Long-lasting, monovalent-selective capacitive deionization electrodes

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Emerging water purification applications often require tunable and ion-selective technologies. For example, when treating water for direct use in irrigation, often monovalent Na$^+$ must be removed preferentially over divalent minerals, such as Ca$^{2+}$, to reduce both ionic conductivity and sodium adsorption ratio (SAR). Conventional membrane-based water treatment technologies are either largely non-selective or not dynamically tunable. Capacitive deionization (CDI) is an emerging membraneless technology that employs inexpensive and widely available activated carbon electrodes as the active element. We here show that a CDI cell leveraging sulfonated cathodes can deliver long-lasting, tunable monovalent ion selectivity. For feedwaters containing Na$^+$ and Ca$^{2+}$, our cell achieves a Na$^+$/Ca$^{2+}$ separation factor of up to 1.6. To demonstrate the cell longevity, we show that monovalent selectivity is retained over 1000 charge–discharge cycles, the highest cycle life achieved for a membraneless CDI cell with porous carbon electrodes to our knowledge, while requiring an energy consumption of ~0.38 kWh/m$^3$ of treated water. Furthermore, we show substantial and simultaneous reductions of ionic conductivity and SAR, such as from 1.75 to 0.69 mS/cm and 19.8 to 13.3, respectively, demonstrating the potential of such a system towards single-step water treatment of brackish and wastewaters for direct use in irrigation.

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INTRODUCTION

The problem of hazardous compounds present at low concentrations in water sources spans domains from public health to heavy industry. Toxic heavy metals such as lead, cadmium, and arsenic enter water sources due to both anthropogenic activity and natural processes, while boron, present in seawater and ground-water, can have detrimental health effects when present at sufficiently large concentrations. Selective ion removal is an emerging water treatment approach for removing contaminants from water supplies while retaining desirable dissolved species, and it is a relevant technique for the environmental pollutants listed above as well as for recovery of high-value materials, such as Li$^+$ for use in lithium-ion batteries.

A promising application of selective removal is the treatment of water for agricultural irrigation, which accounts for 80% of total water consumption in the United States. Two important factors in determining irrigation water quality are conductivity and relative Na$^+$ content, the latter of which is quantified via the sodium adsorption ratio (SAR). Crop yields and health usually suffer from excessive conductivity and high SAR, the latter of which also negatively impacts soil permeability and water infiltration rate. Conductivity and SAR tolerances are known to vary between crops. For example, a 6-year study conducted by Singh et al. found that pearl-millet tolerated water with SAR < 10 while wheat tolerated SAR < 20. Furthermore, the same study found that long-term use of high-conductivity and high-SAR waters lead to increased salt accumulation and clay dispersion, which necessitate occasional treatment to restore soil quality. A system performing selective removal can reduce both conductivity and SAR in incoming water by selectively removing Na$^+$ ions while permitting important minerals to remain in the treated water. This can increase crop yields and can reduce costs incurred either by restorative soil treatments or by use of non-selective water treatment necessitating water remineralization.

Conventional membrane-based technologies have limited capability to remove specific ions, and when they are selective, they tend to reject divalent ions, thus potentially increasing the SAR of the feedwater. Reverse osmosis (RO) typically removes >95% of all dissolved ions indiscriminately when used for seawater desalination and requires additional post-treatments, including pH adjustment, to remove boron and remineralization to reintroduce divalent ions. Electrodialysis membranes are typically divalent selective. The emerging use of a thin polyelectrolyte layer with a charge opposite to that of the membrane bulk can inhibit divalent counterion passage and thus enable monovalent selectivity, typically at the cost of lifetime and membrane resistivity. Nanofiltration is perhaps the most promising membrane technology for selective separation, where addition of either a positively or negatively charged layer to the membrane feed side can enhance selectivity of either monovalent or divalent ions. However, membrane charge density is not dynamically tunable, and thus adjusting such a system to varying feedwater composition or changes in effluent requirements is challenging. Further, membrane fouling, concentration polarization, and often-high membrane costs are significant issues in membrane separation methods. The need for tunable, highly selective technologies relying on inexpensive active elements has spurred exploration into next-generation water treatment processes and materials.

CDI is a water treatment technology that does not require membranes and is promising for selective ion removal. Typical CDI operation involves applying a constant voltage or current between two electrodes, which are often inexpensive and widely available activated carbons, in the presence of a feed...
stream (Fig. 1a). The electric field applied to the system causes the ions in the feed stream to electromigrate into the electrodes, where they are stored electrostatically in electric double layers (EDLs) formed in electrode nanopores (Fig. 1b). Once fully charged, the electrodes are regenerated electrically, by either short-circuiting or reversing the applied current, and the discharged ions are released from the EDLs to form a concentrated brine waste stream. CDI has been identified as a promising emerging water treatment technology for brackish water desalination and ion-selective separations, but a major cause of electrode degradation is short electrode lifetime. A major cause of electrode degradation is Faradaic side reactions which oxidize the anode and reduce dissolved oxygen at the cathode. In addition to activated carbons, alternate electrode materials that rely on intercalation processes and redox conversion reactions to deionize water are currently under investigation for selective ion removal. Use of stable sulfonic (SO₃⁻) groups and charging to short times minimizes Faradaic side reactions and thus electrode degradation.

