# Direct Observation of the Local Microenvironment in Inhomogeneous CO<sub>2</sub> Reduction Gas Diffusion Electrodes via Versatile pOH Imaging

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## Abstract

We report how the micrometer-scale morphology of a carbon dioxide reduction ( $CO_2R$ ) gas diffusion electrode (GDE) affects the mass transport properties and with it, the local CO<sub>2</sub>R performance. We developed a technique to probe the microenvironment in a CO<sub>2</sub>R GDE via local pOH imaging with timeand three-dimensional spatial micrometer-scale resolution. The local activity of hydroxide anions ( $OH^{-}$ ), represented by the pOH value, around a GDE in contact with an aqueous electrolyte is a crucial parameter that governs the catalytic activity and CO<sub>2</sub>R selectivity. Here, we use fluorescence confocal laser scanning microscopy (CLSM) to create maps of the local pOH around a copper GDE by combining two ratiometric fluorescent dyes, one of which is demonstrated as a pOH sensor for the first time in this work. We observe that the local pOH decreases when current is applied due to the creation of OH<sup>-</sup> as a byproduct of  $CO_2R$ . Interestingly, the pOH is lower inside microtrenches compared to the electrode surface and decreases further as trenches become narrower due to enhanced trapping of OH<sup>-</sup>. We support our experimental results with multiphysics simulations that correlate exceptionally well with measurements. These simulations additionally suggest that the decreased pOH inside microcavities in the surface of a  $CO_2R$  GDE leads to locally enhanced selectivity towards multicarbon ( $C_{2+}$ ) products. This study suggests that narrow microstructures on the length scale of 5 µm in a GDE surface serve as local  $CO_2R$  hotspots and highlights the importance of a GDE's micromorphology on the  $CO_2R$  performance.

# Introduction

It is crucial to store renewable energy effectively for long time periods in order to reach the negative  $CO_2$  emissions required to mitigate the consequences of climate change. Accordingly, the concept of electrochemical  $CO_2$  reduction ( $CO_2R$ ) is compelling because it enables the storage of renewable energy in the form of chemical bonds.  $CO_2R$  establishes  $CO_2$  and water ( $H_2O$ ) as the primary sustainable feedstocks to form useful chemicals and fuels, thereby closing the carbon cycle.<sup>1-(4)</sup>

Though there have been many encouraging advances in the field of electrochemical  $CO_2R$ , the process is highly complex and there are still challenges that must be overcome before this technology is energy-efficient and selective enough to be used at an industrial scale.<sup>1,(3),(5)</sup> One promising approach is the use of gas diffusion electrodes (GDEs).<sup>(6)-(8)</sup> This type of electrode addresses the problem of mass transfer limitations that are encountered in conventional catalytic setups by delivering  $CO_2$  in the gas phase to a catalyst in contact with a liquid electrolyte. Most GDEs consist of a macroporous gas diffusion

layer topped with a hydrophobic microporous layer that is then subsequently coated with a catalyst.<sup>(9)-(11)</sup> (**Figure 1c**). In many cases, the catalyst is wetted by a thin layer of aqueous electrolyte to provide ionic conductivity but not limit  $CO_2$  transport to the surface.<sup>(10),(12)</sup> This setup allows for increased current densities by more than one order of magnitude over conventional setups: For a conventional setup in which two electrodes are placed in  $CO_2$ -saturated electrolyte, mass transport to the cathode limits the rate of  $CO_2$  consumption and hence, the maximum current density magnitude does not exceed 30 mA/cm<sup>2</sup>. On the other hand, tailored GDEs with current density magnitudes larger than 1 A/cm<sup>2</sup> have been demonstrated.<sup>(13),(14)</sup>

Various parameters influence the activity and selectivity of a GDE, and it is the interplay between these operating parameters that dictates the CO<sub>2</sub>R performance. It is therefore important to understand the impact of each parameter. One very important parameter that influences the system's performance is the choice of catalyst material.<sup>(15)-(17)</sup> The most commonly studied  $CO_2R$  catalyst is copper, because its moderate binding energy to CO as a reaction intermediate allows for its further reduction to desirable higher-order carbon products such as ethanol, propanol, ethylene, and acetate.<sup>(16),(18),(19)</sup> Furthermore, in determining the selectivity and activity of CO<sub>2</sub>R, the composition and structure of the GDE, i.e. pore size, hydrophobicity, surface depositions or the presence of microstructures, play an ill-understood role.<sup>(9),(10),(20),(21)</sup> Other parameters include the applied potential, the properties of the electrolyte (e.g. the constituent ions and the viscosity), the cell configuration, and the choice of ion-exchange membranes.<sup>(13),(18),(19),(22)-(24)</sup> In addition to these, the local activities of hydroxides,  $a_{OH^-}$ , and protons,  $a_{H^+}$ , for which the pOH and pH values are a measure, are critical because both species are involved in the  $CO_2R$  process<sup>(19),(25)</sup>, as per equations S19-S26. The pOH value is defined as  $pOH = -log_{10}(a_{OH^-})$  and the pH value as  $pH = -log_{10}(a_{H^+})$ . In many cases, hydroxide and proton activity can be approximated as the concentration. At equilibrium,  $pH + pOH = pK_w$ , with pK<sub>w</sub> = 14 under ambient conditions, so if the system is in equilibrium, the pH can be directly inferred from the pOH.<sup>(26)</sup>

For a GDE at open circuit,  $CO_2$  diffuses through the macroporous and microporous layers into the electrolyte where it rapidly reacts with  $OH^-$  to form bicarbonate and carbonate anions, according to the chemical equilibrium reactions listed in equations  $S8 - S12.^{(27)-(29)}$  This decreases the local  $OH^-$  concentration and thereby increases the pOH in the electrolyte. This process is schematically depicted in **Figure1c** (left side). Due to the participation of  $CO_2$  in these buffer reactions, the ability to map the spatially and time-resolved pOH around a GDE enables us to assess the local concentration of  $CO_2$  in the electrolyte around a GDE.

