Membrane-Free Battery for Harvesting Low-Grade Thermal Energy

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ABSTRACT: Efficient and low-cost systems are desired to harvest the tremendous amount of energy stored in low-grade heat sources (<100 °C). An attractive approach is the thermally regenerative electrochemical cycle (TREC), which uses the dependence of electrode potential on temperature to construct a thermodynamic cycle for direct heat-to-electricity conversion. By varying the temperature, an electrochemical cell is charged at a lower voltage than discharged; thus, thermal energy is converted to electricity. Recently, a Prussian blue analog-based system with high efficiency has been demonstrated. However, the use of an ion-selective membrane in this system raises concerns about the overall cost, which is crucial for waste heat harvesting. Here, we report on a new membrane-free battery with a nickel hexacyanoferrate (NiHCF) cathode and a silver/silver chloride anode. The system has a temperature coefficient of −0.74 mV K⁻¹. When the battery is discharged at 15 °C and recharged at 55 °C, thermal-to-electricity conversion efficiencies of 2.6% and 3.5% are achieved with assumed heat recuperation of 50% and 70%, respectively. This work opens new opportunities for using membrane-free electrochemical systems to harvest waste heat.

A vast amount of low-grade heat (<100 °C) exists in industrial processes, the environment, biological entities, and solar-thermal and geothermal energy.¹⁻³ Conversion of this low-grade heat to electricity is difficult due to the distributed nature of these heat sources and the low temperature differential. Different technologies, such as solid-state thermoelectric energy conversion⁴⁻⁷ and organic Rankine cycles,¹⁻⁸ are being actively investigated but face their own challenges in energy conversion efficiency, cost, and system complexity. Thermally regenerative electrochemical cycle (TREC) is an alternative approach based on the temperature dependence of cell voltage of electrochemical systems.⁹⁻¹² For a full cell reaction A + B → C + D (discharge), the temperature coefficient α is defined as

\[ \alpha = \frac{\partial E}{\partial T} = -\frac{1}{nF} \frac{\partial \Delta G}{\partial T} = \frac{\Delta S}{nF} \]

where \( E \) is the full cell voltage, \( T \) is the temperature, \( n \) is the number of electrons transferred in the reaction, \( F \) is Faraday’s constant. \( \Delta G \) and \( \Delta S \) are the change of partial molar Gibbs free energy and partial molar entropy in the full cell reaction.¹²⁻¹⁴ If \( \alpha \) is negative, net electricity can be produced by discharging the cell at low temperature \( T_L \) and recharging at high temperature \( T_H \) with lower voltage (Figure 1a). If \( \alpha \) is positive, electricity is produced by discharging at \( T_H \) and recharging at \( T_L \). In both cases, the work generated originates from heat absorbed at \( T_H \) indicated as entropy increase in the temperature–entropy \( (T–S) \) diagram (Figure 1b). The \( T–S \) diagram clearly shows that TREC is an Ericsson cycle with isothermal charge/discharge processes and isobaric heating/cooling processes. The corresponding heat-to-electricity conversion efficiency \( (\eta) \) of TREC can be expressed as

\[ \eta = \frac{W}{Q_H+Q_{HR}} = \frac{\Delta S \Delta T - E_{\text{loss}}}{T_H \Delta S + (1 - \eta_{HR}) C_p \Delta T} \]

where \( W \) is the difference between discharge and charge energy in a cycle. \( Q_H \) is the heat absorbed at \( T_H \). \( Q_{HR} \) is the extra energy needed to heat the cell up. \( E_{\text{loss}} \) is the energy loss due to internal resistance of the cell. \( \Delta S \) is the entropy change in reaction. \( \Delta T = T_H - T_L \). \( C_p \) is the heat capacity of the cell. \( \eta_{HR} \) is the heat recuperation efficiency, indicating how much energy rejected in the cooling process can be used for the heating process. Theoretically, \( \eta_{HR} \) can reach 100%, and 50—70% has...
been proven to be reasonable with experiments.9 From eq 2, the theoretical efficiency of TREC is the Carnot limit, whereas practical efficiency is mainly limited by the heat capacity of materials, the heat recuperation efficiency,15 and the internal resistance of the cell. This is in contrary to thermoelectric devices, which are mainly limited by transport properties of phonons and electrons.