The treated water has significantly reduced sodium absorption ratio (SAR) and ionic conductivity, rendering it suitable for direct use in irrigation. A major cause of equipment degradation in CDI, but are gradually exchanged with divalent ions in the electrode nanopores as the system approaches equilibrium. Seo et al. observed the dynamic replacement of Na⁺ by Mg²⁺ and Ca²⁺ during a CDI cycle. Zhao et al. found that Na⁺ and Ca²⁺ were initially stored in proportion to their bulk concentrations, yet Ca²⁺ replaced Na⁺ at longer times, and both ions approached equilibrium values predicted by modified Donnan theory after 5 h of charging, whereas other dynamic models were utilized for selectivity analysis considering other EDL models. Furthermore, Choi et al. observed short cycle time and high charging voltage increase monovalent ion removal relative to divalent ion removal. Thus, connecting between equilibrium nanopore models and cell-level dynamic models is important to fully explore and synergize between parallel ion selectivity mechanisms in CDI. Electrode morphology and surface chemistry strongly influence selectivity in carbon electrodes. Techniques to enhance selectivity broadly follow two main approaches, physical tuning of nanopore size and chemical surface functionalization. Regarding the former approach, sufficiently narrow pores allow small ions to enter while larger ions are physically excluded, an effect known as ion sieving. Regarding the latter approach, chemically deposited surface functional groups have been shown to improve selective removal in CDI by enhancing size-based exclusion effects, and by binding preferentially to specific ions like NO₃⁻ and Pb²⁺, and Pb²⁺. Furthermore, functional groups have been shown to increase salt adsorption capacity, cell charge efficiency, and long-term cycling stability. Surface functional groups behave as either acids or bases and exhibit either weak or strong dissociation behavior. While weak acid groups have been shown to enhance selectivity, their performance can be significantly affected by variations in incoming feed pH. Group stability is also critical, as
Guyes et al.\textsuperscript{43} showed that cathode functionalization with weak acid carboxylic groups improved selectivity of K\textsuperscript{+} over Li\textsuperscript{+}, though significant loss of chemical surface charge was observed after only a few charge–discharge cycles. Functionalization with sulfonic groups can possibly remediate both issues, as their strong-acid cathodes, with electrode and separator thicknesses pristine electrodes (P\textsubscript{Coulombic efficiency})\textsuperscript{64–67}, and pore wettability\textsuperscript{66,68,69} in lab-scale tests on feedwaters containing solely NaCl. However, to our knowledge, the effect of sulfonic groups on ion-selective removal and its long-term stability upon CDI cell charge–discharge cycling has not yet been explored.

In this work, we demonstrate a monovalent-selective CDI cell that can simultaneously reduce feed conductivity and SAR (Fig. 1c), and is stable over 1000 charge–discharge cycles with a Coulombic efficiency greater than 96%. We first employ a combined dynamic-mD model and predict that functionalized electrodes enhance monovalent selectivity at short charging times relative to pristine electrodes. We validate this prediction experimentally using a cell with a sulfonated cathode, demonstrating enhanced separation factor and reductions of both SAR and feedwater conductivity compared to a cell with a pristine cathode. Moreover, we show that short charging times enable charging the cell at extended voltage windows of up to 2.5 V, well beyond those where electro-corrosion of the anode normally occurs. We show that such high-voltage operation leads to further reductions in conductivity and SAR, while maintaining a Coulombic efficiency greater than 90%, indicating a low rate of Faradaic side reactions, though separation factor is not improved. We compare flush and no-flush CDI operation in the extended voltage range and show that despite gains in productivity and charge efficiency achieved with flushing, desalinated water SAR, Ca\textsuperscript{2+} selectivity, and conductivity increase relative to no-flush operation, indicating that possible inclusion of a flush step for a selectivity process must be carefully evaluated.

Finally, we cycled a cell with a sulfonated cathode 1000 times with a charging voltage of 1.2 V, achieving excellent electrochemical stability and reduced effluent SAR with increasing cycle number. Overall, we demonstrate that sulfonated electrodes can serve as the foundation for long-lasting, monovalent selective, membraneless CDI systems, which potentially can be utilized to improve irrigation water quality.

**RESULTS AND DISCUSSION**

**Simulations**

Zhao et al.\textsuperscript{40} described a time-dependent selectivity between Na\textsuperscript{+} and Ca\textsuperscript{2+}, where at short charging times, Na\textsuperscript{+} is adsorbed selectively due to its larger concentration in the feed, but at longer times approaching equilibrium, Ca\textsuperscript{2+} is preferred due to the stronger electric force acting on the higher-valence ion. However, the latter study did not develop a dynamic model for a full CDI cell, incorporate the effects of ion size, or study the impact of chemically charged electrodes on selectivity. Building on our previous work in dynamic CDI modeling\textsuperscript{43,70}, here we develop a model which addresses the latter elements by capturing the dynamics of a complete CDI cell with an electrolyte containing competing smaller-sized Na\textsuperscript{+} and larger Ca\textsuperscript{2+}, and includes a nanopore chemical charge concentration, \(\sigma\)\textsubscript{chem}, such as that created by strong-acid (e.g. sulfonic) groups. We consider two electrode types, a pristine electrode in which \(\sigma\)\textsubscript{chem} = 0, and a chemically functionalized electrode in which \(\sigma\)\textsubscript{chem} is constant and equal to \(-2\mathrm{M}\), such as from strong-acid sulfonic groups. We simulate two CDI cell configurations under an applied charging voltage of \(V\textsubscript{ch} = 1.0\mathrm{~V}\), the first comprised of a pristine anode and pristine cathode (P–P cell), and the second consisting of a pristine anode and functionalized cathode (F–P cell). Both electrodes have thickness \(l\) and are electrically isolated by a separator with thickness \(l\textsubscript{sp}\) while the cathode is placed upstream.

To gain insight into ion adsorption dynamics, in Fig. 2b we plot ion adsorption per unit electrode mass, \(\Gamma\) (defined in “Simulations” in “Methods”, Eq. (6)), for the P–P cell and F–P cell, scaled by \(\Gamma\textsubscript{eq,i}\) the adsorption of the P–P cell at equilibrium, where the subscript \(i\) denotes either Na\textsuperscript{+} or Ca\textsuperscript{2+}. On the horizontal axis is time scaled by the cell diffusion timescale, \(t \equiv D\textsubscript{Na} l^2 / l\textsubscript{sp}^2\), where \(t\) is time and \(D\textsubscript{Na}\) is the Na\textsuperscript{+} ion diffusivity at infinite dilution. Our dynamic model displays an expected time-dependent selectivity as \(\Gamma\textsubscript{Na} / \Gamma\textsubscript{Ca} = 1\textsuperscript{0.5}\textsuperscript{1}\) for the P–P cell (solid lines) reaches a maximum of 2.25 at \(t = 0.87\) and then decreases. Meanwhile, \(\Gamma\textsubscript{Na} / \Gamma\textsubscript{Ca} = 1\textsuperscript{0.5}\textsuperscript{1}\) for the F–P cell (dotted lines) reaches a slightly lower maximum of 2.20 at \(t = 0.90\). Turning our attention to Ca\textsuperscript{2+}, \(\Gamma\textsubscript{Ca} / \Gamma\textsubscript{P}\) of both the P–P and F–P cells is negative for very early times, indicating net Ca\textsuperscript{2+} discharge.

