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Achievement of high-cyclability and high-voltage Li-metal batteries by heterogeneous SEI film with internal ionic conductivity/external electronic insulativity hybrid structure

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Abstract

Establishing electronic hinder/ionic transfer pathway in SEI film is key issue for high-performance Li-metal an-odes (LMA), which requires the SEI with high ionic conductivity to enable fast Li^+ diffusion and regulated Li deposition behavior, and poor electronic conductivity to block the electrolyte consumption. Herein, we propose a strategy to construct heterogeneous SEI via selective reduction of electrolytes components to improve Li stability and suppress dendrite growth. The inner N-rich sub-layer of SEI film enables fast Li^+ transportation for nodule-like Li deposition while the outer C-rich sub-layer of SEI film exhibits an electronic insulation property to block electrolyte decomposition. This hybrid SEI endows the LMA with high Coulombic efficiency (99.0%), long lifespan, and dendrite suppression. Theoretical calculations, XPS and AFM were employed to examine the heterogeneous SEI structure and clarify its formation mechanism. A high-capacity retention of 91.6% after 160 cycles at 0.5 C in LiCoO_2/Li pouch cells with ultra-thin Li anodes (25 nm) and low N/P ratio (1.67), and an excellent performance with 85.7% capacity retention after 300 cycles at higher charge potential (4.5 V) was also obtained. The insight in heterogeneous SEI formation provides new opportunities for rational electronic/ionic transfer pathway construction for achieving high-performance Li-metal batteries.

Keywords:

Lithium-metal anode Heterogeneous SEI Internal ionic conductivity External electronic insulativity Lithium dendrites

1. Introduction

Rechargeable lithium metal batteries are a transformative technology that could revolutionize the energy storage sector because of their high theoretical specific capacity (3860 mAh g^{-1}) and low redox potential (-3.04 V vs. standard hydrogen electrode) of the Li-metal anode [1]. The Li-metal anode is capable of coupling with a variety of cathodes, including traditional layered oxides (440 Wh Kg^{-1}), sulfur (650 Wh Kg^{-1}) and oxygen (950 Wh Kg^{-1}), [2] but continues to exhibit two challenging issues: (1) limited cyclability and Coulombic efficiency (CE) due to the continuous interaction between Li metal and electrolyte and (2) formation of uneven dendrites and instable Li/electrolyte interface. The latter results in active Li loss and possible penetration of dendrites through the separator, leading to an internal short circuit of batteries and the possibility of catching fire [3].

Tremendous efforts have been made to address the obstacles in application of Li metal anodes. Strategies such as artificial protective films [4-9], stable porous host skeletons [10-13], electrolytes optimization [14-15], solid-state electrolytes [16-19] etc. were studied to suppress dendrite growth and enhance the cycling performance of Li metal batteries. Electrolyte optimization has proven itself to a simple but effective method to create high-performance Li metal batteries. It was reported that increasing the concentration of Li salt can improve the compatibility between Li and electrolytes [20-21]. However, high viscosity and low ionic conductivities impede the charge and mass transportation, which results in sluggish reaction kinetics and hinders its practical application. Another approach is to adjust the chemical and physical properties of the Li electrolyte interface via film-forming electrolyte components or functional additives [22-25].

A fast Li-ionic transfer ability for SEI film on the Li anodes can lead to a homogeneous Li ion flux and increase the ionic concentration beneath of the SEI film, which aids with uniform Li deposition [26]. While the surface electron leakage would result in the electron acquisition of electrolyte components, which give rise to the further reduction of electrolyte [27]. It is reported that only 2 nm LiF or 3 nm Li_2CO_3 are thick enough to block electron tunneling, which significantly attributes to block pathway for SEI generation, leading to less irreversible capacity loss at initial cycles [28]. Thus, a desirable SEI film for Li metal anodes needs the combination of enhanced ability for Li ion transfer and good passivated properties. However, the investigation of rational design of SEI film with heterogenous electronic insulated/ionic conductive structure is still missing.

Batteries with high-energy-density commercial cathodes (high-voltage LiCoO_2 , etc.) matching with ultra-thin Li anodes can be one of the most promising approaches for the practical application of Li-metal batteries. However, combination of high-voltage tolerance and stable Li interface stability still is the main obstacle. Commercial carbonate electrolytes (i.e. 1 M LiPF_6 in EC/DMC/EMC 1/1/1 v/v/v) deliver high-voltage stable potential window, while poor Li stability [29]. And ester-based electrolytes (i.e. 1 M LiTFSI in DME/DOL) are more compatible with Li, however, a narrow stable potential window limits their application. Tetra ethylene glycol dimethyl ether (TEGDME) is one of the promising candidates for electrolyte solvent in Li-metal-based batteries, which were able to deliver an excellent cycling performance in Li- O_2 batteries with high-voltage tolerance (4.5 V) [2]. Unfortunately, TEGDME cannot form efficient SEI films on the Li anodes due largely to the porous Li structure formation. This leads to the continuous reaction with lithium and the formation of thick SEI film. Consequently, applying a TEGDME-based electrolyte results in a low Li CE and short battery lifetime [30]. Herein, rational design of the SEI film with heterogeneous electronic insulated/ionic conductive structure can remarkably enhance the Li stability of TEGDME-based electrolyte with a major leap in Li CE from less than 20.0% to 99.0%.

In our contribution, we propose a novel strategy to rationally design electronic/ionic transfer pathway in a heterogeneous SEI film via selective reduction of LiNO_3 and EC components in electrolyte, which can efficiently regular Li deposition behavior and suppress interface side-reaction. Owing to the thermodynamically feasible reaction between metallic Li and LiNO_3 , the LiNO_3 additive in electrolyte will fast consumption during repeating cycles. Besides, narrow band gap of LiNO_3 -induced products, Li_xNO_y , cannot prohibit the leaking electron, leading to the constantly electrolyte decomposition. Thus, the performance of the batteries in LiNO_3 -only electrolyte suffers fast degradation. Differently, some typical SEI constitutions, like LiF, Li_2CO_3 etc. has a wide band gap, meaning the poor electronic conductivity, which can be desirable components to passivate SEI surface to prohibit the leaking electron. However, poor ionic conductivity of these components

cannot provide enough ionic transfer pathway to regular the Li ion flux, leading to the dendrite growth, and unstoppable breaking/repairing of SEI film. Aiming to suppress both dendrite and electrolyte consumption, one desirable SEI film can be a heterogeneous hierarchical structure with internal ionic conductivity and external electronic insulation. The internal sub-layer with fast ionic transfer pathway regular the Li deposition and suppress the dendrite, while external sub-layer with electronic insulation prevents the leaking electron and blocks the electrolyte de-composition. To our knowledge, LiNO_3 has a smaller lowest unoccupied molecular orbital (LUMO) energy than ethylene carbonate (EC), meaning the prior reduction of LiNO_3 with Li metal. Thus, we combined the LiNO_3 additive and EC co-solvent to take advantage of the high-voltage stability in TEGDME, leading to a desirable electrolyte system comprised of both Li stability and anti-oxidation properties. The components of internal SEI sub-layer, which dominated by LiNO_3 reduction products, Li_xNO_y or Li_3N , can provide enough Li ion diffusion pathway, and the components of external SEI sub-layer, mainly constituted by EC or Li salt reduction products, Li_2CO_3 , LiF etc. prohibits the decomposition of electrolyte. SEM results illustrate that the cycled Li anodes with a heterogeneous SEI film display nodule-like Li particles with less active Li loss and a tight cycled Li layer (6 μm), which is much thinner than that in the bare TEGDME electrolyte (136 μm) and EC or LiNO_3 -only electrolytes. The design of heterogeneous SEI film allows Li CE in EC/TEGDME-50/50 (v/v) with 0.1 M LiNO_3 additive (with 0.5 M LiTFSI /0.5 M LiBOB as lithium salts) to reach 99.0 %. Stability of electrolyte was also investigated in pouch cell with high-loading LiCoO_2 cathodes (3 mAh cm^{-2}) and ultra-thin Li foils (25 μm), corresponding to a low N/P (negative/positive) ratio of 1.67, that can deliver an excellent cycling performance with a capacity retention of 91.6% after 160 cycles at 0.5 C, holding great potential for practical application. A cycling stability with a capacity retention of 85.7% after 300 cycles was also observed in high-loading (2.5 mAh cm^{-2}) LiCoO_2/Li full cells under high charge voltage of 4.5 V, suggesting desirable Li utilization and anti-oxidation property of EC/TEGDME/ LiNO_3 electrolyte. These results demonstrate that the heterogeneous SEI structure with different electronic/ionic diffusion abilities can be generated according to the selective reduction of electrolyte components, leading to the combination of anode stability and high-voltage tolerance.

