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# Lithium Metal Stripping Mechanisms Revealed Through Electrochemical Liquid Cell Electron Microscopy

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**Keywords:** electrochemical liquid cell transmission electron microscopy, lithium-ion battery, lithium metal anode, lithium stripping, solid-electrolyte interphase

**Abstract.** Understanding of lithium stripping is as important as that of lithium plating to achieve significant advances in using lithium metal anodes for high-energy rechargeable batteries. However, there have been limited studies on lithium stripping compared to lithium plating. Here we report the lithium stripping mechanisms revealed through *in-situ* electrochemical liquid cell transmission electron microscopy (TEM). We directly observe and compare the stripping behaviors of the *in-situ* grown lithium dendrites and lithium nanograins covered by a lithium fluoride-rich solid-electrolyte interphase (SEI). We find the sporadic lithium stripping behavior and three important modes that can describe the stripping of individual lithium deposits, regardless of their morphology: (i) symmetric stripping, (ii) surface-preferred asymmetric stripping, and (iii) interface-preferred asymmetric stripping. In addition, SEI chemical mapping with high spatial resolution shows a remarkable SEI loss at the end of the lithium metal stripping, which illustrates the importance of SEI protection in the subsequent cycles.

## 1. Introduction

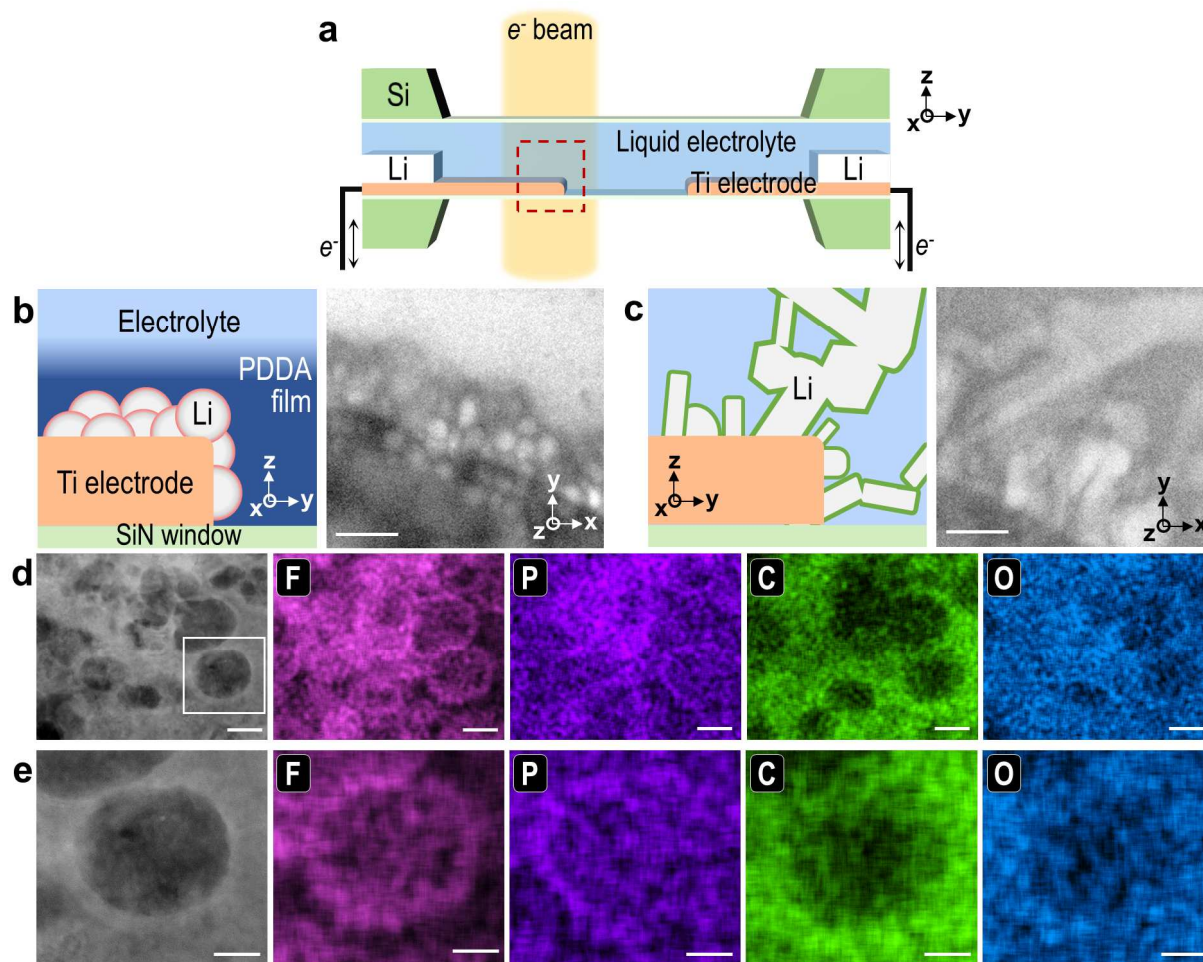
Increasing demands for energy storage devices with much higher energy density have attracted a revival of interest in lithium metal anodes for future rechargeable batteries.[1, 2] However, it remains a great challenge to use lithium metal as an anode because of safety issues associated with lithium dendrite formation, which may result in short-circuiting of the batteries.[3] In addition, the formation of “dead lithium” during the stripping cycles, which refers to fragmentation of lithium deposits detached from the electrode, may also result in capacity loss.[4-6] These challenges have led to much research effort aimed at understanding and controlling lithium dendrite growth.[3, 7] For example, the formation of large lithium dendrites under a polymer electrolyte has been observed using x-ray techniques.[8] Moreover, the atomic structures of lithium dendrites and solid-electrolyte interphase (SEI) has been revealed by cryo-electron microscopy.[9, 10] Recently, with *in situ* electrochemical liquid cell transmission electron microscopy (TEM), lithium nanogranules instead of lithium dendrites were obtained through modifying the SEI composition by applying a thin cationic polymer film on the electrode.[11] Extensive studies have shown that SEI on lithium metal deposits plays a key role in transforming the lithium growth behavior.[12-14] In contrast, there have been limited studies on the stripping mechanisms of lithium, even though it is as important as lithium plating for improving the performance of lithium metal batteries.[15-21] For example, fast accumulation of voids under high stripping rates,[16] different lithium plating/stripping behaviors from initially plated and stripped lithium electrode,[17] and critical stripping current density causing lithium dendrite formation during the following cycle[18] were investigated at microscales by scanning electron microscopy studies. Previous studies suggested potential models of dead lithium formation, however, the limited spatial resolution hinders the in-depth understanding of lithium stripping mechanisms and the relevant dead lithium formation. In addition, although an elaborate *in-situ* optical microscopy showed intact SEI after the lithium stripping,[19] providing a very rare study on lithium SEI, technical

