Improvement of Cycle Capability of Fe-Substituted Li₂S-Based Positive Electrode Materials by Doping with Lithium Iodide

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For an attempt to improve the cycle capability of the Li₈FeS₅ cells, we have prepared LiI-doped Li₈FeS₅ composite positive electrode materials. The obtained Li₈FeS₅ · xLiI sample cells showed the improved cycle capability, though the initial capacity value decreased with proportional to the mass of LiI. The improved cycle performance was attributable to the suppressed resistance rise of the cells, due probably to the suppression of high-resistive surface precipitates formed by the reaction between the active material and the electrolyte. The dopant LiI would stabilize the local structure against Li extraction/insertion reactions, as well as suppress the reaction with the electrolyte, leading to the improved cycle performance.

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Lithium sulfide (Li₂S) is one of the promising cathode active materials for high-energy rechargeable lithium batteries, because it has a high theoretical capacity (ca. 1170 mAh·g⁻¹) and can be used with a variety of anode materials without lithium sources (such as graphite, silicon) in practical battery systems. However, this material shows high electrical resistivity, which gives rise to poor material utilization in the cells. In order to enhance the conductivity of Li₂S, several attempts, such as forming composites with metals (Li₂S-Fe, Li₂S-Cu) or carbon (Li₂S-C), have been performed. Recently, we have prepared Li₂S-FeS (x = 1, 2) composites using a combination of thermal heating (spark plasma sintering; SPS) and mechanical milling. The composite samples consisted of Fe-containing low-crystalline Li₂S and showed relatively high specific capacity particularly in the case of the Li₂S-rich composition, typically ca. 730 mAh·g⁻¹ for the Li₂S-FeS composite (Li₈FeS₅) cell. Although relatively high initial discharge capacity was attained, the Li₈FeS₅ sample cell showed rapid capacity degradation with cycling; the capacity retention after 30 cycles was ca. 36% of the initial value.

In traditional Li-S batteries, there still suffers from a rapid capacity fading with cycling, which are mainly related to the shuttle effect of polysulfides and the insulating nature of sulfur as well as the reduced products of Li₂S and/or Li₂S₂. Since both the ionic and electronic channels are important to ensure superior cyclic stability of Li-S batteries, developing sulfur-based cathode materials with high ionic and electronic conductivity might be one of the effective ways to improve their electrochemical performances. Recently, Hakari et al. applied Li₈S-Lix (X = Cl, Br, and I) solid solutions, showing higher ionic conductivity (10⁻⁶ - 10⁻⁷ S·cm⁻¹) as compared to Li₂S (ca. 10⁻¹³ S·cm⁻¹), to all-solid-state cells and found the improved utilization of the active materials with superior cycle capability. In particular, the Li₈S-LiI sample cell exhibited nearly theoretical capacity value, due mainly to the enhanced electrochemical reaction sites. On the other hand, addition of LiI to liquid electrolyte in Li-S batteries has been reported to result in the improved capacity and cycle stability due to the formation of the “protective” layer on the surface of both electrodes; such Li-permeable layer prevents the dissolution of polysulfides on the cathode side and the reduction of polysulfides on the anode side.

In the present work, we have prepared LiI-doped Li₈FeS₅ positive electrode materials (Li₈FeS₅·xLiI) for an attempt to improve the cycle capability of the Li₈FeS₅ cells. We have examined the effect of LiI from the standpoint of the structural change of the active material itself, as well as the formation of some protective layer on the surface of both electrodes.

Experimental

First, a blended powder of Li₈S and FeS in a 4: 1 molar ratio underwent SPS process (SPS-3.20 MK-IV, Fuji Electronic Industrial, Japan) at 600 °C for 3 min under an argon atmosphere, and the product was blended with LiI and acetylene black (AB) powders so as to the molar ratio of 4Li₂S·FeS: LiI: Li = 1: x(0 ≤ x ≤ 1) and weight ratio of (4Li₂S·FeS + xLiI): AB = 9: 1, and then mechanically milled for 8 h using a pulverizer (Model No. MC-4A, Ito Seisakusho Ltd., Japan) at a rotation speed of 1000 rpm, to yield the Li₈FeS₅·xLiI samples. Most of the procedures were carried out in an argon-filled glove box, because Li₈S and LiI·xLiI are very sensitive to atmospheric moisture.