2. Results and discussion

2.1. The mechanism of heterogeneous SEI film formation

Tuning of SEI structure could create high-performance Li anodes and excellent stability in Li-metal-based full cells configuration. In order to consider the high oxidation potential window and Li anode stability, TEGDME was applied to serve as the bulk solvent of electrolyte to acquire high-voltage electrolyte system, and heterogeneous SEI structure with internal ionic conductivity/external electronic insulation was directionally designed by controlling the reduction sequences of electrolyte components to achieve stable Li electrolyte interface and dendrite-free morphology. Herein, EC and LiNO_3 were adopted as the source of the formation of heterogeneous SEI structure. The LiNO_3 can be reduced prior to the EC, generating a Li_xNO_y -rich SEI near the surface of Li metal to create a fast ion transfer pathway. And EC further is decomposed in the outside shell of SEI film to block the electron transfer and passivate the SEI film.

The density functional theoretical (DFT) calculations were performed to evaluate the reducing activity of electrolyte components. As presented in Fig. 1a, the lowest unoccupied molecular orbital (LUMO) energy of LiNO_3 is -1.421 eV. Thus, another film-forming electrolyte component should have a higher LUMO energy, which enables LiNO_3 to preferentially reduce on Li anodes. EC, a desirable film-forming molecule on anodes, which exhibits a LUMO energy of -0.215 eV, can be reduced after LiNO_3 to form an outer passivated layer on the Li anode. In particular, compared with EC, another film-forming additive, FEC (LUMO energy of -0.8256 eV), displays a higher reduction potential than EC, which would not reduce after LiNO_3 . Besides, main compounds decomposed by FEC is mainly dominated by LiF , which trends to guide filamentous Li morphology. Thus, the EC may be a better choice for this system. The TEGDME-based electrolytes combining both co-solvent EC and additive LiNO_3 were thoroughly investigated. Among several different compositions of electrolyte, 50 Vol% TEGDME, 50 Vol% EC and 0.1 M LiNO_3 additive (EC/TEGDME-50/50+0.1 M LiNO_3) demonstrated the most interesting performance of Li anodes. The Li deposition curves in EC/TEGDME-50/50 and EC/TEGDME-50/50+0.1 M LiNO_3 were

analyzed to investigate the reduction sequence of EC and LiNO₃ (Fig. 1b). The deposition curve in EC/TEGDME-50/50 electrolyte exhibits a plateau around 0.5 V, which can be associated with the reduction of the EC on the Li surface. After adding LiNO₃, besides the EC reduction plateau, a new plateau appears at approximately 0.8 V, corresponding to the reduction of LiNO₃. The higher reduction potential of LiNO₃ indicates that the SEI film formation of LiNO₃ can occur before EC, proving that a heterogeneous SEI film can be directionally generated by selective reduction of the LiNO₃ and EC.

The mechanism of forming heterogeneous SEI film is illustrated by schematic diagrams in Fig. 1c. The internal SEI sub-layer dominated by ionic conductor to provide a fast Li ion transfer channel, which enables uniform Li nucleus formation. For the external SEI sub-layer, electronic insulated constitutions block the electronic transfer, prohibits the electrolyte components capture the electron and be reduced by Li metal. Thus, uniform Li⁺ flux can be realized on the Li surface and guides the Li deposition behavior, leading to a nodule-like Li morphology.

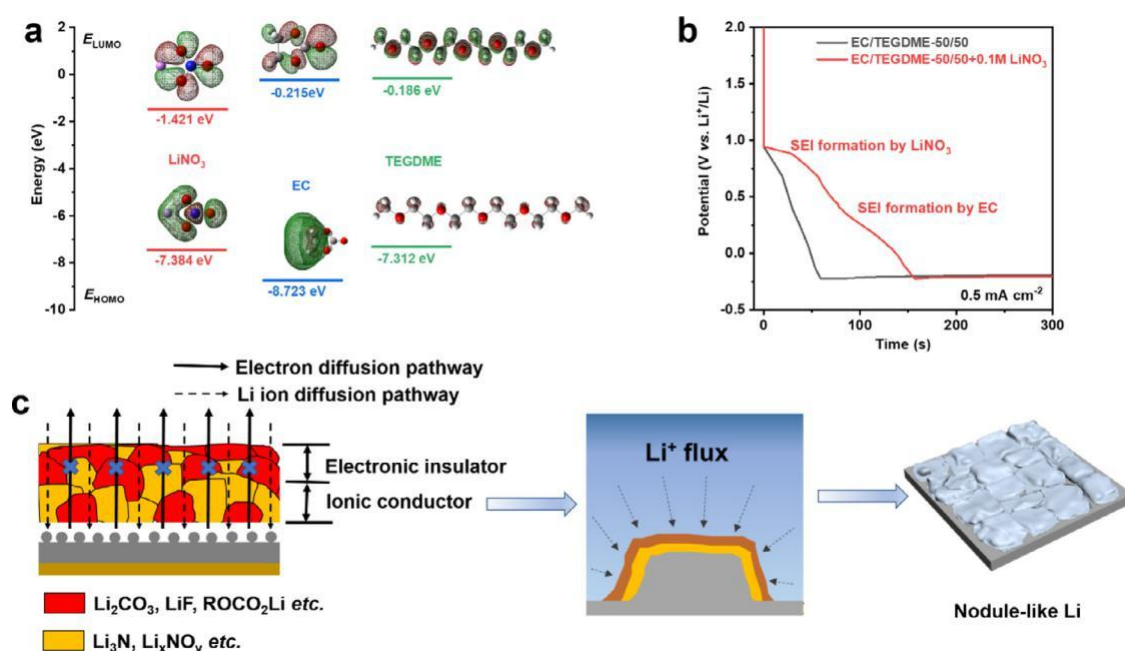


Fig. 1. Schematic diagram of SEI formation. a) Highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) energies and b) the typical Li deposition curves in EC/TEGDME-50/50 and EC/TEGDME-50/50+0.1 M LiNO₃ electrolytes. c) The schematic diagrams for the heterogeneous SEI structure.

2.2. Identification of heterogeneous SEI structure

The chemical and physical properties of SEI films play crucial roles in regulating morphology and elevating the electrochemical performance of Li metal anodes. To analyze the SEI layer composition and structure, the XPS in-depth analysis and Atomic Force Microscope (AFM) were carried on the Li electrode cycled in different electrolytes.

The high-resolution C 1 s spectra, obtained on the surface Li metal cycled in EC/TEGDME-50/50+0.1 M LiNO₃ (Fig. 2a), shows the main components of the SEI layer including peaks corresponding to -C-O-C- bonding, which can corresponds to poly(ethylene oxide) (PEO),

$\text{CH}_2\text{-CH}_2\text{-O})_n$) and/or $\text{R}-\text{C}-\text{O}-\text{CO}_2\text{Li}$ at ~ 286.5 eV, $-\text{C}=\text{O}$ bonding at ~ 288.6 eV and carbonate groups ($-\text{CO}_3$ and/or $\text{RC}-\text{O}-\text{CO}_2\text{Li}$) at ~ 289.6 eV). The presence of these C peaks originates from the electrochemical reduction of EC [31,32], however, the high intensity of carbon peak at ~ 286.5 eV can be related to the reductive decomposition of TEGDME [33]. The presence of hydrocarbon (~ 285 eV (C-C)) can be related to the typical surface contamination usually present on the electrode surface. The in-depth composition (after sputtering of the SEI layer, Fig. 2a) shows the decrease intensity of the higher binding energy peaks (at B.E. higher than 286 eV) and increase of the hydrocarbon peak at 285.0 eV. Moreover, a new peak appears at lower binding energy (of around 283.5 eV), which can be attributed to carbides (C-Li). [34,35]

The SEI layer carbon-oxygen containing species was also confirmed by analysis of O 1 s peak (Fig. 2b), showing three peaks at ~ 530.5 eV corresponding to $-(\text{C}-\text{O})$, at 532.5 eV to $-\text{C}-\text{O}-\text{C}-$ (including $\text{RC}-\text{O}-\text{CO}_2\text{Li}$ and/or PEO $(\text{CH}_2\text{-CH}_2\text{-O})_n$) and the peak at ~ 534.0 eV, which can be attributed to $\text{poly}(\text{CO}_3)$, Li_2CO_3 and/or $\text{RC}-\text{O}-\text{C}-\text{O}_2\text{Li}$ [32]. The O 1 s peak at 532.5 eV can be also overlaid with the peak corresponding to $\text{N}-\text{O}$ bonding [36]. As discussed in the literature one of the possible SEI layer components formed on Li anode in the presence of carbonate electrolytes can be the polycarbonates, showing the high O 1 s and C 1 s binding energy peaks (at around 534 eV and over 291 eV, respectively). [24,37] However, here, no high binding energy C 1 s peaks were observed, thus the presence of polycarbonates can be discarded (Fig. 2a). Concerning the O 1 s peak, the low binding energy peak at around 529 eV corresponding to Li_2O [38,39] can be also observed. The in-depth analysis (Fig. 2d) shows the enrichment of the bulk SEI layer in Li_2O as well as PEO $(\text{CH}_2\text{-CH}_2\text{-O})_n$ components.