![CDI simulation design and predictions](image-url)

**Fig. 2** CDI simulation design and predictions. **a** Schematic of the CDI system used in simulations with either pristine (P) or functionalized (F) cathodes, with electrode and separator thicknesses \(l\) and \(l\textsubscript{sp}\) respectively. Feedwater with ion concentrations \(c\textsubscript{feed}\) enters the cell at \(x = 0\) while charging at \(V\textsubscript{ch}\), and the resulting desalinated effluent is collected for various scaled charging times, \(t\), to quantify Na\textsuperscript{+} and Ca\textsuperscript{2+} concentration, \(c\textsubscript{eff}\) and separation factor, \(\beta\textsuperscript{eq}\). **b** Predictions of scaled Na\textsuperscript{+} ion adsorption, \(\Gamma\textsubscript{Na}/\Gamma\textsubscript{P}\textsuperscript{eq}\) (left axis, green lines), Ca\textsuperscript{2+} ion adsorption, \(\Gamma\textsubscript{Ca}/\Gamma\textsubscript{P}\textsuperscript{eq}\) (left axis, red lines), and \(\Gamma\textsubscript{P}\) (right axis, blue lines), as functions of \(t\). Calculations are for a model cell with either two pristine electrodes (P–P cell, solid lines) or a cell with a chemically functionalized cathode (F–P cell, dotted lines).

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desorption, followed by monotonically increasing Ca$^{2+}$ adsorption, reaching values of 0.46 and 0.41 at $t = 2$, respectively. Figure 2b also shows the predicted separation factor $\beta_{\text{Ca}^{2+}}^{\text{Na}^+}$, which quantifies selective adsorption of Na$^+$ over Ca$^{2+}$ for the P–P cell and the F–P cell. For both cell designs, we see $\beta_{\text{Ca}^{2+}}^{\text{Na}^+}$ approaches a negative asymptote at very short charging times as Ca$^{2+}$ is not desorbed from the cell. This is followed by very high positive values of $\beta_{\text{Ca}^{2+}}^{\text{Na}^+}$ at early times of $t \ll 1$, indicating strong monovalent selectivity, which sharply decreases to near-zero values for $t \approx 1$. In the P–P cell, $\beta_{\text{Ca}^{2+}}^{\text{Na}^+}$ drops below unity much earlier, at $t = 0.053$, as compared to the F–P cell which remains monovalent-selective until $t = 0.52$. Thus, our model predicts that the F–P cell can deliver improved monovalent selectivity for longer charging times relative to the P–P cell, and that charging for longer times enables greater ion storage. Thus, we conclude that functionalizing the cathode is predicted to enhance Na$^+$ removal from the feedwater.

**Electrode characterization**

Given the prediction from our dynamic theory that functionalizing the cathode with a strong-acid group enables improved monovalent selectivity (Fig. 2), we prepared a sulfonated activated carbon electrode (see “Methods”). Transmission Fourier Transform Infrared (FTIR) spectra of the pristine and sulfonated electrode materials are shown in Fig. 3a. The sulfonated electrode spectrum displays distinctive peaks at 1162 and 1157 cm$^{-1}$, corresponding to the R$\text{SO}_3^-$ group, and multiple peaks in the band 1315–1220 cm$^{-1}$, corresponding to the RO$\text{SO}_4^-$ group, thus confirming the presence of sulfonic functional groups on the electrode surface. We further quantified the concentration of nanopore chemical charge, $\sigma_{\text{chem}}$, by fitting experimental results of direct titrations of the carbon material (Fig. 3b) immersed in a strong basic solution (see “Methods”). Results show the pristine material is mildly charged with a significant pH dependence of the measured chemical charge, indicating the presence of weak acidic and basic functional groups. By contrast, the sulfonated electrode is strongly negatively charged, presenting $\sigma_{\text{chem}}$ values between approximately $-1.25$ and $-2.5$ mC for the tested pH range, indicating the existence of fixed negative charges due to the presence of strongly acidic functional groups. Thus, sulfonated electrodes possess high negative nanopore chemical charge concentration regardless of local cell pH environment, a required condition for enhanced monovalent selectivity, as identified by the model shown in Fig. 2b.

**CDI experiments**

Figure 4a schematically shows the differences between pristine and sulfonated cathode nanopores. The pristine nanopore contains a small amount of native chemical charge, while the sulfonated nanopore contains a high concentration of negatively charged sulfonic groups (Fig. 3b). To verify the theoretical predictions of improved monovalent selectivity via functionalized cathodes (Fig. 2b), we built and characterized a CDI cell with either a pristine cathode and a pristine anode (P–P cell) or a sulfonated cathode and a pristine anode (S–P cell). All experiments involved equally long charging and discharging steps and a feed solution consisting of 14 mM NaCl and 0.5 mM CaCl$_2$. Results presented are from the limit cycle, which is the cycle at which the conductivity profile reaches a dynamic steady state. Figure 4b depicts the measured Na$^+$ separation factor, $\beta_{\text{Na}^+}^{\text{Na}^+}$, calculated via Eq. (1), vs. charging voltage, $V_{\text{ch}}$, for a discharge voltage of 0 V and for full-cycle times (FCTs) of 6, 15, and 30 min (which have charging steps corresponding to $t$ of 0.67, 1.67, and 3.33, respectively). The results at 15 and 30 min FCT are very similar for P–P and S–P cells, with a strong Ca$^{2+}$ preference for both charging times. For example, at 30 min FCT and 0.4 V, $\beta_{\text{Na}^+}^{\text{Na}^+} = 0.1$ in the P–P cell, and at 1.2 V it rises to 0.49. However, the two cells exhibit significantly different trends at 6 min FCT. The P–P cell separation factor ranges between 0.49 at 0.4 V and 0.85 at 1.2 V, whereas in the S–P cell the increase is much sharper, from 0.25 at 0.4 V to 0.96 at 1.2 V. This latter result indicates that the sulfonated cathode enables stronger divalent selectivity at low $V_{\text{ch}}$, yet stronger monovalent selectivity at high $V_{\text{ch}}$, relative to a pristine cathode (see Supplementary Fig. 1 for measured Na$^+$ and Ca$^{2+}$ adsorption). These findings are consistent with the theoretical trends of Fig. 2b, which predicts similar selectivity between the two cells for $t > 1$ but enhanced monovalent selectivity for the cell with a functionalized cathode when $t < 1$.

Separation factor, defined by Eq. (1), is a fundamental metric which can be compared across different water treatment technologies. However, separation factor alone is not sufficient to evaluate a technology’s suitability for treating water for irrigation applications. Additionally, the application-specific metrics of ionic conductivity, $\kappa$, and SAR are also important due to their importance to soil quality and crop health. The latter is defined as:

$$\text{SAR} \equiv \frac{c_{\text{Na}^+}}{\sqrt{c_{\text{Ca}^{2+}} + c_{\text{Mg}^{2+}}}},$$

where the ion concentrations $c_{\text{Na}^+}$, $c_{\text{Ca}^{2+}}$, and $c_{\text{Mg}^{2+}}$ have units of mM. In this work, $c_{\text{Mg}^{2+}} = 0$ and so SAR $= c_{\text{Na}^+}/\sqrt{c_{\text{Ca}^{2+}}}$.