Phase purity of the samples was checked by X-ray diffraction (XRD) (RINT TTR-III, Rigaku, Japan) using monochromatic Cu Kα radiation within a 2θ range of 10–125°. Before the measurements, each sample was covered with Kapton film in an argon-filled glove box, and measurements were carried out within 1 h to minimize reaction with atmospheric moisture. Structural refinement by X-ray Rietveld analysis was carried out using the RIETAN-2000 program. The local atomic structure of each sample powder was examined by taking high-energy X-ray total scattering measurements, which were carried out at BL28XU of SPring-8. The incident X-ray energy was 38.0 keV, and scattering patterns measured with a Q-range from 0.3 to 17 Å⁻¹ were analyzed using pair distribution function (PDF) analyses. A vacuum chamber was used to avoid air scattering the sample. The measured datasets were corrected for absorption, background, and polarization effects, and were normalized to obtain the reduced pair distribution function G(r), according to the procedure detailed elsewhere. The elemental composition of the sample was examined by inductively coupled plasma – atomic emission spectroscopy (ICP-AES, Thermo Fisher Scientific, Inc., ICP 6500 DUO). The valence state and local structure of I atom for the sample electrodes were examined by I L₃-edge X-ray absorption fine structure (XAFS) measurements, which were carried out at BL-3 of the Synchrotron Radiation Center, Ritsumeikan University. The...
conventional transmission mode was used and the incident X-ray beam was monochromatized with a Si(220) crystal (2θ = 3.840 Å).26,29 All the samples were sealed in an argon-filled aluminum-laminated bag. The surface analyses of the sample electrodes were carried out by X-ray photoelectron spectroscopy (XPS) measurements (Quantera SXM, Ulvac-PHI) using monochromatic Al Kα1,2 radiation (1486.6 eV). The typical analysis area was 200 μm in diameter, and binding energy was calibrated with C1s peak (284.6 eV).

Electrochemical lithium insertion/extraction reactions were carried out using lithium coin-type cells. The working electrode consisted of a mixture of 11.1 mg of the Li8FeS5-LiI sample (containing 10% (1.1 mg) AB) and 3.9 mg of additional AB powder with 0.5 mg of Teflon powder pressed into a 15 mm diameter tablet under a pressure of 10 MPa. The electrochemical test cell was constructed in a stainless steel coin-type configuration. The negative electrode was a 0.5 mg of Teflon powder pressed into a 15 mm diameter tablet under a pressure of 10 MPa. The electrochemical test cell was constructed in an argon-filled glove box, and electrochemical measurements were carried out at 30 °C.

Results and Discussion

The SPS-treated 4Li2S + FeS blended powder was grayish black in color,12 and it changed to a black powder after the addition of LiI + AB and milled for 8 h (Li6FeS5 · xLiI). Figure 1 shows the XRD patterns of the obtained Li6FeS5 · xLiI samples with different LiI content (x). The LiFeS5 · xLiI samples mainly consisted of low-crystalline Li2S. As discussed previously,14 the LiFeS5 sample (x = 0) could be best described as having a Li12S-based anti-fluorite structure in which Fe ions partially occupy interstitial sites close to Li ions in the Fm3m unit cell. For samples with x ≥ 0.1, any peaks ascribed to LiI were not detected, indicating that the added Li and I ions were incorporated into Li2S lattice or LiI was present in an amorphous phase. The lattice parameters estimated from the Li2S peaks were a = 5.7181(14) − 5.7384(8) Å, which were larger than that of the pristine Li8FeS5 (a = 5.7048(10) Å). This can be explained by the partial substitution of the larger I− ions (2.20 Å) for S2− ions (1.84 Å).30 Small amounts of FeS and unknown phase for x = 1 sample (Li6FeS5 · LiI) suggest that it was beyond the solid solution limit in the Li8FeS5-LiI system. These results are consistent with the previous report on the Li1-S-LiI solid solution,15 in which the lattice parameter increased with LiI content up to the solid solution limit of the composition 80LiI · S:20LiI (the above mentioned Li1FeS2 · LiI corresponds to 83Li1.8Fe0.2S:17LiI). The detailed atomic structure will be discussed later using PDF analyses.