At a non-zero current, a portion of the CO<sub>2</sub> molecules is reduced to form products such as carbon monoxide, formic acid, methane or ethylene, among others<sup>(11),(18),(19),(30)-(32)</sup>, as per equations S19 – S26. During this non-equilibrium process, one OH<sup>-</sup> is created, or one buffer species is deprotenated for each electron involved.<sup>(33)</sup> The same holds true for the competing hydrogen evolution reaction (HER) that must be suppressed to maximize the CO<sub>2</sub>R yield. As a result, at sufficiently high current densities, we expect that the pOH will decrease, as is depicted schematically in **Figure1c** (right side). This effect is in competition with an increase in pOH caused by unreduced CO<sub>2</sub> molecules that undergo reactions to form bicarbonate and carbonate anions. As a result, a low observed pOH indicates the presence of CO<sub>2</sub> reduction activity.

In addition, it has been shown that the pH and pOH themselves have a substantial impact on the reactivity and selectivity of CO<sub>2</sub>R. It has been demonstrated that high pH (corresponding to low pOH) suppresses the parasitic  $HER^{(25),(34)}$  and shifts the CO<sub>2</sub>R selectivity towards C<sub>2+</sub> products.<sup>(8),(25),(32),(35)</sup> The

reason for the latter is that  $OH^-$  actively suppresses the creation of single carbon products (C<sub>1</sub>) and hydrogen molecules (H<sub>2</sub>), while it does not affect the C<sub>2+</sub> current density. This leads to an increased Faradaic efficiency (FE) towards C<sub>2+</sub> products at a given current density<sup>(35),(36)</sup>.

As a result, knowledge of the pOH in the vicinity of an operating GDE is critical and can aid researchers in better optimizing the performance of a GDE. Notably, the pOH is expected to vary as a function of distance from the electrode surface. In addition, it also varies in the plane parallel to the surface if there are any inhomogeneities such as microcavities present on the surface. For these reasons, a measuring technique that allows one to map to pOH around an operating GDE with high resolution in three dimensions is desired.

Previous efforts towards this end have included theoretical studies of the pH around GDEs performing  $CO_2R$ .<sup>(10),(37),(38)</sup> Experimental approaches have involved scanning electrochemical microscopy (SECM)<sup>(39),(40)</sup>, surface-enhanced Raman (SERS)<sup>(41)</sup>, electrochemical atomic force microscopy (EC-AFM)<sup>(42)</sup> and surface-enhanced infrared spectroscopy (SEIRAS).<sup>(34)</sup> While all these techniques are powerful and can reach spatial resolutions on the nanometer scale, they do not have the ability to map the operando pH/ pOH of an entire macroscopic sample in three spatial dimensions. Fluorescent confocal laser scanning microscopy (CLSM), a well-known technique in the context of electrochemistry, (43)-(48) is our method of choice to address these concerns in the context of CO<sub>2</sub>R because it enables time-resolved measurements in three spatial dimensions. The spatial resolution of this technique can reach 250 nm under ideal conditions. The time resolution is, depending on the size and spatial resolution of the frame of interest, microseconds to several seconds. The combination of time resolution with sub-micrometer spatial resolution in three spatial dimensions with high accuracy is what makes this pOH-mapping technique unique. Importantly, it enables the map the local pOH value under operating conditions over a wide current density scale, from 0 to at least 200 mA/cm<sup>2</sup> in magnitude. To map the operando pOH in three dimensions with high resolution is important because it allows us to probe the pOH within microstructures on the surface of a sample of interest such as an active GDE. No other technique allows to probe the operando pOH within cavities in the surface of a GDE. This capability presents the unique opportunity to correlate the microstructure geometry of a GDE with its CO<sub>2</sub>R performance. Understanding the local chemical microenvironment near a CO<sub>2</sub>R catalysts is key for understanding selectivity and performance of these devices, as well as to validate continuum level theories of CO<sub>2</sub>R in porous electrodes.<sup>(58)</sup> Furthermore, our novel technique might also prove to be useful for the investigation of other electrochemical applications beyond CO<sub>2</sub>R.

In previous experiments,<sup>(49)</sup> we used the commercially available ratiometric fluorescent photoacid dye 6,8-dihydroxypyrene-1,3-disulfonic acid disodium salt (DHPDS), which is sensitive to pH values between 6 and 10. The term 'ratiometric fluorescent dye' refers to a probe with a spectrum that exhibits at least two distinct peaks with intensities changing in opposite directions upon varying the parameter to which the probe is sensitive. By evaluating the ratio between the peak intensities, the observed signal becomes independent of the local dye concentration.<sup>(50)-(52)</sup> The sensing mechanism of the photoacid DHPDS involves proton-transfer reactions in its electronic ground state to perturb its absorption spectrum. This is the mechanism used by nearly all fluorescent pH indicators in biological studies at near-neutral-pH conditions.<sup>(53)</sup> With DHPDS, we observed that the local pH at the GDE surface increased from pH 6.8 to greater than 10 as the magnitude of the current density was increased from 0 to -28 mA/cm<sup>2</sup> in 100 mM KHCO<sub>3</sub> electrolyte. We showed that the pH increased locally inside trenches 5 – 20 µm wide in the GDE surface. Furthermore, the pH increased as the trench width diminished which indicated that narrow trenches exhibit higher CO<sub>2</sub>R activity than wider trenches and planar surfaces.<sup>(49)</sup> While these results were insightful, they were limited by the pH range in which DHPDS is sensitive. We observed that

