Simultaneous Energy Storage and Seawater Desalination using Rechargeable Seawater Battery: Feasibility and Future Directions

Moon Son, Sanghun Park, Namhyeok Kim, Anne Therese Angeles, Youngsik Kim,* and Kyung Hwa Cho*

1. Introduction

There is great interest in securing alternative water resources at an affordable price, consequently accelerated by the imbalance in accessible water due to climate change.[1] In areas adjacent to seawater, freshwater is partially supplied through seawater desalination.[2–4] However, most conventional processes, which utilize either high temperature (evaporation; >8 kWh m\(^{-3}\)) or high applied pressure (membrane separation; >3 kWh m\(^{-3}\)), are known to be energy intensive.[3,4] Fortunately, a new concept introduced in the year 2018 has the potential in addressing high desalination energy requirements,[5] causing excitement in the water and energy societies. Seawater battery desalination (SWB-D) uses rechargeable seawater battery (SWB) to save energy used during seawater desalination. It is a multi-functional process as it simultaneously stores electricity induced by sodium ion movement while removing salt ions from seawater.[5–8] So far, the cost for unit desalinated water from SWB-D remains unexplored. This is primarily due to the lack of information from previous literatures regarding the energy prices of stand-alone SWBs. There has been a study that compared the energy price of an entire SWB unit with other energy storage systems (ESS) in a plant-scale,[9] but the price of a unit cell or the detailed energy price divided by cell compartment has not been reported yet. Therefore, this perspective discusses the feasibility of SWB-D system based on the unit cost per desalinated water ($ m\(^{-3}\)), which was calculated by the amount of produced water divided by the unit price of SWB-D. In addition, the future direction for seawater desalination research is also discussed. To accomplish these goals, the fundamental working mechanism of SWB will be first discussed as it works as an anode in SWB-D system.

2. Feasibility of Seawater Battery Desalination System

2.1. Seawater Battery: What It Is?

This perspective aims to provide comprehensive understandings and future directions of a new concept of simultaneous energy storage and seawater desalination using SWB. To understand this
concept, the working principles of SWB must be discussed first. Unlike lithium ion batteries (LIBs), SWB utilizes sodium ions instead of lithium ions as charge carriers (Figure 1a,b).[10,11] SWB is composed of three parts: 1) Open-cathode compartment for seawater exposure, 2) sodium super-conducting separator (ceramic based Na$_3$Zr$_2$Si$_2$PO$_12$; NASICON), and 3) closed-anode compartment.[11] Water on Earth is 97.5% seawater wherein 30.5% of its total salt is sodium.[12] Thus, when SWB is immersed into the seawater, the free and abundant sodium ions in the catholyte can migrate into the anode compartment during charging, storing it as sodium metal. Then, discharging converts the sodium metal back to sodium ions, releasing it into the seawater. Throughout this perspective, we assume that sodium metal anode is used because it is one of the most widely used anodes in SWB research. The continued research during the past years reveal that SWB is rechargeable (85% energy efficiency over 300 cycles) similar to other ESS technologies.[12] To simplify our discussion, we are going to discuss about this rechargeable SWB (invented in the year 2014).[14] otherwise noticed, although there were non-rechargeable SWBs (or primary SWBs), which was developed in the year 1943.[12]

It is worth noting here that SWB is different from sodium ion battery (SIB)[15] or desalination battery (DB).[16-18] SIBs have almost the same structure as LIBs but only use sodium ions and accordingly tuned materials.[15] DBs use sodium-intercalating electrodes (cathode and anode) and a salt solution flowing along the surface of the electrodes.[16-18] The water flowing channel in the DB is often divided by ion-exchange membranes in between two electrodes, and redox solution is used as the catholyte and the anolyte. Unlike SIB and DB, SWB system has a unique structure of an open-cathode compartment with a closed-anode compartment. Thus, SWB can be operated by immersing it directly into the seawater without using a redox chemical and its associated chambers. In addition, the use of sodium metal anode in SWB enabled a higher capacity compared to DB with sodium intercalation material as anode ($\approx 300$ mAh g$^{-1}$ for sodium metal$^{[19]}$ and $\approx 35$ mAh g$^{-1}$ for Na$_2$-$x$Mn$_5$O$_{10}$ electrode$^{[14]}$). In fact, there is another study of SWB where the configuration is similar to DB since ion intercalation electrodes were used. However, relatively low discharging voltage of $<1.2$ V was reported in that study,$^{[20]}$ which is significantly lower than that of typical output voltage of SWB ($>3.0$ V).$^{[12]}$ The reaction mechanisms for each component will not be discussed in this perspective but are available in previous literatures if further understanding is needed.$^{[11,12]}

