Triphenylboroxine and triphenylborane as anion acceptors for electrolyte in fluoride shuttle batteries

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For liquid-based fluoride shuttle batteries, electrolyte composed of organic solvent and supporting electrolyte salt is developed. To increase the solubility of supporting electrolyte salt in organic solvent, anion acceptors (triphenylboroxine or triphenylborane) are added. The addition of anion acceptor greatly increases the solubility of supporting electrolyte salt, and discharge-charge reaction of BiF₃ electrode is confirmed in the prepared electrolytes.

Keywords: Fluoride shuttle battery, Anion acceptor, Bismuth fluoride

Currently, lithium ion batteries (LIBs) are widely used as power sources for electronic devices, and recent developments of devices have necessitated the improvement of energy density for batteries.1–3 Therefore, many research institutions develop high-energy-density electrode materials for LIBs4–6 and high-energy-density next-generation batteries.7–10 Recently, new all-solid-based and liquid-based batteries employing metal fluoride as an active material have been proposed.11–14 We have focused on liquid-based fluoride shuttle batteries (FSBs).15 A schematic image of FSB is shown in Figure 1. For positive electrode, metal was formed (MF₃ + xe → M + xF³⁻) during the discharge process, and metal fluoride is formed (M + xF³⁻ → MF₃ + xe⁻) during the charge process. To facilitate the electrochemical reaction in a liquid-based electrolyte over a wide potential range, an electrolyte composed of an organic solvent and a supporting electrolyte salt such as a fluoride compound is used. However, fluoride compounds dissolved sparingly in the organic solvent.15 It is reported that boron-based compounds are used as anion acceptors (AAs), and the addition of AAs in the organic solvent was effective in dissolving insoluble lithium salt for LIBs.16–18 We previously found fluorobis(2,4,6-trimethylphenyl)borane (FBTMBPh) significantly increased the solubility of cesium fluoride (CsF) in bis[2-(2-methoxyethoxy)ethyl]ether (tetruglyme: G4) as an AA, and discharging and charge reactions of a BiF₃ electrode were progressed using the G4 containing CsF and FBTMBPh as an electrolyte.19 In order to extend the usability of the FSB, the potential of other compounds as AA should be investigated for the electrolyte. In this work, two boron-containing compounds, triphenylboroxine (TPhBX) and triphenylborane (TPhB), were selected as potential AAs. The anion-accepting tendency of boron-containing compounds has been reported in the context of LIBs owing to the fact that the positively charged boron ions in the AA attract negatively charged fluoride ions, thus, increasing the solubility of fluoride compounds in organic solvents.19–24 TPhBX contains a boroxine ring and three phenyl groups, whereas TPhB contains a boron atom bound to three phenyl groups.21–24 Herein, the ability of TPhBX and TPhB to play the role of AAs for electrolytes in FSBs and the electrochemical performance of the BiF₃ electrode were investigated using prepared electrolytes.

The electrolytes for FSBs were prepared by dissolving 0.45 mol dm⁻³ CsF (Tokyo Chemical Industry Co., Ltd) and 0.5 mol dm⁻³ TPhBX (Tokyo Chemical Industry Co., Ltd) or 0.5 mol dm⁻³ TPhB (Tokyo Chemical Industry Co., Ltd) in G4 (KISHIDA CHEMICAL Co., Ltd). These electrolytes are denoted as CsF-TPhBX-G4 and CsF-TPhB-G4, respectively, in the following text. Cyclic voltammetry (CV) experiments were conducted to estimate the range of working potential of these electrolytes. The three-electrode electrochemical cell (EC FRONTIER CO., Ltd, VB7) setup of the CV experiment comprised a platinum foil as the working electrode, a platinum mesh as the counter electrode, and a silver nitrate and 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).20 CV experiments on the prepared electrolytes were performed at room temperature using a multipotentiostat (Biologic VMP-300) over a −4.0 to 2.0 V (vs. ref.) potential range at a sweep rate of 0.1 mV s⁻¹. BiF₃ (Fluorochem Ltd.) was used as the active material, and was mixed with acetylene black and polyvinylidene difluoride (60:25:15 wt%) to prepare the electrode. The discharge and charge capacities of the prepared electrode were measured by using it as the working electrode in the three-electrode electrochemical cell with electrolytes CsF-TPhBX-G4 and CsF-TPhB-G4, separately. The whole setup containing the electrochemical cell was housed in an argon atmosphere in a glovebox maintained at room temperature. The charge-discharge measurements were performed using a multipotentiostat (Biologic VMP-300) at 0.025C (1C = 302 mA g⁻¹). The electronic state of Bi from each working electrode containing BiF₃ was analyzed via X-ray photoelectron spectroscopy (XPS; Ulvac Phi Quantera SXM™) using Al Kα X-ray radiation in ultra-high vacuum conditions. After discharging and charging the BiF₃...
 electrode, it was washed with G4, followed by dimethyl carbonate to remove any residual electrolyte. Next, the BiF₃ electrodes were transferred into an XPS chamber without air exposure using transfer vessels filled with high-purity argon. In the XPS chamber, the surface of the BiF₃ electrode was etched to an estimated depth of 12 nm by an Ar⁺ beam to remove any potential contaminants adsorbed during the sample transfer. The binding energy of the obtained spectra was calibrated to that of the Bi 4f₇/₂ peak of Bi metal at 156.9 eV. The intensity of the spectra was normalized to the peak area of the Bi 4f₅/₂ peaks.

