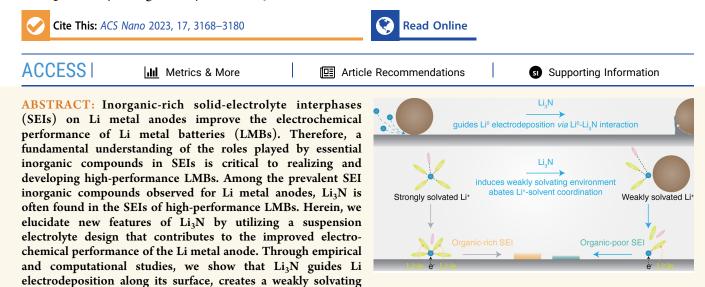


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# Revealing the Multifunctions of Li<sub>3</sub>N in the Suspension Electrolyte for Lithium Metal Batteries

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environment by decreasing Li<sup>+</sup>-solvent coordination, induces organic-poor SEI on the Li metal anode, and facilitates Li<sup>+</sup> transport in the electrolyte. Importantly, recognizing specific roles of SEI inorganics for Li metal anodes can serve as one of the rational guidelines to design and optimize SEIs through electrolyte engineering for LMBs.

**KEYWORDS:** *lithium*—metal battery, *lithium* metal anode, *suspension electrolyte*, *lithium nitride*, *solid-electrolyte interphase*, *lithium solvation environment*, *electrolyte engineering* 

# **INTRODUCTION**

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Reversibly utilizing Li metal (Li<sup>0</sup>) anodes is a primary step in successfully developing practical high-energy Li metal batteries (LMBs).<sup>1,2</sup> Although the Li<sup>0</sup> anode involves a classic electroplating/stripping mechanism, achieving highly reversible and electrochemically stable Li<sup>+</sup> migration at the Li<sup>0</sup>/ electrolyte interface becomes challenging due to the inevitable formation of solid-electrolyte interphases (SEIs) on Li<sup>0</sup> anodes with electrolytes.<sup>3</sup> Despite the desirable properties of Li<sup>0</sup> anodes, such as a high theoretical specific capacity of 3,861 mAh g<sup>-1</sup> and a low standard electrolytes thermodynamically unstable against Li<sup>0</sup> that decompose electrolyte species to form SEIs, in which the properties of SEIs largely influence the electrochemical performance of Li<sup>0</sup> anodes.

During LMB cycles, solvated Li<sup>+</sup> must reversibly migrate across the electrolyte and SEIs on Li<sup>0</sup> anodes, in which characteristics of SEIs predominantly affect Li<sup>+</sup> transport behavior.<sup>3</sup> Essentially, SEIs on Li<sup>0</sup> anodes affect Li<sup>0</sup> electroplating/stripping and Li<sup>+</sup> desolvation processes, thereby dictating the electrochemical performance of Li<sup>0</sup> anodes.<sup>7–9</sup> Despite the critical functions of SEIs on Li<sup>0</sup> anodes, understanding the roles played by SEI components is still at an early stage to rationally design and manipulate SEIs via electrolyte engineering. As SEI evolutions on Li<sup>0</sup> anodes are directly linked to the solvation structures of Li<sup>+</sup> in electrolytes,<sup>3</sup> various modifications to the Li<sup>+</sup> solvation environment and trends have been investigated to correlate SEI properties that positively impact the electrochemical performance of Li<sup>0</sup> anodes. In particular, weakly solvating electrolytes and

Received:December 16, 2022Accepted:January 23, 2023Published:January 26, 2023





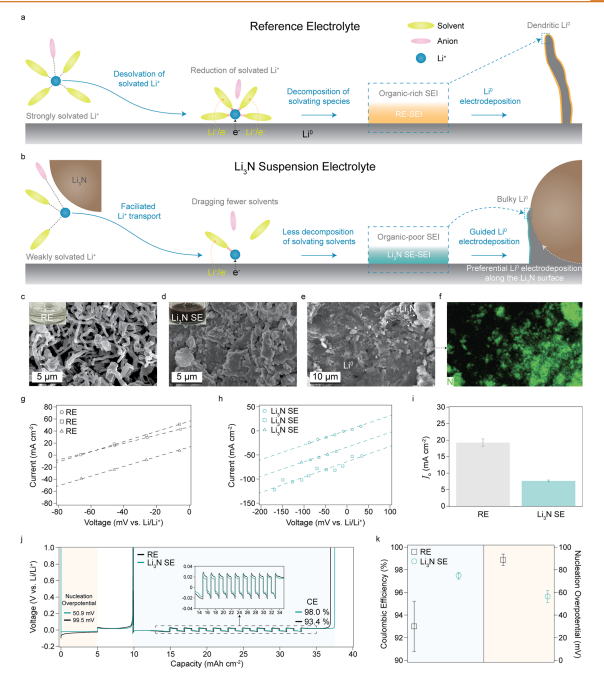


Figure 1. Features of Li<sub>3</sub>N for Li<sup>0</sup> anodes. (a) Schematic illustration of strongly solvated Li<sup>+</sup> in RE impacting the SEI evolution on Li<sup>0</sup> anodes and Li<sup>0</sup> electrodeposition. The first step shows Li<sup>+</sup> desolvation and Li<sup>+</sup> reduction processes from the solvated Li<sup>+</sup> in RE. The next step represents the SEI formation on Li<sup>0</sup> anodes by decomposing solvating species that yield organic-rich SEI. The final step delineates the morphology of dendritic Li<sup>0</sup> electrodeposits influenced by RE-SEI. (b) Schematic illustration of weakly solvated Li<sup>+</sup> in Li<sub>3</sub>N SE impacting the SEI evolution on  $Li^0$  anodes and  $Li^0$  electrodeposition. The first step represents the weakly solvated  $Li^+$  by  $Li_3N$  in the electrolyte, in which fewer solvents are coordinated with Li<sup>+</sup> relative to that in RE. The next step shows facilitated Li<sup>+</sup> transport by the decreased Li<sup>+</sup>-solvent interaction. Then, the weakly solvated Li<sup>+</sup> suppresses decomposition of solvents by dragging relatively fewer solvent molecules close to the Li<sup>0</sup> surface that forms organic-poor SEI. The final step shows bulky Li<sup>0</sup> electrodeposition allowed by the preferential Li<sup>0</sup> deposition across the Li<sub>3</sub>N surface and Li<sub>3</sub>N SE-SEI. (c) SEM image of Li<sup>0</sup> electrodeposits on Cu from LilCu cell with RE at 1 mA cm<sup>-2</sup> and 1 mAh cm<sup>-2</sup>. The inset represents the physical image of RE. (d) SEM image of Li<sup>0</sup> electrodeposits on Cu from LilCu cell with Li<sub>3</sub>N SE at 1 mA cm<sup>-2</sup> and 1 mAh  $cm^{-2}$ . The inset represents the physical image of Li<sub>3</sub>N SE (e<sub>j</sub>f), SEM image (e), and corresponding nitrogen elemental mapping (f) of Li<sup>0</sup> electrodeposits on Cu from LilCu cell with exposed Li<sub>3</sub>N from Li<sub>3</sub>N SE. (g,h) Representative kinetically controlled region from the reverse sweep of ultramicroelectrode cyclic voltammograms for RE (g) and Li<sub>3</sub>N SE (h). Three measurements were performed for RE and Li<sub>3</sub>N SE. The scan rate and voltage window were 20 V s<sup>-1</sup> and -1.75 to 2.75 V vs Li/Li<sup>+</sup>. (i) Calculated exchange current density ( $J_0$ ) from g and h for RE (19.27 mA cm<sup>-2</sup>  $\pm$  1.13 mA cm<sup>-2</sup>) and Li<sub>3</sub>N SE (7.64 mA cm<sup>-2</sup>  $\pm$  0.28 mA cm<sup>-2</sup>). (j) Representative voltage profiles of CE measurement from LilCu cells with RE and Li<sub>3</sub>N SE. The inset represents the zoomed-in voltage profiles from 13 mAh cm<sup>-2</sup> to 35 mAh cm<sup>-2</sup>. The background colors represent where the CE and nucleation overpotential were measured. The applied current density was 0.5 mA cm<sup>-2</sup>. (k) Averaged CE and nucleation overpotential values with standard errors calculated from three identical LilCu cells with RE (93.01%  $\pm$  2.21% and 88.77 mV  $\pm$  5.11 mV) and Li<sub>3</sub>N SE (97.48%  $\pm$  0.31% and 56.43 mV  $\pm$  5.31 mV).

inorganic-rich SEIs were found to be beneficial to the electrochemical performance of  ${\rm Li}^0$  anodes.<sup>3,6,10-14</sup>

It has been found that SEIs on Li<sup>0</sup> anodes can swell with electrolytes,<sup>15</sup> indicating that electrolyte species still exist within the SEI domain. This important outcome further justifies the aging of SEIs on Li<sup>0</sup> anodes.<sup>16</sup> In other words, SEIs are still electrolyte penetrable up to a certain degree, and the electrochemical performance of Li<sup>0</sup> anodes are, therefore, affected by both SEIs and the electrolyte species. Furthermore, a modified Li<sup>+</sup> solvation environment by the SEI inorganic compound in the electrolyte has been discovered, which sheds light on the specific features of SEI inorganics that affect SEI evolution and the electrochemical performance of Li<sup>0</sup> anodes.<sup>6</sup> These combined results suggest that not only can the Li<sup>+</sup> solvation environment within the SEI region be different from the Li<sup>+</sup> solvation environment in the electrolyte, but also the electrochemical performance of SEIs on Li<sup>0</sup> anodes is determined by the distinct SEI inorganic species that sets the specific Li<sup>+</sup> solvation environment within the SEI layer where the actual Li<sup>+</sup> desolvation occurs. These insights lead to a critical hypothesis that the Li<sup>+</sup> solvation shells, not just Li<sup>+</sup>, can migrate across SEIs, in which the solvated Li<sup>+</sup> in the bulk electrolyte further gets modified by the specific inorganic species present in the SEIs on Li<sup>0</sup> anodes. Thus, revealing the roles of inorganic compounds in SEIs is considered one of the fundamental studies to design high-performance SEIs via electrolyte engineering for Li<sup>0</sup> anodes that will potentially lead to the development of advanced electrolytes for developing reliable LMBs.

