Impact of Low Temperatures on the Lithiation and Delithiation Properties of Si-Based Electrodes in Ionic Liquid Electrolytes

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ABSTRACT: Lithium-ion batteries are used in various extreme environments, such as cold regions and outer space; thus, improvements in energy density, safety, and cycle life in these environments are urgently required. We investigated changes in the charge and discharge properties of Si-based electrodes in ionic liquid electrolytes with decreasing temperature and the cycle life at low temperature. The reversible capacity at low temperature was determined by the properties of the surface film on the electrodes and/or the ionic conductivity of the electrolytes. The electrode coated with a surface film formed at a low temperature exhibited insufficient capacity. In contrast, a Si-only electrode precoated with the surface film at room temperature exhibited a cycle life at low temperatures in ionic liquid electrolytes longer than that in conventional organic liquid electrolytes. Doping phosphorus into Si led to improved cycling performance, and its impact was more noticeable at lower temperatures.

1. INTRODUCTION

Since the commercialization of lithium-ion batteries (LIBs), the importance of rechargeable batteries continues to increase; for example, LIBs are widely used as power sources in devices ranging from portable electronics to electric vehicles (EVs), mounted on microsatellites, and used as a power source for space applications. The EV range in cold regions drops by half compared to their range at normal temperatures. Additionally, LIBs will contribute to the electrification of aircraft in the future. Thus, LIBs will be used in extreme environments, including those that have low and high temperatures, are under vacuum, and experience radiation exposure; therefore, it is essential to investigate the electrochemical performances of LIBs in various environments and improve their energy density, safety, and cycle life.

Silicon (Si) is an up-and-coming material for negative electrodes in next-generation LIBs because of its theoretical capacity (3580 mA h g⁻¹ for Li₁₄Si) is higher than that of currently used graphite (372 mA h g⁻¹ for Li₀.₁₇C). Nevertheless, the poor cycling performance of Si-based electrodes is an obstacle to their practical use, which can be attributed to following: the significant volume change during lithiation and delithiation which generates considerable stress and high strain in active materials; high electrical resistivity; and a low Li⁺ diffusion coefficient. Various attempts have been made to address such issues, for example, synthesizing nanosized Si materials to prevent the occurrence of surface cracking and fracture, reducing the electrical resistivity of Si by coating it with carbon materials; fabricating composite electrodes to cover the shortcomings of Si; doping Si with impurities, such as phosphorus (P), boron, antimony, or arsenic, to adjust its properties, including its electrical resistivity, Li distribution, crystallinity, and morphology; preparing silicides to give ductility and electronic conductivity specific to metals; and the prelithiation of Si to increase the initial Coulombic efficiency.

Electrolytes dominate the charge and discharge properties and safety of rechargeable batteries. A battery with an enhanced energy density increases the risk of ignition and explosion, and therefore, nonflammable electrolytes are required. In particular, the electrolyte viscosity increases with decreasing temperature, leading to an increase in internal resistance. Additionally, low temperatures result in higher rates of Li dendrite formation, decreasing the safety of a battery. To improve battery safety, fluorine-containing organic liquid electrolytes have been applied to graphite negative electrodes. In contrast, ionic liquid electrolytes that have superior physicochemical properties have been studied for Si-based electrodes. A certain ionic liquid electrolyte contributes to...
an improvement in not only the safety but also the lithiation–delithiation properties of a battery. Consequently, LIBs using ionic liquid electrolytes have been increasingly studied as next-generation rechargeable batteries.

While the electrochemical performances of Si-based electrodes at room temperature in various ionic liquid electrolytes and at low temperature in conventional organic liquid electrolytes have been investigated, there are few reports on their performances at low temperatures in the ionic liquid electrolytes. Considering the use of LIBs in cold regions and outer space, it is important to understand and improve their low-temperature characteristics. In the present study, we investigated changes in the lithiation and delithiation properties of a Si electrode with decreasing temperature in ionic liquid electrolytes and in the cycle life at low temperatures. We also demonstrated that a P-doped Si electrode that exhibited superior electrochemical performance at room temperature showed superior low-temperature characteristics. Herein, we used lithium bis(fluorosulfonyl)amide (LiFSA) dissolved in N-methyl-N-propylpyrrolidinium bis(fluorosulfonyl)amide (Py13-FSA) and LiFSA dissolved in 1-ethyl-3-methylimidazolium bis(fluorosulfonyl)amide (EMI-FSA) as ionic liquid electrolyte. The Si electrodes exhibited superior cycle life and better rate performance in the former and latter electrolytes, respectively.

