UC Berkeley UC Berkeley Previously Published Works

Title

Engineering Electrode Rinse Solution Fluidics for Carbon-Based Reverse Electrodialysis Devices.

Permalink https://escholarship.org/uc/item/1dg1558g

Journal ACS Applied Materials and Interfaces, 15(41)

Authors

Platek-Mielczarek, Anetta Lang, Johanna Töpperwien, Feline <u>et al.</u>

Publication Date

2023-10-18

DOI

10.1021/acsami.3c10680

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at <u>https://creativecommons.org/licenses/by/4.0/</u>

Peer reviewed

Engineering Electrode Rinse Solution Fluidics for Carbon-Based Reverse Electrodialysis Devices

Anetta Platek-Mielczarek, Johanna Lang, Feline Töpperwien, Dario Walde, Muriel Scherer, David P. Taylor, and Thomas M. Schutzius*



ABSTRACT: Natural salinity gradients are a promising source of so-called "blue energy", a renewable energy source that utilizes the free energy of mixing for power generation. One promising blue energy technology that converts these salinity gradients directly into electricity is reverse electrodialysis (RED). Used at its full potential, it could provide a substantial portion of the world's electricity consumption. Previous theoretical and experimental works have been done on optimizing RED devices, with the latter often focusing on precious and expensive metal electrodes. However, in order to rationally design and apply RED devices, we need to investigate all related transport phenomena—especially the fluidics of salinity gradient mixing and the redox electrolyte at various concentrations, which can have complex intertwined effects—in a fully functioning and scalable system. Here, guided by fundamental electrochemical and fluid dynamics theories, we work with an iron-based redox electrolyte with carbon electrodes in a RED device with tunable microfluidic



environments and study the fundamental effects of electrolyte concentration and flow rate on the potential-driven redox activity and power output. We focus on optimizing the net power output, which is the difference between the gross power output generated by the RED device and the pumping power input, needed for salinity gradient mixing and redox electrolyte reactions. We find through this holistic approach that the electrolyte concentration in the electrode rinse solution is crucial for increasing the electrical current, while the pumping power input depends nonlinearly on the membrane separation distance. Finally, from this understanding, we designed a five cell-pair (CP) RED device that achieved a net power density of 224 mW m⁻² CP⁻¹, a 60% improvement compared to the nonoptimized case. This study highlights the importance of the electrode rinse solution fluidics and composition when rationally designing RED devices based on scalable carbon-based electrodes.

KEYWORDS: reverse electrodialysis, RED, electrode rinse solution, ERS, blue energy, salinity gradient power, microfluidics, carbon electrodes, redox electrolyte

INTRODUCTION

The global demand for energy requires constant deployment of renewable energy sources. However, one such source that is often overlooked is the natural mixing of fresh (ca. 0.012% NaCl) and saltwater (ca. 3.5% NaCl)^{1,2}—like rivers flowing into the sea. The theoretical Gibbs free energy of mixing one cubic meter of fresh water with one cubic meter of saltwater gives 2 MJ, and if scaled globally has a yearly potential of 1000 TW h.^{3,4} Switzerland consumes yearly, on average, 225 TW h, according to the Federal Department of Foreign Affairs (FDFA).⁵ The two most important systems that can leverage this at the pilot-scale are pressure-retarded osmosis and reverse electrodialysis (RED).^{6–8} The former uses semipermeable membranes, which allow the fresh water to pass to the saltwater side through osmosis, building up significant pressures that can power turbines.⁹ The latter system uses alternating cation- and anion-exchange membranes (CEM and AEM, respectively), creating channels where on either side of each membrane freshwater and saltwater are passed.^{10,11} This

arrangement allows salt ions to diffuse across the membranes in opposite directions^{12,13}—due to the salinity gradient between the feed solutions—creating an electrical potential (Nernst) that drives reversible Faradaic reactions at the electrode/electrolyte interface¹⁴ and generates electricity.^{3,15} To facilitate this energy conversion process, an electrode rinse solution (ERS), consisting of two redox species—forming a redox pair—and a supporting electrolyte, is pumped, circulating it from one electrode to the other. The simplest RED system consists of one membrane, with freshwater and saltwater on either side, and this is defined as a cell pair (CP), which is in between two electrodes.^{16,17} In this way, the

Received: July 21, 2023 Accepted: September 22, 2023 Published: October 9, 2023







Figure 1. RED device design and characterization. (a) Schematic showing the RED device, which consists of alternating CEM and AEM separated by a distance, *h*, and filled with LC and HC salinity water solutions, which are pumped in between the membranes. An individual membrane–water contact area for one side of the membrane is defined as *A*. Each pair of CEM and AEM with an LC and a HC compartment in between forms a "cell pair" (CP) that creates Nernst potential called the electromotive force, EMF, and this device consists of n = 5 CPs what gives SEMF voltage output. Each end of the device is connected to a carbon electrode inside a compartment where the ERS is pumped and circulated through both carbon electrodes. The electrodes are connected to a copper current collector, which is connected to an external load for indirect current, *I*, measurements by closing the circuit with selected resistors by reading the voltage, *U*. We can control and measure the flow rate of the HC, LC, and ERS solutions as well as the pressure drop along the HC and LC flow paths and the pressure drop across the ERS pump. Electrical resistance values of various device components, R_i , are also shown. These values are calculated using the following assumptions: $h = 300 \,\mu$ m, HC = 1 M NaCl, LC = 0.01 M NaCl, ERS = 20 mM FeCl₂ = FeCl₃ in 0.01 M NaCl and carbon-felt electrodes (see Supporting Information Note 1). Performance of the RED device with n = 5 at a fixed flow rate of saline solutions (0.21 cm³ s⁻¹) and ERS (0.05 cm³ s⁻¹) using external load resistors: (b) Voltage, *U*, vs current, *I* and (c) net power density, $Pd_{net} = UI/(An)$, vs *I* and vs external load resistance, *R*, with a fixed pumping power density, $Pd_{in} = (\Delta p_{LC} \dot{V}_{LC} + \Delta p_{HC} \dot{V}_{HC} + \Delta p_{ERS} \dot{V}_{ERS})/(nA)$. This is the configuration used throughout the study, where individual parameters are varied, but *n* and *A* are kept constant. Error bars are calculated standard deviations of at least three independent RED device e

chemical potential between the feed solutions is directly fully converted into electrical energy, without being converted into thermal or mechanical energy.^{18,19} The RED device output voltage (U_{max}) is directly proportional to the number of CPs (n) connected in series, and the typical number of CPs used in RED devices is n = 50,^{20,21} but applied pilot plant studies going beyond n = 100 are also reported.^{22,23} One of the main failure reasons of RED devices is membrane fouling, addressed in fundamental studies of different materials—improved ones significantly increase the overall cost of a final device.^{24,25}

The most straightforward method to improve the power output of a real RED device is to modify its structure, especially membrane spacing defined as a membrane separation distance, $^{20,26-28}$ the number of CPs, $^{29-31}$ and the flow rate of the low- and high-salinity solutions.^{30,32-34} Typically, spacers are inserted between the anion- and cation-exchange membranes to form compartments filled with saline solutions.^{14,30,33,35} Nevertheless, depending on the properties of the spacers, they can induce more flow resistance and deteriorate the RED device performance.³⁶ This is why spacer-less systems or modified textured membranes are also introduced.^{9,29,37} Various membranes' modification and optimization for enhanced mass transport are reported.³⁸⁻⁴⁰ Several significant implications are outlined in theoretical predictions, investigating the impact of intermembrane distance on the power density of a RED system, but they remain unverified for these specific conditions.⁴¹ Universal findings discussed in the literature can be summarized as follows: the gross power density is higher for smaller membrane separation distances;²⁶ the pressure drop, Δp , across the device is strongly dependent on the membrane separation distance;^{20,26} the most common membrane separation distances are 100, 200, or 300 μ m,^{31,42,43} which seem to achieve a good balance between the gross power output and the pumping power input; the use of mixing promoters can be beneficial for systems operating with laminar flows;⁴⁴ textured membranes appear to be a better choice for decreasing the membrane separation distance than thin spacers $(<100 \ \mu m)$;^{20,27,44,45} the membrane thickness should be as low as possible to achieve high gross power density;⁴⁶ and the residence time of saline solutions-crucial for ensuring high ion flux through the membrane—can be tuned either by a membrane separation distance or feed solution flow rate.^{10,43,46} From this summary, it is clear that work has been done on optimizing the gross power output from RED devices for a given pumping power input-the difference between the two being the net power output-by focusing on tuning the fluidics between the membranes.⁴⁷ However, what has been neglected in these studies is the pumping power needed to circulate the ERS and its relationship to the net power output, which is nontrivial, taking into account both losses as well as the ability to transport redox species to the electrode interfaces driving the necessary reactions.