the DHPDS signal saturated at current density magnitudes larger than -20 mA/cm<sup>2</sup> in 100 mM KHCO<sub>3</sub>. However, there is an interest in operating GDEs at high current densities to increase the CO<sub>2</sub>R rate up to industrially relevant-levels ( $|J| > 100 \text{ mA/cm}^2$ ).<sup>(54),(55)</sup> Therefore, for this study, we developed a new sensing protocol based on our recent discovery of an excited-state proton-transfer mechanism for weak photoacids, such as 8-aminopyrene-1,3,6-trisulfonic acid trisodium salt (APTS). Unlike DHPDS, at nearneutral-pH and alkaline-pH conditions APTS does not undergo proton-transfer reactions in its electronic ground state. Instead, the fluorescent signal of APTS is altered via quenching of its thermally equilibrated electronic excited state by direct proton transfer to aqueous OH<sup>-.(56)</sup> We also show herein that aqueous dissolved inorganic carbon species do not interfere with this process (Figure S1). Owing to our newly characterized mechanism, the aromatic amine form of APTS can be used as a probe for the pOH value and is sensitive to pOH values between 0 and 2.8, compared to pOH 4 - 8 for DHPDS. It must be emphasized that this is the first demonstration of APTS as a sensor for the local pOH. APTS exhibits a novel sensing mechanism and greatly enhances the capabilities of our technique to map the local pOH value with CLSM. By combining DHPDS with APTS, we are now able to cover a pOH range from 0 to 8 (with a gap between 2.8 and 4) and investigate operating GDEs under current densities as large in magnitude as -200 mA/cm<sup>2</sup>. Additionally, we explore the influence of different bicarbonate concentrations in the electrolyte and of different microstructure geometries. All told, the work crucially enables a correlation between the bulk electrolyte properties and electrode structure with the pH present at the electrochemical active site across a broad range of operating current densities. We also evaluate the CO<sub>2</sub>R performance of our GDEs with gas chromatography measurements and supplement our experimental analysis with continuum modeling to enables a better understanding of the correlation between local pH and CO<sub>2</sub>R performance.

# **Experimental**

CLSM with the two fluorescent dyes DHPDS and APTS was used to map the pOH around an operating GDE performing CO<sub>2</sub>R over a wide pOH range. An electrochemical cell compatible with CLSM was developed, depicted schematically in Figure 1a as a cross section and Figure 1b as a top view. The electrochemical cell comprises of an electrolyte chamber with two perpendicular electrolyte inlets and outlets and is open at the top which allows the water immersion objective of the confocal microscope to be immersed into the electrolyte. For experiments at constant current densities, a leakless silver/silver chloride reference electrode and a platinum mesh counter electrode are submerged into the electrolyte. This cell enables us to study a GDE that is fed with gaseous  $CO_2$  through a gas chamber. The GDE is made of a Sigracet 22 BB carbon paper substrate covered with 300 nm Cu as well as a carbon black, graphite and Nafion coating. The substrate's surface is covered by an irregular pattern of trenches 5 – 30  $\mu$ m wide which is shown schematically as a cross section in Figure 1c as well as in a top-down scanning electron microscopy (SEM) image in **Figure 1d**. The trenches cover approximately 6% of the sample surface. Energy dispersive X-ray spectroscopy (EDS) measurements reveal that copper covers not only the planar carbon paper surface but also the trench walls and bottoms (Figure S2). The maximum spatial resolution of the confocal microscope is 250 nm in the x-y plane and 500 nm in the z-direction, however, electrolyte flow introduces noise, so the resolution is estimated to be on the order of one micron. We investigate features that are no smaller than 5  $\mu$ m, so this resolution is satisfactory.

We first used the dye APTS in the absence of electrical current to investigate the diffusion of  $CO_2$  through a GDE. For this, 1 M KOH electrolyte (pOH 0) with 100  $\mu$ M APTS was used. The fluorescence signal was mapped in the plane perpendicular to the electrode surface as a function of time with one frame captured approximately every four seconds. After one minute of continuous measurements, a 10 standard

cubic centimeters per minute (SCCM)  $CO_2$  gas stream was fed into the gas chamber of the electrochemical cell and the change in pOH in the electrolyte was observed. The measurements were performed both with and without circulating the electrolyte at a rate of 6 mL/min.

In addition, we used both dyes, DHPDS and APTS, to map the pOH around an operating GDE performing  $CO_2R$  at current densities between 0 mA/cm<sup>2</sup> and -200 mA/cm<sup>2</sup>. Measurements were performed in the plane parallel to the electrode surface, at 20 µm above the surface, at the surface, and at 20 µm below the surface inside a trench. A  $CO_2$ -saturated aqueous KHCO<sub>3</sub> solution with KHCO<sub>3</sub> at concentrations of 100 mM, 200 mM and 400 mM spiked with 100 µM DHPDS or 200 µM APTS was used as the electrolyte and pumped through two perpendicular inlets at a rate of 6 mL/min. The  $CO_2$  gas flow through the gas chamber was set to 10 SCCM. The electrolyte with dilute dye was removed from the cell and replaced after each measurement. Every measurement was performed at least three times.

We further evaluated the  $CO_2R$  performance of equivalent copper on carbon paper GDEs with gas chromatography. We used an electrochemical cell that was tailored for use with gas chromatography. More details about the sample fabrication and characterization, dye calibration, and measuring procedure can be found in the Supporting Information, section **S1** and **Figures S2 – S4**.

The development of the multiphysics model, including the physics, parameters, and assumptions employed, is described in the Supporting Information, section **S2** and **Figure S11**.



**Figure 1: (a)** Cross section of the electrochemical cell with water-immersion objective. The reference electrode is a leakless Ag/AgCl electrode, the counter electrode is a Pt mesh. **(b)** Top view of the electrochemical cell showing two perpendicular electrolyte inlets and outlets. **(c)** Schematic cross section of the carbon paper GDE (not to scale), the left hand side depicts the pOH increase if no current is passing, the right hand side shows the pOH decrease with applied current. **(d)** Top-down SEM image of the GDE surface (Sigracet 22BB carbon paper with 300 nm Cu).

