Electrochemical Regeneration of Spent Alkaline Absorbent from Direct Air Capture

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ABSTRACT: CO₂ capture from the atmosphere (or direct air capture) is widely recognized as a promising solution to reach negative emissions, and technologies using alkaline solutions as absorbent have already been demonstrated on a full scale. In the conventional temperature swing process, the subsequent regeneration of the alkaline solution is highly energy-demanding. In this study, we experimentally demonstrate simultaneous solvent regeneration and CO₂ desorption in a continuous system using a H₂-recycling electrochemical cell. A pH gradient is created in the electrochemical cell so that CO₂ is desorbed at a low pH, while an alkaline capture solution (NaOH) is regenerated at high pH. By testing the cell under different working conditions, we experimentally achieved CO₂ desorption with an energy consumption of 374 kJ mol⁻¹ CO₂ and a CO₂ purity higher than 95%. Moreover, our theoretical calculations show that a minimum energy consumption of 164 kJ mol⁻¹ CO₂ could be achieved. Overall, the H₂-recycling electrochemical cell allowed us to accomplish the simultaneous desorption of high-purity CO₂ stream and regeneration of up to 59% of the CO₂ capture capacity of the absorbent. These results are promising toward the upscaling of an energy-effective process for direct air capture.

1. INTRODUCTION

Climate change is one of the most critical global challenges nowadays. Increasing atmospheric CO₂ concentration brought by anthropogenic emissions has been recognized as the primary driver of global warming.¹⁻⁵ The global average surface temperature has risen by 0.85 °C in the period from 1880 to 2012, and over 85% of this temperature rise occurred in the past century can be attributed to industrialization.⁶ Besides, most of the CO₂ in the atmosphere will ultimately be absorbed by the oceans causing ocean acidification, which has been proved harmful to marine organisms.⁷ The ocean pH has been already reduced by 0.1 units compared to the preindustrial level,⁸ and it could eventually be 0.7 units lower with continuous emission of CO₂ from the combustion of fossil fuels.⁹

In this context, it has been widely recognized that capturing CO₂ from emission points and even directly from the air provides a potential solution to mitigate CO₂ emissions and ultimately reduce the atmospheric CO₂ concentration.¹⁰,¹¹ In particular, direct air capture (DAC) technologies aim to extract CO₂ directly from ambient air and hence offer the possibility to achieve CO₂ capture from distributed sources.¹²⁻¹⁴ Therefore, DAC is not restricted by location as conventional carbon capture technologies that are designed to be coupled with large CO₂ emission points. The thermodynamic minimum energy requirement for DAC is relatively higher compared to carbon capture from flue gas, due to the low CO₂ concentration in air.¹³ Nonetheless, further studies on DAC are essential for reducing climate risks.

Various technologies have been proposed for DAC,¹⁴⁻¹⁶ and one of the most extensively studied approaches is wet scrubbing with alkaline hydroxide solutions (typically NaOH or KOH).¹⁶⁻¹⁸ When air is in contact with an alkaline solution (e.g., NaOH), CO₂ is absorbed, and a sodium carbonate precipitate is thermally treated at 700 °C to produce CO₂ and calcium oxide. Finally, calcium hydroxide can be regenerated by rehydration of calcium oxide.¹⁹

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2NaOH + CO₂ → Na₂CO₃ + H₂O
ΔH⁰ = −109.4 kJ mol⁻¹
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Conventionally, the sorbent (NaOH) is regenerated by exchanging sodium and calcium ions by dosing calcium hydroxide. Subsequently, the resulting calcite (CaCO₃) precipitate is thermally treated at 700 °C to produce CO₂ and calcium oxide. Finally, calcium hydroxide can be regenerated by rehydration of calcium oxide.
Since the chemical reaction between NaOH and CO₂ is spontaneous, the regeneration of the alkaline sorbent (coupled with desorption of CO₂ gas) consumes the largest fraction of energy in the conventional wet scrubbing process (i.e., a minimum of 179 kJ·mol⁻¹ CO₂ at standard temperature and pressure²⁸,²⁹). Alternative regeneration processes have been developed to reduce energy consumption. For instance, Mahmoudkhani and Keith proposed a titration cycle that halved the energy requirements compared to the conventional regeneration process, while Kim et al. suggested an electrochemical method for the regeneration of LiOH from used CO₂ adsorbents. The use of electrochemical systems as the CO₂ desorption step includes a number of advantages such as (i) potentially low energy consumption (since the process occurs at room temperature), (ii) better operational control (by exploiting the electric field), and (iii) easy integration with renewable energy sources (since the process requires only electricity as energy input).²¹−²⁸