The gross power output from a RED device can also be increased by engineering the composition and concentration of the ERS and the composition and texture of the electrode interface, preferably sustainable and scalable.^{48–50} The lifecycle of materials used in the energy storage/conversion systems and their environmental impact are of high importance;^{51–53} therefore, carbon attracts scientists' attention in various electrochemical applications.^{48,54–57} Previous works have used carbon-based and metallic electrodes in RED devices: the former being carbon felt that is compatible with more

benign redox electrolytes^{58,59} and the latter being platinum, titanium, ruthenium, or iridium for their catalytic activity. 14,34,44,60,61 For carbon-based electrodes, previous works have used a reversible iron-based redox pair, Fe^{2+}/Fe^{3+} (concentration ranges from 20 to 50 mM), ^{58,62} with the supporting electrolyte NaCl or Na2SO4.63 For metallic electrodes, which have been studied to a higher extent than their carbon-based counterparts, many RED devices use $Fe(CN)_6^{3+}/Fe(CN)_6^{4+}$ as the redox pair with a range of concentrations from 50 to 400 mM, $^{27,34,35,64-66}$ with the supporting electrolyte NaCl. In each of these studies, regardless of the chosen electrode material and ERS, various ERS flow conditions have been implemented. Depending on the electrode size and texture, these ERS flow rates have ranged from 0.03 to 30 L min⁻¹. D'Angelo et al. studied the effect of the ERS flow rate on the gross power output,⁶⁷ and Tedesco et al. scaled up a RED device considering all circulating solutions (freshwater, saltwater, and ERS).^{68,69} From this, we see that a range of redox species concentrations and ERS flow rates have been investigated;⁷⁰ however, a RED device performance for net power output was not optimized. We find that a holistic view on a RED device net power output, accounting for the pumping power density input of the saline solutions and ERS for a range of redox species concentrations, is missing, especially for carbon-based electrodes.

Here, guided by fluid dynamics and electrochemical theories, we systematically investigate the effects of saline and ERS fluidics on gross and net power densities from a RED device. To illustrate this approach, we use carbon-based electrodes, which are scalable, cheap, and compatible with benign redox species, Fe²⁺/Fe³⁺. First, we explore experimentally the effect of engineering the membrane separation distance together with implementing ERS in the RED stack. We report a gross power density and a pumping power density, resulting in a net power density output, which provides a clear evaluation of the proposed systems. Guided by fundamental electrochemical characterization, we discuss the ERS composition and its further performance in a RED system. We explain the physics and chemistry behind RED operation by exploring the effect of electrolyte optimization, i.e., the concentration of the redox pair and supporting electrolyte, flow conditions, and fluidics engineering, to enhance energy conversion. Finally, to show the promise of this approach, we demonstrate a net power density of 224 mW m⁻² CP⁻¹ for a n = 5 CP RED device, a 60% improvement compared to the nonoptimized system defined as a control case. These results are underpinned by comparison of the fluidic and adsorption-reaction time scales.

RESULTS AND DISCUSSION

The RED device used in this study is built with n = 5 CPs connected in series with carbon-felt electrodes at either end of the device. Each CP consists of a CEM and an AEM separated by a distance *h*. Placing the CPs in series forms compartments where the low concentration (LC; 0.01 M NaCl) and high concentration (HC; 1 M NaCl) salinity solutions are pumped through. The values of *h* are varied between h = 100, 125, and 300 μ m. A schematic of the RED device is presented in Figure 1a with the calculated electrical resistance values, *R*, for each component (assuming $h = 300 \ \mu$ m; see Supporting Information Note 1). The concentration difference between the feed solutions is what generates the Nernst potential across the membrane (Supporting Information Note 2). When feed

solutions flow through the intermembrane compartments, the voltage spontaneously builds up depending on the membranes' permselectivity. Finally, an ERS, consisting of Fe^{2+}/Fe^{3+} redox species, is circulated over the electrodes, performing reversible reactions and generating current that can power external devices.

To measure the power produced by the RED device, the circuit was closed using resistors with variable resistance, and the voltage drop over the resistor was measured, enabling the calculation of steady-state current output, I, from the device, Figure 1b. In order to compare different devices, the operational parameter—power density, Pd, should be reported, which depends on the number of CPs, n_i and the area of the membrane/saline solution interface, A. The gross power density out of the device is therefore $Pd_{gross} = IU/(nA)$ (see Supporting Information Note 3). However, this parameter does not account for the power density needed to pump the saline solutions and ERS, $Pd_{in} = (\Delta p_{LC}\dot{V}_{LC} + \Delta p_{HC}\dot{V}_{HC} +$ $\Delta p_{\text{ERS}} \dot{V}_{\text{ERS}} / (nA)$, where Δp and \dot{V} are the pressure difference and volumetric flow rates, respectively, across the HC, LC, and ERS flow paths. Differential pressure sensors are introduced at the inlet and the outlet of the LC and HC flow paths, and Δp_{ERS} is measured with a pressure sensor across the pump that is circulating the ERS. With this, one can define the net power density, $Pd_{net} = Pd_{gross} - Pd_{in}$, Figure 1c. The maximum Pd_{net} recorded for the current configuration, which is based on the conditions most commonly used in RED studies, ^{10,26,30} is 159 mW CP⁻¹ m⁻² (LC: 0.01 M NaCl with $\dot{V}_{LC} = 0.21$ cm³ s⁻¹; HC: 1 M NaCl with $\dot{V}_{HC} = 0.21 \text{ cm}^3 \text{ s}^{-1}$; ERS: 20 mM Fe²⁺ = Fe³⁺ in 0.01 M NaCl with $\dot{V}_{ERS} = 0.05 \text{ cm}^3 \text{ s}^{-1}$, n = 5, A = 110 cm^2).

Various membrane separation distances h were tested in order to find the optimal configuration for further tests in terms of Pd_{net} h = 100, 125, and 300 μ m, Figure 2. The recorded U_{max} values for those configurations are comparable, ca. 0.9 V (see Figure S1); however, I depends on h. This correlation is almost proportional, with three times decrease in the channel height ($h = 300 \ \mu m$ to $h = 100 \ \mu m$) resulting in a 2-fold increase of Pd_{gross} : 159 mW CP⁻¹ m⁻² to 319 mW CP⁻¹ m^{-2} . This is mainly caused by manipulating the resistance of the LC compartment, R_{LC} , by changing the *h*. Interestingly, if a similar correlation is done for Pd_{net} the opposite trend is observed. The RED device with $h = 100 \ \mu m$ exhibits the highest Pd_{gross} but is not capable of producing enough power to maintain its own operation, $Pd_{net} < 0$. This is the result of the high pumping power requirements and a high value of Pd_{in} . Figure 2c shows a plot of experimental data of maximum Pd_{net} Pd_{in} , and Pd_{gross} for $h = 100 \ \mu m$, 125 μm , and 300 μm and their estimated values. We assume that Pd_{in} is proportional to A/h^3 , a Hagen-Pouiseulle flow, and is justified on the basis of the Reynolds number for all experiments conducted, Re < 1 (see also Supporting Information Note 4 and Figure S2 for the relationship between the flow rate and the pressure drop). We assume that Pd_{gross} is proportional to B/h, attributed to gradients in salinity concentration within the liquid, and following a Fickian diffusion process normal to the flow direction. Assuming these forms, a best fit is performed on the experimental data with $Pd_{net} = B/h - A/h^3$ (with constant values $A = 2.25 \times 10^8$; B = 33,030). From this, we see that by only analyzing the fluidics between the membranes, and using this simplified analysis, one can expect that we are operating close to the optimum condition ($h = 150 \ \mu m$). Based on what has been proposed in the literature,⁷¹ passive mixers' influence