The concept of TREC was developed a few decades ago. It focused on high temperature applications (500−1500 °C) and showed efficiencies up to 40−50% of the Carnot limit. However, low temperature TREC did not received as much attention since electrode materials with low polarization and high charge capacity at low temperature were limited.16 There was little data presented on charge/discharge voltage profiles and efficiencies. Recently, a new thermally regenerative electrochemical cycle (TREC) based on a copper hexacyanoferrate (CuHCF) cathode and a Cu/Cu2+ anode was demonstrated for harvesting low-grade heat.9 The low polarization of electrodes, moderate temperature coefficient, high charge capacity, and low heat capacity led to a high efficiency. An efficiency of 5.7% was demonstrated when the cell was operated between 10 and 60 °C, assuming a heat recuperation efficiency of 50%.9

One potential issue of the CuHCF//Cu2+/Cu system is the use of an ion-selective membrane to allow NO3− anion passing through but not Cu2+ cations to avoid side reaction between CuHCF and Cu2+ (Figure 1c). Ion-selective membranes are currently expensive and difficult to completely block penetration of Cu2+ in long-term operation. Finding membrane-free systems to lower the cost and facilitate long-term operation would make TREC systems more practical. To address this issue, we apply a criterion that any soluble chemical species in electrolyte should not induce adverse side reactions other than the desired two half-cell reactions. In this paper, a membraneless electrochemical system with a nickel hexacyanoferrate (NiHCF, KNiIIFeIII(CN)6) cathode and a silver/silver chloride anode is demonstrated, where no adverse side reaction is introduced due to solutes in electrolyte (Figure 1d). The reactions of the two half cells are

\[
\text{Cathode: } \text{KNi}^{II}\text{Fe}^{III}(\text{CN})_6 + \text{K}^+ + e^- \rightarrow \text{K}_2\text{Ni}^{II}\text{Fe}^{II}(\text{CN})_6
\]

\[
\text{Anode: } \text{AgCl} + e^- \rightarrow \text{Ag} + \text{Cl}^-
\]

In this system, ions involved in each electrode do not have side reactions with each other, so the ion-selective membrane is unnecessary and can be replaced by an inexpensive porous separator. The full cell has a temperature coefficient of −0.74 ± 0.05 mV K−1 and reaches thermal-to-electricity conversion efficiencies of 2.6% and 3.5% when cycled between 15 and 55 °C with assumed heat recuperation efficiency of 50% and 70%, respectively.

NiHCF nanoparticles were synthesized using a simple solution approach by dropping 50 mM Ni(NO3)2 aqueous solution into 25 mM K3Fe(CN)6 aqueous solution at 50 °C. The average size of as-synthesized particles is ∼50 nm (Supporting Information Figure S1). The small particle size enhances surface area for reaction and reduces distance for ionic transport in solid, leading to fast kinetics and lower overpotential. The porous Ag/AgCl electrode was made by charging Ag film in KCl aqueous solution. Ag/AgCl has an oxidation/reduction voltage gap due to a combination of

Figure 1. Schematic of thermally regenerative electrochemical cycle (TREC). (a) The voltage−capacity plot of a TREC. Net energy is generated because the discharge voltage is higher than charge voltage. The case with negative α is presented. (b) The corresponding temperature−entropy (T−S) plot. (c) A TREC with an ion-selective membrane to block certain ions to avoid side reactions. (d) The membrane-free NiHCF/Ag/AgCl system with no unwanted reaction between electrodes and ions.
considerable nucleation barrier and AgCl self-resistance.\textsuperscript{19} This issue is tackled by using a porous electrode with high surface area. The electrolyte was 3 M KCl with 0.2 M Ni(NO\textsubscript{3})\textsubscript{2} aqueous solution. Ni\textsuperscript{2+} was used to stabilize NiHCF at high temperature based on the common ion effect.\textsuperscript{20} The pH of the electrolyte was tuned to 2 by HNO\textsubscript{3} to optimize the performance of NiHCF.\textsuperscript{17} A silver wire with preoxidized AgCl was used as the reference electrode. Glassfiber filter was used as the separator to prevent battery shorting. The NiHCF and AgCl electrodes were assembled in a pouch cell configuration (Supporting Information Figure S2). Details on electrode preparation and assembly of the pouch cell are described in the Supporting Information.