The effects of $V_{\text{ch}}$, FCT, and cell type on product water ionic conductivity and SAR are shown in Fig. 4c–e, with dashed horizontal and vertical lines representing the feed conditions ($V_{\text{ch}} = 1.75$ mV/cm, SAR$_{\text{feed}} = 19.8$). At the longest FCT tested of 30 min, Fig. 4c shows that for both the P–P and S–P cells, increasing $V_{\text{ch}}$ decreases effluent conductivity, but for every $V_{\text{ch}}$, SAR is increased above its feed value by up to several units. This is the expected result when approaching equilibrium, as the nanopore will select the ion with higher valence due to a higher electric force exerted on such ions (Fig. 2b). For an FCT of 15 min (Fig. 4d), both cells deliver a reduction in conductivity as compared to the feed, but the output SAR remains at or above the feed value at all tested conditions. By contrast, the FCT of 6 min is sufficiently short to selectively store Na$^+$ and reduce SAR below the feed value. Figure 4e shows monotonous reductions of SAR and conductivity with increasing $V_{\text{ch}}$ for the P–P cell, with output values of 17.3 and 1.22 mS/cm at 1.2 V, respectively. The S–P cell achieves improved performance with an output SAR of 16.0 and a conductivity of 1.12 mS/cm at 1.2 V. These findings are again consistent with the trends predicted in Fig. 2b and show experimentally that use of a strong-acid functionalized cathode allows for enhanced monovalent selectivity.

With the promising results obtained with short cycle times in Fig. 4e, we hypothesized that short charging times would also limit electrode degradation. Thus, we explored extending the charging voltage window to further reduce SAR and conductivity of the treated water. Figure 5a shows that increasing $V_{\text{ch}}$ to 1.5, 2, and 2.5 V with a 6 min FCT in the S–P cell achieved this goal, culminating at 2.5 V with $\beta_{\text{Na}^+}^{\text{Na}^+}$ of 0.89, SAR of 14, and conductivity of 0.81 mS/cm, a decrease of 6 SAR units and a 54% conductivity reduction from the feed solution. Furthermore, a second pass of product water from the 1.2 V and 6 min FCT experiment resulted in an even lower SAR of 13.3 and conductivity of 0.69 mS/cm, also shown in Fig. 5a, with $\beta_{\text{Na}^+}^{\text{Na}^+} = 0.91$. We also explored incorporating a flush step into the CDI cycle, where the flush step involved pumping feedwater through the cell for 1 min at 5 mL/min after each half-cycle while the cell was held at zero current. Such flush steps prevent resalination of the product water and desalination of the brine. Figure 5a shows that flushing increased SAR of the product water significantly, but still resulted in a net reduction.
from the feed SAR at 2.5 V. Further, the flush step increased product water conductivity relative to no-flush operation, as some feedwater was flushed into the product water. This served to raise productivity from ~46 L/h/m² without flushing to ~93 L/h/m² with flushing.

To evaluate further the strategy of extended voltage windowing and flushing, Fig. 5b shows the measured Coulombic and charge efficiencies from the experiments in Fig. 5a. Coulombic efficiency is an indicator of the charge magnitude lost to Faradaic side reactions, and is defined as $\eta_{\text{coul}} \equiv \frac{q_{\text{dl}}}{q_{\text{ch}}}$, where $q_{\text{dl}}$ is the electric charge transferred between the electrodes during the charge step and $q_{\text{ch}}$ is the charge stored in the electrodes and subsequently released during the discharge step. Coulombic efficiencies of approximately 100% imply that side reactions are nearly absent, while lower values reveal that significant charge is lost to side reactions, and that degradation of the electrodes likely occurs during cycling. Figure 5b shows that Coulombic efficiency differs by no more than 5% between the flush and no-flush modes for each $V_{\text{ch}}$. Furthermore, the Coulombic efficiency remains greater than 93% up to 2 V, which shows that extending the voltage window is a promising strategy for improved monovalent selectivity. At 2.5 V, Coulombic efficiency drops to 85% (flush) and 88% (no flush), indicating that this $V_{\text{ch}}$ may be less suitable for long-term CDI operation. Figure 5b also shows the flush step increases the charge efficiency by ~20–40% relative to the no-flush mode. Charge efficiency for this system is
The results of Figs 4 and 5 show that over many experimental conditions, our cell can simultaneously reduce SAR and conductivity of the feedwater, and that extending the voltage window can be an effective strategy. To further evaluate our results, Fig. 6a shows the energy consumption per unit product water, $E_p$, for the results shown in Figs 4 and 5 (excluding the second pass results). Energy consumption increases by over an order of magnitude, from ~0.01–0.1 kWh/m³ at separation factors of order 0.1, to ~0.3 kWh/m³ for separation factors of ~1. Higher energy consumptions of ~1 kWh/m³, obtained with applied voltages of 2 V or greater, did not further improve separation factor. In context of the performance evaluation of Hawks et al., we find typical energy usage of 0.2–0.4 kWh/m³ for Cl⁻ concentration reductions of 4–6 mM and a throughput of ~46 L/h/m² (Supplementary Table 1). For comparison, Cerón et al. reported 0.21–0.24 kWh/m³ for a concentration reduction of 4–6 mM using a 20 mM NaCl feed and 17–19 L/h/m² productivity. Thus, we conclude that our system consumes similar levels of energy but at higher throughput than Cerón et al.

To understand possible sources of energy loss, we calculate the resistive losses per unit product water, approximated via $E_{\text{res}} = (P^2) \cdot \text{FCT} \cdot R_L \cdot \sqrt{S}$, where $(P^2) = (1/\text{FCT}) \int_{\text{FCT}} P^2 dt$ is the time-averaged squared current over a full charge–discharge cycle and $R_L$ is the series resistance for one separator layer, the latter of which is determined from electrode impedance spectroscopy measurements (see "Methods"). The inset in Fig. 6a reveals that series resistive losses are a significant fraction of overall energy input, between 24 and 56% of the overall energy usage, and that the lowest fractions of resistive losses occur when achieving high separation factors in the S–P cell. In this work, no energy recovery was implemented on cell discharge. Energy recovery has been shown experimentally to capture up to 50% of the input energy during the discharge half-cycle, while the highest possible energy recovery fraction is in excess of 80%. The results shown in the inset of Fig. 6a show that such recovery will be an effective strategy, as a majority of the energy input for highly selective conditions was not lost to series resistances, but may instead be stored capacitively. Returning to the main plot of Fig. 6a, the S–P cell tends to acidify the output water to a greater degree than the P–P cell as evidenced by the color of each marker, particularly at separation factors greater than ~0.7. At a separation factor of 0.96, the output pH is reduced to 5.1 from a feed value of 7, even with a Coulombic efficiency of 97.5% (1.2 V, 6 min FCT, S–P cell). At higher $V_{ch}$, acidification is more severe, though flushing somewhat mitigates this (Supplementary Table 1).