Figure 2. Optimizing the net power density output through fluidic engineering. RED device, n = 5, with varied h, h = 100, 125, and 300 μ m: (a) Pd_{gross} vs I and (b) Pd_{net} vs I. Fixed flow rates for the saline solutions ($V_{LC} = V_{HC} = 0.21$ cm³ s⁻¹) and the ERS, 20 mM FeCl₂ = FeCl₃ in 0.01 M NaCl ($\dot{V}_{ERS} = 0.05 \text{ cm}^3 \text{ s}^{-1}$). When operating the RED device for constant flow rates of saline solutions and ERS, for small h (100 μ m), I increases together with the pumping power Pd_{in} (LC, HC, and ERS), leading to negative Pd_{net}. Optimum RED stack design with n=5 for different channel heights, h, e.g., membrane separation distance, using LC-0.01 M NaČl and HČ-1 M NaCl solutions: (c) maximal gross power density, Pd_{gross}, and pumping power, Pd_{in}, where the data points are experimental measurements (each data point represents an average of at least 3 experiments), and the lines are theoretical: Pd_{gross} based on a semiempirical assumption of U_{max} and I_{max} , resulting in a trend $Pd_{\text{gross}} = B/h$; Pd_{in} is computed based on the flow rate and the pressure drop (described above)— Pd_{in} = $\Sigma \dot{V}_i \Delta p_i$, where *i* counts for LC, HC, and ERS, following the correlation $Pd_{in} = A/h^3$.

on Pd_{net} was verified for $h = 300 \ \mu m$; however, we did not observe any improvements for our RED device (see Supporting Information Note 5 and Figure S3). Thus, we found $h = 300 \ \mu m$ as our optimal configuration with respect to



Figure 3. Engineering the ERS fluidics for enhancing the net power output. Electrochemical characterization of redox electrolyte solution with composition 20 mM FeCl₂ = FeCl₃, 0.01 M NaCl: (a) cyclic voltammetry at various scan rates (1, 5, 10, 20, 30, 40, and 50 mV s⁻¹) in the three-electrode setup with Ag wire as a quasi-reference electrode. Potential window is adjusted to reversible oxidation and reduction redox reaction peak, $U_{red-ox}i$ (b) maximum current peak (I) and potential difference ($U_{red-ox}i$) of Fe²⁺/Fe³⁺ redox reaction vs scan rate with U_{max} of RED device with n = 5. (c) Performance of RED device with n = 5 using LC and HC at flow rates $\dot{V}_{HC} = \dot{V}_{LC} = 0.21$ cm³ s⁻¹ in $h = 300 \,\mu$ m channels with ERS in the carbon electrode compartment (20 mM FeCl₂ = FeCl₃, 0.01 M NaCl) at different flow conditions resembling 2, 5, and 10 mV s⁻¹ (see Supporting Information Note 6): Pd_{gross} vs I_i (d) Pd_{net} vs I with pumping power values (LC + HC + ERS).

the fluidic design, for which Pd_{net} exceeds the theoretical prediction; further improvements will be done at these conditions. Next, the focus was on engineering the electrode/electrolyte interface to increase *I*, and, in effect, boost the Pd_{net} of the RED device.

For carbon-based electrodes, a sustainable $\mathrm{Fe}^{2+}/\mathrm{Fe}^{3+}$ redox pair has been proposed.^{58,62} However, considering the ERS composition and flow conditions, it has not been elaborated in the RED device. Previous work has used the following ERS composition with metallic electrodes: 20 mM Fe^{2+} = Fe³⁺. ^{58,67,69} Therefore, we select this specific composition and study the effect of flow rate to identify an optimum based on Pd_{net} , Figure 3. First, the ERS is characterized fundamentally outside of the RED device (see Methods) by applying different scan rates $s = 1 \text{ mV s}^{-1}$ to $s = 50 \text{ mV s}^{-1}$, Figure 3a. The maximum redox peak I_{max} increases with increasing s, according to the charge preservation theory.⁷² What needs to be considered is the U_{max} of the RED device, which is strictly correlated with *n*. Therefore, not all *s* would be applicable in the real RED device, as the limiting factor is its maximum voltage-the potential difference between the oxidation-reduction reactions, $U_{\rm red-ox}$, should be within $U_{\rm max}$. From Figure 3b, one can see that in the case of n = 5, only 2, 5, and 10 mV s⁻¹ can be applied in a way that $U_{\text{red-ox}} \leq U_{\text{max}}$. The

residence time of an ERS within a carbon electrode, τ , can be calculated based on $\tau = U_{red-ox}/s$ (see Supporting Information Note 6). Then, knowing our electrode volume, $V = 28 \text{ cm}^3$, and the time that ERS should be spent within its bulk for selected scan rate $s = 2 \text{ mV s}^{-1}$, $s = 5 \text{ mV s}^{-1}$, and s = 10 mV s^{-1} , $\tau = U_{red-ox}/s$, we can calculate $\dot{V}_{ERS} = V/\tau$, and we can correlate it with the s that was applied in fundamental cyclic voltammetry tests. We therefore tested various $\dot{V}_{
m ERS}$ near this estimated value of $V/\tau = 0.28 \text{ cm}^3 \text{ s}^{-1}$ that was calculated for s = 10 mV s⁻¹, so 0.21 cm³ s⁻¹, s = 5 mV s⁻¹: 0.14 cm³ s⁻¹, and s= 2 mV s⁻¹: 0.05 cm³ s⁻¹. The scan rate s of 10 mV s⁻¹ is limited not only by $U_{\text{red-ox}}$ for the RED with n = 5 but also by the pumping system's upper limitations (lower limitation of the applied flow rate is $0.05 \text{ cm}^3 \text{ s}^{-1}$). The highest flow rate, $V_{\text{ERS}} = 0.21 \text{ cm}^3 \text{ s}^{-1}$, is predicted to be the best among all ERS flow rates tested in the RED device, according to the high current recorded in the fundamental studies. From Figure 3c, it is clear that Pd_{gross} is not affected by the flow rate, indicating that we are in some type of adsorption-reaction-limited regime. Therefore, further increasing the flow rate should increase Pd_{in} and result in a decrease in Pd_{net} which is seen in Figure 3d. In contrast to metallic electrodes, when carbon-based electrodes are flushed with ERS, no increase in Pd_{gross} is observed (because of overlapping U-I curves with increasing V_{ERS} —

Figure S4). This clearly indicates that such an electrode/ electrolyte system is adsorption-reaction-limited, as the time scale for Fe²⁺ or Fe³⁺ to diffuse toward redox-active sites at the carbon interface is much longer⁷³ than the one obtained with the lowest \dot{V}_{ERS} . Based on the diffusion coefficient of Fe²⁺, Fe³⁺ in water, $D_{\text{Fe}^{2+}} = 7.19 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$, $D_{\text{Fe}^{3+}} = 6.04 \times 10^{-6} \text{ cm}^2$ s^{-1} , and electrode thickness, 0.25 cm, the average diffusion time equals 49,000 s-ca. 13.5 h. The electrolyte solution can search for easy passages within carbon felt electrodes to pass through, as it has substantial electric resistance (423 m Ω -Figure 1). When metal electrodes are in use, no limitations are observed-diffusion and adsorption, reaction-and the flow rate of ERS directly increases the recorded current.⁶⁷ Therefore, engineering of carbon-based electrodes should be the focus of future research to enhance wetting and ERS passages through porous electrode's bulk. Thus, to optimize Pd_{net} of a RED device with n = 5 with carbon-based electrodes, slow \dot{V}_{ERS} are recommended, and in our study $\dot{V}_{\text{ERS}} = 0.05 \text{ cm}^3$ s⁻¹ was further used.