The SEI layer formed on the Li negative electrode cycled in EC/TEGDME-50/50+0.1 M LiNO_3 is also enriched in the N-containing species as shown by N 1 s spectra in Fig. 2c. The N 1 s peak at around 399 eV is quite large (FWHM of more than 2 eV), thus, the presence of different products of LiNO_3 decomposition can be present such as Li_3N and Li_xNO_y as previously demonstrated on Li electrode cycled in FEC/ LiNO_3 electrolyte [40]. As shown there, $\text{Li}_x\text{NO}_y/\text{Li}_3\text{N}$ contributes to formation of uniform lithium deposition and ionic conductivity of the SEI layer [41]. The second N 1 s peak at higher binding energy of around 401 eV can be attributed to LiNO_2 [36]. The in-depth analysis (Fig. 2d) shows the increase of N-like components while a decrease of C-like components (as discussed) above can be observed in the inner part of the SEI layer.

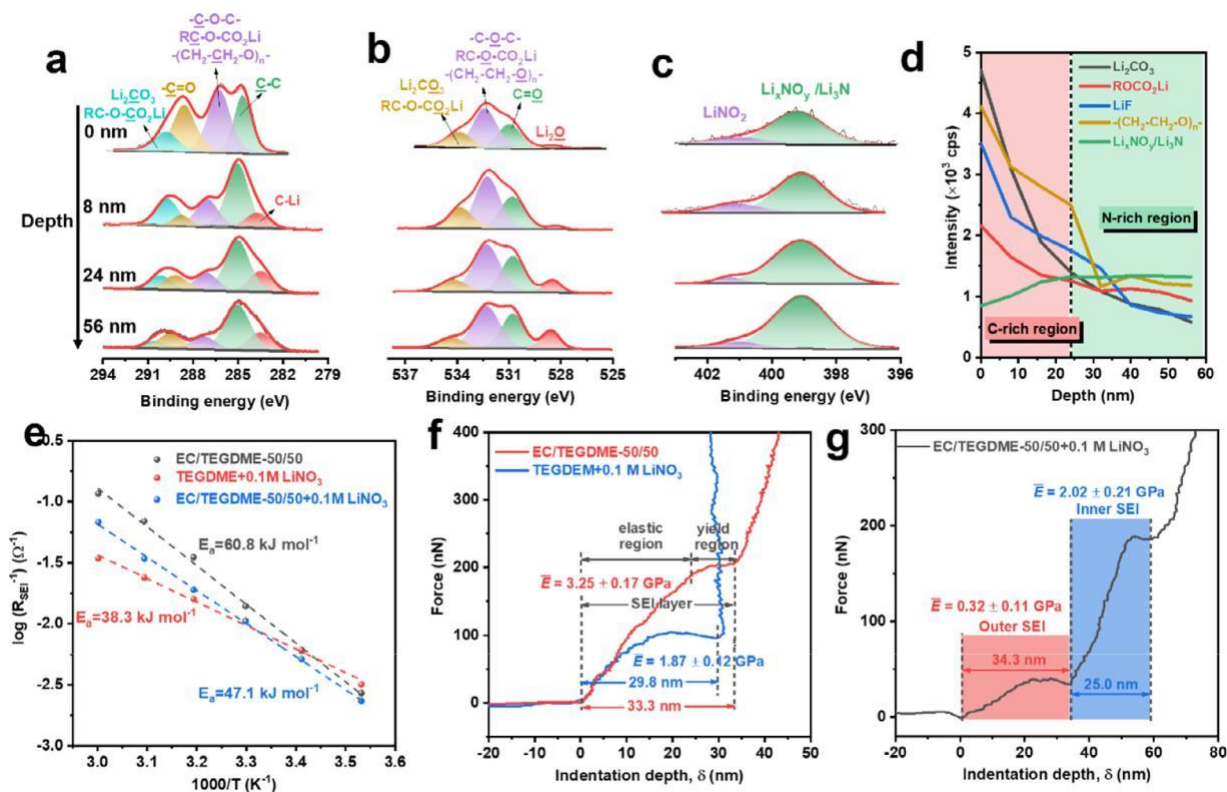


Fig. 2. Chemical and physical analyzation of SEI. The in-depth XPS spectra of SEI layer formed on Li metal cycled in EC/TEGDME-50/50+0.1 M LiNO₃ electrolyte:

a) C 1 s spectra, b) O 1 s spectra, c) N 1 s spectra and d) corresponding depth profiles showing a distribution of SEI components from the surface to the bulk. e) The Arrhenius behavior of the reciprocal and the activation energy (E_a) derived for the ion diffusion through the SEI formed in different electrolytes. The typical force curves obtained by AFM showing mechanical responses of SEI film generated in f) EC/EC/TEGDME-50/50, TEGDME+0.1 M LiNO₃ and g) EC/TEGDME-50/50+0.1 M LiNO₃ electrolytes.

In the carbonate electrolyte system (EC/DMC/EMC), the SEI layer is mainly composed of carbonate and Li-alkyl carbonates (Li₂CO₃/RC-O-CO₂Li) as shown by high intensity of carbon peak at ~289.6 eV (Fig. S1). The presence of high quantity of carbonate and Li-alkyl carbonates on the surface of the negative electrode cycled in carbonate based electrolytes was already confirmed in numerous previous studies. [31,32,42,43] When replaced by the pure TEGDME electrolyte, the lower quantity of carbonates and/or Li-alkyl carbonates can be observed, however, the increased quantity of species composed of -C-O-C-bonding can be clearly seen. Thus, it can be concluded that the SEI layer is principally composed of PEO (CH₂-CH₂-O)_n- and/or RC-O-CO₂Li (Fig. S2).

After introducing LiNO₃ additive into the TEGDME-containing electrolyte, the N 1 s peak can be clearly observed on the surface of the cycled Li electrode (Fig. S3). As it can be observed from sputtering with different times corresponding to different depths a relatively homogenous distribution of N-like components can be observed within the SEI layer (Fig. S3a-d). However, the TEGDME decomposition product as observed from high intensity of C 1 s and O1s peaks corresponding to PEO -(CH₂-CH₂-O)_n- is still significant, indicating that both TEGDME and LiNO₃ participate in the formation of the SEI layer.

A relatively uniform in-depth distribution of the SEI layer can be also observed in the case of EC-only TEGDME electrolyte (EC/TEGDME-50/50) (Fig. S3e-h). The presence of EC shows more important enrichment in carbonate and Li-alkyl carbonates (Li₂CO₃/RC-O-CO₂Li) than it can be observed in the TEGDME-LiNO₃ electrolyte (Fig. S3a-d) and the presence of TEGDME related components is less obvious. It is worth noting by making a comparison of depth profiles obtained in different electrolytes that the unique heterogeneous SEI structure can be observed in EC/TEGDME-50/50+0.1 M LiNO₃ electrolyte (Fig. 2d). As aforementioned LiNO₃ leads to formation of a Li_xNO_y and/or Li₃N component on the Li anode (Fig. 2c); the content of Li_xNO_y and/or Li₃N component gradually increases with sputtering depth (<24 nm) and maintains stable in the inner part of the SEI layer (>24 nm). The in-depth increase of Li_xNO_y and/or Li₃N component is accompanied by the suppression of C-species components. These results are consistent with the selective reduction of LiNO₃ and EC due to their stepwise increase of LUMO energies.

The kinetics and structure of various SEIs formed in several electrolytes were studied through the electrochemical impedance spectra (EIS) of Li/Li symmetric batteries with a temperature range of 10 to 60 °C (Fig. S4). The activation energy (E_a) for Li ion diffusion through SEI films can be calculated using the Arrhenius equation: $R_{SEI}^{-1} = A \exp(-\frac{E_a}{R \cdot T})$. [26,44] A lower activation energy (E_a) corresponds to faster kinetics of Li ion diffusion through the SEI film. This results in a larger Li ion concentration beneath SEI, which is a key factor for the generation of big Li particles instead of Li dendrites [26]. As one can observe from the Fig. 2e, the E_a of Li⁺ ion diffusion through SEI film generated in EC/TEGDME-50/50 electrolyte is 60.8 kJ mol⁻¹, while the E_a of TEGDME+0.1 M LiNO₃ is 38.3 kJ mol⁻¹. These results

indicate that the N-rich SEI film generated in LiNO_3 -contained electrolyte has a higher Li^+ ion conductivity. Consequently, a smaller E_a (47.1 kJ mol^{-1}) was observed in EC/TEGDME-50/50 with LiNO_3 as additive, indicating a higher Li ion concentration beneath SEI, benefiting large Li particle formation.