advances allowing further quantitative interpretations of SEI behaviors during the lithium stripping are needed.

Here, we investigate lithium stripping behavior using a state-of-the-art *in-situ* electrochemical liquid cell TEM.[22] The *in-situ* liquid cell TEM allows direct observation of the electrochemical plating/stripping processes of lithium metal in a common liquid electrolyte for lithium ion batteries. We compare the stripping process of lithium dendrites and lithium nanogranular deposits. We focus in particular on dead lithium formation during lithium stripping. By correlating the lithium morphology evolution during stripping with the chemistry of SEI, lithium stripping mechanisms are elucidated.

## **2. Results and Discussion**

### **2.1. Electrochemical liquid cell TEM experiments for lithium stripping studies**

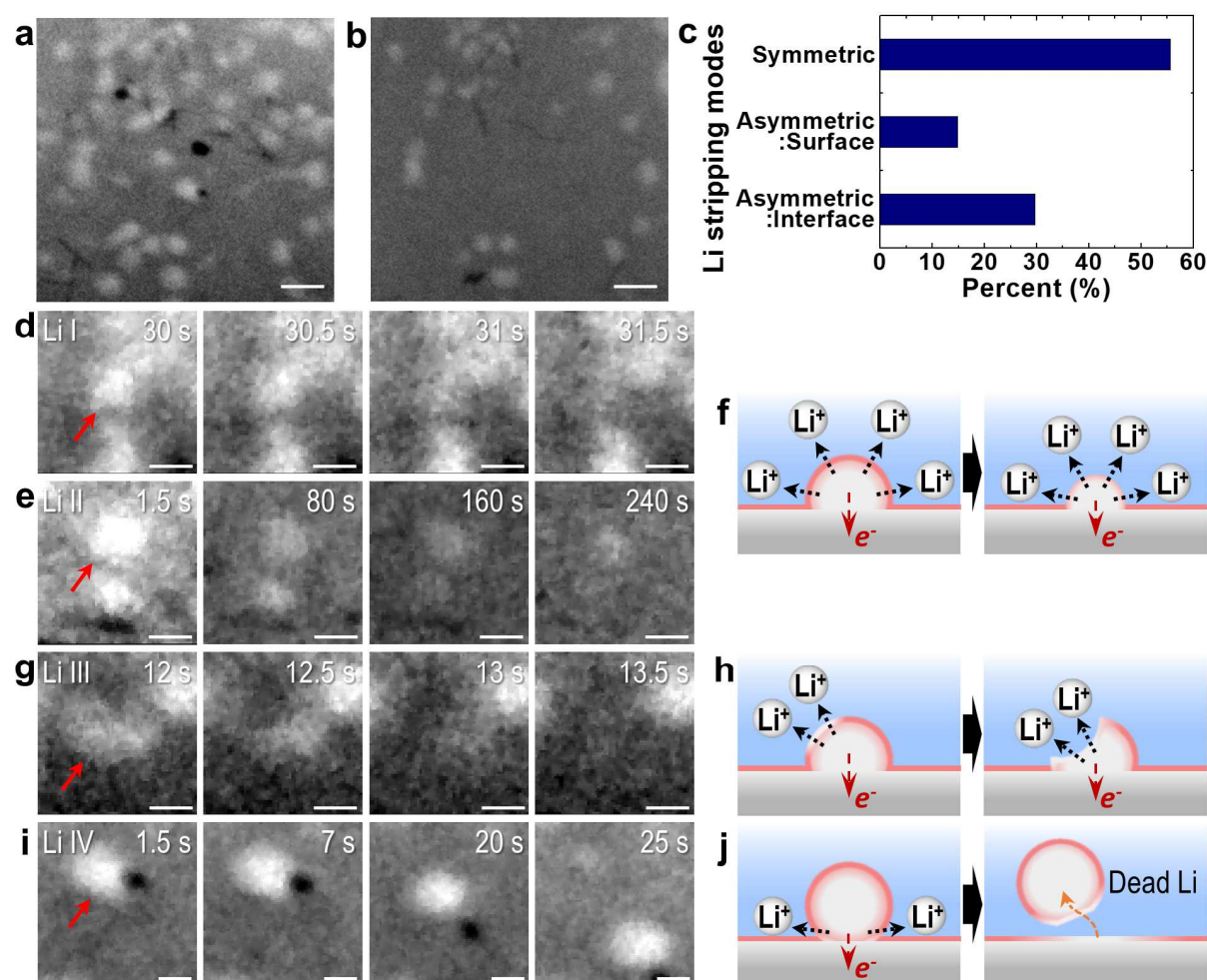


**Figure 1.** Schematic drawings of electrochemical liquid cell transmission electron microscopy (TEM) experimental designs for Li stripping studies and chemical images of *in-situ* grown lithium deposits with LiF-rich solid-electrolyte interphase (SEI). (a) A schematic drawing of an electrochemical liquid cell for *in-situ* TEM lithium plating/stripping experiments. (b, c) Illustrations (left panels) and TEM images (right panels) of the *in-situ* grown (b) lithium nanogranules with LiF-rich SEI and (c) lithium dendrites at the area corresponding to the red square in (a). Scale bars are 500 nm. (d) An annular dark-field (ADF) scanning TEM (STEM) image and energy-dispersive x-ray spectroscopy (EDS) elemental maps of lithium nanogranules with LiF-rich SEI. Scale bars are 200 nm. (e) A magnified ADF-STEM image and EDS elemental maps of a lithium nanogranule marked in (d). Scale bars are 100 nm.