After selection of the most optimal ERS flow conditions for the RED device with n = 5 and carbon-based electrodes, we focused on the optimization of ERS composition-concentration of redox species, Fe^{2+}/Fe^{3+} , Figure 4, and the supporting electrolyte, NaCl. The initial supporting electrolyte concentration equals LC-0.01 M-to avoid any cross contamination of ERS in the LC compartment. Thus, we also optimize the device's performance with respect to the supporting electrolyte concentration (Figure S5). From fundamental studies of the scan rate on the redox activity of the Fe^{2+}/Fe^{3+} redox pair, the highest s that could have been applied in the RED device (within the electrode compartment) is 10 mV s⁻¹. Thus, cyclic voltammetry with s = 10 mV s⁻¹ is conducted for ERS with Fe^{2+}/Fe^{3+} concentrations from 5 to 300 mM, the concentrations that are shown in previous studies of RED devices.^{58,62,67,69} With the evaluation of $U_{\rm red-ox}$ and I, one can see that only a concentration up to 50 mM lies in the U_{max} region of the RED device with n = 5 and $h = 300 \ \mu\text{m}$. This is why we find it important to rationalize selected conditions for application of ERS within the RED device to fully exploit its power generating potential. By increasing the supporting electrolyte concentration, a delicate difference in the U-I curves can be observed for 20 mM Fe^{2+} = Fe^{3+} . However, considering the increase of redox species concentration, one can notice a big improvement of Pd_{gross} (see Figure S5) and Pd_{net} . Flow rates of LC, HC, and ERS are the same for each experiment; therefore, an increase of Pd_{net} results directly from increased I for ERS with 50 mM Fe^{2+} = Fe^{3+} , reaching a maximum value of 224 mW CP⁻¹ m⁻², Figure 4c. This indicates that, by applying a mesoporous carbon-based electrode, one can tune Pd_{net} by applying more concentrated ERS. The optimal concentration is correlated with the number of CPs, n, in the RED device—thus, both factors should always be considered and tested together. We prove that for sustainable carbon electrode/ERS interface-with FeCl₂ and $FeCl_3$ —one can easily manipulate Pd_{net} by increasing the concentration of redox species in accordance with U_{max} of RED device related to n while keeping a low Pd_{in} penalty for pumping the ERS. Even if the pumping power used for ERS circulation is lower than those for LC and HC, it should not be neglected. Thus, our work has been compared with state-ofthe-art publications in Table 1, highlighting various in-house experimental conditions and RED performance metrics obtained for device laboratory-scale studies.



Figure 4. Rationalization of the ERS composition for carbon electrodes, guided by fundamental studies: redox pair and supporting electrolyte concentration in the specific RED device (n = 5). Electrochemical characterization of the redox pair (Fe^{2+}/Fe^{3+}) and the supporting electrolyte (NaCl) with various concentrations: (a) cyclic voltammetry at 10 mV s⁻¹ for concentrations from 50 to 300 mM of redox species with 0.01 M NaCl; (b) maximum current peak (*I*) and potential difference (U_{red-ox}) of the Fe²⁺/Fe³⁺ redox reaction vs redox pair concentration. Performance of the RED device with three optimized compositions of the ERS, with n = 5, $h = 300 \ \mu$ m, $\dot{V}_{LC, HC} = 0.21 \ cm^3 \ s^{-1}$, and $\dot{V}_{ERS} = 0.05 \ cm^3 \ s^{-1}$: (c) Pd_{net} vs *I* with pumping power values.

CONCLUSIONS

Our investigation of the RED device with five CPs reveals that fluidic engineering (such as height and mixing conditions) does not offer significant room for further improvement when commercially available membranes and carbon-based electrodes are used. However, we discovered that the composition and flow conditions of the ERS, which are often overlooked in

material	и	LC	$\dot{V}_{\rm LC}$	HC	\dot{V}_{HC}	ERS	\dot{V}_{ERS}	$Pd_{\rm in} =$	$Pd_{ m gross_max}$	$Pd_{\mathrm{net_max}}$	refs
carbon-based	S	0.01 M	$0.21 \text{ cm}^3 \text{ s}^{-1}$	1 M	$0.21 \text{ cm}^3 \text{ s}^{-1}$	50 mM FeCl ₂	$0.05 \text{ cm}^3 \text{ s}^{-1}$	$Pd_{inLC} + Pd_{inHC} + Pd_{inERS} = 39 mW CP^{-1} m^{-2}$	$263 \text{ mW CP}^{-1} \text{ m}^{-2} = 1.32 \text{ W m}^{-2}$	224 mW CP ⁻¹ m ⁻² = 1.12 W m^{-2}	this work
						50 mM FeCl ₃ 0.5 M NaCl					
	50	0.017 M		0.5 M		50 mM FeCl ₂					62
						50 mM FeCl ₃ 0.5 M NaCl-HCl					
	100	tap water	1.66 cm ³ s ⁻¹	natural seawater	$1.66 \ cm^3 \ s^{-1}$	NaCl	$1.66 \ cm^3 \ s^{-1}$		0.72 W m^{-2}		77
	7	0.017 M	$1.66\ cm^{3}\ s^{-1}$	0.6 M	$1.66 \ cm^3 \ s^{-1}$	50 mM Fe(SO ₃) 50 mM Fe ₂ (SO ₃) ₃	$5 \ cm^3 \ s^{-1}$		1.34 W m ⁻²		63
	29	0.017 M	$1.66\ cm^3\ s^{-1}$	0.6 M	$0.66 \ cm^3 \ s^{-1}$	1.2 M IVACI NaCl			$259 \text{ mW CP}^{-1} \text{ m}^{-2} =$ 7.5 W m^{-2}		78
precious metal-based	25	0.02 M	$12.5 \ cm^3 \ s^{-1}$	4 M	$12.5 \ cm^3 \ s^{-1}$	0.1 MK ₃ Fe(CN) ₆			4.78 W m ⁻²		27
						0.1 MK4Fe(CN) ₆ 2 M NaCl					
	S	1.6	5 cm s^{-1}	5 M	3 cm s ⁻¹	0.1 M K ₃ Fe(CN) ₆ 0.1 M K ₄ Fe(CN) ₆ 0.6 M NaCl	1 cm s ⁻¹	$Pd_{inLC} + Pd_{nHC}$	10.5 W m ⁻²		66
	50	0.1 M	1 cm s^{-1}	5 M	1 cm s^{-1}	0.1 M K ₃ Fe(CN) ₆	$30 \text{ L } \text{h}^{-1} = 8.33$ $cm^3 s^{-1}$	$Pd_{inLC} + Pd_{inHC}$		$4.5 \text{ W} \text{ m}^{-2}$	64
						0.1 M K4Fe(CN)6 2.5 M NaCl					
	50	$1 g L^{-1} = 0.017 M$		$30 g L^{-1} = 0.5 M$		50 mM K ₄ Fe(CN) ₆		$Pd_{inLC} + Pd_{inHC}$		$0.71 \mathrm{ W m^{-2}}$	10
						50 mM ₃ Fe(CN) ₆ 0.25 M NaCl					
	50	$1 g L^{-1} = 0.017 M$	0.4 L min ⁻¹ = $6.7 \text{ cm}^3 \text{ s}^{-1}$	$30 g L^{-1} = 0.5 M$	0.4 L min ⁻¹ = $6.7 \text{ cm}^3 \text{ s}^{-1}$	$50 \text{ mM K}_4 \text{Fe}(\text{CN})_6$	$60 \text{ mL } \min^{-1} = 1 \text{ cm}^3 s^{-1}$	1 Pd _{gross}	0.93 W m^{-2}		21
						50 mM K ₃ Fe(CN) ₆ 1 M NaCl					

RED studies, can significantly affect the overall RED performance and should not be ignored. By conducting fundamental three-electrode studies of the electrochemical behavior of the ERS, we were able to preselect the composition and operational conditions that led to the optimization of the RED device using three different electrolytes (vary in the concentration of redox species and the supporting electrolyte). We show that rational selection of ERS and its flow conditions can affect the maximum current and redox reaction voltage difference, which need to be adjusted to the system's operational metrics, specifically, the maximum voltage of the RED device. Considering it, Pd_{net} can be increased by keeping a low Pd_{in} penalty for the ERS flow and a high redox activity by using redox species with higher concentrations.