In an effort to estimate the physical properties of SEI films formed in the target electrolyte, typical force curves showing mechanical responses of SEI films were measured using Atomic Force Microscope (AFM). To prevent the growth of Li dendrites, the SEI films generated on Li surface must to possess a high Yang's module ($>1 \text{ GPa}$) [45]. A typical mechanical response of SEI film can be divided into two regions: elastic region and yield region (Fig. 2f). The Yang's module of SEI film can be obtained with data in the elastic region. [46,47] Yang's module of SEI film formed in TEGDME+0.1 M LiNO_3 ($1.87 \pm 0.12 \text{ GPa}$) is smaller than that formed in EC/TEGDME-50/50 ($3.25 \pm 0.17 \text{ GPa}$), which may be a result of the large amount of stiff inorganic components (Li_2CO_3 , ROCO_2Li etc.) generated by EC component. In contrast, the SEI film formed in EC/TEGDME-50/50+0.1 M LiNO_3 electrolyte displays a typical heterogeneous SEI structure, which corresponds well to the results revealed by XPS (Fig. 2g). The inner sub-layer SEI film with a thickness of $\sim 25.0 \text{ nm}$, shows a Yang's module of approximately 2.02 GPa , associating well with the N-richness. However, in the outer C-rich sub-layer, a soft SEI film ($\sim 0.32 \text{ GPa}$) with a thickness of $\sim 34.3 \text{ nm}$ is detected. This unique soft-rigid heterogeneous SEI structure can collaborate well in regulating Li morphology and suppressing electrolyte decomposition.

To explain outer C-rich and inner N-rich sub-layers of SEI on Li metal anode, we simulated the interactions between electrolyte molecules and Li metal surface by density functional theory (DFT) as shown in Fig. 3. Fig. 3a demonstrates that LiNO_3 molecules undergone strongly chemical reaction with Li metal surface. The outer Li-ions on Li metal surface enter the region of LiNO_3 molecules and come into being amorphous form, and inner Li-ions within Li metal also undergo large displacement. However, Li metal surface with EC and TEGDME molecules can maintain stable molecular geometries and Li-metal configuration, indicating that there is no strong interaction between carbon-based molecules and Li metal. Thus, we hypothesized that inner N-rich layer of SEI (Li_3N , $\text{Li}_x\text{N}_y\text{O}_z$ compounds) are more likely to contribute the high reactivity of LiNO_3 , which is easily reduced with Li metal.

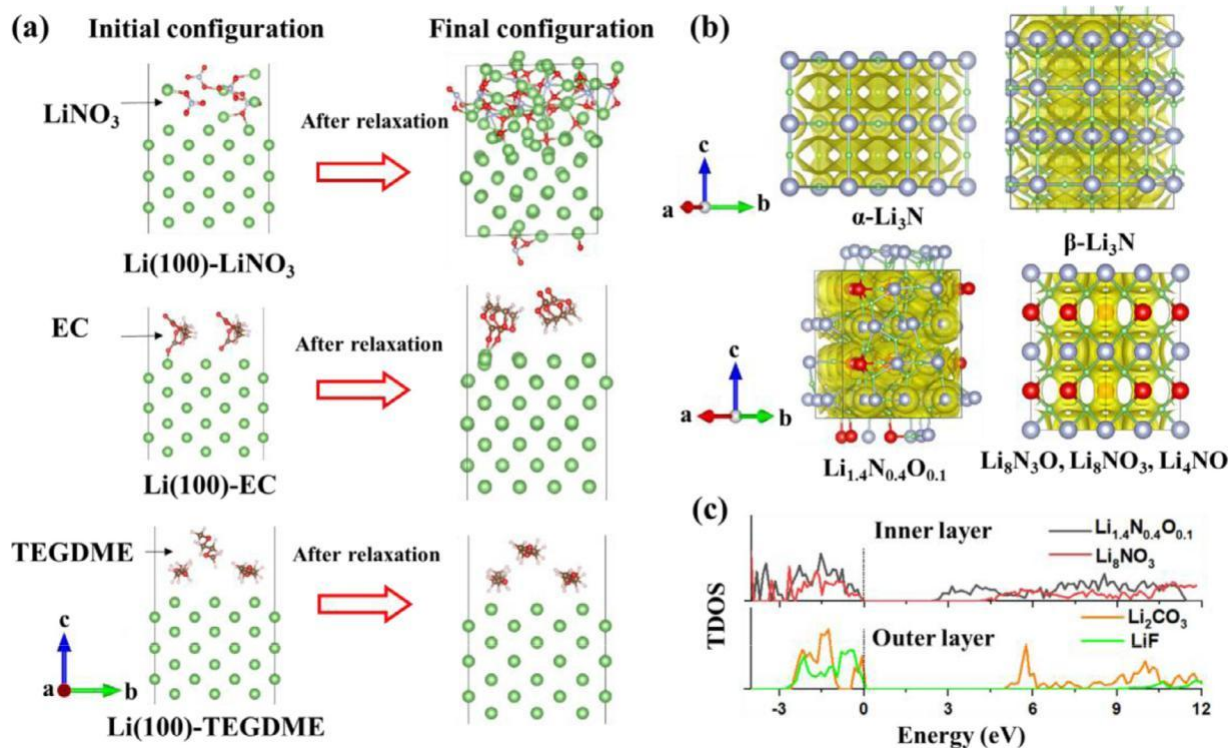


Fig. 3. Theoretical calculation of the SEI properties. a) The interactions of electrolytes and Li metal surface. The stability of electrolytes with Li metal was assessed by relaxing the configurations. The right insets suggested that LiNO_3 molecules undergone strongly interfacial chemical reaction. However, Li metal surface with EC and TEGDME molecules maintain stable after relaxation. b) The potential diffusion pathway in $-\text{Li}_3\text{N}$, $-\text{Li}_3\text{N}$, $\text{Li}_{1.4}\text{N}_{0.4}\text{O}_{0.1}$, and $\text{Li}_x\text{N}_y\text{O}_z$ (Li_8NO_3 , $\text{Li}_8\text{N}_3\text{O}$, and Li_4NO belong to the same space group) illustrated by the bond valence mismatch method. Green, red, while, dark brown, and light blue spheres represent Li, O, H, C, and N atoms, respectively. c) The total density of states (TDOS) of $\text{Li}_{1.4}\text{N}_{0.4}\text{O}_{0.1}$ and Li_8NO_3 . Energies are referenced to the Fermi level.

It has been accepted that Li-ion diffusion barrier governs the structure of the plated/stripped surface, that is, the higher the diffusivity of the Li ions on SEI, the lower the tendency to form Li dendrites. We next checked the Li-ion diffusivity in $-\text{Li}_3\text{N}$, $-\text{Li}_3\text{N}$, $\text{Li}_{1.4}\text{N}_{0.4}\text{O}_{0.1}$, and $\text{Li}_x\text{N}_y\text{O}_z$ compounds (Li_8NO_3 , $\text{Li}_8\text{N}_3\text{O}$, and Li_4NO belong to the same

space group) by bond valence mismatch method as shown in Fig. 3b. All three configurations offer connected Li-ion diffusion channels and short Li-Li distance. More importantly, $-\text{Li}_3\text{N}$ and $-\text{Li}_3\text{N}$ have been proven to be fast Li-ion conductors and can suppress dendrite growth. In Fig. 3c, for the compounds of the inner shell of SEI film, such as $\text{Li}_{1.4}\text{N}_{0.4}\text{O}_{0.1}$ and Li_8NO_3 , have narrow band gap ranging from 2.0 to 4.5 eV. For outer sub-layer of SEI film, such as Li_2CO_3 and LiF , their band gap as large as 5.0 and 9.5 eV, respectively, indicate that they are good electron insulators to avoid the electrolyte components capture the electron and be reduced. Consequently, a heterogeneous SEI structure with high Li-ions conductivity in the inner shell while with electronic insulation in the outer shell is generated.