Also of interest is the thermodynamic energy efficiency (TEE), a useful measure for quantifying the input energy lost to irreversibilities such as resistive losses and parasitic side reactions. For our CDI system, which separates feedwater into brine and product streams, TEE is calculated by dividing $E_c$ by the minimum thermodynamic desalination energy $E_{\text{min}}$, given by

$$E_{\text{min}} = RT \sum_{i} \left[ \frac{C_{f,i}}{V}\ln \left( \frac{C_{b,i}}{C_{f,i}} \right) - \frac{C_{p,i}}{V}\ln \left( \frac{C_{b,i}}{C_{p,i}} \right) \right].$$

Here $i$ includes Na⁺, Ca²⁺, and Cl⁻, and the subscripts F, B, and P represent feed, brine, and product streams, respectively. Water recovery is $WR = V_F/V_P$, where $V_F$ and $V_P$ are the volumes of the product and influent feed streams, respectively. Further details on the derivation and use of Eq. (3) are given in the Supplementary Discussion. Equation (3) has been applied by a number of authors to CDI operation, though to our knowledge there are no previous studies evaluating TEE for a process investigating monovalent ion selectivity in CDI. Figure 6b displays the TEE of the results in Fig. 6a, and we see that TEE is not correlated with separation factor, but is generally higher for the S–P cell. This can be explained by the fact that the S–P cell generally exhibits higher Coulombic and charge efficiencies (Supplementary Table 1). We emphasize that values reported in Fig. 6 for energy consumption and TEE are for zero energy recovery since discharge occurs at 0 V, and furthermore for a largely unoptimized cell. TEEs of up to ~9% have been reported for optimized CDI cells, and we expect that future cell optimization will significantly reduce the series resistances.

Returning to the results of Fig. 4e, the significant SAR and conductivity reductions at 1.2 V and 6 min FCT in the S–P cell are accompanied by a Coulombic efficiency of 97.5% (Supplementary Table 1). The relatively insignificant faradaic losses suggest that the electrodes are stable under long-term cycling at these conditions. To evaluate this, we performed a 1000-cycle experiment at these experimental conditions and evaluated water quality at the beginning (limit cycle), middle, and end of cycling. The results in Fig. 7a reveal that the Na⁺ separation factor, $p_{\text{Na}^+}$, increases from 0.83 to 1.6, indicating that the electrodes are unequivocally selective for Na⁺ over the majority of tested cycles. Coulombic efficiency remains between 96 and 98% from the limit cycle (cycle 11) until the end of the experiment. The inset in Fig. 7a surprisingly reveals that SAR of the produced water continually improves during cycling, from 17.7 at the limit cycle to 15.8 at the end of the experiment. This improvement is due to a drastic reduction in Ca²⁺ adsorption from 2.6 to 1.1 µmol/g, whereas Na⁺ adsorption only modestly decreases from 65 to 55 µmol/g.
(Supplementary Table 1). We further see that the effluent conductivity is quite stable and increases by only 8%, from 1.12 to 1.20 mS/cm, from the limit cycle to the final cycle. Furthermore, the effluent pH becomes more neutral, rising from 5.11 to 6.57 (Supplementary Table 1). We therefore conclude that the CDI system is not only electrochemically stable and monovalent selective at the given conditions, but the output water quality actually improves with repeated cycling due to a decrease in SAR and a more-neutral pH.

In Fig. 7b we compare our results with other CDI longevity studies that used capacitive carbon electrodes without membranes\textsuperscript{27–29,57,85–89}. Since the other studies presented utilize NaCl alone in the feedwater whereas we use NaCl and CaCl\textsubscript{2}, we employ chloride adsorption, \( \Gamma_{\text{Cl}} = \Gamma_{\text{Na}} + 2\Gamma_{\text{Ca}^2+} \), as the comparison metric. The vertical axis position indicates \( \Gamma_{\text{Cl}} \) at the beginning of the experiment, and marker color indicates the percentage of \( \Gamma_{\text{Cl}} \) retention at the final cycle (studies with \( \Gamma_{\text{Cl}} \) retention of 100% or greater are shown as white open markers\textsuperscript{27,28,86,87}). Our CDI system surpasses the next-longest cycling study by \( \sim \)400 cycles, and to our knowledge is the CDI cell filled with salt water containing Na\textsuperscript{+}, Ca\textsuperscript{2+}, and Cl\textsuperscript{−} (Fig. 2a). The mass balance equation for each of the dissolved ions is of the form

\[
\rho_{\text{Na}} \frac{\partial \Gamma_{\text{Na}}}{\partial t} + \rho_{\text{Cl}} \frac{\partial \Gamma_{\text{Cl}}}{\partial t} = -\frac{\partial}{\partial x} \left( \rho_{\text{Na}} \frac{\partial \Gamma_{\text{Na}}}{\partial x} + \rho_{\text{Cl}} \frac{\partial \Gamma_{\text{Cl}}}{\partial x} \right) - U \frac{\partial \Gamma_{\text{Na}}}{\partial x}.
\]

