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Electrochemical performance of a lead fluoride electrode mixed with carbon in an

electrolyte containing triphenylboroxine as an anion acceptor for fluoride shuttle

batteries

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Keywords

fluoride shuttle battery; anion acceptor; triphenylboroxine; lead fluoride





Abstract

In fluoride shuttle batteries (FSBs), the addition of an anion acceptor (AA) is required to dissolve the supporting electrolyte salt in an organic solvent. Based on theoretical calculations and practical experiments, the effectiveness of triphenylboroxine (TPhBX) as an AA for FSB was verified. Using an electrolyte with TPhBX as an AA, the specific capacities of the following two types of lead fluoride (PbF₂) electrodes were evaluated: (i) PbF₂ pulverized using a ball mill, (ii) pulverized PbF₂ mixed with carbon using a ball mill. The experimental results indicate that mixing PbF₂ with carbon using a ball mill increases the specific capacity of PbF₂ electrode.

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1. Introduction

Batteries, which utilize the migration of cations such as lithium, sodium, and

magnesium ions to advance the electrochemical reactions, have been widely developed

[1–6]. Reddy et al. (2011) reported an all-solid-based battery that uses the migration of

fluoride ions to advance the electrochemical reaction [7]. Recently, several groups began

to develop batteries that use anion migration [8-14]. In these batteries, metal fluoride

(MF_x; M: metal) is used as an active material, and the discharge/charge reactions proceed

as follows:

Discharge: $MF_x + xe^- \rightarrow M + xF^-$

Charge: $M + xF^- \rightarrow MF_x + xe^-$

The abovementioned batteries, due to the use of a solid electrolyte, can operate at high

temperatures. Contrastingly, the electrochemical performance of batteries with a liquid-

based electrolyte is generally higher than the performance of batteries with a solid-based

electrolyte [15, 16]. Therefore, we paid attention to liquid-based fluoride shuttle batteries

(FSBs) [17–20]. The construction of a liquid-based FSB requires an electrolyte composed

of organic solvents and a supporting electrolyte salt. The supporting electrolyte salt, such

as cesium fluoride (CsF) is difficult to dissolve in the organic solvent (bis[2-(2-

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methoxyethoxy)ethyl] ether; tetraglyme: G4), therefore, an anion acceptor (AA) was added to the G4 [17]. Although the AA addition improved the solubility of CsF, it resulted in capacity degradation during cycling due to dissolution of the active material during charging [18]. To resolve this problem, we modified the CsF concentration in the electrolyte and found that increasing the CsF concentration (saturated state) reduced the dissolution of the active material and suppressed the capacity degradation during cycling [18].

Previous studies have reported that a boroxine-based compound acts as an AA for lithium-ion batteries (LIBs) [21]. In this study, we first evaluate triphenylboroxine (TPhBX)'s ability as an AA via theoretical calculations. Secondly, we investigate TPhBX's suitability as an AA for an FSB electrolyte with practical experiments. Finally, using the electrolyte with TPhBX as an AA, we evaluate the discharge/charge capacities of the lead fluoride (PbF2) electrode. When metal halides were used as an active material in LIBs, previous studies have reported that when mixed with carbon using a ball mill, the specific capacity of the electrodes increases [22–28]. Therefore, in this study, PbF2 electrodes were prepared by two methods: (i) PbF2 was pulverized using a ball mill and (ii) pulverized PbF2 was mixed with carbon using a ball mill.





2. Theoretical calculations and experimental methods

To evaluate the effectiveness of TPhBX as an AA, the adsorption energy (Ea) of F^- on TPhBX was evaluated using the density functional theory (DFT) calculation as mentioned in a previous study [17]. Briefly, Ea was calculated using the following equation:

$$Ea = E(TPhBX-F^{-}) - [E(TPhBX) + E(F^{-})]$$

where $E(TPhBX-F^-)$, E(TPhBX), and $E(F^-)$ are the total gas-phase energies of the TPhBX-F⁻ complex, TPhBX molecule, and F⁻ anion, respectively.