Among the frequently observed SEI inorganic compounds (e.g.,  $Li_2O$ ,  $Li_2CO_3$ , LiF,  $Li_3N$ ,  $Li_2S$ , ...),  $Li_3N$  was particularly found in almost all the high-performance SEIs of  $Li^0$  anodes.<sup>3</sup> The beneficial features of  $Li_3N$  for  $Li^0$  anodes from prior studies include thermodynamic stability,<sup>17,18</sup> high  $Li^+$  conductivity,<sup>19</sup> and  $Li^0$  dendrite suppression.<sup>20–23</sup> Although various forms of  $Li_3N$  (pinhole-free film,<sup>23</sup> nanoflowers,<sup>24</sup> and nanosheets<sup>25</sup>) and electrolyte additives ( $LiN_3^{20}$  and  $LiNO_3^{26}$  to derive  $Li_3N$ ) have been applied to improve the electrochemical performance of  $Li^0$  anodes, there is still an inadequate understanding of the functional mechanism of  $Li_3N$  for  $Li^0$  anodes that rationalize the working mechanism of  $Li_3N$  for  $Li^0$  anodes and SEIs.

Herein, we designed a Li<sub>3</sub>N suspension electrolyte (Li<sub>3</sub>N SE) by mixing 80–100 nm Li<sub>3</sub>N nanoparticles with a reference electrolyte (RE) of 1 M LiPF<sub>6</sub> in ethylene carbonate (EC) and diethyl carbonate (DEC) (1:1 v/v) with 10 vol % fluoroethylene carbonate (FEC), in which RE serves as one of the most widely studied electrolytes for the SEI analyses of Li<sup>0</sup> anodes.<sup>6,8,9,15</sup> Also, the electrolyte blend of RE does not contain any nitrogen species that could form Li<sub>3</sub>N through the reduction and/or decomposition of RE species, which will help to closely examine the effect of Li<sub>3</sub>N in RE. Moreover, utilizing the suspension electrolyte design allows for revealing the functional mechanism of a particular SEI inorganic compound for LMBs.<sup>6</sup> Thus, Li<sub>3</sub>N SE was investigated in detail to identify the Li<sup>+</sup> solvation environment change, SEI evolution on Li<sup>0</sup> anodes, and electrochemical performance. By combining the experimental assessments in tandem with first-principles results, we were able to identify new features of Li<sub>3</sub>N in the liquid electrolyte for  $Li^0$  anodes: (i) the surface of  $Li_3N_2$ , especially (001) and (002), is lithiophilic, which guides Li<sup>0</sup>

electrodeposition through a favored Li<sup>0</sup>-Li<sub>3</sub>N interaction, (ii) Li<sub>3</sub>N creates a weakly solvating environment by decreasing Li<sup>+</sup>solvent interaction in the electrolyte, (iii) Li<sub>3</sub>N suppresses solvent decomposition to form organic-poor SEI on Li<sup>0</sup> anodes, and (iv)  $\mathrm{Li}_3N$  facilitates  $\mathrm{Li}^{\scriptscriptstyle +}$  transport in the electrolyte and Li<sup>0</sup> interphase. These findings support the progress of comprehending the improved Li<sup>0</sup> anode performance in connection with the previously reported Li<sub>3</sub>N studies and the inorganic-rich SEI concept, particularly having high Li<sub>3</sub>N content in the SEIs, of Li<sup>0</sup> anodes. Significantly, beneficial SEI inorganic compounds induce a weakly solvating Li<sup>+</sup> environment by modifying Li<sup>+</sup>-solvent and/or Li<sup>+</sup>-anion coordination,<sup>6</sup> in which Li<sub>3</sub>N is responsible for adjusting Li<sup>+</sup>-solvent coordination, which affects the electrochemical performance of and SEI evolution on Li<sup>0</sup> anodes. Consequently, understanding the unique features of SEI inorganic species can potentially provide scientific and rational guidelines to control and design SEIs on Li<sup>0</sup> anodes by optimizing electrolyte chemistries to improve the electrochemical performance of LMBs.

# **RESULTS AND DISCUSSION**

Analyses of Li<sub>3</sub>N in the Electrolyte. Deriving inorganicrich SEIs by engineering electrolytes became one of the practical approaches to significantly improve the electrochemical performance of Li<sup>0</sup> anodes for LMBs.<sup>12,13,27,28</sup> Since the SEI evolution on Li<sup>0</sup> anodes is directly associated with the electrolyte, <sup>3,12,13,28-30</sup> especially its Li<sup>+</sup> solvation environment, understanding the roles played by the major SEI inorganic components, such as Li<sub>2</sub>O, Li<sub>2</sub>CO<sub>3</sub>, LiF, Li<sub>3</sub>N, and Li<sub>2</sub>S,<sup>3</sup> can serve as a bridge to connect the mechanism of SEI species and electrochemical performance of Li<sup>0</sup> anodes.<sup>6</sup> Furthermore, the swelling of SEIs on Li<sup>0</sup> anodes has recently been identified.<sup>15</sup> It has been shown that the degree of SEI swelling correlates to the electrochemical performance of Li<sup>0</sup> anodes, in which inorganic-rich or organic-poor SEIs promote superior electrochemical performance of Li<sup>0</sup> anodes and less SEI swelling.<sup>15</sup> This discovery also leads to an important hypothesis that electrolytes are still present within the SEI layer, where the inorganic compounds in SEIs play an important role in adjusting the Li<sup>+</sup> solvation environment at the interphase of and SEI evolution on Li<sup>0</sup> anodes.<sup>6</sup> Since Li<sub>3</sub>N is a prominent inorganic that exists in the SEIs of many high-performance  $\text{Li}^0$  anodes, <sup>3,20,26,31–39</sup> this study aims to unveil the important roles of Li<sub>3</sub>N in terms of Li<sup>+</sup> solvation effects as well as Li<sup>0</sup> electrodeposition stability (Figure 1a,b).

Hypotheses of Li<sup>+</sup> solvation environment, SEI evolution, and Li<sup>0</sup> electrodeposition behavior with RE (Figure 1a) and Li<sub>3</sub>N SE (Figure 1b) are illustrated to delineate the features of  $Li_3N$ . In Figure 1a, the first step represents the desolvation process of strongly solvated Li<sup>+</sup> by solvents. Then, the solvated Li<sup>+</sup> in RE gets reduced at the surface of Li<sup>0</sup> anodes to form organic-rich SEI. Eventually, the SEI derived from RE (RE-SEI) induces dendritic Li<sup>0</sup> electrodeposition morphology. With Li<sub>3</sub>N, the solvated Li<sup>+</sup> becomes weakly solvated relative to that of RE due to the decrease in the Li<sup>+</sup>-solvent interaction by Li<sub>3</sub>N in the electrolyte (Figure 1b). Please note that the Li<sup>+</sup>-anion coordination was not affected by the Li<sub>3</sub>N in the electrolyte. The weakly solvated Li<sup>+</sup> in Li<sub>3</sub>N SE facilitates the transport of solvated Li<sup>+</sup> through the electrolyte and Li<sup>0</sup> interphase. This modified environment of solvated Li<sup>+</sup> in Li<sub>3</sub>N SE then suppresses solvent decomposition to form relatively organicpoor SEI on Li<sup>0</sup> anodes. Furthermore, the lithiophilic surface of

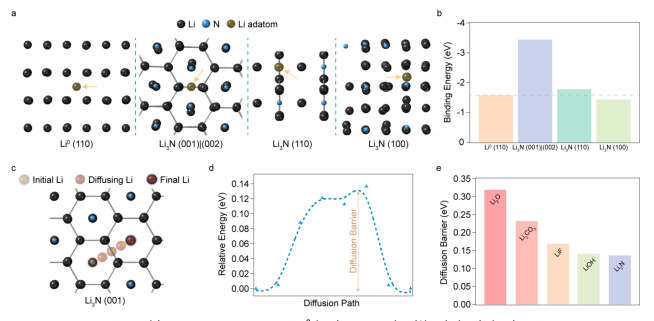


Figure 2. DFT analyses of  $Li_3N$ . (a) Li adatom binding sites of  $Li^0$  (110) and  $Li_3N$  (001)l(002), (110), (100) surfaces. The arrows indicate surface adsorbed Li adatom onto the corresponding surfaces of  $Li^0$  and  $Li_3N$ . The color codes for Li of  $Li_3N$ , N, and Li adatom are black, skyblue, and gray, respectively. (b) Calculated Li adatom binding energies for  $Li^0$  (110),  $Li_3N$  (001)l(002),  $Li_3N$  (110), and  $Li_3N$  (100) surfaces. The dashed gray line indicates the binding energy of  $Li^0$  (110). (c) The Li adatom diffusion pathway across  $Li_3N$  (001) surface. (d) Relative energies based on the Li adatom diffusion trajectories across  $Li_3N$  (001) surface. The diffusion barrier is represented by the magnitude between the highest and lowest relative energies. (e) Li adatom surface diffusion barriers for  $Li_2O$  (0.319 eV),  $Li_2CO_3$  (0.232 eV), LiF (0.169 eV), LiOH (0.141 eV), and  $Li_3N$  (0.133 eV). The diffusion barriers of  $Li_2O$ ,  $Li_2CO_3$ , LiF, and LiOH were retrieved from the literature.<sup>44</sup>

 $Li_3N$  and the organic-poor SEI derived from  $Li_3N$  SEI ( $Li_3N$  SE-SEI) allow bulky  $Li^0$  electrodeposition. The rest of the discussions are devoted to corroborating these features of  $Li_3N$  for  $Li^0$  anodes.