2.2. Is Seawater Battery Competitive to Lithium-Ion Battery?

To rationalize the use of SWB-based system in seawater desalination, a brief comparison between SWB and LIB is necessary. For SWB systems, the charging process involves two redox reactions: oxygen evolution reaction on the cathode ($E^0 = 0.77$ V vs SHE) and sodium solidification on the anode ($E^0 = -2.71$ V vs SHE).$^{[12]}$ Thus, the overall cell reaction in SWB during charging requires $3.48$ V (vs Na/Na$^+$). Such high voltage of the SWB cell enabled the first coin-type SWB (SWB$_{coin}$) competitive to lead-acid battery (70 Wh L$^{-1}$ of SWB$_{coin}$ vs 80 Wh L$^{-1}$ of lead-acid battery) (Figure 2).$^{[21]}$ However, there is still a considerable difference in the energy density between the second generation of rectangular-type SWB (SWB$_{Rect}$) and conventional LIB (201 Wh L$^{-1}$ of SWB$_{Rect}$ vs 450 Wh L$^{-1}$ of LIB).$^{[10]}$ Although the theoretical maximum energy density of SWB is higher than that of LIB (3051 Wh L$^{-1}$ of SWB vs 1901 Wh L$^{-1}$ of LIB),$^{[10]}$ more investigations should be carefully carried out to determine the achievable energy density of SWB, and then a reasonable comparison with the corresponding LIB must be made in future research.

To our knowledge, only one discussion was reported about the energy and power costs of SWB compared to other battery systems.$^{[9]}$ According to that study, the energy cost for a plant-size
SWB (or full-size ESS system) is 187 $ kWh^{-1}. Although the energy cost is highly dependent on the size of the battery used in the calculation and the life cycle of the total system, the plant-size calculation of SWB was lower than that of LIB (996–2126 $ kWh^{-1}). The low energy cost determined from this study is most likely from the SWB’s use of seawater as the catholyte, and the absence of expensive lithium ion. However, the detailed cost analysis for each compartment at a unit cell level and its associated future research directions were not provided in that study. In contrast, a different study comparing SIB and LIB revealed that the cell material cost of SIB (101 $ kWh^{-1} for $\beta$-NaMnO$_2$) is slightly higher than that of LIB (88 $ kWh^{-1} for LiMn$_2$O$_4$). In terms of cathode material, changing lithium to sodium marginally decreased the total material cost by 3.8%. Conversely, switching from a lithium-specific anode to a sodium-specific anode increased the total cost owing to its lower energy density than a lithium-specific anode. For SIB, the cell material cost was 101 $ kWh^{-1}, while the total cost including additional items such as casings, battery management system, and battery thermal management unit was 287 $ kWh^{-1}. Based on the above-mentioned literatures, the significant discrepancy of costs calculated for different scales and materials implies that the compartment cost (or cell material cost) at a unit cell level must be provided for a fair comparison between SWB and LIB as an ESS system. In addition, by using the cost-related information of SWB for each compartment, we are extending this to SWB-D systems to determine the unit cost per desalinated water. We are hoping that our calculations and methods can serve as a reference for future SWB studies and their relevant expansion to other technologies.