Excess CsF (Tokyo Chemical Industry) was added in G4 (KISHIDA CHEMICAL) containing 0.5 mol dm⁻³ TPhBX (Tokyo Chemical Industry), which was then used as an electrolyte. The CsF concentration in the electrolyte was evaluated using atomic absorption spectrometry (AAS; Hitachi ZA3000). A three-electrode electrochemical cell (EC FRONTIER VB7) was prepared with a platinum (Pt) foil as the working electrode, a Pt mesh as the counter electrode, and a silver rod immersed in acetonitrile containing 0.1 mol dm⁻³ silver nitrate and 0.1 mol dm⁻³ tetraethylammonium perchlorate as the reference electrode (0.587 V vs. standard hydrogen electrode) [29]; a multipotentiostat (Biologic VMP-300) was used to perform cyclic voltammetry (CV) measurement of a Pt electrode in the prepared electrolyte. Three cycles were performed



between potentials of -4.0 and 2.0 V (vs. the Ag/Ag⁺ reference electrode) at a sweep rate of 0.1 mV s⁻¹.

The active material was prepared according to the following method. PbF₂ (Aldrich) was pulverized in a planetary ball mill (FRITSCH PULVERISETTE 7) at 1100 rpm for 1 h. The pulverized PbF₂ is denoted by PbF₂(P). Then, 80 wt% PbF₂(P) and 20 wt% acetylene black (AB) were mixed in a planetary ball mill at 1100 rpm for 1 h, and the obtained material is denoted by PbF₂(C). A scanning electron microscope (SEM; Hitachi SU6600) equipped with an energy dispersive X-ray (EDX) analyzer (HORIBA EX-350x-act) was used to evaluate the distributions of PbF₂ and AB in PbF₂(C) powder.

The PbF₂(P) and PbF₂(C) electrodes were prepared according to the following methods. PbF₂(P) was mixed with AB and polyvinylidene difluoride (PVDF) in a proportion of 60: 25: 15 wt%. Since PbF₂(C) was composed of PbF₂ and AB (80: 20 wt%), PbF₂(C) was mixed with AB and PVDF in a proportion of 75: 10: 15 wt%. Using these electrodes as the working electrode in a three-electrode electrochemical cell, the specific capacities of PbF₂(P)/PbF₂(C) electrodes were measured over the potential range from – 2.2/–2.0 to –0.6 V (vs. ref.) at 0.025C (1C = 219 mA g⁻¹). The specific capacities of both electrodes were obtained by dividing the capacity by the weight of the PbF₂. The electrolyte preparation, sealing of powders in the planetary ball mill, and electrochemical



measurements were performed at room temperature in a glove box filled with high-purity Ar.

The electronic states of Pb in each electrode during the first cycle were analyzed using X-ray photoelectron spectroscopy (XPS; ULVAC PHI Quantera SXMTM) with Al K α X-ray radiation under ultrahigh vacuum conditions. XPS spectra of the electrodes were obtained after Ar⁺ sputtering (Ar⁺ energy: 2 keV, ion current: 1.3 μ A, and time: 16 min). The binding energies were calibrated against the Pb 4f_{7/2} peak of PbF₂ at 139.0 eV, and the intensities were normalized by the peak areas of the Pb 4f_{7/2} peaks.

3. Results and discussion

First, the effectiveness of TPhBX as an AA was evaluated using DFT calculations. Fig. 1 shows the optimized molecular structure of TPhBX–F⁻. The most stable binding site for F⁻ is at the boron atom, and the resultant *Ea* of F⁻ on TPhBX is – 335 kJ mol⁻¹. The *Ea* is close to that of F⁻ on fluorobis(2,4,6-trimethylphenyl)borane (– 312.3 kJ mol⁻¹), which is used as an AA for FSB [17], suggesting that TPhBX is effective as an AA. Second, the effectiveness of TPhBX as an AA was experimentally verified. The solubility of CsF in G4 before and after addition of 0.5 mol dm⁻³ TPhBX was investigated using AAS, and the results indicate that the solubility of CsF in G4 increases



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from 0.00017 to 0.51 mol dm⁻³ after TPhBX addition. Thus, the effectiveness of TPhBX as an AA for FSB is verified by both calculations and experiments.