2. EXPERIMENTAL SECTION

2.1. Materials. Elemental Si (99.9%) was obtained from FUJIFILM Wako Pure Chemical Corporation, Ltd. and was mechanically milled by a planetary high-energy ball mill (P-6, Fritsch). The active material of 100 ppm P-doped Si powder (Silgrain e-Si, Elkem ASA) was used as received. LiFSA (99%), 1 mol dm⁻³ (M) lithium hexafluorophosphate (LiPF₆) dissolved in 1:1 (by volume) mixture of ethylene carbonate (EC) and diethyl carbonate (DEC) and 1 M lithium bis(trifluoromethanesulfonyl)amide (LiTFSA) dissolved in propylene carbonate (PC) were purchased from Kishida Chemical Co., Ltd. Ionic liquids of Py13-FSA (97%) and EMI-FSA (97%) were supplied by Kanto Chemical Co., Inc. The chemical structures of the above Li salts, ionic liquids, and organic liquids are displayed in Figure S1 (Supporting Information).

2.2. Electrode Fabrication and Cell Assembly. Si- and P-doped Si electrodes were fabricated by the gas-deposition (GD) method without any binder and/or conductive assistance. We used copper foil as the current collector. The loading amount of active material on the Cu substrate was 30 ± 3 μg. Other detailed GD conditions are described in our previous papers.

We assembled coin cells (2032-type) using the Si- or P-doped Si electrode as the working electrode. Details of the counter electrode and separator are reported in our previous papers. We used 1 M LiFSA/Py13-FSA and 1 M LiFSA/EMI-FSA as the ionic liquid electrolyte and 1 M LiPF₆/EC + DEC (1:1, vol %) and 1 M LiTFSAC/PC as the organic liquid electrolyte. We assembled the cell and prepared the electrolyte in an Ar-filled glovebox (Miwaf MFG, DBO-2.5LNK-TS) with an oxygen content less than 1 ppm and a dew point below −90 °C.

2.3. Charge–Discharge Testing. Charge–discharge testing was galvanostatically performed using an electrochemical measurement system (HJ-1001S58, Hokuto Denko Co., Ltd.) and a thermostatic chamber (SH-242, ESPEC Corp.).

Precycling was conducted as follows to form a good surface film on the electrode; the Si- or P-doped Si electrode was charged from open circuit voltage (OCV) to 0.500 V vs Li⁺/Li at 0.1C, maintained at 0.500 V vs Li⁺/Li for 12 h, and then discharged to 2.000 V vs Li⁺/Li at 0.1C (1C: 3600 mA g⁻¹). That is, the charge and discharge procedures of the precycling were conducted by the constant current (CC) and constant voltage (CV) mode and CC mode, respectively. Charge–discharge testing was performed at various temperatures between 303 and 233 K, whereas low-temperature testing was conducted at 263 K. The charge capacity limit was determined by controlling the limitation time to approximately 17 and 8 min for the 1000 and 500 mA h g⁻¹ capacity limitation, respectively. The C rate was set at 1C and the current density per area of 1C was 108 μA cm⁻². Other detailed conditions are described in our previous paper.

2.4. Ionic Conductivity Measurement. The ionic conductivity (σ) of the various electrolytes was investigated by electrochemical impedance spectroscopy (EIS, Compact-Stat, Ivium Technologies B.V.) in the temperature range of 303 to 233 K. A cell equipped with two Pt electrodes was used in an Ar atmosphere. The cell constant was calibrated using a KCl standard solution. The cell was placed in the thermostatic chamber at each temperature for at least 1 h. EIS measurements were then performed by applying a sine wave with an amplitude of 10 mV over a frequency range of 10 kHz to 50 Hz.