2.3. Seawater Battery Desalination System: Energy Storage System during Desalination

The unique sodium adsorption property of SWB led to the development of its expansion systems such as SWB-D. The proposed SWB-D system can be divided into two parts: Three chambers for charging (desalination) and two chambers for discharging (salination) (Figure 3). Unlike SWB, there is a desalination compartment between the anode and cathode compartments for charging of the system (Figures 1c and 3). The desalination compartment is separated from the cathode compartment by an anion exchange membrane (AEM). Upon charging, sodium ions present in the seawater migrate to the anode compartment and solidify to sodium metal (typically when sodium metal anode is used). Meanwhile, its anion pair (primarily chloride ions) transfers to the cathode compartment through the AEM to maintain charge neutrality in the desalination compartment. The migrated chloride ions can be further removed (partially) through chlorine gas evolution ($2Cl^- \rightarrow Cl_2(g) + 2e^- \ E^0= -1.36 \text{ V vs SHE}$). Thus, SWB-D is a multi-functional system to store renewable energy during desalination because the partial amount of the energy used during desalination is directly stored into the SWB. The discharging part of the SWB-D system often shares the anode compartment with the charging part to utilize the energy stored during the charging of the system (Figure 3). Upon discharging of the SWB, the stored energy is released while the dissolved oxygen is reduced ($O_2 + H_2O + 2e^- \rightarrow HO_2^- + OH^- \ E^0= 0.21 \text{ V vs SHE}; O_2 + 2H_2O + 4e^- \rightarrow 4OH^- \ E^0= 0.77 \text{ V vs SHE}$). Consequently, sodium ions...
are migrated from the desalination compartment of the charging part to the cathode compartment of the discharging part (rightest part of Figure 3) during simultaneous operation of the system. Thus, this proposed concept can prevent the salination of the desalinated water during discharging. A proof-of-concept of this complete SWB-D system was proposed in the year 2018\cite{5} and was successfully demonstrated in the previous study.\cite{7}

It is worth mentioning here that nomenclatures of cathode and anode in SWB-D system are opposite to those in other electrochemical desalination technologies such as capacitive deionization and electrodialysis (ED). For example, sodium ions are attracted to the anode in SWB-D system, whereas those are adsorbed onto the cathode in other systems. This difference in nomenclature has arisen from the different purposes of the operation. The purpose of battery systems such as SWB-D is to provide electrons through the outer circuit connected to the cell. In contrast, other electrochemical cells focus more on the electrochemical reactions inside the cell.

2.4. Seawater Battery Desalination versus Other Seawater Desalination Technologies

To compare SWB-D with other seawater desalination technologies, a plot describing salt removal as a function of specific energy consumption (SEC) was created (Figure 4). Since, SWB-D is an electrochemical method, it was compared with three other electrochemical desalination technologies (membrane capacitive deionization, MCDI; flow-electrode capacitive deionization, FCDI; and ED). RO was also included in the plot as a reference since it is one of the dominant seawater desalination technologies in the market.

When RO was first commercialized back in the 1970s, its SEC was reported as 20 kWh m$^{-3}$ (orange trapezoid in Figure 4).\cite{21} The development of pressure exchangers and highly permeable membranes enabled to reduce this SEC to ≈3.5 kWh m$^{-3}$ starting approximately in the year 2010 (at 50% water recovery).\cite{24,25}

This number is close to the theoretical minimum energy requirement for seawater desalination (≈1.1 kWh m$^{-3}$ at 50% water recovery; Intersection of black and blue dash lines in Figure 4).\cite{26} Thus, RO has been recognized as one of the most promising seawater desalination technologies even though it is still energy intensive compared to wastewater treatment processes for clean water production. In contrast, the two representative capacitive deionization technologies, MCDI and FCDI, have shown relatively high SEC of 83.2 kWh m$^{-3}$ (MCDI)\cite{27} and
43.0 kWh m$^{-3}$ (FCDI)\cite{28} for seawater desalination. Although MCDI and FCDI have been extensively investigated for brackish water desalination,\cite{29-31} the application of these processes to seawater desalination has been limited primarily because their SECs proportionally increase with feed salinity, and the salinity of seawater is often $\approx 35,000$ ppm (total dissolved solids; TDS). For ED, several studies have been reported for seawater desalination, which showed SECs of 6.6 kWh m$^{-3}$ (99% salt removal),\cite{32} 15 kWh m$^{-3}$ (87% salt removal),\cite{33} 16.2 kWh m$^{-3}$ (98% salt removal),\cite{34} or 18.1 kWh m$^{-3}$ (98% salt removal).\cite{35} These values are distinctly higher than that of RO (3.5 kWh m$^{-3}$ for $\approx 100%$ salt removal). Note that the SEC of RO varies from 2.5 to 5 kWh m$^{-3}$ depending on the size of the plant, the number of stages (e.g., single pass), and feed water quality.\cite{36-38} An SEC of 3.5 kWh m$^{-3}$ was selected as a baseline since this is a common value that often appears in the references.\cite{36-38}