Here \( \rho_{\text{Na}} \) and \( \rho_{\text{Cl}} \) are the macro pore and nanopore porosities, respectively; \( \Gamma_{\text{Na}} \) and \( \Gamma_{\text{Cl}} \) are the ion concentrations in the macropores and nanopores, respectively; and the subscript \( \text{i} \) refers to the \( i \)-th ion. Also, \( J_i \) is the superficial ionic flux, given by the extended Nernst–Planck equation, \( U \) is the superficial velocity, and \( \Gamma_{\text{Na}} \) is the macropores’ tortuosity described by the Bruggeman equation, \( \Gamma_{\text{Na}} = \rho_{\text{Na}}^{1/3} \). Further, \( D_i \) and \( z_i \) are the molecular diffusion coefficient at infinite dilution and ionic valence, respectively, \( \phi_{\text{Na}} \) is the bulk electric potential scaled by the thermal potential \( V_t \equiv RT/\mathcal{F} \), where \( R \) is the ideal gas constant; and \( T \) is absolute temperature. We invoke electroneutrality in the bulk solution, \( \sum_j z_j \Gamma_j = 0 \). We find the bulk potential from the charge balance equation,

\[
\rho_{\text{Na}} \frac{\partial \phi_{\text{Na}}}{\partial t} + \rho_{\text{Cl}} \frac{\partial \phi_{\text{Cl}}}{\partial t} = -\frac{\partial}{\partial x} \left( \rho_{\text{Na}} \frac{\partial \phi_{\text{Na}}}{\partial x} + \rho_{\text{Cl}} \frac{\partial \phi_{\text{Cl}}}{\partial x} \right).
\]

Here \( J_i \equiv \sum_j z_j \Gamma_j \) is the ionic current, \( \phi_{\text{on}} \equiv \sum_j z_j \phi_{\text{Na}} \) is the ionic charge concentration in the nanopores, and the summation is performed over all dissolved ions Na\textsuperscript{+}, Ca\textsuperscript{2+}, and Cl\textsuperscript{−}.

Nanopore concentrations are calculated by applying the modified Donnann (mD) model\textsuperscript{10,26}, \( \phi_{\text{Na}} = \phi_{\text{Na}}^0 \exp \left( -z \Delta \phi_{\text{D}} - \Delta \phi_{\text{m}} \right) \), where \( \Delta \phi_{\text{D}} \) is the Donnann potential drop. Further, \( \Delta \phi_{\text{m}} \) is the excess potential, accounting here for the hydrated size of ions, utilizing the Boublik–Mansoori–Camahan–Starling–Leland equation of state\textsuperscript{23,26,90,91}. The Donnann potential drop is defined from \( \phi_{\text{Na}} - \phi_{\text{Na}}^0 = \Delta \phi_{\text{D}} = \Delta \phi_{\text{D}}^0 + \Delta \phi_{\text{D}}^D \), where \( \Delta \phi_{\text{D}}^D \) is the scaled electrode potential and \( \Delta \phi_{\text{D}}^0 \) is Stern potential drop given by \( \Delta \phi_{\text{D}}^0 = F \Delta \phi_{\text{D}}^0 / V_{\text{Na}} \). Here, \( C_s \) is the Stern layer capacitance and \( \phi_{\text{on}} \) is the nanopore electronic charge concentration found from nanopore charge balance, \( \phi_{\text{on}} + \phi_{\text{chem}} + \phi_{\text{elec}} = 0 \).

The boundary conditions employed to solve Eq. (4) are \( J_{i,x=0} = U_{\text{elec,0}} \) and \( J_{i} - L_{i} = U_{\text{elec,0}} \). The lateral boundary conditions are appropriate for the case of high Peclet numbers, where the reservoir is long compared to the cell (\( \Delta \text{z} / L \gg 10^{3} \)). The boundary conditions employed for Eq. (5) are zero current at the electrodes’ edges, \( J_{i,x=0} = J_{i,x=L_{i}} = 0 \). For Eq. (4), the initial conditions are \( \phi_{\text{Na}}(x, t = 0) = \phi_{\text{Na,0}} \), while for Eq. (5), the initial condition is constant potential, \( \phi_{\text{Na}}(x, t = 0) = \phi_{\text{D}} \). To calculate \( \phi_{\text{D}} \), we introduce the total charge balance of the system at equilibrium,

\[
J_{i,x \geq 0} \phi_{\text{Na}} dV_{A} + J_{i,x \leq 0} \phi_{\text{Cl}} dV_{C} = 0,
\]

where \( V_{A} \) and \( V_{C} \) are the anode and cathode volumes, respectively.
and the cathode volumes, respectively, and $\sigma^{\text{elec, A}}$ and $\sigma^{\text{elec, C}}$ are the nanopore electronic charge concentrations before charging, at the anode and the cathode, respectively. The concentrations $\sigma^{\text{elec, A}}$ and $\sigma^{\text{elec, C}}$ are functions of $\phi_{i}$ and therefore solving the charge balance equation results in finding $\phi_{i}$. The set of equations was solved by COMSOL Multiphysics using the parameters presented in Supplementary Table 2.

The adsorption per unit electrode mass of ion $i$, either Na$^+$, Ca$^{2+}$, or Cl$^-$, is defined as

$$
\Gamma_{i} = (\sigma_{\text{chem, i}} - \sigma_{\text{eff, i}}) / \theta \cdot m,
$$

where $\theta$ is the collected effluent water volume, $m$ is the combined mass of the dry anode and cathode, and $\sigma_{\text{chem, i}}$ and $\sigma_{\text{eff, i}}$ are the ion concentrations of the feedwater and collected effluent, respectively.

### Electrode material

The pristine electrode material used in this work is activated carbon cloth (ACC-5002-15; Kynol Europa GmbH, Germany) with $\sim 600$ µm thickness and $\sim 1400$ m$^2$/g surface area, with the latter determined from N$_2$ gas sorption measurements.

The sulfonated electrode material was prepared as follows.

Pristine carbon cloth weighing about 4 g was inserted into a flask containing at least 28 mL of 20% fuming sulfuric acid (H$_2$SO$_4$). The flask was then closed with a stopper. After 24 h, the cloth was removed from the flask and soaked with an excess of hexane for 5–10 min. The cloth was then washed by soaking in deionized (DI) water at room temperature three additional times, each time for 30 min, and then dried in air at 80 °C for 12 h.

The pristine and sulfonated materials were cut into electrode squares with a cross-sectional area of 6.25 cm$^2$. Electrodes were rinsed with deionized (DI) water (18.2 MΩ, Synergy Water Purification System, Merck Millipore, USA) using the transmission mode. Raw data were smoothed, baseline-calibrated and normalized with the peak at 804 cm$^{-1}$.

### Fourier transform infrared spectroscopy

The pristine and sulfonated materials were characterized with FTIR to determine functional group bonds. Potassium bromide (KBr) was dried in a furnace before usage, and then about 400 mg was ground into powder before mixing with about 0.05 mg of electrode. Approximately 150 mg of the well-mixed powder was transferred to a KBr pellet die and pressed with 5 t of force for about 1 min. The prepared KBr pellet was then tested with FTIR (Bruker Tensor 27, Bruker Corporation, USA) using the transmission mode. Raw data were smoothed, baseline-calibrated and normalized with the peak at 804 cm$^{-1}$.