The experimental setup of the electrochemical liquid cell TEM for *in-situ* lithium plating/stripping is shown in **Figure 1a-c**. A liquid electrolyte (1M LiPF<sub>6</sub> in propylene carbonate (PC)) was filled in a Si/SiN<sub>x</sub> electrochemical nano-cell where titanium was deposited as current collectors for the working/counter electrodes (Figure 1a).[23] Lithium flakes were attached on the rear side of the both Ti electrodes to allow reversible electrochemical reactions and supply lithium ions in case there is no source at the front lines.

Experimental details are described further in the *Materials and Methods* section. Potential electron beam effects during the *in-situ* lithium plating/stripping experiments were comprehensively investigated and elaborated in Supplementary Material (Supplementary Note 1 and Figure S1). We studied stripping behaviors of lithium deposits grown *in-situ* in the electrochemical liquid cells with dendritic or nanogranular structures, as shown in Figure 1b and 1c. The different morphologies of the lithium deposits were achieved by modifying the electrode environment with a poly(diallyldimethylammonium chloride) (PDDA) cationic polymer film coating. As elaborated in our previous study[11], by applying a thin layer of the PDDA cationic polymer film on the electrode,  $\text{PF}_6^-$  salt anions are attracted to the electrode surface by electrostatic force, promoting the formation of a lithium fluoride (LiF)-rich SEI on the lithium deposits. The LiF-rich SEI inhibits dendritic growth of lithium, resulting in lithium nanogranular growth instead.[24, 25] The LiF-rich SEI on lithium nanogranules can be identified in the annular dark-field (ADF) scanning TEM (STEM) images and corresponding EDS elemental maps (Figure 1d and 1e). An additional ADF-STEM image and EDS elemental maps at lower magnification are provided in Supplementary Material (Figure S2) for better understanding of the experimental results. The ADF-STEM image in Figure 1d shows a cluster of lithium nanogranules. The darker contrast of the nanogranules is due to the low atomic number of lithium, resulting in lower scattering angles of the incident electrons compared to those from other elements, e.g. carbon, oxygen, fluorine, phosphorus, etc.[26] In the EDS maps, both of fluorine and phosphorus are concentrated on the surface of lithium nanogranules but fluorine is covered much more uniformly than phosphorus. Carbon and oxygen are broadly distributed in the outer layer of SEI and the residual electrolyte around the lithium nanogranule. The LiF-rich SEI on lithium nanogranules compared with lithium dendrites was confirmed by x-ray photoelectron spectroscopy (XPS), as shown in our recent publication.[11]

## 2.2. Stripping modes of individual LiF SEI-rich lithium nanogranules



**Figure 2.** Stripping modes of lithium nanogranules on top of the titanium electrode with LiF-rich SEI. (a, b) TEM images of the electrode area captured (a) before and (b) after the *in-situ* lithium stripping (Video S1). The images are a top view of a portion of the Ti electrode. Scale bars are 500 nm. (c) Relative frequency distribution of each stripping mode for lithium nanogranules with LiF-rich SEI. (d, e, g, i) Sequential TEM images of *in-situ* stripping and (f, h, j) corresponding schematic illustrations of lithium stripping modes: (d, e, f) symmetric (Video S2 and S3), (g, h) surface-preferred asymmetric (Video S4), and (i, j) interface-preferred asymmetric stripping modes (Video S5). Scale bars are 200 nm.

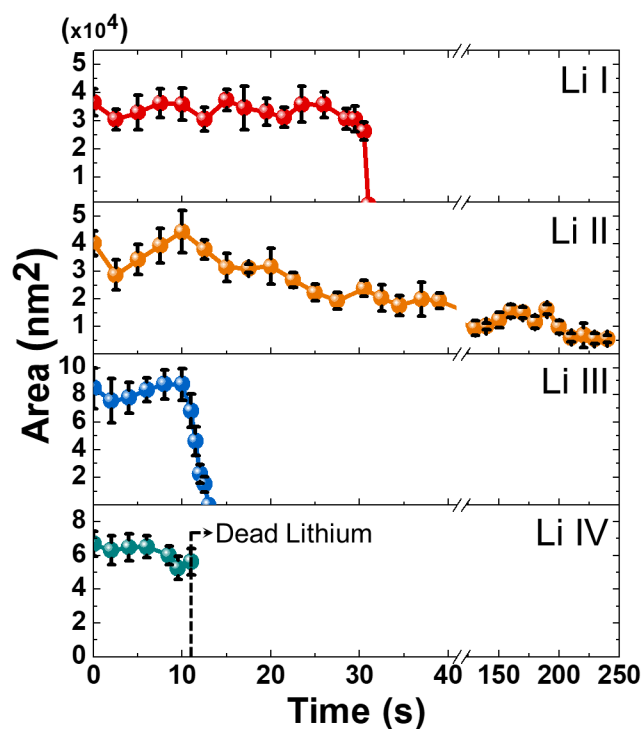
TEM images of lithium nanogranules with LiF-rich SEI on the Ti electrode before and after the *in-situ* lithium stripping are shown in **Figure 2a** and **2b** (also see Video S1). The TEM images show part of the Ti electrode viewed from above with lithium metal deposits, bright contrasts in the bright-field TEM images, laying on top of the Ti electrode. The black nanowires and nanoparticles are Sn@SnO<sub>2</sub> nanostructures, which were used to assist PDDA