Observing Li<sup>0</sup> electrodeposition morphologies became a convenient way to anticipate electrochemical characteristics, such as Coulombic efficiency (CE) and Li nucleation overpotential, of  $Li^0$  anodes.<sup>1,40,41</sup> Scanning electron microscopy (SEM) images were examined first to analyze morphological features of Li<sup>0</sup> electrodeposits with RE and  $Li_3N$  SE (Figure 1c-f). High surface area and dendritic  $Li^0$ morphologies were observed with RE (Figure 1c). On the other hand, the Li<sup>0</sup> electrodeposits with Li<sub>3</sub>N SE exhibited nondendritic morphology (Figure 1d). Notably, Li<sub>3</sub>N in Li<sub>3</sub>N SE were embedded into the Li<sup>0</sup> electrodeposits, alluding to preferential Li<sup>0</sup> electrodeposition along the surfaces of Li<sub>3</sub>N (Figure 1d,e). The elemental mapping of nitrogen shown in Figure 1e via energy-dispersive X-ray spectroscopy (EDXS) confirmed that the observed particles were Li<sub>3</sub>N as Li<sub>3</sub>N was the only nitrogen-containing species in Li<sub>3</sub>N SE (Figure 1f). To justify the guided Li<sup>0</sup> electrodeposition toward Li<sub>3</sub>N, the low-magnification SEM image of the electrodeposited Li<sup>0</sup> with Li<sub>3</sub>N SE is shown in Figure S1a. Figure S1a vividly depicts rough surface features due to the Li<sub>3</sub>N network formed by the Li<sub>3</sub>N SE, in which the nitrogen mapping indicates that the topmost part is Li<sub>3</sub>N (Figure S1b). The carbon and oxygen mapping in Figure S1c,d indirectly illustrate Li<sup>0</sup> electrodeposits, as they are the major elements for SEIs on Li<sup>0</sup>. Based on Figure S1a-d, the region that exhibits the height variation due to Li<sub>3</sub>N was selected and shown in Figure S1e, in which the rough surface region was purposefully chosen to demonstrate the preferred Li<sup>0</sup> electrodeposition toward Li<sub>3</sub>N. Figure S1e evidently shows that Li<sup>0</sup> was electrodepositing along the surfaces of Li<sub>3</sub>N as the bottommost surface was free

of Li dendrites, and the  $Li_3N$  was well covered by  $Li^0$  electrodeposits. Hence, the SEM/EDXS analysis suggest that  $Li^0$  tends to get electrodeposited along the surfaces of  $Li_3N$ .

To electrochemically probe possible Li<sup>0</sup>-Li<sub>3</sub>N interactions in  $Li_3N$  SE, the exchange current densities ( $J_0$ ) for RE and  $Li_3N$ SE were measured with the ultramicroelectrode under a fast scan  $(20 \text{ V s}^{-1})$  cyclic voltammetry (CV) (Figure 1g-i). The fast scan CV with ultramicroelectrode allows for closely investigating electron transfer kinetics of Li with respect to a particular electrolyte by minimizing the effect of mass transports across electrolytes and SEIs on Li<sup>0</sup> electrode.<sup>42</sup> Since  $J_0$  reflects the intrinsic rate of electron transfer at the Li<sup>0</sup>/ electrolyte interface, any changes in  $J_0$  for Li<sub>3</sub>N SE would clearly indicate the existence of Li<sup>0</sup>-Li<sub>3</sub>N interaction, mainly due to the identical liquid electrolyte used for RE and Li<sub>3</sub>N SE. By analyzing the kinetically controlled region<sup>42</sup> in the voltammograms of RE (Figure 1g) and Li<sub>3</sub>N SE (Figure 1h), the averaged  $J_0$  for RE and Li<sub>3</sub>N SE were calculated as 19.27 mA cm<sup>-2</sup>  $\pm$  1.13 mA cm<sup>-2</sup> and 7.64 mA cm<sup>-2</sup>  $\pm$  0.28 mA  $cm^{-2}$ , respectively (Figure 1i). The noticeable changes in  $J_0$  for Li<sub>3</sub>N SE compared to that of RE confirmed the Li<sup>0</sup>-Li<sub>3</sub>N interaction, in which the electron transfer was also occurring at the Li<sup>0</sup>/Li<sub>3</sub>N interface for Li<sub>3</sub>N SE. Also, the Li<sup>0</sup>-Li<sub>3</sub>N interaction probed by the  $J_0$  supports the lithiophilic feature of Li<sub>3</sub>N, which is discussed further in the later section. Moreover, the lower value of  $J_0$  for Li<sub>3</sub>N SE signifies that the rate of electron transfer with Li<sub>3</sub>N SE was slower compared to that with RE. This is obvious due to the additional electron transfer happening at the Li<sup>0</sup>/Li<sub>3</sub>N interface that should intrinsically be slower than that at the Li<sup>0</sup>/electrolyte interface. Since  $Li_3N$  is a good  $Li^+$  conductor,<sup>19</sup> the charge transfer occurring at the Li<sup>0</sup>/Li<sub>3</sub>N interface further suggests that Li<sup>0</sup> can be electrodeposited and grow from the surfaces of Li<sub>3</sub>N once Li<sub>3</sub>N is in contact with Li<sup>0</sup>. In other words, the heterojunction

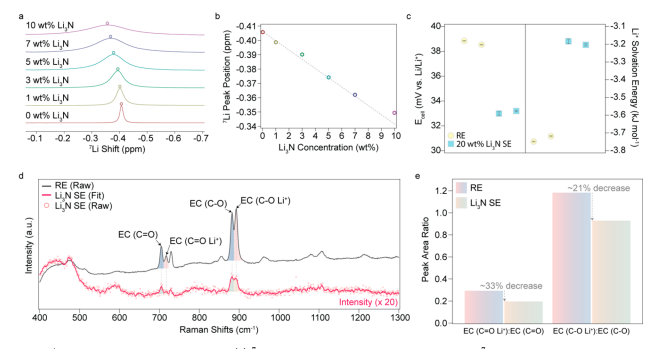


Figure 3. Li<sup>+</sup> solvation environment of Li<sub>3</sub>N SE. (a) <sup>7</sup>Li NMR spectrum of 0–10 wt % Li<sub>3</sub>N in RE. <sup>7</sup>Li NMR peaks for the corresponding weight content of Li<sub>3</sub>N in RE are indicated by the hexagonal symbols. (b) <sup>7</sup>Li NMR peak positions based on 0–10 wt % Li<sub>3</sub>N in RE. The dashed gray line represents the linear fit of the peaks. (c) The cell potential  $(E_{cell})$  and Li<sup>+</sup> solvation energy measurements for RE and 20 wt % Li<sub>3</sub>N SE. Two measurements were taken for each of the electrolytes. The averaged  $E_{cell}$  and solvation energy values with standard errors for RE (1st measurement:  $38.850 \pm 0.024$  mV vs Li/Li<sup>+</sup> |  $-3.748 \pm -0.002$  kJ mol<sup>-1</sup> and 2nd measurement:  $38.540 \pm 0.037$  mV vs Li/Li<sup>+</sup> |  $-3.719 \pm -0.003$  kJ mol<sup>-1</sup>) and 20 wt % Li<sub>3</sub>N SE (1st measurement:  $32.992 \pm 0.139$  mV vs Li/Li<sup>+</sup> |  $-3.183 \pm -0.013$  kJ mol<sup>-1</sup> and 2nd measurement:  $33.197 \pm 0.061$  mV vs Li/Li<sup>+</sup> |  $-3.203 \pm -0.006$  kJ mol<sup>-1</sup>) were obtained by recording the measurements up to 3 s after the cell potentials were stabilized. (d) Raman spectrum for RE and Li<sub>3</sub>N SE. Four major peaks were assigned for EC (C=O), EC (C=O Li<sup>+</sup>), EC (C-O), and EC (C-O Li<sup>+</sup>). (e) Raman peak area ratios of RE and Li<sub>3</sub>N SE for EC (C=O Li<sup>+</sup>) and EC (C-O):EC (C=O Li<sup>+</sup>) retrieved from (d).

formed at the Li<sup>0</sup>/Li<sub>3</sub>N interface can be depicted as the current collector (Li<sup>0</sup> as the electron conductor) and solid electrolyte (Li<sub>3</sub>N as the Li<sup>+</sup> conductor) model, in which the charge transfers occurring at the heterojunction allow electro-deposition of Li<sup>0</sup> on the surfaces of Li<sub>3</sub>N. Hence, the  $J_0$  measurement substantiates the Li<sup>0</sup> electrodeposition at the Li<sup>0</sup>/Li<sub>3</sub>N heterojunction (as observed in Figure 1d,e) by electrochemically probing the Li<sup>0</sup>-Li<sub>3</sub>N interaction. We note that this outcome was particularly crucial as this serves as the first demonstration that  $J_0$  can also be impacted by the suspension electrolyte design, meaning the  $J_0$  of a particular liquid electrolyte can still be modified without changing the liquid components.