METHODS

RED Device. The in-house-made RED stack consists of two PLEXIGLAS end plates, in between which the electrodes—filled with

Table 2. Physicochemical Characterization of Saline Solutions (HC and LC) and Redox Electrolyte (ERS: 20 mM FeCl₂ = FeCl₃ in 0.01 M NaCl) before and after Operation of the RED Device with n = 5, $h = 300 \,\mu\text{m}$, $\dot{V}_{\text{HC}} = \dot{V}_{\text{LC}} = 0.21 \,\text{cm}^3 \,\text{s}^{-1}$, $\dot{V}_{\text{ERS}} = 0.05 \,\text{cm}^3 \,\text{s}^{-1}$, at the Inlet and Outlet

	inlet	outlet
aqueous solutions	$\sigma~({ m mS~cm^{-1}})$	$\sigma~({ m mS~cm^{-1}})$
HC	87	44
LC	1	53
ERS	14	10

ERS, membranes, and fluid compartments of LC and HC solutions (channels formed by gaskets—a membrane separation distance) are pressed together with 10 screws. Each end plate has an inlet and an outlet for the ERS. The front end plate has inlets for LC and HC saline solutions in the counter-flow arrangement, and the back end plate has the corresponding outlets. In the electrode compartment, adhesive copper foil playing the role of a current collector is directly connected to a carbon felt electrode (SGL Carbon GFD 2.5 EA) and connected to external wiring. ERS co-flows-within the carbon electrode's bulk-with the neighboring LC and HC solutions. To avoid ionic short-circuiting, the length of the connecting tube between the two electrodes (ERS must pass one over each other before making the closed loop) is chosen to be 65 cm long with a tubing resistance of 142 M Ω . As the tubing resistance is higher by an order of magnitude than Ri-the internal resistance of the RED device-no short-circuit losses are expected.⁷⁴ Commercial ion-separation membranes are selected: AEM FAS-PET-75-anion-exchange membrane and CEM FKS-PET-75-cation-exchange membrane (Fumasep) with a thickness of 75 μ m. Membranes tend to expand when in contact with water, so they are presoaked in DI water for 72 h, and later holes for the screws are punched. Three different materials are utilized as the gasket—NBR with a thickness of $h = 300 \ \mu m$, MDT foil (3M) with h= 125 μ m, and Folex foil (PE) with $h = 100 \mu$ m. No additional spacers are used. MDT foil is a three-layer adhesive tape that after operation within a RED device tend to self-detach its layers. Thus, we believe that its low mechanical integrity causes big experimental deviations for RED device runs with $h = 125 \ \mu m$ visible in the biggest error bars. Passive micromixers are laser cut (Trotec Laser Cutter, Speedy 300) from Folex foil using the following parameters: power, 10; velocity, 1%; and frequency, 1 kHz. To reconstruct the h = 300 μ m channel, either one NBR gasket or three Folex foils, F-F-F, are placed between two membranes.

Redox Electrolyte. FeCl₂, FeCl₃, and NaCl are purchased from Sigma-Aldrich with ACS quality min. 95%. DI water is used to prepare





Figure 5. Physico-chemical characterization of ERS with $Fe^{2+} = Fe^{3+} = 0.02$, 0.05, 0.1, 0.2, 0.3, 0.5, 1, and 2 M in 0.5 M NaCl supporting electrolyte: (a) photo of ERS solutions (~20 mL); (b) conductivity and pH vs concentration of the Fe^{2+}/Fe^{3+} redox pair.

all solutions. ERS is always done in the same order: dissolution of NaCl, FeCl₃, and FeCl₂ in appropriate mass defined by the molar concentration of the final solution. For one experimental run of a RED operation, a loop of 250 mL of ERS is employed for a RED stack with n = 5 and A = 110 cm². ERS is prepared freshly once per week and kept without light exposure to avoid any side reactions or decomposition of iron-based species. ERS composition optimization is done and verified within a RED device, $n = 5 \mu m$ and $h = 300 \mu m$, for three formulations

 $20 \text{ mM FeCl}_2 = \text{FeCl}_3, \qquad 0.01 \text{ M NaCl} \qquad (i)$

 $20 \text{ mM FeCl}_2 = \text{FeCl}_3, \qquad 0.5 \text{ M NaCl}$ (ii)

$$50 \text{ mM FeCl}_2 = \text{FeCl}_3, \quad 0.5 \text{ M NaCl}$$
 (iii)

Conductivity and pH Tests. LC and HC solutions are verified before and after operation using electrochemical impedance spectroscopy at 100 kHz in a Teflon ring (10 mm in diameter and 4.4 mm in height) placed in a Swagelok Teflon body with stainless-steel current collectors, Table 2. The extrapolated real part of the impedance, Z(Re), where the imaginary Z'(Re) = 0, is used for conductivity calculations, that is, $\sigma = 1/Z(Re)$.

Characterization of the Redox Electrolyte. Fundamental characterization of the ERS is done in a three-electrode setup with an excess of electrolyte using the cyclic voltammetry technique (with scan rates ranging from 1 to 50 mV s⁻¹). All measurements (at different scan rates, ERS concentrations, and ERS compositions) are done using a potentiostat/galvanostat SP-200 (Biologic). 20 mL of solution is tested using two carbon felt electrodes (counter electrode twice the size of the working one, 2×112 to 2×56 mm²), and a Ag wire is employed as a quasi-reference electrode, after verifying its stable potential in studied solutions. All ERS solutions have pH < 3, as shown in Figure 5. According to the Pourbaix diagram of iron species in aqueous solutions,⁷⁵ the potential stability of the Fe²⁺/Fe³⁺ redox pair is constant for pH < 3.

The highest conductivity of ERS with 0.5 M NaCl supporting electrolyte has been found for a concentration of Fe^{2+}/Fe^{3+} redox pair equal to 0.75 M, Figure 5. After exceeding this concentration, deterioration of the ERS performance, especially its reversibility, is expected. Such behavior is in accordance with physicochemical properties of aqueous electrolyte solutions.⁷⁶

Characterization of the RED Device. After building the RED device with n = 5, all of the fluid compartments are flushed with DI water for 5 min to get rid of trapped air bubbles and possible contamination. ERS is first connected to the RED stack, and electrode compartments are flushed with it as long as ERS is collected at the outlet (observed in the visible yellow color of ERS solution compared to colorless DI water), usually for an additional 5-10 min. This way, electrodes are fully filled with the redox electrolyte before connecting the LC and HC saline solutions. After the ERS is reaching the outlet of the second electrode, the ERS outlet is connected to the first electrode inlet reservoir preserving a closed ERS loop. From this point, the voltage is recorded until reaching the plateau of U_{max} using a Keithley 2000 multimeter. After $U_{\rm max}$ is reached and stabilized (ca. 1 V), the voltage drop across a series of external load resistors is measured, namely, 1.88, 5.26, 7.72, 10.44, 15.17, 18.00, 22.08, 26.76, 38.70, and 56.21 Ω (TRU components, Vishay Intertechnology). Each resistor is inserted as long as the voltage reaches the plateau and/or maximally for 2 min.

ASSOCIATED CONTENT

Data Availability Statement

All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supporting Information. Additional data related to this paper are available on request.

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.3c10680.

Resistance calculations; electromotive force and voltage calculations; gross, pumping, and net power density calculations; fluidics; passive mixers design; ERS fluidics; performance of the RED device n = 5 with different channel heights h; U-I curve; engineering the RED device n = 5 with different channel heights h; pressure drop of HC and LC vs \dot{V} , vs h; enhanced mixing at the membrane/saline solution interface; engineering ERS fluidics in RED device n = 5 with $h = 300 \ \mu$ m; and engineering ERS composition in RED device n = 5 with $h = 300 \ \mu$ m (PDF)

AUTHOR INFORMATION

Corresponding Author

Thomas M. Schutzius – Laboratory for Multiphase Thermofluidics and Surface Nanoengineering, Department of Mechanical and Process Engineering, ETH Zurich, Zurich CH-8092, Switzerland; Department of Mechanical Engineering, University of California, Berkeley, Berkeley, California 94720, United States; oricid.org/0000-0003-3309-3568; Email: tschutzius@berkeley.edu

Authors

- Anetta Platek-Mielczarek Laboratory for Multiphase Thermofluidics and Surface Nanoengineering, Department of Mechanical and Process Engineering, ETH Zurich, Zurich CH-8092, Switzerland; ◎ orcid.org/0000-0001-6231-3908
- Johanna Lang Laboratory for Multiphase Thermofluidics and Surface Nanoengineering, Department of Mechanical and Process Engineering, ETH Zurich, Zurich CH-8092, Switzerland
- Feline Töpperwien Laboratory for Multiphase Thermofluidics and Surface Nanoengineering, Department of Mechanical and Process Engineering, ETH Zurich, Zurich CH-8092, Switzerland
- Dario Walde Laboratory for Multiphase Thermofluidics and Surface Nanoengineering, Department of Mechanical and Process Engineering, ETH Zurich, Zurich CH-8092, Switzerland
- Muriel Scherer Laboratory for Multiphase Thermofluidics and Surface Nanoengineering, Department of Mechanical and Process Engineering, ETH Zurich, Zurich CH-8092, Switzerland; orcid.org/0000-0003-3777-8206
- **David P. Taylor** Laboratory of Thermodynamics in Emerging Technologies, Department of Mechanical and Process Engineering, ETH Zurich, Zurich 8092, Switzerland

Complete contact information is available at: https://pubs.acs.org/10.1021/acsami.3c10680

Author Contributions

T.M.S. and A.P.-M. conceived the project and planned the experiments. J.L., F.T., D.W., and M.S. carried out the experiments. T.M.S., A.P.-M, D.T., and J.L. analyzed the data and developed the theoretical analysis. A.P.-M. and T.M.S. wrote the paper. All authors proofread the paper, made comments, and approved the manuscript.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors acknowledge the European Commission and the European Research Council for financial support by the

Starting Grant project (GA 853257) under the European Union's Horizon 2020 Research and Innovation Programme. Dimos Poulikakos is acknowledged for his strong support. Jovo Vidic is acknowledged for the programming of the voltage/ current measurements with the LabView support for Keithley Power Supply. Peter Fuesi is acknowledged for his technical support and spare parts manufacturing. Julian Schmid and Tobias Armstrong are acknowledged for fruitful discussions and input.