#### **Results and Discussion**

First, we investigated how the local pOH in the electrolyte surrounding a GDE changed upon exposure to CO<sub>2</sub> at open circuit. This allowed us to draw conclusions about the diffusion of CO<sub>2</sub> molecules through a GDE. Our developed measuring technique is ideal for this study as it combines time resolution with the capability to spatially resolve the local pOH inside inhomogeneities in the surface of a GDE.  $CO_2$ reacts with OH<sup>-</sup> and water molecules in the electrolyte to form bicarbonate and carbonate anions. This increases the pOH which becomes an indicator for the CO<sub>2</sub> diffusion pattern. Figure 2a shows color-coded maps of the local pOH in the plane perpendicular to the electrode surface as a cross-section through a trench as well as the above electrolyte for different times without electrolyte flow. We define t = 0 s as the time when the CO<sub>2</sub> flow through the gas chamber of the electrochemical cell was turned on. A distinct change in pOH is immediately visible in the first frame after starting the CO<sub>2</sub> flow but it is restricted to the trench. The pOH changes at the electrode surface and in the bulk electrolyte as time proceeds. In Figure 2c, the average pOH at three different positions is tracked as a function of time. To this end, we draw a line 20  $\mu$ m above the electrode surface (+20  $\mu$ m), at the electrode surface (0  $\mu$ m), and 20  $\mu$ m below the electrode surface inside the trench (-20 µm) as indicated in the first panel of Figure 2a. The pOH is averaged along these lines as a function of time. The vertical gray line in Figure 2c indicates the time when the CO<sub>2</sub> gas supply was turned on. This data confirms that the pOH first changes inside the trench, then at the GDE surface and approximately five seconds later at 20 µm above the surface, which is consistent with the gaseous CO<sub>2</sub> feed coming from the backside of the GDE. After 20 s, the pOH values converge at all positions monitored and remain constant for the duration of the experiment. An analogous measurement was performed in a region of the electrode void of trenches (Figure S5) and we observed that the pOH increases at a slower rate in comparison to measurements at a location with a trench. This demonstrates that microstructures such as trenches promote faster CO<sub>2</sub> transport through a GDE substrate.

The experiment displayed in Figure 2a was repeated under an electrolyte flowrate of 6 mL/min (Figure 2b). This has a dramatic effect on the pOH around the GDE: The first change in pOH is only visible after 19 s, compared to less than 5 s without electrolyte flow. A significant increase in pOH caused by the diffusion of  $CO_2$  is solely observed inside the trench. Figure 2d tracks the pOH over time in a similar manner as Figure 2c. This confirms that the pOH increases little at the GDE surface as well as 20 µm above the surface, and even within the trench, the pOH remains below the value obtained without electrolyte flow. A steady state, with constant pOH up to 20  $\mu$ m above the electrode surface, is reached after 40 s. However, the pOH at the three locations never converges to a single value, in contrast to the measurements performed with stationary electrolyte. Electrolyte flow results in a well-defined boundary layer around the GDE surface which leads to a  $CO_2$  concentration gradient. The boundary layer is taken into consideration in the multiphysics model. In the absence of electrolyte recirculation, diffusion is the dominant transport mechanism for CO<sub>2</sub> and subsequent CO<sub>2</sub>R products. When the electrolyte is circulated however, convection dominates the mass transfer and causes OH<sup>-</sup> to be more quickly removed from the GDE surface. This prevents the pOH from rising as much as it would without electrolyte flow. As the electrolyte is more stationary inside trenches, a larger pOH increase can be observed in these confined spaces.



**Figure 2**: Changing pOH in the electrolyte due to  $CO_2$  diffusion around a trench in a carbon paper GDE at open circuit, without electrolyte flow (**a**, **c**) and with electrolyte flow (**b**, **d**). Measurements performed with 1M KOH electrolyte with 100  $\mu$ M APTS, 10 SCCM  $CO_2$  turned on at the backside of the electrode at t = 0 s. Frames in (**a**) and (**b**) show pOH maps as a cross section through a trench in the plane perpendicular to the electrode surface for different times. The scale bars apply to both (**a**) and (**b**). Panels (**c**) and (**d**) track the average pOH at locations 20  $\mu$ m above the surface (+20  $\mu$ m), at the surface (0  $\mu$ m) and 20  $\mu$ m below the surface in the trench (-20  $\mu$ m), as specified in the first frame of (**a**), as a function of time. The vertical gray lines indicate the start of the  $CO_2$  flow.

An additional study was conducted to compare the  $CO_2$  diffusion through homogeneous GDE substrates made of laminated polytetrafluoroethylene (PTFE) with pore sizes that are specified by the manufacturer as 0.1 to 0.2 µm and 0.45 µm, respectively. **Figure S6** shows a schematic representation of the structure of both a carbon paper GDE and a PTFE GDE together with SEM images. **Figure S7** compares the flow patterns through PTFE GDEs with different pore sizes. We observe that, as expected, the pOH in the electrolyte above the surface increases faster for PTFE with a larger pore size. After 45 seconds it converges to the same value for both substrates. This result gives evidence that, not only microcavities in a GDE's surface, but also larger pore size can promote faster  $CO_2$  transport. However, there is a critical pore size that should not be exceeded, otherwise the pores will be flooded with liquid electrolyte during an experiment. We observed this for a laminated PTFE substrate with pore sizes of 5 µm or greater. Flooding prevents effective  $CO_2$  transport and inhibits  $CO_2$  diffusion to catalytic sites.<sup>(10),(12)</sup> These results suggest that consideration of the influence of microstructures, even in commercially available GDE substrates, is a critical parameter to understand device performance.