Since lower surface area and more compact Li<sup>0</sup> electrodeposition morphologies indicate high CE and low Li nucleation overpotential,<sup>1,40,41</sup> CE and Li nucleation overpotential were measured from LilCu cells with RE and Li<sub>3</sub>N SE to cross-validate the features attained from Figure 1c-f (Figure 1j,k). Figure 1j shows the representative voltage profiles of CE and Li nucleation overpotential measurements for RE and Li<sub>3</sub>N SE. From the voltage profile analyses, it is apparent that Li<sub>3</sub>N SE exhibited a higher CE and lower Li nucleation barrier than those of RE. Moreover, the inset of Figure 1j showed reduced overpotentials for Li<sub>3</sub>N SE, suggesting facilitated Li<sup>+</sup> transport. Duplicate measurements were taken with three identical LilCu cells containing each of the electrolytes to calculate averaged CE (97.48%  $\pm$  0.31% for  $\rm Li_3N$  SE and  $93.01\% \pm 2.21\%$  for RE) and Li nucleation overpotentials  $(56.43 \text{ mV} \pm 5.31 \text{ mV} \text{ for } \text{Li}_3\text{N} \text{ SE and } 88.77 \text{ mV} \pm 5.11 \text{ mV}$ for RE) with standard errors (Figure 1k). Based on the

morphological and electrochemical analyses from Figure 1c–k, three noticeable features of  $Li_3N$  were observed: (i) guided  $Li^0$  electrodeposition along the surface of  $Li_3N$ , (ii) existence of  $Li^0$ -Li<sub>3</sub>N interaction in  $Li_3N$  SE, (iii) improved electrochemical performance of  $Li^0$  anodes, and (iv) facilitated  $Li^+$  transport with  $Li_3N$  in the liquid electrolyte.

Analyses of Li<sub>3</sub>N via First-Principles Calculations. To construe the guided Li<sup>0</sup> electrodeposition along the surfaces of Li<sub>3</sub>N (Figure 1d,e), binding energies and diffusion barriers were computed for Li<sub>3</sub>N via density functional theory (DFT). Based on X-ray powder diffraction (XRD) of Li<sub>3</sub>N, four orientations were found: (001), (100), (002), and (110), of which (001) was the dominant surface of  $Li_3N$  (Figure S2). To relatively compare Li adatom binding energies and diffusion barrier of Li3N, the same analyses were executed with Li0 (110) as (110) surface is most commonly observed for Li<sup>0</sup> anodes.<sup>34,43</sup> Figure 2a shows thermodynamically favorable Li adatom binding sites on Li<sup>0</sup> (110), Li<sub>3</sub>N (001)I(002), Li<sub>3</sub>N (110), and  $Li_3N$  (100) surfaces with the corresponding binding energies of -1.59 eV, -3.44 eV, -1.78 eV, and -1.44 eV, respectively (Figure 2b). The Li adatom binding energy trend in Figure 2b suggests that Li<sub>3</sub>N (001)I(002) surfaces (please note that the (002) surface is equivalent to the (001) surface of  $Li_3N$  were the most favorable Li adatom binding surfaces. This means that Li thermodynamically prefers to get adsorbed onto Li<sub>3</sub>N (001)l(002) surfaces instead of the Li<sup>0</sup> (110) surface. The favored Li adsorption onto Li<sub>3</sub>N surfaces further justifies the change in J<sub>0</sub> observed for Li<sub>3</sub>N SE (Figure 1i), as the DFT results (Figure 2b) indicate that it is thermodynamically more favorable for Li to

bind with Li<sub>3</sub>N rather than with Li<sup>0</sup>. Thereby, the results in Figure 2b rationalize the Li<sup>0</sup>-Li<sub>3</sub>N interaction, illustrating favored Li growth at the Li<sup>0</sup>/Li<sub>3</sub>N heterojunction. Additionally, the binding energies of Li<sub>3</sub>N (001)l(002) surfaces were significantly higher, about two folds, than that of Li<sup>0</sup> (110), Li<sub>3</sub>N (110), and Li<sub>3</sub>N (100) surfaces, in which the binding energies of Li<sub>3</sub>N (110) and (100) were comparable to that of Li<sup>0</sup> (110). Since the Li<sub>3</sub>N (001) surface was experimentally found to be the primary surface (Figure S2),<sup>34,43</sup> this supports the claim that Li<sub>3</sub>N, specifically Li<sub>3</sub>N (001)l(002) surfaces, is lithiophilic, which explains the guided Li<sup>0</sup> electrodeposition along the surfaces of Li<sub>3</sub>N.

With the aim of further supporting the stabilized Li<sup>0</sup> electrodeposition with Li<sub>3</sub>N, the surface diffusion barrier of Li adatom on the  $Li_3N$  (001) surface was calculated as the low diffusion barrier which facilitates surface diffusion of Li to promote uniform  $Li^0$  electrodeposition (Figure 2c).<sup>43-45</sup> In Figure 2c, Li was diffusing from the nitrogen to the adjacent nitrogen. From this diffusion trajectory, a total of eight relative energies, which represent threshold energies needed to adsorb Li adatom across the diffusion pathway, were computed to figure out the diffusion barrier. The diffusion barrier of Li adatom on Li<sub>3</sub>N (001) surface was 0.133 eV (Figure 2d). Then, the diffusion barrier of  $Li_3N$  (0.133 eV) was compared with other prominent SEI inorganic compounds, such as Li<sub>2</sub>O (0.319 eV), Li<sub>2</sub>CO<sub>3</sub> (0.232 eV), LiF (0.169 eV), and LiOH (0.141 eV), to comparatively understand the diffusion barrier for  $\text{Li}_3 \text{N}_{\cdot}^{44}$  According to Figure 2e,  $\text{Li}_3 \text{N}$  has the lowest diffusion barrier, claiming Li<sub>3</sub>N as the most favorable SEI inorganic in terms of Li surface diffusion characteristics. We note that the low diffusion barrier would be an important parameter for ionically conductive inorganics, such as Li<sub>3</sub>N, stabilizing the Li<sup>0</sup> electrodeposition at the Li<sup>0</sup>/inorganic heterojunction. Furthermore, the surface energy of Li<sub>3</sub>N (001) surface (3.60 meV atom<sup>-1</sup> Å<sup>-2</sup>) was higher than that of  $\text{Li}^0$  (110) surface (2.58 meV atom<sup>-1</sup> Å<sup>-2</sup>), in which the higher surface energy of Li<sub>3</sub>N at the Li<sup>0</sup>/Li<sub>3</sub>N heterojunction suppresses dendritic Li<sup>0</sup> growth (Figure S3).<sup>46</sup> Hence, the high binding energy, low diffusion barrier, and high surface energy of Li<sub>3</sub>N support the observed guided and stabilized Li<sup>0</sup> electrodeposition along the surface of Li<sub>3</sub>N observed in Figure 1d,e and Figure S1.

Li<sup>+</sup> Solvation Environment of Li<sub>3</sub>N. <sup>7</sup>Li nuclear magnetic resonance (NMR), relative Li<sup>+</sup> solvation energy, and Raman analyses were performed to investigate the Li<sup>+</sup> solvation environment influenced by  $Li_3N$  in the electrolyte (Figure 3). Figure 3a shows <sup>7</sup>Li NMR spectra of RE (0 wt % Li<sub>3</sub>N) and Li<sub>3</sub>N SE (1–10 wt % Li<sub>3</sub>N in RE). <sup>7</sup>Li NMR peaks were shifted in the downfield direction, meaning the peaks were moving to the higher parts per million (ppm), as the Li<sub>3</sub>N content increased. The peak shifts directly reflect the changes in the Li<sup>+</sup> solvation environment, in which the downfield peak shifts signify decreased electron density around the nuclei of solvated Li<sup>+</sup> in the electrolyte.<sup>47</sup> As the solid particles in the liquid sample agitate the magnetic field during the NMR measurement, the retrieved signals become heterogeneous, which broadens the <sup>7</sup>Li NMR peak. This means that the peak becomes broader as the suspension concentration increases as shown in Figure 3a. Despite the peak broadening as a function of suspension content, all the peak shapes are symmetric, which confirms that the peak position is shifting with respect to the changes in the suspension content. To verify the <sup>7</sup>Li shift stems from dissociated  $Li^+$  from LiPF<sub>6</sub>, not Li<sub>3</sub>N, the <sup>7</sup>Li

NMR spectrum of RE without LiPF<sub>6</sub>, supernatant of Li<sub>3</sub>N SE without LiPF<sub>6</sub>, and Li<sub>3</sub>N SE without LiPF<sub>6</sub> were measured (Figure S4). The spectra in Figure S4 do not exhibit any peaks, meaning that the dissociated Li<sup>+</sup> from Li<sub>3</sub>N was too low to be detected by the <sup>7</sup>Li NMR. Therefore, the <sup>7</sup>Li shift observed in Figure 3a is due to the change in the electron density of dissociated  $Li^+$  nuclei from  $LiPF_6$ , validating that  $Li_3N$  is modifying the solvation environment of Li<sup>+</sup> in the suspension electrolyte. The downfield peak shifts observed in Figure 3a illustrate that the solvated Li<sup>+</sup> in Li<sub>3</sub>N SE became less coordinated with solvating electrolyte species. Since the peak position varied with respect to the Li<sub>3</sub>N contents, the peak positions were plotted in Figure 3b. The peak positions as a function of the Li<sub>3</sub>N content were almost linear up to 7 wt %. Then above 7 wt % Li<sub>3</sub>N, the peak shift was less in magnitude. Based on the suspension electrolyte design, the dispersity of the suspension also matters to obtain reliable electrochemical performance. Despite the <sup>7</sup>Li downfield peak shift observed for 10 wt % Li<sub>3</sub>N in Figure 3b, particle aggregation above 7 wt % was observed (see Figure S4). Since the <sup>7</sup>Li peak shift depends on the surface area to volume ratio<sup>6</sup> of the Li<sub>3</sub>N suspension, Figure 3b correctly shows that the peak shift becomes nonlinear and shifts less above 7 wt %, indicating that the suspension starts to aggregate (reducing the surface area to volume ratio of the suspension) as observed in Figure S5. Hence, 7 wt % Li<sub>3</sub>N content was used to analyze the electrochemical performance, as this weight content was the most reliable for making Li<sub>3</sub>N SE. Please note that optimizing the suspension content for the suspension electrolytes depends on several parameters (i.e., suspension types, dimensions, geometries, electrolyte formulations, and so on), in which the reported optimized content can be changed when the parameters vary.