In this work, we developed an electrochemical process for regenerating the alkaline solution in the wet scrubbing process for DAC application; meanwhile, high-purity CO₂ gas stream could be produced that is suitable for storage or utilization. The process is based on a H₂-recycling electrochemical system (HRES) coupled to a membrane contactor that was originally developed for nitrogen recovery from wastewater.²⁹,³⁰ In HRES, alkaline and acidic conditions are created in two adjacent compartments, respectively. Additionally, CO₂ equilibria are highly dependent on the pH of the solution (eqs 2−4, values taken from refs 31, 32). Accordingly, we adapted the system operation for DAC application. The electrochemical cell consisted of three compartments, i.e., an anode compartment, an “acidifying” compartment, and a cathode compartment. During the operation, protons (H⁺) produced from the H₂ oxidation at the anode are transported to the acidifying compartment where the spent solution coming from the air contactor is fed. The decreasing pH of the acidifying solution leads to the conversion of (bi)carbonate ions to carbonic acid (eqs 2 and 3). When the solution is saturated in carbonic acid (H₂CO₃*), including dissolved CO₂, a further pH decrease leads to the desorption of CO₂ gas (eq 4). The oversaturation point is determined by the CO₂ partial pressure in the membrane contactor. Meanwhile, NaOH is regenerated in the cathode due to the production of hydroxides (OH⁻) and can be reused as a DAC absorbent.

\[
\begin{align*}
\text{HCO}_3^- & \rightleftharpoons \text{CO}_3^{2-} + \text{H}^+ & \text{pK}_{a1} = 10.33 \\
\text{H}_2\text{CO}_3 & \rightleftharpoons \text{HCO}_3^- + \text{H}^+ & \text{pK}_{a1} = 6.35 \\
\text{CO}_2 + \text{H}_2\text{O} & \rightleftharpoons \text{H}_2\text{CO}_3 & K_{H} = 0.00339 \text{ mol·L}^{-1} \cdot \text{atm}^{-1}
\end{align*}
\]

In this work, we show a proof of principle of such an electrochemical method for NaOH regeneration and CO₂ desorption using an HRES. Different operational conditions were studied to identify suitable conditions to regenerate the absorbent at low energy consumption.

2. MATERIALS AND METHODS

2.1. Experimental Methodology. 2.1.1. Experimental Setup. A schematic representation of the experimental setup is shown in Figure 1. The main process streams are the influent flowing into the acidifying compartment and then the membrane contactor, the overflow of excess acidifying solution flowing into the catholyte tank, and the effluent from the cathode compartment. To increase the flow speed inside the cell, two recirculation loops were added for the acidifying and cathode compartments, respectively. The anode was a membrane electrode assembly (MEA) consisting of a 15 cm × 15 cm Nafion N117 cation exchange membrane (CEM) integrated with a 10 cm × 10 cm gas diffusion layer (GDL) coated by a Platinum—Vulcan (carbon) catalyst (0.5 mg Pt·cm⁻²) (FuelCellsEtc, TX). The GDL side of the MEA was facing the anode compartment, while the CEM side was facing the acidifying compartment. The flow channel of the acidifying compartment was created using a polymeric (nitril) spacer (500 µm, Sefar, Switzerland). The acidifying compartment and the cathode compartment were separated by a CEM (Nafion N117, 15 cm × 15 cm, FuelCellsEtc, TX). Two Ru/Ir-coated titanium mesh electrodes (9.8 cm × 9.8 cm, Magneto Special Anodes BV, The Netherlands) were used in the cell, one as cathode and the other one as the current collector for the anode. The whole cell was closed between two poly(methyl...
The Na+ in the system has mainly two roles: (i) supporting the system can operate without any net H2 consumption by being transported through the membrane to sustain the ionic current and fed with a 25 mM H2SO4 aqueous solution. The gas-liquid separation was conducted by a membrane contactor (type MM 1.7 × 8.75, 3M, Germany) whose microporous hollow fiber membrane provided a large surface area for contact between gas and liquid phases. The produced gas first passed through a Nafion tubing (TUB-0003, CO2Meter.com, FL) to remove water vapor. Next, the amount of gas produced was measured by a mass flow meter (ELFLOW Prestige FG-111B, Bronkhorst, The Netherlands). Finally, the gas composition was checked by micro-gas chromatograph (Varian CP-4900, Agilent, CA).