The focus of our study is to investigate the local pOH around an operating GDE performing CO<sub>2</sub>R over a wide range of current densities. Our newly developed technique allows to perform pOH measurements with superior three-dimensional resolution inside cavities in an electrode under operating conditions. To ensure that our GDE samples (300 nm Cu on carbon paper) perform CO<sub>2</sub>R, we evaluated their CO<sub>2</sub>R performance with gas chromatography. We performed chronopotentiometry experiments for different current densities. Our results are consistent with copper GDEs previously reported in literature<sup>(15),(18),(57)</sup>. H<sub>2</sub> production dominates for low current density magnitudes. As the magnitude of the current density increases, we observe more CO<sub>2</sub>R products, and the selectivity shifts towards C<sub>2+</sub> products. We observe ethylene for current densities of -50 mA/cm<sup>2</sup> or higher in magnitude and ethanol for -200 mA/cm<sup>2</sup> (**Figure S8**). We can therefore assume that the analogous electrodes we use for our pOH imaging experiments behave as one would expect from literature.

By combining the two pOH-dependent fluorescent ratiometric dyes, DHPDS and APTS, we are able to cover the pOH range from 0 to 8. In the context of  $CO_2R$ , a low local pOH under operation indicates high  $CO_2R$  activity and is desirable because it has been shown that high pH can help suppress the parasitic HER and favor the formation of  $C_{2+}$  products.<sup>(8),(25),(32),(35),(36)</sup> **Figure 3** displays pOH maps in the plane parallel to the electrode surface in 100 mM KHCO<sub>3</sub> electrolyte for different current densities between 0 mA/cm<sup>2</sup> and -100 mA/cm<sup>2</sup> and at three different z-positions: 20 µm above the electrode surface, at the electrode surface, and 20 µm below the electrode surface inside a trench. The panels on the left-hand side were captured using DHPDS, and the ones on the right-hand side were captured using APTS.

We observed that APTS degrades and slowly loses its fluorescence for high current densities. We hypothesize that this is connected to the reduction of APTS at the electrode surface. A more detailed analysis of this effect can be found in the Supporting Information, section S1. Despite the slow degradation of APTS, the pOH measurements obtained with APTS are reliable because APTS is a ratiometric dye. The pOH value is calculated from the ratio between two signals that are captured independently in two different wavelength intervals. As some dye molecules degrade, both signals become weaker. This leads to a decreased signal-to-noise ratio, however, the ratio between both signals remains unchanged (see Figure S9). The dye degradation effect could be mitigated by using an APTS concentration of 200  $\mu$ M (compared to DHPDS, where 100  $\mu$ M is sufficient), removal and replacement of the electrolyte after each measurement, and the introduction of two perpendicular electrolyte inlets to ensure the transportation of fresh APTS to the GDE surface. Some of the panels under high current densities still appear relatively dark and noisy. This is especially evident inside the trench as the electrolyte is more stationary in confined spaces and this encumbers the transport of fresh, undegraded dye. However, since APTS is a ratiometric dye, the pOH it predicts does not depend on concentration. At magnitudes of the current density as high as -200 mA/cm<sup>2</sup>, we were still able to collect enough fluorescence signal to determine the pOH.

For a current density of J = 0 mA/cm<sup>2</sup>, the pOH equals 7.2 everywhere (pOH of CO<sub>2</sub>-saturated 100 mM KHCO<sub>3</sub>). When the current is non-zero, the local pOH decreases because OH<sup>-</sup> is created as a byproduct of CO<sub>2</sub>R. The pOH is lower at the electrode surface than 20  $\mu$ m above the surface because CO<sub>2</sub> is reduced at the electrode surface and for the pOH to decrease at +20  $\mu$ m, OH<sup>-</sup> has to diffuse away from the surface. Due to electrolyte flow, a concentration gradient is created. Furthermore, the pOH inside the trench is lower than at the surface. This can be seen especially well in the panels for current densities of -2 mA/cm<sup>2</sup> and -20 mA/cm<sup>2</sup>.

The surface morphology of copper does not change during CO<sub>2</sub>R experiments. We performed SEM as well as EDS measurements for samples before and after CO<sub>2</sub>R. We observed potassium deposits on the sample after CO<sub>2</sub>R that originate from KHCO<sub>3</sub> molecules in the electrolyte, but the copper catalyst doesn't change in appearance, neither in the trenches nor on the planar electrode surface (see **Figure S2**).



**Figure 3**: pOH maps in the plane parallel to the electrode surface 20  $\mu$ m above the surface, at the surface and 20  $\mu$ m below the surface inside a trench in 100 mM KHCO<sub>3</sub> electrolyte for different current densities between 0 mA/cm<sup>2</sup> and -100 mA/cm<sup>2</sup>, obtained with 100  $\mu$ M DHPDS (left) and 200  $\mu$ M APTS (right). Note the different color legends for the panels obtained with DHPDS and APTS.

**Figure 4** shows the average pOH as a function of current density, both from experiments and from multiphysics simulations. For the experimental values, the pOH is averaged over at least three independent measurements. The data for low current densities was obtained with DHPDS, while the data for higher current densities was obtained with APTS. The red shaded areas indicate the pOH range that is inaccessible by either dye. The insets are a zoom-in on the experimental data for low current densities.

Simulations were performed to understand better the variations of pOH with respect to position, electrolyte concentration and trench geometry. A 2-dimensional continuum model of a periodic trench in a GDE was developed. The multiphysics simulations solve for the concentration profiles of all species involved, along with the liquid electrolyte potential in a single, periodic trench of a given width and depth. Concentration-dependent Tafel kinetics are employed to simulate the electrochemical reduction of CO<sub>2</sub> to various products on the Cu electrode surface.<sup>(35),(36)</sup> Greater detail regarding the physics, parameters, and assumptions employed in the continuum model (shown schematically in **Figure S11**) can be found in the Supporting Information, section **S2**. The model replicates the experimental results with outstanding accuracy. This is true both for the pOH values observed in all panels in **Figure 4** as well as for the product distribution that was measured with gas chromatography. The agreement between model and experiment gives us confidence that the model is a good representation of the real world and that predictions that go beyond what can be observed in experiments can be trusted.