Table I lists the atomic content and the resulting composition of the Li8FeS5 · xLiI samples, estimated by ICP-AES measurements. Each sample showed similar atomic ratio to that of the starting composition; for example, the x = 0.5 sample, initially Li8.9Fe0.5S4.5, was estimated to Li8FeS5 · xLiI. Thus, the initial atomic ratio was nearly unchanged during the present preparation process, as was in the case of LiFeS5.12

Contrary to the reduced capacity value, the cycle performance was much improved by the addition of LiI. Figure 2b, particularly in the Li8FeS5 · 0.5LiI sample cells, the capacity retention after 30 cycles was ca. 72%, which was much higher than that of the pristine Li8FeS5 sample cells (ca. 36%). Slight decrease in capacity retention beyond 30 cycles was attributable to the degradation of the cell components, such as electrolyte and lithium anode, due to the incomplete sealing of the present coin-type cells, as was observed also in the conventional Li/LiCoO2 cells. We also carried out the same electrochemical test for the cell with the blended powder of Li8FeS5 · 0.5LiI, and found similar cycle degradation to that of the pristine Li8FeS5 sample cell, as shown also in Fig. 2b. Therefore, a simple blending of LiI has nearly no effect on the improvement of the cycle performance in the present Li8FeS5 cell.

Wu et al.19 reported that the addition of LiI to liquid electrolyte in Li-S batteries improved the cycle capability due to the formation of the
“protective” later on the surface of both electrodes via the reaction between the electrolyte solvent and I⁻ radicals, and such Li-permeable layer prevented the dissolution of polysulfides on the cathode side and the reduction of polysulfides on the anode side. In order to check similar effect in the present Li/Li₈FeS₅ₓLiI cell, we carried out the surface analyses of both electrodes by XPS measurements. The measured XPS spectra for the Li/Li₈FeS₅₀.5LiI and Li/Li₈FeS₅ cells after 15 cycles showed the presence of main elements Li, C, O, and F in both cathode and anode, suggesting the presence of some surface precipitates originated possibly from the decomposition of electrolyte, as in the case of Li-S batteries. Small amounts of sulfur (< 1 atomic%) were detected in both electrodes, as shown in Fig. 3 for the S₂p spectra. The cathode spectra consisted of mainly characteristic two peaks; the peak at 161–162 eV was assigned to Li-S band, while that at 163–164 eV to S-S band. In the present Li/Li₈FeS₅ₓLiI cells, the Li-S band could be possibly originated from active materials, while the S-S band possibly from organosulfur compounds precipitated via the reaction between the active materials and electrolyte. The lower peak height of the S-S band in the Li/Li₈FeS₅₀.5LiI cell suggests that the cathode surface was covered with less amounts of precipitates, as compared to the Li/Li₈FeS₅ cell. Such precipitates would partly migrate during electrochemical cycling to deposit on the anode surface, and they were detected in the anode S₂p spectra, Fig. 3. Less amounts of S-S band component in the anode of the Li/Li₈FeS₅₀.5LiI cell would be originated from less amounts of S-S band component in the cathode. Since any obvious peaks ascribed to the Li-S band were not detected in the anode for both sample cells, reduction of polysulfides on the anode were hardly occurred in the Li/Li₈FeS₅ₓLiI cells, which makes a clear contrast to the Li-S batteries. Therefore, the surface precipitates on the electrodes did not act as protective layer, but were responsible for the cycle degradation of the cells.

The above-mentioned surface precipitates on the electrodes were also suggested by the impedance measurements. Figure 4 shows the Nyquist plots for the Li/Li₈FeS₅₀.5LiI and Li/Li₈FeS₅ cells after the 1st and 20th charging. Both plots consisted of low-frequency spikes and high-frequency deformed semi-circles. According to the previous report on Li-S cells, the deformed semi-circles were resolved into two semi-circles, as shown by the dashed line in Fig. 4, and the data were analyzed as being composed of the electrolyte resistance (Rₑ) and the charge-transfer resistance on anode (Rₐ) and cathode (Rₖ). Each resistance showed similar value for both cells after the 1st charging. However, it showed a significant increase after the 20th charging in the Li/Li₈FeS₅ cell; particularly, Rₖ value, which was a major contributor to the total resistance value, became 10 times higher. This makes a clear contrast to that of the Li/Li₈FeS₅₀.5LiI cells.

Figure 2. (a) Charge and discharge profiles for the Li₈FeS₅ₓLiI sample cells at 46.7 mA · g⁻¹ (corresponding to 0.04C for 2e⁻/Li₂S). The solid and dashed lines represent the profiles with the capacity values normalized by the mass of Li₈FeS₅ₓLiI and Li₈FeS₅, respectively. (b) Cycle performances for the Li₈FeS₅ₓLiI sample cells at 46.7 mA · g⁻¹. The capacity values are normalized by the mass of Li₈FeS₅ₓLiI.

Figure 3. S₂p spectra for cathode and anode in the Li/Li₈FeS₅₀.5LiI and Li/Li₈FeS₅ cells after 15 cycles.