cationic polymer coating in the electrochemical liquid cell.[11] The Sn@SnO<sub>2</sub> nanostructures did not contribute to the electrochemical reactions, probably due to their high contact resistance in the electrochemical liquid cell, which can be confirmed by the fact that their volume did not change.[11] Most lithium nanogranules on the electrode were stripped after applying a positive electric potential for a period of time, as shown in Figure 2a and 2b. Control experiments ruled out potential effects of the electron beam on the lithium stripping (Figure S1). Different stripping behaviors were observed for individual lithium nanogranules grown simultaneously under the same experimental condition as shown in Figure 2c-j.

We found three stripping modes of individual lithium nanogranules (Figure 2c): (i) symmetric stripping (Li I, II; Figure 2d-f), (ii) surface-preferred asymmetric stripping (Li III; Figure 2g,h), and (iii) interface-preferred asymmetric stripping (Li IV; Figure 2i,j). A number of factors may have contributed to the observed different stripping behaviors of lithium nanogranules, such as variations in the local SEI conditions, different morphology of lithium deposits, local inhomogeneity of the surrounding electrolyte, the status of electrical contact, etc. In the symmetric stripping mode, the lithium nanogranule shrinks inward radially, preserving its round shape. The corresponding uniform decreases of size and contrast are clearly visible in Figure 2d and 2e (also see Video S2 and S3). In contrast to the symmetric lithium stripping, surface-preferred asymmetric stripping is defined as the case that the stripping starts from a certain side position at the surface of lithium nanogranule. Figure 2g shows an example of this surface-preferred asymmetric stripping mode (also see Video S4). The upper-left corner of the lithium nanogranule is first stripped at 12.5 s. Interestingly, stripping continues rapidly from a neighboring position (at 13 s, Figure 2g). This shows that the collapsed SEI layer at the lithium-stripped position may ease further lithium stripping through that position (Figure 2h). The surface-preferred asymmetric stripping mode was not frequently observed (Figure 2c), which shows that most of the lithium nanogranules were in a fairly homogeneous environment. We also note that lithium stripping triggered at specific



points does not always lead to the rapid complete stripping of lithium deposits, as partially stripped lithium nanogranules were observed after the *in-situ* stripping as shown in Figure S3. The interface-preferred asymmetric stripping mode refers to preferential lithium stripping at the interface between the lithium deposit and the electrode (current collector), which contributes significantly to “dead lithium” formation. Figure 2i shows an example of the interface-preferred asymmetric stripping mode (also see Video S5). After several seconds of stripping, the lithium nanogranule marked in Figure 2i became dead lithium floating in the liquid electrolyte. This indicates preferential lithium stripping at the Li/electrode interface resulting in the separation of the lithium deposit from the electrode (Figure 2j), which is distinct from coincidental desorption of lithium nanogranules from the electrode as further described in Supplementary Material. It could be influenced by a higher energy of the electrolyte-lithium-electrode triple point than that of other surface points, which might explain why there are twice as many interface stripping events as surface stripping events (Figure 2c). It shows a drawback of tiny lithium deposits (e.g. nanogranules) with small contact areas to the electrode, which leads to higher chance of losing the electrical connection during their stripping, resulting in the dead lithium formation.



**Figure 3.** A comparison of lithium nanogranule stripping rates. Size reduction of individual lithium nanogranules labeled as Li I (Figure 2d; symmetric stripping), Li II (Figure 2e; symmetric stripping), Li III (Figure 2g; surface-preferred asymmetric stripping) and Li IV (Figure 2i; interface-preferred asymmetric stripping).

Stripping rates of individual lithium nanogranules are compared in **Figure 3**. Area changes of the individual lithium nanogranules shown in Figure 2d (Li I; symmetric stripping), Figure 2e (Li II; symmetric stripping), Figure 2g (Li III; surface-preferred asymmetric stripping), and Figure 2i (Li IV; interface-preferred asymmetric stripping) were tracked as a function of time. The measurement details are further discussed in Supplementary Material. The plot shows that the stripping kinetics of individual lithium deposits, including reaction initiation time and reaction speed, are heavily influenced by local environments. Mostly, lithium nanogranules were stripped very rapidly following the initiation, as shown in cases of Li I and Li III. The stripping reactions were completed within several seconds. However, some lithium nanogranules had a considerably slower stripping rate. As shown in cases of Li I and Li II, there is a significant difference of the lithium stripping rate even in the same stripping mode.

Note that the plot for Li IV shows there was almost no area change of Li IV before it lost the contact with the electrode and became the dead lithium.