Electronic device

Positive electrode

Metal fluoride

Electrolyte
Supporting electrolyte salt, anion acceptor, and organic solvent

Negative electrode

Figure 1. A schematic image of FSB.

Positive electrode: \( \text{MF}_x + \text{xe}^+ \rightarrow \text{M} + \text{xF}^- \)

Negative electrode: \( (x/y)\text{M}' + \text{xF}^- \rightarrow \text{xe}^- + (x/y)\text{M}'\text{F}_y \)

Figure 2. Cyclic voltammograms of a Pt electrode in CsF-TPhBX-G4 and CsF-TPhB-G4. Sweep rate: 0.1 mA s⁻¹. Sweep range: -4.0 - 2.0 V (vs. ref.).

The abilities of TPhBX and TPhB as AAs were investigated by observing the change in the solubility of CsF in G4. Previously, we reported that the solubility of CsF in the G4 is very low (2.6 × 10⁻⁵ mol dm⁻³) without any AAs in the solution. After adding 0.5 mol dm⁻³ TPhBX to G4, up to 0.45 mol dm⁻³ CsF dissolved in G4. This nearly four-order-of-magnitude increase in the solubility of CsF in G4 suggests strong interaction of TPhBX with fluoride ions in G4. Similar results were obtained for TPhB, suggesting the efficacy of both TPhBX and TPhB as AAs for G4.

The potential window of the prepared electrolytes was measured by CV (Figure 2). The reduction peaks in the CV appeared below -1.4 and -1.6 V for CsF-TPhBX-G4 and CsF-TPhB-G4, respectively. With CsF-TPhBX-G4 as the electrolyte, weak oxidation peak was observed due to the decomposition of the electrolyte above 1.0 V. With CsF-TPhB-G4 as the electrolyte, oxidation peaks were observed above 0.2 V. These results indicate that CsF-TPhBX-G4 and CsF-TPhB-G4 are stable between -1.4 and 1.0 V, and between -1.6 and 0.2 V, respectively.

Figure 3. Discharge and charge curves of BiF₃ electrode with electrolyte (a) CsF-TPhBX-G4 within potential range from -1.2 to -0.2 V (vs. ref.) and (b) CsF-TPhB-G4 within potential range from -1.4 to -0.2 V (vs. ref.) in the first, second, and third cycle.