The electrochemical stability of the prepared electrolyte during cycling was evaluated by CV (Fig. 2). A reduction peak appears below –2.2 V and a negligible oxidation peak is observed in the first cycle. The shape of CV curves changes only slightly during three cycles, indicating that the electrolyte is stable between –2.2 and 2.0 V during cycling.

Next, the extent of mixing of PbF₂ and carbon in the PbF₂(C) powder was evaluated. Previously, particles of PbF₂(C) powder were observed using SEM; however, it was difficult to evaluate the distributions of PbF₂ and AB in PbF₂(C) powder from the secondary electron image [19]. Therefore, the distributions of Pb, F, and C were evaluated using EDX in this work. Fig. 3 shows the secondary electron image and EDX mapping of Pb, F, and C. The bright area observed in the center of Fig. 3(a) is attributed to PbF₂(C), since Pb, F, and C are detected (Figs. 3(b), (c), and (d)). The bright area in which C is detected (Fig. 3(d)) overlaps with the areas in which Pb and F are detected (Figs. 3(b) and (c)), indicating that PbF₂ is adhered to AB in PbF₂(C).

Then, using the prepared electrolyte, we investigated the effects that mixing with AB have on the electrochemical performance of the PbF₂ electrode. Fig. 4 shows the



discharge and charge curves of PbF₂(P) and PbF₂(C) electrodes. Since the discharge reaction of the PbF₂(P) electrode barely proceeded above -2.0 V, the PbF₂(P) electrode was discharged to -2.2 V, which is close to the electrochemical window of the electrolyte. For the PbF₂(P) electrode (Fig. 4(a)), the first discharge capacity is similar to the theoretical capacity (219 mAh g⁻¹). However, the subsequent discharge/charge capacities are much lower than the first discharge capacity. For the PbF₂(C) electrode (Fig. 4(b)), discharge capacity in the first cycle is similar to the theoretical capacity and higher reversible capacities remain during subsequent cycles. The reversible discharge/charge capacities for PbF₂(C) (Fig. 4(b)) are much higher than those for the PbF₂(P) electrode (Fig. 4(a)). These results indicate that mixing PbF₂ with AB using a ball mill increases the reversible capacity of the PbF₂ electrode.

To clarify discharge and charge reaction differences between the PbF₂(P) and PbF₂(C) electrodes, the electronic state of Pb in each electrode during the first cycle was investigated using XPS (Fig. 5). For the PbF₂(P) electrode (Fig. 5(a)), in the pristine state, two large peaks are observed at 139.0 and 143.9 eV and two small peaks are observed at 136.6 and 141.5 eV. Four peaks are indexed to the Pb 4f_{7/2} and 4f_{5/2} peaks of PbF₂ (Pb²⁺) and those of metallic Pb, respectively [30, 31]. Upon changing from the pristine state to the fully discharged state, peaks indexed to metallic Pb increase and peak intensities



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indexed to PbF₂ decrease. From the fully discharged state to the fully charged state, the intensities of the peaks assigned to PbF₂ increase slightly and those assigned to metallic Pb decrease slightly. These results indicate that the discharge reaction partially progresses, and the charge reaction proceeds less than the discharge reaction. The first discharge capacity observed for the PbF₂(P) electrode (Fig. 4(a)) is partially attributed to side reactions such as decomposition of the electrolyte, since the discharge potential is close to the electrochemical window of the electrolyte. For the PbF₂(C) electrode (Fig. 5(b)), from the pristine state to the fully discharged state, peaks assigned to metallic Pb appear and the intensities of the peaks assigned to PbF₂ greatly decrease. From the fully discharged state to the fully charged state, peak intensities assigned to PbF₂ increase and those assigned to metallic Pb decrease. The change in the electronic state of Pb observed for the PbF₂(C) electrode was much larger than that for the PbF₂(P) electrode, indicating

4. Conclusion

the $PbF_2(P)$ electrode.