NOMENCLATURE

Α	area of the membrane/saline solution interface
	(one-side), m ²
А	constant for Hagen-Pouiseulle flow
а	width of the grooves, m
В	constant for the Fickian diffusion process normal
	to the flow direction
h	width of ridges m
0	concentration of the cation or anion in the UC
L	concentration of the cation of allon in the FIC
(TP)	and LC, M
CP	cell pair
d	depth of the grooves, m
$\Delta p_{\rm LC/HC/ERS}$	pressure drop across the device, respectively, for LC, HC, and ERS, bar
$d_{\rm h}$	characteristic length of a fluid channel with a
	rectangular cross section equals the hydraulic
	diameter, m
EME	electromotive force V
EDS	electroniotive force, v
LKS	electrode fillse solution
Jobs	Destruction factor
F	Faraday constant, C mol
h	membrane separation distance/channel height,
	μ m
HC	high-concentration feed solution
Ι	current output, mA
L	channel length, m
LC	low-concentration feed solution
п	number of cell pairs
N	number of grooves per half-cycle
P	asymmetric factor equals w/w -
Dd	nower density $W CP^{-1} m^{-2}$
DJ	prover density; w Cr m
Pu_{gross}	gross power density, w CP in $W CP^{-1} m^{-2}$
Pa_{in}	pumping power density, w CP m
Pa _{net}	net power density, W CP m 2
P _{wet}	cross-sectional perimeter wetted by the shear
	stress, m
r	resistance of one cell pair, Ω
R	universal gas constant, J mol ⁻¹ K ⁻¹
$R_{AEM/CEM}$	resistance of the anion-/cation-exchange mem-
	brane, respectively, Ω
Re	Revnolds number (for LC, HC, and ERS), -
RED	reverse electrodialysis
R ,	resistance of the electrode compartment (con-
elec	sisting of the shielding membrane the ERS and
	the electrodes themselves) O
л	the electrodes themselves), s2
$K_{\rm HC/LC}$	resistance of the HC/LC compartment, respec-
	tively, W
<i>R</i> _i	internal resistance of RED stack, Ω
R _{non-ohmic}	non-Ohmic resistance of RED stack, Ω
R _{ohmic}	Ohmic resistance of RED stack, Ω
5	scan rate, mV s ⁻¹
Т	temperature, K

www.acsami.org

Research Article

τ	residence time of ERS within the electrode
	compartment, s
$U_{\rm max}$	RED device voltage output, V
$U_{\rm red-ox}$	potential difference of the oxidation-reduction
	reaction of ERS, V
V	electrode volume, m ³
\dot{V}	volumetric flow rates for LC, HC, and ERS,
	respectively, $m^3 s^{-1}$
w	channel width, m
w_{l}	width of the long arm, m
w _s	width of the short arm, m
z	valence of the ion, -
$\alpha_{ m CEM/AEM}$	apparent permselectivity of AEM and CEM,
	respectively, -
γ	activity coefficient of the cation or anion in the
	HC and LC, -
η	fluid dynamic viscosity, Pa s
θ	groove intersection angle, $^\circ$
ρ	fluid density, kg m ⁻³
$\sigma_{ m HC/LC}$	ionic conductivity of respectively, HC and LC
	compartments, respectively, S m^{-1}
υ	fluid velocity, m s ⁻¹

REFERENCES

(1) Norman, R. S. Water Salination: A Source of Energy. Science 1974, 186 (4161), 350-352.

(2) Cosenza, A.; Campisi, G.; Giacalone, F.; Randazzo, S.; Cipollina, A.; Tamburini, A.; Micale, G. Power Production from Produced Waters via Reverse Electrodialysis: A Preliminary Assessment. *Energies* **2022**, *15* (11), 4177.

(3) Siria, A.; Bocquet, M.-L.; Bocquet, L. New avenues for the largescale harvesting of blue energy. *Nat. Rev. Chem* **2017**, *1* (11), 0091.

(4) Tufa, R. A.; Noviello, Y.; Di Profio, G.; Macedonio, F.; Ali, A.; Drioli, E.; Fontananova, E.; Bouzek, K.; Curcio, E. Integrated membrane distillation-reverse electrodialysis system for energyefficient seawater desalination. *Appl. Energy* **2019**, *253*, 113551.

(5) F.D.o.F.A. FDFA, Energy. https://www.eda.admin.ch/ aboutswitzerland/en/home/wirtschaft/energie/energie---fakten-undzahlen.htmwebl. (accessed July 10, 2023).

(6) Logan, B. E.; Elimelech, M. Membrane-based processes for sustainable power generation using water. *Nature* **2012**, 488 (7411), 313–319.

(7) Biesheuvel, P. M.; Porada, S.; Elimelech, M.; Dykstra, J. E. Tutorial review of reverse osmosis and electrodialysis. *J. Membr. Sci.* **2022**, 647, 120221.

(8) Choi, J.; Oh, Y.; Chae, S.; Hong, S. Membrane capacitive deionization-reverse electrodialysis hybrid system for improving energy efficiency of reverse osmosis seawater desalination. *Desalination* **2019**, *462*, 19–28.

(9) Yip, N. Y.; Elimelech, M. Thermodynamic and Energy Efficiency Analysis of Power Generation from Natural Salinity Gradients by Pressure Retarded Osmosis. *Environ. Sci. Technol.* **2012**, *46* (9), 5230–5239.

(10) Veerman, J.; Saakes, M.; Metz, S. J.; Harmsen, G. J. Electrical Power from Sea and River Water by Reverse Electrodialysis: A First Step from the Laboratory to a Real Power Plant. *Environ. Sci. Technol.* **2010**, *44* (23), 9207–9212.

(11) Rijnaarts, T.; Moreno, J.; Saakes, M.; de Vos, W. M.; Nijmeijer, K. Role of anion exchange membrane fouling in reverse electrodialysis using natural feed waters. *Colloids Surf.*, A **2019**, *560*, 198–204.

(12) Biesheuvel, P. M.; Dykstra, J. E.; Porada, S.; Elimelech, M. New parametrization method for salt permeability of reverse osmosis desalination membranes. *J. Membr. Sci. Lett.* **2022**, *2* (1), 100010.

(13) Wu, Q. Y.; Wang, C.; Wang, R.; Chen, C.; Gao, J.; Dai, J.; Liu, D.; Lin, Z.; Hu, L. Salinity-Gradient Power Generation with Ionized Wood Membranes. *Adv. Energy Mater.* **2020**, *10* (1), 1902590.

(14) Veerman, J.; Post, J. W.; Saakes, M.; Metz, S. J.; Harmsen, G. J. Reducing power losses caused by ionic shortcut currents in reverse electrodialysis stacks by a validated model. *J. Membr. Sci.* **2008**, *310* (1–2), 418–430.

(15) Chae, S.; Kim, H.; Gi Hong, J.; Jang, J.; Higa, M.; Pishnamazi, M.; Choi, J.-Y.; Chandula Walgama, R.; Bae, C.; Kim, I. S.; Park, J.-S. Clean power generation from salinity gradient using reverse electrodialysis technologies: Recent advances, bottlenecks, and future direction. *Chem. Eng. J.* **2023**, 452 (4), 139482.

(16) Veerman, J.; Vermaas, D. A. 4 - Reverse electrodialysis: Fundamentals. *Sustainable Energy from Salinity Gradients;* Woodhead Publishing, 2016; pp 77–133.

(17) Abdullah Shah, S.; Haider, Z.; Shahbabaei, M.; Kim, D. Development of an Efficient System for Blue Energy Production Based on Reverse Electrodialysis (RED) by Optimizing Electrolyte Composition: Experimental and Theoretical Simulations. *Energy Fuels* **2022**, *36* (12), 6353–6361.