**Figure 4a** compares the pOH in 200 mM KHCO<sub>3</sub> electrolyte at positions around a GDE analogous as shown in **Figure 3**. As the current density increases, the pOH decreases accordingly, due to the generation of OH<sup>-</sup> from the CO<sub>2</sub>R reactions. At first, the slope of the pOH decrease is steep, then gradually tapers as the current density increases. This can be explained by the logarithmic nature of the pOH scale. A decrease in pOH from 4 to 3 requires the OH<sup>-</sup> concentration to increase from 0.1 mM to 1 mM, while a decrease in the pOH from 2 to 1 requires the OH<sup>-</sup> concentration to rise from 10 mM to 100 mM, a much larger absolute change in concentration. The pOH in trenches is lower than at the surface, which again is lower than 20  $\mu$ m above the surface for all current densities; this confirms our previously described results. The simulations agree well with the CLSM experiments, particularly for data at the electrode surface (z = 0  $\mu$ m) and in the trench (z = -20  $\mu$ m). Agreement between experiments and simulations is similarly strong for bulk electrolyte concentrations of 100 mM and 400 mM (**Figure S10a-c**). The greatest discrepancy between experiments and simulations is apparent in the z = +20  $\mu$ m data, which more closely represents the pOH of the bulk electrolyte. The discrepancy is likely due to the static nature of the boundary layer in the simulations, which is set to a constant value of 150  $\mu$ m, regardless of the applied current density. Previous experimental studies have shown that the bulk electrolyte pH, and thus, the boundary layer thickness is a function of current density at higher current densities.<sup>(25),(34)</sup> However, implementing an adaptive, 2-D mesh that grows with current would be very computationally intensive, and for the present study, the pOH in the trench, where CO<sub>2</sub>R occurs, is more critical to simulate accurately.

Figure 4b displays a study of the influence of KHCO<sub>3</sub> concentration in the electrolyte on the pOH at the GDE surface. The trend observed at the electrode surface is the same as it is 20 µm above the electrode surface and 20 µm below in the trenches (Figure S10d-f). Measurements for higher concentrations could be carried out at higher current densities due to the enhanced electrical conductivity of the electrolyte. Again, the model generally agrees with observed experimental trends that show that the pOH decreases more slowly as electrolyte concentration is increased. This is because KHCO<sub>3</sub> acts as a buffer and, as the KHCO<sub>3</sub> concentration increases, the buffering capacity is enhanced, which restrains the decrease in pOH. However, at higher current densities, the simulated pOH becomes more independent of current density whereas the experimental pOH remains impacted by buffer capacity. This discrepancy at higher current densities can be attributed to a multitude of factors, such as the large uncertainty in the rate constants for the homogeneous-phase buffer reactions used in the simulations, (28), (58), (59) and the fact that dilute-solution theory was employed in the simulations whereas concentrated solution theory may be necessary to simulate adequately the behavior at higher current density.<sup>(60)</sup> Additionally, the simulations assume that there are no mass transport losses of gas phase CO<sub>2</sub> throughout the GDE itself, and thus the CO<sub>2</sub> concentration is at a constant value of 34 mM at the electrode surface. This assumption is likely unrealistic at high current densities and could potentially impact the simulated pOH due to the participation of  $CO_2$  in homogeneous buffer reactions. Nonetheless, the simulations are capable of generally capturing the impact of buffer composition, and the incorporation of these second-order phenomena is beyond the scope of the present study.

Lastly, the influence of the trench geometry on the local pOH was investigated (**Figure 4c**). Eighteen trenches with different widths in 100 mM KHCO<sub>3</sub> were taken into consideration. The average pOH inside the trenches, i.e. 20  $\mu$ m below the surface, was evaluated at different current densities. The trenches were divided into widths of 5 - 10  $\mu$ m, 10 - 15  $\mu$ m, 15 - 20  $\mu$ m, and >20  $\mu$ m and the average pOH was calculated for each. Both measurements and simulations agree that the pOH decreases as trench width diminishes for all current densities investigated. This behavior is due to the increased surface-areato-volume-ratio of the thinner trenches (**Figure S13**). The increased surface-area-to-volume ratio means that more OH<sup>-</sup> is generated via CO<sub>2</sub>R on the trench walls relative the electrolyte volume contained within a trench. In addition, the mass transfer out of trenches is poor compared to a planar surface, which leads to trapping of OH<sup>-</sup>. Thus, the OH<sup>-</sup> concentration is increased for thinner trenches as a result of both the increased surface-area-to-volume-ratio as well as mass transport effects. This result is in line with previously reported electrochemical studies that have demonstrated that smaller pore radii in meso- or micro-structured electrodes enhance local OH<sup>-</sup> trapping.<sup>(54),(62),(63)</sup> Interestingly, the measured pOH is less

sensitive to trench width at higher current densities than the simulated pOH. A possible explanation for this discrepancy could be the lack of bubble-induced convection in the present simulations. At high current densities, the latter could ameliorate  $OH^-$  trapping within GDE trenches and reduce sensitivity to trench geometry. The influence of trench width on the pOH further emphasizes the importance to understand how a GDE's micrometer-scale geometry affects  $CO_2R$ .

When we combine the results from **Figure 4 (a)** and **(c)**, we find that the pOH is decreased inside trenches compared to the planar electrode surface and furthermore, this effect is stronger for narrower trenches. This observation is caused by the altered mass transport inside narrow trenches. Our experiments therefore suggest that the micrometer-scale morphology of a gas diffusion electrode plays an important role in determining the mass transport properties of a GDE, and with it the  $CO_2R$  performance.