Temperature-dependent electrochemical characteristics were measured with a homemade temperature cycler (Figure 2a and Supporting Information Figure S3). The thin pouch cell was sandwiched between two thermoelectric plates and thermocouples were attached to the surface of the pouch cell. Thermal paste was applied to all interfaces to ensure good thermal contact. Temperatures of thermocouples were acquired by a data acquisition (DAQ) board and controlled with fluctuations less than 0.1 °C. This compact design allows the temperature to be switched in less than 3 min so that other effects, such as self-discharge and dissolved oxygen, can be minimized. An example of measuring electrode voltage at different temperatures is shown in Supporting Information Figure S4. Each temperature step lasted for 8 min and the voltage became steady after 3 min in each step, indicating that the system quickly reached equilibrium and there was no obvious effect due to self-discharge.

The effect of temperature on the full cell voltage is investigated first (Figure 2b–d). Figure 2b shows the dependence of full cell voltage on temperature at different states of charge (SOC) for the 3 M KCl cell. The number on the right side is in the unit of milliamperes hours per gram. The fully discharged state corresponds to 0 mAh g\textsuperscript{−1}. For all measurements, the full cell voltage is linearly related to the temperature in the range of 15 to 55 °C, indicating that α is a constant in the temperature range. α of the full cell at different SOC with [KCl] from 1 to 4 M are presented in Figure 2c. For all concentrations, α shows an inverse bell shape against SOC. α is flat in the middle of the voltage curve (10–50 mAh g\textsuperscript{−1}), but its absolute value becomes smaller when the system approaches a fully charged or a fully discharged state. Moreover, lower [KCl] always leads to a more negative temperature coefficient for all SOC, which is likely a result of changes in K\textsuperscript{+} and Cl\textsuperscript{−} activity. For instance, a plot of α at 50% SOC against [KCl] shows a trend consistent with the derivation of α from Nernst equation (Figure 2c)

\begin{equation}
E = E_0 + \frac{RT}{F} \ln\left(\frac{[K^+]_0}{[Cl^-]_0}\right)
\end{equation}

\begin{equation}
\alpha = \alpha_0 + \frac{R}{F} \ln\left(\frac{[K^+]_0}{[Cl^-]_0}\right)
\end{equation}

where \(E\) is the electrode potential, \(R\) is the ideal gas constant, and \(F\) is the Faraday constant (96485 C mol\textsuperscript{−1}). \(E_0\) and \(\alpha_0\) are the electrode potential and temperature coefficient with unit activity of ions for a certain SOC. The activities of solid phases are assumed to be 1 so they are not shown in the equation. The activity coefficients of ions are assumed to be 1 so that activities of ions are replaced by concentration. The deviation from Nernst equation may arise from the activity coefficient and influence of Ni\textsuperscript{2+}, which can also be inserted into NiHCF.\textsuperscript{20} The temperature coefficient mainly comes from the half cell of NiHCF, as previous studies show that the temperature coefficient of Ag/AgCl is 0.22–0.26 mV K\textsuperscript{−1} with 1 M KCl\textsuperscript{12} and 0.12 mV K\textsuperscript{−1} with 4 M KCl.\textsuperscript{9} The measured temperature coefficient is slightly more negative than CuHCF,\textsuperscript{7} which is the reason why we use NiHCF instead of CuHCF. In our measurement, although the reference electrode and anode are the same, a small difference in electrode potential is observed (∼0.04 mV/K). The origin is unclear and may be related to the preparation process of reference electrode, which exposed the electrode in air and light and may result in a difference in chemical composition. In order not to overestimate α and energy conversion efficiency, α of the full cell is calculated based on the potential of NiHCF cathode and Ag/AgCl anode and its absolute value is less than that between the cathode and the reference electrode.

The dependence of α on [KCl] leads to a trade-off between voltage gap (\(\Delta T\)) and heat capacity. Because K\textsuperscript{+} and Cl\textsuperscript{−} are stored in the electrolyte, higher [KCl] indicates a smaller amount of KCl electrolyte is required and, thus, less energy is needed to heat the system up. However, it also reduces the absolute value of α and the voltage gap between discharge and charge (\(\alpha \Delta T\)). After estimating the conversion efficiency at different concentrations, 3 M was chosen as the optimal concentration and used in following tests (Supporting Information Table S1).