(18) Yip, N. Y.; Elimelech, M. Comparison of Energy Efficiency and Power Density in Pressure Retarded Osmosis and Reverse Electrodialysis. *Environ. Sci. Technol.* **2014**, *48* (18), 11002–11012.

(19) Vermaas, D. A.; Saakes, M.; Nijmeijer, K. Capacitive Electrodes for Energy Generation by Reverse Electrodialysis. *Procedia Eng.* **2012**, 44, 496–497.

(20) Cipollina, A.; Micale, G.; Tamburini, A.; Tedesco, M.; Gurreri, L.; Veerman, J.; Grasman, S. 5 - Reverse Electrodialysis: Applications. *Sustainable Energy from Salinity Gradient*; Woodhead Publishing, 2016, 135–180..

(21) Veerman, J.; Saakes, M.; Metz, S. J.; Harmsen, G. J. Reverse electrodialysis: Performance of a stack with 50 cells on the mixing of sea and river water. *J. Membr. Sci.* **2009**, 327 (1–2), 136–144.

(22) Nam, J.-Y.; Hwang, K.-S.; Kim, H.-C.; Jeong, H.; Kim, H.; Jwa, E.; Yang, S.; Choi, J.; Kim, C.-S.; Han, J.-H.; Jeong, N. Assessing the behavior of the feed-water constituents of a pilot-scale 1000-cell-pair reverse electrodialysis with seawater and municipal wastewater effluent. *Water Res.* **2019**, *148*, 261–271.

(23) Tian, H.; Wang, Y.; Pei, Y.; Crittenden, J. C. Unique applications and improvements of reverse electrodialysis: A review and outlook. *Appl. Energy* **2020**, *262*, 114482.

(24) Ali, A.; Tufa, R. A.; Macedonio, F.; Curcio, E.; Drioli, E. Membrane technology in renewable-energy-driven desalination. *Renewable Sustainable Energy Rev.* **2018**, *81*, 1–21.

(25) Tufa, R. A.; Curcio, E.; Brauns, E.; van Baak, W.; Fontananova, E.; Di Profio, G. Membrane Distillation and Reverse Electrodialysis for Near-Zero Liquid Discharge and low energy seawater desalination. *J. Membr. Sci.* **2015**, 496, 325–333.

(26) Vermaas, D. A.; Saakes, M.; Nijmeijer, K. Doubled Power Density from Salinity Gradients at Reduced Intermembrane Distance. *Environ. Sci. Technol.* **2011**, 45 (16), 7089–7095.

(27) Hulme, A. M.; Davey, C. J.; Tyrrel, S.; Pidou, M.; McAdam, E. J. Transitioning from electrodialysis to reverse electrodialysis stack design for energy generation from high concentration salinity gradients. *Energy Convers. Manage.* **2021**, *244*, 114493.

(28) Hu, J.; Xu, S.; Wu, X.; Wu, D.; Jin, D.; Wang, P.; Leng, Q. Multi-stage reverse electrodialysis: Strategies to harvest salinity gradient energy. *Energy Convers. Manage.* **2019**, *183*, 803–815.

(29) Pawlowski, S.; Crespo, J. G.; Velizarov, S. Pressure drop in reverse electrodialysis: Experimental and modeling studies for stacks with variable number of cell pairs. *J. Membr. Sci.* **2014**, *462*, 96–111.

(30) Altiok, E.; Kaya, T. Z.; Guler, E.; Kabay, N.; Bryjak, M. Performance of Reverse Electrodialysis System for Salinity Gradient Energy Generation by Using a Commercial Ion Exchange Membrane Pair with Homogeneous Bulk Structure. *Water* **2021**, *13* (6), 814.

(31) Zhang, W.; Han, B.; Tufa, R. A.; Tang, C.; Liu, X.; Zhang, G.; Chang, J.; Zhang, R.; Mu, R.; Liu, C.; Song, D.; Li, J.; Ma, J.; Zhang, Y. Tracing the impact of stack configuration on interface resistances in reverse electrodialysis by in situ electrochemical impedance spectroscopy. *Front Environ. Sci. Eng.* **2022**, *16* (4), 46.

(32) He, Z.; Gao, X.; Zhang, Y.; Wang, Y.; Wang, J. Revised spacer design to improve hydrodynamics and anti-fouling behavior in reverse

(33) Moreno, J.; Slouwerhof, E.; Vermaas, D. A.; Saakes, M.; Nijmeijer, K. The Breathing Cell: Cyclic Intermembrane Distance Variation in Reverse Electrodialysis. *Environ. Sci. Technol.* **2016**, *50* (20), 11386–11393.

(34) Ortiz-Imedio, R.; Gomez-Coma, L.; Fallanza, M.; Ortiz, A.; Ibañez, R.; Ortiz, I. Comparative performance of Salinity Gradient Power-Reverse Electrodialysis under different operating conditions. *Desalination* **2019**, 457, 8–21.

(35) Choi, J.; Kim, W.-S.; Kim, H. K.; Yang, S. C.; Han, J.-H.; Jeung, Y. C.; Jeong, N. J. Fouling behavior of wavy-patterned pore-filling membranes in reverse electrodialysis under natural seawater and sewage effluents. *npj Clean Water* **2022**, *5* (1), 6–12.

(36) Ma, T.; Balanzat, E.; Janot, J.-M.; Balme, S. B. Nanopore Functionalized by Highly Charged Hydrogels for Osmotic Energy Harvesting. ACS Appl. Mater. Interfaces **2019**, *11* (13), 12578–12585.

(37) Pawlowski, S.; Crespo, J. G.; Velizarov, S. Profiled Ion Exchange Membranes: A Comprehensible Review. *Int. J. Mol. Sci.* **2019**, 20 (1), 165.

(38) Rouilly, M. V.; Kötz, E.; Haas, O.; Scherer, G. G.; Chapiro, A. Proton exchange membranes prepared by simultaneous radiation grafting of styrene onto Teflon-FEP films. Synthesis and characterization. J. Membr. Sci. **1993**, 81 (1–2), 89–95.

(39) Abidin, M. N. Z.; Nasef, M. M.; Veerman, J. Towards the development of new generation of ion exchange membranes for reverse electrodialysis: A review. *Desalination* **2022**, *537*, 115854.

(40) Fan, H.; Yip, N. Y. Elucidating conductivity-permselectivity tradeoffs in electrodialysis and reverse electrodialysis by structure-property analysis of ion-exchange membranes. *J. Membr. Sci.* **2019**, *573*, 668–681.

(41) Ciofalo, M.; La Cerva, M.; Di Liberto, M.; Gurreri, L.; Cipollina, A.; Micale, G. Optimization of net power density in Reverse Electrodialysis. *Energy* **2019**, *181*, 576–588.

(42) Han, J.-H.; Hwang, K.-S.; Jeong, H.; Byeon, S.-Y.; Nam, J.-Y.; Kim, C.-S.; Kim, H.; Yang, S.; Choi, J. Y.; Jeong, N. Electrode system for large-scale reverse electrodialysis: water electrolysis, bubble resistance, and inorganic scaling. *J. Appl. Electrochem.* **2019**, *49* (5), 517–528.

(43) Veerman, J.; Saakes, M.; Metz, S. J.; Harmsen, G. J. Reverse electrodialysis: A validated process model for design and optimization. *Chem. Eng. J.* **2011**, *166* (1), 256–268.

(44) Vermaas, D. A.; Saakes, M.; Nijmeijer, K. Enhanced mixing in the diffusive boundary layer for energy generation in reverse electrodialysis. *J. Membr. Sci.* **2014**, *453*, 312–319.

(45) Mehdizadeh, S.; Yasukawa, M.; Abo, T.; Kakihana, Y.; Higa, M. Effect of spacer geometry on membrane and solution compartment resistances in reverse electrodialysis. *J. Membr. Sci.* **2019**, *572*, 271–280.

(46) Brauns, E. Salinity gradient power by reverse electrodialysis: effect of model parameters on electrical power output. *Desalination* **2009**, 237 (1–3), 378–391.

(47) Kim, H.; Yang, S.; Choi, J.; Kim, J.-O.; Jeong, N. Optimization of the number of cell pairs to design efficient reverse electrodialysis stack. *Desalination* **2021**, *497*, 114676.

(48) Acevedo, B.; Barriocanal, C.; Lupul, I.; Gryglewicz, G. Properties and performance of mesoporous activated carbons from scrap tyres, bituminous wastes and coal. *Fuel* **2015**, *151*, 83–90.