Figure 4. Nyquist plots for the Li/Li₈FeS₅₀.5LiI and Li/Li₈FeS₅ cells after the 1st and 20th charging.
cell, where approximately doubled resistance value was observed. The $R_c$ component would be associated with the surface precipitates on the cathode, possibly consisted of high-resistive organosulfur compounds formed by the reaction between the active materials and electrolyte, and is responsible for the cycle degradation of the cells; less amounts of surface precipitates in the Li/Li$_8$FeS$_5$·0.5LiI cell showed improved cycle capability with suppressed resistance rise. Such improvement effect seemed to increase with the LiI content, but excess amount of LiI resulted in the formation of some impurity phase, as is evident in Fig. 1 ($x = 1$), which might inhibit the structural reversibility of Li$_8$FeS$_5$ for Li extraction/insertion reactions with cycling. Indeed, the capacity retention increased with the LiI content up to $x = 0.5$, while it decreased beyond $x = 0.5$, Fig. 2b. Probably these conflicting two factors were balanced around the composition at $x = 0.5$ in the present Li$_8$FeS$_5$·xLiI system.

For clarifying the mechanism of suppressing the growth of surface precipitates, as well as for comprehensive understanding of the charge/discharge mechanism, we examined the structural change of the Li$_8$FeS$_5$·0.5LiI cathode material. We disassembled the Li/Li$_8$FeS$_5$·0.5LiI cells after the charge and discharge at several steps denoted as closed circles in Fig. 5a, and carried out ex situ XRD measurements. Figure 5b shows the ex situ XRD patterns of the Li$_8$FeS$_5$·0.5LiI positive electrode during electrochemical cycling. In the 1st charge, the peaks ascribed to Li$_2$S decreased in its intensity and finally disappeared (sample C). There observed no definite peaks ascribed to, for example, elemental sulfur, which was anticipated to appear as the charged product in the conventional Li-S cells. In the subsequent discharge, the peaks ascribed to Li$_2$S appeared again (sample F). In the following cycling, the Li$_2$S peaks repeated the disappearance/appearance regularly, indicating that the Li$_8$FeS$_5$·0.5LiI sample shows reversible structural change for Li extraction/insertion reactions with maintaining the anti-fluorite structure, similarly to Li$_8$FeS$_5$.12

To examine the local structure of the Li$_8$FeS$_5$·0.5LiI sample in detail, we carried out PDF analysis of the high-energy X-ray total scattering data, which is particularly useful for determining the structure of amorphous or low-crystalline samples.24–27 First, we examined the local structure before electrochemical test (sample A). Figure 6a shows the experimentally obtained X-ray PDF data (reduced pair distribution function $G(r)$), which shows the probability of finding a pair of atoms, weighted by their scatter power, at distance $r$ for the Li$_8$FeS$_5$·0.5LiI sample as well as those of the Li$_8$FeS$_5$, Li$_2$S, LiI, and FeS powders. As reported previously,14 the PDF of the Li$_8$FeS$_5$ sample has a similar profile to that of the Li$_2$S powder, and can be best explained as having a Li$_2$S-based anti-fluorite structure in which Fe ions partially occupy interstitial sites close to Li ions in the $Fm\overline{3}m$ unit cell. The PDF of the Li$_8$FeS$_5$·0.5LiI sample also has a very similar profile to that of the Li$_8$FeS$_5$ sample, indicating that the basic local structure was maintained after the addition of LiI. Since the characteristic peaks observed in the LiI powder, such as 3.01 Å for Li – I, 4.26 Å for I – I, and 5.22 Å for Li – I, were not observed obviously, the added LiI was not present as its amorphous phase with maintaining the original crystallographic structure in the Li$_8$FeS$_5$·0.5LiI sample.

More detailed structural analysis could be examined by XAFS measurements. Figure 7 shows the ex situ L$_3$-edge X-ray absorption near-edge structure (XANES) spectra for the Li$_8$FeS$_5$·0.5LiI sample during electrochemical cycling. The spectrum before electrochemical test (sample A) was rather similar to that of FeI$_2$, indicating that the valence state and local structure around I ions in the Li$_8$FeS$_5$·0.5LiI sample was rather similar to those of FeI$_2$. Possibly the I ions would occupy some interstitial sites near Fe ions in the $Fm\overline{3}m$ unit cell so as to have similar local structure to that of FeI$_2$, though the atomic position of I ions cannot be determined definitely.

