
<tr>
<td>PEM</td>
<td>Polymer (e.g. Nafion)</td>
<td>65-80%</td>
<td>1.5-4 A cm\textsuperscript{-2}</td>
<td>10-150 bar</td>
</tr>
</tbody>
</table>

In addition to being difficult to regenerate and replace, the MEA is also highly sensitive to impurities. Even ppm level concentrations of calcium or magnesium ions can cause the membrane pores to clog and deactivate.\textsuperscript{20,21} To mitigate the risk of contamination and maintenance, upstream water purification systems are required, adding to the overall capital costs of the system. Even in the absence of impurities, the electrocatalyst and support material can corrode and deposit in the membrane, further accelerating the deactivation of the electrolyzer and reducing its effective lifetime.\textsuperscript{22} The shortfalls of membranes in PEM electrolyzers also largely apply to the diaphragm separator for alkaline electrolyzers. The diaphragm itself is inherently resistive, and its presence between the electrodes decreases the energy efficiency of the device. Over the lifetime of the diaphragm, contaminants can clog its pores and further increase the ohmic resistance. If an electrolyzer could be designed to operate without a membrane or diaphragm, many of these efficiency losses and hidden capital costs could be avoided altogether.

In this thesis, we review the design of electrolyzers which operate without the use of a membrane or separator. Historically, the first demonstrations of water electrolysis were without a membrane. Early reactor designs for industrial electrolysis date back to the end of the 19\textsuperscript{th} century, when DC power stations were first being constructed.\textsuperscript{23} It was not until around the 1920s-1940s that asbestos separators for alkaline electrolyzers were commercialized.\textsuperscript{23} Membrane technology
continued to mature throughout the 20th century, but in the 2000s membraneless electrochemical cell designs began to re-emerge in research for fuel cells24,25 and flow batteries.26,27 This research was largely motivated by the capital cost and performance constraints imposed by membrane separators. In the proposed membraneless electrolyzers, adequate separation of the fuel streams was maintained by controlling the flow properties of the electrolyte.

Hashemi, et. al extended the concept of membraneless flow cells to electrolyzers for water splitting.28 Whereas earlier demonstrations of membraneless flow cells were designed to prevent mixing of the inlet fuel streams for fuel cell applications, flow electrolyzers are designed to prevent mixing of the product gases. In the study by Hashemi, et. al, this was accomplished by using parallel plate, flow-by electrodes in a microfluidic cell.28 Gillespie, et al. first reported on the use of mesh, flow through electrodes, where product separation was achieved by pumping electrolyte through the void spacing in the mesh.29

Schematics of example membraneless flow electrolyzers are shown in Figure 1.2. An example of the flow-by design is shown in Figure 1.2a. The laminar flow profile between the electrodes exerts a force on the H2 and O2 bubbles that causes them to remain near the walls as they’re pushed downstream and into their respective collection channels. Figure 1.2b shows a schematic of a flow-through electrolyzer reported by O’Neil, et al.30 In this design, H2 and O2 bubbles are generated on the metal surfaces of the mesh wires as aqueous electrolyte continuously flows through the gap spacing, pushing the generated bubbles into their collection channels. The flow of the electrolyte both ensures continuous replenishment of the reactants while also removing product gas bubbles occupying reaction sites on the electrode surface.
Figure 1.2: Reactor designs for membraneless electrochemical cells. a) A flow-by electrolyzer design which uses solid electrodes mounted to the walls of a laminar flow channel b) A flow-through electrolyzer where porous mesh electrodes extend into the center of a flow channel. In these devices, electrolyte flows through the gap spacing in the mesh and pushes the product bubbles downstream.

A membraneless architecture can simplify the cost of assembly for an electrolyzer as well as reduce the overall capital costs. The study by O’Neil, et al. demonstrated that a membraneless flow through electrolyzer can be assembled out of as few as three parts: an anode, a cathode, and a plastic chassis to facilitate fluid flow and product collection.\textsuperscript{30} The absence of a membrane in these devices improves their overall durability and tolerance to impurities in the electrolyte, but the same design tradeoffs exist between the ohmic resistance and the purity of the product gas streams. Ideally, the electrodes should be placed as close together as possible to minimize ohmic resistance losses, but doing so also increases the likelihood of cross-over – an event defined by an electrochemically generated bubble crossing over the flow channel and entering the incorrect collection chamber. The transport mechanisms for bubble cross-over can be complicated, and involve an understanding of how bubbles detach from the electrode, how they transport in response
to electrolyte convection, and how they equilibrate with gas dissolved in solution. A better understanding of these loss mechanisms would give insight into the true cost and performance limitations of a membraneless electrolyzer. Furthermore, a key challenge for membraneless electrolyzers is to demonstrate that they can generate H$_2$ gas with purities similar to membrane and diaphragm cells.

### 1.5 Integrating membraneless electrolyzers with solar power

Up until this point, the discussion has largely assumed that the proposed electrolyzers are connected to a grid infrastructure which supplies clean energy. However, if membraneless electrolyzers are to be used as a means for solar fuels production, it is important to understand how they would be directly integrated with a solar powered system. One way is by directly coupling the electrolyzer to a PV system, as shown in Figure 1.3a. Another way is by using a photoelectrochemical (PEC) cell, shown in Figure 1.3b. A PEC is an all-in-one device which uses semiconducting electrode to both extract energy from light and catalyze the electrochemical reaction. Hydrogen can also be produced using photocatalytic suspension reactors,$^{31}$ however this area of research is largely outside the scope of this dissertation.

A key metric for comparing solar water splitting devices is the solar-to-hydrogen efficiency, that is, the efficiency by which solar power is converted to storable chemical energy. PV-electrolysis systems have the advantage of using two relatively mature technologies which can be separately optimized. For a PV-electrolysis system, the record solar to hydrogen efficiency is 30%, which was achieved for a multi-junction PV cell connected to a PEM electrolyzer.$^{32}$ For a PEC, the record solar to hydrogen efficiency is 14%.$^{33}$ Although PV-electrolyzers have so far been demonstrated to be more efficient, they have the drawback of requiring capital investment into two separate pieces of equipment. Using a membraneless electrolyzer design could help reduce the
capital costs for a PV-electrolyzer pair. A PEC also has the potential to reduce the capital costs, given that it would only require investment into one single device. PEC research is still in its early stages, however, and information regarding scalability is limited. The simplicity of a membraneless architecture can make it an ideal candidate for scaled up PEC designs, where the transport of the reactant and product species can be directly tied to the maximum achievable photocurrent.

Figure 1.3: Integrated designs for direct production of solar fuels. a) PV-electrolyzer b) Photoelectrochemical cell.

1.6 Dissertation overview

The research objective of this dissertation is to improve the understanding of mass transport processes in membraneless electrolyzers in order to engineer lower-cost devices that can operate safely at higher current densities, higher efficiencies, and with excellent durability that may allow them to compete with conventional electrolyzer designs. Chapter 2 reviews established concepts of electrochemical engineering to characterize the performance and efficiency of membraneless electrolyzers. Chapter 3 explores the use of high speed video (HSV) as a new method of characterizing the multiphase flows of electrochemically generated bubbles as they depart from
the electrodes. The findings in this chapter largely explain the physical processes taking place in a membraneless electrolyzer.

After establishing an understanding of the modeling and transport processes in membraneless electrolyzers, the second half of this thesis is devoted to improving their performance and integrating them with solar powered systems. Chapter 4 explores how the electrode design can be leveraged to optimize the efficiency of a mesh flow through electrode while maximizing the reliability of product collection. A complete membraneless electrolyzer is then integrated into an array of PV cells to directly generate H\textsubscript{2} using light energy. Chapter 5 explores the application of membraneless electrolyzers to a photoelectrochemical (PEC) cell. Concluding remarks and future directions for membraneless electrolyzers are presented in Chapter 6.

1.7 References


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CHAPTER 2

DESIGN PRINCIPLES FOR MEMBRANELESS ELECTROLYZERS

Chapter 1 outlined three important metrics used for evaluating the performance of an electrolysis system: efficiency, capital cost, and durability. Although a full technoeconomic analysis of membraneless electrolyzers is outside the scope of this dissertation, one could imagine that each of these metrics is governed by an optimizable objective function. The equations that form the basis of this objective function are highly coupled, meaning that designing an electrolyzer with the aim of improving one metric can easily cause another metric to become worse off. For example, using a more efficient electrocatalyst material may cause the efficiency of the electrolyzer to increase, but if the material is more expensive it will also increase the capital costs. Behind the overall objective function is the governing physics of the electrochemical system. Recognizing this, Chapter 2 provides an overview of the thermodynamics, kinetics, and transport phenomena that governing the performance and design considerations for membraneless electrolyzers. At the end of this chapter, these concepts are built into a framework for modeling the performance of a membraneless electrochemical cell.

2.1 Thermodynamics of the water electrolysis reaction

The efficiency of an electrolyzer is calculated by comparing the actual power consumed to the theoretical minimum power based on thermodynamics. For any electrical system, the power consumed is the product of current and voltage. The current of an electrolyzer is directly proportional to the rate of reaction, which can be calculated using Faraday’s law shown in Equation 2.1. The voltage, on the other hand, describes the change in energy state of the reactants
and products. Equation 2.2 shows that the change in Gibbs free energy at standard state can be used to calculate the cell potential $U_{\text{cell}}^0$.

$$I = r \cdot n \cdot F$$ \hspace{1cm} 2.1

$$U_{\text{cell}}^0 = -\frac{\Delta G^0}{nF}$$ \hspace{1cm} 2.2

Typically, in chemical engineering processes, quantities are expressed using moles. In Equation 2.1, the reaction rate $r$ is expressed in dimensions of moles per unit time. Similarly, the Gibbs free energy of formation, $G^0$, has dimensions of energy per mole. In electrochemistry, however, it is typically more convenient to convert from units of moles to units of charge. This is achieved by either multiplying or dividing by number of electrons participating in the overall reaction, $n$, and the Faraday number, $F$. In this way, the reaction rate can be described using the current, $I$, which has dimensions of charge per unit time. Equation 2.2 shows that the cell potential, $U_{\text{cell}}^0$, is the change in free energy per unit charge passed through the electrolyzer. In the previous chapter, we established that for the water splitting reaction, $U_{\text{cell}}^0 = -1.23 \, V$.

However, a water electrolysis system does not always operate at standard state. Under non-standard conditions, the reduction potential at each electrode must be recalculated using the Nernst Equation. Equations 2.3 and 2.4 show the non-standard reduction potentials for HER and OER, respectively:

$$U_{H_2|H^+} = U_{H_2|H^+}^0 - \frac{RT}{2F} \ln \left( \frac{P_{H_2}}{1 \text{ bar}} \right) \left( \frac{c_{H^+}}{1 \text{ M}} \right)^2$$ \hspace{1cm} 2.3

$$U_{H_2O|O_2} = U_{H_2O|O_2}^0 + \frac{RT}{2F} \ln \left( \frac{c_{H^+}}{1 \text{ M}} \right)^2 \frac{P_{O_2}}{1 \text{ bar}}$$ \hspace{1cm} 2.4
where $U_j$ is the reversible potential at non-standard conditions for each half reaction $j$, $R$ is the gas constant, $T$ is the temperature, $P_{H_2}$ is the partial pressure of H$_2$, $P_{O_2}$ is the partial pressure of O$_2$, and $c_{H^+}$ is the concentration of protons in the electrolyte. The dependence on concentration and partial pressure for each equation is based on the activity ratio of the respective half reaction. Each species activity is raised to the power of its stoichiometric coefficient. The activity of the solvent water is assumed to be 1. When the system is at standard state, $c_{H^+} = 1$ M and $P_{H_2} = P_{O_2} = 1$ bar, and therefore $U_j = U_j^0$ for both half reactions.

Operating at elevated pressures will cause the value of $U_{cell} = U_{H_2|H^+} - U_{H_2O|O_2}$ to increase. Likewise, allowing for a concentration gradient across the electrodes can also cause the value of $U_{cell}$ to increase. One should note that the value of $c_{H^+}$ in Equations 2.3 and 2.4 should be evaluated locally at each electrode. If $c_{H^+}$, and consequently the pH, is the same at both electrodes, then the Nernstian shift for $U_{H_2|H^+}$ and $U_{H_2O|O_2}$ will cancel each other. However, over long periods of operation, a concentration gradient can form if there is no significant mixing in the electrolyte.

One motivation for operating an electrolyzer in strongly acidic or strongly alkaline electrolytes is that the local change in pH across the cell will be negligible at practical current densities. This prevents additional voltage penalties described by the Nernst equation. However, when using an unbuffered pH-neutral electrolyte such as sodium sulfate, a concentration gradient will form across the cell. In a flow-through membraneless electrolysis system reported by Talabi, et al., this concentration gradient was deliberately formed as a means for simultaneous production of acid and base. At the cathode, protons were locally consumed according to Equation 1.1, causing the local pH to become more alkaline. At the anode, protons were locally generated according to Equation 1.2, causing the local pH to become more acidic. Along with the
electrochemically generated H₂ and O₂, the resulting acid and base streams were collected. Although the resultant concentration gradient caused the total voltage required for electrolysis to increase, the extra voltage penalty can be justified if the acid and base streams are able to be collected at quantities and concentrations that are practical for commercial use.

In summary, a thermodynamic analysis is necessary for determining the minimum theoretical voltage required for water electrolysis. The Nernst equation can be used to determine the value of the minimum cell voltage \( U_{\text{cell}} \) at non-standard conditions. In reality, an applied voltage that exceeds \( U_{\text{cell}} \) is required to overcome losses due to kinetics, ionic conduction, and mass transport. Being able to calculate \( U \) for each electrode half reaction is also important for understanding the relationship between the external voltage applied to the electrode and the kinetic rate of reaction. In the next section, this relationship is discussed in greater detail in the context of membraneless electrolyzer designs.

2.2 Kinetics

One reason that designing a membraneless electrolyzer is challenging is that it requires careful consideration of phenomena that occur across a wide range of length scales. The performance of an electrolyzer device, which typically occupies space on the cm to m length scale, can be determined in part by the kinetics of the reaction, which are engineered at the Å to nm scale. The kinetics of the surface reaction are often dictated by the properties of the electrode material and the contents of the surrounding electrolyte. In an acidic electrolyte, the state-of-the-art electrode materials are platinum (Pt) for HER and iridium (Ir) for OER. These materials are scarce and expensive, and research in this area focuses on developing alternative electrode materials which are low-cost, active, and stable over long periods of operation. Although improved
Electrocatalyst materials have the potential to lower the overall capital costs and increase efficiency, research in this area is largely outside the scope of this dissertation.

Nonetheless, understanding the kinetic processes at the electrode/electrolyte interface is essential for designing an electrochemical cell. In order to better understand the relationship between the electrode surface and its catalytic properties, we first draw attention to the thermodynamic driving force of the electrode half reaction: the surface overpotential, $\eta_s$. In the previous section, we detailed how to calculate the reversible potential $U$ of an electrode at thermodynamic equilibrium. An overpotential at the electrode surface is the difference between the externally applied potential of the electrode, $V$, and the reversible potential of the half reaction, shown in Equation 2.5.

$$
\eta_s = V - U \tag{2.5}
$$

It is important to note that $V$ in Equation 2.5 is not the overall voltage applied to the electrochemical cell, but rather the local thermodynamic potential of the electrode in contact with the electrolyte. When $V > U$, $\eta_s$ is positive and the driving force for the reaction favors oxidation. When $V < U$, $\eta_s$ is negative and the reaction equilibrium shifts towards reduction. Lastly, when $V = U$, $\eta_s$ is zero and no current passes through the electrode because it is at equilibrium.

The relationship between current density and surface overpotential is described by the Butler Volmer equation, which is derived for a single electron transfer reaction (Equation 2.6):\(^1\)

$$
i = i_0 \left[ \exp \left( \frac{\alpha_A F \eta_s}{RT} \right) - \exp \left( -\frac{\alpha_C F \eta_s}{RT} \right) \right] \tag{2.6}
$$

where $i$ is the current density of the electrode half reaction, $i_0$ is the exchange current density, and $\alpha_A$ and $\alpha_C$ are the apparent transfer coefficients. By convention, positive values of $i$ refer to oxidation currents whereas negative values refer to reduction currents. The exchange current density $i_0$ is a property of the electrode material, and is descriptive of the rate of the forward and
reverse reactions when they are at equilibrium with each other. The magnitude of $i_0$ correlates directly to the catalytic activity of the electrode material. When the overpotential is sufficiently large, it can be shown that the Tafel equation applies, and is reported below in Equations 2.7 and 2.8. Equation 2.7 is the Tafel equation for anodic currents at positive overpotentials, and Equation 2.8 is the Tafel equation for cathodic currents at negative overpotentials.

$$i = i_0 \exp \left[ \frac{\alpha F \eta_s}{RT} \right]$$  \hspace{1cm} 2.7

$$i = -i_0 \exp \left[ -\frac{\alpha C F \eta_s}{RT} \right]$$  \hspace{1cm} 2.8

The Tafel equation is particularly useful for describing the kinetics of the half reactions in electrolyzers, where high current densities and therefore large overpotentials are necessary to produce practical quantities of H$_2$. This relationship is also convenient because it is easy to extract kinetic rate parameters from experimental data. Equation 2.9 shows the linear form of the Tafel Equation, which is used for a so-called Tafel analysis.

$$\eta_s = \frac{RT}{\alpha F} \ln(|i|) - \frac{RT}{\alpha F} \ln(i_0)$$  \hspace{1cm} 2.9

A linear plot of $\eta_s$ vs $\ln(|i|)$ can be used to determine the values of $\alpha$ and $i_0$ for a given electrode half reaction on a specific material. The quantity $\frac{RT}{\alpha F}$ is often referred to as the Tafel slope, and $i_0$ can be calculated based on the intercept. O’Neil et. al performed a Tafel analysis on Pt-coated titanium (Ti) mesh flow through electrodes which were used in a membraneless electrolyzer, the results of which are reproduced in Table 2.1. These electrodes are similar to the electrodes used in chapters 3 and 4 of this dissertation.
Table 2.1: Tafel kinetic parameters for HER and OER on Pt mesh flow through electrodes, reproduced from O’Neil, et al.³

<table>
<thead>
<tr>
<th></th>
<th>$i_0$ / A cm$^2$</th>
<th>$\frac{RT}{\alpha F}$ / mV</th>
</tr>
</thead>
<tbody>
<tr>
<td>HER on Pt</td>
<td>6.6 E-04</td>
<td>38.8 ± 2.2</td>
</tr>
<tr>
<td>OER on Pt</td>
<td>1.8 E-07</td>
<td>133.2 ± 9.0</td>
</tr>
</tbody>
</table>

The Tafel analysis for Table 2.1 was carried out in 0.5 M sulfuric acid. $i_0$ for HER on Pt is several orders of magnitude larger than $i_0$ for OER. Although Pt is the state-of-the-art electrode material for HER, it is not very efficient as an OER catalyst. In many cases, at least 500 mV of overpotential at the anode are required before a sufficient onset of current density can be observed. Thus, even though a minimum voltage of 1.23 V is thermodynamically predicted for electrolysis, practical current densities for a pair of Pt electrodes in 0.5 M H$_2$SO$_4$ are not realized until a total voltage of at least 1.8 V is applied.

An optimized combination of electrode materials can decrease the surface overpotential to improve the efficiency of the electrolyzer. For the experiments presented in this dissertation, Ti coated Pt electrodes were used because of their stability in 0.5 M sulfuric acid at both reducing and oxidizing potentials. While the catalytic activity of Pt for HER in acid is excellent, its activity for OER in acid is at least tolerable for the purposes of characterizing new electrolyzer designs. A comprehensive study by McCrory et al benchmarks the catalytic activity of several electrode materials in both acidic and alkaline electrolytes.⁴ Operating in alkaline conditions has the advantage of offering a wider selection of catalyst materials. Nickel, for example, corrodes in acid but in an alkaline electrolyte it is an active and stable anode material for OER. A drawback of alkaline electrolytes, however, is their lower conductivity relative to acidic electrolytes. This lower
conductivity increases the ohmic resistance of the electrolyzer, potentially offsetting any advantages from decreasing the activation overpotential.

Ohmic resistances can also have a strong impact on the overall efficiency of an electrolyzer. Conceptually, it is useful to think of the relationship between current density and overpotential using a charge transfer resistance, $R_{CT}$. At large overpotentials, the charge transfer resistance can be calculated by evaluating the derivative of $\eta_s$ from the Tafel equation at the average current density, shown in Equation 2.10.

$$R_{CT} = \frac{\partial \eta_s}{\partial i} \bigg|_{i = i_{avg}} = \frac{RT}{aF|i_{avg}|}$$  \hspace{1cm} 2.10

At large current densities, the charge transfer resistance becomes negligible, and the relationship between current and voltage is no longer described by the Tafel equation. Rather, the performance of the electrolyzer is limited by the conduction of ions in the electrolyte. Ionic conduction is governed by Ohm’s law, which will be discussed in greater detail in the next section.

### 2.3 Ohmic Resistance

When an electric field is applied across an electrolyte, the ions will flow in response. This flow of ions is necessary to complete the circuit of an electrochemical cell. The magnitude of ionic flow, or current, depends on the strength of the electric field and the ohmic resistance of the electrolyte. This relationship is widely known as Ohm’s law, and the differential form is shown in Equation 2.11:

$$i = \kappa \nabla \phi$$  \hspace{1cm} 2.11

where $\phi(x, y, z)$ is the local potential of the electrolyte and $\kappa$ is the conductivity. If the potential field across an electrolyte domain is known, the integrated form of Ohm’s law can be solved for, shown in Equation 2.12.$^1$
\[ \Delta V = IR_\Omega \]

where \( \Delta V \) is the voltage drop between electrodes, \( I \) is the total current, and \( R_\Omega \) is the overall ohmic resistance of the electrolyte. The ohmic resistance is a function of the electrolyte conductivity \( \kappa \), and the geometry of the electrodes. For a pair of planar electrodes enclosed by insulating walls, \( R_\Omega \) can be solved for directly using the electrode separation distance, \( L \), and the nominal area of the electrodes \( A_c \). This relationship is shown in Equation 2.13.

\[ R_\Omega = \frac{L}{A_c \kappa} \]

The cross-sectional area of the electrolyte channel is assumed to be equal to the area of the electrodes. In some derivations of the overall ohmic resistance, numerator of Equation 2.13 may also include a cell constant, \( \beta_r \), to account for geometries where the electrodes are not parallel, have non-planar shapes, or have unequal areas. In these cases, \( \beta_r \) serves as a dimensionless shape factor with a magnitude typically on the order of 1. For a pair of large, parallel electrodes, \( \beta_r = 1 \). When the electrodes are not parallel, the value of \( \beta_r \) will depend on how \( L \) is defined. If available, an analytical solution can calculate an exact value of \( \beta_r \), and is typically a dimensionless ratio of important length scales in the reactor design. \( \beta_r \) can also be obtained from simulations, which will be explained in greater detail in section 2.5 of this chapter. Newman, for example, has shown both analytically and using simulations that \( \beta_r = \frac{\pi}{4} \) for the ohmic resistance between a rotating disc electrode and a faraway reference electrode.

Ohm’s law reveals the trade-off between productivity and efficiency for an electrolyzer. For example, a water electrolyzer produces the most \( \text{H}_2 \) when the current is maximized. However, Ohm’s law clearly indicates that increasing the current will also increase the voltage penalty.
required to conduct ions between the electrodes. Therefore, to minimize this efficiency loss, the ohmic resistance should be as low as possible when designing an electrolyzer.

One way to decrease the ohmic resistance is to increase the conductivity, \( \kappa \), of the electrolyte. Typically, \( \kappa \) can be calculated using Equation 2.14, and depends directly on the concentration \( c_j \), mobility \( u_j \), and valence charge \( z_j \) of each ion \( j \) present in the electrolyte.\(^1\) For the majority of demonstrations of membraneless electrolyzers in this dissertation, 0.5 M H\(_2\)SO\(_4\) is used and has a conductivity of 0.226 S/cm.\(^7\) The conductivity of an electrolyte typically scales linearly with its ion concentration, but the relationship becomes non-linear at high concentrations. Using higher acid concentrations also introduces hazards due to the general corrosiveness of the electrolyte. Concentrated acids can be dangerous to handle and may require additional safety protocols if used on an industrial scale. They may also require the electrolyzer to be constructed out of more expensive, corrosion resistant parts.

\[
\kappa = F^2 \sum_j z_j^2 u_j c_j \tag{2.14}
\]

Another way to minimize the ohmic resistance is to decrease the separation distance \( L \) between the electrodes. This is largely the motivation for using an ion conducting solid electrolyte such as a Nafion membrane. Nafion membranes used in PEM electrolyzers typically have thicknesses between 100 and 200 \( \mu \)m.\(^2\) Demonstrations of microfluidic membraneless electrolyzers have also achieved electrode separation distances as narrow as 150 \( \mu \)m, but the lack of a physical barrier presents challenges with managing the product gas flows of H\(_2\) and O\(_2\). Related to this complicated interplay of ionic transport and gas phase transport is the area of the electrodes themselves. Increasing the electrode area can also decrease the ohmic resistance of the electrolyzer, but this also raises questions on how to precisely control fluid flow and maintain effective separation of anode and cathode products over larger length scales. For example, the
electrode area for a parallel plate design such as in Figure 1.2a can be increased simply by making the electrodes longer. However, as bubbles detach and migrate away from the channel wall, their probability of crossover will likely increase with longer travel distances and residence times spent in the channel. A major focus of this dissertation is to better understand the physical mechanisms of product gas transport in order to improve the product purity of collected gases and demonstrate safe operation at close electrode separation distances.

The discussion in this section has focused mostly on the ohmic resistance as it relates to the electrolyte of a membraneless electrolyzer. However, the contacts between the electrolyzer and the external circuit can also contribute to the overall resistance. Maintaining high quality electrical connections is important for ensuring efficient operation throughout the life of the electrolyzer. Degradation and corrosion of the contacts over time can increase the overall ohmic resistance. In a laboratory setting, if a higher than normal ohmic resistance is measured, a good first step is to check that the external contacts are connected properly and free of corroded surfaces. Sometimes, a source of ohmic resistance is the electrode material itself. Typically, this is not considered as metallic electrodes have a conductivity that is several orders of magnitude higher than that of the electrolyte. However, Equation 2.13 also applies to the calculation of electronic conduction in an electrode material. At large electrode length scales, the ohmic drop within the electrode may be relevant to the overall resistance.