3. RESULTS AND DISCUSSION

3.1. Lithiation–Delithiation Properties of a Si Electrode at Various Temperatures. Figure 1a shows the schematic control flow for charge–discharge testing with decreasing temperature. We performed testing for five cycles...
after precycling at room temperature (303 K). Then the temperature was decreased by 10 K in 1 h and held for 5 h at 293 K to bring the temperature of the whole cell to 293 K. After that, we conducted charge–discharge testing for five cycles at 293 K. The set of the processes indicated by the asterisk in Figure 1a was repeated until reaching 233 K. Finally, the testing was again carried out at 303 K. Figure 1b shows changes in the discharge capacity of a Si electrode with a decrease in temperature in various electrolytes. The electrode could maintain a reversible capacity of 1000 mA h g\(^{-1}\) in all electrolytes until reaching 293 K. It was expected that the Si electrode at low temperatures in PC-based electrolytes would show superior charge and discharge properties because PC remains in a liquid state between 224 and 515 K.\(^{52-54}\) However, contrary to our expectations, the capacity in LiTFSA/PC was the lowest at a relatively high temperature of 283 K. The discharge capacity of 1000 mA h g\(^{-1}\) in LiFSA/Py13-FSA and LiFSA/EMI-FSA or LiPF\(_6\)/EC+DEC began to decrease at 273 and 263 K, respectively. The Si electrode exhibited almost no reversible capacity at 243 and 233 K regardless of the electrolyte, whereas the capacity recovered to 1000 mA h g\(^{-1}\) when returned to 303 K, implying no electrode degradation.

Figure 2 provides each fifth differential capacity (dQ/dV) plot at various temperatures in the ionic liquid electrolyte (1 M LiFSA/Py13-FSA) and organic liquid electrolyte (1 M LiTFSA/PC). The corresponding charge/discharge curve is shown in Figure S2.

Figure 3 and Table 1 display the \(\sigma\) and activation energy \(E_a\) values of various electrolytes. Because the ionic liquid is not in the liquid state but in a gel-like state when evaluated at the lowest temperature (233 K), the \(\sigma\) at 233 K is not shown in Figure 3a. Additionally, the \(\ln \sigma\) at 243 K was out of correlation of Figure 3b; thus, we show the Arrhenius plot between 303 and 253 K. \(E_a\) was calculated from the equation \(\ln \sigma = -E_a/RT + C\), where \(R\) is the gas constant and \(C\) is a constant. Although the obtained \(\sigma\) at room temperature was in close agreement with the reported values, the \(E_a\) estimated from room temperature to low temperatures was several kJ mol\(^{-1}\) higher than the value evaluated from room temperature to high temperatures, implying slower ion conduction at low temperatures.\(^{53,56}\) Additionally, the \(E_a\) for the organic liquid electrolytes was higher than that for the ionic liquid electrolytes. This result indicated the faster ion conductive behavior of the organic liquid electrolytes over the temperature range due to their lower viscosities.\(^{60}\) The \(\sigma\) of LiTFSA/PC was almost the same as that of LiFSA/Py13-FSA at 283 K. Thus, the capacity decay in LiTFSA/PC at 283 K (Figure 1b) did not result from \(\sigma\) but should be attributed to the difference in properties of the surface film formed on the electrode, that is, the higher interfacial resistance in LiTFSA/PC.

Figure 2. Each fifth dQ/dV plot of the Si electrodes at various temperatures in (a) 1 M LiFSA/Py13-FSA and (b) 1 M LiTFSA/PC. The corresponding charge/discharge curve is shown in Figure S2.