The above-mentioned local structure changed irreversibly during electrochemical cycling. As shown in Fig. 6b, the PDF data changed with Li extraction/insertion reactions; the peak at 4.0 Å attributable to S – S interatomic distance showed splitting during charging (denoted as blue arrows), and it was not reformed to the original one in the subsequent discharging. This suggests an irreversible structural
change around S ions, resulting in inhomogeneity with different S – S distance. Such irreversible change would be responsible for the relatively large capacity loss from the 1st to 2nd cycle, Fig. 2b. In addition, shorter interatomic distance of less than 2 Å appeared during charging (denoted as a red arrow), and disappeared during the subsequent discharging. This would be originated from a shorter S – S distance caused by the delithiation of Li8FeS5 · 0.5LiI, as is often observed in metal polysulfides. After the 2nd cycle, the PDF data showed rather reversible changes for Li extraction/insertion reactions, as shown in Fig. 6c, though all the peaks could not be assigned definitely. Thus, the Li8FeS5 · 0.5LiI sample converted to delithiated phase with inhomogeneous S – S distance during the initial charge, and then it converted to lithiated phase with still inhomogeneous S – S distance during the subsequent discharge, and afterward they mutually repeated well-reversibly with little loss of capacity value.

Accompanying with such structural change, the local structure around I ions also showed an irreversible change. As shown in Fig. 7, the XAFS spectra of the Li8FeS5 · 0.5LiI sample showed a change during the charge, and it was not recovered to the original one after the subsequent discharge. This is consistent with the results of the above PDF analyses. Hakari et al. reported that the 80Li2S – 20LiI solid solution may be decomposed to form LiI dispersed in Li2S matrix on charge process. Similar decomposition may proceed in the Li8FeS5 · 0.5LiI sample during the 1st cycling; rather homogeneously distributed I ions forming a FeI2-like local structure in the Li8FeS5 · 0.5LiI sample were rearranged during the 1st cycling, resulting in forming an inhomogeneous cluster dispersed in the Li8FeS5 matrix. Actually, the peak at 2.9Å in Fig. 6b, which was very close to the I – Fe interatomic distance (2.885Å) in FeI2 and overlaps with rather shorter Fe – Li (2.864Å) and Li – Li (2.852Å) in Li8FeS5, remained nearly unchanged for samples A – C and shifted to shorter r-value for samples D – F (denoted by purple dashed line). At the same time, a new peak at 3.2Å appeared in sample C and shifted to shorter r-value for samples D – F (denoted by brown dashed line), approaching to an interatomic distance for Li – I in LiI (3.01Å). These are suggestive of the local structural change from FeI2-like to LiI-like ones during the 1st cycling. In the subsequent cycling, these characteristic two peaks remained nearly unchanged, Fig. 6c, suggesting a rather steady local structure around I ions. Thus, the initial Li8FeS5 · 0.5LiI converted to a delithiated one (Li2FeS3 · 0.5LiI; y ∼ 0.2) accompanying with a structural rearrangement resulting in inhomogeneous distribution of, particularly, S ions, and then converted to a lithiated one (LiFeS2 · 0.5LiI; y ∼ 0.9) accompanying with a local structural rearrangement around, particularly, I ions. Another notable point is that the edge positions of the XANES spectra, Fig. 7, showed little changes during the electrochemical cycling, indicating that the valence state of I ions did not change for Li extraction/insertion reactions. This result signifies that I ions contributed little to the capacity value of the Li/Li8FeS5 · 0.5LiI cell, which is consistent with the decrease in capacity value with the LiI content, Fig. 2a. This is again indicative of rather steady state around I ions for the Li extraction/insertion reactions, which would fasten the local structure around I ions and

Figure 6. X-ray radial pair distribution function (PDF) data for (a) Li8FeS5 · 0.5LiI sample, as well as for Li4FeS4, Li2S, LiI, and FeS powders, and (b) (c) those during electrochemical cycling. Sample identification is the same as denoted in Fig. 5a. The reduced pair distribution function G(r) indicates the probability of finding a pair of atoms, weighted by their scatter power, at distance r.

Figure 7. 1L3-edge XANES spectra for the Li8FeS5 · 0.5LiI samples during electrochemical cycling. Sample identification is the same as denoted in Fig. 5a. LiI and FeI2 were used as references.
Conclusions

We have successfully improved the cycle capability of Fe-substituted Li$_2$S-based positive electrode material (Li$_8$FeS$_5$) by doping with LiI. The improved cycle performance was attributable to the suppressed resistance rise of the sample cells, due probably to the suppression of high-resistive surface precipitates formed by the side-reaction between the active material and the electrolyte. The dopant LiI probably stabilizes the local structure of Li$_8$FeS$_5$ against Li extraction/reinsertion reactions, as well as suppresses the side-reaction with the electrolyte, resulting in the improved cycle performance.

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