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Temperature coefficient of the NiHCF/Ag/AgCl system. (a) A schematic of the experimental setup for measuring temperature coefficient. (b) The temperature-dependent voltage of a full cell at different states of charge (SOC). The number on the right indicates capacity charged in the unit of mAh g\textsuperscript{−1}. The point at 0 mAh g\textsuperscript{−1} refers to the fully discharged state. The electrolyte is 3 M KCl. (c) The temperature coefficient at different SOC and concentrations of KCl electrolyte. (d) The dependence of temperature coefficient on the concentration of KCl solution at 50% SOC. The red curve is the fitting curve based on eq 3. For all measurements, 0.2 M Ni(NO\textsubscript{3})\textsubscript{2} was added to stabilized NiHCF and HNO\textsubscript{3} was used to tune the pH of the solution to 2.}
\end{figure}

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\textsuperscript{6580}
Figure 3. Thermal cycle of a NiHCF/Ag/AgCl/3 M KCl full cell. (a) Voltage vs time plot showing the charge and discharge curves of both NiHCF cathode (blue) and Ag/AgCl anode (black). The reference electrode is the Ag/AgCl electrode in the pouch. The current rate is 40 mA g\(^{-1}\). (b) and (c) Voltage vs specific capacity plot at (b) 40 and (c) 20 mA g\(^{-1}\). The dashed lines at the inner side of voltage profiles of NiHCF in (b) and (c) are the full cell voltage. Magnified voltage profiles of electrodes and full cells are shown in Supporting Information Figure S5. All currents are based on the mass of NiHCF.

Table 1. Calculation of Conversion Efficiency at Different Current Rates\(^d\)

<table>
<thead>
<tr>
<th>current rate</th>
<th>(V_{dis}) (mV)</th>
<th>(V_{ch}) (mV)</th>
<th>(Q_{dis}) (mAh g(^{-1}))</th>
<th>CE (%)</th>
<th>(W) (mWh g(^{-1}))</th>
<th>(Q_{ch}) (mAh g(^{-1}))</th>
<th>(C_p\Delta T) (mWh g(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>1C</td>
<td>561.9</td>
<td>546.4</td>
<td>35.4</td>
<td>99.72</td>
<td>0.50</td>
<td>8.62</td>
<td>31.6</td>
</tr>
<tr>
<td>C/2</td>
<td>565.7</td>
<td>545.8</td>
<td>36.0</td>
<td>99.72</td>
<td>0.65</td>
<td>8.76</td>
<td>32.1</td>
</tr>
</tbody>
</table>

\(^d\)All symbols are the same as those in eq 2, 4, and 5.

Figure 3a shows the thermal cycle of a NiHCF/Ag/AgCl full cell with 3 M KCl/0.2 Ni(NO\(_3\))\(_2\) electrolyte. The temperature is well controlled with a fluctuation less than 0.1 °C. At the end of each discharge or charge, the cell was rested for 3 min to allow the temperature to change and the system to reach equilibrium. Voltage vs capacity at currents of 1C (40 mA g\(^{-1}\)) and C/2 (20 mA g\(^{-1}\)) are presented as Figure 3b and c, respectively. All electrode potentials discussed below are versus Ag/AgCl reference electrode exposed to the same electrolyte in the pouch cell. The current rate and specific capacity are based on the mass of NiHCF. At both 1C and C/2 rates, the battery was heated up to 55 °C and charged to 640.0 mV. Then it was cooled down to 15 °C, which increased the open circuit voltage (OCV) to 660.1 mV for 1C and 661.6 mV for C/2. Next, the cell was discharged to 485 mV at 15 °C and then heated up to 55 °C again. The electricity produced in one cycle (W) normalized to the mass of NiHCF can be written as

\[
W = Q_{dis}V_{dis} - Q_{ch}V_{ch} = Q_{dis}\left(V_{dis} - \frac{V_{ch}}{CE}\right)
\]

where \(Q\) and \(V\) are the specific capacity normalized to the mass of NiHCF and average full cell voltage, respectively. The subscripts "dis" and "ch" indicate discharge and charge, respectively. CE is the Coulombic efficiency, which is defined as \(Q_{dis}/Q_{dis}\). From eq 4 we can see that in addition to large specific discharge capacity and voltage gap between discharge and charge, high Coulombic efficiency is also important to achieve high energy output and conversion efficiency. \(V_{dis} - V_{ch}\) and \((V_{dis} - V_{ch}/CE)\) are defined as the apparent and effective voltage gap, respectively, as the latter one directly determines the energy difference between discharge and charge.

