A molten battery consisting of Li metal anode, AlCl₃-LiCl cathode and solid electrolyte

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A B S T R A C T
High performance rechargeable batteries are urgently needed to address the demands of grid-scale stationary energy storage. High temperature battery systems, such as Na-S battery, Na-NiCl₂ battery (ZEBRA battery) and liquid metal electrode (LME) battery, exhibit advantages like high power density and high cyclic stability, but also suffer from the high operating temperature. We recently invented new concept of molten lithium metal batteries, consisting of liquid lithium anodes, alloy (Sn, Bi, Pb) liquid cathodes and lithium ion conductor as solid electrolytes. Here we demonstrate a molten metal chloride battery that operates at a relatively low temperature of 210 °C. The battery has been designed to include molten (AlCl₃-LiCl) cathode, solid electrolyte (garnet-type Li₆.4La₃Ta₀.₆Zr₁.₄O₁₂ (LLZTO) ceramic tube) and molten lithium anode. The assembled AlCl₃-LiCl||LLZTO||Li full cell has an average discharge voltage of 1.55 V and energy efficiency of 83% and has been successfully cycled for 100 cycles (800 h) without capacity fade. The theoretical specific energy of cell is 350 Wh/kg and the estimated cost is $11.6/kWh based on the weight of electrode materials. Considering the high performance, high safety, low operating temperature and low cost of raw materials, our new type of molten-electrode battery system opens up new opportunities for stationary energy storage.

Large-scale electrochemical energy storage system is critical for the renewable energy and smart grid technologies [1–3]. In particular, rechargeable batteries with low cost, long lifespan, good safety and high power density are required for stationary energy storage [4–6]. Many types of batteries technologies are being developed, examples including traditional lead-acid batteries [7], Li-ion batteries [8–10], Al-ion batteries [11,12], and flow batteries [13,14]. However, such room-temperature batteries suffer from high cost or limited lifespan, which seriously restricts their applications in stationary energy storage [15,16]. Recently, high-temperature batteries, including Na-S battery [17] and liquid metal electrode (LME) battery [18], have become an area of focus since they potentially meet the performance requirements for the smart electric grid. However, the exceedingly high operating temperature (>300 °C), which results in safety hazards and high maintenance cost, still remains as serious problem. The ZEBRA battery (Na-NiCl₂) can be operated at a relatively low temperature (250 °C), however, the high cost of Ni cathode severely limits its practical applications [19,20]. Therefore, a new molten rechargeable battery for large-scale energy storage, which can be operated at a relative low temperature with an acceptable cost, is still challenging.

Recently, our team developed a new-type LME battery using LLZTO ceramic tube as solid electrolyte, which can operate at a relative low temperature of 240 °C [21]. In the first demonstration, we show a battery chemistry system consisting of molten Bi-Sn-Pb alloying cathodes and molten Li metal anodes with theoretical specific energy limited to 230 Wh/kg. Herein, we designed a new high temperature battery chemistry with much higher theoretical specific energy and lower cost, which consists of AlCl₃-LiCl molten salt cathode, solid electrolyte tube (garnet-type Li₆.4La₃Ta₀.₆Zr₁.₄O₁₂ (LLZTO) ceramic tube) and liquid lithium anode. The AlCl₃-LiCl||LLZTO||Li full cell delivered a high cyclic performance and an acceptable power/energy density with a relative low cost. Besides, the relative low operating temperature and low...
flammability of the cathode materials ensure the high safety of the full cell. Therefore, the new-designed AlCl₃-LiCl||LLZTO||Li full cell is a promising candidate for stationary energy storage.

We design the battery with AlCl₃-LiCl cathode, LLZTO solid electrolyte and lithium anode. We selected the AlCl₃-LiCl as cathode with the following design thinking in mind: 1) The Li-AlCl₃ redox couple is a promising candidate owing to its high theoretical specific capacity (520.7 mAh g⁻¹). 2) The AlCl₃-LiCl system have a low melting point of 110 °C. 3) The AlCl₃ materials have much lower cost compared with other metal chlorides, like NiCl₂. The price of Al is only 15% of the price of Ni. 4) The corrosion of the above molten salt system is much lower than the system containing Ni²⁺ and Fe³⁺, which provides more economical material choices used for cathode current collectors and cell cases [22]. The overall redox reaction of the AlCl₃-LiCl||LLZTO||Li battery is as follow:

\[
\text{LiAlCl}_4 + 3 \text{Li} \leftrightarrow \text{Al} + 4 \text{LiCl} \quad (E_0 = 1.68 \text{ V at 210 °C})
\]