Three peristaltic pumps (Masterflex L/S, Metrohm Applikon BV, The Netherlands) were used in the system, one for the inlet, one for the effluent, and one for the recirculation of solution in acidifying and cathode compartments (150 mL min-1). The total volume of the acidifying compartment recirculation was 180 mL including the flow channels of the acidifying compartment (~5 mL), membrane contactor (~70 mL), and tubing (~105 mL). The cathode compartment recirculation had a total volume of 120 mL.

The anode (MEA) potential and cathode potential were measured by two Ag/AgCl reference electrodes (+0.2 V vs NHE, QM711X, QIS-Prosense BV, The Netherlands) placed at the inlets of the acidifying and cathode compartments, respectively. Two high-impedance preamplifiers (Ext-Ins Technologies, The Netherlands) were connected to the reference electrodes to avoid current leakage. The conductivities of the solutions in the acidifying and cathode compartments were measured using a conductivity sensor (Memosens CLS82D, Endress+Hauser BV, The Netherlands). The pH of the solution in the acidifying compartment was measured using a pH sensor (Orbisint CPS11D, Endress+Hauser BV, The Netherlands). A constant current was applied to the electrochemical cell by a power supply (ES 030-5, Delta Elektronika BV, The Netherlands). The pH of the solution in the acidifying compartment reached equilibrium. As a result, the spent absorbent solution consists of 0.175 M Na2CO3 and 0.150 M NaHCO3, based on Henry’s law and the CO2 equilibria shown in eqs 2–4. The spent absorbent solution has a pH of 9.7 and a conductivity of 30.6 mS·cm-1.

First, experiments at different Na+ values were performed while keeping the current density constant. Then, the effect of current density was investigated at constant Na+ values. The experiments were performed until the system reached a steady state (i.e., by monitoring the outlet conductivity). Table S1 shows an overview of all of the experiments that were performed.

2.1.4. Figures of Merit. Here, we define the figures of merit associated with energy consumption and the regeneration of the alkaline absorbent. The specific energy consumption of the system (EC, J mol-1) is defined as

\[
EC = \frac{V_{cell} \cdot I + P_H_2}{j_{CO_2}}
\]  

(6)

where \(V_{cell}\) is the cell voltage (V), \(I\) is the applied current (A), \(P_H_2\) is the power for production of 10% excess H2 in the electrolyzer (W), and \(j_{CO_2}\) is the measured CO2 production rate (mol·s-1). \(P_H_2\) is included in the calculations to compensate for possible small H2 gas leakage in practice. Note that we assumed the H2 production at the cathode fully compensates the H2 requirement at the anode in optimized cell designs (i.e., gas-tight).

The primary purpose of the system is to regenerate the alkaline absorbent. As the maximum total carbon concentration in the spent absorbent is determined by the CO2 partial pressure in the air, the regenerated CO2 capture capacity of the absorbent equals the amount of total carbon removed during the regeneration. The total carbon removed during the regeneration is in the form of desorption of CO2(g).

Therefore, we define a “carbon removal efficiency” (\(\xi_{removal}\)) to quantify the efficiency of the absorbent regeneration step, and it is expressed as

\[
\xi_{removal} = \frac{C_{s,CO_2}}{C_{s,CO_2} \cdot \tau - \frac{Q \cdot F}{c_{Na^+}}}
\]

(5b)
where $c_{i,2}$ and $c_{i,0}$ are the total carbon concentration in the cathode out and influent, respectively (mol·m$^{-3}$).

Finally, we describe the potential drop over the cell as a sum of equilibrium potential ($V_{eq}$), anode overpotential ($\eta_{anode}$), cathode overpotential ($\eta_{cathode}$), ionic losses ($V_{ionic}$), and transport losses ($V_{transport}$). Further details on the calculations can be found in the Supporting Information.