Lastly, the ohmic resistance of a membraneless electrolyzer can also increase due to the presence of the electrochemically generated gas bubbles. Gases behave as insulators, and when they occupy area on an electrode surface, the effective cross-sectional area available for ionic conduction decreases. In the bulk electrolyte, the presence of bubbles can be modeled using an effective conductivity of the electrolyte. Equation 2.15 is known as the Bruggeman correlation,\textsuperscript{6,8}
and is an empirical correlation between the effective conductivity of the electrolyte \( \kappa_{eff} \) and the void fraction of gas bubbles in the liquid electrolyte, \( \epsilon \).

\[
\kappa_{eff} = \kappa(1 - \epsilon)^{1.5}
\]

As the void fraction of gas bubbles increases, the effective conductivity decreases to account for the decrease in accessible volume for ions to conduct through. In a membraneless electrolyzer, using flowing electrolyte can help remove gas bubbles and thereby decrease the bubble void fraction, resulting in decreased overall ohmic resistance of the device. The next section reviews transport processes which take place in electrochemical cells.

2.4 Transport

In the previous section, we addressed the efficiency losses due to the conduction of ions between electrodes. Although ionic conduction is necessary for a complete circuit in an electrochemical cell, it is not the only mode of transport in an electrolyte. Equation 2.16 shows the Nernst-Planck Equation for the total molar flux of all species present in the electrolyte:\(^1\)

\[
N_i = -z_i u_i F c_i \nabla \phi - D_i \nabla c_i + c_i \mathbf{v}
\]

For a given species \( i \), \( z_i \) is the valance charge, \( u_i \) is the ionic mobility, \( c_i(x, y, z) \) is the local concentration, and \( D_i \) is the diffusion coefficient. \( \phi(x, y, z) \) is the local potential of the electrolyte and \( \mathbf{v} \) is the velocity vector. The first term in Equation 2.16 is simply Ohm’s law, describing how ions move in response to the local potential gradient, \( \nabla \phi \). This term is often referred to as the migration current. The second term refers to the diffusion of species throughout the electrolyte in the presence of a concentration gradient. The last term on the right is the convection term. The total ionic current can be calculated by summing up the fluxes of all species, as shown in Equation 2.17:
\[ i = F \sum_i z_i N_i \tag{2.17} \]

An important constraint in electrochemical systems is the electroneutrality condition, which states that the total sum of all charges at any location in the electrolyte must be zero (Equation 2.18). When Equation 2.16 is summed up using Equation 2.17, it can be shown that the convection term for ionic currents can be neglected due to electroneutrality. The resulting combined equation for the ionic current is Equation 2.19.

\[ \sum_i z_i c_i = 0 \tag{2.18} \]

\[ i = -F \sum_j \left( z_j^2 u_j F c_j \nabla \phi + z_j D_j \nabla c_j \right) \tag{2.19} \]

Concentration gradients typically form as the operating current of the electrolyzer approaches the mass transfer limited current, \( i_{\text{lim}} \). An electrochemical device reaches \( i_{\text{lim}} \) when the limiting process is no longer the reaction kinetics, but rather the mass transport in the electrolyte. When a diffusing species carries an ionic charge, Equation 2.19 shows that it can be affected by both the electric field and the concentration gradient. Being able to solve Equation 2.19 in its entirety can be complicated and difficult to achieve convergence. Good physical intuition of the system can help guide decisions on how to simplify the model and obtain a solution.

For an initial guess of the mass transport limited current, the electric field can be temporarily neglected in the case where there is excess supporting electrolyte. In such a case, one must solve for the diffusive flux of the limiting reagent across a diffusion boundary layer. For diffusion to a planar electrode across a boundary layer of thickness \( \delta \), the concentration profile across the boundary layer is linear and the limiting current can be obtained from Fick’s law (Equation 2.20):
\[ i_{\text{lim}} = nFD\frac{c_{i,\text{bulk}}}{\delta} \]  \hspace{1cm} (2.20)

where \( c_{i,\text{bulk}} \) is the bulk concentration of reagent \( i \) and \( n \) is the number of electrons participating in the electrode half reaction. The best way to determine the thickness and profile of \( \delta \) is by first determining the dominant mode of transport in the electrolyte. This can be calculated using the Péclet number (\( Pe \)), defined in Equation 2.21. \( Pe \) is the dimensionless ratio of the convective transport rate to the diffusive transport rate, where \( U \) is the characteristic velocity, \( L \) is the characteristic length, and \( D \) is the diffusion coefficient of the limiting reagent.

\[ Pe = \frac{UL}{D} \]  \hspace{1cm} (2.21)

When the value of \( Pe \gg 1 \), convection is the dominant mode of transport. In such a case, the diffusion boundary layer can be solved based on the developed flow profile near the electrode. Several empirical correlations for \( \delta \) exist in the literature, where \( \delta \) is a function of the Reynolds number \( Re \) and the Schmidt number \( Sc \).\(^1\)

As \( Pe \to 0 \), bulk diffusion becomes the dominant mode of transport. In such a case, a diffusion cloud will grow at the electrode surface, and \( \delta \) will increase as a function of time, such as in the Cottrell equation.\(^5\) After long periods of operation, a steady state diffusion boundary layer can also be achieved. However, a purely diffusion limited current is rarely expected to arise in a membraneless electrolyzer, where the product gases are separated using bulk convection. Even in cases when there is no pumped electrolyte, the free convection of the departing bubbles can be expected to have a large impact on the diffusion boundary layer thickness. Free convection is particularly relevant to the studies presented in Chapters 3 and 4.

Regardless of the thickness of the diffusion boundary layer, the bulk reactant concentration in Equation 2.20 is also a driving force of the limiting current density. The relevant concentration
is determined by the limiting reagent of an electrode half reaction. A membraneless electrolyzer operating in 0.5 M H$_2$SO$_4$ will be limited by the concentration of protons at the cathode and by the concentration of the solvent water at the anode. In reality, these concentrations are quite large, and therefore neither electrode half reaction is expected to be limited by mass transport of the reactant species at practical current densities. However, for electrolysis reactions such as CO$_2$ reduction in an aqueous electrolyte, the bulk concentration of CO$_2$ is limited by its solubility. In these cases, understanding and modeling the diffusion boundary layer thickness may be more relevant to the overall performance of the device. This topic of discussion is explored in greater detail in Chapter 5, where forced convection is used to enhance the rate of mass transport in a device with dilute reactant concentrations.

It is important to note that although the convection term does not appear explicitly in Equation 2.19, the bulk fluid velocity is an important component for calculating the diffusion boundary layer. Equation 2.19 also assumes that the concentration of ions is dilute relative to the concentration of the supporting electrolyte. Thus, the ionic current between the electrodes is not expected to directly influence the macroscopic flow properties of the electrolyte, and in many cases, the two can be modeled independently.

To model the transport of non-charged species, such as the solvent water or the electrochemically generated gas bubbles, the Navier-Stokes equations are used. Modeling multiphase flows can be particularly difficult, and the computational cost depends on the total number of bubbles present, the size distribution of the bubbles, and the size of the bubbles relative to the critical dimensions of the electrolyzer cell. Understanding these transport processes as they relate to the collection of product gases in membraneless electrolysers is a topic of ongoing research. In Chapters 3 and 4, both quantitative and qualitative observations are made to
characterize the relationship between the operating parameters of the electrochemical cell and the transport of the product gases.

2.5 Bubble dynamics in an electrochemical system

Although a fully defined model of the gas bubble transport is outside the scope of this dissertation, a qualitative understanding of the theory is still relevant for guiding the design of membraneless electrolyzers. Gas evolving electrodes are ubiquitous in electrochemical systems, and a significant body of research has been devoted to their study. The force balance on a bubble governs the critical diameter at which it departs from the electrode surface. Upon departure, the diameter of the bubble can strongly affect its trajectory in the bulk electrolyte. For membraneless electrolyzers, this flow behavior can determine the purity of the collected H₂ and O₂ gases. High purity gas is desirable because it reduces separation cost. It is also important to have highly pure gases to avoid creating an explosion hazard in the downstream gas collection reservoirs. For H₂ gas, the upper flammability limit is 75% in air and 94% in pure O₂. The lower flammability limit for H₂ is 4%.

The schematics of the membraneless electrolyzers shown in Figure 1.2a and Figure 1.2b illustrate an idealized reactor where pure streams of H₂ and O₂ gases can be collected. In these designs, one can imagine the momentum of the incoming electrolyte being used to push and direct the bubbles into their respective collection chambers. In reality, the product gases that are collected are not 100% pure, and the mechanisms by which gas bubbles cross over into the opposing collection chamber are not well understood. Experimentally, cross-over can be determined by measuring the amount of hydrogen collected in each chamber. In this dissertation, the percentage cross-over is generally defined as the amount H₂ collected in the anode chamber relative to the total amount of H₂ collected in both chambers.
When H$_2$ is evolved during electrolysis, it can either enter the gas phase as a bubble or remain dissolved in the electrolyte. In order to enter the gas phase, the bubble must first nucleate on the electrode surface. The size of the bubble is largely determined by the Young Laplace equation for a spherical bubble, shown in Equation 2.22: $\Delta P = \frac{2\gamma}{R}$

where $\Delta P$ is the pressure difference between the gas and liquid phases, $\gamma$ is the surface tension, and $R$ is the bubble radius. The smaller the bubble radius, the larger the relative pressure inside the gas bubble. According to Henry’s law, the pressure of H$_2$ is directly proportional to the concentration of H$_2$ dissolved in the electrolyte. Therefore, the bubble diameter on an electrode surface is likely to be governed by the degree of H$_2$ super saturation in the surrounding electrolyte. Figure 2.1 shows a simple schematic of the forces acting on a H$_2$ bubble growing on an electrode surface.

![Diagram of H$_2$ bubble on electrode surface]
In this diagram, the buoyancy force $F_B$ lifts the bubble upwards while the surface tension force $F_S$ pulls to keep the bubble attached to the electrode. Equation 2.23 shows the buoyancy force is a function of the bubble radius:

$$F_B = \frac{4}{3} \pi R^3 \Delta \rho g$$  \hspace{1cm} (2.23)$$

where $\Delta \rho$ is the difference in density between the gas and the electrolyte and $g$ is the gravitational constant. As the bubble radius increases, the buoyancy force eventually exceeds the surface tension force and the bubble departs from the electrode surface. Also shown in Figure 2.1 is the contact angle between the bubble and the electrode surface, $\theta$. The contact angle can be used to calculate the surface tension force, and is related to the hydrophilicity of the electrode surface. When $\theta < 90^\circ$, the electrode is hydrophilic and prefers contact with electrolyte. When $\theta > 90^\circ$, the surface is hydrophobic and prefers contact with the gas bubble. The contact angle is a property of the electrode material and the composition of the gas and surrounding electrolyte.

The force balance shown in Figure 2.1 is simplified, and there are in reality many other forces which act on the bubble. In the presence of forced convection, the drag forces and lift forces acting on the bubble can also affect the departure diameter. A more complete force balance diagram on a bubble is offered by Taqieddin, et al.\textsuperscript{20} Although detailed models exist for a single bubble force balance and departure diameter, gas evolving electrodes typically produce a wide distribution of bubble sizes, many of which are significantly smaller than what would be predicted based on the force balance. The departure of a neighboring bubble induces turbulence, causing the surrounding bubbles to depart even if they have diameters smaller than the predicted departure diameter.\textsuperscript{20} The surface tension forces surrounding a bubble are also not constant, and can change as a function of the dissolved gas concentration.\textsuperscript{20}
Once a bubble detaches from the electrode surface, it is exposed to lift and drag forces which cause it to permeate throughout the electrolyte.\textsuperscript{21} Hashemi, et al. proposed that a balance between lift forces and wall forces, known as the Segré-Silberberg effect, can be engineered to predict and control where the bubbles migrate in a membraneless flow channel.\textsuperscript{22} In the context of a membraneless electrolyzer, these forces are still not fully understood and it is challenging to relate stochastic bubble phenomena with the overall performance of the device. As a part of the work of this dissertation, observations of the bubble dynamics in a membraneless electrolyzer are related to the predicted current distributions typically solved for in electrochemical systems. In the next section, these current distributions and the governing equations behind them are explored in greater detail.

### 2.6 Approach to modeling the ionic current in an electrolyzer

Having established the fundamentals of thermodynamics, kinetics, and mass transport for electrochemical cells, this last section of the chapter outlines how to connect these concepts into a computational model. The ultimate goal of this modeling framework is to be able to model or predict the current distribution on electrodes (for cases where there is non-uniform current distribution) and the overall current-voltage curve for an electrolysis device of known geometry. Knowing the current-voltage curve for the electrolysis device, it is easy to compute the electrolyzer efficiency by Equation 1.5. A visual framework for the approach taken in this dissertation is shown in Figure 2.2, which is largely inspired by the approach given in Chapter 6 of *Electrochemistry and Electrochemical Engineering*.\textsuperscript{5} The starting point for this analysis is the Nernst-Planck Equation (Equation 2.16). As mentioned previously, solving Equation 2.16 without simplification can be tedious, computationally expensive, and difficult to achieve convergence. Rather, simplifying the governing physics and judiciously neglecting terms in Equation 2.16 can yield
tractable problems with easily interpretable results. At each decision node in Figure 2.2, there is a calculable quantity to be evaluated. If simplifications can be made, each branch of the decision tree will lead to a blue box describing how to model the electrolyte phase and a green box describing what boundary conditions to impose.

Figure 2.2: Framework for the simulation of the current density in a membraneless electrochemical cell.
The first decision node in Figure 2.2 checks to see if concentration gradients can be ignored. This is determined by calculating the dimensionless quantity $i/i_{lim}$, where $i$ is the designed operating current density. This dimensionless grouping can be thought of as a Damkohler number, which compares the kinetically limited rate of reaction to the mass transfer limited rate of reaction. In the previous section, we reviewed a simplified approach for calculating $i_{lim}$ assuming that electrical migration can be neglected. For common acidic and alkaline electrolytes used for water electrolysis, $i_{lim}$ is often times expected to be very large therefore can be assumed that $i/i_{lim} \to 0$. A sample calculation of the limiting current density in an electrolyzer in 0.5 M H$_2$SO$_4$ is given in Chapter 3, Appendix A, section 3.5.2.

2.6.1 Modeling the electric field

As long as the operating current is significantly below the mass transfer limited current, then the electrolyte can be assumed to be well mixed and we are able to neglect the effect of concentration gradients. The Nernst-Planck equation then simplifies to differential form of Ohm’s law (Equation 2.11). It can be shown that by applying a conservation of charge and electroneutrality, the potential field must obey Laplace’s equation, Equation 2.24:

$$\nabla^2 \phi = 0$$

2.24

By imposing Equation 2.24 for $\phi$ everywhere in the electrolyte domain, the potential field ($\phi(x, y, z)$) may be solved. Software packages such as COMSOL can be useful for drawing out the domain and boundaries for a model electrolyzer in a 2D or 3D simulation. Once the electrolyte domain and conductivity are defined, regions along the boundary walls must be specified as either an electrode surface or an insulating wall, and the appropriate boundary conditions must be imposed. At an insulating wall, the potential gradient is set to zero, as shown in Equation 2.25. In
the context of Ohm’s law, Equation 2.25 specifies that there can be no ionic current flow into an insulating surface. The vector $\mathbf{n}$ refers to the normal direction of the insulating surface.

**Insulating walls:** $\mathbf{n} \cdot \nabla \phi = 0 \quad 2.25$

The boundary conditions at the cathode and anode depend on the desired level of complexity for the model. The simplest set of boundary conditions ignore the kinetic processes at the electrodes and model only the ohmic resistance in the electrolyte. This model is often times referred to as the primary current distribution. For the primary current distribution, the potential of the electrolyte at the boundary is equal to the externally applied potential to the electrode, shown in Equations 2.26 and 2.27:

Cathode boundary: $\phi(x_{cath}, y_{cath}, z_{cath}) = V_{cath} \quad 2.26$

Anode boundary: $\phi(x_{anode}, y_{anode}, z_{anode}) = V_{anode} \quad 2.27$

where $x_{cath}, y_{cath}, \text{ and } z_{cath}$ are the coordinates of the cathode and $x_{anode}, y_{anode}, \text{ and } z_{anode}$ are the coordinates of the anode. $V_{cath}$ and $V_{anode}$ are the externally applied potentials of the cathode and anode electrodes, respectively. Solving the primary current distribution is computationally inexpensive for most geometries and is convenient for calculating the total ohmic resistance of an electrolyzer. For a given electrolyte conductivity $\kappa$ and total voltage $\Delta V = V_{anode} - V_{cath}$, a simulation of the primary current distribution can calculate the total current expected to pass through the cell and the ohmic resistance between the electrodes.

A simulation of the primary current also serves as a limiting case estimate for the maximum degree of nonuniformity of the local current density across an electrode surface. It is well known in electrical circuits that the current will follow the path of least resistance. This is also true for ionic conduction in the electrolyte. Depending on the geometry of the electrode, non-uniform current density profiles can arise if some regions of the electrode surface are more easily accessible.
to conducting ions than others. The primary current distribution is usually expressed as a non-
dimensional quantity, where the local current density is divided by the average current density
along the electrode. For a given cell geometry, the current distribution is the same regardless of
the chosen conductivity of the electrolyte or total voltage applied at the boundary conditions. The
shape of the primary current distribution depends only on the geometry of the electrodes.

When the kinetics are accounted for at the electrode boundaries, the ionic current passing
between the electrodes is said to obey the secondary current distribution. Any valid kinetic model
can be applied at the electrode boundary, including the Butler-Volmer equation and the Tafel
equation. Equation 2.28 shows how the anodic Tafel equation, originally shown in Equation 2.7,
can be applied to an anode boundary. In the form presented in Equation 2.28, the Tafel equation
is now coupled to the surrounding potential field. Notice that the definition of the overpotential $\eta_s$
is modified in Equation 2.29 and now contains a term for the electrolyte potential adjacent to the
electrode surface.

$$ \mathbf{n} \cdot \kappa \nabla \phi = i_0 \exp \left[ \frac{\alpha_a F}{RT} \eta_s \right] $$  \hspace{1cm} 2.28

$$ \eta_s = V_{\text{anode}} - E - \phi(x_{\text{anode}}, y_{\text{anode}}, z_{\text{anode}}) $$  \hspace{1cm} 2.29

Using this definition of the overpotential, the independent parameter of interest, $\phi$, is
implicit in Equation 2.28. Solving for the secondary current distribution is more computationally
demanding than the primary current distribution, and in some cases can be more difficult to achieve
convergence. The secondary current distribution is generally more uniform than the primary
current distribution because it accounts for the charge transfer resistance. To determine whether or
not the charge transfer resistance is relevant to the current distribution at the electrode surface, the
Wagner number, $Wa$, is calculated.
\( Wa \) is the ratio of the charge transfer resistance to the ohmic resistance in the vicinity of the electrode. The charge transfer resistance is defined for a system with Tafel kinetics using Equation 2.10. In the context of the Wagner number, the ohmic resistance is defined using Equation 2.30:

\[
R_\Omega' = \frac{l}{\kappa} \tag{2.30}
\]

where \( l \) is the characteristic length scale. Equation 2.30 is slightly different from the total ohmic resistance given by Equation 2.13 in that the relevant length scale in the vicinity of the electrode being modeled is often times not the electrode separation distance, \( L \). Rather, it is more likely to be related to the length of the electrode itself. Equation 2.30 does not include the electrode cross sectional area in its calculation because it is built into the variable \( R_\Omega' \). Dimensionally, \( R_\Omega' \) defined in Equation 2.30 is consistent with \( R_{CT} \) defined in Equation 2.10. Using these definitions, \( Wa \) is calculated using Equation 2.31 for an electrode using Tafel kinetics:

\[
Wa = \frac{R_{CT}}{R_\Omega'} = \frac{RT\kappa}{\alpha F |i_{avg}| l} \tag{2.31}
\]

As the value of \( Wa \) approaches zero, the ohmic resistance dominates and the current distribution on the electrode surface follows the primary current distribution. This is most likely to happen at large average current densities and large electrode sizes. Thus, the primary current distribution is typically the most relevant current distribution profile for a scaled up device. As the value of \( Wa \) approaches 1, the current distribution obeys the secondary current distribution. Depending on the choice of length scale, the current distribution may be nearly uniform as the value of \( Wa \) reaches or exceeds 1.

Although the primary current is more likely to describe the current distribution profile for scaled up devices when \( Wa \ll 1 \), using the boundary conditions for the secondary current can still
calculate the same result. The reverse is not true, and the primary current does not apply as $Wa \to 1$. Additionally, the kinetic boundary conditions must be used in order to simulate IV performance curves of the electrolyzer. As the voltage becomes sufficiently large, however, the mass transport limiting current will be approached and the concentration gradient in the electrolyzer must be modeled.

2.6.2 Modeling the concentration gradient

As $i/i_{lim} \to 1$, concentration gradients begin to form in the electrolyzer and the diffusion $(\nabla c)$ term in the Nernst Planck equation can no longer be neglected. Instead, the Nernst Planck equation can be simplified for conditions where the effects of the electric field on species transport can be neglected. The electric field must be present in the electrolyte such that the ions can conduct between the electrodes, but in many cases, this migration current is carried by spectator ions which otherwise do not participate in the electrochemical reaction. In order to determine if the electric field is relevant to the mass transport limited current, the transference number for the limiting reagent, $t_i$, must be calculated as shown in Equation 2.32:

$$t_i = \frac{z_i^2 u_i c_i}{\sum_k z_k^2 u_k c_k}$$  \hspace{1cm} 2.32

Conceptually, the transference number refers to the fraction of the total ionic current which is carried by a given species. It is related to the calculation of the conductivity of the electrolyte, shown in Equation 2.14. All parameters in the numerator in Equation 2.32 refer to the limiting reagent, designated as species $i$, whereas the denominator is a summation over all ionic species $k$. Calculation of the transference number requires a knowledge of all the ionic species present in the electrolyte, their bulk concentrations, and conductivities. As the value of $t_i \to 0$, a minimal fraction of the ionic current between the electrodes is carried by the limiting reagent, and it can be
assumed that species transport of the limiting reagent by migration can be ignored. As the value of \( t_i \to 1 \), the limiting reagent is significantly affected by the electric field, and the migration current cannot be neglected. When the migration current is significant, further simplifications cannot be applied to the Nernst-Planck equation, and both the concentration gradient and the potential field must be modeled. For some special cases, such as for a binary electrolyte, it can be shown that the electric field can also be ignored if corrective factors are applied to the diffusion coefficient.\(^1\)

If the electric field is neglected, it can be shown that the concentration gradient can be solved for using the equation of convective diffusion, Equation 2.33.\(^1\) At the electrode boundary, the current can be solved for based on the diffusive flux of the limiting reagent, shown in Equation 2.34.

\[
\frac{\partial c_i}{\partial t} + \mathbf{v} \cdot \nabla c_i = D_i \nabla^2 c_i \tag{2.33}
\]

\[
i = nF D_i \nabla c_i \tag{2.34}
\]

In the previous section, the potential field was solved by applying the appropriate boundary conditions at the cathode and anode. For solving concentration fields using the equation of convective diffusion, however, the analysis and subsequent boundary conditions are typically only applied in the region near the electrode surface. In the bulk electrolyte, \( c_i \) is determined by its bulk concentration, \( c_{i,\text{bulk}} \). At the limiting current density, the concentration \( c_i \) at the electrode surface is assumed to be zero. As previously discussed, the governing equations of the surrounding concentration profile is largely determined by calculating \( Pe \), which can help determine if the transport is predominantly diffusion or convection. Once \( Pe \) is known, the concentration field can largely be solved for using the strategies given in section 2.4 of this chapter.
When $Pe >> 1$, convection is the dominant mode of transport, and the concentration gradient can be described using a boundary layer such as in Equation 2.20. When $Pe << 1$, convection can be ignored, and Equation 2.33 is simplified to Equation 2.35:

$$\frac{\partial c_i}{\partial t} = D_i \nabla^2 c_i$$  \hspace{1cm} 2.35

In the case of the limiting current where $c_i$ at the electrode surface is fixed to zero, Equation 2.35 can be used to derive the Cottrell equation. The Cottrell equation is time dependent, and describes a limiting current which decreases over time. As previously discussed, a purely diffusion limited current is less likely to exist in membraneless electrolyzers, where convective transport is dominant. Diffusion limited currents such as in Equation 2.35 may be relevant for the application of downstream sensor electrodes to detect dissolved gases. However, this discussion is largely outside the scope of this dissertation. Rather, the continuum level models discussed in this chapter are applied to describe the overall performance of the device.

In summary, the relationship between thermodynamics, kinetics, and transport can be highly coupled in any electrochemical system. This chapter has reviewed each of these concepts as they relate to the study of membraneless electrolyzers. For a number of limiting cases, Figure 2.2 provides a general approach to determine the overall current in an electrolysis cell. In the next chapter, the predicted current distribution is compared to high speed images of the bubbles departing from a flow through electrode in a membraneless electrolyzers. Bubble nucleation and growth is explored in greater detail in Chapter 4. Lastly, Chapter 5 explores the mass transport limited current density in a membraneless photoelectrochemical cell.

2.7 References


11. Vogt, H. On the supersaturation of gas in the concentration boundary layer of gas evolving


CHAPTER 3
HIGH SPEED VIDEO INVESTIGATION OF BUBBLE DYNAMICS AND CURRENT DENSITY DISTRIBUTIONS IN MEMBRANELESS ELECTROLYZERS

Gas evolving electrodes are found in many electrochemical systems, such as water electrolysis, where the dynamics of multiphase flow at these electrodes can strongly impact cell performance. This is especially true for membraneless water electrolyzers that have emerged in recent years as a promising approach to low cost hydrogen production. However, experimental and modeling efforts to characterize the multiphase dynamics in these systems can be non-trivial due to the complexity of the coupled chemistry and physics that underlie their operation. In this chapter, we utilize a high speed video (HSV) camera as a non-invasive analytical tool to better understand bubble dynamics in membraneless electrolyzers and to better quantify the void fraction of gas bubbles in the region directly downstream of the mesh flow-through electrodes. By detecting and quantifying the void fraction of bubbles immediately downstream of an operating electrode, the local current density distribution can be determined along the length of the electrode. This HSV-measured void fraction distribution is in good agreement with the modeled primary current distribution. This study also highlights the ability to utilize in situ HSV analysis to monitor gas evolution efficiency yields and bubble size distributions under varying operating conditions.
3.1 Introduction

Water electrolysis is a promising route for carbon-free hydrogen (H₂) production, but the capital costs of conventional electrolyzer designs are expected to limit their market penetration.¹,² Membraneless electrolyzers have been proposed as simplified, low-cost alternatives,²–¹⁴ but the absence of a membrane or porous diaphragm between the electrodes requires that careful consideration be given to how the product gases are safely and efficiently collected. Forced convection of liquid electrolyte is often used to impart separation of product gases,¹¹–¹³ although a
passive design based on buoyancy induced product separation has also been demonstrated. Questions still remain on the effectiveness of gas collection in these systems, especially as devices are scaled up and a wider range of operating conditions are explored.