2.3. Effect of heterogeneous SEI film on Li dendrites and Li loss

The formation of Li dendrites is the one of the main challenges that threatens the practical application of Li metal batteries. To explore the effect of heterogeneous SEI film on Li dendrite formation and Li loss, scanning electron microscopy (SEM) was employed to visualize the evolution of Li metal after 30 cycles at a current density and Li capacity of 2.0 mA cm^{-2} , 2.0 mAh cm^{-2} in different electrolytes. In EC/DMC/EMC electrolyte, Li surface becomes porous and heterogenous after cycling with numerous filament-like Li growth (Fig. S5a) and the thickness of the porous Li layer is $\sim 53 \text{ m}$ (Fig. S5b). The surface of Li foil cycled in the TEGDME electrolyte displays a nodule-like morphology and easily peels off during cycling, resulting a large quantity of dead Li formed on the separator (Fig. S5c, black color). The introduction of LiNO_3 additive into TEGDME electrolyte shows a similar morphology compared with the bare one (Fig. 4a). However, the thickness of porous Li layer decreases from 136 m (Fig. S5d) to 60 m (Fig. 4b), showing the efficient film-forming additive, LiNO_3 , contributes to a compact Li deposition. However, the LiNO_3 -only TEGDME electrolyte cannot avoid the active Li loss during long-term cycling processes as evidenced by the presence of some dead Li deposited on the separator (inserted in Fig. 4b, black color). After replacing the electrolyte with EC/TEGDME-50/50, the Li surface exhibits a hybrid morphology with both filament-like and nodule-like Li growth (Fig. 4c), and the formation of dead Li on separator is effectively suppressed (inserted in Fig. 4c, black color)). The thickness of porous Li layer is reduced to only 38 m (Fig. 4d), proving that the mixture of EC and TEGDME aids in forming a tighter Li layer. Differently, big Li particles with brick-shape are obtained by the heterogeneous SEI film generated in EC/TEGDME-50/50+0.1 M LiNO_3 electrolyte. The Li surface here is much flatter than others, and there is no dead Li formed on the separator (Fig. 4e). Moreover, the thickness of porous Li layer is only $\sim 6 \text{ m}$, embedded with packed Li-particles (Fig. 4f). The larger particle size of Li mitigates an internal short circuit and improve safety and reduces the parasitic interfacial reactions by decreasing the contact area between Li and electrolyte. It allows for a higher Li Coulombic efficiency and longer lifespan of Li metal anodes [48].

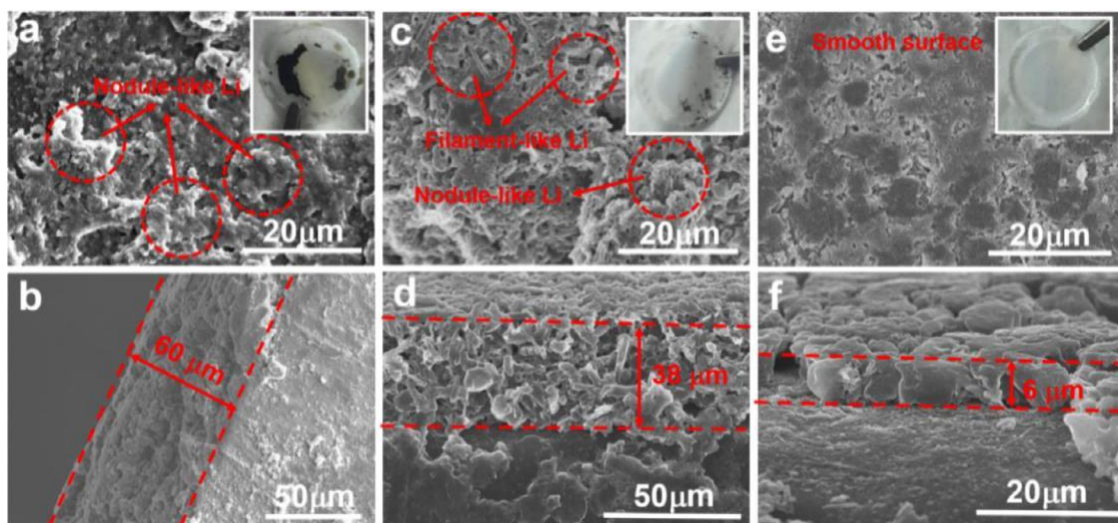


Fig. 4. Elucidation of the morphological evolution of lithium metal anodes. The SEM top- and side-view images of the Li foils cycled in different electrolytes for 30 cycles at 2.0 mA cm^{-2} , 2.0 mAh cm^{-2} : a, b) TEGDME+0.1 M LiNO_3 , c, d) EC/TEGDME-50/50, e, f) EC/TEGDME-50/50+0.1 M LiNO_3 . Corresponding optical photographs of separators after 30 cycles inserted in Figure. 4a, c, e.

2.4. Effect of heterogeneous SEI film on Li/electrolyte interface

We then attempted to identify the heterogeneous SEI structure and investigation of its electrochemical properties. Commercialized carbonate electrolyte exhibits a low average Li Coulombic efficiency of only 69.7% at 0.5 mA cm^{-2} , 0.5 mAh cm^{-2} (Fig. S6), implying significant Li loss due to the severe corrosion between Li and electrolyte. The TEGDME electrolyte also suffers poor Li utilization and low Li coulombic efficiency of only 14.6% (Fig. S7). This is attributed to the Li loss caused by loose Li deposition. Poor Li performance in TEGDME electrolyte can be significantly enhanced by the addition of film-forming electrolyte components. The Li/Cu cell in TEGDME+0.1 M LiNO_3 increase the Li coulombic efficiency to over 90%, compared to efficiency of 14.6% using a LiNO_3 -free TEGDME electrolyte (Fig. 5a). This improvement is a direct result of LiNO_3 forming a relatively stable SEI film on the Li anodes. However, deterioration of the Li performance in TEGDME+0.1 M LiNO_3 occurs after only 25 cycles due to the depletion of LiNO_3 additive. According to the narrow band gap of LiNO_3 reductive products, Li_xNO_y , the fast degradation of Li CE in TEGDME+0.1 M LiNO_3 can be ascribed to the electronic conductive SEI film cannot prohibit the constant electrolyte consumption. A feasible solution is to introduce EC as co-solvent to form a passive outer layer which stabilizes the Li electrolyte inter-face. We study the systematic results on the role of EC/TEGDME ratio in the electrochemical performance. A volumetric percentage increase of EC also increases the ionic conductivity of electrolyte from 2.84 mS cm^{-1} (10 Vol% EC) to 6.21 mS cm^{-1} (60 Vol% EC) (Fig. S8). Further increasing the percentage of EC to 70 Vol% prevents the electrolyte from maintaining liquid status (Fig. S9). Improvement in ionic conductivity of electrolytes with different amount of EC components can be explained by a lower viscosity and strong solvation effect of EC. As we can observe from Fig. S10, the TEGDME electrolyte displays a viscosity of 15 mPa s, which decreases to 11.3 mPa s after introducing 50 Vol.% EC. A less viscous electrolyte decreases the transfer resistance of Li ions. This enables a higher ionic conductivity and faster charge-discharge processes in full cells. We conclude that the optimum EC volume percentage is 50%, which results in a Li Coulombic efficiency of 97.5% (Fig. 5a and Fig. S11). The EC/TEGDME-50/50 electrolyte alone, is still far from satisfying the requirements in a practical battery system. Combining EC and LiNO_3 in TEGDME-based electrolyte elevates the Li Coulombic efficiency. An enhanced Li average coulombic efficiency of 99.0% for

200 cycles at 0.5 mA cm^{-2} , 0.5 mAh cm^{-2} was achieved with further 0.1 M LiNO_3 addition (Fig. 5b), indicating the heterogeneous SEI structure is effective in stabilizing Li electrolyte interface. The superior performance of EC/TEGDME-50/50+ 0.1 M LiNO_3 electrolyte was observed under higher current density and Li capacity cycled for each cycle. For in-stance, 98.1% at 1.0 mA cm^{-2} , 3.0 mAh cm^{-2} (Fig. 5c), 98.2% at 3.0 mA cm^{-2} , 3.0 mAh cm^{-2} (Fig. 5d) and 97.2% at 3.0 mA cm^{-2} , 3.0 mAh cm^{-2} (Fig. 5e).