(49) Fic, K.; Platek, A.; Piwek, J.; Frackowiak, E. Sustainable materials for electrochemical capacitors. *Mater. Today* **2018**, *21* (4), 437–454.

(50) Cataldi, P.; Athanassiou, A.; Bayer, I. Graphene Nanoplatelets-Based Advanced Materials and Recent Progress in Sustainable Applications. *Appl. Sci.* **2018**, *8* (9), 1438.

(51) Piątek, J.; Afyon, S.; Budnyak, T. M.; Budnyk, S.; Sipponen, M. H.; Slabon, A. Sustainable Li-Ion Batteries: Chemistry and Recycling. *Adv. Energy Mater.* **2021**, *11* (43), 2003456.

(52) Sougrati, M. T.; Arayamparambil, J. J.; Liu, X.; Mann, M.; Slabon, A.; Stievano, L.; Dronskowski, R. Carbodiimides as energy materials: which directions for a reasonable future? J. Chem. Soc., Dalton Trans. 2018, 47 (32), 10827-10832.

(53) López, F. A.; Rodríguez, O.; Alguacil, F. J.; García-Díaz, I.; Centeno, T. A.; García-Fierro, J. L.; González, C. Recovery of carbon fibres by the thermolysis and gasification of waste prepreg. *J. Anal. Appl. Pyrolysis* **2013**, *104*, 675–683.

(54) Śliwak, A.; Díez, N.; Miniach, E.; Gryglewicz, G. Nitrogencontaining chitosan-based carbon as an electrode material for highperformance supercapacitors. *J. Appl. Electrochem.* **2016**, *46* (6), 667– 677.

(55) Seredych, M.; Hulicova-Jurcakova, D.; Lu, G. Q.; Bandosz, T. J. Surface functional groups of carbons and the effects of their chemical character, density and accessibility to ions on electrochemical performance. *Carbon* **2008**, *46* (11), 1475–1488.

(56) Brousse, T.; Bélanger, D.; Chiba, K.; Egashira, M.; Favier, F.; Long, J.; Miller, J. R.; Morita, M.; Naoi, K.; Simon, P.; Sugimoto, W. 16 - Materials for Electrochemical Capacitors. *Springer Handbook of Electrochemical Energy*; Springer, 2017; pp 495–561.

(57) Cataldi, P.; Dussoni, S.; Ceseracciu, L.; Maggiali, M.; Natale, L.; Metta, G.; Athanassiou, A.; Bayer, I. S. Carbon Nanofiber versus Graphene-Based Stretchable Capacitive Touch Sensors for Artificial Electronic Skin. *Adv. Sci.* **2018**, *5* (2), 1700587.

(58) Scialdone, O.; Guarisco, C.; Grispo, S.; Angelo, A. D.; Galia, A. Investigation of electrode material – Redox couple systems for reverse electrodialysis processes. Part I: Iron redox couples. *J. Electroanal. Chem.* **2012**, *681*, 66–75.

(59) Manohar, A. K.; Kim, K. M.; Plichta, E.; Hendrickson, M.; Rawlings, S.; Narayanan, S. R. A High Efficiency Iron-Chloride Redox Flow Battery for Large-Scale Energy Storage. *J. Electrochem. Soc.* **2016**, *163* (1), A5118–A5125.

(60) Binninger, T.; Fabbri, E.; Patru, A.; Garganourakis, M.; Han, J.; Abbott, D. F.; Sereda, O.; Kötz, R.; Menzel, A.; Nachtegaal, M.; Schmidt, T. J. Electrochemical Flow-Cell Setup for In Situ X-ray Investigations: I. Cell for SAXS and XAS at Synchrotron Facilities. *J. Electrochem. Soc.* **2016**, *163* (10), H906–H912.

(61) Davi, M.; Keßler, D.; Slabon, A. Electrochemical oxidation of methanol and ethanol on two-dimensional self-assembled palladium nanocrystal arrays. *Thin Solid Films* **2016**, *615*, 221–225.

(62) Veerman, J.; Saakes, M.; Metz, S. J.; Harmsen, G. J. Reverse electrodialysis: evaluation of suitable electrode systems. *J. Appl. Electrochem.* **2010**, 40 (8), 1461–1474.

(63) Lee, J.-Y.; Kim, J.-H.; Lee, J.-H.; Kim, S.; Moon, S.-H. Morphologically Aligned Cation-Exchange Membranes by a Pulsed Electric Field for Reverse Electrodialysis. *Environ. Sci. Technol.* **2015**, 49 (14), 8872–8877.

(64) Tedesco, M.; Brauns, E.; Cipollina, A.; Micale, G.; Modica, P.; Russo, G.; Helsen, J. Reverse electrodialysis with saline waters and concentrated brines: A laboratory investigation towards technology scale-up. J. Membr. Sci. **2015**, 492, 9–20.

(65) Güler, E.; Elizen, R.; Vermaas, D. A.; Saakes, M.; Nijmeijer, K. Performance-determining membrane properties in reverse electrodialysis. *J. Membr. Sci.* **2013**, 446, 266–276.

(66) Abdullah Shah, S.; Cucchiara, R.; Vicari, F.; Cipollina, A.; Tamburini, A.; Micale, G. Energetic Valorisation of Saltworks Bitterns via Reverse Electrodialysis: A Laboratory Experimental Campaign. *Membranes* **2023**, *13* (3), 293.

(67) D'Angelo, A.; Tedesco, M.; Cipollina, A.; Galia, A.; Micale, G.; Scialdone, O. Reverse electrodialysis performed at pilot plant scale: Evaluation of redox processes and simultaneous generation of electric energy and treatment of wastewater. *Water Res.* **2017**, *125*, 123–131.

(68) You, X.; Ye, Q.; Cheng, P. The Dependence of Mass Transfer Coefficient on the Electrolyte Velocity in Carbon Felt Electrodes: Determination and Validation. *J. Electrochem. Soc.* **2017**, *164* (11), E3386–E3394.

(69) Tedesco, M.; Scalici, C.; Vaccari, D.; Cipollina, A.; Tamburini, A.; Micale, G. Performance of the first reverse electrodialysis pilot plant for power production from saline waters and concentrated brines. *J. Membr. Sci.* **2016**, *500*, 33–45.

(70) Luque Di Salvo, J.; Cosenza, A.; Tamburini, A.; Micale, G.; Cipollina, A. Long-run operation of a reverse electrodialysis system fed with wastewaters. *J. Environ. Manage.* **2018**, *217*, 871–887.

(71) Choudhary, R.; Bhakat, T.; Singh, R. K.; Ghubade, A.; Mandal, S.; Ghosh, A.; Rammohan, A.; Sharma, A.; Bhattacharya, S. Bilayer staggered herringbone micro-mixers with symmetric and asymmetric geometries. *Microfluid. Nanofluidics* **2011**, *10* (2), 271–286.

(72) Conway, B. E. Electrochemical Supercapacitors Scientific Fundamentals and Technological Applications; Springer, 1999.

(73) Stefanoni, M.; Zhang, Z.; Angst, U.; Elsener, B. The kinetic competition between transport and oxidation of ferrous ions governs precipitation of corrosion products in carbonated concrete. *RILEM Technol. Lett.* **2018**, *3*, 8–16.

(74) Veerman, J. Reverse electrodialysis design and optimization by modeling and experimentation: design and optimization by modeling and experimentation. Ph.D. Dissertation, University of Groningen, 2010. https://www.osti.gov/etdeweb/biblio/2141143web0 (accessed Aug 20, 2023).

(75) Pourbaix, M. Atlas of Electrochemical Equilibria in Aqueous Solutions; Pergamon: Oxford, 1966.

(76) Zhang, W.; Chen, X.; Wang, Y.; Wu, L.; Hu, Y. Experimental and Modeling of Conductivity for Electrolyte Solution Systems. *ACS omega* **2020**, *5* (35), 22465–22474.

(77) Jwa, E.; Jeong, N.; Nam, J.-Y.; Han, J.-I. Sustainable energy harvesting and on-site disinfection of natural seawater using reverse electrodialysis. *Water Res.* **2022**, *220*, 118681.

(78) Suda, F.; Matsuo, T.; Ushioda, D. Transient changes in the power output from the concentration difference cell (dialytic battery) between seawater and river water. *Energy* **2007**, 32 (3), 165–173.