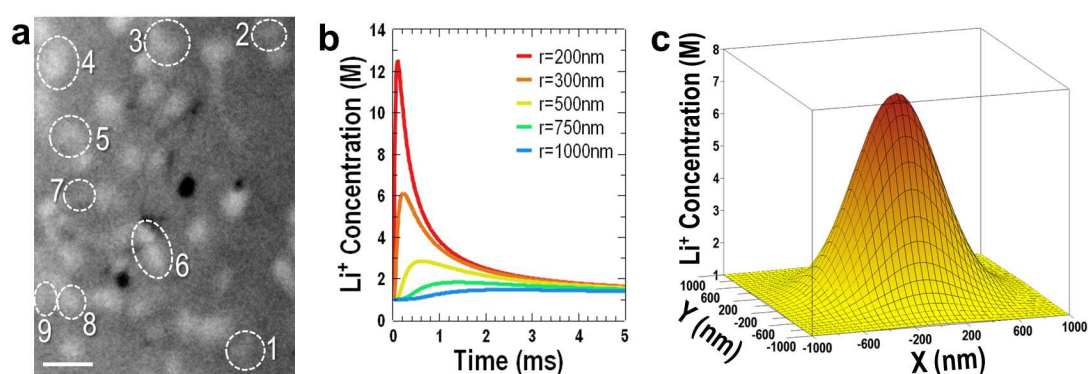
### 2.3. Sporadic stripping behavior of lithium metal deposits

We also found an interesting overall lithium stripping behavior; neighboring lithium deposits are stripped sporadically rather than simultaneously. In other words, lithium deposits in immediate proximity to the rapidly stripped lithium deposit are not stripped immediately. This behavior is shown in **Figure 4a**. The numbers labeled on the TEM image indicate the stripping sequence of individual lithium nanogranules in about  $10 \mu\text{m}^2$  area. The start and finish times of each stripping event for different lithium nanogranules are shown in Table 1. TEM images before and after each stripping event are shown in Figure S5. The labeled lithium deposits were stripped rapidly, but their neighboring ones were not stripped immediately, which was generally observed regardless of the symmetric or asymmetric stripping behavior of individual lithium deposits (Video S1). We propose that the rapid stripping of one lithium deposit impedes stripping of adjacent deposits by increasing the local concentration of lithium ions. The lithium ion concentration changes as a result of a single lithium deposit stripping are shown in Figure 4b and 4c. These were calculated using Equation (1), assuming that the rapidly stripped lithium deposit is an instantaneous point source for the lithium-ion diffusion.[27]

$$C(x, y, t) = \frac{M}{4\pi DL_z t} e^{-\frac{x^2+y^2}{4Dt}} + C_0 \quad (1)$$

In Equation (1),  $C(x, y, t)$  refers to the  $\text{Li}^+$  concentration of position  $(x, y)$  at time  $t$ ,  $M$  is total mol number of the  $\text{Li}^+$  point source,  $D$  is the  $\text{Li}^+$  diffusion coefficient,  $L_z$  is the thickness of the liquid cell compensating the missing dimension, and  $C_0$  is the base  $\text{Li}^+$  concentration from the electrolyte. The calculation details are described further in Supplementary Material. As shown

in Figure 4b, stripping of a lithium deposit causes instantaneous great increases of  $\text{Li}^+$  concentration in its neighborhood. Figure 4c shows that the  $\text{Li}^+$  concentration at positions about 500 nm away can be doubled within 0.5 milliseconds after the stripping of the lithium deposit. This increased local  $\text{Li}^+$  concentration may increase the potential required for the stripping of these adjacent lithium deposits and slow down their dissolution rates momentarily, facilitating the stripping of lithium deposits elsewhere.