Gas evolving electrodes are ubiquitous in electrochemistry, with common examples including PEM and alkaline electrolyzers, electrowinning cells, the chlor alkali process, electrodialysis cells, photoelectrochemical cells, the Hall-Heroult process, and as side reactions in many batteries. Being able to directly and continuously monitor gas evolution behavior during operation through in situ imaging is of particular interest for the study of membraneless electrolyzers, where insight into the downstream transport and collection of gas bubbles is of particular interest. Membraneless electrolyzers are particularly amenable to such in situ studies since their simple architecture and void space between electrodes allows for direct optical imaging through windows placed on the side of modified electrolysis cells. Observations of bubble dynamics in membraneless electrolyzers have been used in several previous studies to qualitatively characterize device performance. In their demonstration of a microfluidic membraneless electrolyzer, Hashemi et al. used in situ imaging to visualize gaseous product separation through the Segre-Silberberg effect, whereby gas bubbles remain pinned close to the channel walls in the presence of forced convection and laminar fluid flow. In a bench scale demonstration, O’Neil et al. used in-situ imaging to observe non-uniform bubbling along the length of angled mesh electrodes. Gillespie et al. visualized a phenomena they referred to as void fracture, whereby evolved bubbles merge to form a gaseous void that bridges the inter-electrode gap and greatly reduces electrolyzer performance. In all of these studies, in situ imaging of membraneless electrolyzers has proven to be very useful for tracking cross-over events and observing bubble dynamics in flowing electrolytes, but has been largely limited to qualitative analyses.
In situ imaging has also been used to analyze bubble dynamics in electrochemical systems beyond membraneless electrolyzers. Many qualitative imaging studies have characterized the formation and detachment of bubbles or droplets from the electrode surface.\textsuperscript{18–22} Quantitative image analysis has been used to investigate the contact angles,\textsuperscript{23} bubble size statistics,\textsuperscript{24–26} and velocity profiles\textsuperscript{26–28} of electrogenerated bubbles. Modelling studies generally acknowledge that the distribution of the local density of bubbles, also known as void fraction, strongly affects the local current density on the electrode.\textsuperscript{27,29–34} However, none of these studies use imaging methods to directly quantify the local void fraction. Riegel et al. calculated the local void fraction indirectly based on local ohmic resistance measurements and a relationship derived by Maxwell.\textsuperscript{35} Jayaprakash et al. reported on a method of determining local void fraction from image analysis, which was used to study cavitation pressure waves.\textsuperscript{36}

In this chapter, we describe the use of high speed videography (HSV), combined with image processing algorithms and electrochemical engineering principles, to quantitatively characterize key performance metrics of a membraneless, flow-through electrolyzer for water electrolysis. Analysis of HSV output is based on detecting the size and position of bubbles as they detach from the electrode surface. When this information is obtained in consecutive frames, basic electrochemical engineering principles can be employed to quantify key operating parameters such as local void fraction, gas evolution efficiency, bubble size distribution, current distribution, and more. In order to demonstrate these capabilities, \textit{in situ} HSV was recorded during operation of a membraneless electrolyzer containing two mesh flow-through electrodes that were positioned at 180 degrees with respect to each other (Figure 3.1a). The first portion of this chapter presents a detailed description of the electrolyzer and experimental HSV setup used to monitor gas bubbles that evolve from its electrodes. Next, the image analysis and bubble detection procedures are
described and applied to experimental HSV data to generate bubble size distributions and determine gas evolution efficiencies as a function of operating time, current density, and choice of purge gas. The volume of gas detected using HSV is directly compared to the current output from the potentiostat to demonstrate the influence of bubbles on the rate of reaction. Lastly, time-averaged void fractions are calculated immediately downstream of the electrode, and the spatial variation in void fraction is compared to the primary current distribution. We expect that the experimental methods and image analysis procedures described herein will find broad utility in helping to diagnose inefficiencies and guide the design of membraneless electrolyzers.
Figure 3.1: a) Schematic top-view of membraneless electrolyzer based on two flow-through mesh electrodes placed at an angle of 90° with respect to the direction of fluid flow. The close-up photo on the right shows a magnified view of the front of a woven mesh electrode used in this study. b) Exploded diagram of the membraneless cell used in this study. c) Schematic of the assembled flow cell.

3.2 Materials and Methods

Chemicals – All solutions were prepared using 18.2 MΩ cm deionized water. Electrolyte solutions were prepared using concentrated sulfuric acid (Certified ACS plus, Fischer Scientific). Electrolytes were purged with either argon (Ar) (PurityPlus, 99.998% purity) or H₂ (PurityPlus, 99.999% purity) gas.
Electrode fabrication – The electrodes were fabricated using Titanium (Ti) woven mesh sheets (40 wires per inch; 0.007” wire diameter, Unique Wire Weaving Co., Inc.). Based on the wire diameter and number of wires per inch, 51% of the nominal area of the mesh was calculated to be void space. The measured nominal thickness of the mesh was 0.014”, an equivalent of two wire diameters. The wires were coated with 50 nm thick layers of platinum (Pt) by electron-beam evaporation. The nominal dimensions of the electrodes were a width of 0.30 cm and a length of 2.0 cm, resulting in a geometric area of 0.6 cm$^2$. The average current density of the electrode was calculated by dividing the applied current by the geometric area.

Electrolyzer fabrication – The body of the electrolysis cell was printed using polylactic acid (PLA) (Makergear LLC). Design files have been made available at http://echem.io. A 0.25” thick glass sheet is clamped to the front of the device to allow for visual observation with a high speed camera. A glass window is epoxied (J.B. Weld) to the backside of the device to allow for backlighting and improved contrast for the HSV experiments. The electrolyzer has a channel height of 0.5 cm and a full channel width of 4.5 cm. A 2 mm wide separator tab is inserted into the device to split the channel into two equal streams with a width of 2.2 cm. The central 3D printed tabs are standalone pieces allowing them to be easily interchanged to evaluate the effect of tab length ($L_{tab}$) on the measured current density distribution. A Viton gasket is inserted above and below the tab to ensure a mechanical seal when the device is clamped shut. 1 mm thick PLA baffles running parallel to the fluid flow are positioned in the regions far downstream of the electrodes as well as upstream of the electrodes to dampen any large convective cells which may form during electrolysis.

The electrodes are inserted into slots in the side of the electrolysis device and epoxied into place. The ends of the electrodes (< 1 mm) are also coated with epoxy to eliminate any
electrochemical activity at the interface between the tip of the electrode and the location where they rest on the central tab.

**Device characterization and operation** – The electrolyzer is oriented vertically such that the buoyant force acting on the electrogenerated bubbles is normal to the electrode surface. Flexible silicone tubing (Masterflex L/S 16, Cole Parmer) is fitted to the electrolyzer inlets and outlets to allow for electrolyte circulation. The tubing is connected to a peristaltic pump (Cole Parmer) which circulates the electrolyte at a total flow rate of 1 mL s\(^{-1}\). After splitting into two separate effluent streams, the average flow velocity through each sub channel of the device is 0.5 cm s\(^{-1}\). A pulse dampener (Cole Parmer) is connected between the pump and the flow cell to dampen oscillations in flow rate that are inherent to the flow from the peristaltic pump. The ends of inlet and outlet tubes are submerged in an electrolyte reservoir containing 0.5 M H\(_2\)SO\(_4\), which was continuously purged with either Ar or H\(_2\) gas. Experiments were carried out at room temperature and atmospheric pressure.

All electrochemical measurements were carried out with a Biologic SP-300 bi-potentiostat. High-speed videos of bubble evolution were recorded using an Edgertronic SC1 high-speed camera operating at 500 frames per second (fps) and a resolution of 1280 x 1024 pixels. The camera was fitted with a FotodioX 52 mm reverse mount macro adapter ring and a Nikon AF Nikor 50 mm f/1.8D lens. Each video recording captured the flow of bubbles over 10 s of operation.

**HSV processing** – Data files from HSV experiments were analyzed using MATLAB Image Processing Toolbox (R2017b). Videos were cropped to an area of analysis spanning the width of the channel (2 cm) and a vertical height of 150 px (3.2 mm). This area was located as close to the electrode as possible to capture the bubbles as they detach. More precisely, the analysis area was
located ≈ 1 mm above the gas film on the electrode, such that the large stationary bubbles adhered to the electrode did not overlap with the analysis area. The mean bubble diameter detected was 112 microns, meaning that ≈ 4% of the pixels in the analysis window were located within a distance of one mean bubble radius from the edge. It is possible that bubbles detected within this region could be detected with less accuracy since a portion of the bubble will be out of frame. However, the Hough transform algorithm is able to resolve the presence of partially visible bubbles by detecting the boundary between the bubble edge of the background. If the bubble center point is calculated to be outside the region of interest, however, the bubble will not be detected in the current frame. Another possible source of error results when the algorithm misinterprets a cluster of bubbles as a single, larger bubble. This error overestimates the total volume of gas collected, and is more likely to occur at higher applied current densities when larger quantities of bubbles are generated.

Bubble detection was carried out over 100 frames of video, with each frame separated by a time difference of 0.1 s. Each frame was converted to a binary image using an adaptive thresholding algorithm developed by Bradley. In MATLAB this algorithm is included in the ‘imbinarize’ function, and a sensitivity parameter of 0.7 was specified. After binarizing the image, a circular Hough transform algorithm was used to detect circles of varying radii and spatial position. The Hough transform is included in the ‘imfindcircles’ function, which requires an estimated size range of bubbles to detect. Table 3.1 in the Appendix A, Section 3.5.1, lists the size range and sensitivity parameter inputs used for circle detection. The ‘imfindcircles’ command was run five consecutive times to account for detection of bubbles in different size ranges. The algorithm searches for the largest bubbles before searching for smaller bubbles. Within each step, the Hough transform is capable of identifying multiple circles which partially overlap each other.
In some instances, if the bubbles are too close together, the algorithm may interpret them as 1 larger bubble. Once a bubble is detected, it is temporarily “deleted” from the image before searching for more bubbles. Thus, if there is a smaller bubble present directly in the foreground of a larger bubble, the smaller bubble will likely not be counted due to “shadowing” by the larger bubble. The parameters in Table 3.1: Input parameters for Hough Transform algorithm used in this study for bubble radius detection range and sensitivity were optimized to minimize these errors.

**Finite element modeling** – The potential field in the electrolyte was modeled using the COMSOL Electrochemistry Multiphysics package. The current distribution along the electrode was determined by numerically by solving Laplace’s equation, $\nabla^2 \phi(x, z) = 0$, for a two dimensional system where $\phi$ is the local potential as a function of position. This equation is valid only when there are no concentration gradients present in the cell. In 0.5 M H$_2$SO$_4$, the operating current was determined to be well below the maximum possible current for electrodes with mixing due to free convection, and thus it is reasonable to assume that concentration gradients are negligible. For the electrolyzer in this report, the limiting current was estimated to be $\approx 3.7$ A cm$^{-2}$, which was determined by solving for the effective diffusion of protons across a boundary layer. A more detailed overview of this calculation, which uses a mass transfer correlation for a system with free convection, can be found in the Appendix A, Section 3.5.2. At an applied current density of 100 mA cm$^{-2}$, the electrolyzer is operating at approximately 2.7% of the calculated limiting current value.

The spatial coordinates of the electrodes and insulating walls were specified and the appropriate boundary conditions were imposed. At an insulating boundary, the current normal to the insulator was fixed to be zero. At the counter electrode, a constant potential boundary condition was applied where $\phi = 0$. A constant potential boundary condition was also applied to the working
electrode, $\phi = V$, where $V$ is the applied voltage across the cell. Charge transfer resistance effects were not considered in this model due to their negligible magnitude relative to the ohmic resistance. Numerically, the ratio of charge transfer resistance to ohmic resistance is known as the Wagner number ($Wa$), and for this system is calculated to be less than 0.11 for all operating conditions tested in this cell (see Appendix A, Section 3.5.3). Experiments which attempt to quantify the current distribution were carried out at a nominal current density of 50 mA cm$^{-2}$, and were found to have a $Wa$ value of 0.04. After solving for the potential as a function of position, the local current density is determined at the electrode boundary from Ohm’s law, $i = \kappa \nabla \phi$, where $\kappa$ is the conductivity of the electrolyte.

3.3 Results and Discussion

3.3.1 Description of the electrolyzer and HSV setup

A top view schematic of the electrolysis cell used in this study is shown in Figure 3.1. In all experiments, the electrolyzer was oriented vertically and the 0.5 M H$_2$SO$_4$ electrolyte was fed into the inlet port located at the bottom of the device. Upon entering the electrolyzer, the flowing electrolyte diverges as the channel expands to a width of 4.5 cm before impinging on the two mesh electrodes that span the channel width and are located 5.3 cm downstream from the inlet port. An insulating tab separates the two electrodes at the center of the channel and causes the flowing electrolyte to split into two streams. During operation, a voltage is applied between the cathode and anode mesh electrodes, where hydrogen (H$_2$) and oxygen (O$_2$) evolve, respectively. Product gases can exist either in the dissolved phase or as bubbles which grow on the electrode surface. As the electrolyte flows through the mesh electrodes, it promotes detachment of gas bubbles and sweeps both dissolved and gaseous species upwards into the two effluent ports. In all
measurements carried out in this study, the electrolyte passes through the electrolyzer at an average velocity of 0.5 cm s\(^{-1}\).

The woven pattern of the mesh electrodes is shown in the inset in Figure 3.1a. The void spaces in the mesh enable the electrodes to span the width of the channel (\(L_e\)) while also allowing electrolyte to continuously flow through the electrodes. Ionic transport occurs in the region upstream of the electrodes. For larger values of \(L_e\), the ionic conduction path increases and a larger overall ohmic resistance to the outer edges of the cell is expected. A tab of variable length \(L_{tab}\) extends into the upstream portion of the cell (the region below the electrodes in Figure 3.1a). A tab of this nature can help prevent cross-over of product species between the anode and cathode, but does so at the expense of increasing the average ionic conduction path between the two electrodes. As a result, changes in the tab length are expected to both increase the ohmic resistance of the cell and alter the current distribution.

The current distribution of the cell arrangement illustrated in Figure 3.1a is also strongly dependent on the ohmic resistance of the electrolyte to ion transport (\(R_\Omega\)) and the charge transfer resistance at the electrode surface due to reaction kinetics (\(R_S\)). The ratio \(R_S/R_\Omega\) is known as the Wagner number (\(Wa\)), the magnitude of which can be used as a useful guide for modeling non-uniform current distributions. For the length scale and current densities being passed through the electrolysis cell in this study, \(Wa \ll 1\) (see Appendix A, Section 3.5.3), meaning that the current distribution along the length of the electrodes is expected to depend strongly on the ohmic resistance and the geometric orientation of the electrodes relative to the insulating boundaries.

An exploded view of the electrolyzer is shown in Figure 3.1b. The front side window is clamped to the body of the cell and allows for a clear viewing path for the high speed camera. A permanent window is epoxied to the backside of the device to ensure adequate lighting. The
separator tab is printed as an insert-able piece, allowing for different tab lengths to be tested in one cell. The mesh electrodes are inserted through slits in the sides of the cell and are epoxied into place. Figure 3.1c shows the fully assembled device.

**3.3.2 Description of image analysis procedure**

Once the cell is assembled, sulfuric acid is pumped through the device and a current is applied to drive the electrolysis reaction at a specified rate. Using HSV, the electrogenerated bubbles are detected immediately downstream of the electrode. Figure 3.2a shows the region downstream of the cathode which is recorded using HSV, while Figure 3.2b contains a representative still frame image taken during electrolysis. During HSV post processing, individual frames are cropped down to the region of interest, typically spanning the width of the cathode channel and a vertical height of 3.2 mm, corresponding to 150 pixels (Figure 3.2c). Using the MATLAB Image Processing Toolbox, the grayscale image is converted into a binary black and white image (Figure 3.2d). A Hough transform algorithm is used to determine the size and positions of circles in the frame.\(^{38}\) The algorithm is carried out in five successive steps as described in the Appendix A, Section 3.5.1. In order to verify the accuracy of the circle detection algorithm, the recorded bubble size and coordinate data is used to digitally super-impose red circles on top of the binarized image. As shown in Figure 3.2e, good agreement exists when the circles are able to outline all the bubbles in the image. In order to determine the local void fraction of gas, the image can be further discretized into finite volumes. Although 5,000 frames of images are recorded over 10 s, only 100 evenly spaced frames were typically analyzed to reduce computational cost. A sample video showing 100 frames of analysis for a cathode operating at 50 mA cm\(^{-2}\) is available in the online publication of *J. Electrochem. Soc*, **166**, F312-F321 (2019).
Figure 3.2: Procedure used for processing HSV images recorded during electrolysis. a) Schematic showing the region of the electrolyzer recorded with HSV. b) Still frame from a HSV showing H₂ bubbles evolving from a mesh cathode operating with an average current density of 50 mA cm⁻² in 0.5 M H₂SO₄. c) The still frame is cropped to limit analysis to a narrow section of the channel located immediately downstream of the cathode. d) Conversion of the cropped still frame into a binary image to reduce background interference. e) The size and position of bubbles are determined using a circle detection algorithm, with detected bubbles shown by red circles that are overlaid with the binary image from (d). The local density of bubbles, used to estimate the local current density, is determined by discretizing the analysis area into equal control volume.

In addition to determining the size and position of the bubbles in each frame, HSV images were used to determine their terminal velocities. Often times, trends in bubble velocity can give
insights into the flow properties of a fluidic system. Because of the vertical orientation of the electrolysis cell and the slow rate of electrolyte pumping, the bubble velocities depend predominantly on buoyancy forces. In general, bubbles leaving the electrode surface were found to rise vertically with minimal motion in the horizontal direction. During post processing of HSV experiments, the terminal velocities of bubbles of varying size were determined by directly measuring the time required to rise vertically across a set distance (6.4 mm). Figure 3.3a shows the terminal velocities of several H\textsubscript{2} bubbles as a function of bubble radius in an electrolyzer operating at 20 mA cm\textsuperscript{-2}. The linear trend in terminal velocity with the radius of the bubble suggests that the velocity is governed by the buoyancy force, which scales with the volume of gas contained.\textsuperscript{40} The Reynolds numbers of the bubbles in Figure 3.3a were found to be between 0.6 and 150 depending on the diameter of the bubble. Given the predictable relationship between bubble radius and velocity, an empirical fit of the trend was incorporated in the analysis algorithm to calculate bubble velocities without the need to track bubbles between frames. For example, if a bubble is detected with a radius of 0.2 mm, then a vertical velocity vector of 5 cm s\textsuperscript{-1} can be reasonably assumed based off of the fit in Figure 3.3a. A similar analysis comparing the total bubble volume detected to the unique bubble volume detected as a function of the size of the time step is shown in Figure 3.10 (see Appendix A, Section 3.5.5) for an operating current density of 100 mA cm\textsuperscript{-2}.
Figure 3.3: a) Terminal velocity of H₂ bubbles rising off of the cathode as a function of bubble radius. Individual bubble velocities were determined in an electrolyte of 0.5 M H₂SO₄ purged with Ar and pumped at an average velocity of 0.5 cm s⁻¹. Bubbles were generated at an operating current density of 20 mA cm⁻². b) Total volume of gas detected over the duration of a 10 s HSV as a function of time step between image frames. Image analysis was conducted for the same
experimental conditions as in (a). Also marked on the plot is the maximum residence time for a bubble traveling across the analysis area. The red star corresponds to a time step of 0.1 s, which was the time step used for the remainder of the analyses in this paper. c) Cumulative volume of H$_2$ bubbles detected during a 10 s long HSV. Image analysis was conducted for the same experimental conditions as in (a). The upper trace corresponds to the total volume of all bubbles detected in the image frame, whereas the lower trace uses an algorithm that ensures that each individual bubble is only counted once.

3.3.3 **Determining gas evolution efficiency and bubble size distribution**

After determining the size, position, and velocity of all detected bubbles in each frame, the total volume of gas detected can be calculated as a function of time. This is calculated by summing up the total volume of bubbles detected in each frame. However, for a given size of the analysis area and time step between video frames (Δt), individual bubbles can be counted multiple times depending on their velocity and distance traveled between frames. An algorithm for avoiding counting a bubble multiple times was implemented to ensure accurate measurements of the total volume of gas detected. As shown in Figure 3.9 (See Appendix A, Section 3.5.4), the velocity vector of each detected bubble can be used to infer where a bubble was located in a previous frame. If the bubble is calculated to be out of frame in the previous time step, then it is counted as a uniquely detected bubble. Figure 3.3b shows the total volume of gas bubbles summed across all frames of a 10 s HSV as a function of Δt. This analysis was conducted for the same experimental conditions as in Figure 3.3a. Decreasing Δt increases the total number of frames, and thus total number of bubbles detected over the duration of a 10 s video. The plot compares the sum of all bubbles detected to the total sum when applying the unique bubble algorithm. At large Δt, the two traces converge, suggesting that every bubble in the analysis area is being detected for the first time. Δt is considered large when it exceeds the maximum residence time of a bubble in the analysis area ($t_{max}$). This value is based on the time required for the smallest and therefore slowest
moving bubbles to cross the analysis area, and is calculated to be approximately 0.3 s. When $\Delta t < \tau_{\text{max}}$, the trace for uniquely detected bubbles begins to diverge and plateau as duplicate bubbles are screened out from the total sum. In the interest of the computational time for analysis, a time step of 0.1 s was chosen for this study, and is marked as a red star on Figure 3.3b. Choosing an even smaller $\Delta t$ can potentially result in a greater volume of unique bubbles detected, but this is dependent on the precision of the algorithm itself. Any errors in determining the size and velocity of a bubble can increase the uncertainty of the unique detection algorithm, and thus bubbles can still be counted multiple times. A more thorough error analysis can help better determine the optimal choice of time step, but is beyond the scope of this report.

Having chosen a time step of 0.1 s, Figure 3.3c shows the cumulative sum of H$_2$ bubble volumes detected as a function of time for electrolysis at 20 mA cm$^{-2}$. The linear trend in cumulative volume suggests a constant rate of H$_2$ production, which is expected due to the constant applied current density. A fit of the slope can be taken to calculate the average volumetric flow rate of H$_2$ coming off of the electrode. The second trace in Figure 3.3c shows the total volume of uniquely detected bubbles for the same HSV recording. By employing the algorithm for determining uniquely detected bubbles, we avoid overestimating the volumetric flow rate of H$_2$ evolved by a factor of two.

The volume detection analysis can be applied to higher current densities as well. As shown in Figure 3.4a, the slope of the cumulative volume detected as a function of time increases in proportion to the current density applied. This trend is unsurprising, as higher current densities should correspond to higher rates of gas production according to Faraday’s law. However, the relationship between current density and volumetric flow rate of gas is not necessarily perfect. The empirically fitted volumetric flow rate based on uniquely detected bubbles can be compared to the
maximum theoretical value based on the applied current, Faraday’s law and the density of H\textsubscript{2} gas. This ratio between the experimental and theoretical values of product species entering the gas phase is referred to as the gas evolution efficiency (\(GEE\)), and can be calculated using Equation 3.1:

\[
GEE = \frac{\hat{V}_{HSV} \cdot n \cdot F}{i \cdot A_e \cdot MW_{H_2} \cdot \rho_{H_2}}
\]  

3.1

where \(\hat{V}_{HSV}\) is the empirically fitted volumetric flow rate of H\textsubscript{2} based on the uniquely detected bubbles from HSV, \(n\) is the number of electrons transferred per molecule of H\textsubscript{2}, \(F\) is the Faraday constant, \(i\) is the current density, \(A_e\) is the geometric area of the cathode, \(MW_{H_2}\) is the molecular weight of H\textsubscript{2}, and \(\rho_{H_2}\) is the density of H\textsubscript{2}. Typically, the gas evolution efficiency is expected to be less than 100\% due to dissolution of the products into the aqueous phase.

Figure 3.4: a) Total volume of H\textsubscript{2} detected over the cathode during a 10 s long HSV. Each trace corresponds to bubbles detected for a constant average applied current density of 20, 50, or 100 mA cm\textsuperscript{-2}. In each trial, the electrolyte was pre-saturated with Ar. b) Gas evolution efficiency of the cathode as a function of time. Each data point corresponds to the average gas evolution efficiency over the course of a 10 s HSV. Experiments were carried out for a cathode operating at 20 mA cm\textsuperscript{-2} in 0.5 M H\textsubscript{2}SO\textsubscript{4}, and the effect of Ar and H\textsubscript{2} saturation gases is compared. The total volumetric flow rate of electrolyte through the cell was set to 1 mL s\textsuperscript{-1}.
Using this methodology, Figure 3.4b shows the gas evolution efficiency of the cathode as a function of time. For this experiment, a fixed current density of 20 mA cm\(^{-2}\) was applied for 30 minutes. Each data point corresponds to the average gas evolution efficiency calculated from 10 s long HSVs that were recorded every 5 minutes. The plot also compares the effect of saturating the electrolyte feed with different dissolved gases. For both H\(_2\) and Ar, there was no clear time dependence of gas evolution efficiency. In general, a lower gas evolution efficiency of 40-60\% is observed when the electrolyte feed is saturated with inert Ar. When the saturation gas is changed to H\(_2\), the gas evolution efficiency increases to 80-90\%. This trend is expected, as losses in gas evolution efficiency are typically attributed to solvation of the product gas.\(^{41}\) By pre-saturating the electrolyte feed with H\(_2\), its capacity to solvate electrogenerated H\(_2\) at the cathode is diminished. Although the applied current density during these experiments is constant and continuous, the rate of bubble detachment, particularly large bubbles, is not. The departure events of large bubbles can be missed if they occur in the time between video recordings. For comparison, the electrolysis experiment spanned 30 minutes, but the image analysis was only carried out for only 60 s of recorded footage. Thus, in the 29 minutes of electrolysis between recorded HSVs, large bubble detachment events can occur undetected.