To further support the changes in the Li<sup>+</sup> solvation environment of Li<sub>3</sub>N, relative Li<sup>+</sup> solvation energies were potentiometrically probed for RE and 20 wt % Li<sub>3</sub>N SE. With symmetric Li<sup>0</sup> electrodes and asymmetric electrolytes configured in H-cell, it became possible to electrochemically measure the relative Li<sup>+</sup> solvation energies of electrolytes (Figure 3c).<sup>11</sup> The reason for the high weight content of  $Li_3N$ used in the Li<sup>+</sup> solvation energy measurement was to vividly observe the direction of the Li<sup>+</sup> solvation energy change for Li<sub>3</sub>N SE. From Figure 3c, 20 wt % Li<sub>3</sub>N SE exhibited lower cell potential  $(E_{cell})$  and more positive Li<sup>+</sup> solvation energy than those of RE. These results were critical as the lower  $E_{cell}$  and higher Li<sup>+</sup> solvation energy cross-check the changes in the Li<sup>+</sup> solvation environment with Li<sub>3</sub>N in the electrolyte as observed from <sup>7</sup>Li NMR measurements (Figure 3a).<sup>6,11</sup> Also, the lower  $E_{cell}$  and higher Li<sup>+</sup> solvation energy of 20 wt % Li<sub>3</sub>N SE translate to the weakly solvating  $Li^+$  environment.<sup>11</sup> This interchangeably means that the interaction between solvated Li<sup>+</sup> and solvating electrolyte species has been weakened by the presence of Li<sub>3</sub>N in the electrolyte, which agrees with <sup>7</sup>Li NMR analysis in Figure 3a. Thereby, we concluded that Li<sub>3</sub>N in the electrolyte modifies Li<sup>+</sup> solvation environment by decreasing the interaction between Li<sup>+</sup> and solvating electrolyte species, which creates a weakly solvating environment. Also, it is worth mentioning that the weakly solvating environment allows facilitated Li<sup>0</sup> electrodeposition and Li<sup>+</sup> desolvation kinetics, demonstrating beneficial features of Li<sub>3</sub>N.<sup>42</sup>

To verify the decreased interaction between  $Li^+$  and the solvating electrolyte species observed in <sup>7</sup>Li NMR and the relative  $Li^+$  solvation energy analyses in Figure 3a-c, Raman

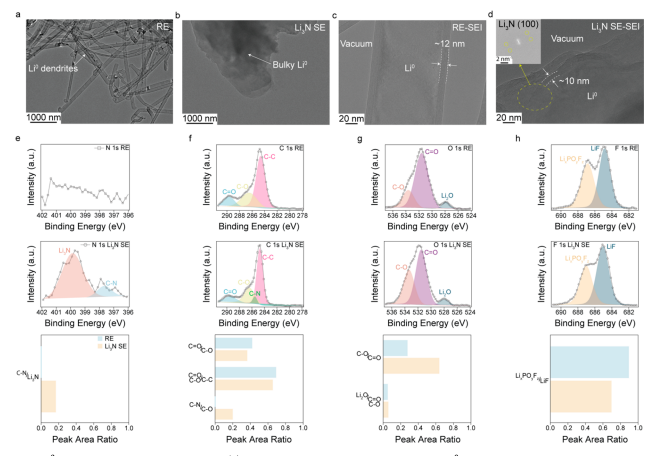


Figure 4.  $Li^0$  interphases with RE and  $Li_3N$  SE. (a) A low magnification cryo-TEM image of  $Li^0$  electrodeposits on Cu TEM grid with RE. The arrow indicates dendritic  $Li^0$  electrodeposits with RE. (b) A low magnification cryo-TEM image of  $Li^0$  electrodeposits on Cu TEM grid with  $Li_3N$  SE. The arrow indicates bulky  $Li^0$  electrodeposits with  $Li_3N$  SE (c) A high magnification cryo-TEM image of  $Li^0$  dendrite and RE-SEI, indicated by the arrows. (d) A high magnification cryo-TEM image of bulky  $Li^0$  and  $Li_3N$  SE-SEI, indicated by the arrows. The inset shows the fast Fourier transform pattern of the image. From the inverse fast Fourier transform of the inset, the dashed yellow circle indicates the position of  $Li_3N$  lattice. (e) N 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. The bottom plot represents the peak area ratio of C–N: $Li_3N$  based on N 1s XPS spectra of RE and  $Li_3N$  SE. (f) C 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. (g) O 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. The bottom plot represents the peak area ratio of C–O:C=O C–O:C–C, and C–N:C–O based on C 1s XPS spectra of RE and  $Li_3N$  SE. (g) O 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. (h) F 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. (h) F 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. The bottom plot represents the peak area ratio of C–O:C=O and  $Li_2O:C=O$  C–O based on O 1s XPS spectra of RE and  $Li_3N$  SE. (h) F 1s XPS spectra of 1 mAh cm<sup>-2</sup> electrodeposited  $Li^0$  on Cu at 1 mA cm<sup>-2</sup> from LilCu cells with RE and  $Li_3N$  SE. The bottom plot represents the peak on F 1s XPS spectra of RE and  $Li_3$ 

spectra of RE and Li<sub>3</sub>N SE were examined to understand Li<sup>+</sup> to solvent coordination (Figure 3d). Due to the solid nanoparticles in the liquid electrolyte (Li<sub>3</sub>N SE), the retrieved intensity for Li<sub>3</sub>N SE was relatively lower than that for RE. Despite the low signal, reliable peaks for alkoxy, EC (C-O), and carbonyl, EC (C=O), groups for EC were observed for  $Li_3N$  SE (Figure 3d). The peaks around 892 cm<sup>-1</sup>, 882 cm<sup>-1</sup>, 718 cm<sup>-1</sup>, and 705 cm<sup>-1</sup> represent solvated alkoxy groups of EC with  $Li^+$ , EC (C-O  $Li^+$ ), nonsolvated alkoxy groups of EC, EC (C–O), solvated carbonyl groups of EC with  $Li^+$ , EC (C= O Li<sup>+</sup>), and nonsolvated carbonyl groups of EC, EC (C=O), respectively.48,49 To qualitatively compare the Li<sup>+</sup>-EC coordination between RE and Li<sub>3</sub>N SE, the corresponding peaks were integrated and normalized to investigate the relative Li<sup>+</sup>-EC coordination, which was denoted as the EC (C=O  $Li^+$ ):EC (C=O) and EC  $(C-O Li^+)$ :EC (C-O) ratios. By calculating the peak area ratios in Figure 3e, the decrease in Li<sup>+</sup>-EC coordination was clearly observed for Li<sub>3</sub>N SE, in which both EC (C=O  $Li^+$ ):EC (C=O) and EC (C-O  $Li^+$ ):EC (C-O) ratios for Li<sub>3</sub>N SE showed approximately 33% and 21% relative

decrease compared to those for RE. The decrease in Li<sup>+</sup>-EC coordination for Li<sub>3</sub>N SE cross-validates the results from <sup>7</sup>Li NMR (the decrease in electron density around the nuclei of solvated Li<sup>+</sup> in Li<sub>3</sub>N SE) and the relative Li<sup>+</sup> solvation energy (weakly solvating Li<sup>+</sup> environment for Li<sub>3</sub>N SE) measurements. Thusly, these results reveal the essential roles of Li<sub>3</sub>N in terms of the Li<sup>+</sup> solvation environment: Li<sub>3</sub>N changes Li<sup>+</sup> solvation environment by decreasing the interaction between Li<sup>+</sup> and solvating electrolyte species, and specifically Li<sub>3</sub>N weakens the Li<sup>+</sup>-solvent coordination that induces the weakly solvating environment in the electrolyte.