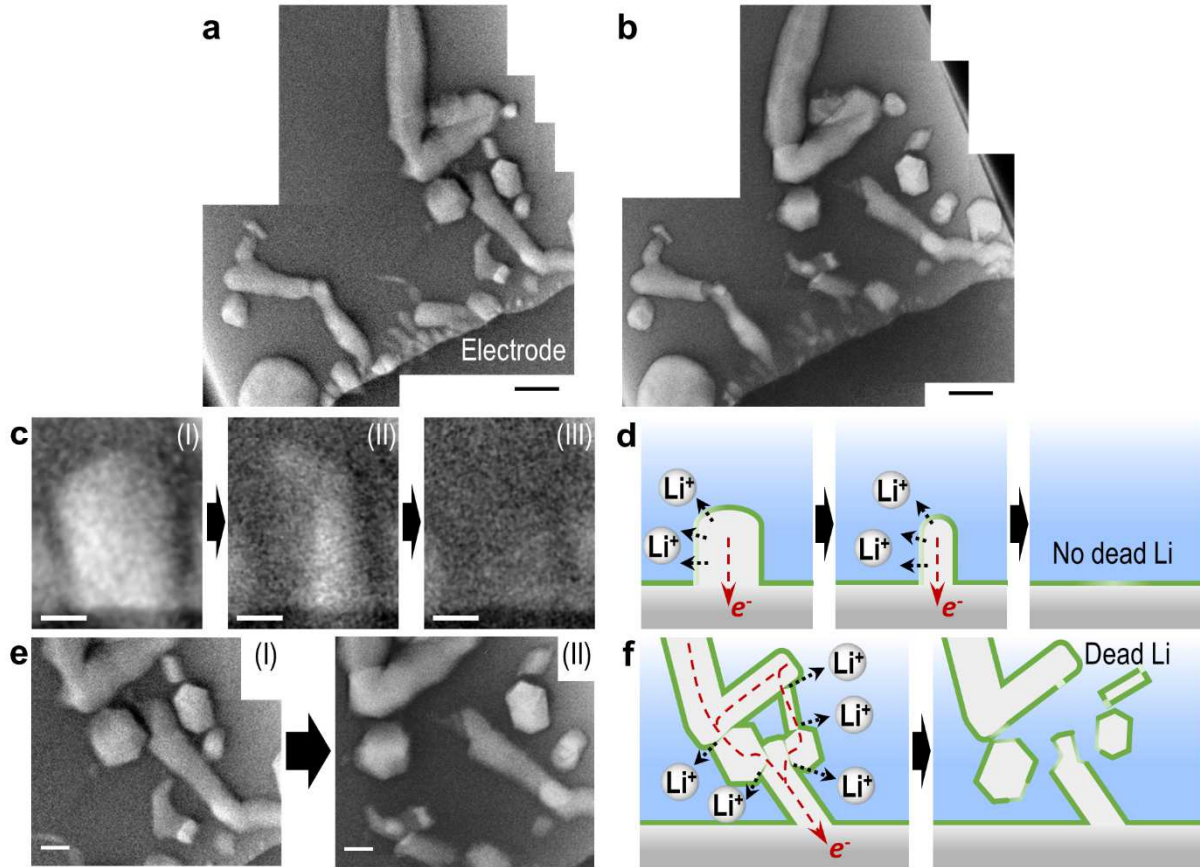


**Figure 4.** Spatially scattered stripping behavior of lithium deposits. (a) A TEM image marked with the stripping sequence of lithium nanogranules and their positions. The image is a top view of a portion of the Ti electrode. Scale bar is 500 nm. (b, c) Estimation of surrounding Li-ion concentration changes as a result of a single lithium deposit stripping. (b) Calculations of Li-ion concentration changes with time at certain distances (denoted as “r”) from a stripped lithium deposit. (c) Calculations of Li-ion concentration distribution around a lithium deposit after 0.5 milliseconds of its stripping.

**Table 1.** Stripping start and end times of individual lithium nanogranules labeled on a TEM image in Figure 4a.

Li deposit	Stripping start time [s]	Stripping end time [s]
1	4.0	6.0
2	5.5	7.0
3	7.5	9.5
4	11.5	13.0
5	17.0	18.0
6	21.0	22.5
7	25.0	25.5
8	30.0	31.0

## 2.4. Stripping modes of individual lithium dendrites

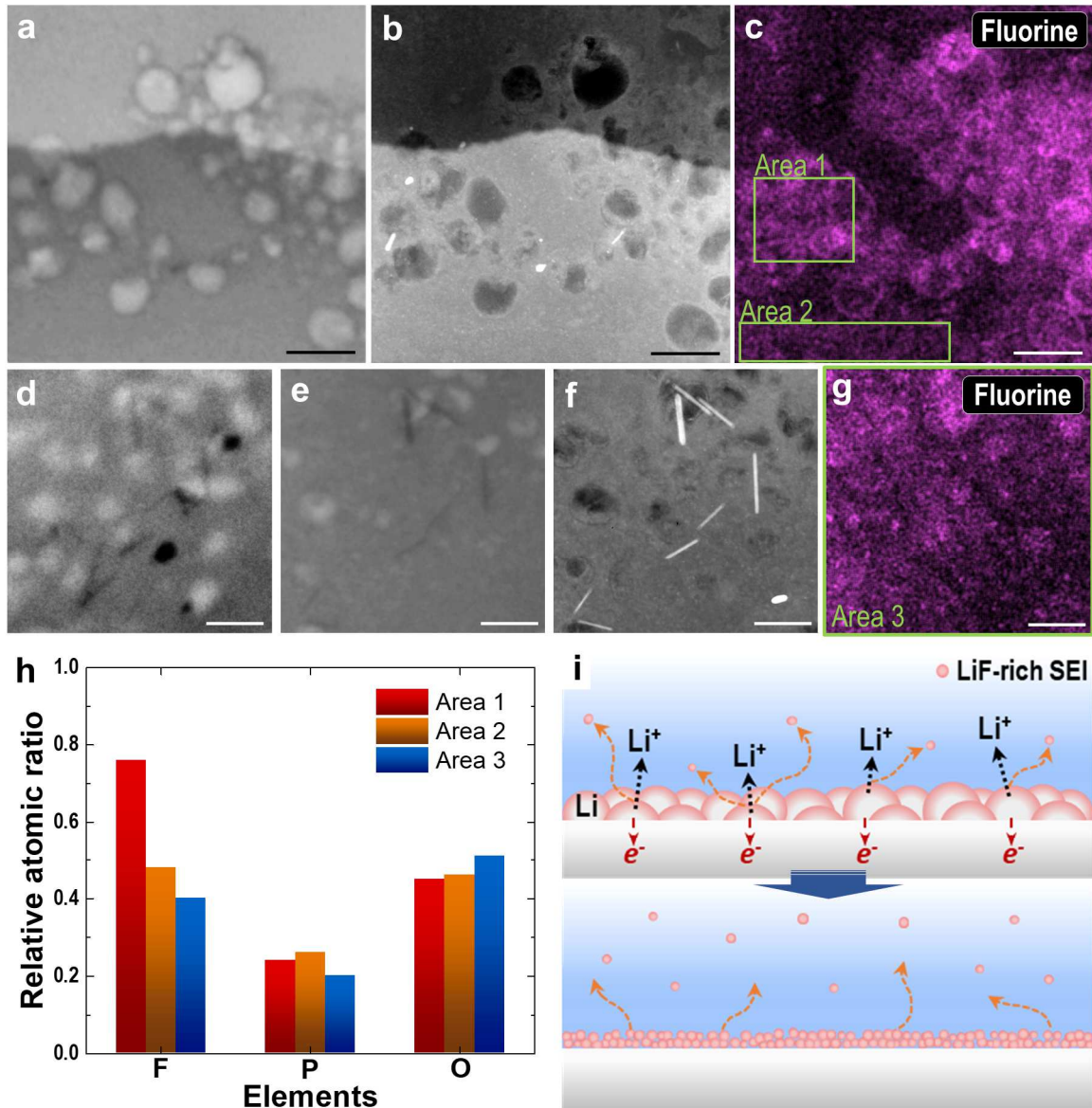


**Figure 5.** Stripping modes of lithium dendrites. (a, b) TEM images captured (a) during *in-situ* lithium plating and (b) after *in-situ* lithium stripping (Video S6). (c, e) Sequential TEM images of stripping and (d, f) corresponding schematic illustrations of lithium stripping modes: (c, d) surface-preferred (Video S7) and (e, f) interface-preferred asymmetric stripping modes. Scale bars: (a, b) 1 μm, (c) 200 nm, (e) 500 nm.