The first reported SEC for SWB-D is 53.9 kWh m$^{-3}$ (97.5% salt removal) in the year 2020,\cite{39} which is similar to MCDI and FCDI. According to literature, the energy consumption of SWB-D is almost proportional to the amount of salt removed.\cite{6,8} Thus, when 100% of salt removal is assumed, the SEC of SWB-D could proportionally increase to 71.9 kWh m$^{-3}$, not including the increase in salt solution resistance as salt removal reaches 100%. Note that although we ruled out the change of the solution resistance during operation to simplify our discussion, the practical SEC can further increase. However, one of the important aspects of SWB-D that can distinguish it from other desalination technologies is its energy storing ability during desalination. In practical application, $\approx 80%$ of energy recovery (or energy storing) is often reported for SWB.\cite{12} This means that 80% of the energy used for charging can be reused for discharging. In this case, the SEC of SWB-D can decrease to 10.8 kWh m$^{-3}$. When 90% of energy recovery is assumed, which is achievable by reducing the voltage gap during cycling,\cite{12,19} the energy consumption further decreases to 5.4 kWh m$^{-3}$. In other literatures, the use of intercalating material as a cathode or redox electrolyte solution as catholyte minimized the voltage gap of SWB.\cite{39-41} Thus, similar approaches can be adopted for SWB-D to maximize the energy recovery during desalination. In conclusion, in order for SWB-D to have similar energy consumption and salt removal rate as RO, it must have an energy recovery rate of 95% (3.6 kWh m$^{-3}$ at 100% salt removal; Open red star in Figure 4) if SWB-D can be operated solely without any pre- and post-treatment of seawater.

Since SWB-D system was newly proposed, insufficient information has been reported regarding the influence of feed water quality on the overall system performance. However, from the general engineering point of view, similar pre- and post-treatment of seawater could be required for a full-scale operation of SWB-D system. This is because untreated seawater contains several organics/inorganic matters and suspended solid/particles, which can easily decrease the overall performance of the system due to channel-clogging and contamination (also known as fouling) of the materials. Thus, for the SWB-D system, additional energy consumption for additional treatment such as intake and pre-treatment of seawater, and distribution of the produced clean water could be further accounted. Note that these are found to be 0.19 kWh m$^{-3}$ (intake), 0.39 kWh m$^{-3}$ (pre-treatment), and 0.18 kWh m$^{-3}$ (distribution) for RO system when the energy consumption was divided by the amount of produced water.\cite{16} To simplify our discussion, however, those additional energy consumptions were not accounted in this study.

Another factor that should be considered is the salt removal rate of SWB-D system. Although the maximum salt removal of SWB-D system remains unexplored primarily due to the voltage threshold of the system, the current design of SWB-D system can mostly remove Na$^+$ and Cl$^-$ ions. This is because NASICON allows only Na$^+$ ions to pass through and AEM has no specific selectivity toward anions. Thus, the maximum salt removal of the current SWB-D system is $\approx 85\%$, which is the NaCl portion in seawater.\cite{12} The remaining ions, primarily divalent sulfate (7.6%), magnesium (3.7%), and calcium (1.2%) should be further treated with other methods. In this regard, post-treatment using nanofiltration (NF), which is effective to remove divalent ions, can be considered. Thus, an energy consumption of 0.5 kWh m$^{-3}$ for NF post-treatment was further added (see the Experimental Section/Methods for details). The result showed that the energy consumption of this system (denotes SWB-D-NF) is competitive to RO when 96% of energy recovery is achieved (3.4 kWh m$^{-3}$). Thus, the energy recovery ratio of well above 90% should be targeted to render SWB-D system competitive to RO in terms of energy consumption. Alternatively, energy-free ion movement, such as diffusion between compartments, particularly between the desalination and cathode compartment, must be considered to reduce the energy used per ion removed. For example, if the ion transportation between the desalination and cathode compartment is promoted via diffusion, the desalination kinetics can be significantly improved without additional energy consumption, thereby lowering the overall SEC used for desalination. Therefore, future studies would need to focus on achieving SECS lower than RO in order for SWB-D to be competitive in the market of desalination.

2.5. Cost Breakdown of Seawater Battery Desalination and Seawater Battery

Similar with other processes, the price of the system dictates the price of the product. To estimate the price of clean water produced, the system price of SWB at a unit cell level was calculated. The energy normalized material cost (unit of $\text{kWh}^{-1}$) was calculated based on the assessable retail price (Figure 5) although material cost could be significantly reduced by bulk purchasing of chemicals. Thus, the material cost presented in this section aims to provide a simplified number for future reference, not to feature the exact number for the current energy price of SWB based systems.