### Titrations

Approximately 0.2 g of electrode material was ground to a powder and added to a vessel containing 35 mL of 0.05 M NaOH and 35 mL of 18.2 MΩ DI water (Synergy Water Purification System; Merck Millipore, USA). The solution was bubbled with N$_2$ gas under stirring for 20 min, then sealed and equilibrated under stirring for 24 h. The solution was then transferred to a closed vessel in the titration system (iAquatrode Plus and 904 Titrand, Metrohm AG, Switzerland) and bubbled with N$_2$ for an additional 16 min. The nitrogen flow was then reduced to one bubble every several seconds to prevent air intrusion in the vessel. The solution was stirred for 4 min, then automatically titrated with 70 mL of 0.05 M HCl with a drift criterion of 0.5 mV/min. A blank sample without electrode was titrated in the same manner without the initial 20-min bubbling and 24 h equilibration steps. The titration curves are shown in Supplementary Fig. 3.

#### Titrination modeling

The chemical charge concentrations of Fig. 3b were obtained from a model-to-data fitting process using the equilibrium acid–base model of Guyes et al. with an additional strong acid term. Briefly, the model considers an electrode of mass $m$ and nanopore volume $V_{\text{np}}$, immersed in strong base (NaOH) of initial volume $V_{0}$ and concentration $c_{\text{NaOH}}$, which is titrated with volume $V$ of a strong-acid (HCl). Two phases are considered, the electrode nanopores and the bulk solution. No external electric potential is applied to the electrode, but the presence of charged surface groups causes a Donnan potential term $\Delta \phi_{\text{D}}$, to form between the nanopores and the bulk. The surface groups considered are a single weak acid group, A, a single strong-acid group assumed to be a sulfonic group, $-\text{SO}_3$, and a single weak base group, B. The groups A, $-\text{SO}_3\text{H}$, and B dissociate according to their equilibrium constants $K_{\text{A}} = c_{\text{A}} / c_{\text{HA}}$, $K_{\text{B}} = c_{\text{B}} / c_{\text{BH}}$, and $K_{\text{BH}} = c_{\text{BH}} / c_{\text{B}}$, respectively, where $c_{\text{A}}$, $c_{\text{BH}}$, and $c_{\text{B}}$ are the concentrations of the groups in charged form; $c_{\text{A}}$, $c_{\text{BH}}$, and $c_{\text{B}}$ are the concentrations of the groups in uncharged form; and $c_{\text{A}}$ and $c_{\text{BH}}$ are the micropore hydronium concentration (all concentrations are per unit nanopore volume).

In the absence of an applied potential, the nanopore charge balance is

$$
\sigma_{\text{chem}} + \sigma_{\text{elec}} = 0,
$$

and the nanopore chemical charge is

$$
\sigma_{\text{chem}} = F \left( -\frac{c_{\text{A}} + c_{\text{B}}}{1 + c_{\text{BH}} / K_{\text{B}}} - \frac{c_{\text{SO}_3 +} + c_{\text{SO}_3 \text{H}}}{1 + c_{\text{BH}} / K_{\text{D}}} + \frac{c_{\text{A}} + c_{\text{BH}}}{1 + c_{\text{BH}} / K_{\text{D}}} \right),
$$

and the nanopore ionic charge is $\sigma_{\text{ionic}} = -2(c_{\text{A}} + c_{\text{B}}) \sinh(\Delta \phi_{\text{D}})$. Ion size effects are neglected for simplicity. The nanopore charge balance Eq. (7) is solved numerically along with Na$^+$ and Cl$^-$ balances between the bulk and nanopores,

$$
V_{\text{NaOH}} v_{\text{NaOH}} c_{\text{NaOH}} \exp(-\Delta \phi_{\text{D}}) - (V_{0} + V - \nu_{\text{HCl}}) c_{\text{Na}^+} = 0,
$$

and

$$
V_{\text{HCl}} v_{\text{HCl}} c_{\text{HCl}} \exp(\Delta \phi_{\text{D}}) - (V_{0} + V - \nu_{\text{NaOH}}) c_{\text{Cl}^-} = 0.
$$

where $c_{\text{Na}^+}$ and $c_{\text{Cl}^-}$ are bulk concentrations, and electroneutrality in the bulk,

$$
\nu_{\text{Na}^+} c_{\text{Na}^+} + \nu_{\text{Cl}^-} c_{\text{Cl}^-} - K_{\text{w}} c_{\text{H}^+} c_{\text{OH}^-} = 0.
$$

The model was fitted to the data using the fitting parameters given in Supplementary Table 3. For further details, the reader is referred to Guyes et al. and Hemmatifar et al.
CDI characterization

Cell design. The custom-built FTE CDI cell consists of two electrodes electronically isolated by a separator (Whatman 2 cellulose filter paper, GE Life Sciences, 2.7 cm × 2.7 cm, 190 μm thickness). Graphite current collectors (FC-GR, Graphistore.com) contact the posterior side of each electrode. A 9 × 9 grid of 1.5 mm diameter holes was milled into each collector to allow water passage. The cell is enclosed on both sides with milled PVDVF blocks, which each contain one fluid flow line and a valve line for air removal. A funnel was cut into the downstream-side block to facilitate water flow toward the fluid line. Compressible, laser-cut ePTFE gaskets (Gore-Tex NSG16X-6; W.L. Gore & Associates, ~1.4 mm uncompressed thickness) provide sealing. The cell was assembled and fastened with stainless steel bolts. The internal cell volume is approximately 2 mL. A stainless-steel schematic is provided in Supplementary Fig. 4.

Electrochemical impedance spectroscopy (EIS). The separator resistance \( R_{sepa} \), the external electronic resistance \( R_{sepa} \), and the series resistance \( R_s \) were determined from a method based on Hawks et al.\(^\text{16} \) and used as simulation parameters. EIS spectra were taken over the frequency range 700 kHz to 40 mHz in the CDI cell with a potentiostat (Reference 3000, Gamry, USA). A two-point configuration was used, where the cathode was connected to the counter and reference wires and the anode was connected to the working and sensing wires. Both P–P and S–P configurations were evaluated, and feed solution was flowed through the cell at 1 mL/min. Spectra were taken with different separator thicknesses, achieved by stacking 1–4 separators together (Supplementary Fig. 5a, b). The minimum \( R(2) \) on each spectrum is the point of minimum phase angle, which we treat as \( R_s \). Supplementary Fig. 5c shows \( R_s \) vs. separator thickness with a linear fit, where the vertical intercept is \( R_{ER} \) (1.35 Ω for the P–P cell and 0.721 Ω for the S–P cell), and the slope is \( R_p \) (7.28 Ω/separat or for the P–P cell and 7.75 Ω/separat or for the S–P cell).