### 2.2. Equilibrium Model

An equilibrium model was developed for the acidifying and cathode compartments, based on mass balance and transport equations of Na$^+$ and H$. The model considers steady-state conditions, well-mixed compartments, and ideal CEM behavior (i.e., a complete rejection of co-ions). The anode was excluded from this framework to simplify the system. Hence, we also assume that protons are produced in the acidifying compartment and consumed at the cathode (i.e., assuming hydrogen evolution from protons instead of from water). Furthermore, the recirculation of each compartment is not considered in the model, and the outlet from the acidifying compartment flows directly to the cathode.

Neglecting any net transport of water through the membrane, the mass balance for sodium in the acidifying compartment can be written as

$$J_{Na^+} = \frac{Q}{A}(c_{Na^+,0} - c_{Na^+,1})$$

where $J_{Na^+}$ is the molar flux of Na$^+$ through the CEM, and $c_{Na^+,0}$ and $c_{Na^+,1}$ are the Na$^+$ concentrations at the inlet and outlet of the acidifying compartment, respectively.

The mass balance of total carbon species is related to the CO$_2$ production in the acidifying compartment. The total carbon concentration in the anode outflow ($c_{a,1}$) in equilibrium with the gas phase in the membrane contacting (1 atm pure CO$_2$) is thus a function of sodium concentration ($c_{Na^+,1}$) and CO$_2$ partial pressure. If $c_{a,1}$ is lower than the total carbon concentration in the anode inflow ($c_{a,0}$), the reduction of total carbon in the liquid phase is desorbed into the gas phase as CO$_2$ gas. Therefore, the specific CO$_2$ production rate ($J_{CO_2}$) can be defined as

$$J_{CO_2} = \frac{Q}{A}(c_{CO_2,0} - c_{CO_2,1})$$

Note that $J_{CO_2}$ is normalized for the active area of the cell. If $c_{CO_2,1}$ is calculated to be higher than $c_{CO_2}$ we consider the acidifying compartment as a closed system, which has no gas production and $c_{CO_2,1} = c_{CO_2}$.

In the cathode compartment, since ideally no carbon species can pass through the membrane, the total concentration of carbon species remains constant, i.e., $c_{c,2} = c_{c,1}$ and $c_{Na^+,2} = c_{Na^+,0}$ where the subscript “2” indicates the cathode out.

The total ionic current through the CEM is given by

$$J = J_{Na^+} + J_{H^+}$$

where $J_{H^+}$ is the molar flux of H$^+$ through the membrane. Both cation fluxes can be calculated by considering the Nernst–Planck equation. The electroneutrality condition inside the CEM gives

$$c_{Na^+} + c_{H^+} = c_X$$

where $c_X$ is the fixed charge density of the cation exchange membrane. Therefore, the Nernst–Planck equation for $J_{Na^+}$ and $J_{H^+}$ can be rewritten as

$$J_{Na^+} = -D_c X(c_{Na^+}) \left( \frac{df_{Na^+}}{dx} + f_{Na^+} \frac{df}{dx} \right)$$

$$J_{H^+} = -a D_c X(c_{H^+}) \left( \frac{df_{H^+}}{dx} + f_{H^+} \frac{df}{dx} \right)$$

where $D_c X$ is the diffusion coefficient of Na$^+$ inside the membrane, $a$ is the ratio between the diffusion coefficient of H$^+$ and Na$^+$, $f$ the electric potential, $x$ the position inside the membrane, and $f_{Na^+}$ and $f_{H^+}$ are the concentration fractions of Na$^+$ and H$^+$, respectively ($f_{Na^+} = c_{Na^+/c_{Na}}$). The ion concentration in the bulk solution is the same as the ion concentration on the membrane surface since we assumed perfectly stirred compartments (i.e., no concentration polarization phenomena). By combining eqs 7 and 8 into eq 5a, the total ionic current is equal to

$$J = \frac{-D_c X}{\delta} \left( (a-1)(f_{H^+,c} - f_{H^+,a}) + \frac{1 + (a-1)}{2} \frac{f_{H^+,c} + f_{H^+,a}}{2} df \right)$$

where $\delta$ is the CEM thickness and the subscripts $a$ and $c$ indicate the acidifying and cathode sides of the membrane, respectively. Furthermore, the transport number of Na$^+$ ($t_{Na^+}$) is defined as

$$t_{Na^+} = \frac{J_{Na^+}}{J}$$

Finally, the specific energy consumption (EC$^0$) is calculated by

$$EC^0 = \frac{V_{total}\frac{J}{J_{CO_2}}}$$

where $V_{total}$ is the total cell voltage, which is calculated as the sum of ohmic losses ($V_{ohmic}$), potential drop due to the pH difference in the cell ($V_{pH}$), and membrane potential ($V_{membrane}$).