At 1C rate (Figure 3b), the average discharge voltages of NiHCF and Ag/AgCl were 566.27 and 4.37 mV, respectively, and the capacity is 35.4 mAh g\(^{-1}\) based on the mass of NiHCF. The average charge voltages of NiHCF and Ag/AgCl were 542.42 and -3.93 mV, respectively, and the capacity is 35.5 mAh g\(^{-1}\). As a result, the apparent and effective voltage gaps are 15.8 and 14.2 mV, respectively. The total specific discharge and charge energy are 19.90 mWh g\(^{-1}\) and 19.40 mWh g\(^{-1}\), respectively; thus 0.50 mWh g\(^{-1}\) of heat energy was converted to electricity. Based on eq 2 and procedures discussed in our previous publication,\(^9\) The heat-to-electricity conversion efficiency (\(\eta\)) was calculated as

\[
\eta = \frac{W}{T_i\Delta S + Q_{HR}} = \frac{W_{discharge} - W_{charge}}{\int dtT_iQ_{C} + (1 - \eta_{HR})C_p\Delta T}
\]

where \(W\) is the difference between discharge and charge energy in a cycle. \(Q\) is the discharge capacity at \(T_{HR}\) \(\eta_{HR}\) is the heat recuperation efficiency, and 50–70% are reasonable.\(^9\) Other parameters are defined as same as those in eq 2. As \(W\) is calculated based on experimentally measured voltage profiles (Figure 3b and c), changes in concentration of K\(^+\) and Cl\(^-\) are taken into account. In the process described above, \(Q_{HR}\Delta T = 8.617 mWh g^{-1}\) with an \(\alpha\) of -0.74 mV K\(^{-1}\). The total heat capacity of electrolytes and electrodes are 2.84 J g\(^{-1}\) K\(^{-1}\) and \(C_p\Delta T = 31.6 mWh g^{-1}\) for cycles between 15 and 55 °C. Based on these values, \(\eta\) reaches 1.3%, 2.1%, and 2.8% for \(\eta_{HR}\) of 0%, 50%, and 70%. The calculation above is also presented as Table 1 and the details are described in the Supporting Information.

Similarly, the voltage curves at C/2 rate (20 mA g\(^{-1}\)) tested in the same voltage range are plotted as Figure 3c. For the full cell, the average discharge voltage increases to 565.7 mV and the average charge voltage decreases to 545.8 mV, as lower current leads to smaller overpotential. The specific capacity also increases to 36.0 and 36.1 mAh g\(^{-1}\) for discharge and charge, respectively. Consequently, the discharge and charge energy are 20.35 mWh g\(^{-1}\) and 19.71 mWh g\(^{-1}\), respectively. The energy converted to electricity reaches 0.65 mWh g\(^{-1}\), 29% higher than that at 1C rate, and the corresponding \(\eta_{HR}\) are 1.6%, 2.6%, and 3.5% for \(\eta_{HR}\) of 0%, 50%, and 70%.

For long-term operation, cycle life of TREC is crucial. The specific capacity, Coulombic efficiency, average charge/
discharge voltage, and thermal-to-electricity efficiency are plotted against cycle number as Figure 4a, b, and c in sequence. The cell was cycled at 1C rate for the first 35 cycles, then at C/2 for 50 cycles. The capacity fading rate is on average 0.10% and 0.18% per cycle at 1C and C/2, respectively. The higher capacity fading at C/2 is likely due to a longer operation time at 55 °C. The Coulombic efficiency is 99.2% at the beginning but quickly increases to ∼99.5–99.7% after five cycles. The average charge/discharge voltage shows a steady increasing trend of ∼0.1 mV per cycle (Figure 4b). The origin of this is not yet clear. Figure 4b also shows that the apparent voltage gap between charge and discharge (\( V_{\text{aq}} - \bar{V}_{\text{aq}} \)) is about 4 mV higher at C/2 compared to 1C rate, as a result of lower overpotential at smaller current. Moreover, the effective voltage gap (\( V_{\text{aq}} - \bar{V}_{\text{aq}}/CE \)) is obviously lower than the apparent voltage gap (\( V_{\text{aq}} - \bar{V}_{\text{aq}} \)) due to non-100% Coulombic efficiency. The difference is ∼5 mV at the beginning and decreases to ∼2 mV as the Coulombic efficiency gradually increases and stabilizes around 99.7%. The absolute conversion efficiency (\( \eta \)) is a synergistic result of the three factors above based on eq 4 (Figure 4c). At 70% heat recuperation, \( \eta \) is 2.2% in the first cycle due to low Coulombic efficiency and gradually increases to 2.9% after 30 cycles. The following cycles at C/2 shows \( \eta \) of 3.5% at the beginning and it decreases slowly to 2.9% after 50 cycles. This is because CE is steady in this part and the major fading factor is the decreasing capacity and apparent voltage gap. The fading rate is much smaller than our previous work4 as evaporation is fully eliminated by employing pouch cell configuration. In addition, it should be noticed that AgCl has a noticeable solubility in KCl solution, which is ∼1 mM in 3 M KCl solution at room temperature.21 This may lead to degradation of the anode and accumulation of Ag at cathode. The effect of soluble Ag ions on long-term cycling needs further investigation.