The ability of TPhBX as an AA was verified by theoretical calculations, and the results indicate that TPhBX can act as an AA. Then, TPhBX's effectiveness to increase

that the discharge/charge reactions of the PbF₂(C) electrode progress more than those of



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the solubility of CsF in G4 was confirmed with practical experiments. Using the electrolyte with TPhBX, the electrochemical performance of PbF₂ electrode was investigated. The discharge and charge reactions for PbF₂ can proceed using the TPhBX-containing electrolyte and the reversible capacity of PbF₂ is further improved by mixing with carbon using a ball mill. The discharge and charge reaction progress of the PbF₂ is confirmable using XPS.

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Figures

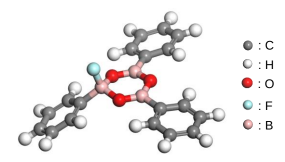


Fig. 1. Optimized molecular structure of TPhBX-F⁻.



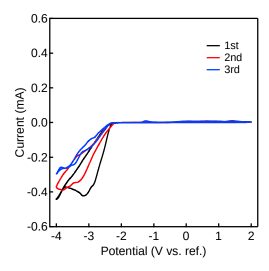


Fig. 2. Cyclic voltammograms of a Pt electrode immersed in an electrolyte during three cycles.

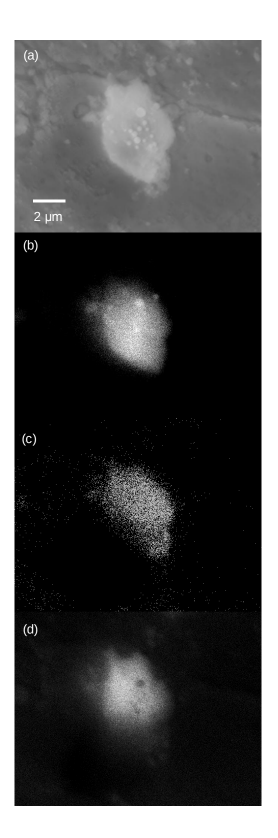


Fig. 3. (a) Secondary electron image and EDX mapping of (b) Pb, (c) F, and (d) C for PbF₂(C) powder.

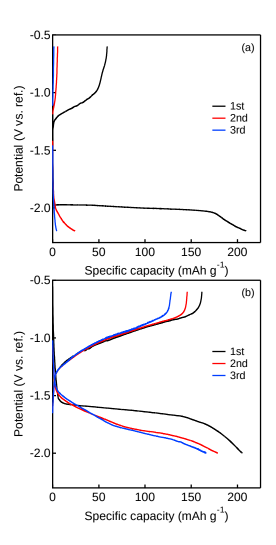


Fig. 4. Discharge and charge curves of the (a) $PbF_2(P)$ and (b) $PbF_2(C)$ electrodes during three cycles.

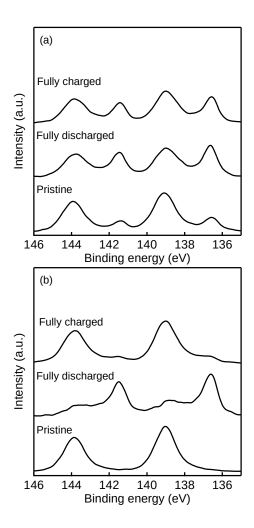


Fig. 5. Pb 4f XPS spectra of (a) PbF₂(P) and (b) PbF₂(C) electrodes in the pristine, fully discharged, and fully charged states during the first cycle.