$$V_{total} = V_{ohmic} + V_{pH} + V_{membrane}$$

$V_{ohmic}$ is the potential loss related to the ohmic resistance of the acidifying and cathode compartments and can be expressed as

$$V_{ohmic} = l \left( \frac{d_a}{\sigma_a} + \frac{d_c}{\sigma_c} \right)$$

where $d_a$ and $d_c$ are the thicknesses of acidifying and cathode compartments, respectively, $\sigma_a$ and $\sigma_c$ are the conductivities of acidifying solution and catholyte (calculated by OLISynt version 9.5 (OLI Systems, Inc., NJ) based on the solution composition, respectively.

$V_{pH}$ is determined by the pH difference between anode and cathode considering a potential drop of 59.2 mV per pH unit, and $pH_{nada} = 1$ due to the high proton concentration in the MEA.
shown in eqs 2, the accumulation of H2CO3 and CO3$^-$ is observed. Based on the CO2 equilibria 0.4, (ii) acidifying solution is higher than 7.3 (Figure 2b). Concomitantly, the CO2 partial pressure in the gas phase of the membrane (i.e., the change caused by H+ produced at the anode is neutralized by the OH$^-$ produced at the cathode).

In the second regime at $L_{Na^+}$ between 0.4 and 1.0, the carbon removal efficiency and CO2 production rate increase with $L_{Na^+}$ due to the lower pH in the acidifying compartment, showing that the carbon species in the spent solution are removed as desorbed CO2. Besides, the conductivity of the acidifying solution keeps decreasing as a higher amount of Na+ is transported to the cathode and CO2/HCO3$^-$ are neutralized by H+. Due to the decrease of ionic conductivity of the acidifying solution, the ohmic resistance of the cell increases with the load ratio. As a result of the trade-off between the increasing CO2 production rate and the increasing ohmic resistance, we experimentally reached a minimum specific energy consumption of 520 kJ mol$^{-1}$ CO2 at $L_{Na^+}$ = 0.8 and $j_c$ = 150 A m$^{-2}$.

In the third regime at $L_{Na^+}$ > 1.0, the carbon removal efficiency reaches a plateau, and the CO2 production rate starts to decrease after it reaches a maximum at $L_{Na^+}$ = 1.0. By definition, $L_{Na^+}$ > 1.0 implies that the applied current is sufficient to transport all of the Na+ from the acidifying compartment to the cathode compartment. As a result, the pH of the acidifying solution is so low that most of the carbon species in the aqueous solution are in the form of H2CO3$^*$. Therefore, the carbon removal efficiency is mostly limited by the CO2 partial pressure in the gas phase of the membrane contactor, which is constant at 1 atm. Moreover, the CO2 production is limited by the amount of total carbon feeding into the system and not by the pH of the acidifying solution. The CO2 production rate decreases at $L_{Na^+}$ > 1.0, since less influent and hence less total carbon is supplied than those at $L_{Na^+}$ < 1.0. Furthermore, the transport number of Na+ decreases at $L_{Na^+}$ > 1.0 (Figure S1), and some H+ ions in the acidifying solution are transported through the CEM rather than contributing to the CO2 desorption, as shown by the decreasing CO2 production rate in the last regime. Finally, the conductivity in the acidifying compartment is as low as 0.43 mS cm$^{-1}$ ($L_{Na^+}$ = 1.0) and 0.08 mS cm$^{-1}$ ($L_{Na^+}$ = 1.2). As a

$$V_{pH} = 0.0592 \cdot (pH_{cathode} - 1)$$

(19)

$$V_{membrane} = \frac{R \cdot T}{F} \ln \frac{c_{Na^+\ 2}}{c_{Na^+\ 1}}$$

(20)

The model requires as input the properties of the influent (i.e., $c_{Na^+\ 0}$, $c_{CO_3}$ and $Q$) and the Na+ concentration of the outlet acidifying solution ($c_{Na^+\ 1}$), and then calculates the current density, Na+ load ratio, CO2 production rate, and outlet concentration as output.