We also investigated stripping behaviors of typical lithium dendrites and found that the stripping behaviors of lithium dendrites share many similarities with those of lithium nanogranules. **Figure 5a** and **5b** show TEM images of the lithium dendrites before and after the *in-situ* stripping experiment. The lithium dendrites were *in-situ* plated and stripped under the same experimental conditions as those for the lithium nanogranular growth but in the absence of the polymer film coating. Detailed stripping modes of lithium dendrites are shown

in Figure 5c-f. We found that the stripping modes of lithium dendrites follow those of lithium nanogranules. Figure 5c shows the stripping of a short lithium dendrite with a length of 700 nm and a width of 400 nm. We note that rapid lithium self-discharge contributed to the stripping of the lithium deposit from stage (I) to (II) in Figure 5c, as further discussed in the Supplementary Material (Figure S6). The small lithium deposit was stripped preferentially from the left side by the surface-preferred asymmetric stripping mode. The lithium deposit was fully stripped within one second during the *in-situ* lithium stripping (see Video S7). This shows that short and straight lithium deposits can be stripped easily even under the asymmetric stripping mode, as shown in Figure 5d. However, majority of lithium deposits have long branched dendrite structures, which preferentially separate at branch-branch interface during the stripping. This causes significant dead lithium formation. This asymmetric stripping of the branched dendrite is displayed in Figure 5e. This might have originated from the SEI inhomogeneity at the Li/Li interface that can lead to the easier lithium-ion extraction at this position. Or, it could be due to the higher energy of the interface compared to other positions, which eases the charge transfer reaction. This stripping process is schematically illustrated in Figure 5f. It clearly shows a critical origin of dead lithium formation from lithium dendrites, which had not been demonstrated before.

## **2.5. Impact of lithium stripping on SEI layer**



**Figure 6.** A comparison of SEI layers before and after the lithium stripping. (a) A TEM image, (b) a corresponding ADF-STEM image, and (c) a corresponding EDS fluorine map of *in-situ* plated lithium nanogranules with LiF-rich SEI. Lower half area with different contrasts in the TEM/STEM images corresponds to Ti electrode. (d, e) TEM images of the electrode area captured (d) before and (e) after *in-situ* lithium stripping. The images are a top view of a portion of the Ti electrode. (f) An ADF-STEM image and (g) an EDS fluorine map of *in-situ* lithium stripped area corresponding to (e). Bright objects in the STEM image are assistant nanorods used for the polymer film coating. All scale bars are 500 nm. (h) Atomic ratios of fluorine ( $n_F/n_{Ti}$ ), phosphorus ( $n_P/n_{Ti}$ ), and oxygen ( $n_O/n_{Ti}$ ) relative to titanium electrode at selected areas marked in (c) and (g), calculated from the EDS results. (i) Schematic illustration of stripping behaviors of lithium deposits and their SEI layers.

We further trace the impact of lithium stripping on SEI layer. This is essential to determine whether the SEI formed in the previous cycle can be beneficial for the following cycle. With

the stripping of lithium nanogranules as an example, SEIs on the electrodes before and after stripping were measured using STEM-EDS. The EDS fluorine maps before and after the stripping of lithium nanogranules are shown in **Figure 6** and additional EDS elemental maps are shown in Figure S7. Figure 6a-c shows *in-situ* plated lithium nanogranules and corresponding EDS fluorine map. Consistent with the EDS elemental maps in Figure 1d, fluorine-rich SEI is clearly visible for individual lithium nanogranules (Figure 6c). There is an obvious concentration difference of fluorine between areas with and without lithium nanogranules. Figure 6d and 6e show TEM images obtained before and after the *in-situ* lithium stripping experiments. Most lithium nanogranules were stripped leaving only small remains and traces behind, shown as dark contrast at the corresponding positions in the ADF-STEM image (Figure 6f). Interestingly, the fluorine concentration differs between areas with and without the remaining lithium nanogranule after the stripping (Figure 6g).

The SEI concentration ratio before and after lithium stripping were quantitatively compared by EDS spectra collected from different electrode areas: (i) densely covered by plated lithium nanogranules (Area 1 in Figure 6c), (ii) free from lithium after the lithium plating experiment (Area 2 in Figure 6c), and (iii) after lithium stripping (Area 3 in Figure 6g). Figure 6h shows the relative atomic ratios of representative elements of the SEI, i.e. fluorine ( $n_F/n_{Ti}$ ), phosphorus ( $n_P/n_{Ti}$ ), and oxygen ( $n_O/n_{Ti}$ ), scaled relative to titanium for the electrode as reference. For example, in Area 1, the concentration of fluorine is equivalent to 76% of Ti concentration. As titanium electrode is uniformly deposited on the bottom chip of each liquid cell and it is electrochemically inactive during the *in-situ* experiment, the EDS signal of titanium concurrently measured at each area can be a good reference for the quantitative comparison. There is little difference on the amount of phosphorus and oxygen at all areas, however, the fluorine concentration significantly decreased on the electrode after the lithium stripping (Area 1 vs. 3). The fluorine amount after the lithium stripping (Area 3) is



comparable to that of the electrode area where lithium was not deposited during the lithium plating experiment (Area 2). This indicates that the collapsed SEI layers (mostly LiF in this case) drift away in the electrolyte during lithium stripping. These results suggest that “good SEI” formed during the prior lithium plating step will be hardly beneficial to the next cycle unless it is tightly fastened by other supporting materials. The SEI behavior during lithium stripping is schematically drawn in Figure 6i. Our discovery of the substantial SEI loss, possibly during every cycle, provides valuable information for the future development of lithium metal batteries. LiF within SEI has been known to be promising for lithium dendrite suppression and the high Li/LiF interfacial energy is considered an important factor.[28] However, the high interfacial energy can also lead to small spherical lithium deposits with excess SEI rendering them sensitive to SEI loss every cycle. This illustrates that LiF-rich SEI conditions must be carefully designed to induce large and flat lithium growth for minimizing SEI areas while suppressing dendrite formation. It furthermore will reduce the interface-preferred lithium stripping and the severe SEI fragmentation, both of which will be beneficial to the coulombic efficiency. In this regard, rigid artificial SEIs or host structures, which transform the lithium growth behavior,[29-31] would also be helpful to seize SEIs during lithium stripping and encourage their functionality in the next cycle.