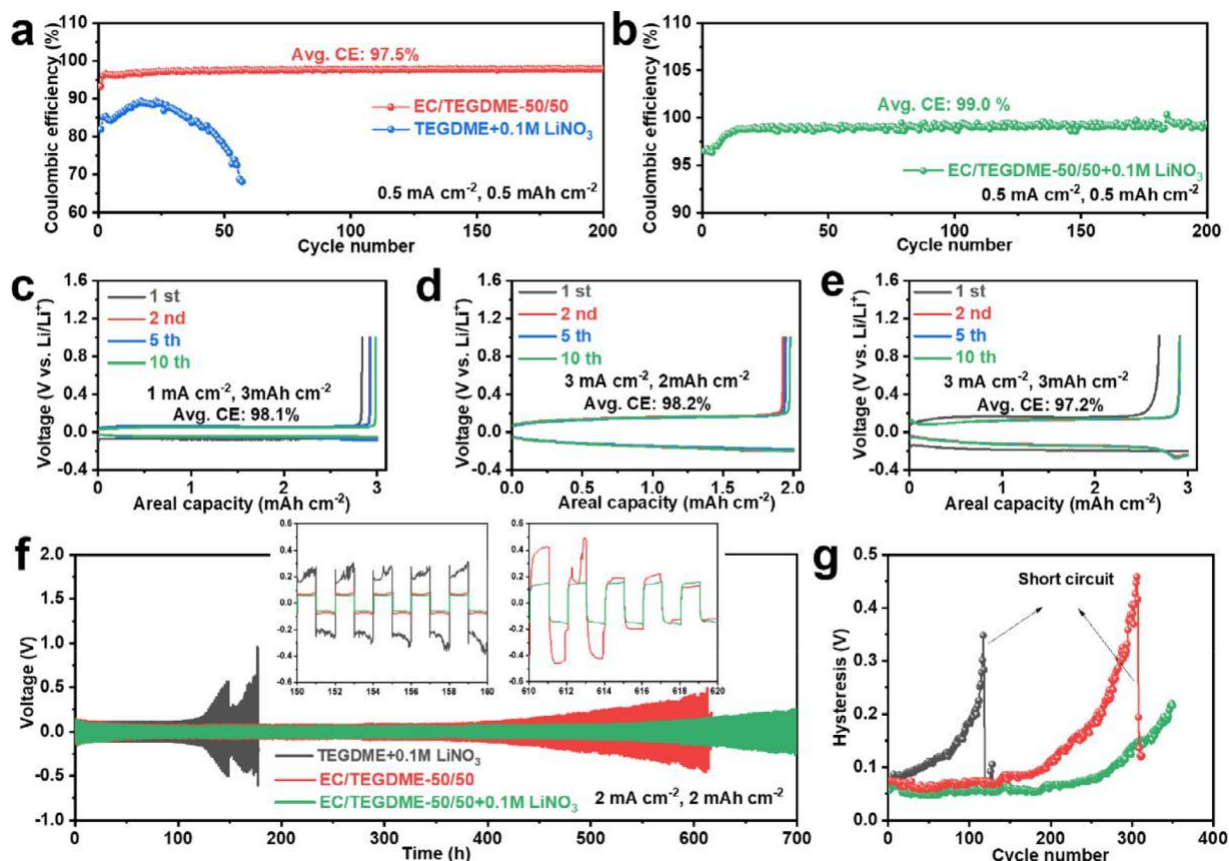


Fig. 5. Coulombic efficiency and cycling stability of lithium metal anodes. Li Coulombic efficiency of Li/Cu half-cells with different electrolytes at 0.5 mA cm^{-2} , 0.5 mAh cm^{-2} : a) EC/TEGDME-50/50 and TEGDME+ 0.1 M LiNO_3 , b) EC/TEGDME-50/50+ 0.1 M LiNO_3 . The typical charge/discharge curves of Li/Cu half-cells with EC/TEGDME-50/50+ 0.1 M LiNO_3 electrolyte at different conditions: c) 1.0 mA cm^{-2} , 3.0 mAh cm^{-2} , d) 3.0 mA cm^{-2} , 2.0 mAh cm^{-2} , e) 3.0 mA cm^{-2} , 3.0 mAh cm^{-2} . f) The voltage-time curves and g) corresponding voltage hysteresis of the symmetric Li/Li cells with different electrolytes at 2 mA cm^{-2} , 2 mAh cm^{-2} .

The long-term stability of the Li electrolyte interface was compared to the voltage-time profiles of symmetric Li/Li cells. As shown in Fig. S12, instability of the Li electrolyte interface in TEGDME electrolyte revealed an initial voltage hysteresis of 225 mV, which significantly deteriorated to 3.9 V after only 40 cycles. In contrast, a similar initial volt-age hysteresis of about 70 mV was measured with different electrolytes (Fig. 5f) at 2.0 mA cm^{-2} , 2.0 mAh cm^{-2} . Gradual increase followed by sudden subsequent drop in voltage was observed after 120 cycles in TEGDME+ 0.1 M LiNO_3 electrolyte and 300 cycles in EC/TEGDME-50/50 (Fig. 5g), corresponding an internal short circuit of the cells [49]. On the contrary, the EC/TEGDME-50/50+ 0.1 M LiNO_3 electrolyte delivers a lower hysteresis without short circuiting after 300 cycles (600 h), proving the heterogeneous SEI film can efficiently suppress the electrolyte decomposition and stabilize the Li electrolyte interface. In addition, the rate performance of Li/Li symmetric cells also compared in Fig. S13. The cell with EC/DMC/EMC exhibits high polarization and in-crease sharply with the current densities enhanced, while the cell with EC/TEGDME-50/50 + 0.1 M LiNO_3 delivers slowly elevation in over-potential with the current densities increase, indicating an excellent rate performance of Li

anodes. Furthermore, short circuit occurs at 3 mA cm^{-2} for Li/Li cell with EC/DMC/EMC reveals the uncontrollable dendrite growth in EC/DMC/EMC electrolyte.

2.5. High-cyclability and high-voltage Li metal batteries

After identifying the optimum electrolyte composition, its electro-chemical performance in both coin cells and pouch cells with commercial cathodes (LiCoO₂, LCO) and a thin Li foil (25 or 200 m) were studied. We first investigated the cyclability LCO/Li coin cells (loading: $19.64 \text{ mg cm}^{-2}/3 \text{ mAh cm}^{-2}$, Li: 200 m, operating voltage region: 2.8–4.25 V) in different electrolytes at 0.2 C and 0.5 C. In the bare TEGDME electrolyte, the specific capacity of LCO/Li cell drops to less than 25 mAh g^{-1} after only 10 cycles at 0.2 C (Fig. S14). Similar degradation also occurs on the LCO/Li cells with commercial electrolyte (EC/DMC/EMC) (Fig. S14), a result of the large amount of dead Li formation during cycling. As shown in Fig. S15, LiNO₃-only TEGDME (TEGDME+0.1 M LiNO₃) electrolyte exhibits an improvement in cycling performance over the bare TEGDME electrolyte, but still suffers rapid capacity decay in the following cycles. The lifespan of the LCO/Li coin cells with an EC-only TEGDME electrolyte (EC/TEGDME-50/50) can be enhanced to 106 cycles at 0.2 C and 75 cycles at 0.5C with a 80% ca-capacity retention, demonstrating that EC can generate efficient passive layer to deter the formation of dead Li (Fig. S15 and S16). The heterogeneous SEI film derived through EC/TEGDME-50/50+0.1 M LiNO₃ dramatically improves the cycling stability of the LCO/Li cell. This cell can deliver an initial capacity of 158.2 mAh g^{-1} (3.1 mAh cm^{-2}), and 138.6 mAh g^{-1} after 130 cycles at 0.2 C, corresponding to a capacity retention of 87.6% (Fig. S15). Similarly, a desirable capacity retention of 86.1% after 100 cycles can be achieved at 0.5 C (Fig. S16). The effect of LiNO₃ concentration in EC/TEGDME-50/50 electrolytes was further investigated. As we can see from Fig. S17, low concentration of LiNO₃ (0.05 M) delivers a relatively low Li CE (~98%) than that with 0.1 M LiNO₃ (~99%), while exhibits no obvious elevation at high concentration of LiNO₃ (0.5 M). However, the cycling performance of LCO/Li cells with high-concentration LiNO₃ electrolyte (EC/TEGDME-50/50+0.5 M LiNO₃) would cause the declination of lifespan to only around 60 cycles (Fig. S18), which indicates the negative impact of LiNO₃ on cathode interface at high concentration, meaning the 0.1 M LiNO₃ can be suitable concentration for the LMBs.

To achieve a higher energy density than traditional LIBs, the cell requires the combination of high-areal-capacity cathodes ($>2.5 \text{ mAh cm}^{-2}$) and ultra-thin Li foils ($<60 \text{ m}$). Hence, cathodes were fabricated by casting LiCoO₂ on both sides of Al foils with an areal capacity of 3 mAh cm^{-2} for each side, and two pieces of ultra-thin Li chips with a thickness of only 25 m were employed as anodes, corresponding to a low N/P (negative/positive capacity) ratio of 1.67, (Fig. 6a). As the performance shown in Fig. 6b, the pouch cells with this configuration were fabricated with a initial capacity of 150 mAh in total. An unprecedented capacity retention of 91.6% after 160 cycles was achieved with EC/TEGDME-50/50+0.1 M LiNO₃ as electrolyte, which demonstrates the promising practical application of the target electrolyte.