### Table 1. Activation Energy of Various Electrolytes Estimated by Figure 3b

| electrolyte     | \(E_a\) (kJ mol\(^{-1}\)) |
|-----------------|--------------------------|
| LiFSA/Py13-FSA  | 30.6                     |
| LiFSA/EMI-FSA   | 28.0                     |
| LiPF\(_6\)/EC+DEC | 20.1                   |
| LiTFSA/PC       | 22.8                     |
3.2. Cycle Life of the Si Electrode at Low Temperature. Figure 4a,b shows the schematic procedure for the cycle life test of the Si electrode and the obtained cycle life, respectively. Precycling was performed at (i) 303 and (ii) 263 K. Figure S3 displays the relationship between the potential and capacity when precycling at each temperature. While the charge capacity was approximately 870 and 570 mA h g$^{-1}$ in the CC and CV modes at 303 K, respectively, it was 350 and 570 mA h g$^{-1}$ in the CC and CV modes at 263 K, respectively. Precycling at 303 K enabled a discharge capacity of 500 mA h g$^{-1}$ to be maintained over 500 cycles. In contrast, the capacity did not reach 500 mA h g$^{-1}$ and faded with precycling at 263 K. These results revealed that the surface film formed at room temperature gave the Si electrode superior lithiation and delithiation properties, whereas the surface film formed at the low temperature had a low σ and an inhomogeneous thickness and/or components. The same should apply to the Li metal as the counter electrode. When the cycle test at the low temperature was performed with five cycles at 263 K after precycling at 303 K, the cycling performance was poor (data not shown). This result indicates that the five cycles at 303 K after precycling contribute to stabilizing the surface film; in other words, the slight formation of a surface film should occur during those five cycles.\(^{50}\) Hence, we investigated the cycle life of Si-based electrodes at the low temperature after precycling and five cycles at 303 K.

Figure 5 provides the cycle life of a Si electrode at 263 K in organic and organic liquid electrolytes with a charge capacity limitation of 500 mA h g$^{-1}$. In organic liquid electrolytes, an initial discharge capacity of 500 mA h g$^{-1}$ was not obtained, which is an expected based on Figures 1b and 3 and Table 1. In contrast, the Si electrode maintained a reversible capacity of 500 mA h g$^{-1}$ for over 150 and 500 cycles in 1 M LiFSA/EMI-FSA and 1 M LiFSA/Py13-FSA, respectively. We demonstrated that the cycle life of Si-based electrodes in FSA-based organic liquid electrolytes was superior to that in conventional liquid electrolytes at room temperature.\(^{17,20,21,50}\) In organic liquid electrolytes, a surface film with an inhomogeneous thickness forms on the electrode surface, and preferential Li$^+$ storage into Si occurs through thinner parts of the film. Inhomogeneous Li$^+$ storage prompts the local formation of a Li-rich Si alloy phase, which causes local changes in the Si volume and electrode disintegration.\(^{23}\) Conversely, in ionic liquid electrolytes, a surface film that includes LiF, which contributes to its mechanical and/or structural stability, forms by the rapid decomposition of the FSA anion;\(^{61}\) thus, the surface film does not grow and remains uniform and thin at room temperature. The results indicate that homogeneous formation of a Li-poor Li$^-$Si alloy phase occurs over the entire electrode, preventing the local accumulation of high strain.\(^{23}\) Thus, severe Si electrode disintegration is suppressed. Additionally, the surface film formed in 1 M LiFSA/EMI-FSA should be thicker than that derived from the Py13-based electrolyte because EMI cation decomposition continually takes place in addition to FSA anion decomposition.\(^{17,62,63}\) While we conducted the cycle test at the low temperature, surface film formation was performed at room temperature. Consequently, it was concluded that the cycle life at the low temperature was similar to that at room temperature and that the Si electrode exhibited a superior cycle life in 1 M LiFSA/Py13-FSA.