Analyses of Li<sup>0</sup> Interphase. Li<sup>0</sup> electrodeposition morphologies and SEIs with RE and Li<sub>3</sub>N SE were investigated via cryogenic transmission electron microscopy (cryo-TEM) to consolidate observed features of Li<sub>3</sub>N in Figures 1–3 (Figure 4a–d). With RE, dendritic Li<sup>0</sup> electrodeposition morphology was observed (Figure 4a), as seen in Figure 1c. On the contrary, a bulky Li<sup>0</sup> electrodeposit was observed with Li<sub>3</sub>N SE (Figure 4b), indicating a more compact Li<sup>0</sup> electrodeposition with Li<sub>3</sub>N in the electrolyte (Figure 1d,e and Figure S1). To

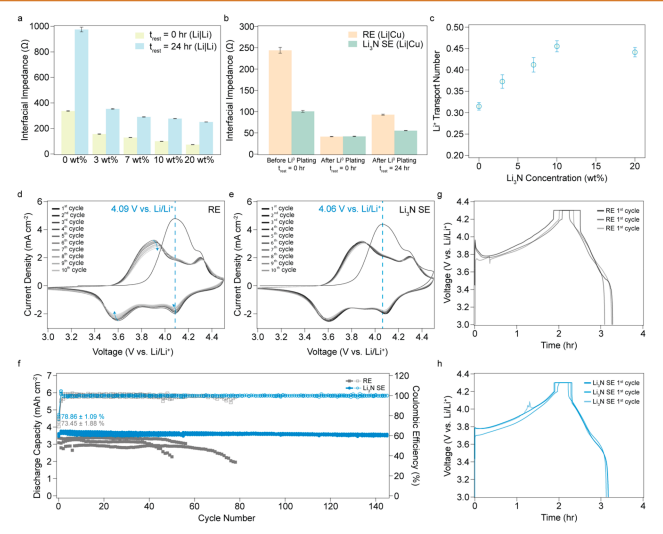


Figure 5. Electrochemical characteristics of Li<sub>3</sub>N SE for Li<sup>0</sup> anodes. (a) Measured interfacial impedances from LilLi cells with 0–20 wt % Li<sub>3</sub>N in RE with respect to the aging time ( $t_{rest} = 0$  and 24 h). Three identical LilLi cells were used for each electrolyte to obtain the averaged interfacial impedances with standard errors. (b) Measured interfacial impedances from LilCu cells at  $t_{rest} = 0$  h and after Li<sup>0</sup> electrodeposited on Cu at  $t_{rest} = 0$  and 24 h with RE and Li<sub>3</sub>N SE. Three identical LilCu cells were used for each condition to obtain the averaged interfacial impedances with standard errors. The current density of 1 mA cm<sup>-2</sup> and capacity of 1 mAh cm<sup>-2</sup> were used for plating Li<sup>0</sup>. (c) Li<sup>+</sup> transport number measured from LilLi cells with 0–20 wt % Li<sub>3</sub>N in RE. Three identical LilLi cells were used for each electrolyte to obtain the averaged Li<sup>+</sup> transport number with standard errors. (d) First 10 cycle cyclic voltammograms of the LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cell with RE. The voltage window and scan rate used are 3.0 to 4.5 V vs Li/Li<sup>+</sup> and 0.1 mV s<sup>-1</sup>. The dashed line indicates the peak position during the first charge cycle of the LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cell with Li<sub>3</sub>N SE. The voltage window and scan rate used are 3.0 to 4.5 V vs Li/Li<sup>+</sup> and 0.1 mV s<sup>-1</sup>. The dashed line indicates the peak position during the first charge cycle of the LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cell with Li<sub>3</sub>N SE. The voltage window and scan rate used are 3.0 to 4.5 V vs Li/Li<sup>+</sup> and 0.1 mV s<sup>-1</sup>. The dashed line indicates the peak position during the first charge cycle of the LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cell with Li<sub>3</sub>N SE. The voltage window and scan rate used are 3.0 to 4.5 V vs Li/Li<sup>+</sup> and 0.1 mV s<sup>-1</sup>. The dashed line indicates the peak position during the first charge cycle of the LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cell with Li<sub>3</sub>N SE. The voltage window and scan rate used are 3.0 to 4.5 V vs Li/Li<sup>+</sup> and 0.1 mV s<sup>-1</sup>. The dashed line indicates the peak position during the first charge cycle

look closely into the SEIs on  $Li^0$  with RE (RE-SEI) and  $Li_3N$  SE ( $Li_3N$  SE-SEI), the thickness of RE-SEI (Figure 4c) and  $Li_3N$  SE-SEI (Figure 4d) was estimated with the high magnification cryo-TEM images. The measured SEI thicknesses of RE-SEI and  $Li_3N$  SE-SEI were about 12 and 10 nm. Furthermore, the fast Fourier transform (FFT) pattern in the Figure 4d inset exhibits the lattice of  $Li_3N$ , indicating the presence of  $Li_3N$  in the  $Li^0$  electrodeposits. From the inverse FFT of the Figure 4d inset, the observed  $Li_3N$  lattice was mapped in Figure S6 for visualization, in which the detected  $Li_3N$  lattice spot is demarcated by a dashed yellow circle in Figure 4d. Therefore, embedded  $Li_3N$  into the  $Li^0$  electrodeposit was found for  $Li_3N$  SE, in which the presence of

embedded  $Li_3N$  in  $Li^0$  electrodeposits further confirmed guided  $Li^0$  electrodeposition behavior along the  $Li_3N$  surface as observed in SEM (Figure 1d,e) and DFT (Figure 2) analyses.

X-ray photoelectron spectroscopy (XPS) was performed to analyze chemistries of  $Li^0$  interphases formed by RE and  $Li_3N$ SE (Figure 4e-h). Based on N 1s narrow XPS scans,  $Li_3N$  and C-N were detected on the  $Li^0$  interphase derived from  $Li_3N$ SE, whereas no N 1s signals were retrieved from the  $Li^0$ interphase derived from RE as RE contains no nitrogen species (Figure 4e). The existence of  $Li_3N$  can be explained by the guided  $Li^0$  electrodeposition along the surface of  $Li_3N$  that gets engulfed during the  $Li^0$  plating (Figure 1d,e, Figure S1, and Figure 4d). We hypothesize that C–N bonds were formed during the reduction of solvent species at the interface junction between Li<sub>3</sub>N and Li<sup>0</sup>, at which the electrolyte should also be reduced at the surface of Li<sub>3</sub>N due to the plating potential below 0 V vs Li/Li<sup>+,3</sup> The ratio of C–N to Li<sub>3</sub>N (C–N/Li<sub>3</sub>N) was calculated to understand which species were dominantly present at the Li<sup>0</sup> interphase. The C–N/Li<sub>3</sub>N ratio was around 0.18, indicating that Li<sub>3</sub>N more dominantly existed at the Li<sup>0</sup> interphase formed by Li<sub>3</sub>N SE (Figure 4e). From C 1s narrow XPS scans (Figure 4f), there were three distinct differences observed between the Li<sup>0</sup> interphase formed by RE and Li<sub>3</sub>N SE: (i) a relative increase of C–O, (ii) lower C–C to C–O + C=O ratio (C-C/C-O C=O), and (iii) C-N species for the Li<sup>0</sup> interphase of Li<sub>3</sub>N SE. It has been found that enrichment of C-O (alkoxides and ethers) over C=O (carbonates) groups in the organic parts of the SEIs on Li<sup>0</sup> provides improved passivity, enhanced Li<sup>+</sup> transport, and physicochemical stability, implying beneficial features of C–O over C=O species in the SEIs.<sup>50–53</sup> Also, C=O + C–O over C-C ratio (C-O C=O/C-C) represents a relative solvent decomposition at the interphase of Li<sup>0</sup> anodes, in which C-C and C-O C=O contents should increase and decrease, respectively, upon a more solvent decomposition and complete reduction of solvent molecules to form Li bound species from C-O and C=O groups.<sup>50-52</sup> The presence of C-N and the low C-N/C-O ratio in Figure 4f also cross-check the observed C-N in Figure 4e. Hence, the lower C=O/C-O and C=O C-O/C-C ratios for  $Li_3N$  SE than those for RE suggest a better SEI passivity and a less solvent decomposition on Li<sup>0</sup> anodes. O 1s narrow XPS scans (Figure 4g), which exhibited higher C-O/C=O and Li<sub>2</sub>O/C=O C-O ratios for Li<sub>3</sub>N SE, showed the same aspects of Li<sup>0</sup> interphase formed by  $Li_3N$  SE as observed in Figure 4f. Moreover, a lower  $Li_xPO_vF_z/$ LiF ratio for Li<sub>3</sub>N SE than that for RE was observed from F 1s narrow XPS scans (Figure 4h). The lower  $\text{Li}_{r}\text{PO}_{v}F_{z}/\text{LiF}$  ratio for Li<sub>3</sub>N SE indicates a reduced solvent decomposition as the oxygen in the  $Li_x PO_v F_z$  complex originates from the solvent decomposition with anions. To further verify the reduced solvent decomposition at the Li<sup>0</sup> interphase of Li<sub>3</sub>N SE, elemental ratios of P/C, F/C, and O/C were calculated (Figure S7). The lower P/C, F/C, and O/C ratios were achieved for Li<sub>3</sub>N SE than those for RE, meaning reduced solvent-derived species for the Li<sup>0</sup> interphase formed by Li<sub>3</sub>N SE. Overall, XPS analyses manifest that less solvent decomposition and better Li<sup>0</sup> interphase passivation characteristics were achieved with Li<sub>3</sub>N SE. This also confirms the impact of the modified Li<sup>+</sup> solvation environment by Li<sub>3</sub>N as observed in Figure 3, in which the decrease in the Li<sup>+</sup>-solvent coordination of solvated Li<sup>+</sup> with Li<sub>3</sub>N in the electrolyte suppressed solvent decomposition and promoted a complete solvent decomposition.