The linear sweep voltammetry (LSV) curves suggest that the TEGDME can achieve a stable anodic potential as high as 4.8 V, and only slightly decrease to 4.5 V after introducing 50 Vol.% EC, indicating the potential of utilizing EC/TEGDME-50/50 electrolyte in high-voltage battery systems. Further LiNO₃ addition would not affect the stable potential window of electrolyte, which also enables the operation at 4.5 V (Fig. S19). To further evaluate the anti-oxidation property of the optimum electrolyte, high-loading ($\sim 13 \text{ mg cm}^{-2}$, $\sim 2.5 \text{ mAh cm}^{-2}$) LiCoO₂ cathodes were matched with thin Li foils (200 m) to assemble the LCO/Li coin cells, which underwent an elevated charging voltage of 4.5 V (Fig. 6c and d). Herein, 1 C is defined as 180 mA g^{-1} . The cell cycled in TEGDME+0.1 M LiNO₃ electrolyte had a high initial capacity of 187.9 mAh g^{-1} (2.44 mAh cm^{-2}) but suffers a sharp capacity decay in the following cycles. From the corresponding charge-discharge curves we can know that the polarization of the cells significantly increase after 20 cycles, associating a destructive capacity degradation, and only left $\sim 20 \text{ mAh g}^{-1}$ after 30 cycles (Fig. 6e). With the EC/TEGDME-50/50 electrolyte, LCO/Li coin cells achieve 148 cycles and 157 cycles at 0.2 C and 0.5 C, respectively. An excellent cycling performance was obtained in EC/TEGDME-50/50+0.1 M

LiNO₃ with a capacity retention of 91.5% after 180 cycles at 0.2 C (Fig. 6c). At higher current rate (0.5C), a remarkable long-term cycling performance over 300 cycles with a capacity retention of 85.7% is possible when using EC/TEGDME-50/50+0.1 M LiNO₃ electrolyte (Fig. 6d). As revealed in Fig. 6f and g, the LCO/Li cell cycled in EC/TEGDME-50/50+0.1 M LiNO₃ displays smaller polarization and less capacity decay than that cycled in EC/TEGDME-50/50. And even after 300 cycles, there is only slight increase in polarization, indicating the desirable heterogenous SEI structure significantly stabilize the Li interface. Additionally, the rate capability of LCO/Li cells with different electrolytes was compared (Fig. S20). For the commercial EC/DMC/EMC electrolyte, the LCO/Li cell delivers a poor rate performance with nearly 0 mAh cm⁻² at 3 C, which results from the poor performance of Li anodes. Thus, to eliminate the influence of Li anodes, the EC/DMC/EMC electrolyte with 5% FEC as additive was applied to evaluate the real rate capability of commercial electrolyte. Significantly, EC/DMC/EMC with 5% FEC can deliver an excellent rate capability with ~2 mAh cm⁻² at 3 C rate. Owing to the relatively higher viscosity of TEGDME-based electrolyte, EC/TEGDME-50/50+0.1 M LiNO₃ delivers a relatively poor performance at high rate (3 C). However, EC/TEGDME-50/50+0.1 M LiNO₃ also can perform a similar capacity at 2 C compared with EC/DMC/EMC with 5% FEC, which can satisfy the practical application.

The evolution of SEI properties in full cells can be monitored by *in-situ* EIS. Herein, a three-electrodes pouch cell with LCO as work electrode, Li foil as counter electrode, and Li wire as reference electrode was adopted to avoid the effect of cathode resistance on the anode. Fig. S21 reveals the evolution of resistance of Li anodes during a whole charge-discharge cycle in LCO/Li cells in EC/DMC/EMC and EC/TEGDME-50/50+0.1 M LiNO₃. The Nyquist plots were further fitting analysis according the equivalent circuit model shown in Fig. S22. For the EC/DMC/EMC electrolytes, the R_{SEI} decrease during the charging process, which results from the porous Li deposition enlarge the surface area of anodes. While R_{SEI} further significantly increase in following discharging process, accompanied with improvement of R_{ct} , which means the thick SEI generation significantly enhances the resistance of SEI film and retard the charge transfer processes. On the contrary, the Li anodes cycled in EC/TEGDME-50/50+0.1 M LiNO₃ delivers a relatively small and stable SEI resistance during the whole charging/discharging process. Meanwhile, low R_{ct} also are delivered in EC/TEGDME-50/50+0.1 M LiNO₃, indicating the a stable and fast ionic conductive SEI was generated by EC/TEGDME/LiNO₃ electrolyte system.

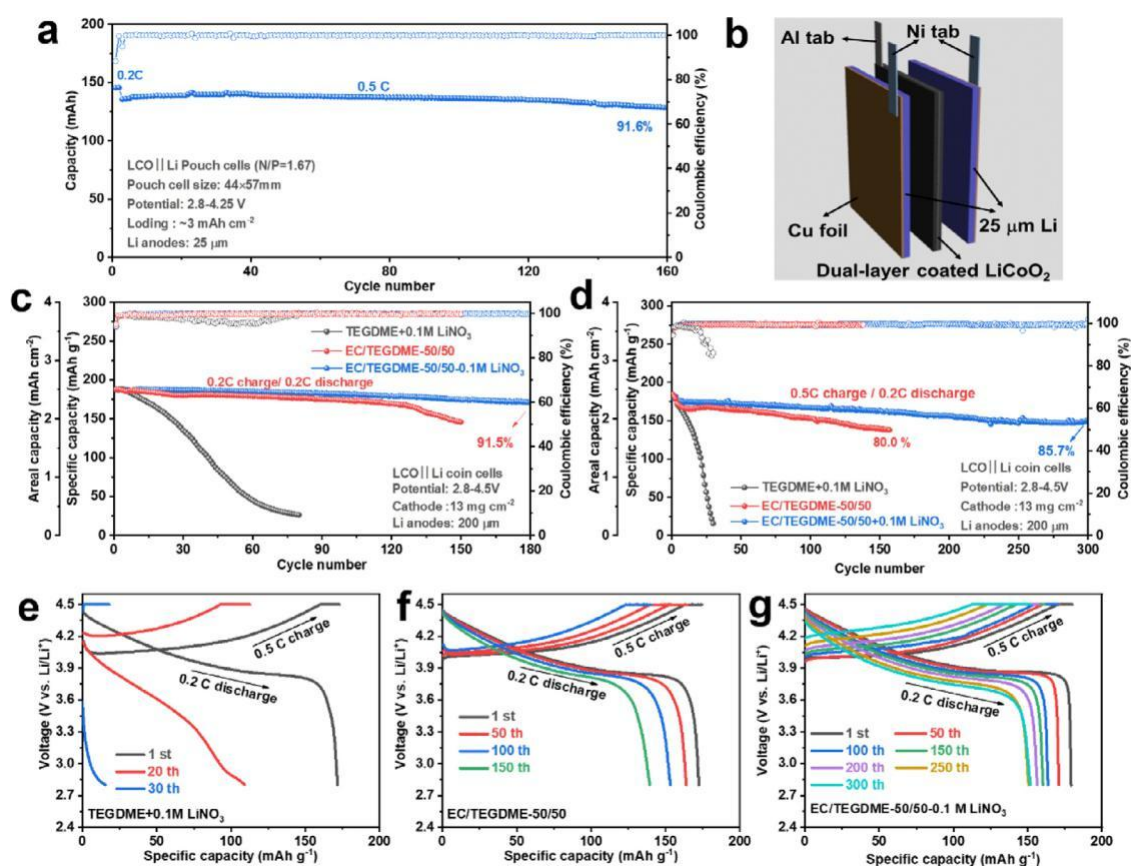


Fig. 6. Electrochemical performance under practically relevant conditions. a) The structure of pouch cells and b) Cycling performance of the pouch cell with ultra-thin Li foil (25 m). The cycling performance of the LiCoO₂/Li coin cells with different electrolytes at c) 0.2 C and (d) 0.5 C at a cut-off voltage range of 2.8–4.5 V. The typical charge-discharge curves of the 4.5 V LCO/Li cells in different electrolytes: e) TEGDME+0.1 M LiNO₃, f) EC/TEGDME-50/50 and g) EC/TEGDME-50/50+0.1 M LiNO₃.

3. Conclusions

A novel and straightforward method to directionally design heterogeneous SEI with inner-outer hybrid electronic/ionic conductivity via selective reduction of electrolyte components is found to be capable of achieving both high Li utilization and high oxidation tolerance for Li metal battery. TEGDME-based electrolyte suffers poor performance on Li metal anodes, resulting in the constant decomposition of TEGDME by Li metal. LiNO₃ efficiently stabilizes the Li electrolyte interface; however, the batteries still suffer sharp degradation with the consumption of LiNO₃. Introducing 50 Vol.% EC to replace part of the TEGDME in electrolyte significantly suppresses the exhaustion of LiNO₃ additive. Consequently, a remarkable Li coulombic efficiency of 99.0% is attained using EC/TEGDME-50/50+0.1 M LiNO₃. DFT calculations and XPS depth analysis demonstrate that the LiNO₃ is reduced by Li metal prior to the EC. A heterogeneous SEI film with an outer C-rich region and an inner N-rich region is generated on the Li surface. In the film, the inner region exhibits high Li⁺ conductivity, leading to the round-shape Li de-position, while the outer region with electronic insulation serves as a passive layer delivering less electrolyte consumption. The AFM SEI film study verified the formation of the heterogeneous SEI structure and their physical properties, consisting of an inner rigid region (~2.02 GPa) and outer soft region (~0.32 GPa). Benefiting from this novel heterogeneous SEI structure, the morphology of Li metal becomes nodule-like and the thickness of the porous Li layer after cycling was only 6 m, which is significantly thinner than that in bare TEGDME electrolyte (136 m). The LCO/Li full cells were also studied for the practical application of EC/TEGDME-55+0.1 M LiNO₃ with excellent results. Pouch cells with ultra-high-loading (3 mAh cm⁻²) LiCoO₂ electrodes and ultra-thin Li foils (25 m) deliver an outstanding cycling performance with a capacity retention of 91.6% after 160 cycles. The high-voltage LCO/Li (4.5 V) batteries with modified configuration deliver an 85.7% capacity retention after 300 cycles, demonstrating superb Li utilization and high voltage tolerance of electrolytes. This approach can be considered as a suitable strategy to generate a desirable SEI structure on the Li surface to achieve rechargeable Li metal batteries, which would be compatible with most of the existing Li battery systems in the pursuit of high energy densities.