Apparent losses in gas evolution efficiency can also be attributed to small bubbles that are not detected by the camera. The minimum bubble diameter detected in this work was approximately 85 \(\mu\)m (4 px). Bubbles smaller than this radius could go undetected and thus unaccounted for in the volumetric measurements. The smallest bubble sizes for electrochemically generated H\(_2\) have been proposed to be as small as nanometers in diameter.\(^{42}\) The stability of these nanobubbles is debated in literature,\(^{43}\) but it is well known that the concentration of dissolved H\(_2\) in the vicinity of the electrodes is often at super saturation levels.\(^{44}\) In either case, the presence of
H₂ in the electrolyte at concentrations exceeding the solubility limit would manifest itself in an even lower gas evolution efficiency. Using higher optical magnification, beyond that which is available in this study, is one approach to overcoming this limitation.

From a product collection perspective, achieving a 100% gas evolution efficiency is desirable. Flowing the electrolyte can help separate the electrogenerated H₂ and O₂ bubbles, but flowing too fast can cause dissolution losses and decrease the gas evolution efficiency. A drawback of the present calculation procedure for gas evolution efficiency is that it assumes that the dominant mode of bubble transport is buoyancy, and thus bubble velocity can be inferred from its size. Knowing this velocity is a critical aspect of the unique bubble counting method that prevents overestimation of the gas evolution efficiency. At higher flow rates, however, the velocity of the bubbles is determined by the drag and slip relationship with the surrounding liquid velocity profile. Because the bubbles are being detected in the immediate wake of the mesh flow through electrodes, the liquid phase velocity profile is not readily known or easily solved for without using advance computational fluid dynamics (CFD) tools. An alternative approach could be to use a method which tracks bubbles between frames to directly calculate their velocities.

Ideally, the HSV analysis should be able to guide the design and choice of operating conditions for membraneless electrolyzers such that the gas evolution efficiency and the collection of the products is maximized. When considering the transport and collection of product gas bubbles, it can be advantageous to know or be able to predict the sizes of bubbles produced by the electrolyzer. Figure 3.5a shows the number and volumetric probability distributions of bubble diameters generated by a cathode in H₂ saturated electrolyte. For the number probability distribution, each detected bubble is weighted equally and the height of each bar corresponds to the number of bubbles detected in a bin range relative to the total number of bubbles detected
across all bins. Using this methodology, ~99% all the bubbles detected were less than 250 μm in diameter. The volumetric probability distribution was generated by weighting each bubble by its volume relative to the total volume of gas detected. Although only 1% of bubbles detected were larger than 250 μm, these bubbles contained ~25% of the generated volume. It can be convenient to re-plot the size distribution data as a cumulative volume distribution function (CVDF) as shown in Figure 3.5b. For a given bubble diameter \(d_i\), the cumulative volume fraction refers to the summation of volumes detected for all bubbles with a diameter less than or equal to \(d_i\). The volume fraction is normalized to the total cumulative volume detected for all bubbles. The data in Figure 3.5b corresponds to the same experiments as reported in Figure 3.4b for an electrolyzer operating at 20 mA cm\(^{-2}\) in Ar saturated electrolyte. In these experiments, the arithmetic mean bubble diameter was found to be 112 ± 0.5 μm. When weighted by the gas of volume contained, the average bubble diameter was found to be 177 ± 1.7 μm. The error bars associated with the mean bubble diameter represent 95% confidence intervals based on analysis of 600 still frames of the HSV recordings. Approximately 25% of the total detected gas volume comes from bubbles smaller than 100 μm for all of the 10 s HSV clips analyzed, although the volume distribution of larger bubbles can vary greatly between experiments. We attribute the large variation in the shape of the cumulative volume fraction curves in Figure 3.5b at large bubble diameters to the fact that the frequency of larger bubble detachment events is similar to the 10 s window of the HSV recording. Not surprisingly, the reported gas evolution efficiency values in Figure 3.4b show some common trends with the bubble size statistics reported in Figure 3.5b for the Ar-saturated electrolyte. In particular, the trials in which larger bubbles are detected – such as the videos recorded at the 10 and 15 minute mark – tend to exhibit higher gas evolution efficiencies (~60%) in Figure 3.4b. HSV recordings characterized by bubble distributions skewed to smaller sizes – such as at the 5, 20, and
30 minute marks – the gas evolution efficiency is found to be at or below 50%. Again, these trends do not necessarily imply that larger bubbles cause an increase in the average gas evolution efficiency. These quantities are nonetheless correlated in our experiments, and this is likely due to the fact that large bubbles detach less frequently than smaller bubbles, on average. An exception to this trend is the video at the 25 minute mark, for which a large bubble detachment event was recorded but a relatively low gas evolution efficiency was still calculated. This suggests that the variation in gas evolution efficiency with time cannot be entirely explained by infrequent large bubble detachment events.
Figure 3.5: Analysis of H₂ bubble sizes determined from HSV recorded during a 10 s of electrolysis in 0.5 M H₂SO₄ with electrolyte pumped through the electrolysis cell at a rate of 1 mL s⁻¹. a) Probability histogram of bubble diameters evolved from the cathode during electrolysis at 20 mA cm⁻² in Ar saturated electrolyte. HSV was recorded after 5 minutes of electrolysis. Number probabilities were calculated based on the number of bubbles counted, while volumetric probabilities were calculated based on the total volume of gas generated. b) Cumulative volume fraction as a function of bubble diameter for H₂ bubbles evolving from a cathode operating at 20 mA cm⁻² in Ar saturated electrolyte. Each trace corresponds to a different start time of HSV and represents the same data set reported in Figure 3.4b. c) Cumulative volume fraction as a function of bubble diameter for different operating current densities in Ar saturated electrolyte. HSVs were recorded after 5 minutes of electrolysis. d) Comparison of cumulative volume fraction as a function of bubble diameter for a cathode operating at 50 mA cm⁻² in different saturation gases. HSVs were recorded after 5 minutes of electrolysis.
It should be noted that the presented analysis on bubble size distributions is not exhaustive, and many factors can contribute to the observed profiles. Figure 3.5c shows the effect of current density on the CVDF, and suggests that higher current densities promote the formation and detachment of more large bubbles from the electrode. In reality, the frequency of detachment for all bubbles increases with increasing current density. For large bubbles with long periods between isolated detachment events, an increase in frequency means a higher likelihood of large bubbles being detected by the camera during the 10 s window of recording. Figure 3.5d shows that saturating the electrolyte with H₂ gas results in larger bubbles being formed compared to saturating the electrolyte with Ar gas, which may be attributed to the higher frequency of gas bubble detachment from the electrodes due to the higher gas evolution efficiency in H₂ saturated electrolyte.

Despite the complexities of modelling and quantifying the distribution of bubble sizes, this information is highly relevant for membraneless electrolyzers that rely on transport of multiphase electrolyte fluid to separate the electrogenerated gases. One question is if there is an optimal bubble size for maximum product collection efficiency. Intuitively, a multiphase mixture containing a large volume fraction of small bubbles may not separate as readily as larger bubbles. The shape of the electrodes can also affect the size and frequency of detaching bubbles. For example, on a large planar electrode, bubbles adhere for longer before growing to a size in the millimeter scale when they can detach. On the mesh electrodes, however, smaller bubbles readily form and detach. Given that the mesh wire diameter in this study is approximately 180 μm, it is not surprising that the majority of bubbles detected have a diameter of less than 200 μm. Many plots in Figure 3.5 commonly show an uptick in bubbles detected at d = 425 μm, which may be a result of the gap spacing distance in between wires (approximately 460 μm). Surface treatment of the electrodes
could also affect the bubble detachment size by altering the average contact angle of the bubbles attached to the surface, and therefore the average force of adhesion.

### 3.3.4 Comparing HSV-derived current transients to measured current transients

Most of the discussion up until this point has focused on the response of bubble behavior to changes in the electrochemical properties of the system. However, the presence of bubbles can also affect the observed electrochemical properties. For example, the electrogenerated bubbles can behave as resistors to ionic flow as well as block active sites on the electrode surface. The relationship between the effective conductivity and the void fraction of bubbles in the electrolyte has been widely observed, and can be described by the Bruggeman equation. An important consideration for the electrolysis cell presented herein is that the detached bubbles in the electrolyte exist downstream of the electrodes, whereas the ionic transport and potential field gradient exists only upstream of the electrodes. Thus, any effect of the bubbles on the electrochemical properties of this electrolyzer will result from bubbles that are attached to the electrode.

To view the impact of bubbles on the electrochemical performance of this electrolyzer, HSV was recorded simultaneously with electrical current during a chronoamperometry experiment where a constant 2.7 V was applied between the anode and the cathode. The relationship between the volumes of bubbles detected and the current readout from the potentiostat is shown in Figure 3.6 over a 10 s period. The current measured by the potentiostat varies between 60-65 mA, with the largest jumps in the current occurring at t = 0.8 and 3.8 s. These step changes in current correspond precisely with the timing of large bubble departure events measured using HSV. The increase in current at these points is likely due to improved ionic conductivity of the electrolyte between the electrodes in the absence of the detached bubbles. A smaller step change in the current density at t=6.5 s also coincides with a large bubble detachment event. However, some changes in
the current versus time plot have no direct correlation to large bubble departure events, such as at \( t = 2.5 \) and 8.5 s. One explanation for these changes is that only the cathode was filmed for large bubble departure events, but bubble departures at the anode can also decrease the ohmic resistance of the ionic conduction path. Because the experiment was conducted without a reference electrode, bubble departure events at the anode cannot be isolated from the current response.

Figure 3.6: Comparison of the total electrolysis current recorded by the potentiostat to the volume of gas detected during a 10 s HSV. The HSV was recorded after five minutes of electrolysis at a constant applied voltage of 2.7 V. Electrolysis was carried out in 0.5 M \( \text{H}_2\text{SO}_4 \) with electrolyte pumped through the cathode at a rate of 0.5 mL s\(^{-1}\).

One might expect to see a direct correlation at a given time between the measured current and the detected volume of \( \text{H}_2 \). In reality, there are temporal differences between the rate of reaction and the rate of hydrogen bubble accumulation, growth, and departure. As mentioned in
the discussion of Figure 3.4a and b, when the volume H₂ detected is averaged over time, a direct relationship to the applied current can be calculated using Faraday’s law. When comparing the measured current to the volume of gas in an individual video frame, however, the relationship with Faraday’s law is less straightforward. Large bubbles are the result of hydrogen accumulation over the course of several seconds or even minutes. For an experiment with a constant applied potential, the departure of a large hydrogen bubble causes an increase in current because it is no longer able to insulate the electrode from ions conducting to it. However, it is not necessarily true that a sudden increase in current causes a large bubble to detach.

### 3.3.5 HSV-derived current distributions

Because the rates of gas bubble generation are proportional to current density, as seen in Figure 3.4a, HSV recordings of gas-evolving electrodes can also be used to measure the current distribution of an electrochemical cell. A non-uniform current distribution, or variation in local reaction rate, can arise from differences in ohmic resistance, charge transfer resistance, or mass transport resistance along the length of the electrode.⁴⁷,⁴⁸ In our system, mass transport effects can be neglected because the typical operating current densities used in this study are well below the mass transport-limited current for the water splitting half reactions in 0.5 M sulfuric acid. In electrochemical systems without mass transport limitations, the governing physics of the current distribution can be determined by calculating $W a$ – the ratio of charge transfer resistance to ohmic resistance.³⁹ In most electrochemical systems, ohmic resistance is dominated by the ionic transport in the electrolyte, and depends on the conductivity and path length for ion transport between electrodes. The charge transfer resistance is determined by the kinetics of the electrode reaction. Equation 3.2 shows the calculation for $W a$ when the charge transfer resistance is modeled using Tafel kinetics:³⁹
\[ Wa = \frac{R \cdot T \cdot \kappa}{\alpha_{CT} \cdot F \cdot |i_{avg}| \cdot l} \]

where \( R \) is the gas constant, \( T \) is the temperature, \( \kappa \) is the conductivity of the electrolyte, \( \alpha_{CT} \) is the charge transfer coefficient, \( F \) is the Faraday constant, \( i_{avg} \) is the average current density of the electrode, and \( l \) is a characteristic length, which in this study is taken to be the length of the electrode. As \( Wa \) approaches zero the current distribution is governed by the ohmic resistance and the kinetics of reaction do not need to be modelled. Such systems are said to obey the primary current distribution. For larger values of \( Wa \), the reaction kinetics at the electrode surface also become important to describing the variation in current along an electrode. A higher relative charge transfer resistance generally causes the current distribution to be more uniform, resulting in what is referred to as the secondary current distribution.

When the electrolyzer is operated at a nominal current density of 50 mA cm\(^{-2}\), the calculated value of \( Wa \) is 0.04, indicating that the current distribution can be modeled as a primary current distribution. The current distribution can be determined after solving the potential field, which is governed by Laplace’s equation, \( \nabla^2 \phi = 0 \), where \( \phi \) refers to the local potential of the electrolyte. Non-uniformities in the primary current distribution arise due to the geometry of the electrolyzer and the spatial position of the electrodes. In this work, the lowest ohmic resistances to ion transport, and hence the highest local current densities, are expected in the center of the electrolyzer where the shortest distance for ion transport between electrodes is possible. Moving outward from the central baffle, longer ion transport distances are required, leading to higher ohmic overpotentials and small local current densities that decrease to a minimum at the outer edges of the channel walls. Figure 3.7a shows the simulated local current density of the cathode as a function of position along the length of the electrode. Each trace corresponds to a different tab length, \( L_{\text{tab}} \). When the insulating tab extends only 2 mm upstream of the electrodes, the current
distribution is very non-uniform, with the modeled local current density at the inner edge of the electrode being 3 times the average current density ($i_{avg}$). The current density decreases along the length of the electrode, with the local value tapering off to approximately half of $i_{avg}$ at the outer edge.
Figure 3.7: Comparison of the predicted current distribution to the local volume fraction along the length of the electrode, $x/L_e$. All experiments were carried out at an operating current density of 50 mA cm$^{-2}$. a) Simulated primary current distribution of the electrolysis cell for different separator tab insertion lengths between electrodes. b) Relative gas volume fraction along the length of the electrode for a cathode operating in Ar-saturated H$_2$SO$_4$ and tab insertion length of 2 mm. c)
Relative gas volume for a cathode operating in Ar-saturated H₂SO₄ and a tab insertion length of 10 mm. Error bars were determined by calculating the 95 % confidence interval based on averaging the volume fraction over 100 frames of data.

Since the tab is an insulator that directly protrudes into the path of ion flow, increasing $L_{\text{tab}}$ increases the average distance of ion transport and alters the current distribution. For longer tab insertion lengths, the current distribution becomes more uniform. When the tab extends 10 mm between the electrodes, the local current density at the interior edge of the electrode is predicted to be only $1.2 \times l_{\text{avg}}$. As the vertical conduction path around the upstream end of the tab increases, the ohmic drop in the horizontal direction becomes less relevant. In the limit of an infinitely long tab, the system behaves as a pair of parallel electrodes directly in contact with perpendicular insulating walls, which is expected to have a uniform current distribution. This is a special case of the primary current distribution, and the predicted uniform current distribution is entirely a result of the reactor geometry. Although the current distribution is more uniform at longer tab insertion lengths, this comes at the cost of a higher overall ohmic resistance in the cell. The primary value of having a tab extending upstream of the electrodes is to prevent bubbles from crossing over between the anode and cathode. In the electrochemical cell employed in this study, varying the tab length can be useful for modulating the current distribution and comparing the modeled distribution to that estimated from HSV-derived current distributions.

In this work, experimental current distributions were represented by the HSV derived void fraction of gas ($\alpha$) determined along the length of the electrode. This was done by calculating the average volume of gas detected within discrete control volumes along the length of the electrode for 100 frames of video recorded over 10 s. Figure 3.7b shows the local values of HSV-derived $\alpha$ for electrolysis at 50 mA cm⁻² in Ar-saturated electrolyte normalized to the average void fraction ($\alpha_{\text{AVG}}$) detected for all control volumes along the electrode for $L_{\text{tab}} = 2$ mm. Also plotted is the
simulated primary current distribution normalized to the average applied current density. The experimental and modeled current distributions show reasonable agreement, with the local void fraction being highest at the interior edge of the electrode. When $L_{\text{tab}}$ is extended further upstream, the void fraction profile responds accordingly. Figure 3.7c shows the void fraction profile during electrolysis at 50 mA cm$^{-2}$ in an Ar-saturated electrolyte with a 10 mm tab length bisecting the anode and cathode. Under these conditions, the HSV-derived void fraction exhibits significantly lower variation along the length of the electrode, consistent with the simulated primary current distribution. A large bubble detachment event at $x/L_e = 0.8$ causes a large local spike in void fraction which does not correspond with the predicted local current density. This suggests that although small bubble departure events accurately reflect the local current density, larger bubbles in the gas film may depend on other mechanisms for growth and detachment and lead to significant error in measuring current distributions by our method. For example, a disturbance in the flow may cause a large bubble to detach from a random position. The time period between large bubble detachment events is also likely to occur at intervals longer than the video recording time of 10 s, yielding uncertainty as to where on an electrode a large bubble will detach. In general, the deviations between the simulated current distribution and the void fraction distribution are likely due to the dynamics of bubble growth and detachment. It may be possible that the role of charge transfer resistance may also help resolve these deviations, but a comparison of the primary and secondary current distributions in Figure 3.11 (See Appendix A, Section 3.5.6) shows that adding kinetic parameters to the boundary conditions modifies the local distribution by no more than 11% at any point along the electrode.

In light of these uncertainties, more can be done to resolve differences between theory and experimental data gathered from HSV. Deviations from the predicted current distribution also exist
in Figure 3.7b at the ends of the electrodes where they approach the inner insulating divider and the outer channel wall. It is possible that both flow and electric field effects contribute to these deviations at the boundaries. For example, fluid velocity gradients caused by the separator tab and the exterior channel walls may affect bubble transport in ways that are not accounted for in our analysis method. An algorithm which tracks bubbles in between frames could better determine the velocity profile and in turn help incorporate a theoretical model which accounts for velocity field effects.

3.4 Conclusions

This Chapter has described and demonstrated methods of using HSV to quantitatively characterize the behavior of electrochemically-generated bubbles in a membraneless electrolyzer. A circle detection algorithm was employed to determine the size and position of H₂ bubbles as they detached from a mesh flow-through cathode. By ensuring that bubbles were not counted multiple times, average gas evolution efficiencies were calculated as a function of time and for different electrolyte conditions. The gas evolution efficiency was found to be 40-60% in Ar saturated electrolyte, but increased to almost 90% when the electrolyte was pre-saturated with H₂. These results suggest that dissolution of the electrogenerated H₂ can be a major source of product loss in this membraneless electrolysis system. Statistical analysis of the bubble size distribution indicates that inaccuracies in gas evolution efficiency can be attributed to the infrequent detachment of larger bubbles. Although only 1% of the bubbles detected were larger than 250 μm, they contained over 20% of the generated volume. During experiments for which a constant potential applied, large bubble departure events were associated with a temporary increase in the current passed through the cell. This study has also demonstrated that in situ HSV imaging can be used to determine the local void fraction of bubbles immediately downstream of a flow-through electrode. When the
void fraction is plotted as a function of position along the length of the electrode, the profile was found to correspond to the primary current distribution. Overall, we show that HSV can be a versatile non-invasive imaging tool to quantitatively analyze bubble evolution dynamics and electrochemical properties of membraneless electrolyzers.

3.5 Appendix A

3.5.1 Parameters used for Hough transform

The imfindcircles function in the MATLAB Image Processing Toolbox uses a Hough transform algorithm that is able to analyze a still frame image and return the radius and (X,Y) center point locations of all circular features present. Circles are determined by first identifying bubble edges, which are defined by dark pixels that are directly adjacent to white background pixels. The center point for each circle is determined statistically based on the coordinates of the detected edges and the anticipated range of bubble radii. The function requires that the user specify the size range of circles to be detected, as well as a sensitivity parameter which sets the minimum threshold probability value for whether or not a grouping of boundary pixels constitute a circle with a common center. The sensitivity parameter must be specified within the range of 0 to 1, with a value of 1 corresponding to the highest sensitivity. Increasing the sensitivity allows for the detection of overlapping or poorly defined circles, but setting it too high increases the likelihood of false positive detection. Setting the sensitivity parameter too low can result in not detecting any bubbles. In order to avoid false positive detection of smaller bubbles, it is also recommended that the size range be specified such that the maximum anticipated radius size does not exceed three times the magnitude of the minimum radius size to be detected. Because of the wide range of possible bubble sizes present in the videos imaged in this study, the imfindcircles function was used to analyze each image frame five different times, where successively smaller bubble size
ranges were analyzed after artificially removing the larger bubbles that were identified in the prior analyses. Circles were removed from the image using the MATLAB insertShape function to replace the detected circles with pixels matching the background color. Table 3.1 provides a summary of the parameters specified for each successive scan applied to every frame of a HSV.

Table 3.1: Input parameters for Hough Transform algorithm used in this study

<table>
<thead>
<tr>
<th>Analysis number</th>
<th>Lower radius / pixels</th>
<th>Upper radius / pixels</th>
<th>Sensitivity / unitless</th>
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</tr>
<tr>
<td>5</td>
<td>2</td>
<td>4</td>
<td>0.99</td>
</tr>
</tbody>
</table>

3.5.2 Limiting current density and I-V curve of electrolyzer

Due to electroneutrality, the average limiting current of a binary electrolyte can be treated as a convection diffusion problem with correction factors calculated for the diffusion coefficient. In an electrochemical system with a boundary layer defined by free convection, the limiting current can be calculated as shown in Equation 3.3:

$$i_{avg} = -C \frac{z_+ \nu_+ c_\infty F D}{1 - t_+} \left[ \frac{g (\rho_\infty - \rho_0)}{\rho_\infty D \nu L} \right]^{1/4}$$

3.3

$C = 0.6532$, A dimensionless modeling parameter based on $Sc$

$Sc = \frac{\nu}{D} = 366$, Schmidt number

$z_+ = 1$, Valence of a proton

$\nu_+ c_\infty = 1$ M, Total concentration of protons in the bulk electrolyte

$F = 96485$ C mol$^{-1}$, Faraday number

$D = 2.6E-05$ cm$^2$ s$^{-1}$, Effective diffusion coefficient of $H^+$ assuming full dissociation of $H_2SO_4$
$t_+ = 0.814$, Transference number of protons assuming full dissociation of H$_2$SO$_4$

$g = 980 \text{ cm s}^{-2}$, gravitational constant

$\rho_\infty = 1030 \text{ kg m}^{-3}$, density of 0.5 M H$_2$SO$_4$

$\rho_0 = 0.8 \text{ kg m}^{-3}$, density of H$_2$ at standard conditions

$\nu = 0.0095 \text{ cm}^2 \text{ s}^{-1}$, kinematic viscosity of 0.5 M H$_2$SO$_4$

$Sc = \frac{\nu}{D} = 366$, Schmidt number

$L = 2 \text{ cm}$, Length of electrode

Using these parameters, the calculated limiting current density is 3.7 A cm$^{-2}$. The largest current density applied in our experiments was 100 mA cm$^{-2}$, or 2.7% of the calculated limiting current density. The limiting current density is likely higher since electrolyte is also being pumped at a rate of 0.5 mL s$^{-1}$ to the cathode, replenishing protons and decreasing the diffusion boundary layer thickness. Figure 3.8 shows a current voltage (I-V) curve of the electrolyzer. No evidence of mass transport limitations are observed up to the maximum recorded current density of 300 mA cm$^{-2}$. If concentration gradients were present, the slope of the IV curve would become shallower until it eventually reached the mass transfer limiting current.
Figure 3.8: Current-voltage curve for an electrolyzer with a tab insertion length $l_{\text{tab}}$ of 2 mm. 0.5 M H$_2$SO$_4$ was pumped to the cathode at a rate of 0.5 mL s$^{-1}$. The LSV was measured at a scan rate of 20 mV s$^{-1}$.

3.5.3 Calculation of Wagner Number

The Wagner number is defined as the ratio of charge transfer resistance to ohmic resistance. At large overpotentials, the charge transfer resistance can be described using Tafel kinetics, and the Wagner number is defined using Equation 3.4:

$$Wa = \frac{RT\kappa}{\alpha_{CT}F|I_{avg}|l}$$  \hspace{1cm} 3.4
where $R$ is the gas constant, $T$ is the temperature, $\kappa$ is the conductivity of the electrolyte, $\alpha_{CT}$ is the transfer coefficient, $F$ is Faraday’s constant, $i_{avg}$ is the average current density of the electrode, and $l$ is a characteristic length. For a 0.5 M H$_2$SO$_4$ electrolyte, the conductivity $\kappa$ is 0.226 S/cm.$^{50}$

The grouping $\frac{RT}{\alpha_{CT}F}$ can be determined from the slope of a Tafel plot. In a previous study for HER in 0.5 M H$_2$SO$_4$ on a Pt coated Ti mesh, the value of this grouping was determined to be 0.039 V.$^{13}$ The characteristic length of the system was taken to be the length of the electrode, 2.0 cm. The value of $i_{avg}$ used must be normalized to the active area of the electrode, and not the geometric area. For example, a current density of 20 mA cm$^{-2}$ in the main text was calculated by dividing an applied current of 12 mA by the geometric area of the electrode, $0.3 \text{ cm} \times 2.0 \text{ cm} = 0.6 \text{ cm}^2$. Based off of the wire diameter and woven pattern of the mesh electrodes, however, 51% of the electrode area is actually void area. Thus, the corrected value of $i_{avg}$ relative to the active surface area would be 40.8 mA cm$^{-2}$, and the calculated value of $W\alpha$ for these conditions would be 0.11. Figure 3.7 in the main text shows the current distributions for an electrolysis cell operating at 50 mA cm$^{-2}$ of geometric area, and the calculated value of $W\alpha$ is 0.04.

**3.5.4 Description of algorithm for detecting unique bubbles**

For a given image frame taken from a HSV for analysis, the size, position, and velocity of each detected bubble was determined using the Hough transform algorithm (See Appendix A, Section 3.5.1) and the bubble size velocity correlation reported in Figure 3.3a. In order to determine if a bubble appearing in a given frame was already detected in the prior frame, the position of the bubble in the prior frame was first estimated by multiplying its velocity (determined from Figure 3.3a) by the time step between frames. Figure 3.9 shows a visualization of the logic arguments used for the unique bubble detection algorithm. If the calculated bubble position in the prior frame
(N-1) falls within the area of analysis, the bubble is not counted because it was already counted in the prior frame. If the position in frame N-1 falls upstream of the analysis area, then the bubble is counted because it was not present in the previously analyzed frame.