3.3. Impact of P-Doping into Si on Its Charge–Discharge Properties at Low Temperature. We previously reported that the electrochemical performance of Si electrodes was improved by P-doping into Si\(^{19–21}\). The effect of P-doping on the performance improvement is explained as follows: (1) it suppresses the phase transition from Si into Li$_x$Si$_{1-y}$ due to shrinking the Si crystal lattice as a result of the replacement of some Si atoms with smaller P atoms; (2) it homogenizes the Li distribution because of faster Li diffusion within P-doped Si as a result of low electrical resistivity; and (3) it suppresses the Si volume expansion during its lithiation, which results from the above two effects.\(^{19,21,23}\) We investigated the charge–discharge properties of a P-doped Si electrode at low temperatures. Figures S4 and S5 show changes in the reversible capacity of the 100 ppm P-doped Si electrode with decreasing temperature in various electrolytes and the corresponding charge–discharge curves, respectively. Compared to the Si electrode (Figure 1b), the capacity decay of the P-doped Si electrode with decreasing temperature was suppressed.

Figure 6 provides the cycle life of 100 ppm P-doped Si and Si electrodes at (a) 263 and (b) 303 K in 1 M LiFSA/Py13-FSA with a charge capacity limitation of 500 mA h g$^{-1}$. There was no significant difference between the charge–discharge properties of the Si electrode after 1000 cycles. In contrast, the capacity of the Si electrode after 1000 cycles was 300 mA h g$^{-1}$ in 1 M LiFSA/Py13-FSA. The cycle life of 100 ppm P-doped Si and Si electrodes at 263 K was similar to that of Si electrodes at 303 K.
profiles of the electrodes at any temperature and cycle (Figure S6). The metallic luster of the counter electrode (Li metal) seemed to be slightly lost after the charge−discharge testing at 303 K, whereas white precipitates covered the electrode at 263 K. The appearance of the electrode after the testing differs depending on the temperature, but there is no difference at the same temperature (Figure S7); thus, the effect of the stripping and precipitation of the Li metal should be ignored. At 263 K, capacity fading of the Si- and P-doped Si electrodes occurred at approximately the 550th and 1710th cycles, respectively. Conversely, at 303 K, the Si- and P-doped Si electrodes exhibited capacity decay at approximately the 2000th and 3100th cycles, respectively. P-Doping into Si improved the cycle life of the electrode by approximately 3.1 times at low temperatures, whereas it was approximately 1.6 times at room temperature: the impact of P-doping into Si on the cycling performance improvement was more noticeable at low temperatures. Lithium diffusion changes within P-doped Si; thus it is considered that the shrinkage of the Si crystal lattice by P-doping contributes to the above improvement because the difference in temperature did not influence on the crystal lattice.

Figure 7 shows the cycle life of the 100 ppm P-doped Si electrode in various electrolytes at a lithiation capacity limitation of 500 mA h g\(^{-1}\). In LiPF\(_6\)/EC+DEC, an initial discharge capacity of 500 mA h g\(^{-1}\) was obtained, whereas the Si electrode did not exhibit 500 mA h g\(^{-1}\) (Figure 5). In LiFSA/EMI-FSA, P-doping into Si improved the cycle life of the electrode by approximately 2.7 times, which is almost the same as the observed improvement in LiFSA/Py13-FSA. Hence, P-doping into Si enhanced the cycle life regardless of the electrolyte.

4. CONCLUSIONS
To use LIBs in various extreme environments, such as cold regions and outer space, we investigated the lithiation and delithiation properties of Si-based electrodes at low temperature. The reversible capacity could not be obtained with a surface film formed at low temperatures. The electrodes exhibited superior cycle life in the ionic liquid electrolytes compared to conventional organic liquid electrolytes. The P-doped Si electrode had a cycle life at a low temperature (263 K) 3.1 times longer than that of the Si-only electrode, while it showed 1.6 times longer cycle life at room temperature (303 K). The effect of P-doping into Si on the cyclability improvement was more noticeable at low temperatures. Li diffusion changes within P-doped Si; thus, the shrinkage of the Si crystal lattice by P-doping may contribute to the above improvement because the difference in temperature did not influence the crystal lattice. Additionally, P-doping into Si enhanced the cycle life irrespective of the electrolyte. The obtained findings are essential for the practical application of LIBs using ionic liquid electrolytes.

ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c00947.

Chemical structures, charge−discharge curves, temperature dependence of the discharge capacity, dQ/dV plots, and photographs of the counter electrodes (PDF)

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