**Electrochemical Performance Analyses.** To understand the electrochemical properties of  $Li^0$  interphase formed by RE and  $Li_3N$  SE, interfacial impedances were measured for RE (0 wt %  $Li_3N$ ) and 3–20 wt %  $Li_3N$  SE from LilLi cells before and after aging<sup>16</sup> the cell for a day (Figure 5a, Figure S8, and Table S1). The interfacial impedances of  $Li^0$  with  $Li_3N$  SE electrolytes were much lower than that with RE, which can be expected from the weakly solvating environment and organic-poor  $Li^0$  interphase<sup>3</sup> with  $Li_3N$ . Also, the temporal characteristics of the interfacial impedance changes for  $Li_3N$  SE were more stable than that for RE. Moreover, the interfacial stability of electrodeposited  $Li^0$  with RE and  $Li_3N$  SE was

investigated by measuring the interfacial impedances before and after Li<sup>0</sup> plating as well as aging the electrodeposited Li<sup>0</sup> for a day from LilCu cells (Figure 5b, Figure S9, and Table S2). The magnitude of the interfacial impedance changes before and after Li<sup>0</sup> plating as well as after aging was lower for  $Li_3N$  SE, demonstrating stable electrodeposition of  $Li^0$  with and temporally stabilized Li<sup>0</sup> interphase formed by Li<sub>3</sub>N SE. The altered Li<sup>+</sup> solvation environment with Li<sub>3</sub>N in the electrolyte contributes to the stabilized temporal characteristics of Li<sup>0</sup> interphase with Li<sub>3</sub>N SE, as the decrease in Li<sup>+</sup>-solvent interaction helps to create organic-poor SEI that mitigates SEI swelling.<sup>3,15</sup> Remarkably, the Li<sup>+</sup> transport number of Li<sub>3</sub>N SE increased with the presence of Li<sub>3</sub>N in the electrolyte (Figure 5c and Table S3). Having a higher Li<sup>+</sup> transport number can reduce the overpotential, as a larger fraction of the applied current is getting used to migrate Li<sup>+</sup>. To cross-validate the effect of the improved Li<sup>+</sup> transport number, the ionic conductivity was measured for RE (0.272  $\pm$  0.008 mS cm<sup>-1</sup>) and Li<sub>3</sub>N SE (0.261  $\pm$  0.013 mS cm<sup>-1</sup>), see Figure S10. The ionic conductivities of RE and Li<sub>3</sub>N SE were comparable to each other, indicating that the improved Li<sup>+</sup> transport number with Li<sub>3</sub>N SE enables reducing the cell overpotential as the electrolyte conductivity remains similar for RE and Li<sub>3</sub>N SE. Accordingly, the decrease in interfacial impedance, increase in Li<sup>+</sup> transport number, and no change in electrolyte ionic conductivity justify the reduced Li<sup>0</sup>-plating/stripping cell overpotential with Li<sub>3</sub>N SE, as observed in Figure 1j.

To verify the electrochemical activity of Li<sub>3</sub>N in Li<sub>3</sub>N SE, cyclic voltammetry scans (from 3 to 4.5 V vs Li/Li<sup>+</sup>) and linear sweep voltammetry (LSV, from 3 to 5 V vs Li/Li<sup>+</sup>) were performed with LilLi1Ni0.8Co0.1Mn0.1 and LilAl cells (Figure 5d,e, Figure S11, and Figure S12). By comparing the representative cyclic voltammograms of LilLi1Ni0.8Co01Mn01 cells with RE (Figure 5d) and Li<sub>3</sub>N SE (Figure 5e), a lower first charging redox potential and less cyclic voltammogram hysteresis,43,54 signifying facilitated Li<sup>+</sup> transport and reversible redox reactions in the full cell with Li<sub>3</sub>N SE, were observed for Li<sub>3</sub>N SE than those for RE. Additional cyclic voltammograms are shown in Figure S11. LSV measurements in Figure S12 revealed that the Li<sub>3</sub>N in the electrolyte remains electrochemically stable as no current overshoot was observed. Also, the cyclic voltammograms of LilNMC811 with Li<sub>3</sub>N SE in Figure 5e did not exhibit additional redox peaks up to 4.5 V vs Li/Li<sup>+</sup>. We hypothesize that oxidizing dispersed Li<sub>3</sub>N in the liquid electrolyte may require a higher voltage to oxidize Li<sub>3</sub>N as the surfaces of Li<sub>3</sub>N are well-wetted by the liquid electrolyte, which can hinder direct contact of dispersed Li<sub>3</sub>N with the cathode materials. To confirm the cycling characteristics of Lil Li<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> full cells with RE and Li<sub>3</sub>N SE, cycling and corresponding voltage profiles are illustrated in Figure 5fh. The full cells with Li<sub>3</sub>N SE yielded prolonged cycle life with a higher first cycle CE (Figure 5f). By comparing the first cycle voltage profiles of RE (Figure 5g) and Li<sub>3</sub>N SE (Figure 5h) retrieved from Figure 5f, a smaller voltage overshoot at the beginning of the first charging was observed for the Li<sub>3</sub>N SE, demonstrating better Li<sup>+</sup> transport and stabilized electrochemical reactions in the full cells.

# CONCLUSIONS

By combining empirical and first-principles calculation analyses of  $Li_3N$  SE, we showed that  $Li_3N$  guides  $Li^0$  electrodeposition along its surface via thermodynamically favored  $Li^0$ - $Li_3N$ interactions, creates the weakly solvating  $Li^+$  environment by decreasing the interaction between Li<sup>+</sup> and solvating electrolyte species, suppresses solvent decomposition on Li<sup>0</sup> anodes, induces organic-poor SEI on Li<sup>0</sup> anodes, facilitates Li<sup>+</sup> transport in the electrolyte, and promotes reversible electrochemical reactions in the full cells. The outcome of this study can potentially serve as one of the scientific guidelines (i.e., inorganics that promote the weakly solvating Li<sup>+</sup> environment) to rationalize the roles played by inorganic compounds in the SEIs on Li<sup>0</sup> anodes. Importantly, it is crucial to realize that SEIs on Li<sup>0</sup> anodes are still electrolyte permeable, as evidenced by SEI swelling properties,<sup>15</sup> in which solvated Li<sup>+</sup> electrolyte species are present within the SEI layer. Therefore, we must understand how the Li<sup>+</sup> solvation environment is influenced and controlled by SEI inorganic species at the SEI/Li<sup>0</sup> interface that dictates Li<sup>+</sup> transport as well as charge transfer behavior and impacts SEI evolution on Li<sup>0</sup> anodes. We believe that the systematic revealment of SEI inorganic features would ultimately allow sophisticated electrolyte engineering strategies that contribute to designing high-performance SEIs on Li<sup>0</sup> anodes for realizing reliable yet practical energy-dense LMBs.

### **METHODS**

**Materials.** The electrolytes were prepared and handled in an Arfilled glovebox with an O<sub>2</sub> concentration below 0.2 ppm and H<sub>2</sub>O concentration below 0.01 ppm. The reference electrolyte (RE) was made with 1 M LiPF<sub>6</sub> in EC:DEC (1:1 v/v) (Gotion LP40) with 10 vol % FEC (Gotion). The Li<sub>3</sub>N suspension electrolyte (Li<sub>3</sub>N SE) was prepared by mixing indicated amount (3–20 wt %) of Li<sub>3</sub>N nanoparticles (80–100 nm in diameter, Nanoshel) with RE. The default weight content of Li<sub>3</sub>N for Li<sub>3</sub>N SE was 7 wt %, otherwise indicated. High purity Li foil (750  $\mu$ m thick, 99.9%, Alfa Aesar), Cu foil (Pred Materials), 11  $\mu$ m thick polyethylene separator (W-Scope), and ~4 mAh cm<sup>-2</sup> NMC811 (Targray) were used to make 2032-type coin cells for LilLi and LilCu, and LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> configured cells.

DFT Calculation. The calculations were performed with the DFT code GPAW.<sup>55-57</sup> The electron exchange-correlation was described by Perdew-Burke-Ernzerhof functional within the generalized gradient approximation.<sup>58</sup> The valence electrons are expressed in plane wave basis sets and the core electrons with projector augmented wave (PAW) method. The energy cutoff of 500 eV was applied to all the calculations. The k-point grid of  $3 \times 3 \times 1$  was used for all Li<sub>3</sub>Nslab calculations, and  $5 \times 3 \times 1$  was used for Li slab calculations. The energy and forces per atom were converged to within  $10^{-4} \ eV$  and  $10^{-3}$  eV/Å, respectively. For the slab calculations, the vacuum of 15 Å thickness is used to reduce periodic artificial interactions. The Li<sub>3</sub>Nslab consists of six layers of unit cells (spanning approximately 15 Å) in the z-direction. During geometry optimization, the top 4/5/4/4layers are allowed to relax for Li (110)/Li<sub>3</sub>N (001)l(002)/Li<sub>3</sub>N (100)/Li<sub>3</sub>N (110) adsorption calculations. The diffusion pathways were calculated via the machine-learning nudged elastic band (ML-NEB) using the CatLearn package,<sup>59</sup> with nine images between the initial and the final configurations.

**XRD.** X-ray diffraction (PANalytical Empyrean with a Cu(K $\alpha$ ) X-ray source) was used to examine the crystal structural of Li<sub>3</sub>N nanopowder. The Li<sub>3</sub>N nanopowders were placed on a glass slide followed by Kapton taping to seal the nanopowders. Then, the diffraction patterns were collected from 10° to 70° using a step size of 0.01.

**SEM/EDXS.** All the samples were rinsed with pure diethyl carbonate (DEC) and dried inside the Ar glovebox before microscopy. SEM was performed using a Thermo Fisher Scientific Apreo S LoVac Scanning Electron Microscope. Micrographs were taken at a potential of 5 kV and beam current of 50 pA, and the electron signal was collected via the Everhart–Thornley Detector (ETD). Energy dispersive X-ray spectroscopy data was collected with

a Bunker XFlash 6–60 detector at a potential of 10 kV and beam current of 800 pA.