3. RESULTS AND DISCUSSION

3.1. Effect of Na+ Load Ratio. We first tested the cell under constant current operations and investigated the effect of the Na+ load ratio ($L_{Na^+}$) by varying the influent flow rate. Figure 2 shows both the experimental and theoretical results obtained with $L_{Na^+}$ ranging from 0.2 to 1.2 at a constant current density (150 A m$^{-2}$).

The carbon removal efficiency as a function of load ratio clearly shows three different regimes (Figure 2a): (i) $L_{Na^+}$ < 0.4; (ii) $L_{Na^+}$ = 0.4 – 1, and (iii) $L_{Na^+}$ > 1. For $L_{Na^+}$ < 0.4, no carbon removal is observed. Based on the CO2 equilibria shown in eqs 2–4, we calculated that CO2 can only be desorbed into the gas phase when the pH is lower than 7.3 (i.e., assuming that the acidifying solution is in equilibrium with 1 atm pure CO2 in the gas phase of the membrane contactor). In fact, in this regime at $L_{Na^+}$ < 0.4, the pH of the acidifying solution is higher than 7.3 (Figure 2b). Consequently, the CO2 production rate is zero (Figure 2c). The invested energy is, however, essential for achieving the suitable pH conditions to desorb CO2 at higher $L_{Na^+}$. The reason is that the H+ produced in this first regime is used to convert CO2$^2-$ to HCO3$^-$ and a small amount of H2CO3$^*$ so that further accumulation of H2CO3$^*$ can lead to CO2 desorption. Finally, Figure 2d shows a linear decrease of the electrical conductivity of the acidifying solution, since Na+ is transported to the cathode and CO3$^2-$/HCO3$^-$ are neutralized by H+. Meanwhile, the electrical conductivity of catholyte is constant correspond-

Figure 2. Effect of Na+ load ratio on the cell performance at a constant current density (150 A m$^{-2}$). (a) Carbon removal efficiency, (b) pH, (c) specific energy consumption and CO2 production rate, and (d) electrical conductivity. Symbols (●▲): experimental data; lines: model results.
The carbon removal efficiency shown in Figure 2a also indicates the regeneration efficiency of the CO₂ capture capacity of the absorbent. At \( L_{\text{Na}^+} = 0.8 \), the removal efficiency is 57% implies that 57% of the CO₂ capture capacity of the absorbent is regenerated. The outlet catholyte reaches a pH of 13.19 and a conductivity of 62.58 mS cm⁻¹, which leads to a final composition of 0.22 M NaOH and 0.14 M Na₂CO₃ (calculated by OLISstudio based on measured conductivity). Given this, the OH⁻/CO₃²⁻ molar ratio, we believe that such a solution can be directly reused as an absorbent for DAC. In comparison, Keith et al. used an aqueous solution of 1.0 M KOH and 0.5 M K₂CO₃ as an absorbent in their DAC pilot plant that has a very close OH⁻/CO₃²⁻ molar ratio to our regenerated absorbent.16

Furthermore, Figure 2 also shows a good agreement between the model simulations and experimental data. In particular, the model predicts the same trend for carbon removal efficiency, pH, and conductivity (Figure 2a,b,d), thus proving the validity of the mass balance calculations in the system. Regarding the CO₂ production rate (Figure 2c), the model overestimates by 12–20% the experimental results, likely due to some CO₂ leakage occurring in the experimental setup. The residence time of the solution in the system increases at a higher load ratio due to the lower influent flow rate, so the risk of gas leakage also increases. Despite the overestimation by the model, the simulation results correctly predict a maximum for CO₂ production rate at \( L_{\text{Na}^+} = 1.0 \), where the limiting factor for CO₂ desorption shifts from the solution pH to the amount of carbon available in the bulk solution.

Regarding the energy consumption, the model correctly predicts the experimental trend, but it gives lower values compared to experimental data (Figure 2c). Such difference can be due to some of the simplifying assumptions, i.e., the absence of gas leakage (CO₂ and H₂) and of electrode overpotentials. In particular, the model predicts a minimum energy consumption of 195 kJ·mol⁻¹ CO₂ and a carbon removal efficiency of 59% at \( L_{\text{Na}^+} = 0.8 \). We believe that these values predicted by the model can be experimentally achieved by further optimization of the system (i.e., minimizing the gas leakage and electrode overpotentials).