Based on the improved solubility of CsF in G4 by addition of TPhBX or TPhB and electrochemical window measurements, the discharge and charge capacities of BiF₃ were measured by using CsF-TPhBX-G4 and CsF-TPhB-G4.
as the electrolytes. The discharge cut-off potential was limited to 0.2 V from the potential window (CsF-TPhBX-G4: −1.2 V and CsF-TPhB-G4: −1.4 V). The charge cut-off potential was limited to −0.2 V. Without AA, the discharge capacity of the BiF$_3$ electrode is very low (2.0 × 10$^{-5}$ mAh g$^{-1}$). By using CsF-TPhBX-G4, clear improvements in the electrochemical performance were observed, as evident from the first discharge process in Figure 3(a). Here, a plateau at around −0.85 V was observed. The potential of the plateau is almost the same as that for the plateau observed in the electrolyte prepared by CsF, FBTMPbB, and G4 (CsF-FBTMPbB-G4). In CsF-FBTMPbB-G4, the reduction of BiF$_3$ to Bi metal was observed in the discharge process. This suggests that the reduction of BiF$_3$ can also proceed in CsF-TPhBX-G4. A discharge capacity of 318 mAh g$^{-1}$ was observed in the first discharge process. The observed capacity is higher than the theoretical capacity (302 mAh g$^{-1}$) of the BiF$_3$ electrode, suggesting the reaction of the electrolyte with Bi metal, which forms during the discharge process. Similar behavior was also observed in CsF-FBTMPbB-G4. In the next charge process, a capacity of 130 mAh g$^{-1}$ was obtained. Although the charge capacity was lower than the discharge capacity, the charge capacity was clearly observed. By increasing the cycles, the observed discharge/charge capacities of BiF$_3$ in the CsF-TPhBX-G4 were changed to 318/130, 38/46, and 7/14 mAh g$^{-1}$ in the first, second, and third cycles, respectively. The discharge and charge capacities decreased as the cycling progressed; however, the discharge and charge reactions progressed from the first to the third cycles. Figure 3(b) shows the discharge and charge curves of the BiF$_3$ electrode in CsF-TPhB-G4. The discharge/charge capacities of BiF$_3$ in CsF-TPhB-G4 were 364/184, 123/119, and 78/95 mAh g$^{-1}$ in the first, second, and third cycles, respectively. The charge of AA from TPhBX to TPhB improved the discharge and charge capacities and cycling performance. These results indicated that the discharge and charge reactions of the BiF$_3$ electrode progressed in the electrolyte containing TPhBX or TPhB, and the nature of AA affected the electrochemical performance of the BiF$_3$ electrode for the FSB.

To confirm the progress of discharge (BiF$_3$ + 3e$^-$ → Bi + 3F$^-$) and charge (Bi + 3F$^-$ → BiF$_3$ + 3e$^-$) reactions in the prepared electrolytes, the changes in the electronic state of Bi in the BiF$_3$ electrode during the discharge and charge processes were investigated by XPS. The Bi 4f spectra of the BiF$_3$ electrode in the pristine, fully discharged, and fully charged states prepared in the CsF-TPhBX-G4 and CsF-TPhB-G4 are shown in Figure 4. In Figure 4(a), in the pristine state, large peaks were observed at 160.5 and 165.8 eV, which are assigned to Bi 4f$_{5/2}$ and 4f$_{7/2}$ spectra of Bi metal, respectively. This is due to the reduction of BiF$_3$ by the Ar$^+$ sputtering.

From the pristine to fully discharged state, the peaks assigned to Bi metal enlarged and the peaks assigned to Bi$^{3+}$ diminished, indicating that Bi is reduced during discharging. From the fully discharged to fully charged states, the peaks assigned to Bi$^{3+}$ appeared and the peaks assigned to Bi metal became smaller. This indicates that Bi is oxidized during charging. As shown in Figure 4(b), the changes in the spectra from the pristine to fully discharged, followed by fully charged state are similar for CsF-TPhB-G4 and CsF-TPhBX-G4 (Figure 4(a)). This indicates that Bi is reduced during discharging and oxidized during charging when using CsF-TPhB-G4. These results support the progress of discharge and charge reactions in both electrolytes.

In summary, the addition of TPhBX and TPhB greatly improved the solubility of CsF in G4. The discharge and charge reactions of the BiF$_3$ electrode can be progressed using an electrolyte containing TPhBX or TPhB. The cycling performance of the BiF$_3$ electrode in the electrolyte with TPhB was better than that with TPhBX. The boron-containing compounds can play the role of AA, and the G4 containing these compounds can be used as electrolytes in
The electrochemical performance of the BiF₃ electrode was affected by the nature of AA in the electrolyte. Future work will attempt to investigate the effect of AA nature on the electrochemical performance of the BiF₃ electrode and to improve the cycling performance of the BiF₃ electrode.

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References and Notes