Cryo-TEM. Cu TEM grids were utilized as the working electrode for Li<sup>0</sup> electrodeposition from LilCu TEM grid cells. 0.1 mAh cm<sup>-2</sup> amount of Li was electrodeposited galvanostatically at 1 mA  $\rm cm^{-2}.$ After the electrodeposition of Li<sup>0</sup>, the coin cells were disassembled in an argon-filled glovebox. Then, the electrodeposited TEM grids were rinsed with 60  $\mu$ L of DEC and dried before freezing the sample with liquid nitrogen. After, the sample was instantly submerged in liquid nitrogen to rapidly expose the sample to cryogen without the ambient exposure. For cryo-TEM characterization, while immersed in liquid nitrogen, the sample was loaded into the Gatan side-entry cryotransfer holder (Gatan model 626) and inserted into the TEM column. The cryo-TEM holder uses a specialized cryoshutter to prevent air exposure and ice condensation onto the sample. Once inside the TEM column, the temperature was maintained at approximately -178 °C. Cryo-TEM experiments were performed on a Thermo Fisher Titan 80-300 environmental scanning transmission electron microscope operated at an accelerating voltage of 300 kV. The instrument was equipped with an aberration corrector in the image-forming lens, which was tuned before each sample analysis. Cryo-TEM images were acquired by a Gatan K3 IS direct-detection camera in the electroncounting mode. The images were taken with an electron dose rate of around 100  $e^{-}/Å^{2}/s$ , a total of 5 frames were taken with 0.1 s per frame for each image.

Li<sup>+</sup> Solvation Energy Measurements. The Li<sup>+</sup> solvation energies of RE and Li<sub>3</sub>N SE were probed by Biologic VMP3 using H-cell that contains control electrolyte (1 M LiFSI in DEC)lsalt bridge (1 M LiFSI in DME/DOL)ltesting electrolytes (RE and 20 wt % Li<sub>3</sub>N SE) with submerged Li metal into the control electrolyte and testing electrolytes. The cell potentials and solvation energies for each of the testing electrolytes were recorded for 3 s to obtain averaged values. The detailed apparatus setup can be found in the literature.<sup>11</sup>

<sup>7</sup>Li NMR. An NMR tube with a coaxial insert was used for 1D <sup>7</sup>Li NMR experiments. The outer tube contained freshly prepared RE and (1-10 wt %) Li<sub>3</sub>N SE. The inner tube contained deuterated solutions as external references. <sup>7</sup>Li NMR was performed on a Varian Inova 500 MHz NMR spectrometer, and the chemical shifts were referenced to 1 M LiCl in D2O at 0 ppm.

**Raman.** Raman spectra were collected with a 532 nm green laser at a Horiba XploRA+ confocal microscope. The electrolytes, RE and  $Li_3N$  SE, were sealed in quartz cuvettes under Argon environment.

**XPS.** The XPS scans were performed on a PHI Versaprobe I with a monochromatized Al K $\alpha$  X-ray source. The XPS chamber was kept at a pressure of about 10<sup>-7</sup> Pa during the measurements. For the sample preparation, electrodeposited Li<sup>0</sup> on Cu with the capacity and current density of 1 mAh cm<sup>-2</sup> and 1 mA cm<sup>-2</sup> was rinsed with 60  $\mu$ L of DEC in an Ar filled glovebox. A vacuum transfer vessel was used to transfer the samples directly from the Ar glovebox to the vacuum transfer chamber of the instrument to prevent air exposure. High-resolution spectra were calibrated using the C 1s peak at 284.6 eV.

Electrochemical Measurements. The RE (1 M LiPF<sub>6</sub> in EC:DEC (1:1 v/v) with 10 vol % FEC) and Li<sub>3</sub>N SE (1–20 wt % Li<sub>3</sub>N in RE) were used for the electrochemical measurements. Twenty  $\mu$ L of specified electrolytes were used for LilCu, LilLi, Lil  $Li_1Ni_{0.8}Co_{0.1}Mn_{0.1}$  cells. The cyclic voltammetry with ultramicroelectrodes for measuring the exchange current density of RE and 20 wt % Li<sub>3</sub>N SE was measured by following the literature.<sup>42</sup> In brief, three electrodes were used: Li metal as the reference/counter electrodes and Cu ultramicroelectrode as the working electrode. The scan rate was 20 V s<sup>-1</sup>. The voltage window was -1.75 to 2.75 V vs Li/Li<sup>+</sup>. Three measurements were performed to measure averaged exchange current density with statistics. The exchange current density of RE and  $Li_3N$  SE was calculated by analyzing the kinetically controlled region in the cyclic voltammograms.<sup>42</sup> The interfacial impedances were measured by Biologics VMP3 with the frequency ranging from 100 mHz to 7 MHz. For the Li<sup>+</sup> transport number measurements, Lil Li symmetric cells were precycled at 0.5 mA cm<sup>-2</sup> current density and 1 mAh cm<sup>-2</sup> areal capacity for 5 cycles. A constant voltage bias of 10 mV was applied for 1 h to obtain initial  $(I_0)$  and static  $(I_s)$  currents.

The interfacial impedance measurements were carried out before and after the constant voltage step to obtain interfacial resistance before  $(R_0)$  and after  $(R_s)$ . The ionic conductivity of electrolytes was measured by Biologics VMP3 with the frequency ranging from 1 MHz to 100 MHz using symmetric stainless-steel (SS) cells with the 11  $\mu$ m thick separator (SSISS cells). The area of 2 cm<sup>2</sup>, distance of 11  $\mu$ m, and measured solution resistance were used to calculate the ionic conductivity of specified electrolytes. For cycling LilCu, LilLi, and Lil Li<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cells, the measurements were recorded by Arbin and Land instruments. For CE and nucleation overpotential measurements with LilCu cells, current density of 0.5 mA cm<sup>-2</sup> was applied with cutoff voltage of 1 V. For cyclic voltammetry, Lil  $Li_1Ni_{0.8}Co_{0.1}Mn_{0.1}$  full cells were cycled at 0.1 mV s<sup>-1</sup> with the voltage window of 3.0 to 4.5 V vs Li/Li<sup>+</sup> with Biologics VMP3. For the linear sweep voltammetry, the current profiles were obtained from LilAl cells with RE and Li<sub>3</sub>N SE, in which the used voltage window and sweep rate were 3.0-5.0 V vs Li/Li<sup>+</sup> and 1 mV s<sup>-1</sup>. For cycling Lil Li1Ni0.8Co0.1Mn0.1 full cells, specified charging and discharge C rates were applied with the voltage window of 3.0-4.3 V, where the constant current goes up to 4.2 V and the constant voltage was held at 4.3 V with the cutoff current value of 0.05 C.

# ASSOCIATED CONTENT

### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsnano.2c12470.

Morphologies of electrodeposited Li<sup>0</sup>, XRD measurements of Li<sub>3</sub>N suspension, surface energies of Li<sub>3</sub>N (001) and Li<sup>0</sup> (110) surfaces, <sup>7</sup>Li NMR spectra of RE and Li<sub>3</sub>N SE without LiPF<sub>6</sub>, pictures of 7, 10, and 20 wt % Li<sub>3</sub>N SE, overlaid inverse FFT cryo-TEM image of Figure 4d inset, calculated P/C, F/C, and O/C ratios from XPS for electrodeposited Li<sup>0</sup> with RE and Li<sub>3</sub>N SE, Nyquist plots of LilLi and LilCu cells, ionic conductivities of RE and Li<sub>3</sub>N SE, cyclic voltammograms of LilLi<sub>1</sub>Ni<sub>0.8</sub>Co<sub>0.1</sub>Mn<sub>0.1</sub> cells with RE and Li<sub>3</sub>N SE, linear sweep voltammetry of RE and Li<sub>3</sub>N SE, averaged values of the interfacial impedances for LilLi and LilCu cells, and averaged Li<sup>+</sup> transport numbers (PDF)

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# **Author Contributions**

M.S.K. and Y. Cui conceived the idea and conceptualized the work. M.S.K. performed the experiments and analyzed the data with the guidance from Y. Cui. M.S.K. and Y.Cui wrote the manuscript. Z.Z. performed cryo-TEM analysis. J.W. performed DFT calculations. S.T.O. performed XPS analysis. Z.Y. measured the transport number of the electrolytes. Y. Chen performed <sup>7</sup>Li NMR of the electrolytes. S.C.K. measured the solvation energy of the electrolytes. D.B. performed transient voltammetry measurements with ultramicroelectrodes. Y.Y. helped electrochemical measurements. Z.H. performed XRD measurements. P.S. provided helpful discussions. S.F.B. reviewed the manuscript. Y. Cui, Z.B., and J.Q. supervised the overall studies. All the authors discussed the manuscript and provided comments.

### Notes

The authors declare no competing financial interest.

# ACKNOWLEDGMENTS

We acknowledge support from the Assistant Secretary for Energy Efficiency and Renewable Energy, Office of Vehicle Technologies of the US Department of Energy under the Battery Materials Research (BMR) Program and Battery 500 Consortium. Z.Z. acknowledges the support from Stanford Interdisciplinary Graduate Fellowship. S.T.O. acknowledges support from the Knight Hennessy scholarship for graduate studies at Stanford University. J.W. acknowledges computational resources from the Sherlock cluster at Stanford University.

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