The material cost for two different sizes of SWB (SWB$_{\text{coin}}$ and SWB$_{\text{Rec}}$) was first calculated. For SWB$_{\text{coin}}$, the overall material cost is 150.8 $\text{kWh}^{-1}$ in which the separator (NASICON) occupied 122 $\text{kWh}^{-1}$ followed by cathodic current collector (cathodic C. C.; 23 $\text{kWh}^{-1}$), anode (4 $\text{kWh}^{-1}$), and anolyte (2 $\text{kWh}^{-1}$). For SWB$_{\text{Rec}}$, which has a larger cathodic C. C. than SWB$_{\text{coin}}$ (2$\text{cm}^2$ for SWB$_{\text{coin}}$ and $\approx 68\text{cm}^2$ for SWB$_{\text{Rec}}$), the overall material cost is 216.4 $\text{kWh}^{-1}$. The material cost for the cathodic C. C. (108 $\text{kWh}^{-1}$) is dominant for SWB$_{\text{Rec}}$ followed by the separator (86 $\text{kWh}^{-1}$), anolyte (18 $\text{kWh}^{-1}$), and anode (4 $\text{kWh}^{-1}$). The material cost of LIB was reported in the range of $\approx$88 to $\approx$200 $\text{kWh}^{-1}$.\cite{12,42,43} Thus, the material cost for SWB$_{\text{Rec}}$ was calculated.
3. Future Directions for Practical Applications

The SWB-D process has been tested only in a steady-state (or batch) mode so far, so there was not much consideration for limiting factors such as the diffusion of ions and conductivity of influent solution. Thus, in order to fairly compare SWB-D technology with other seawater desalination technologies, the development of a continuous process such as a flow-cell must be investigated. When manufacturing a flow-cell, an architecture design that can maximize the contact area between the SWB and the desalination compartment is required. It is known that the time taken for desalination in SWB-D is proportional to the amount of current flowing through the system.\[^{4}\] For the current SWB-D system, it takes >30 h to desalinate (≥75%) ≈3.4 mL of seawater.\[^{6}\] Therefore, if a flow-cell, which maximizes the contact area between the anode and seawater, is proposed, the time required for desalination can be drastically reduced. In a batch mode system, solution resistance increases as the desalination progresses resulting to a rapid increase in cell voltage. In contrast, a continuous mode system continuously feeds the cell with high concentration influent, which prevents the increase in solution resistance and reduces the rate of increase in cell voltage. In this way, the amount of current applied per volume of seawater can be maximized in a flow-cell. For this purpose, a rectangular or pouch type developed for SWB system can be directly used for SWB-D system.\[^{21}\]

Since most RO processes have been operated under the cross-flow mode, the development of a flow-cell particularly for the desalination compartment could lead to a fair comparison between SWB-D and RO in future studies.

As reported in the previous literature,\[^{6,8}\] SWB-D system can be applied to hypersaline water treatment owing to a great ion deposition capacity of the SWB anode. However, the energy recovery efficiency in the actual SWB-D system remains unknown, whereas it is well known in SWB system, which does not possess a desalination compartment. For SWB-D, it is common to charge with SWB-D cells (composed of three compartments) and discharge with SWB cells (composed of two compartments) because a flow-cell (or continuous flow) for SWB-D has not yet been developed. In this regard, the energy recovery of 100% was assumed in the hybrid process,\[^{6}\] which is difficult to achieve in the practical operation. Therefore, the charging and discharging performance of SWB-D has to be further conducted in the same flow cell to confirm the practical energy efficiency of SWB-D system.