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Supplementary material

Supplementary data associated with this article is available.

References

- [1] J.M. Tarascon, M. Armand, *Nature* 414 (2001) 359–367.

- [2] X. Gao, Y. Chen, L.R. Johnson, Z.P. Jovanov, P.G. Bruce, *Nat. Energy*. 2 (2017) 17118.
- [3] H. Kim, G. Jeong, Y.U. Kim, J.H. Kim, C.M. Park, H.J. Sohn, *Chem. Soc. Rev.* 42 (2013) 9011–9034.
- [4] Q. Xu, J. Lin, C. Ye, X. Jin, D. Ye, Y. Lu, G. Zhou, Y. Qiu, W. Li, *Adv. Energy Mater.* 10 (2019) 1903292.
- [5] K. Liao, S. Wu, X. Mu, Q. Lu, M. Han, P. He, Z. Shao, H. Zhou, *Adv. Mater.* 30 (2018) 1705711.
- [6] M. Bai, K. Xie, K. Yuan, K. Zhang, N. Li, C. Shen, Y. Lai, R. Vajtai, P. Ajayan, B. Wei, *Adv. Mater.* (2018) 1801213.
- [7] E.K. Jang, J. Ahn, S. Yoon, K.Y. Cho, *Adv. Funct. Mater.* 29 (2019) 1905078.
- [8] Y.T. Weng, H.W. Liu, A. Pei, F. Shi, H. Wang, C.Y. Lin, S.S. Huang, L.Y. Su, J.P. Hsu, C.C. Fang, Y. Cui, N.L. Wu, *Nat. Commun.* 10 (2019) 5824.
- [9] S.J. Zhang, Z.G. Gao, W.W. Wang, Y.Q. Lu, Y.P. Deng, J.H. You, J.T. Li, Y. Zhou, L. Huang, X.D. Zhou, S.G. Sun, *Small* 14 (2018) 1801054.
- [10] S. Liu, X. Xia, Y. Zhong, S. Deng, Z. Yao, L. Zhang, X.B. Cheng, X. Wang, Q. Zhang, J. Tu, *Adv. Energy Mater.* 8 (2018) 1702322.
- [11] H. Shi, C.J. Zhang, P. Lu, Y. Dong, P. Wen, Z.S. Wu, *ACS Nano* 13 (2019) 14308–14318.
- [12] C. Sun, A. Lin, W. Li, J. Jin, Y. Sun, J. Yang, Z. Wen, *Adv. Energy Mater.* (2019) 1902989.
- [13] Q. Xu, X. Yang, M. Rao, D. Lin, K. Yan, R. Du, J. Xu, Y. Zhang, D. Ye, S. Yang, G. Zhou, Y. Lu, Y. Qiu, *Energy Stor. Mater.* 26 (2020) 73–82.
- [14] Z. Peng, X. Cao, P. Gao, H. Jia, X. Ren, S. Roy, Z. Li, Y. Zhu, W. Xie, D. Liu, Q. Li, D. Wang, W. Xu, J.G. Zhang, *Adv. Funct. Mater.* 30 (2020) 1606422.
- [15] X. Cao, X. Ren, L. Zou, M.H. Engelhard, W. Huang, H. Wang, B.E. Matthews, H. Lee, C. Niu, B.W. Arey, Y. Cui, C. Wang, J. Xiao, J. Liu, W. Xu, J.G. Zhang, *Nat. Energy*. 4 (2019) 796–805.
- [16] T. Jiang, P. He, G. Wang, Y. Shen, C.W. Nan, L.Z. Fan, *Adv. Energy Mater.* 10 (2020) 1903376.
- [17] L. Chen, Y. Li, S.P. Li, L.Z. Fan, C.W. Nan, J.B. Goodenough, *Nano Energy* 46 (2018) 176–184.
- [18] J. Hu, P. He, B. Zhang, B. Wang, L.Z. Fan, *Energy Stor. Mater.* 26 (2020) 283–289.
- [19] L. Chen, W. Li, L.Z. Fan, C.W. Nan, Q. Zhang, *Adv. Fuct. Mater.* 29 (2019) 1901047.
- [20] J. Qian, W.A. Henderson, W. Xu, P. Bhattacharya, M. Engelhard, O. Borodin, J.G. Zhang, *Nat. Commun.* 6 (2015) 6362.
- [21] X. Dong, Y. Lin, P. Li, Y. Ma, J. Huang, D. Bin, Y. Wang, Y. Qi, Y. Xia, *Angew. Chem. Int. Ed. Engl.* 58 (2019) 5623–5627.
- [22] X. Shangguan, G. Xu, Z. Cui, Q. Wang, X. Du, K. Chen, S. Huang, G. Jia, F. Li, X. Wang, D. Lu, S. Dong, G. Cui, *Small*. 15 (2019) 1900269.
- [23] P. Shi, L. Zhang, H. Xiang, X. Liang, Y. Sun, W. Xu, *ACS Appl. Mater. Interfaces* 10 (2018) 22201–22209.
- [24] J. Zheng, M.H. Engelhard, D. Mei, S. Jiao, B.J. Polzin, J.G. Zhang, W. Xu, *Nat. Energy* 2 (2017) 17012.
- [25] H. Ye, Y.X. Yin, S.F. Zhang, Y. Shi, L. Liu, X.X. Zeng, R. Wen, Y.G. Guo, L.J. Wan, *Nano Energy* 36 (2017) 411–417.

- [26] X.R. Chen, Y.X. Yao, C. Yan, R. Zhang, X.B. Cheng, Q. Zhang, *Angew. Chem. Int. Ed. Engl.* 132 (2020) 7817–7821.
- [27] J. Fu, X. Ji, J. Chen, L. Chen, X. Fan, D. Mu, C. Wang, *Angew. Chem. Int. Ed.* 59 (2020) 22194–22201.
- [28] Y.X. Lin, Z. Liu, K. Leung, L.Q. Chen, P. Lu, Y. Qi, J. *Power Sources* 309 (2016) 221–230.
- [29] C.C. Su, M. He, R. Amine, Z. Chen, R. Sahore, N.D. Rago, K. Amine, *Energy Stor. Mater.* 17 (2019) 284–292.
- [30] J. Heine, P. Hilbig, X. Qi, P. Niehoff, M. Winter, P. Bieker, *J. Electrochem. Soc.* 162 (2015) A1094–A1101.
- [31] C. Pereira-Nabais, J. Świątowska, A. Chagnes, F. Ozanam, A. Gohier, P. Tran-Van, C.-S. Cojocaru, M. Cassir, P. Marcus, *Appl. Surface Sci.* 266 (2013) 5–16.
- [32] J. Świątowska, V. Lair, C. Pereira-Nabais, G. Cote, P. Marcus, A. Chagnes, *Appl. Surface Sci.* 257 (2011) 9110–9119.
- [33] M. Carboni, A.G. Marrani, R. Spezia, S. Brutti, *J. Electrochem. Soc.* 165 (2018) A118–A125.
- [34] J. Zheng, P. Yan, D. Mei, M.H. Engelhard, S.S. Cartmell, B.J. Polzin, C. Wang, J.G. Zhang, W. Xu, *Adv. Energy Mater.* 6 (2016) 1502151.
- [35] D. Aurbach, K. Gamolsky, B. Markovsky, Y. Gofer, M. Schmidt, U. Heider, *Electrochim. Acta* 47 (2002) 1423–1439.
- [36] C. Yan, Y.X. Yao, X. Chen, X.B. Cheng, X.Q. Zhang, J.Q. Huang, Q. Zhang, *Angew. Chem. Int. Ed. Engl.* 57 (2018) 14055–14059.
- [37] V. Shutthanandan, M. Nandasiri, J. Zheng, M.H. Engelhard, W. Xu, S. Thevuthasan, V. Murugesan, *J. Electron Spectros. Relat. Phenomena* 231 (2019) 2–10.
- [38] S. Tanaka, M. Taniguchi, H. Tanigawa, *J. Nucl. Mater.* 283-287 (2000) 1405–1408.