The theoretical specific energy based on the active materials is 686 Wh/kg. We emphasize that our battery system exhibits high safety owing to the relatively low operating temperature and the non-flammable cathode materials. In a Na-S battery, the failure of Al₂O₃ electrolyte or internal shorting of battery will lead to severe safety hazard caused by the short-time high temperature and flammability of molten sulphur. As a comparison, the AlCl₃-LiCl||LLZTO||Li battery system would exhibit much improved safety. Even the ceramic separators have mechanical failure, there is still no serious safety hazard. In that case, the liquid cathode salt will get into contact with the molten Li and react to form non-flammable and non-explosive LiCl and Al, which will block further reaction.

Fig. 1a shows the configuration schematic of the battery. A stainless steel cylinder was used as negative current collector, molten Li as anode, LLZTO ceramic tube (Fig. 1b,c and Fig. S1) as electrolyte, AlCl₃-LiCl (60:40 mole %) as cathode and aluminum foil as positive current collector. Both anode and cathode are liquid state when the cell operates at 210 °C, which ensures the good interface contact between the molten liquid electrodes and the solid electrolyte. At such temperature, the solid electrolyte also exhibits high ionic conductivity of 65 mS cm⁻¹ at 210 °C (Fig. 1d), which is greater than that of most liquid organic electrolytes (ex. ~10 mS cm⁻¹, 1 M LiPF₆ in carbonate-based solvent at room temperature) [23]. Besides, the LLZTO tube possesses a high relative density of 99%, which can effectively prevent any leakage or crossover of the liquid electrode materials and the self-discharge [21,24–27]. The existing of LiCl in the molten salt not only lowers the melting point of the cathode materials, but also functions as a secondary electrolyte to transfer lithium ions.

The voltage profile of the first discharge was shown in Fig. 2a. There is a short plateau at ~1.8 V, which is possibly caused by the side reactions in the interfaces. The main discharge plateau was at ~1.65 V, corresponding to the cathode specific capacity of ~200 mAh g⁻¹. After the plateau, the discharge voltage rapidly fell down to the cutoff voltage (1.2 V). There was ~50% of AlCl₃ contributing to the reaction. As the reaction progressed, the mole ratio of the AlCl₃ and LiCl continuously
decreased, leading to the precipitation of LiCl and the reduction of the liquid phase, therefore the discharge voltage started to fall down when the specific capacity of cathode reached 230 mAh g$^{-1}$. Based on the weight of electrode materials, the specific energy of cell is 350 Wh/kg and the energy density per unit volume is 584.1 Wh/L. The estimated cost is $11.6/kWh.

Long lifespan is a critical factor for large-scale energy storage applications. To test the cyclic performance, an AlCl$_3$-LiCl||LLZTO||Li full cell was cycled at a current density of 10 mA cm$^{-2}$ with a specific discharge/charge capacity of 100 mAh g$^{-1}$ for 100 cycles (800 h). The phase change of the cathode materials during cycling was shown in Fig. 2b. During the discharge process, part of solid LiCl precipitated. Before cycling, the cell was activated to lessen the influence of interface side reactions. As shown in Fig. 2c and d, the cell exhibited a stable cycling performance. The voltage profiles of the 1st, 10th, 50th and 100th cycles are highly consistent (Fig. 2d). The voltage upper and lower limit stabilized at around 2.2 V and 1.4 V, respectively (Fig. 2c). The molten salt cathode delivered an average discharge specific energy of 155 Wh kg$^{-1}$ (based on the weight of cathode materials) with an average energy efficiency of 83% (Fig. 2e). Both the charge energy density and the discharge energy density remained stable during cycling, which also indicated the high cyclic performance of the whole cell. The XRD results (Fig. 1c) indicate the high
stability of LLZTO tube during cycling as there is no obvious phase change before and after cycling. To test the safety of the cell in case the LLZTO tube breaks, a puncturing experiment was conducted. The solid electrolyte tube was full of molten AlCl3-LiCl salt (2 g), and the stainless steel case was full of molten lithium (1 g). After heating to 210°C, the LLZTO tube was punctured, and molten lithium and molten salt was partly contacted. After puncturing, the surface temperature of the stainless steel case increased by only about 50°C.