Figure 3.9: Detection of unique bubbles in a single image analysis frame. Green arrows show the path traveled by bubbles from the previous frame to the current frame. Red circles correspond to a previous bubble position which was already in the window of analysis. Black circles correspond to bubbles which were previously out of frame, and thus have not yet been detected.
3.5.5 Relationship between time step and total volume of bubbles detected at 100 mA cm$^{-2}$

![Graph showing total volume of gas detected over time compared to unique volume detected]

Figure 3.10: Total volume of gas detected over the duration of a 10 s HSV as a function of time step between image frames. Experiment was carried out in 0.5 M H$_2$SO$_4$ purged with Ar and pumped at an average velocity of 0.5 cm s$^{-1}$. Bubbles were generated at a current density of 100 mA cm$^{-2}$.

3.5.6 Comparison of the primary and secondary current distributions

The simulated current distribution shown in Figure 3.7 is based on the primary current distribution, which only accounts for nonuniformity due to the ohmic resistance in the electrolyte. In Appendix A, Section 3.5.3, the choice of solving the primary current distribution was justified by calculating $W_a$, which was found to be 0.04. In this case, effects of charge transfer are thought to be negligible.

Figure 3.11 below compares the simulated primary current distribution to the secondary current distribution, which includes the physics for charge transfer at the electrode surface.
The current distribution was modeled for an insulating tab insertion length of 2 mm. The electrolyte was assumed to be 0.5 M H$_2$SO$_4$ with a conductivity 0.226 S cm$^{-1}$. The average current density at the cathode and anode boundaries was fixed to be 50 mA cm$^{-2}$ on a geometric basis. Similar to the primary current distribution simulation, the solution was determined by solving Laplace’s equation for the potential field in the electrolyte, $\nabla^2 \phi = 0$. While constant potential boundary conditions were applied for the primary current distribution, Tafel kinetic boundary conditions were applied for the secondary current distribution. Kinetic parameters were obtained using Tafel analysis from a previous study. At the anode, the oxygen evolution reaction (OER) on Pt was modeled: the equilibrium potential was specified to be 1.23 V, the exchange current density $i_{0,\text{OER}} = 1.87\times10^{-7}$ A m$^{-2}$, and a Tafel slope $\beta_{\text{OER}} = 133$ mV/decade. At the cathode kinetic parameters for the hydrogen evolution reaction (HER) were used: the equilibrium potential was specified to be 0.0 V, the exchange current density $i_{0,\text{HER}} = 6.6\times10^{-4}$ A m$^{-2}$, and a Tafel slope $\beta_{\text{HER}} = -39$ mV/decade. At insulating boundaries the gradient in the potential field normal to the surface was specified to be zero.
Both the primary and secondary simulation curves in Figure 3.11 show similar trends as a function of position along the electrode. As one would intuitively expect, the local current density is highest at the end of the cathode adjacent to the central tab (x/L=0), where the conduction path between the electrodes is shortest and thus the lowest ohmic resistance is expected to be. The primary current distribution can be thought of as a limiting case where the current distribution is most nonuniform. When charge transfer kinetics are included at the operating conditions specified in this report, the current distribution is calculated to be slightly more uniform. Whereas the local current density at the central cathode edge is expected to be $2.86 \cdot i_{avg}$ based on the primary current distribution, it is predicated to be $2.53 \cdot i_{avg}$ (11.5% lower) based on the secondary current distribution.
distribution At the outer edge of the electrode (x/L=1.0), the local current density tapers off to a value of $0.52 i_{avg}$ in the primary current distribution case, and $0.58 i_{avg}$ for the secondary current distribution. Overall, the current distribution is slightly more uniform for the secondary current case, but the differences are small enough to warrant our choice of using the primary current distribution for comparison to experimental measurements in this study.

3.6 Acknowledgements

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3.7 References


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Chapter 4 describes the design and performance of a scalable, stand-alone photovoltaic (PV) electrolysis device used for hydrogen (H₂) production. The electrolyzer component of this device is based on a simple, membraneless design that enables efficient operation with high product purity and without active pumping of the electrolyte. Key to the operation of this PV-electrolyzer is a novel electrode configuration comprised of mesh flow-through electrodes that are coated with catalyst on only one side. These asymmetric electrodes promote the evolution of gaseous H₂ and O₂ products on the outer surfaces of the electrodes, followed by buoyancy-driven separation of the detached bubbles into separate overhead collection chambers. The successful demonstration of this concept was verified with high-speed video and analysis of product gas composition with gas chromatography. While the device based on asymmetric electrodes achieved product cross-over rates as low as 1%, a control device based on mesh electrodes that were coated on both sides with catalyst had cross-over rates typically exceeding 7%. The asymmetric electrode configuration was then incorporated into a standalone, floating PV-electrolyzer and shown to achieve a solar-to-hydrogen efficiency of 5.3% for 1 sun illumination intensity. The simplicity of this membraneless prototype, as characterized by the lack of a membrane, scaffolding, or actively pumped electrolyte, makes it attractive for low-cost production of hydrogen.
4.1 Introduction

Hydrogen is an attractive carbon-free energy carrier that can be sustainably produced by water electrolysis if the electricity is provided by a renewable energy source. The commercial market for electrolyzers that are used to drive water electrolysis are currently dominated by two technologies: polymer electrolyte membrane (PEM) electrolyzers\(^1\) and alkaline electrolyzers.\(^2\) In both technologies, a membrane or diaphragm is positioned between hydrogen (H\(_2\)) and oxygen (O\(_2\)) evolving electrodes and serves a crucial role in device operation by separating product gases while enabling efficient transport of ions between the electrodes. However, membranes can be costly, prone to degradation and failure, and susceptible to cross-over issues.\(^3\) An electrolyzer design that is able to operate with similar energy and collection efficiencies as the conventional technology but without a membrane is highly attractive.\(^4\)\(^-\)\(^7\)

While membrane-free electrolysis has the potential to reduce the capital costs of the electrolyzer, its integration with a renewable energy source must be carefully considered in order for carbon-free H\(_2\) generation from water electrolysis to become commercially viable. In the case of photovoltaic (PV)-electrolysis, integration of PV-cells with the electrolysis cells in a single
device offers an opportunity to reduce the complexity and cost of the system. The current study is motivated by the idea of incorporating PV-electrolysis reactors into a large-scale solar-hydrogen plant that operates on open water where sunlight, water, and non-agricultural space are extremely abundant. Such “solar fuels rigs” might be reminiscent, in some respects, to “deep sea rigs” used to harvest fossil fuels today. Figure 4.1a shows an idealized rendering of a floating solar fuels rig, where PV panels extending from the rig are integrated with low-cost membraneless electrolysis cells, which utilize the solar-generated electricity to split water into oxygen and hydrogen. Although seawater is cheap and abundant, there are substantial barriers to utilizing commercially available, membrane-based electrolyzers for H₂ production from seawater electrolysis. Microorganisms in seawater can lead to biofouling, and trace amounts of magnesium and calcium ions can form hydroxides that clog the pores of the membrane and increase cell resistance. In contrast to conventional membrane-based electrolyzers, a membraneless electrolyzer is expected to have less stringent requirements for electrolyte purity, and thus may be uniquely suited for seawater electrolysis.
Figure 4.1: a) Schematic of a deep sea “Solar Fuels Rig” based on a utility-scale, floating photovoltaic (PV)-electrolysis platform which uses sunlight to generate the solar fuel, H$_2$. On this rig, PV arrays mounted on a floating platform harvest sunlight and produce electricity that is sent to durable membraneless electrolyzers that split water into O$_2$ and H$_2$. Hydrogen generated by the electrolyzers is collected and stored in tanks above the surface where they wait to be shipped back to shore. b) Schematic of novel membraneless electrode assemblies that are the basis of the current study and might one day be key components of the solar fuels rig. Alternating sets of membraneless electrolyzers wired in parallel use electricity supplied from the PV cells to split water into H$_2$ and O$_2$. c) Close up schematic of a passive membraneless electrolyzer employing buoyancy-based product separation. The mesh electrodes are oriented at an angle $\theta$, resulting in an average separation distance $L$. The electrocatalyst is deposited on the outward facing sides of the mesh in order to constrain product gas nucleation and growth to this region only. When the gas bubbles become large enough to detach, they float directly upward for collection or venting to the atmosphere.
In this chapter, we describe and demonstrate a prototype membraneless PV-electrolyzer design (Figure 4.1b, c) that might one day be scaled up and incorporated into a large scale, floating solar fuels rig such as that shown in Figure 4.1a. The primary novelty of this device is that efficient separation and collection of the H\textsubscript{2} and O\textsubscript{2} product gases may be achieved without the use of a membrane, diaphragm, or active pumping of the electrolyte. In previous studies, membraneless separation of product species was demonstrated in electrochemical membraneless flow cells based on flow-by electrodes\textsuperscript{5,21–26} although these microfluidic devices possess significant challenges for scale-up\textsuperscript{27}. More recently, membraneless electrolyzers based on mesh flow-through electrodes have been demonstrated\textsuperscript{6,28}, including by our research group\textsuperscript{4,29}. In all of these previous studies on membraneless electrolyzers, an essential feature was “flow-induced product separation”, whereby convective forces associated with flowing electrolyte continuously remove product species before they can cross-over to the opposing electrode. Thus, the reduced complexity that is achieved by using a membraneless architecture is partially offset by the need to actively pump the electrolyte. By contrast, we show that membraneless electrolyzers based on mesh flow through electrodes can operate in a stagnant electrolyte in the absence of forced advection. Figure 4.1c shows a side-view schematic of a single electrolysis cell consisting of angled mesh electrodes separated by a baffle. Two key features of this device are i.) the slight angle between the electrodes, and ii.) the selective placement of the active electrocatalysts only on the outer surfaces of the mesh electrodes. We herein refer to these electrodes as “asymmetric electrodes”, as opposed to symmetric electrodes with catalyst coated on both sides. Importantly, the product gas bubbles that form on an asymmetric electrode predominately nucleate, grow, and detach from the outer (catalytic) side of the mesh electrodes such that there is minimal cross-over of the product species between the two electrodes. If the two mesh electrodes are oriented vertically and perfectly parallel to each other,
detached bubbles can still cross-over from one electrode to the other by colliding with objects directly above them, such as other bubbles or mesh wires of the electrode itself. However, by positioning the electrodes at a slight angle, separation is improved by facilitating bubbles to float upwards unimpeded. Whereas previous membraneless electrolyzers employed "flow-induced separation", this study presents electrolyzers that utilize asymmetric mesh electrodes for "buoyancy induced separation" in which growing gas bubbles passively detach from the active electrode surfaces and float into separate collection manifolds or chambers.

The electrode configuration utilized in this study consists of titanium mesh electrodes coated with platinum catalyst that was deposited by physical vapor deposition on the outer surfaces of the mesh. These electrodes were integrated into a 3D-printed floating PV-electrolysis device with onboard collection of the generated H₂. The device was fabricated using 3D printing, or additive manufacturing, which has numerous advantages over traditional manufacturing processes and has found increasing use for (photo)electrochemical applications.⁴,³⁰⁻³⁷ Among these advantages are rapid prototyping, efficient materials utilization, and fabrication capabilities that are simply not possible with traditional tools. In this work, we have 3D-printed an electrochemical device from PLA filament, which may be derived from renewable resources, and exhibits reasonable chemical stability for the purpose of prototyping demonstrations in acidic and neutral electrolytes.

In the remainder of this chapter, we present a detailed description of the novel electrode configuration, followed by physical and electrochemical characterization of the electrodes in 2-electrode measurements. For these proof of principle demonstrations, most characterization is performed in 0.5 M H₂SO₄, a commonly employed laboratory electrolyte. Using a combination of high speed video and gas chromatography measurements, product gas cross-over is characterized
under varied electrode angles and operating current densities. After demonstrating the efficacy of passive membraneless electrolysis based on asymmetric flow-through electrodes, this concept is applied to integrated, standalone PV-electrolysis cells. Finally, we discuss challenges and opportunities for application of membraneless electrolyzers for H₂ production from sea water electrolysis.

4.2 Experimental

*Chemicals* – Solutions of sulfuric acid were prepared as described in Chapter 3. pH-neutral electrolytes were prepared using sodium chloride (ACS Reagent grade, Sigma Aldrich), and universal pH indicator (pH 4-10, Sigma Aldrich). Prior to experiments, electrolytes were de-aerated by purging with nitrogen gas.

*Electrode fabrication* – Titanium (Ti) mesh sheets (40 wires per inch; 0.007” wire diameter, Unique Wire Weaving Co., Inc.) were coated with 50 nm thick layers of platinum (Pt) by electron-beam evaporation. Electrodes were designated as “asymmetric” if Pt was deposited on only one side of the mesh sheet, and “symmetric” if Pt was deposited on both sides. The electrodes were cut into rectangles with a width of 1.5 cm and a length of 6.0 cm. When incorporated into the electrolyzer, only the bottom 2.0 cm of the mesh was submerged in electrolyte such that the geometric area of the electrode involved in electrochemical reactions was 3.0 cm².

*Electrolyzer fabrication* – The gas collection lid was printed out of Formlabs Grey photopolymer resin, and was fitted with rubber septa for gas sampling. The electrodes were mounted in the cell through slotted holes in the lid. For volumetric collection experiments, a specialized lid was printed with a viewing window and graduated lines to read the height of the electrolyte level. A specialized lid was also printed to measure the crossover of dissolved species.
electrochemically. For these experiments, a separate electrode slot and guide rail was included to allow for the insertion of an auxiliary sensor electrode.

The body of the floating PV-electrolysis module was assembled from several 3D-printed components: the electrode scaffolding, a PV support raft, and a gas collection lid. The electrode scaffolding was printed out of PLA and contained slits to incorporate two pairs of mesh electrodes. Glass windows were epoxied to the walls of the scaffolding opposite of the electrodes to allow for visual observation of accumulating H₂ gas. The scaffolding also contained connection ports to form a snap fit with the PV support raft. In addition to housing the PV panels, an important function of the PV support raft is to provide floatation for the whole device. Thus, the PV support raft was printed at low density, filling only 15% of the interior volume with PLA. The PV support raft contained insets for four polycrystalline Si PV cells (Sundance Solar Products Inc.), each rated with an open circuit voltage of 3.0 V and a short circuit current of 70 mA. Each panel comprised of six smaller PV cells connected in series which together occupied an area of 14.4 cm². The gas collection lid was printed using Formlabs Tough photopolymer resin and included a port on which a rubber septum was placed to withdraw the collected gas with a syringe.

Device characterization and performance – The electrolyzer and PV panels were characterized using linear sweep voltammetry (LSV) and electrochemical impedance spectroscopy (EIS). All current densities reported are normalized to the 2D rectangular area of the mesh electrodes immersed in the electrolyte (3.0 cm²). For PV-electrolysis experiments, the illumination source was a 500 W halogen work lamp (Utilitech) calibrated to standard air mass (AM) 1.5 intensity using a silicon reference cell (VLSI Standards, SRC-1000-RTD-QZ). During PV-electrolysis experiments, the current was recorded by wiring the potentiostat in series and operating it as a zero resistance ammeter. High-speed videos of bubble evolution from the mesh
electrodes were recorded using the same procedure outlined in Chapter 3. Color videos observing the cross-over of dissolved species were recorded using a Casio EX-ZR700 camera recording at 30 frames per second and a resolution of 1920 x 1080 pixels.

**Analysis of product gas composition** – The collected gas was characterized using gas chromatography (Agilent 7890B GC) with a thermal conductivity detector (TCD) using He as a carrier gas. For experiments with the electrolyzer device, the lid of the cell contained two isolated gas collection chambers located directly above the anode and cathode. Each chamber was fitted with a rubber septum to allow for sampling with a gas-tight syringe (Hamilton). Product cross-over was calculated according to previous reports\(^4,5\) by measuring the amount of H\(_2\) collected in the anode chamber (located above the anode) and dividing by the total amount of H\(_2\) measured in both the anode and cathode chambers. Although the reported quantities of cross-over are for single trials, three additional replicates were conducted for one set of conditions (asymmetric electrodes, 30° separation angle, 40 mA cm\(^{-2}\)), for which an average percent cross-over of H\(_2\) was found to be 2.1% with standard deviation of 0.8%. When operating the floating PV-electrolysis cell, only the headspace above the cathode is analyzed, while O\(_2\) was vented to the laboratory.

4.3 Results and discussion

4.3.1 Description and demonstration of a passive membraneless electrode assembly

The membraneless mesh electrode assembly was first tested in an electrochemical test cell, independent from connection to PV cells, in order to demonstrate the concept of buoyancy induced separation. The body, or base, of this electrolyzer is shown in Figure 4.2a and consists of a rectangular container that holds the electrolyte and has two glass windows epoxied to the front and back of the device in order to allow for side-view *in situ* high speed video imaging of the gas bubbles evolving from the electrodes. The second part of the test cell, which is placed onto the cell
body, is a lid component through which the mesh electrodes are inserted and which contains two 3 mL gas collection chambers positioned immediately above the anode and cathode and separated by a 1 mm wide insulating baffle. The two opposing mesh electrodes span the cell from top to bottom, although electrolysis only takes place in the lower portion of the cell where the electrodes are submerged in electrolyte. Each collection chamber is capped with a rubber septum to seal off the headspace from the surrounding air. Prior to operation, the electrolyte is drawn up into the collection chambers with a syringe. As the electrolysis reaction proceeds, the product gas bubbles float up into the collection chambers and displace the electrolyte.

Figure 4.2: a) Photo of the electrolysis cell with interior components shown through the viewing window. b) Still frame image of bubble growth on an asymmetric electrode operating at 150 mA cm$^{-2}$ in 0.5 M H$_2$SO$_4$. c) Still frame image of bubble growth on a symmetric electrode at the same conditions as in (b). The full videos can be found in the online publication of Int. J. Hydrogen Energy, 43, 1224–1238 (2018).

The basic operating principle of buoyancy-induced separation with an asymmetric mesh electrode assembly was visualized using a high speed video (HSV) camera. Operation of the "asymmetric" electrodes at 150 mA cm$^{-2}$ is shown in Figure 4.2b, where the sides of the electrodes
coated with the Pt electrocatalyst face outwards. So long as the electrolyte is sufficiently conductive and the gap between the two electrodes is small, low solution resistance can be achieved with this electrode geometry because dissolved ionic species—which do not experience buoyancy forces—may easily pass through the holes in the mesh electrode. Figure 4.2b shows that bubble nucleation is primarily confined to the outside faces of the asymmetric electrodes. Once a bubble grows to a size where the buoyancy force exceeds the adhesion forces that keep it attached to the electrode surface, the bubble detaches from the electrode and floats upward into the overhead collection chamber. By this means, a passive membraneless electrolyzer based on asymmetric mesh electrodes can operate with minimal product “cross-over”, which we define here to be a process or event in which product generated at one electrode is transported across the electrode gap and collected in the opposite overhead collection chamber. Figure 4.2c shows a still frame image of bubble formation on a symmetric electrode, which has Pt catalyst deposited on both sides of the electrode. Whereas bubble growth on the asymmetric electrodes is primarily limited to the outside surfaces, bubbles readily grow and detach from both sides of the symmetric electrodes. When bubbles reach the interior region between the electrodes, variations in local convection can cause the bubbles to float up into either collection chamber, resulting in higher cross-over rates and decreased product purity. The HSVs of asymmetric and symmetric electrodes in operation are available in the online publication of Int. J. Hydrogen Energy, 43, 1224–1238 (2018).

The voltage efficiency associated with electrolysis can be determined from the 2-electrode current-voltage (IV) curves. In order for the reaction to proceed, the applied cell voltage must be greater than the difference in the reversible potentials of the oxygen evolution reaction (OER) and hydrogen evolution reaction (HER), ΔE₀, which is equal to 1.23 V at standard conditions when
based on the lower heating value (LHV) of \( \text{H}_2 \) (-237.1 kJ mole\(^{-1}\)). The voltage efficiency of the electrolyzer is equal to \( \Delta E^0 \) divided by the operating voltage. Figure 4.3a compares the IV characteristics of parallel (\( \theta = 0^\circ \)) asymmetric and symmetric electrodes operating in 0.5 M H\textsubscript{2}SO\textsubscript{4}. At an operating current density of 100 mA cm\(^{-2}\), the electrolysis efficiencies for the asymmetric and symmetric electrodes are 50.6% and 53.4%, respectfully. The slightly higher efficiency for the symmetric electrodes can be attributed to the fact that they have twice as much electrochemically active surface area. When accounting for the differences in platinum surface area, the IV-curves for the asymmetric and symmetric electrodes collapse on top of each other (Figure 4.3b). Figure 4.3c shows the IV characteristics for asymmetric electrodes at varying angles of separation. When operating at a current density of 100 mA cm\(^{-2}\), the electrolyzer efficiency decreases to 48.8% for an electrode angle of 30\(^\circ\). This loss in efficiency with increasing electrode separation angle can be attributed to an increase in the solution resistance, which increases as the average electrode separation distance increases. Figure 4.3d shows the ohmic resistance of the cell determined by EIS as a function of electrode separation angle. If the LSVs are corrected for ohmic resistance using these values, the traces collapse into a single curve, as expected (See Appendix B, Section 4.5.1. Figure 4.11).
Figure 4.3: LSV analysis for the passive membraneless electrolyzer for various electrode configurations. Experiments were carried out for a two-electrode setup in 0.5 M sulfuric acid and a scan rate of 100 mV s\(^{-1}\). a) IV curves for asymmetric and symmetric electrodes held at \(\theta=0\degree\). Current density is normalized relative to the geometric area of the mesh electrodes. b) Re-plotting the data in (a) correcting for IR losses and accounting for the fact that the symmetric electrodes have twice as much electrochemically active surface area. c) IV curves for asymmetric electrodes at varying separation angles. The current density is normalized relative to the geometric area of the mesh electrodes. d) Ohmic resistance measured across the cell for asymmetric electrodes as a function of electrode angle. Values for the ohmic resistance were determined using electrochemical impedance spectroscopy (EIS).

4.3.2 Analyzing product gas cross-over

In order to perform water electrolysis safely, every measure must be taken to ensure that an explosive mixture of \(\text{H}_2\) and \(\text{O}_2\) gases is not created. An explosive mixture is formed when \(\text{H}_2\) is present at a concentration of at least 4% in \(\text{O}_2\) (lower flammability limit) or if \(\text{O}_2\) is present at a
concentration of at least 5% in \( \text{H}_2 \) (upper flammability limit). Therefore, it is crucial to demonstrate that the proposed electrolyzer design is able to efficiently collect the product gasses with minimal cross-over of those products into the opposing collection chambers. To quantify cross-over experimentally, the cell was operated at a constant current until \( \approx 3 \) mL of product was collected in the cathode chamber. The collected gas was then analyzed using gas chromatography as described in Appendix B, Section 4.5.2. HSV was also recorded during operation to visually inspect the dynamics of bubble evolution.

Cross-over behavior was systematically studied as a function of electrode angle and applied current density for both symmetric and asymmetric electrodes. Figure 4.4 compares the percent cross-over of \( \text{H}_2 \) into the anode collection chamber for electrolysis experiments run at various electrode angles and current densities. In Figure 4.4a, the percent \( \text{H}_2 \) cross-over values for asymmetric and symmetric electrodes are compared for different separation angles operating at a current density of 20 mA cm\(^{-2}\), where the current density was calculated by dividing the total applied current divided by the 2D area of the immersed electrode. As expected, lower cross-over rates were measured for electrolysis with the asymmetric electrodes at all current densities, and both electrode configurations showed lower rates of \( \text{H}_2 \) cross-over as the electrode separation angle was increased. Figure 4.4b shows the percent \( \text{H}_2 \) cross-over recorded for the asymmetric electrodes in response to the electrode separation angle and applied current density. Overall, the same trend exists where increasing the electrode angle decreases the percentage of \( \text{H}_2 \) cross-over. Interestingly, the percent cross-over was found to decrease with increasing current density over the ranges tested. The lowest percent \( \text{H}_2 \) cross-over achieved for the asymmetric electrodes was 1%, and was recorded at an electrode separation angle of 30° and a current density of 40 mA cm\(^{-2}\). A similar relationship between cross-over, electrode angle, and current density was found for
the symmetric electrodes, with the lowest percent cross-over recorded at 7% (See Appendix B, Section 4.5.2 Figure 4.13).

![Graphs showing percent H₂ cross-over into the O₂ collection chamber of the static electrolyzer measured by gas chromatography.](image)

**Figure 4.4:** Percent H₂ cross-over into the O₂ collection chamber of the static electrolyzer measured by gas chromatography. Experiments were carried out in 0.5 M sulfuric acid at a constant applied current. a) Comparing the percent H₂ cross-over recorded for symmetric and asymmetric electrodes at varying angles and operating at 20 mA cm⁻², where the current density is reported with respect to the geometric area of the electrodes. b) Percent H₂ cross-over measured for asymmetric electrodes at varying electrode angles and current densities.

As indicated by the results in Figure 4.4, asymmetric electrodes can be effective for mitigating cross-over, but they do not eliminate it entirely. The gap spacing in the mesh can still allow small bubbles to pass through, and collisions between rising bubbles combined with bubble-induced local advection can result in lateral motion. Both of these factors can lead to cross-over. By placing the electrodes at an angle to each other, the product bubbles are more likely to float upwards unimpeded without being bumped, for example, by local bubble-induced turbulence into the gap between the two mesh electrodes. Figure 4.5a-c shows schematic side views of three possible mechanisms by which product gas can be transported across the gap between the two angled electrodes. The first case is direct cross-over (Figure 4.5a), whereby a bubble from one electrode travels directly through the opposing mesh electrode and into the opposite overhead
collection chamber. Based on HSV analysis, direct cross-over was seen to occur most frequently when the electrodes were parallel to each other, but was very rarely observed as the angle between the electrodes was increased to 5° or larger. A second mechanism by which cross-over can occur was observed for electrode angles up to 15° and involved indirect cross-over through the formation of a large, merged bubble that becomes pinned on the central baffle that divides the two mesh electrodes (Figure 4.5b). Depending on the electrode type and angle, bubbles of this nature were observed to grow for long time periods before randomly detaching into either of the two adjacent collection chambers. When the asymmetric electrodes are operated at a 30° angle, neither the direct nor indirect bubble cross-over is observed. However, cross-over can still occur in the form of dissolved gas diffusion (Figure 4.5c). Cross-over of dissolved gas is most likely to occur after long periods of operation when the electrolyte becomes saturated with dissolved gas, H$_2$(aq) and O$_2$(aq), which can diffuse freely throughout the cell. The cross-over of dissolved gas is also assisted by the local convection fields formed by the detached bubbles as they float upwards. When the dissolved gas reaches the opposite side of the cell from which it formed, it can equilibrate with the headspace above that chamber and/or the bubbles in the opposite chamber that will eventually float up into the headspace. Although high speed video can be used to observe direct bubble cross-over (Figure 4.5d) and indirect bubble cross-over (Figure 4.5e), it cannot directly observe cross-over of dissolved H$_2$(aq) and O$_2$(aq).
Figure 4.5: Mechanisms of product gas cross-over in a static fluid electrolyzer. a) Direct bubble cross-over: bubbles from one electrode detach and migrate to the opposite collection chamber. b) Indirect bubble cross-over: bubbles from both electrodes accumulate on the central baffle, forming a mixed gas bubble which can detach and travel into either collection chamber. c) Dissolved gas cross-over: the electrolyte becomes saturated with dissolved product gas which can diffuse over to the opposing electrode. The dissolved gas can equilibrate with bubbles as they float upwards to the collection chamber. For simplicity, dissolution and diffusion was only shown for dissolved H₂. Still frame images taken from high speed video measurements showing instances of d) direct bubble cross-over recorded for parallel asymmetric electrodes and e) indirect bubble cross-over recorded for asymmetric electrodes at a 30 degree angle and 100 mA cm⁻².

