Lithium fiber growth on the anode in a nanowire lithium ion battery during charging

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Lithium (Li) dendrite formation has been recognized as one of the major safety concerns for Li metal batteries but not for conventional Li ion batteries (LIBs) where Li metal is not used. With the advanced in situ transmission electron microscopy enabling direct observation of battery operation, we found that Li fibers with length up to 35 μm grew on nanowire tip after charging. The Li fibers growth were highly directional, i.e., nucleating from the nanowire tip, and extending along the nanowire axis, which was attributed to the strong electric field enhancement effect induced by the sharp nanowire tip. This study reveals a potential safety concern of short-circuit failure for LIBs using nanowire anodes. © 2011 American Institute of Physics. [doi:10.1063/1.3585655]

Lithium (Li) dendrite growth on metal Li anodes represents one of the major safety issues that prevented the wide application of rechargeable Li batteries,1–5 although there is renewed interest in developing such high energy density batteries (e.g., 1700 W h kg−1 for the Li-air cells, comparable to that of gasoline).6 The problem of Li dendrite growth is not a serious issue for today’s lithium ion batteries (LIBs), in which metallic Li electrode is not present. Recently, however, nanostructured electrodes are intensively tested targeting more applications such as electric vehicles or tools.7,8 Advantages by using various nanowires and nanoparticles are emerging in terms of enhancing the energy and/or power density and enabling facile stress release.9–11 However, compared to the bulk counterparts, local environment may change significantly around the nanostructured electrodes and particularly, the electrode/electrolyte interface, resulting in unwanted side reactions.8

Herein, we present the direct observation of Li fiber growth from different nanowire anodes during in situ charging nano-LIBs inside a transmission electron microscope (TEM).12 The “open cell” nano-LIB was built using an individual nanowire anode (of Si or SnO2), an ionic liquid electrolyte (ILE, LiTFSI/P14TFSI, see Ref. 13 for details), and a bulk LiCoO2 cathode (Fig. S1). Long Li fibers were found on the tips of the nanowires after charging, which was attributed to the high electric field at the high-curvature tips induced Li+ reduction and Li plating. Besides, lithium fluoride (LiF) thin film was found on the Li fibers from the electron diffraction patterns (EDPs), evidencing decomposition of the electrolyte and consistent with the observed thick solid electrolyte interface (SEI) layers. Figure 1 shows the Li fiber growth from a SnO2 nanowire anode after charging. Figure 1(a) shows the pristine SnO2 nanowire before charging. The nanowire was straight and clean. After being charged at a constant potential of −3.8 V with respect to the LiCoO2 cathode for 50 min, the nanowire elongated and was bent (Fig. 1(b)). When we pulled the charged nanowire out of the ILE, there was a big chunk of jelled electrolyte adhered to the immersed part of the nanowire. Intriguingly, there was a long tail with light contrast (≈35 μm long) attached to the sharp apex of the jelled electrolyte. Figure 1(c) and movie S1 show the pull-out process of the long tail with unique “necklace” (microcrystals decorated fiber) morphology. One of the particles on the necklace is highlighted [Fig. 1(d)], with a well-defined polyhedral shape. There were also some fibers protruding from the thick jelled electrolyte [Fig. 1(e), also marked by the red triangle in Fig. 1(b)]. The electron energy loss spectra (EELS) from these crystals confirmed their main component being metallic lithium [Fig. 1(f)], as multiple plasmon peaks were clearly seen with primary energy loss of 7.2 eV, being consistent with the characteristic peaks from pure Li metal.14

The Li fiber growth was also observed when Si nanowire anodes were charged (Fig. 2 and Figs. S2–S5). Figure 2(a) shows pristine Si nanowire with uniform diameter. After charging at −4 V against LiCoO2 for 95 min, the nanowire swelled and there was also jelled electrolyte formed on the immersed part of the nanowire [Fig. 2(b)]. A Li fiber, about 15 μm thick and 7.5 μm long, was seen after the nanowire was pulled out of the ILE [Fig. 2(b)]. Again the lithium fiber was grown on the sharp apex at the end of the jelled electrolyte. EDPs from such crystals confirmed that they were single crystalline Li metal (body-centered cubic structure, a = 3.501 Å) covered with a thin layer of polycrystalline lithium fluoride (LiF, face-centered cubic structure, a = 4.027 Å) (Figs. S2 and S3). Because LiF was not present in the original electrolyte, it directly evidenced decomposition of the electrolyte (fluorine from the −CF3 groups of LiTFSI and P14TFSI molecules).
Figures 2(c) and 2(d) illustrate the possible mechanism of Li fiber growth on a nanowire anode. From the experimental observations, the jelled electrolyte formation occurred on all the surfaces in contact with the electrolyte, however, Li fiber growth took place predominantly on the tip of the nanowires or the jelled electrolyte. Consider the capacitor approximation of the two electrodes separated by an electrolyte, the surface charge density, \( \sigma \), should be the highest at the nanowire tip. The electric field at the tip is strongly enhanced, \( E = \sigma / \varepsilon \) (\( \varepsilon \) being the dielectric constant of the electrolyte around the nanowire), is thus the highest [Fig. 2(c)]. In the case of a sharp apex against a planar plate with a potential of \( V \), the electric field on the sharp apex is strongly enhanced as compared to that of two parallel plates capacitor, in the former, the effective electric field \( E \) is inversely proportional to the apex radius \( r \), i.e., \( E \propto V/r \). As a result of this very high electric field at the tip or so-called “tip effect,”\(^{15-18} \) Li plating took place. As the Li deposition at the tip continues, the high electric field preserves and sustains the continuous Li growth into a long fiber. Lateral growth is also possible, since charge may be concentrated at local protrusions on the fiber, which accounts for the formation of the necklace-shaped fiber (Fig. 1). Since the electric field is also present on the sidewalls of the jelled electrolyte, it is likely that Li fiber growth occurs at a much higher threshold of electric field than SEI formation. This may explain that Li fiber growth is not severe in conventional LIBs.

Previous studies have shown strong square (\( J^2 \)) dependence of Li dendrite growth on current density, \( J \). Although the actual current was hardly measured in the \textit{in situ} lithiation of individual nanowires due to their finite mass involved,\(^{12} \) the lithiation current \( (i_c) \) could be estimated on the order of \( \sim 1 \) pA based on the observed material conversion [\( \text{M}+x\text{Li}^{+}+xe^-\rightarrow\text{Li}_x\text{M} \), Fig. 2(d)]. The current density at the tip was calculated to be \( \sim 10^4 \) \( \mu \text{A cm}^{-2} \) by assuming cross section area of \( 10^3 \) \( \text{nm}^2 \), significantly higher than the typical values in the previous Li dendrite formation tests, for instance, \( \sim 10^2 \) \( \mu \text{A cm}^{-2} \) for the Li/polymer/Li cells.\(^3 \) At such highly concentrated electric field and current density, it is not surprising that Li deposition and electrolyte decomposition would happen. The formation of Li fiber with the body-centered cubic structure (the room temperature phase) was expected, because the Joule heating effect was negli-
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targets for next-generation lithium ion batteries. By utilizing the importance of basic research on electrochemical processes, large solid materials deposited on the nanowire. Our results illustrate the importance of basic research on electrochemical processes near nanostructured electrodes, for developing next-generation lithium ion batteries.

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13See supplementary material at http://dx.doi.org/10.1063/1.3585655 for experimental details. Li fibers grown on other Si nanowires after charging, and a movie showing Li necklace being pulled out from the electrolyte.