The cell was also assembled in the discharge state and part of LiCl was replaced by NaCl to lower the cost. The pristine cathode was composed of LiCl, AlCl3 and NaCl (mole ratio, 4: 1: 1) and little amount of Li was used to connect the LLZTO and the current collector. The cell was firstly charged at 5 mA cm⁻² with a specific capacity of 200 mAh g⁻¹ (Fig. S2a). After the first charge process, the cell delivered a stable cycling performance (Fig. S2b), which indicates that the future cells can be produced without metallic lithium to be handled. Like ZEBRA batteries, all the required active metal can be inserted as salt, therefore the assembling process of the battery can be conducted just in a drying room. The cost of raw materials can be also significantly reduced when the cell is assembled in the discharge state, because metal Li is not indispensable. The impedance test result (Fig. S4) shows that the resistance of the full cell is small and leads to a promising future for the application of this battery.

To smooth the fluctuation of intermittent renewable energy, high rate capability is desired for grid energy storage. The rate performance of the AlCl3-LiCl ||LLZTO|| Li cell was shown in Fig. 3a. The cell was cycled at various current densities from 5 mA cm⁻² to 40 mA cm⁻² with a constant discharge/charge capacity of 50 mAh g⁻¹. The voltage profiles at the
current density of 10 mA cm\(^{-2}\) showed no obvious changes before and after the cell was cycled at a high current density of 40 mA cm\(^{-2}\), indicating good rate capability of the cell. The discharge plateaus were 1.65, 1.61, 1.54, 1.46 and 1.34 V at current densities of 5, 10, 20, 30 and 40 mA cm\(^{-2}\), respectively (Fig. 3b). Even at a high current density of 40 mA cm\(^{-2}\), the energy efficiency of cell was still higher than 65%. The power density of the AlCl\(_3\)-LiCl||LLZTO||Li cell reached 52 mW cm\(^{-2}\) at the current density of 40 mA cm\(^{-2}\).

The service life of current collector is also an important factor for rechargeable batteries using in the grid energy storage. In the AlCl\(_3\)-LiCl||LLZTO||Li full cell, the Al foil not only functions as positive current collector, but also participates in the reaction. To investigate the stability of the Al current collector, scanning electron microscopy (SEM) was conducted to display the morphology changes of the Al current collector before and after the cycling test. As shown in Fig. 4a, the pristine Al foil was relative smooth with some little protuberances in the surface. After the first discharge/charge cycle, the surface became rough with some corrosion pits (Fig. 4b), which possibly originated from the defects in the surface of the pristine Al foil. After the 100 cycles, there were many particles with the sizes from several micrometers to 30 μm on the surface of the Al current collector (Fig. 4e and d). Energy Dispersive Spectroscopy (EDS) characterization shows that the particles were composed of Al with little amount of oxygen (Fig. S3). The generation of the micron-sized Al particles were partially caused by the inhomogeneous deposition/dissolution of Al during cycles, which possibly led to the loss of the active Al and the corrosion of the current collector. However, the average thickness of the Al foil did not change after the 100 cycles. Even though the thickness of the Al foil used as current collector was only 35 μm, there was no preferable corrosion point. Therefore, the inhomogeneous deposition of Al is not a serious problem for practical applications.

In summary, we demonstrated a newly designed high temperature battery with molten salt cathode-solid electrolyte-molten lithium anode, which can be operated at a relative low temperature of 210 °C with good safety and low cost. The full cell achieved a stable performance for 100 cycles (800 h) with an average energy efficiency of 83% at a current density of 10 mA cm\(^{-2}\). We propose that the cathode material is not limited to AlCl\(_3\)-LiCl in such battery design, and the battery can be further improved by optimize the molten salt cathode systems in the near future. We anticipate that our AlCl\(_3\)-LiCl||LLZTO||Li cells provide new opportunity for large-scale energy storage solutions.

Notes

The authors declare no competing financial interest.

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Appendix A. Supplementary data

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References