In order to visualize cross-over by dissolved H₂ and O₂, electrolysis was conducted in a pH neutral 0.5 M NaCl electrolyte containing 17 % vol. universal pH indicator. The unbuffered, pH-neutral solution allows for visualization of the spatio-temporal evolution of the dissolved-product species because the HER and OER reactions lead to the build-up of hydroxyls at the cathode and
protons at the anode, respectively. When exposed to an alkaline pH, the universal pH indicator dye quickly turns dark purple, and by tracking the movement of this purple plume it is possible to monitor the progression of the dissolved reduction products as they migrate away from the cathode. Technically, the color change is directly tied to OH\(^-\) concentration, but it is a reasonable assumption in these experiments that the plume of H\(_2\) (aq) closely overlays that of the generated hydroxyls. Figure 4.6 contains side-view photographs that show the progression of the alkaline product plume as it spreads away from an asymmetric mesh cathode while operating at 40 mA cm\(^{-2}\) and an electrode separation angle of 30°. Within 10-30 seconds of the start of electrolysis, the product plume reaches the anode, proving that dissolved species transport between the two angled electrodes can occur at reasonably high rates within the time scales of the experiments in Figure 4.5. At these time and length scales, transport must be dominated by convection that results from bubble-induced fluid flow within the electrolyte.
Figure 4.6: Visualization of dissolved species cross-over during electrolysis in 0.5 M NaCl solution with universal pH indicator. The indicator turns from red to dark purple in the presence of the hydroxyls generated at the cathode. The experiment was carried out with asymmetric electrodes operating at a current density of 40 mA cm$^{-2}$.

The pH indicator dye experiment provides strong evidence that the cross-over of dissolved gas is likely responsible for the H$_2$ signal measured by GC in the anode compartment after operation at lower current densities. Cross-over of dissolved H$_2$(aq) may also explain the trend of decreasing cross-over rates with increasing current density that was reported in Figure 4.4b. When lower current densities are applied, a larger fraction of the product gas remains dissolved in the aqueous phase. Operating at a lower current density also requires a longer time for the collection chambers to fill up. While the time to fill the chamber during electrolysis at 40 mA cm$^{-2}$ was only 4 minutes, experiments at 10 mA cm$^{-2}$ required 16 minutes. Longer experiments allow more time for gas to saturate the electrolyte, distribute across the electrolyte reservoir, and equilibrate in the
opposite collection chambers. However, the rate of dissolved gas cross-over is limited by the solubility of the gasses in solution. Thus, operating at high current density at steady state can result in a higher fraction of the products entering the gas phase (i.e. bubbles) rather than the dissolved phase. Additionally, the rate of equilibration between the dissolved gasses and the headspace can be minimized by further optimizing the geometry of the electrolysis cell. For example, the contact area between the electrolyte and the headspace could be decreased. If the electrolyzer was operating in an open body of water, the dissolved anode products would also be able to diffuse away from the device, as opposed to being constrained to the enclosed reservoir of the current design.

In addition to equilibrating with the opposing collection chamber, it is also possible for the dissolved gasses to participate in a back reaction. For example, H₂ evolved at the cathode can be transported over to the anode, where it is oxidized back into protons through the hydrogen oxidation reaction (HOR). Similarly, O₂ evolved at the anode can be reduced back to water at the cathode through the oxygen reduction reaction (ORR). Both of these so-called “back reaction” processes result in lower Faradaic efficiencies for the OER and HER reactions, thereby decreasing electrolysis efficiency. Fortunately, the magnitude of the back reactions is limited by the solubility and mass transport of H₂ and O₂ gasses across the electrolyte gap between the anode and cathode. In order to quantify the rate of the HOR back reaction at the anode used in the passive membraneless electrolyzer, an auxiliary Pt-coated mesh electrode was used as an electrochemical sensor. As illustrated in Figure 4.7a, this sensor electrode was positioned in between the anode and cathode mesh electrodes and held parallel to the anode electrode. During electrolysis, the sensor electrode was held at a potential of 0.8 V vs. Ag/AgCl, where it has ample overpotential to oxidize H₂ crossing over from the cathode, but the overpotential is insufficient to reduce O₂.⁴
means, the auxiliary electrode can serve as a H₂ sensor that can selectively and quantitatively measure the flux of dissolved H₂ that would normally be oxidized at the OER anode.

Figure 4.7: a.) Schematic of a modified electrolysis cell involving a sensor electrode that is used to measure current efficiency losses due to the HOR of H₂ at the anode. b) Average losses in current efficiency due to HOR during constant current electrolysis while holding the applied potential of the sensing electrode at +0.8 V vs. Ag/AgCl. The current efficiency loss was calculated by dividing the total integrated charge recorded by the sensing electrode by the total charge passed between the anode and cathode throughout the electrolysis experiment. Measurements were recorded for a 30° electrode angle in 0.5 M H₂SO₄.

Back reaction experiments utilizing the auxiliary electrode were performed using asymmetric electrodes angled at 30⁰ and operated at constant current densities of 20 mA cm⁻², 30 mA cm⁻², and 40 mA cm⁻² until the H₂ collection chamber was filled. An Ag|AgCl reference electrode was also present in the experimental set-up, and a bipotentiostat was used to apply a constant potential of +0.8 V vs. Ag|AgCl to the sensor electrode while water electrolysis was carried out between the anode and cathode of the electrolyzer cell. The results of the back reaction experiments are shown in Figure 4.7b, where the Faradaic efficiency loss due to oxidation of H₂
that crossed-over from the cathode is plotted as a function of the operating current density of the electrolysis cell. This Faradaic efficiency loss is given by the percentage of H₂ lost due to the HOR back reaction, and was calculated by dividing the net charge passed through the sensing electrode by the net charge passed between the anode and cathode over the duration of the experiment. As expected, the largest fractional loss of H₂, 3.5%, was measured for electrolysis at the lowest operating current density, 10 mA cm⁻², for which the lower rate of H₂ generation allows a higher percentage of the generated H₂ to dissolve into the electrolyte and oxidize at the sensor electrode. As the current density increased to 40 mA cm⁻², the percent loss of H₂ through the back reaction decreased down to 1.5% as a larger fraction of the generated H₂ was converted into gas bubbles that were soon collected in the overhead collection chambers. Depending on the relative prices of electricity and H₂, back-reaction losses of 1.5%-3.5% can be tolerable in real devices if electricity prices and capital costs of the membraneless system are sufficiently low.⁷ Although these experiments only quantified the losses in Faradaic efficiency due to H₂ oxidation at the cathode, it is expected that the losses from oxygen reduction at the anode would be approximately the same magnitude due to the similar transport pathways, solubility and diffusivity of aqueous O₂.

Beyond efficiency losses due to back reaction, bubble cross-over can also impact device performance by increasing the ohmic resistance for ion transport between the two electrodes. When gas phase bubbles enter the spacing between electrodes, they behave as insulators which inhibit ionic conduction.³⁹ Fortunately, the bubbles largely remain on the outside of the asymmetric electrodes where they do not significantly affect ion conduction between the two electrodes.

4.3.3 Measuring product collection efficiencies

The product collection efficiency can be defined as the percentage of generated H₂ that is successfully collected and removed from the electrolyzer. The collection efficiency can be less
than 100\% for several reasons, including dissolution losses, back reaction losses, and losses associated with \text{H}_2 bubbles that never reach the desired collection chamber or apparatus. In this study, the \text{H}_2 collection efficiency of the membraneless electrolyzer was determined by measuring the volumetric rate of \text{H}_2 collected in the overhead collection chambers. These experiments were performed using a specialized lid that was printed with graduated lines to read the height of the electrolyte displaced during electrolysis (Appendix B, Section 4.5.3 Figure 4.14). Figure 4.8a shows the recorded volume of \text{H}_2 and \text{O}_2 collected in the lid as a function of time during a constant current electrolysis at 40 mA cm\(^{-2}\) with asymmetric electrodes held at a separation angle of 30\(^\circ\). The error bars designate a 95\% confidence interval around the average time of collection over three separate trials. As predicted from the stoichiometry, the relative rates of \text{H}_2 and \text{O}_2 collection are at approximately a 2:1 ratio. The solid lines on the plot represent the theoretical rate of volumetric generation predicted from Faraday’s Law and the ideal gas law. When comparing the slopes of the experimental and predicted data, the volumetric collection efficiencies for \text{H}_2 and \text{O}_2 were found to be 83.0\% and 83.8\%, respectively.

![Figure 4.8](image)

**Figure 4.8:** a) Volume of product gas collected as a function of time. Data was recorded for a 30\(^\circ\) electrode separation angle and an operating current density of 40 mA cm\(^{-2}\). The error bars correspond to a 95\% confidence interval for data averaged over three trials. The solid lines represent the theoretical volumetric collection based on Faraday’s law and the Ideal gas law. b)
Mole balance on H\(_2\) generated during electrolysis with an electrode angle of 30\(^\circ\) and operating at 40 mA cm\(^{-2}\) for 3.9 minutes.

By knowing the product collection efficiency, it is furthermore possible to close the mass balance on the H\(_2\) generated by the membraneless electrolyzer. Figure 4.8b shows the resulting balance of H\(_2\) for electrolyzer operation at 40 mA cm\(^{-2}\), which includes the collected H\(_2\), H\(_2\) lost to the back reaction, H\(_2\) loss to cross-over to the anode collection stream, and “uncollected H\(_2\)“ that is still present in the system but did not reach the collection chamber. The percent H\(_2\) collected was determined directly from the volumetric collection efficiency experiments described above, and the percent H\(_2\) loss due to back reaction was determined from the results of the sensor electrode experiments shown in Figure 4.7b. The percent H\(_2\) loss due to cross-over was determined from the gas chromatography experiments shown in Figure 4.4b. From the analysis shown in Figure 4.8b, 14.1\% of the generated H\(_2\) predicted from Faraday’s law remains uncollected. The uncollected H\(_2\) can be primarily attributed to H\(_2\) lost to dissolution in the liquid electrolyte and, to a smaller extent, H\(_2(g)\) bubbles that remain uncollected in the electrolysis cell. Based on the net amount of H\(_2\) generated during the electrolysis experiment, the volume of electrolyte within the cell, and the solubility limit of H\(_2\), up to 12\% of the generated H\(_2\) may have been dissolved in the electrolyte. After experiments, some bubbles remained adhered to the reactor walls and sides of the electrodes, likely accounting for the remaining \(\approx\)2 \% of uncollected H\(_2\). For longer operating times over which large amounts of H\(_2\) are produced, it is expected that the uncollected H\(_2\) losses will go towards 0\% because dissolved H\(_2\) and uncollected gas bubbles will be very small compared to the total amount of H\(_2\) generated. In this case, the collection efficiency should only be limited by the back reaction and dissolved gas cross-over to the opposite gas collection chamber.
4.3.4 Demonstration of a floating PV-electrolysis module

Having demonstrated that buoyancy driven product separation is possible with acceptable rates of cross-over, we devote the remainder of the discussion to integrating the membraneless design into a bench scale floating PV-electrolysis module. As illustrated schematically in Figure 4.9a, this module contains two sets of asymmetric electrodes angled at 30°. The electrodes were wired parallel to each other and connected to four parallel-connected c-Si PV panels, each of which consisted of 6 sub-cells that are connected in series. A wiring diagram of the PV panels and electrolyzers is shown in the Appendix B, Section 4.5.4 Figure 4.15. The entire module was then placed in a 1 L reservoir of 0.5 M H₂SO₄ (Figure 4.9b), where it floats on the surface of the electrolyte thanks to the high buoyancy of the void-containing 3D-printed raft. The purity of the H₂ collected in the interior reservoir was sampled through a sealed port for GC analysis after completion of the electrolysis experiment.
Figure 4.9: a) Schematic side-view of the floating PV-electrolysis module, which is based on two sets of asymmetric mesh electrodes that are wired in parallel to each other and to four PV panels. The module floats in a reservoir of sulfuric acid. Hydrogen evolved at the cathode is collected underneath the PV panels while the oxygen is allowed to vent to the atmosphere. b) Photo of the PV-electrolyzer module floating in a sulfuric acid reservoir. c) I-V curve matching for the electrolyzer and PV cells. The electrolyzer curve is for 6 cm$^2$ of asymmetric mesh electrodes submerged in 0.5 M sulfuric acid and was recorded at a scan rate of 10 mV s$^{-1}$. The PV curve is the combined I-V response for four PVs wired in parallel under a lamp calibrated to the AM 1.5G intensity, and was recorded at a scan rate of 100 mV s$^{-1}$. The intersection of these curves predicts the operating current of the device when the PV panels and electrolyzer are connected to each other. d) Operating current of the floating PV-electrolysis unit as a function of time during unassisted water electrolysis under the same illumination conditions as described for the measurements in c.).

Figure 4.9c shows the predicted performance of the PV-electrolyzer, which can be estimated from the intersection of the individually tested IV curves of the electrolyzer and the PV panels. 

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For these measurements, the two pairs of asymmetric electrodes were connected in parallel, and the four PV panels were also connected in parallel. The performance of the four PV panels connected in parallel was measured under illumination by an unfiltered halogen work lamp with irradiance calibrated to match the solar AM 1.5 power intensity (100 mW cm\(^{-2}\)). The resulting IV curve is characterized by an open circuit voltage (\(V_{oc}\)) of 3.127 V, a short circuit current (\(J_{sc}\)) of 256.2 mA, a fill factor (FF) of 72.1%, and a photovoltaic efficiency (\(\eta_{PV}\)) at max power point of 10.0 %. \(\eta_{PV}\) was calculated based on the cumulative PV cell area of 57.6 cm\(^2\) and incident light intensity of 100 mW cm\(^{-2}\). The 2-electrode IV curve for the two pairs of asymmetric electrodes shows that water electrolysis begins to take place above an applied voltage of \(\approx 1.7 \text{ V}\), followed by an exponential rise in current as voltage is increased further. From the intersection of the two IV curves (\(I= 238.3 \text{ mA}, V= 2.263 \text{ V}\)), the electrolyzer efficiency (\(\eta_e\)), coupling efficiency (\(\eta_c\)), and the solar–to-hydrogen (STH) efficiency (\(\eta_{STH}\)) may be determined. At the predicted operating voltage, the electrolyzer efficiency is calculated to be \(\eta_e = 54\%\). The coupling efficiency, defined as the device power at the operating point divided by the max power that can be produced by the PV cells,\(^{40}\) is calculated to be 98\%. This indicates that the PV and electrode arrangements are almost perfectly matched. Combining the PV, electrolysis, and coupling efficiencies, the solar-to-hydrogen conversion efficiency (\(\eta_{STH}\)) is predicted to be \(\eta_{STH} = \eta_{PV}\eta_e\eta_c = 5.3\%\). It should be noted that \(\eta_{PV}\) was calculated based on the area of the PV cells, not the area of the entire PV-electrolysis module. In an optimized PV-electrolyzer, the percentage of area occupied by PV panels would be maximized such that the PV area is similar to the module area.

After measuring the IV curves for the individual components, the floating PV-electrolysis module was tested under continuous illumination in 0.5 M H\(_2\)SO\(_4\) for 75 min. Figure 4.9d shows the measured operating current of the floating PV-electrolysis device as a function of time,
revealing stable operation. The average current of the device was found to be in good agreement with the predicted operating point in Figure 4.6c. During operation, H₂ was collected underneath the central lid while O₂ was allowed to vent to the atmosphere. After filling the collection chamber, the gas was analyzed with GC. After subtracting out O₂ signal associated with air (based on an N₂:O₂ ratio of 79:21), the purity of the collected product gas was determined to be 98% H₂.

4.3.5 Challenges for Seawater Electrolysis

Although this study has demonstrated that a membraneless, floating PV-electrolysis module can generate and collect H₂ with reasonable efficiency, many challenges for seawater electrolysis must still be overcome before the solar fuels rig concept in Figure 4.1 can be realized. Future demonstrations of devices for seawater electrolysis should report on their stability in electrolytes that more realistically simulate seawater. Laboratory electrolytes are typically prepared from reagent grade NaCl, H₂SO₄, and 18 MΩ DI water, and are free of contaminants which could cause scaling or biofouling by micro- or macro-organisms. Biofouling, or the adhesion of micro- or macro-organisms on a surface, has been found to be highly detrimental to membranes in reverse-osmosis (RO)-based desalination devices. A simple way to overcome this deficiency in an electrolysis device is to operate without a membrane, although the electrodes themselves may be prone to biofouling as well. However, there is evidence to suggest that electrochemical reactions at the anode can generate reactive oxides and hypochlorites that act as a protective antimicrobial agents, and a recent study by our group has demonstrated the possibility of designing “impurity tolerant” catalysts for water splitting. Inorganic scaling can also degrade the performance of the electrodes, with Mg(OH)₂ and Ca(OH)₂ precipitating out of the electrolyte in alkaline conditions and blocking electrochemically active sites. For proper operation of Nafion membranes in chlor-alkali cells, brine purification is required to reduce the concentration of magnesium and
calcium ions below 50 parts per billion.\textsuperscript{50} by comparison seawater typically contains \(\approx 1300\) ppm of \(\text{Mg}^{2+}\) and \(\approx 400\) ppm of \(\text{Ca}^{2+}\).\textsuperscript{51}

In addition to challenges with long term stability, seawater electrolysis faces a number of efficiency drawbacks when compared to concentrated acid or alkaline electrolysis. The electrolysis of a pH neutral electrolyte tends to have more sluggish kinetics,\textsuperscript{52} and additional thermodynamic penalties arise from pH gradients that form across the cell as the reaction proceeds.\textsuperscript{53} Natural seawater is less conductive than acidic solutions such as 0.5 M H\textsubscript{2}SO\textsubscript{4}, which results in a larger solution resistance. All of these factors require that a larger cell voltage be applied between the electrodes to achieve a given current density, which generally requires more PV cells to be connected in series and thereby decreases the solar-to–hydrogen conversion efficiency.\textsuperscript{40}

In order to get an estimate of typical efficiency losses for electrolysis in seawater, 2-electrode IV curves were measured with parallel asymmetric electrodes operating in 0.5 M H\textsubscript{2}SO\textsubscript{4} and a synthetic seawater comprised of 0.6 M NaCl. The salinity of seawater is typically around 3.5\% by mass,\textsuperscript{54} making 0.6 M NaCl a simple electrolyte with similar conductivity to seawater. These IV curves are provided in Figure 4.10, showing that significantly larger voltages are required to achieve the same electrolysis current density in the NaCl solution compared to the sulfuric acid solution. For example, voltages of 2.44 V and 3.68 V are required to operate at a current density of 100 mA cm\textsuperscript{-2} in the 0.5 M H\textsubscript{2}SO\textsubscript{4} and 0.6 M NaCl solutions, respectively. This increase in operating voltage of 1.24 V results in a decrease in the electrolysis efficiency from \(\eta_e = 50.4\%\) in 0.5 M H\textsubscript{2}SO\textsubscript{4} and \(\eta_e = 33.4\%\) in 0.6 M NaCl. Based on the measured solution resistances and electrode arrangement, 0.41 V of this extra voltage requirement can be attributed to the increased solution resistance of the synthetic seawater electrolyte. More voltage is also required to overcome the Nernstian penalty that arises from the pH gradients that develop at the anode and cathode. If
we take the local pH at the anode and cathode to be 3 and 11, respectively (based on the experiments in Figure 4.6), this would mean that the local activities of protons at the anode and hydroxyls at the cathode would be $\approx 1 \times 10^{-3}$ M. Plugging these values into the Nernst Equation and assuming local partial pressures of 1 atm for $H_2$ and $O_2$ at the cathode and anode, respectively, we estimate that $\Delta E^o_{\text{cell}}$ increases from 1.23 V to 1.70 V, consistent with literature.$^{53}$ Finally, we estimate that the increase in kinetic losses incurred during electrolysis at 100 mA cm$^{-2}$ in the 0.6 M NaCl is $\approx 0.36$ V, obtained as the difference between the total change in required voltage (1.24 V) and the additional voltage needed to compensate for the lower solution conductivity (0.41 V) and higher thermodynamic cell voltage (0.47 V).

\[ \text{Figure 4.10: IV characteristics of parallel asymmetric electrodes in 0.5 M H}_2\text{SO}_4 \text{ and 0.6 M NaCl recorded at a scan rate of 100 mV s}^{-1}. \]

Another important consideration for electrolysis in seawater is the presence of chloride ions. Chloride ions in seawater can be particularly corrosive, and are capable of permeation
through protective layers.\textsuperscript{55} Additionally, the oxygen evolution reaction (E\textsuperscript{o} = +1.23 V NHE) must compete with the evolution of chlorine (Cl\textsubscript{2}) gas (E\textsuperscript{o} = +1.36 V NHE). Cl\textsubscript{2} has commercial value, being produced by the Chlor-alkali process at 50-60 mton per year globally,\textsuperscript{56} so there may be some interest in capturing Cl\textsubscript{2} as a value-added side product during seawater electrolysis. However, it is important to note that Cl\textsubscript{2} is a toxic chemical that introduces health and environmental risks.\textsuperscript{57}

In the present PV-electrolyzer design, the O\textsubscript{2} product generated at the anode is simply vented to the atmosphere, but this would not be acceptable in the case of Cl\textsubscript{2}. If it were desirable to avoid Cl\textsubscript{2} evolution, developing an anode catalyst that would be selective for the OER over Cl\textsubscript{2} evolution would be one solution. Fujimera et al. have demonstrated MnO\textsubscript{2} OER electrocatalysts capable of 100\% faradaic efficiency during electrolysis in an alkaline NaCl solution, although it’s behavior was not reported for an acidic electrolyte that would be encountered at the anode during seawater electrolysis.\textsuperscript{58}

Besides developing catalysts that would be compatible with seawater, many other challenges would need to be overcome at the device and plant-level. The corrosive environment of seawater also poses problems for other components of a solar fuels rig (piping, storage vessels, solar panels, pumps, etc), and there are many open questions relating to process engineering and rig construction. What sort of rig size makes economic sense, and at what pressure should the H\textsubscript{2} be stored? Should the rig be anchored to the sea floor, or be a free floating vessel that could move between markets and out of the way of storms? As the concept of seawater electrolysis with stand-alone rigs gains more and more attention, engineering system- and plant-level solutions will be of great importance.
4.4 Conclusions

This study has demonstrated a novel electrode architecture for water electrolysis whereby high purity (≈99% pure) H\textsubscript{2} can be produced without a membrane and without the actively pumped electrolyte. Instead, this design relies on the use of buoyancy-driven separation of H\textsubscript{2} and O\textsubscript{2} gas bubbles that are generated at so called “asymmetric” mesh electrodes with active catalyst deposited onto only their outer surfaces. This arrangement is scalable and enables efficient electrolysis with minimal ohmic resistance losses so long as the electrolyte is sufficiently conductive. By systematically changing the mesh electrode angle and operating current density, the rate of product gas cross-over between the anode and cathode chambers was observed qualitatively using high speed video and quantitatively using gas chromatography. It was shown that the asymmetric mesh electrodes significantly decrease H\textsubscript{2} cross-over rates under all conditions that were tested, with the lowest percent H\textsubscript{2} cross-over found to be 1%. For cases when no direct cross-over of bubbles was observed, \textit{in situ} videos of electrolysis in the presence of color-changing dyes indicated that the convective transport of dissolved H\textsubscript{2} between chambers is the primary mechanism of product cross-over. This study also showed that this membraneless electrode architecture can be easily integrated into a 3D printed, floating PV-electrolysis module, which was used to produce H\textsubscript{2} with a solar-to-hydrogen efficiency of 5.3% in 0.5 M H\textsubscript{2}SO\textsubscript{4}. This demonstration was inspired by the idea of seawater electrolysis, which represents an intriguing approach to the renewable generation of fuel using non-agricultural space and non-drinking water. However, significant challenges in catalysis, reactor design, and plant design will need to be overcome for economical seawater electrolysis to become a reality.
4.5 Appendix B

4.5.1 *iR-corrected IV characteristics of membraneless electrolyzers*

Figure 4.11: iR-corrected IV curves measured for asymmetric Pt/Ti electrodes in 0.5 M H₂SO₄. The curves presented are from the data set as shown in Figure 4.3c, which were shifted to lower voltages by subtracting the iR voltage drop at each current density using the series resistance values obtained from electrochemical impedance spectroscopy (EIS) and provided in Figure 4.3d. IV curves were recorded at 100 mV s⁻¹.

4.5.2 *Calculating the percent cross-over of H₂*

A schematic of the experimental setup used for collection of gasses is shown in Figure 4.2a in the main text. GC measurements were performed using a gas chromatograph using a thermal conductivity detector (TCD) and He as a carrier gas.
Figure 4.12: Sample chromatograph showing the H₂ peak on both the cathode and anode side collected for symmetric electrodes held at 30 degrees and an operating current density of 20 mA cm⁻² in 0.5 M H₂SO₄.

For each chromatograph, the H₂ peak (retention time ≈ 12.11 min) was integrated to obtain the area under the curve, a measure of the total signal associated with the detected H₂. The percent cross-over of H₂ from the cathode to the anode compartment was calculated by dividing the integrated area of H₂ (A_{H₂}) in the anode chamber by the sum of the integrated areas of H₂ recorded in both chambers (Equation B4.1). This procedure follows the method described by Hashemi et al.:⁵

\[
H₂ \text{ Crossover} = \frac{A_{H₂}(\text{Anode Chamber})}{A_{H₂}(\text{Anode Chamber}) + A_{H₂}(\text{Cathode Chamber})} \tag{B4.1}
\]
Figure 4.13: Percent H$_2$ crossover into the O$_2$ collection chamber of the membraneless electrolyzer measured by gas chromatography. Experiments were carried out in 0.5 M sulfuric acid at a constant applied current and using symmetric Pt/Ti electrodes.