### 3.2. Effect of Current Density at a Constant Load Ratio

The tests at different Na⁺ load ratios showed that the lowest energy consumption for the system is achieved at \( L_{\text{Na}^+} = 0.8 \). To further study the effect of current density on the energy consumption, we performed experiments with current density ranging from 25 to 150 A·m⁻² at a fixed load ratio. During the tests, the influent flow rate was modified in line with the current density to maintain a constant \( L_{\text{Na}^+} \) of 0.8. Experimental and theoretical results are shown in Figure 3.

As the model has been validated in the previous section for prediction of carbon removal efficiency, pH, and conductivity, we believe the simulations also predict the behavior of these parameters at different current densities correctly. Figure 3a shows that the carbon removal efficiency remains constant with the change of current density due to an identical steady state obtained at the same value of \( L_{\text{Na}^+} \). This identical steady state is also demonstrated by the constant pH and conductivity (Figure 3b,d). The CO₂ production rate increases linearly with the current density due to the higher influx of H⁺ from the anode and higher total carbon loading, as a result of the increasing influent flow rate (Figure 3c).

Nevertheless, we also noted differences between the simulation and experimental results. In particular, the difference in carbon removal efficiency, conductivity of catholyte, and CO₂ production rate can be explained by the leakage of carbon species from the system and inside the system. The tests at lower current density have a lower influent flow rate, which leads to longer residence time and larger gas leakage. Furthermore, the neutral carbon species (H₂CO₃*), instead of being desorbed as CO₂(g), can be transported from the acidifying compartment to the cathode compartment under a concentration gradient.34 Once in the cathode compartment, the H₂CO₃* can react with OH⁻ in the catholyte to produce CO₃⁻, thus decreasing the ionic conductivity (since CO₃⁻ has a lower ion conductivity than OH⁻). Therefore, the longer residence time at low-current-density tests increases the leakage of carbon species.

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**Figure 3.** Effect of current density and flow rate at a constant Na⁺ load ratio. (a) pH, (b) CO₂ production rate, (c) specific energy consumption and CO₂ production rate, and (d) electrical conductivity. Symbols (●▲): experimental data; lines: model results.
Such leakage is also visible in the increased energy consumption in the test at $j_c = 25$ A·m$^{-2}$ (Figure 3c). With lower current density, we expected lower energy consumption due to minor energy losses from the internal resistance and overpotential, which is in agreement with the tests at 50–150 A·m$^{-2}$. However, we observed a higher energy consumption at $j_c = 25$ A·m$^{-2}$ compared to the test at $j_c = 50$ A·m$^{-2}$. As shown by the similar difference between simulation and experimental results of the CO$_2$ production rate, the leakage of CO$_2$ is at a constant rate. Thus, the leakage has a greater impact on the energy consumption at a lower current density since the fraction of CO$_2$ leaked from the system is larger. The modeling results identify the importance of minimizing the leakage and thus directly point to the need of improving the gas-tightness of the setup in future studies.

We achieved the lowest energy consumption of 374 kJ·mol$^{-1}$ CO$_2$ at a current density of 50 A·m$^{-2}$. In that case, the outlet solution from the cathode compartment reached pH 13.07 and the final composition of 0.167 M NaOH and 0.1665 M Na$_2$CO$_3$, corresponding to a 49% carbon removal efficiency. In addition, the gas desorbed from the membrane contactor contains more than 95% of CO$_2$ in the test at $j_c = 50$ A·m$^{-2}$. This high-purity CO$_2$ gas is suitable for subsequent storage or utilization.\textsuperscript{35} Without the leakage of CO$_2$, as predicted by the model, the specific energy consumption increases gradually with the current density (Figure 3c). At $j_c = 25$ A·m$^{-2}$, a specific energy consumption of 164 kJ·mol$^{-1}$ CO$_2$ is predicted, which represents the lowest value of energy consumption for our system in the investigated operational range. Such an operating condition results in a 58% carbon removal efficiency and a final solution composition suitable for use as a DAC absorbent.