**Figure 4**: The pOH as a function of current density at different positions (a), for different KHCO<sub>3</sub> concentrations (b) and for different trench widths (c), in experiments and simulations. (a) Averaged pOH as a function of current density for 200 mM KHCO<sub>3</sub>, obtained with 100  $\mu$ M DHPDS ( $J \ge -13$  mA/cm<sup>2</sup>)/ 300  $\mu$ M APTS ( $J \le -20$  mA/cm<sup>2</sup>) at three different positions around the electrode surface together with simulation results. (b) Averaged pOH as a function of current density at the electrode surface for three different KHCO<sub>3</sub> concentrations between 100 and 400 mM, obtained with 100  $\mu$ M DHPDS ( $J \ge -15$  mA/cm<sup>2</sup>)/ 200-300  $\mu$ M APTS ( $J \le -20$  mA/cm<sup>2</sup>) together with simulation results. (c) pOH inside a trench 20  $\mu$ m below the surface as a function of current density for different trench widths between 5  $\mu$ m and 30  $\mu$ m. In all panels, the pOH range shaded in red is not accessible by either dye, the inset is a zoom on the data for low current densities.

Beyond elucidating trends observed in the CLSM experiments, continuum simulations can be used to identify pOH variations at resolutions that are not achievable experimentally. **Figure 5a** depicts  $OH^$ transport within a simulated periodic GDE trench operating at a current density of -50 mA/cm<sup>2</sup> ( $OH^$ transport for -1, -10, and -100 mA/cm<sup>2</sup> is depicted in **Figure S15**). As shown in the contour plot, the pOH is lowest, and thus the concentration of  $OH^-$  is greatest deep within the trench. Examination of the  $OH^$ flux rationalizes the diminished pOH within the trench:  $OH^-$  generated via  $CO_2R$  in the trench exits to the bulk electrolyte, but the presence of the trench increases the surface area available for the electrochemical reduction of  $CO_2$  and, accordingly, the generation of  $OH^-$ . The trench also provides a volume where the generated  $OH^-$  can accumulate due to the altered mass transport.  $OH^-$  generated by  $CO_2R$  in the trench is thus effectively trapped within the trench, decreasing the local pOH. Similar  $OH^$ trapping behavior has been observed in prior multi-dimensional continuum studies of mesostructured electrodes<sup>(38),(58)</sup> and has been shown to be beneficial for  $CO_2R$  selectivity.<sup>(38)</sup> To evaluate the influence of the pOH gradient within the trench on local selectivity variations, concentration-dependent Tafel kinetics were employed in the multiphysics model. These concentration dependent Tafel kinetics were fit to experimental data collected on Cu GDEs used in the present study, and are capable of replicating observed partial current densities quite well (**Figure S12**). The study relates local changes in OH<sup>-</sup> activity to changes in current densities for product formation. **Figure 5b** depicts the change in pH as a function of depth into the trench. Note that **Figure 5b** shows the pH value while for the most part, this study focuses on the dynamics of OH<sup>-</sup> and hence, the pOH. Here, we chose to display the pH instead to emphasize the correlation between **Figure 5b** and **5c**. In the model, the relation pH + pOH = 14 holds true everywhere because we only consider the diffuse transport layer while the electrical double layer is not taken into consideration in the model because its length scale is much smaller than the length scales we study. Therefore, within the frame of this model, pH and pOH are directly connected. Consistent with **Figure 5a**, pH increases (pOH decreases) deeper inside the trench (more negative *z*) due to trapping of CO<sub>2</sub>R-generated OH<sup>-</sup>.

The simulations exhibit a direct, albeit modest, relationship between pH and  $C_{2+}$  FE (**Figure 5c**). As the pH increases or the pOH decreases within the trench, the  $C_{2+}$  FE correspondingly increases (variations of  $C_1$ ,  $H_2$ , and local current density within the trench are shown in **Figure S15**). This relationship between  $C_{2+}$  FE and pH has been widely reported, and is connected to the suppression of HER and  $C_1$ products.<sup>(9),(19),(35),(36),(63)</sup> The disproportionate suppression of H<sub>2</sub>, CH<sub>4</sub>, and HCOOH partial current density within the trench can be observed in **Figure S16**. This suppression of  $C_1$  and  $H_2$  in the trench results in an approximately 5% increase in  $C_{2+}$  FE across the depth of the trench, regardless of current density (**Figure 5c**). The enhancement of the performance in the trench is modest, as the change in pH is only about 1 pH unit across the trench depth. Nonetheless, the simulations serve to understand local selectivity trends and provide better understanding of variations in performance within the GDE heterogeneities.

Lastly, performance of CO<sub>2</sub>R within trenches was compared to the performance on a flat surface without trenches (Figure 5d). The simulations demonstrate that the  $C_{2+}$  FE at a given current density is higher within a trench compared to that on a flat surface due to the improved OH<sup>-</sup> trapping. In addition, as the width of the trench decreases, and with it the pOH (Figure 4c and Figure S18), the  $C_{2+}$  FE increases. This is a result of the changed mass transport properties in the confined space as well as the enhanced trench surface-area-to-volume ratio (Figure S13), leading to a lower pOH value. The simulations indicate that the micrometer-scale morphology on a GDE surface has an impact on the product distribution with narrow trenches on the order of 5  $\mu$ m favoring the creation of C<sub>2+</sub> products across the entire current density range (Figure 5d) by approximately 5% compared to the flat Cu case. Within the scope of this work, the connection between surface morphology and CO<sub>2</sub>R selectivity can only be demonstrated via multiphysics simulations because we have no means to locally resolve the product distribution. However, the accuracy of our model has been demonstrated as there is an excellent agreement with the experimentally measured CO<sub>2</sub>R performance as well as the local pOH. Nonetheless, the simulations demonstrate the potential for improved  $C_{2+}$  FE within these microcavities and experimental demonstration of enhanced C<sub>2+</sub> FE inside microcavities in the surface of an operating GDE should be the subject of a future study.