4.5.3 Lid used for volumetric collection efficiency experiments

Figure 4.14: Lid used to record volumetric collection of product gasses. Front end contains epoxied windows with graduated lines reading the level height in 2 mm increments. At the beginning of the experiment, electrolyte is drawn up with a syringe to the top line. As the electrolysis reaction proceeds, gas bubbles displace the electrolyte and the liquid level drops.
4.5.4 Circuit diagram for the PV electrolysis device

Figure 4.15: Schematic of an individual PV panel and its connection to the floating PV electrolysis module. a) A photograph of an individual PV panel, which consists of 6 individual PV cells that are wired in series. b) A total of four PV panels are wired together in parallel. The cathode and anode terminals are connected to two parallel sets of angled electrolyzers. An ammeter (A) measures the current between the positive terminals of the PV panels and the electrolyzers during operation. A schematic of the PV electrolysis device is shown in Figure 4.9a of the main text.

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4.7 References


CHAPTER 5
LIMITING PHOTOCURRENT ANALYSIS OF A WIDE CHANNEL
PHOTOELECTROCHEMICAL FLOW REACTOR

In this chapter, we extend our analysis of reactor designs to photoelectrochemical (PEC) cells – semiconductor devices which are able to both directly capture sunlight and drive an electrolytic process. The development of efficient and scalable PEC reactors is of great importance for the eventual commercialization of solar fuels technology. This chapter presents a systematic study of the influence of convective mass transport and light intensity on the performance of a 3D-printed PEC flow cell reactor based on a wide channel, parallel plate geometry. Using this design, the limiting current density generated from the hydrogen evolution reaction at a p-Si metal-insulator-semiconductor (MIS) photocathode was investigated under varied reactant concentration, fluid velocity, and light intensity. Additionally, a simple model is introduced to predict the range of operating conditions (reactant concentration, light intensity, fluid velocity) for which the photocurrent generated in a parallel plate PEC flow cell is limited by light absorption or mass transport. This model can serve as a useful guide for the design and operation of wide-channel PEC flow reactors. The results of this study have important implications for PEC reactors operating in electrolytes with dilute reactant concentrations and/or under high light intensities where high fluid velocities are required in order to avoid operation in the mass transport-limited regime.
5.1 Introduction

Photovoltaic-electrochemical (PEC) cells are integrated, all-in-one devices that offer a promising means of converting abundant but intermittent solar energy into storable fuels such as hydrogen (H₂). Because a PEC is able to simultaneously harvest sunlight and drive the electrolysis process as a single device, there is opportunity to lower the capital costs for solar hydrogen production. To date, most research in the PEC field has focused on developing materials for stable and efficient photoelectrodes, the “engines” of PEC devices that drive the solar-to-fuel conversion process. These efforts have resulted in significant improvements in photoelectrode performance, with multiple demonstrations of high efficiency (> 8% solar-to-H₂ conversion efficiency) and long term operation (> 100 hours). These results should be applauded, but it is important to recognize that the photoelectrode is only one component of a commercial PEC device. In the same sense that a high performance sports car consists of much more than an engine, a high performance PEC device is much more than its photoelectrode(s). Like a car engine, the photoelectrode must be effectively incorporated into an affordable chassis, or reactor, that effectively facilitates reactant delivery and product removal while minimizing efficiency losses and maximizing operating life.
As photoelectrode technology continues to improve, the task of developing efficient and scalable PEC reactors will become even more crucial to the commercialization of PEC technology.\(^{10}\)

The majority of photoelectrode-based PEC reactor designs that have been reported in literature can be generally classified into one of three broad categories: batch, semi-batch, or continuous. Most bench scale PEC device designs demonstrated to date are operated as semi-batch reactors under 1 sun illumination intensity in a stagnant (non-pumped, non-stirred) electrolyte that may be resupplied or topped off on an as needed basis.\(^{11–17}\) Less commonly explored are PEC continuous flow reactors, in which electrolyte is continuously and actively pumped through the reactor.\(^{18–22}\) PEC flow cells, or reactors, such as that illustrated in Figure 5.1 can offer several advantages over passive semi-batch PEC reactors. First, flow cells can enable more efficient PEC operation by enhancing transport of reactants to the photoelectrode surface.\(^{23}\) This benefit is particularly important for reactions involving low concentrations of reactants such as CO\(_2\), which has a solubility limit in water of only 34 mM at 298 K.\(^{24}\) Similarly, flowing electrolyte can enhance PEC performance by facilitating removal of gaseous product species (i.e. bubbles) that would otherwise accumulate on the photoelectrode surface and decrease its efficiency. Both of these benefits are highly useful, and in many cases required, for PEC operation under concentrated solar illumination whereby photocurrents may exceed the maximum rate of reactant mass transport even for concentrated electrolytes. In solar concentrating reactors, lenses or mirrors are employed to increase the light intensity incident on the photoelectrode and thereby reduce its required area (and associated cost) for a given rate of product generation.\(^{5,6,25}\) Seger \textit{et al.} have demonstrated that flowing electrolyte can be beneficial in a parallel plate-like proton exchange membrane PEC cell,\(^{18,26}\) but there has been a lack of quantitative analysis on the operating limits of PEC flow reactors.
In order to design efficient PEC flow cells and establish their performance limitations under various operating conditions, it is instructive to review the basic processes taking place at and near the photoelectrode surface. Figure 5.1 shows the processes governing the reaction rate at a photoelectrode operated in a PEC flow cell based on a wide channel geometry characterized by a channel height to width ratio $\frac{h}{w} \ll 1$, which is the basis of this work. Electrolyte flows in a narrow channel from left to right between a glass window (top surface) and the photoelectrode (bottom surface), with the counter electrode placed downstream (Figure 5.1a). Under flowing electrolyte conditions, a diffusion boundary layer with thickness $\delta$ develops at the surface of the photoelectrode (Figure 5.1b). For a given set of operating parameters, namely the fluid velocity, bulk reactant concentration, applied electrode potential, and illumination intensity, the reaction rate and associated current density at the photoelectrode surface are determined by the following steps: i) diffusion of reactants from the bulk electrolyte across the boundary layer, ii) adsorption of the reactant onto the electrode surface, iii) absorption of photons to generate electron hole pairs (excitons), iv) collection of the photogenerated minority carriers at the photoelectrode surface, v)
The primary objective of this study is to establish a quantitative understanding of the limiting photocurrent behavior for a photoelectrode operated in a wide channel flow cell under photo- or mass transport-limiting conditions. Such conditions are achieved when sufficient potential is applied or photovoltage generated to ensure that the photoelectrode current is not limited by reaction kinetics. Thus, this study does not delve into the detailed effects of flowing electrolyte on kinetics, but seeks to quantitatively understand the upper-limit of photocurrent that can be generated when kinetics are not limiting. As a basis for this study, water electrolysis was performed using a p-Si photocathode possessing a metal-insulator-semiconductor (MIS) architecture. With this configuration, the photo-absorber, p-type silicon, is separated from the electrolyte by a \( \approx 1.5 \text{ nm} \) thick insulating layer of native silicon dioxide and semitransparent layers of titanium and platinum that were deposited as a continuous bi-layer by physical vapor deposition. The electric field across the MIS junction allows for photogenerated carriers to be collected at the metallic Pt layer, which also catalyzes the hydrogen evolution reaction. Overall, the p-Si based MIS photocathode is a stable photoelectrode capable of high photocurrents, making it an excellent baseline electrode to investigate in PEC flow cells.

The remainder of the chapter is organized as follows. After describing the design and operation of the MIS photocathode in the PEC flow cell, a range of bulk reactant concentrations, flow velocities, and illumination intensities are explored in order to map out the boundaries between mass transport-limited and photo-limited operating regimes. Example measurements are...
provided that clearly illustrate the characteristics of mass transport- and photo-limiting current behavior, as well as the transition between the two operating regimes. Based on the mass transport properties of the wide channel cell, a simple model is presented to predict the limiting current as a function of fluid velocity for higher reactant concentrations and solar insolation. Finally, the implications of this analysis for the design and operation of efficient PEC flow reactors are discussed.

5.2 Experimental

**Chemicals** - Electrolyte solutions were prepared from concentrated sulfuric acid (Certified ACS plus, Fischer Scientific), 18 MΩ cm deionized water, and sodium sulfate (ACS Reagent grade, Sigma Aldrich) supporting electrolyte. The amount of sulfuric acid varied between solutions, but the sodium sulfate concentration was always kept constant at 1 M.

**Photoelectrode fabrication** - All photoelectrodes consisted of an MIS structure in which semitransparent 3 nm Pt and 10 nm Ti layers were deposited sequentially by e-beam evaporation onto Si(100) substrates obtained from WRS Materials. Prime-grade p-Si(100) wafers with resistivity of 1-5 Ω-cm were used for photoelectrodes, while degenerate p-Si(100) wafers with resistivity of < 0.005 Ω-cm were used as metallic control samples. Indium back contacts were soldered onto the back of the samples after fabrication of the front MIS junction, and were verified to have an ohmic resistance of less than 10 Ω as determined by impedance spectroscopy. Exposed electrode areas of ≈ 0.24 cm² were defined by a square opening in 3M Electroplater’s tape. A black metal foil mask having the identical shape and area as the exposed electrode area was aligned on top of the PEC window to ensure that the illuminated area on the electrode was equal to the area exposed to the electrolyte.
Flow cell fabrication - The body of the flow cell consisted of two 3D printed plates (Figure 5.2). The inlet and outlet are located on the top plate, which is mechanically clamped to the bottom plate. A window is epoxied (J.B. Weld) to the top plate to allow for illumination of the photoelectrode. The bottom plate contains a rectangular cavity matching the geometry of the photoelectrode, which is sealed in place using 3M Electroplater’s tape with thickness of 180 μm such that the surface of the photoelectrode is flush with the bottom of the flow channel. A rectangular opening was cut out of the Electroplater’s tape to define the electrode area: 0.4 cm long by 0.6 cm wide. A Ag/AgCl reference electrode (Edaq, ET1073-1) is inserted through a hole on the side of the bottom plate. This reference electrode hole (ϕ = 3 mm) is 14 mm downstream of the photoelectrode. A counter electrode made of coiled Pt wire is placed 17 mm downstream of the working electrode. The two plates of the flow cell are clamped together with a Viton gasket in between, and the height of the flow channel (1.5 mm) is much narrower than its width (10 mm).

(Photo)electrochemical measurements - The PEC flow cell was illuminated using a Newport LCS-100 Solar Simulator fitted with an AM 1.5G filter. The intensity of the lamp was calibrated to 1 sun intensity under air mass (AM) 1.5 conditions using a silicon reference cell (VLSI Standards, SRC-1000-RTD-QZ). For experiments requiring concentrated light intensities, the higher light intensities were calibrated using a secondary cell consisting of a home-made 500 micron diameter silicon photo-diode that was itself calibrated at lower light intensities using the VLSI silicon reference cell. During experiments, the electrolyte was continuously purged with nitrogen in a 600 mL reservoir. Electrolyte was pumped to the flow cell using a Cole Parmer Masterflex L/S peristaltic pump with an Easy Load II pump head. For a given data set, an initial linear sweep voltammogram was created to determine the voltage required to achieve a limiting
current. Three successive chronoamperometry experiments were then held at this potential in order to record an average value of the limiting current.

Figure 5.2: a) Exploded view of 3D printed PEC flow cell assembly. b) Photograph of PEC flow cell. c) Simplified flow diagram of experimental set-up.

5.3 Results and discussion

5.3.1 Description of PEC flow cell and its operation

The wide-channel PEC flow cell and measurement set-up used as the basis for this study are shown in Figure 5.2. An exploded view of the flow cell components is provided in Figure 5.2a and a picture of the assembled device in operation is shown in Figure 5.2b. Within this parallel plate flow cell, the photoelectrode is integrated into the bottom plate, while the reference and
counter electrodes are incorporated downstream of the photoelectrode as shown in Figure 5.2a and described in the experimental section. A glass window in the top plate of the reactor allows for light to be transmitted downward onto the exposed photoelectrode surface, which is mounted horizontally within the cell. The small cross sectional area of the flow channel (0.15 cm²) permits the flow cell to achieve a maximum fluid velocity of 82 cm s⁻¹ with the peristaltic pump used in this study. This fluid velocity corresponds to a Reynolds number (Re) of 1225, meaning that laminar flow was established for all measurements. The rectangular flow channel is 8.5 cm long, with the photoelectrode placed halfway downstream. A parabolic velocity flow profile typical of flow through parallel plates can be assumed for this flow channel given that the channel’s height is much less than its width (h/w = 0.15) and the electrode is placed in the center of the channel, more than 1 characteristic length away from the corners.

Once the photoelectrode was assembled into the flow cell, its performance was evaluated using linear sweep voltammetry (LSV) and chronoamperometry (CA). Figure 5.3 shows a typical LSV curve for the standard p-Si MIS photocathodes measured in 200 mM H₂SO₄ and 1 M Na₂SO₄ under simulated AM 1.5 illumination, illustrating three different operating potential regimes. At positive applied potentials under illumination (Region I in Figure 5.3), no current is measured because too little photovoltage is generated to simultaneously achieve a high carrier collection efficiency and drive HER kinetics. As the potential is scanned more negative, a photo-reduction current becomes apparent around 0.32 V vs. NHE, corresponding to a potential known as the photocurrent onset potential (V_onset). Scanning further, there is a sharp increase in photocurrent until about -0.24 V vs. NHE. In the absence of significant ohmic losses, current in this portion of the LSV curve (Region II) is primarily dominated by reaction kinetics. Eventually, the photocurrent becomes relatively constant, with no substantial change observed as the potential is
scanned more negative (Region III). The current density in this region represents the limiting photocurrent of the photoelectrode, and its magnitude is strongly dependent on the light intensity and mass transport properties of the system. Small deviations in the limiting current arise due to the formation and detachment of bubbles on the photoelectrode surface and/or disturbances in the electrolyte flow.

![Diagram of LSV curves for p-Si MIS photocathode under simulated AM 1.5G illumination in 200 mM H₂SO₄ with 1 M Na₂SO₄. The potential is swept from positive to negative at a scan rate of 20 mV s⁻¹. At positive potentials, the photovoltage is insufficient for HER to proceed, and no photocurrent is recorded (Region I). As the potential is scanned more negative than the photocurrent onset potential (V_{onset}), the current becomes strongly dependent on the reaction kinetics (Region II). Below -0.24 V vs. NHE, the photocurrent becomes independent of the applied potential and is determined by the relative rates of light absorption and mass transfer (Region III).](image_url)

Figure 5.3: Standard LSV curves for the p-Si MIS photocathode under simulated AM 1.5G illumination in 200 mM H₂SO₄ with 1 M Na₂SO₄. The potential is swept from positive to negative at a scan rate of 20 mV s⁻¹. At positive potentials, the photovoltage is insufficient for HER to proceed, and no photocurrent is recorded (Region I). As the potential is scanned more negative than the photocurrent onset potential (V_{onset}), the current becomes strongly dependent on the reaction kinetics (Region II). Below -0.24 V vs. NHE, the photocurrent becomes independent of the applied potential and is determined by the relative rates of light absorption and mass transfer (Region III).

In this study, our focus is on establishing the limiting photocurrent behavior of MIS photocathodes under varied electrolyte flow rates, reactant concentrations, and light intensities. Experimentally, the limiting photocurrent was determined as follows: For a given reactant
concentration and illumination intensity, an LSV was performed to determine the applied potential or voltage that is required for operation in Region III, the limiting current regime. CA was then conducted at a potential that was 0.2 V to 0.3 V negative of the transition between Region II and Region III. Each CA trial lasted for 20 s, and the overall limiting current determined by the average value over three trials. In the following subsections, we explore cases where the photocurrent is limited entirely by light, entirely by mass transport, or a combination of the two.

5.3.2 Light limited photocurrent

PEC device operation is said to be light- or photo-limited when the rate limiting step in the sequence of processes illustrated in Figure 5.1b is the flux of photons to the photoelectrode. Under such conditions, the current density, \( i \), is highly sensitive to the spectral irradiance, \( I_0 \), and often varies linearly with it as described by Equation 5.1:

\[
i = \frac{q}{hc} \int_0^{\lambda_{Eg}} I_0(\lambda) \left(100\% - R(\lambda) - A_m(\lambda)\right) IQE(\lambda) \lambda \, d\lambda
\]

where \( q \) is the elementary charge, \( h \) is the Planck constant, \( c \) is the speed of light, \( \lambda_{Eg} \) is the bandgap energy of the semiconductor, \( R \) is the reflectivity of the photoelectrode surface, \( A_m \) is the absorptance of the metal bi-layer, and \( IQE(\lambda) \) is the internal quantum efficiency, defined as the fraction of photons absorbed by the photoelectrode that result in a collected photocurrent. Equation 5.1 is typically integrated over all wavelengths exceeding the bandgap energy where IQE is non-zero.

In the lab, PECs are typically tested under illumination sources having a spectrum that is modified to match the power density and spectral characteristics of the sun such as the standard air mass 1.5 global (AM 1.5G) spectrum. Assuming that a light source having AM 1.5 spectral
characteristics is being employed, Equation 5.1 can be integrated and simplified to obtain the following expression for photo-limited current density:

\[ i = q\Gamma \phi_{AM\,1.5}X \]  

Where \( \phi_{AM\,1.5} \) is the total photon flux from an AM 1.5 solar source incident on the PEC reactor and \( \Gamma \) is the fraction of incident photons that result in electrochemical current. \( X \) is the solar concentration factor, defined as the ratio of the photon flux incident on the PEC device to that received under one sun illumination at AM 1.5. A practical upper limit on the solar concentrating factor for a PEC has been suggested at 100 suns,\(^6,30\) and a recent technoeconomic analysis of solar concentrated PECs used a base case of 10 suns.\(^30,31\)

The LSVs shown in Figure 5.4a are for a photoelectrode operated in 200 mM sulfuric acid under different light intensities. Under one sun illumination, the photo-limited current was measured to be \( \approx 16 \, \text{mA cm}^{-2} \), substantially less than the theoretical maximum photocurrent of 43 mA cm\(^{-2}\) for crystalline Si under AM 1.5G illumination.\(^32\) As shown in Appendix C, section 5.5.1, Figure 5.7, this difference can be primarily attributed to the high reflectance and absorbance losses associated with the relatively thick Pt/Ti metal bilayer that was deposited on top of the MIS photodelectrodes in this study. In Figure 5.4b, the average limiting currents are plotted as a function of solar concentration factor, revealing that the photo-limited current recorded in 200 mM H\(_2\)SO\(_4\) scales linearly with solar concentration factor. This observation is consistent with the classical behavior predicted by Equation 5.2, and is commonly observed in other studies where the current is photo-limited.\(^3\) By contrast, the photocurrent is nearly independent of light intensity when the sulfuric acid concentration was decreased to 4 mM, where the photocurrent becomes mass transport-limited and primarily depends on the delivery of reactants to the surface. Although not observed in this work, it is worth noting that some lower-quality materials can exhibit sub-linear
dependence of photocurrent on light intensity due to higher carrier recombination rates at high light intensity.\textsuperscript{33}

![Figure 5.4: Photo-limited PEC operation. a) Linear sweep voltammetry (LSV) curves recorded for a p-Si MIS photocathode under varying solar concentrating factors. All scans were swept from positive to negative potential at a rate of 10 mV s\textsuperscript{-1} in 200 mM H\textsubscript{2}SO\textsubscript{4} flowing at an average velocity of 22 cm s\textsuperscript{-1}. b) Limiting photocurrent densities recorded for a p-Si MIS photocathode under varied solar concentration factors in weakly and strongly acidic electrolytes flowing at an average velocity of 22 cm s\textsuperscript{-1}.]

**5.3.3 Mass transport-limited photocurrent**

When the diffusion of reactants across the diffusion boundary layer is the slowest process illustrated in Figure 5.1b, the current generated by the PEC is limited by mass transport. In this case, the maximum flux of reactant species to the photoelectrode surface is less than the flux of photo-generated carriers to the photoelectrode surface. Usually, mass transport becomes limiting when there is a low concentration of reactant in the electrolyte. Such conditions are commonly encountered for photoelectrochemical and electrochemical CO\textsubscript{2} reduction, where the solubility of the reactant, CO\textsubscript{2}, limits its concentration in aqueous electrolytes to approximately 34 mM at 298 K. At such low concentrations, PEC operation in a stagnant electrolyte can greatly reduce the
limiting photocurrent because the concentration of the reactant species is quickly depleted near the photoelectrode surface, resulting in the development of a concentration gradient across the diffusion boundary layer that is insufficient to induce a flux of reactants equivalent to the photon flux. However, flow cells overcome this issue through forced convection of the liquid electrolyte, which decreases the thickness of the diffusion boundary layer across which reactants must diffuse in order to reach the photoelectrode surface. In flow cells, increasing the fluid velocity decreases the diffusion boundary layer thickness, and therefore increases the maximum diffusive flux across it.

Figure 5.5a shows LSV curves for a photoelectrode operating under 1 sun illumination in 4 mM sulfuric acid, a relatively dilute concentration for which mass transport-limited behavior is expected. The mass transport limitations are confirmed by the sensitivity of the limiting current to the electrolyte flow velocity (Figure 5.5b). As the electrolyte flows faster, the limiting current density increases due to the decreasing boundary layer thickness, which in turn causes the diffusive flux across it to increase. Because diffusion is a slow mode of transport relative to convection, the thickness of the diffusion boundary layer strongly affects the mass transport-limited current. The relationship between a forced convection velocity field and the diffusion boundary layer thickness can be described by a mass transport correlation that relates the Sherwood number (Sh=L/δ), Reynolds number (Re = \( \frac{UL}{\nu} \)), and the Schmidt number (Sc = \( \frac{\nu}{D} \)). The Reynolds number is the ratio between inertial and viscous forces, where \( U \) is the average fluid velocity, \( L \) is the characteristic length, and \( \nu \) is the kinematic viscosity. The Schmidt number is the ratio between the kinematic viscosity and the diffusivity \( D \). For a given geometry, the Sherwood number is a function of Re and Sc.
Figure 5.5: Mass transport-limited PEC operation. a) Linear sweep voltammetry for varying fluid velocities in 4 mM sulfuric acid. Scans are from positive to negative potential at a rate of 10 mV s\(^{-1}\). b) Comparing the average limiting current density of an MIS photoelectrode to a metallic electrode (p\(^+\)Si). The limiting current was measured using chronoamperometry.

The development of a convection-diffusion boundary layer for a flat electrode in a wide channel is well studied, and the average mass transport-limited current density can be described as: \(^{34}\)

\[
i = 1.6797 \, nF D^{2/3} C_{\infty} (\frac{U}{3h})^{1/3} L_{c}^{-1/3}
\]

where \(L_c\) is the length of the electrode in the direction of fluid flow, \(h\) is the half channel height, \(U\) is the average fluid velocity, \(n\) is the moles of electrons transferred per mole reactant, and \(C_{\infty}\) is the bulk concentration of the reactant species. As seen in Equation 5.3, the mass transport-limited current should vary linearly with the bulk concentration, and have a cubed root dependence on \(U\).

In order to view photoelectrode operation under mass transport-limiting conditions, the limiting photocurrent of the photoelectrode was measured at different fluid velocities under 1 sun and 2.8 sun light intensities in a dilute electrolyte (11 mM H\(_2\)SO\(_4\)). The relationship between the
experimentally-measured limiting current and $U^{1/3}$ is shown in Figure 5.5b for both cases. Because the reactant concentration is relatively dilute, the diffusive flux across the boundary layer is smaller than the photon flux under both 1 sun and 2.8 sun intensities. As a result, the two curves fall on top of each other and exhibit a strong dependence of limiting current on velocity. The mass-transport-limited behavior of the photoelectrode under these conditions was further confirmed by repeating the measurement in 11 mM H$_2$SO$_4$ with a metallic electrode consisting of the identical Pt layer deposited on a degenerately doped silicon sample ($p^+$ Si). As seen in Figure 5.5b, the dependence of the limiting current of the $p^+$Si electrode on $U^{1/3}$ is identical to that observed for the $p$-Si photoelectrode, verifying that the current is indeed limited by mass transport and independent of light intensity or reaction kinetics.

In higher sulfuric acid concentrations, the diffusive flux across the boundary layer is generally larger than the photon flux at 1 sun. Also shown in Figure 5.5b is the limiting current for the same photoelectrode operated under 1 sun light intensity and 200 mM H$_2$SO$_4$. In contrast to the behavior observed under identical illumination conditions in 11 mM H$_2$SO$_4$, the limiting current is insensitive to fluid velocity (Figure 5.5b), but sensitive to light intensity concentration (Figure 5.4b).

The mass transport-limited current data in Figure 5.5b can also be used to determine the effective diffusion coefficient of the reactants. By fitting the slope of $i_{lim}$ vs. $U^{1/3}$, the diffusion coefficient is calculated using Equation 5.3. In 11 mM H$_2$SO$_4$ with 1 M Na$_2$SO$_4$ supporting electrolyte, the effective diffusion coefficient of protons was found to be 4.1E-06 cm$^2$ s$^{-1}$. Although the proton diffusion coefficient in water is typically found to be around 9E-05 cm$^2$ s$^{-1}$, the presence of excess supporting sulfate ions favors the formation of bisulphate, which has a much slower diffusion coefficient.$^{35}$
The mass transport-limiting current behavior of the parallel plate flow cell is similar to what others have observed for electrochemical proton reduction on a metallic rotating disc electrode (RDE).\textsuperscript{36} While RDEs are not commonly used for studying photoelectrodes, the parallel plate design is convenient for aligning the photoelectrode with the illumination source, an integral part of this study. One can extend this analysis to other device geometries as well, provided that the appropriate mass transfer correlations for that geometry are available and used. Besides the correlation used here for the parallel plate geometry, other correlations are available in literature.\textsuperscript{37}

Finally, we note that this analysis of mass transfer effects on PEC operation has focused on the transport of reactant species to the photoelectrode surface, but the transport and subsequent collection of product species would also be of great importance for practical PEC reactors. In the present parallel plate configuration, one possible solution would be to incorporate a semitransparent counter electrode where the front window is currently located, and utilize flow-induced product separation to sweep the product gasses down two separate effluent channels as has been demonstrated for laminar flow cell fuel cells,\textsuperscript{38,39} batteries,\textsuperscript{40} and electrolyzers.\textsuperscript{41}

### 5.3.4 Predicting limiting photocurrent operating regimes

In the previous two sections, photoelectrode operation was described for operating conditions under which the photocurrent was solely limited by mass transport or illumination intensity. We now focus discussion on identifying and predicting the transition between these two operating behaviors and using that information to map out the operating conditions for which the photocurrent is expected to be limited by mass transport or illumination intensity. The basis of this analysis is the assumption that the transition between mass transport- and photo-limiting behavior occurs when the maximum flux of reactants to the surface is stoichiometrically equal to the maximum flux of photogenerated carriers to the electrode surface. To minimize the presence and
impact of bubbles for this analysis, the limiting current was determined directly from LSVs at a moderately fast scan rate of 40 mV s$^{-1}$ whereby the limiting current can be achieved quickly before the onset of significant product bubble growth.