The energy consumption experimentally measured is higher than the theoretical values (Figure 3d). To investigate this difference, we analyzed the potential losses in the cell at different current densities. Figure 4 shows the contribution of the potential drops to the cell voltage during the tests shown in Figure 3. At a low current density, the cell voltage is clearly dominated by the equilibrium potential ($V_{eq}$) (75% of $V_{cell}$ at 25 A·m$^{-2}$), while its contribution to the cell voltage decreases at a higher current density (38% of $V_{cell}$ at 150 A·m$^{-2}$). The value of $V_{eq}$ is independent of the applied current density due to the constant pH difference between anode and cathode (at constant Na$^+$ load ratio). However, all other potential losses increase by increasing the current density. The overpotential of the anode ($\eta_{anode}$) and the cathode ($\eta_{cathode}$) reach their highest contribution at $j_c = 150$ A·m$^{-2}$, 28 and 13% of the cell voltage, respectively. The Donnan potential given by the CEM of the MEA was included as part of the anode overpotential; thus, the current density has a larger effect on $\eta_{anode}$. Moreover, the transport losses ($V_{transport}$) increase with current density until reaching 20% of the cell voltage at 150 A·m$^{-2}$. In this context, $V_{transport}$ refers to the remaining losses in the system, such as the losses caused by the formation of gas bubbles both in the acidifying compartment (CO$_2$) and in the cathode (H$_2$). Finally, the ionic losses due to the ohmic resistance of the solution ($V_{ionic}$) are negligible, due to the high ionic conductivity at $L_{Na^+} = 0.8$.

The model captures the equilibrium potential, ionic losses, and membrane potential (as a part of the transport losses). This part of potential losses remains stable with the increase of current density, while the remaining losses increase dramatically. Therefore, although the model predicted 80% of the total potential drop at 25 A·m$^{-2}$, only 48% is predicted when the current density rises to 150 A·m$^{-2}$. The difference is mainly due to the overpotential of the electrodes and other nonideal losses, which could be contributed by reaction kinetics, boundary layers, and bubble formation.\textsuperscript{22,25} The deviation between the modeling results and experimental data indicates that further refinement of the model and measurements is needed to identify the contribution of each of these processes and propose improved designs. On a cell design level, future research efforts should be focused on reducing gas leakage. Moreover, different membrane electrode materials could be tested to reduce the electrode overpotential, for instance, using different membrane electrode assemblies or highly electroactive materials for hydrogen evolution. Finally, further studies should be focused on optimizing the electrolyte compositions, aiming to decrease the transport losses in the system and provide more insights into the reaction kinetics.

The proposed H$_2$-recycling electrochemical system shows the ability to regenerate the spent alkaline absorbent from DAC and produce a high-purity CO$_2$ gas stream. We experimentally achieved a minimum energy consumption of 374 kJ·mol$^{-1}$ CO$_2$, while the model simulation has shown energy consumption of the system as low as 164 kJ·mol$^{-1}$ CO$_2$ by further optimization. Such low-energy-consumption values are in line with other regeneration strategies proposed for DAC, thus showing that this electrochemical process can be a competitive alternative among all DAC technologies. For instance, Keith et al. showed an energy consumption of 338 kJ·mol$^{-1}$ CO$_2$ for their thermodynamic cycle,\textsuperscript{16} while Sabatino et al. estimated an energy consumption of 236 kJ·mol$^{-1}$ CO$_2$ for bipolar membrane electrodialysis.\textsuperscript{28} Besides, up to 59% of the CO$_2$ capture capacity of the absorbent could be regenerated during the process, which made the effluent from the system suitable for DAC systems. Furthermore, another advantage of such an electrochemical system for CO$_2$ desorption is that it does not require any heat for the absorbent regeneration (unlike most of the proposed DAC technologies\textsuperscript{16,28}). Instead, it requires only electricity as energy input, so it can be easily coupled to renewable energy sources for a sustainable DAC process. Finally, the desorption of CO$_2$ and regeneration of alkaline absorbent occurring in one step represents a clear advantage for the reduction of the overall operational costs, since it simplifies the process operation compared to energy-intensive processes such as causticization.\textsuperscript{18}
ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.est.0c01977.

List of the performed experiments; calculations of potential drops; definition and results of transport number of Na⁺ and some additional indicators (Figure S1 and Tables S1 and S2) (PDF)

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Notes
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