By multiplying SEC (kWh m\(^{-3}\)) and the material cost ($ kWh\(^{-1}\)) of SWB-D, the water production cost ($ m\(^{-3}\)) can be calculated, which varies depending on the lifespan (or long-term efficiency) of the system. For example, the water cost of SWB-D-NF could be 1.02 $ m\(^{-3}\) to desalinate seawater (0.6 m NaCl at 100% removal of salts), which is competitive to RO (0.60–1.20 $ m\(^{-3}\))\[^{47}\], when the cycling performance is maintained for 1000 cycles (at 96% energy recovery). Conversely, increasing the power density of the SWB-D can be an alternative approach that does not require energy efficiency considerations over massive cycles of operation. Using relatively inexpensive membranes instead of AEM could be an alternative approach to reduce water production cost because AEM cost is 50% of the total material cost in SWB-D. The use of inexpensive membranes might make it possible to divide the desalination compartment of SWB-D into several sub-compartments. This alternative approach could reduce energy consumption per
ion removal because more ions can be transported through ion exchange membranes with the same energy consumption. As proven by ED research,\textsuperscript{45} this approach can achieve low energy consumption for ion separation. Using an alternative membrane to AEM, which could facilitate ion diffusion from desalination to cathode compartment could also be a promising option to improve the desalination kinetics of SWB-D system. Since NASICON has been known as the resistance determining component in SWB-D system,\textsuperscript{8} developing a highly conductive NASICON could lead to lower ohmic and diffusion resistance, thereby using a higher applied current would be possible. Note that the NASICON used in current SWB-D studies has a chemical composition of Na\textsubscript{3}Zr\textsubscript{2}Si\textsubscript{2}PO\textsubscript{12}.\textsuperscript{8} For example, highly conductive NASICONS such as Na\textsubscript{3}\textsubscript{1.1}Si\textsubscript{2.3}Zr\textsubscript{1.55}P\textsubscript{0.7}O\textsubscript{11} or Na\textsubscript{3.4}Zr\textsubscript{2.4}Si\textsubscript{2}P\textsubscript{0.6}O\textsubscript{12}\textsuperscript{47} could be suitable candidates (i.e., the resistance of \(10^{-1} \text{ S cm}^{-1}\) for Na\textsubscript{3}Zr\textsubscript{2}Si\textsubscript{2}PO\textsubscript{12} and \(5 \times 10^{-1} \text{ S cm}^{-1}\) for Na\textsubscript{3.4}Zr\textsubscript{2.4}Si\textsubscript{2}P\textsubscript{0.6}O\textsubscript{12}) for SWB-D application. Another approach that can be considered to improve the desalination kinetics is the use of redox chemistry in cathode or catholyte as shown in the previous SWB study.\textsuperscript{46}

In this case, the sluggish kinetics of oxygen evolution and reduction reactions can be replaced with relatively fast redox reactions, thereby improving the desalination kinetics of SWB-D systems.

Designing a large-scale SWB-D cell with a minimized cost of cathodic C. C. would also be beneficial as the cathodic C. C. cost is \(\approx 50\%\) in SWB\textsubscript{Rect}. The anode compartment design will have a marginal impact on the overall material cost; however, it can render the system more compact, thereby minimizing the capital cost of any SWB-based system including SWB-D system.

Based on the aforementioned discussions, important concluding remarks can be made as follows: The material costs of AEM (50%) and separator (41%) are current hurdles for large-scale application of SWB-D. Energy recovery and cycling efficiency play crucial roles in determining the feasibility of SWB-D compared to other desalination technologies such as RO. When energy recovery of \(96\%\) and stable performance for 1000 cycles are achieved, an equipment cost of \(1.02 \text{ $ m}^{-3}\) can be expected, which is similar to RO (0.60–1.20 \text{ $ m}^{-3}\). Flow-cell (or continuous system) development for SWB-D is urgently required for comprehensive comparisons. Continuous flow could facilitate ion diffusion across the AEM, which could remove more ions without additional energy input. For SWB-D to compete with other seawater desalination processes, particularly RO, in addition to energy aspects, desalination kinetics must be significantly improved. For future studies, a more realistic cost analysis for large-scale SWB-D system can be done when material processing and casing costs are included.

4. Experimental Section

Specific Energy Consumption: Whenever seawater is used for calculation, the salt concentration was assumed as 0.6 m (\(\approx 35 \text{ g L}^{-1}\) NaCl).\textsuperscript{26} The salt removal of RO was assumed 100% because \(\approx 99.7\%\) of salt removal has been often reported.\textsuperscript{49–51} For the calculation of the SEC of SWB-D, it was assumed that salt removal is proportional to energy consumption.\textsuperscript{4} An energy consumption of 0.5 kWh m\textsuperscript{3} was assumed for NF in SWB-D-NF at a feed concentration of 5200 ppm (when SWB-D removed 85% of salt from 35 000 ppm TDS). Moreover, a linear relationship between energy consumption and feed TDS in NF was used.\textsuperscript{52} Note that the numbers used are an approximation and more research is needed to verify the maximum salt removal rate of SWB-D-NF using real seawater.