An example illustrating the transition between mass transport- to photo-limiting behavior for a photoelectrode is provided in Figure 5.6a, which shows the limiting photocurrent of an MIS photocathode in varying H$_2$SO$_4$ concentrations under 1 sun illumination as a function of fluid velocity. In 500 mM H$_2$SO$_4$, equivalent proton concentration of 1000 mM, the current is photo-limited at all flow velocities. As the reactant concentration decreases, higher velocities are required to achieve the photo-limited current. For example, in 33 mM H$_2$SO$_4$, equivalent proton concentration of 66 mM, the limiting photocurrent is highly dependent on flow velocity up until 30 cm s$^{-1}$, indicating that it is primarily limited by mass transport in this region of operation. At flow velocities beyond 30 cm s$^{-1}$, the limiting photocurrent plateaus to the one sun limit, indicating that the photoelectrode has now become limited by light.

Mathematically, the transition between the mass transport and light-limiting photocurrent can be described by setting Equation 5.2 (describing the maximum flux of photo-generated carriers to the electrode surface) equal to Equation 5.3 (describing the maximum reactant flux). For a given light intensity and flow velocity, the bulk reactant concentration, $C_\infty$, at which the photo-limiting and mass transport-limiting processes are balanced is given by Equation 5.4:

$$C_\infty = \frac{0.858}{N_A} \frac{\Gamma \phi_{AM 1.5} X}{n \psi} \quad 5.4$$

where $\psi = \left(\frac{D^2 U}{h_L c}\right)^{1/3}$. The numerator of Equation 5.4, $\Gamma \phi_{AM 1.5} X$, is the maximum photon flux, while the denominator $n \psi$ contains the parameters governing the mass transport properties of the system and has dimensions of velocity. $N_A$ is Avagadro’s number. In Figure 5.6b, $C_\infty$ calculated
from Equation 5.4 is plotted against $\psi$ based on the PEC flow cell geometry and operating parameters employed for the measurements shown in in Figure 5.6a. The value of $\Gamma$ was set to be 0.42, which corresponds to a photo-limited current density of 18 mA cm$^{-2}$ under 1 sun intensity as was observed for the sample used for these measurements. The solar concentration factor $X$ was set to 1 sun. The channel half height $h$ was 0.75 mm, and the electrode length was 4.0 mm. An effective proton diffusion coefficient of 4.1E-06 cm$^2$ s$^{-1}$ was found for H$_2$SO$_4$ in 1.0 M Na$_2$SO$_4$ and was determined experimentally from the slope of the Levich plot in Figure 5.5b. For a given coordinate pair of $\psi$ and reactant concentration, the curve plotted in Figure 5.6b represents a tie line where operating points that lay above the curve are photo-limited, while points that lay below the curve are limited by mass transport. Ideally, a PEC system would be designed to operate just above the tie line to ensure operation in a photo-limited regime. Increasing the fluid velocity, and therefore $\psi$, to excessively high levels can result in efficiency losses due to pump energy requirements as well as dilution of the products. Unnecessarily high flow rates can also result in undesirable mixing of the products downstream as well as dilution of the product species. Therefore, it is advisable to set the operating fluid velocity high enough to ensure that the current is not limited by mass transport, but not so high that large pumping and downstream separation costs will be incurred.
Figure 5.6: Predicting the boundary between photo-limited and mass transport-limited PEC operation. a) The limiting current of a photoelectrode operated in varying H$_2$SO$_4$ concentration under 1 sun illumination. At low fluid velocities the current is mass transport-limited, but increasing the flow velocity allows the current to reach the 1 sun photo-limited value, which is equal to the limiting current in 500 mM H$_2$SO$_4$ ([H$^+$] = 1000 mM). b) Predicted boundary between photo-limited and mass transport-limited current at 1 sun. Individual data points on the plot correspond to the same experimental data points shown in a). The tie line was calculated using Equation 5.4, and $\psi$ is defined in the text. c) Predicted boundary at higher levels of solar concentration for a reaction with $n=1$ mole of electrons transferred per mole reactant. d) Predicted boundary for reactions involving $n=8$ moles of electrons transferred per mole reactant.
Also plotted in Figure 5.6b are the individual operating points corresponding to the electrolyte concentration and flow velocities that were used in the measurements reported in Figure 5.6a. As the bulk proton concentration of the electrolyte increases, lower values of $\psi$, and thus lower flow velocities, are required to achieve a photo-limited current. At a bulk proton concentration of 56 mM, the transition from mass transport-limited to photo-limited operation is expected at $\psi = 0.0029$, corresponding to flow velocities greater than 40 cm s$^{-1}$ and in agreement with Figure 5.6a. Although reasonable agreement between the predicted and observed limiting current transition is also demonstrated at a bulk proton concentration of 66 mM, the transition is not as clear at a proton concentration of 80 mM. This observation of a blurred transition between mass- and light-limited photocurrent most likely results because the predicted transition flow velocity, 13 cm s$^{-1}$, is too slow to remove the hydrogen bubbles adsorbed to the surface, resulting in additional mass transport and optical losses.

The analysis described above based on Equation 5.4 and illustrated in Figure 5.6b can be expanded upon to predict the limiting current over much wider ranges of light intensities, reactant concentrations, and fluid velocities. Figure 5.6c shows the predicted boundary between photo-limited and mass transport-limited currents for four different photo-limited current densities ($qG\phi_{AM\,1.5}X$). If a photoelectrode has an expected photo-limited current density of 15 mA cm$^{-2}$ at one sun intensity, then the photo-limited current density at ten suns can be estimated to be 150 mA cm$^{-2}$. The tie lines in Figure 5.6c assume a reaction stoichiometry where one electron is transferred per mole of reactant. As the solar concentration factor increases, the device requires more concentrated electrolyte to avoid mass transport limitations. For example, under 20 sun illumination, ($qG\phi_{AM\,1.5}X = 300$ mA cm$^{-2}$) a system with a 1000 mM proton concentration is likely to be mass transport-limited. By increasing the proton concentration to 1500 mM, or by increasing
the electrolyte flow velocity to 30 cm s\(^{-1}\) \(\dot{\psi} = 0.0028 \text{ cm s}^{-1}\), the system becomes photo-limited. In the cases where the reactant concentration cannot be increased – for example, due to a solubility limit – increasing the flow rate of the electrolyte can be a practical solution to overcoming mass transport limitations. This analysis can be extended to systems where multiple electrons are transferred per reactant such that the electron transfer number, \(n\), is \(> 1\). Figure 5.6d shows the tie lines for a hypothetical reaction where \(n = 8\). As the value of \(n\) increases, the bulk reactant concentration required to achieve a photo-limited current density decreases because each reactant molecule reaching the photoelectrode surface is capable of accepting multiple photo-generated e\(^-\).

Using an analysis such as that shown in Figure 5.6 to map out the expected limiting cases of operation can be very useful for both the design of PEC reactors and choosing operating conditions. While HER was used as a model reaction to describe the mass transport characteristics of the wide channel cell, these concepts can be applied to other reactions and reactor designs as well. The range of values for \(\dot{\psi}\) will depend on the geometry of the flow cell, as well as the diffusion coefficient(s) of reactant species. Figure 5.6c and Figure 5.6d illustrate that the number of electrons transferred per mole of reactant formed in the reaction is also important for determining the optimal operating conditions for a PEC flow reactor.\(^{42}\) For HER, one electron is transferred per proton, but for \(\text{CO}_2\) reduction, the number of electrons transferred depends on what products are formed – in the case of reduction to methane, 8 mole e\(^-\) are transferred per mole of reactant. Consequently, the system reaches a photo-limited current at lower reactant concentrations.

While the present analysis provides a useful framework for understanding limiting photocurrent behavior of photoelectrodes operating under a large applied bias, it must be recognized that photoelectrodes in stand-alone PEC devices will likely operate at potentials for
which kinetic and ohmic losses will also be important. This is especially true at high reactant concentrations, light intensities, and flow rates for which the high current densities will likely necessitate photoelectrode operation at a potential for which the current is affected by the reaction kinetics and/or ohmic resistances associated with the electrode, electrolyte, and membrane (if present). These ohmic losses are not considered in Figure 5.6c, but are likely very important at solar concentrating factors exceeding 10-20 suns. As an example, consider a PEC flow cell containing a semitransparent counter electrode and a photoelectrode in parallel-plate configuration with a 2 mm separation distance operating in 0.5 M H₂SO₄ (conductivity ≈0.21 S cm⁻¹). If the device current density at 1 sun intensity is 10 mA cm⁻², its operating current density at 50 sun intensity would be expected to be around 500 mA cm⁻². Assuming Tafel kinetics for state-of-the-art Pt hydrogen evolution catalysts and IrO₂ oxygen evolution catalysts,₁¹,₄₃ such a current density would require HER and OER kinetic overpotential losses of 62 mV and 387 mV, respectively. The ohmic voltage drop would be ≈ 475 mV. This simple exercise highlights the challenges associated with operating any PEC reactor at high light intensities. For a stand-alone PEC device capable of unassisted electrolysis, it will be essential that these losses are also accounted for in the design and optimization of both the photoelectrode(s) and PEC reactor.

Another assumption of this analysis is that migration, or ionic transport due to the gradient in the electric field, does not affect the mass transport limited current. This is a good assumption to make in the current study because the relatively low concentrations of H₂SO₄ ensures that most of the migration current is carried by the Na⁺, SO₄⁻², and HSO₄⁻ ions in the supporting electrolyte. At higher H₂SO₄ concentrations, the free protons carry a larger fraction of the migration current, and the effect of migration cannot be entirely ignored. Thus, the predicted boundary between photo-limited and mass transport-limited currents shown in Figure 5.6c at high reactant
concentrations and illumination intensity will likely deviate from what would be observed experimentally.

5.4 Conclusions

This chapter has systematically investigated the limiting photocurrent behavior of a wide channel PEC flow cell containing a p-Si MIS photocathode under varying electrolyte concentrations, flow velocities, and illumination intensities. When the maximum flux of photogenerated carriers to the electrode surface exceeds the flux of reactants across the diffusion boundary layer, increasing the fluid velocity can have a huge effect on the limiting photocurrent, and thus the overall solar-to-fuel conversion efficiency of the PEC device. For this reason, understanding the boundary between mass transport and photo-limited operating regimes is of great importance for optimizing PEC performance. These operating regimes for a given reactor geometry can be predicted and modeled if the relationship between flow velocity and diffusion boundary layer thickness is known. By balancing the photogenerated carrier flux with the diffusive flux of the reactants, one can determine the minimum flow velocity required to achieve a photo-limited current. Importantly, the methodologies and analysis introduced in this paper can be extended to other PEC systems based on different photoelectrodes, optical concentration factors, and dilute reactant concentrations.

5.5 Appendix C

5.5.1 UV-Vis analysis of photoelectrode optical losses

The maximum theoretical photocurrent generated by a single-junction solar cell based on crystalline silicon (c-Si) is 43 mA cm$^{-2}$ under AM 1.5G illumination. In reality, the observed photocurrent is less than this amount due to optical and/or electronic losses. In the metal-insulator-
semiconductor (MIS) photoelectrodes used in this study, the metal bilayer deposited on top of the silicon is directly responsible for decreased photocurrents due to optical absorption in the metal and reflection off of its surface. Reflectance and absorbance of light by the metal bilayer reduce the transmittance (T), defined as the percentage of incident light that is transmitted to the c-Si absorber layer according to Equation C5.5:

\[ T = 100\% - A - R \]  

where \( T \) is the transmittance, \( A \) the absorptance, or percent of incident light absorbed, and \( R \) is the reflectance of light from the front surface of the photoelectrode. In order to measure \( T \) for the metal bilayer that was deposited on the p-Si photoelectrodes studied in this work, an identical (3 nm /10 nm) Pt/Ti bilayer was deposited onto a transparent glass slide. The transmittance spectra for this sample and a bare glass slide were measured by UV-Vis spectroscopy and are shown in Figure 5.7.

![Figure 5.7: Percent transmittance for a glass slide deposited with a 10 nm Ti 3 nm Pt metal bi-layer.](image-url)
The bare glass slide is highly transparent, allowing \(\approx 90\%\) of photons to be transmitted across the visible spectrum. The metal bi-layer causes the transmitted light to drop significantly, and the average transmittance between 300 and 800 nm for the metal-bilayer on glass substrate is 39.5\%. Applying this average transmittance to the AM 1.5 spectrum, the maximum photo-generated current for a silicon absorber is \((39.5\%)\times(43 \text{ mA cm}^{-2})= 17.0 \text{ mA cm}^{-2}\). Comparing this value to the 16 mA cm\(^{-2}\) photocurrent generated at AM 1.5G conditions by the MIS photoelectrodes in this study, it can be concluded that most of the losses in the photolimited current result from optical losses associated with the Pt/Ti bilayer.

It should be noted that the 1 sun photolimited current reported in Figure 5.6 is 18 mA cm\(^{-2}\), which is higher than the photolimited current reported in Figure 5.4. This can be attributed to slight differences in the metal bi-layer thickness of the photoelectrodes used in these experiments. Although the photoelectrodes were created during the same metal deposition process, local variations in metal thickness can still arise, resulting in varied optical losses associated with \(A_m\).

5.6 Acknowledgements

The authors acknowledge Columbia University for funding and Dr. Alan West for helpful discussions.

5.7 References

3. Sun, K. *et al.* A Stabilized, Intrinsically Safe, 10% Efficient, Solar-Driven Water-Splitting


CHAPTER 6

CONCLUSIONS AND FUTURE DIRECTIONS

Membraneless electrolyzers are an emerging reactor design for water electrolysis which have the potential to reduce the capital costs for H₂ production. These devices achieve product separation by carefully regulating the transport of the electrochemically generated gas bubbles as they detach from the electrodes and flow through a liquid electrolyte. The efficiency of a membraneless electrolyzer is strongly dependent on the electrode separation distance. Narrower electrode separation distances decrease the ohmic resistance of the electrolyte, but increase the risk of cross-over of the product gases. Through high speed video (HSV) imaging experiments, the bubble formation processes on the electrode surfaces could be directly observed and quantified. HSV analysis indicates that as much as 50% of the electrochemically generated H₂ remains dissolved in the aqueous electrolyte. Future designs of membraneless electrolyzers need to account for dissolved species transport mechanisms and encourage better phase separation of the H₂ downstream.

Controlling the size and position of the electrochemically generated H₂ bubbles is also important for determining the purity of H₂ collected. In Chapter 4 it was demonstrated that asymmetric electrocatalyst placement on a mesh flow-through electrode can suppress cross-over to be as low as 1% in the absence of forced convection. Direct observation of the dissolved species transport using an indicator dye suggests that dissolved species cross-over is greatly enhanced by free convection caused by local turbulence at the electrode generated by bubble detachment events. As a result, the 1% cross-over rate is thought to be largely attributed to the uncontrolled transport of the dissolved H₂. Using asymmetric electrodes in conjunction with forced convection of the electrolyte could help to suppress the H₂ cross-over percentage to even lower values.
This dissertation also demonstrates how 3D printing can be leveraged to create new device concepts and experimental setups. The 3D printed imaging cell used in Chapter 3 was the first of its kind to combine high speed imaging of bubble flows with an image processing algorithm to reliably quantify the gas evolution efficiency and the current distribution based on the size and positions of the detected bubbles. In Chapter 4, the asymmetric flow-through electrodes were incorporated into a 3D printed collection apparatus for the first ever demonstration of a floating PV-electrolyzer device. This so-called solar fuel rig draws inspiration from the floating oil rig platforms deployed in the oceans today. Membraneless electrolyzers could be particularly well suited for unfiltered seawater electrolysis, where the presence of trace calcium and magnesium ions would readily degrade the performance of a membrane-based device. Lastly, a 3D printed PEC flow cell was demonstrated as a first of its kind parallel plate cell for enhanced mass transport to a photoelectrode. This concept is highly relevant to scaled up PEC reactor designs, where product and reactant transport are not often considered. If a PEC device was to be placed under an array of concentrating mirrors and lenses, extremely high current densities are expected. In these scaled up scenarios, the photocurrent may not be limited by the intensity of light, but rather the transport of the reactants and products. A membraneless PEC flow cell could help facilitate product transport and collection while also allowing the device to operate at the maximum possible photocurrent.

More broadly, the insights for membraneless electrolyzer operation and design that were developed in this dissertation could be applied to other electrochemical systems involving multiphase flows, such as electrochemical CO₂ reduction or ammonia synthesis. However, the initial conclusions of this dissertation also suggest that the phase equilibrium at gas evolving electrodes is nontrivial, and additional research is necessary to fully understand gas bubble
formation and transport throughout the electrolyzer. The remainder of this chapter is devoted to the discussion of future research directions which can build upon the present understanding of gas transport in electrochemical cells.

6.1 **Modeling multiphase flows in membraneless electrochemical cells**

Using computational fluid dynamics (CFD) to model the transport of gas bubbles in a membraneless electrolyzer is a logical extension of the work presented in this dissertation. Past CFD studies of bubbly flows have been conducted for electrochemical systems, but the emphasis of these studies was on the removal of bubbles to improve the overall ohmic resistance of the electrolyzer.\(^1\)\(^-\)\(^7\) A rigorous CFD study relating the forces exerted on bubbles to their downstream separation has yet to be demonstrated.

The focus of a combined CFD and experimental study should be to determine the scaling rules for membraneless electrochemical cells. Theoretically predicted cross-over percentages could be compared to experimental measurements through gas chromatography of the product collection streams. High speed video experiments could also be used to validate CFD predictions for the trajectories and spatial variation of electrochemically generated bubbles flowing in the bulk electrolyte. After establishing a solid intuition of the governing physics of multiphase flows, a successful CFD study should be able to determine the reasonable limits for electrode lengths and separation distances in membraneless electrolyzers.

A significant challenge for multiphase flow models is the amount of computing power required. Ideally, a simulation would account for the total force balance on every bubble in the system, but for an electrolyzer which can generate tens of thousands of bubbles at once, this can be impractical.\(^8\) Grid generation for the simulation can also be challenging when boundaries are moving as a function of time. Nonetheless, computational studies tracking large numbers of
particles do exist. Takeuchi et al. applied a discrete element method to track ~30,000 particles with a uniform diameter of 2.4 mm.\textsuperscript{9} It should be noted that these particles were modeled as rigid spheres, and therefore do not necessarily have the same types of interactions as bubbles. In an electrochemical cell, bubbles can collide, grow, dissolve, coalesce, nucleate, and wet a surface. Eulerian type models can reduce the computational demands of a CFD simulation by assuming the bubbles behave as a pseudo-continuous phase.\textsuperscript{10} However, doing so also compresses information about the behavior of discrete bubbles, and may not be able to characterize stochastic cross-over events. A compromise may be to limit simulations to a handful of bubbles to describe their interactions with each other. Studies of individual or pairs of bubbles can also be useful for describing how they interact with the electrode surface, which is another important question for future research to address.

6.2 Engineering the electrode surface tension

Related to the CFD modeling of membraneless electrolyzers are the physical properties of the bubbles themselves. In Chapter 3, it was reported that a wide distribution of bubble diameters were detected on a mesh flow through electrode. It was hypothesized that these bubble diameters were likely related to both the surface tension and geometry of the mesh wires. Designing electrodes which can produce bubbles of consistent size could help control the flow of bubbles in the bulk electrolyte as well as simplify the analysis in CFD simulations.

Bubble formation and departure is a function of both the flow properties of the electrolyzer, the local current density at the electrodes, and the surface tension on the electrode. Historically, bubbles adsorbed to the electrode surface were of interest because they can increase the overall ohmic resistance of the electrolyzer as well as the kinetic overpotential of the half reaction.\textsuperscript{11} Although these effects can be of interest for describing the overall efficiency, the objective of
studying bubble dynamics in a membraneless electrolyzer is to better understand their impact on cross-over rates, and therefore the purity of collected gas.

Improved understanding of the bubble formation mechanisms can also improve the quality of phase separation. Zhang, et al. have reported on modifying the surface wettability of graphene electrodes to improve the rate of gas evolution.\textsuperscript{12} Yu, et al. reported on a Janus mesh structure, with asymmetric surface wettability on each side of a mesh wire to control how and where bubbles rest on the surface.\textsuperscript{13} These Janus mesh structures are reminiscent of the mesh electrodes with asymmetric catalyst placement reported on in Chapter 4 of this dissertation. One could imagine both of these techniques incorporated into a single mesh electrode to both control where the electrochemical reaction occurs and where bubble nucleation occurs.

Hypothetically, the site of the electrochemical reaction does not have to be the site of bubble nucleation and departure. From a design perspective, it would be easier to model a membraneless electrolyzer where all the electrogenerated product is in the dissolved phase. One could then imagine a downstream process, even if just a few centimeters away from the electrode, where gas bubbles are allowed to nucleate and separate from the electrolyte without risk of cross-over. On the other hand, engineering an electrode surface which has a higher gas evolution efficiency could mitigate inefficiencies and losses due to dissolution of the product gas. In the next section, improvements to the overall phase separation and gas collection in a membraneless electrolyzer are discussed.

6.3 Improving downstream phase separation

Managing the dissolution of \(H_2\) and \(O_2\) gases is a significant challenge for membraneless electrolyzers. The amount of \(H_2\) dissolved in the electrolyte is often times at super saturation levels. In the HSV study in Chapter 3, the gas evolution efficiency for \(H_2\) in the region immediately...
downstream of a flow through electrode was approximately 50%. As the bubbles continue to rise in the effluent stream, however, they are expected to ripen and extract additional H\textsubscript{2} from the super saturated electrolyte. In Chapter 4, the evolved gas was directly collected in the headspace above the cathode, and the gas collection efficiency was measured to be approximately 83%. In a study at similar current densities by O’Neil, et al., the collection efficiency was approximately 90% after letting the H\textsubscript{2} bubbles float upwards in a 12-inch collection column.\textsuperscript{14}

In a scaled-up device, the separation of dissolved H\textsubscript{2} from the electrolyte must be further optimized. Engineered surfaces could encourage nucleation of H\textsubscript{2} to force the gas out of solution. Another option could be to modify the gas solubility of the electrolyte by adjusting the temperature or pressure. Given that the Henry’s law constant for gas solubility is temperature dependent, performing electrolysis at a higher temperature would decrease the saturation concentration of H\textsubscript{2}, thereby promoting additional gas separation from the electrolyte. However, this approach should only be pursued if the enhancement in H\textsubscript{2} capture exceeds the energy penalty for heating the electrolyte.

In any case, the electrolyte must eventually be recycled back to the inlet of the electrolyzer. If the electrolyte is unable to sufficiently degas before recycle, then at steady state the electrolyzer will be saturated with both H\textsubscript{2} and O\textsubscript{2} gases. On the one hand this guarantees that there will be no additional losses due to solvation of the product gases. However, this also guarantees that the dissolved gases will cross-over and mix in the opposing collection chambers, decreasing the overall purity of gas collected. In many scaled up devices some amount of gas crossover is tolerable, and catalytic converters are used to back-oxidize H\textsubscript{2} or reduce O\textsubscript{2} in a controllable manner.\textsuperscript{15} A similar implementation could be used in a membraneless electrolyzer, but more information is needed regarding large scale designs.
6.4 Scale-up of membraneless electrolyzers

Eventually, the research goal for membraneless electrolyzers is to demonstrate H$_2$ production at larger scales without compromising efficiencies and product purities achieved at smaller scales. Before demonstrating a large scale membraneless electrolyzer, it is important to develop an understanding of the product transport fundamentals through CFD modeling. Working with larger scale H$_2$ generation can present serious safety hazards which must be taken very seriously.$^{15}$ CFD modeling can help serve as a guiding tool to understand exactly how a device is supposed to operate without having to engage in the safety risks of an experimental study.

Once CFD simulations are validated for bench scale experiments, larger devices can start to be tested. Large-scale experiments should also be validated using CFD, and some model parameters may need to be adjusted. The performance of a scaled up membraneless electrolyzer should also be compared directly to the performance of a scaled-up PEM electrolyzer. Do both devices operate at comparable efficiency, capacity, and product purity? A more detailed techno-economic analysis based on the results could directly compare the capital costs of the devices and address necessary design changes in order for the technologies to be competitive.

6.5 Membraneless electrolyzers for CO$_2$ reduction

Although the focus of this dissertation has been on membraneless electrolyzers for H$_2$ production, they can also be used for the electrochemical reduction of CO$_2$ to carbon-based fuels. A significant body of work has focused on the electrode materials for CO$_2$ reduction, for which a wide array of products can be made.$^{16-24}$ One challenge for CO$_2$ reduction is achieving high Faradaic efficiency, which becomes challenging due to competition with the hydrogen evolution reaction. Ongoing research is devoted to identifying electrocatalyst materials for CO$_2$ reduction that are active, selective, and stable. Most demonstrations of CO$_2$ electrolysis are limited to C1 or
C2 products such as CO or ethylene, but recent developments have also shown that C3 and even C4 products are possible. However, an important question to raise is the role the reactor design can play in synthesizing longer chain products. Enhanced mass transport in a membraneless electrolyzer could also improve the limiting current of a CO₂ electrolysis device, which depends on the solubility limit of CO₂ in the aqueous electrolyte. Much is unknown as to what a scaled CO₂ reduction reactor would look like, and the design space is wide open. Product separation in a CO₂ reduction reactor is non-trivial, but initial studies in a membraneless electrolyzer could give powerful insight on the relationship between the bulk fluid transport and the overall performance of the device.

More broadly, membraneless electrolyzers have the potential to reduce the capital costs of electrochemical fuel generation. Personal transportation vehicles are beginning to transition to hydrogen and electrical power sources, but longer chain hydrocarbons will continue to be relevant in heavy transportation sectors such as the shipping or airline industry. In a renewable energy economy that relies heavily on electricity from solar or wind energy, membraneless electrolyzers can be a powerful, low cost technology to convert electricity into storable chemical energy.

6.6 References


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