### 3. Conclusion

In conclusion, we revealed the stripping mechanisms of lithium deposits at the nanoscale by *in-situ* liquid cell TEM. It was found that lithium deposits are sporadically stripped and the stripping of individual lithium deposits can be described by one of the three modes regardless of their morphology as follows: (i) symmetric stripping, (ii) surface-preferred asymmetric stripping, and (iii) interface-preferred asymmetric stripping. Importantly, we found that the interface-preferred asymmetric stripping greatly contributes to the formation of dead lithium

from typical lithium dendrites, which has not been understood so far. Furthermore, chemical analysis of SEI with exceptional spatial resolution unveiled that SEI fragments would float in the electrolyte rather than stably adhere to the electrode after lithium stripping. It suggests the necessity of rigid protection layers on the electrode to avoid wasting the “good SEI” formed in the prior cycle. This study provides a comprehensive understanding of the stripping behavior of lithium deposits, which is critical for the development of lithium metal batteries.

#### **4. Material and Methods**

*In-situ electrochemical liquid cell TEM experiments:* Details of the titanium electrode-deposited liquid TEM cell fabrication and the Sn@SnO<sub>2</sub> nanostructure-assisted PDDA cationic polymer synthesis and coating methods are described in our previous papers.[11, 32, 33] Lithium flakes were attached onto both Ti electrodes to construct Li/Li symmetric cell. Liquid electrolyte (1M lithium hexafluorophosphate (LiPF<sub>6</sub>) solution in propylene carbonate (PC)) was loaded dropwise into the cell by a pipette. UV-curing adhesives were used to seal the liquid TEM cell reservoirs, which was fully cured within 30 seconds by a UV flashlight. The liquid TEM cell sample was prepared in an argon-filled glove box. The prepared liquid TEM cell was loaded into our customized *in-situ* TEM holder with electric cables extended to the tip of the holder. Protruding electric wires from the liquid TEM cell were bonded to the cables at the tip of the holder by silver conducting paste to establish electrical connection between the liquid TEM cell and an external potentiostat (CH Instruments). For the *in-situ* TEM lithium stripping experiments, lithium metal was pre-deposited in the liquid TEM cell in the TEM column by voltammetry method (a linear sweep potential from 0 to -4V at 0.1 V/s sweep rate, followed by a constant potential bias of -4V), as shown in our previous paper.[11] A positive potential was applied (a linear sweep potential from 0 to +4V at 0.1 V/s sweep rate, followed by a constant potential bias of +4V) to the liquid TEM cells in the TEM column for

the following *in-situ* TEM lithium stripping experiments. Voltage and current profiles are shown in Figure S8 in Supplementary Material. The reactions were recorded at two frames per second (2 fps) by a charge-coupled device (CCD) camera installed in the TEM. Electron beam with very low electron dose rate of  $\sim 0.2\text{--}0.5\text{ e}^-/\text{\AA}^2\text{s}$  was used during the lithium stripping experiments to avoid unexpected effects by the electron beam. For STEM-EDS measurements, post-mortem analysis was performed after opening the liquid TEM cell to acquire high EDS signal. Before opening the liquid cell, liquid electrolyte was solidified on the *in-situ* grown lithium deposits to make a protection layer. The detailed procedure is described as follows. Electron beam with a low but slightly higher electron dose rate of  $\sim 1\text{ e}^-/\text{\AA}^2\text{s}$  was used to irradiate the *in-situ* experimented liquid TEM cell for more than 20 minutes, to polymerize (solidify) the liquid electrolyte.[34] The polymerized electrolyte acts as a protection layer for the underlying lithium deposits and electrodes. Before the STEM-EDS measurements, the top and bottom chips of the liquid TEM cell were carefully separated to obtain more EDS signals. Both sides (top and bottom) of SiN window membranes remained on the bottom liquid TEM cell, which covered the top and bottom sides of the lithium deposits making double protection layers together with the polymerized electrolyte layer. These layers protected the lithium deposits and the electrodes from air exposure before the cell loading into the TEM column. The *in-situ* liquid cell TEM experiments were performed with JEOL JEM-2100 200 kV LaB<sub>6</sub> TEM instrument and the STEM-EDS experiments were performed with FEI Themis 300 kV field-emission TEM instrument equipped with Bruker SuperX EDS detectors.

*Video/image noise reduction:* Some videos and images were processed for noise reduction. For Video S1 (Figure 2a, 2b, 4a, and 6d), four consecutive frames were averaged to reduce random noise. Subsequently, outliers were removed by replacing each pixel with the median of the surrounding two-pixel radius if it deviates from the median by more than 20 (raw unit). Same procedure was performed for Video S2-S5 (Figure 2d, e, g, i), except two consecutive

frames were averaged to reduce random noise. In addition, total variation denoising (also known as ROF denoising) was performed with the regularization parameter  $\lambda=0$ .<sup>[35]</sup> All noise reduction process was performed using Fiji (ImageJ) software.

### **Conflicts of interest**

There are no conflicts to declare.

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# Graphical Abstract

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## Lithium Metal Stripping Mechanisms Revealed Through Electrochemical Liquid Cell Electron Microscopy