This number Theoretical minimum energy required for seawater desalination was calculated by Gibbs free energy of separation at 50% of water recovery.\textsuperscript{26,51}

Detailed Information for Cost Calculations: The components used in calculations for SWB\textsubscript{coin} were 0.8 g (NASICON), \(\approx 2 \text{ cm}^{2}\) (cathodic C. C.; carbon felt), 15 mL (anolyte; 1 m Biphenyl in diethylene glycol dimethyl ether), and \(\approx 68.3 \text{ cm}^{2}\) (anode; stainless steel mesh). For NASICON price, a mass-based element ratio was applied to each chemical needed to synthesize. Retail prices were used for the chemicals used during NASICON synthesis. The dimensions used in calculations for SWB\textsubscript{rect} were 23.6 g (NASICON), 396 cm\textsuperscript{2} (cathodic C. C.; carbon felt), 6 mL (anolyte; 1 m Biphenyl in diethylene glycol dimethyl ether), and \(\approx 68.3 \text{ cm}^{2}\) (anode; stainless steel mesh). For the material cost of SWB-D, which was calculated based on SWB\textsubscript{coin}, an AEM area of \(\approx 2 \text{ cm}^{2}\) was considered. The AEM price was calculated based on the minimum unit price (200 $ m\textsuperscript{3}) multiplied by the area used (\(\approx 2 \text{ cm}^{2}\)).\textsuperscript{34}

For LIB calculations using retail prices, a coin cell (2325 coin cell; diame\textsuperscript{ter} \(\approx 0.905 \text{ in};\) depth \(\approx 0.098 \text{ in})\) with similar dimensions to SWB-coin was used. Accessible retail prices were used for the chemicals needed for the synthesis of each cell component. The cathode consists of LiNi\textsubscript{0.6}Mn\textsubscript{0.2}Co\textsubscript{0.2}O\textsubscript{2} (also known as NMC622), carbon black, polyvinylidene fluoride, and \(n\)-methyl-2-pyridolone (NMP). Aluminum foil was used as the cathodic C. C. and the electrolyte was a mixture of LiPF\textsubscript{6}, ethylene carbonate, and diethyl carbonate. Polyethylene membrane was used as the separator. The anode consisted of graphite, carboxymethyl cellulose and styrene-butadiene rubber, carbon black, and NMP.

Acknowledgements

This work was supported by the National Research Foundation of Korea (NRF) grant funded by the Korean government (MSIT) (grant number 2020R1A4A1019568).

Conflict of Interest

The authors declare no conflict of interest.

Keywords

cost analysis, desalination, energy storage systems, seawater batteries
Moon Son is a Research Assistant Professor at the School of Urban and Environmental Engineering at Ulsan National Institute of Science and Technology (UNIST), South Korea. He received his Ph.D. (2017, advisor: Prof. Heechul Choi) in environmental engineering from Gwangju Institute of Science and Technology (GIST), South Korea. After that, he joined Prof. Bruce E. Logan’s group at the Pennsylvania State University as a post-doctoral fellow. His current research is focused on renewable energy production, low-energy water treatment, and water desalination by electrochemical- or membrane-based processes.
Youngsik Kim is a Professor at School of Energy and Chemical Engineering at Ulsan National Institute of Science & Technology (UNIST). He received his Ph.D. (2006, advisor: Prof. Steve W. Martin) in Materials Science and Engineering from Iowa State University. Later, he joined Prof. John B. Goodenough’s group at The University of Texas at Austin as a post-doctoral fellow. He conducted research in areas including lithium- and sodium-ion batteries. Particularly, he is an inventor of the “rechargeable seawater battery,” which is developed as an alternative option for grid-scale energy storage. He is also the CEO of energy solution company, 4TOONE Corporation.

Kyung Hwa Cho is an Associate Professor at the School of Energy and Chemical Engineering at Ulsan National Institute of Science and Technology (UNIST), South Korea. He received his Ph.D. (2010) in environmental engineering from Gwangju Institute of Science and Technology (GIST), South Korea. After that, he was appointed as a post-doctoral fellow at the University of Michigan. His current research is focused on water-environmental informatics using artificial intelligence algorithms and modeling approaches. He is also the principal investigator of the seawater battery research team supported by the Korean government (Basic Research Laboratory).