CDI setup and experiments. The CDI experimental setup consists of a 0.5 L glass bottle that served as the feedwater reservoir, a peristaltic pump (Masterflex 07551-30, Cole-Parmer, USA), the CDI cell, a conductivity sensor (TraceAdc 390-70; Innovative Sensor Technologies GmbH, Austria), and a voltage source (2400 Source Meter, Keithley Instruments, USA), the latter of which was connected to the cell’s current collectors. The pump, cell, and conductivity sensor were connected via flexible tubing. Feedwater was drawn from the reservoir and first passed through the cell, and then through the conductivity sensor. The cell was oriented so that water first passed through the cathode (electrode at negative potential) and then the anode (positive potential). The voltage source and pump were remotely controlled with a Matlab script. Current and voltage data were collected in Matlab, and conductivity data were recorded in the conductivity sensor software.

Feedwater comprised of 0.5 mM CaCl\(_2\) and 14 mM NaCl was prepared with CaCl\(_2\) (>99%; Fisher Scientific UK), NaCl (>99.5%, SDFCFL, India) and DI water. Prior to experiments, N\(_2\) gas was bubbled at a high rate in the feedwater reservoir for 20 min to remove dissolved oxygen, and then maintained at a lower rate for the duration of the experiment. The feedwater was then pumped through the system at a flow rate of 1 mL/min and the cell was discharged at 0 V until the current was negligible (<0.1 |mA in the S–P cell or <0.01 |mA in the P–P cell) in order to equilibrate the electrodes with the feedwater before beginning the experiment.

Experiments were carried out under the general framework described by Hawks et al.\(^\text{16} \), and performance metrics and indicators are reported according to the definitions therein. In each experiment, the cell was charged at a constant voltage (0.4–2.5 V) and discharged at 0 V in repeated cycling with a 1 mL/min feedwater flow rate. Each cycle lasted for a full-cycle time (FCT) of 6–30 min, which was divided equally into charge and discharge steps. Conditioning cycles were carried out until the cell reached the limit cycle, which is the cycle where the conductivity curve reaches dynamic steady-state behavior. Desalinated effluent was then collected for the limit cycle while the effluent conductivity was lower than the feedwater conductivity, and water with conductivity above the feed value was discarded to a waste container. For FCT 6 min, the limit cycle and the two following cycles were collected in the same manner and then combined in order to accumulate enough liquid for post-CDI analysis (see the Supplementary Table 1 for a breakdown of the limit cycle according to FCT and cell type). Each collected sample was weighed to calculate its volume with an assumed liquid density of 1 g/mL.

Post-CDI analysis

Sample conductivity and pH determinations. The conductivity sensor was calibrated with KCl 1.41 mS/cm standard (1.01203, Merck) at a 1 mL/min flow rate. Each desalinated water sample was pumped through the conductivity sensor at 1 mL/min until the reading stabilized (~3 min). The sensor and tubing were flushed with DI water between measurements.

The pH electrode and module (Aquamount 110, 904 Titrando, Metrohm AG, Switzerland) were calibrated with standard Metrohm buffers of pH 4, 7, and 9. The pH of each sample was measured under stirring with a drift criterion of 1 mV/min. These measurements were always carried out after conductivity measurements so as not to artificially increase the sample conductivity with the pH sensor filling electrolyte (KCI 3 M). The electrode was thoroughly rinsed with DI water between measurements.

Ca\(^{2+}\) and Na\(^+\) concentration determinations. The Ca\(^{2+}\) concentrations were measured via standard addition. In preparation, 5 mL of sample and 10 mL of 1 M KCl ionic strength adjustor (ISA) were pipetted into a glass beaker. Before each measurement, the Ca\(^{2+}\) ion-selective electrode (6.0510.100, Metrohm) was rinsed for at least 30 s in DI water. The electrode was immersed in the sample/ISA mixture, and 0.1 M CaCl\(_2\) standard (21059, Sigma-Aldrich, USA) was manually pipetted in increments of 0.1, 0.2, and 0.5 mL under stirring. A drift criterion of 1 mV/min was used for each increment. Each measurement was performed at constant temperature (~25 °C), and the DI rinsing water was replaced after 2–3 measurements to prevent contamination. The Ca\(^{2+}\) concentration of each sample was back-calculated to account for the initial dilution with ISA.

The Na\(^+\) concentrations were calculated via

\[ c_{Na^+} (\text{mM}) = C - 2C_{Ca^2+} \left( \frac{10^{-pH_2} + 1}{\Lambda_{Na^+}} \right) \]

where \( C \), \( C_{Ca^2+} \), and \( pH \) are the conductivity, Ca\(^{2+}\) concentration, and pH of the sample, respectively. The equivalent conductivities \( \Lambda_{Na^+} = 115.59 \text{ mS/cm/mM} \) and \( \Lambda_{Ca^2+} = 115.7 \text{ mS/cm/mM} \) are values at 20 mM concentration\(^\text{10} \), while \( C_{Ca^2+} = 349.65 \text{ mS/cm/mM} \) is at infinite dilution\(^\text{10} \). The concentrations \( c_{Na^+} \) and \( c_{Ca^2+} \) were used to calculate experimental separation factor with Eq. (1), SAR with Eq. (2), minimum desalination energy with Eq. (3), ion adsorption with Eq. (6), and additional performance metrics and indicators tabulated in Supplementary Table 1.

DATA AVAILABILITY

Data generated and analyzed during this study are included in this published article and its Supplementary Information files. Raw data used for this manuscript are available from the corresponding author within reasonable request.

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AUTHOR CONTRIBUTIONS
E.N.G. contributed to the research design, performed the CDI and EIS experiments and subsequent analyses, and was the primary manuscript author. A.N.S. developed the CDI model equations and code, performed the theoretical calculations, and wrote the simulations sections in the “Results” and “Methods”. Y.C. performed the electrode sulfonation, pristine and sulfonated electrode titrations, FTIR measurements and analyses, and titration model fitting. C.E.D. contributed the sulfonation protocol, advised on the sulfonation and FTIR methods, and participated in the manuscript editing. M.E.S. contributed to the research conceptualization and design, and to manuscript writing and editing. All authors contributed to the interpretation and discussion of results.

COMPETING INTERESTS
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