The effect of the trench depth on  $CO_2R$  was also evaluated (**Figure S19**), which demonstrates that the ability of a trench to enhance  $C_{2+}$  FE increases as the trench deepens. This is due to the increased surface area available for electrochemical OH<sup>-</sup> generation for deeper trenches. The surface-area-tovolume ratio is far less sensitive to trench depth than to trench width (**Figure S18**) which indicates that compared to the surface area, the volume plays a minor role in determining the  $C_{2+}$  FE. However, enhancements in  $C_{2+}$  FE become marginal beyond trench depths of 50 µm. Finally, the fraction of the surface covered by the trenches was varied in the simulations, to further evaluate the efficacy of trenches. **Figure S20** depicts the pOH and  $C_{2+}$  FE as a function of trench coverage, evaluated at current densities of -20 mA/cm<sup>2</sup> and -50 mA/cm<sup>2</sup> (normalized now to the electrochemical active surface area (ECSA)). This figure aims to deconvolute the effects of roughness from the effects of changes in the microenvironment. As the trench surface coverage increases, the pOH decreases, and the  $C_{2+}$  FE increases, due to the OH<sup>-</sup> trapping ability of the trenches. While the increase in  $C_{2+}$  FE is minor (still approximately 5 % from a flat surface to a completely trenched surface), the enhancement with morphology is similarly observed at both current densities and the results reveal that heterogeneities in commercially available GDEs can serve as local hotspots for enhanced  $C_{2+}$  FE and benefit CO<sub>2</sub>R due to their OH<sup>-</sup> trapping capacity. The simulation results serve as a great support to our novel measurements that allow to probe the microenvironment in a GDE under operando conditions. Both experiment and simulations highlight the importance of the micromorphology of a GDE that influences the mass transport properties. Future work should aim to leverage this insight to develop geometries for which transport is tailored to better trap OH<sup>-</sup>. Techniques like laser ablation can be utilized to realize these tailored electrodes.



**Figure 5**: (a) pOH contour plot and OH<sup>-</sup> flux (white arrows) throughout the simulated domain for a periodic GDE trench (width of 20  $\mu$ m and depth of -100  $\mu$ m) operating at -50 mA/cm<sup>2</sup> of geometric current density (see section **S4** for a description of how geometric current density is calculated). (b) Simulated electrode surface pH (14 – pOH) as a function of depth into the trench for different current densities. (c) Simulated

 $C_{2+}$  FE as a function of depth into the trench for different current densities. **(d)** Average  $C_{2+}$  FE on trench walls as a function of the current density passed solely through the trench.  $Z = 0 \mu m$  represents the top of the trench, and  $Z = -100 \mu m$  represents the bottom of the trench. d and w are the trench depth and width, respectively.

# **Conclusions**

We demonstrate a novel technique to probe the microenvironment in a CO<sub>2</sub>R GDE under operating conditions. We map the local pOH value around a GDE resolved in time and three spatial dimensions with fluorescent confocal laser scanning microscopy (CLSM) with a combination of two ratiometric dyes, DHPDS and APTS, that operate by different sensing mechanisms. The novel application of APTS as a dynamic excited-state pOH probe was demonstrated for the first time. Critically, this work could hold potential applications beyond the investigation of CO<sub>2</sub>R. We were able to capture pOH maps in the pOH range between 0 and 8 and at current densities as large in magnitude as -200 mA/cm<sup>2</sup>.

In a first step, we investigated the diffusion patterns of  $CO_2$  through a GDE. This study revealed that  $CO_2$  diffuses through microcavities first, and that electrolyte flow leads to a  $CO_2$  concentration gradient with the highest concentration confined to trenches. These microcavities in a GDE surface as well as large pore sizes promote faster  $CO_2$  transport, highlighting the importance to understand the influence of the micron-scale morphology of a  $CO_2R$  GDE.

We mapped the local pOH around an operating GDE under applied current. This revealed that the pOH in the vicinity of a GDE decreases as current density is increased. For all current densities covered, the pOH inside trenches is lower than on the planar surface with the lowest pOH observed inside narrow trenches that are approximately 5  $\mu$ m wide. This decrease in pOH can be connected to the trapping of OH<sup>-</sup> in narrow trenches caused by altered mass transport properties.

Multiphysics simulations confirmed all experimental trends, providing pOH maps consistent with measured pOH and also reproduced product distributions with high accuracy. These simulations indicate that there is a correlation between the pOH and the Faradaic efficiency towards  $C_{2+}$  products. Narrow trenches exhibit a decreased pOH and simulations suggest that this results in a locally increased Faradaic efficiency towards  $C_{2+}$  products.

We conclude that we successfully developed a superior technique to probe the microenvironment around a CO<sub>2</sub>R GDE. The micrometer-scale morphology of a GDE has a strong effect on the mass transport properties and with it, on the CO<sub>2</sub>R performance. Microcavities on the order of 5  $\mu$ m on the surface of a copper GDE serve as local hotspots with decreased pOH. We expect that this will inform the design of future CO<sub>2</sub>R devices.

# **Author Contributions**

AB performed dye calibration and all pOH imaging experiments as well as data analysis and wrote the original draft of the manuscript. JCB and AJK developed the multiphysics model with oversight by AZW. JCB ran simulations and performed data analysis on results from the multiphysics model. AF conducted the gas chromatography measurements. RB contributed to conceptualization of the choice of APTS photoacid. CNF investigated possible interference in signal from APTS due to dissolved inorganic carbon

species. AJW developed the experimental setup and designed the electrochemical cell. AB, AF, RB, AJW, SA and HAA contributed to the conceptualization of the study. JCB, AF, RB, AJW, ATB, AZW, SA and HAA reviewed and edited the manuscript. ATB, AZW, SA and HAA supervised the project. All authors discussed the results and contributed to the preparation of the manuscript.

## **Conflicts of Interest**

There are no conflicts to declare.

## **Acknowledgements**

This material is based on work performed within the Liquid Sunlight Alliance, which is supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Fuels from Sunlight Hub under Award Number DE-SC0021266. JCB would like to acknowledge support from the National Defense Science and Engineering Graduate Fellowship (NDSEG) supported by the Army Research Office (ARO). The authors thank Ian Sullivan for 3D-printing the electrochemical cell.

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