As thermoelectrics is a major research focus for waste heat harvesting,4,7 it would be useful to compare the current work and thermoelectrics (TE). We would like to first point out that there are many differences between these two approaches (TE vs TREC). TE is based on transport characteristics, such as thermal and electrical conductivity, whereas TREC relies on thermodynamic properties, such as heat capacity and temperature coefficient.8 Given the low temperature differential for low-grade heat, isothermal operation may be easier than temperature gradient based technologies. However, currently the typical cycle time of TREC is in the order of 1 h, leading to much lower power (∼1 W g⁻¹) than TE, as ionic transport and electrochemical reaction is slower than electronic transport. TREC also faces challenges in long-term operation, which may requires thousands of cycles at elevated temperature. With these differences in mind, a conversion of the efficiencies above to equivalent figure of merit ZT values of thermoelectrics helps evaluate the performance of TREC. At a heat recuperation efficiency of 70%, efficiency achieved at 1C and C/2 are 2.8 and 3.5% for temperature cycle between 15 and 55 °C. A TE device needs to reach ZT of 1.4 and 2.1 to achieve the same efficiency for the same high and low temperatures. If heat recuperation efficiency of 50% is assumed, the corresponding effective ZTs are 0.94 and 1.3 for efficiencies at 1C (2.1%) and C/2 (2.6%), respectively. In contrast, state-of-the-art TE materials have a ZT of 1–1.5 for temperature below 100 °C.22 Again, we want to emphasize that there are remarkable differences between the two approaches and readers should keep this in mind to understand the strengths and limits of both approaches.

A possible concern of the NiHCF/AgCl/Ag system is the cost of Ag. An estimation based on the price of Ag and Nafion ion-selective membrane shows that even with a high mass loading of Ag (∼5 mg cm⁻²), the cost is still less than 20% of Nafion membranes. When power density is considered, the cost of Ag electrode per watt is slightly less than half of membrane used in CuHCF/Cu²⁺/Cu system reported before9 (see Supporting Information for details). Searching for inexpensive electrodes to replace Ag can further reduce the cost.

In summary, a membrane-free electrochemical system with nickel hexacyanoferrate (NiHCF, KNi IIFeIII(CN)₆) cathode and Ag/AgCl anode is demonstrated to convert low-grade heat to electricity. As ions involved in each electrode do not interfere with the opposite electrode, expensive ion-selective membranes are not needed in this system. The system shows a thermo-to-electricity conversion efficiency of 3.5% under 70% heat recuperation when it is cycled between 15 and 55 °C. The system also shows adequate cycle life compared to previous results. We believe that further optimization and searching for new systems will lead to new development and possibly practical deployment of TREC.

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**Figure 4.** Cycling performance of NiHCF/Ag/AgCl system for TREC. (a) Discharge/charge capacity and Coulombic efficiency vs cycle life. (b) Average full cell charge/discharge voltage and their difference vs cycle life. (c) The thermal-to-electricity conversion efficiency vs cycle life.
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REFERENCES

(2) Gur, I.; Sawyer, K.; Prasher, R. Science 2012, 335 (6075), 1454−1455.
(6) Snyder, G. J.; Toberer, E. S. Nat. Mater. 2008, 7 (2), 105−114.