Confining electrodeposition of metals in structured electrolytes

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Electrochemical cells based on alkali metal (Li, Na) anodes have attracted significant recent attention because of their promise for producing large increases in gravimetric energy density for energy storage in batteries. To facilitate stable, long-term operation of such cells a variety of structured electrolytes have been designed in different physical forms, ranging from soft polymer gels to hard ceramics, including nanoporous versions of these ceramics that host a liquid or molten polymer in their pores. In almost every case, the electrolytes are reported to be substantially more effective than anticipated by early theories in improving uniformity of deposition and lifetime of the metal anode. These observations have been speculated to reflect the effect of electrolyte structure in regulating ion transport to the metal electrolyte interface, thereby stabilizing metal electrodeposition processes at the anode. Here we create and study model structured electrolytes composed of covalently linked polymer grafted nanoparticles that host a liquid electrolyte in the pores. The electrolytes exist as freestanding membranes with effective pore size that can be systematically manipulated through straightforward control of the volume fraction of the nanoparticles. By means of physical analysis and direct visualization experiments we report that at current densities approaching the diffusion limit, there is a clear transition from unstable to stable electrodeposition at Li metal electrodes in membranes with average pore sizes below 500 nm. We show that this transition is consistent with expectations from a recent theoretical analysis that takes into account local coupling between stress and ion transport at metal–electrolyte interfaces.

Electrochemical cells based on lithium and sodium metal anodes are considered among the most versatile platforms for high-energy electrical energy storage. Unfortunately, unstable dendritic metal deposition at currents below the diffusion limit lead to premature cell failure by internal short circuits. In a volatile electrolyte, the omnic heat generated by these short poses significant safety risks. The article reports that electrodeposition of metals can be stabilized by confining ion transport to length scales below a few hundred nanometers. It is also shown that dendrite growth can be arrested in electrolytes with mechanical moduli well below that of the metal. This finding contradicts current orthodoxy, which holds that solid-state electrolytes with moduli higher than the metal are required for preventing dendrite growth.

Significance

Rechargeable electrochemical cells based on alkali metal anodes provide opportunities to substantially push the energy storage limits of batteries. Such cells achieve this feat both by increasing the amount of electrical energy that can be stored on a mass or volume basis at the anode and by enabling more flexible material choices for the cathode (1–6). Parasitic reactions between the chemically reactive anode (7–11) and proliferation of rough, dendritic/mossy electrodeposition at the metal anode during battery recharge have been reported to reduce stability of the cells by increasing the likelihood of failure by internal short circuits and by increasing the surface area of reactive metals in contact with electrolytes (12, 13).

The stability analysis of electrodeposition reported by Monroe and Newman (14, 15) (MN) employs continuum modeling of ion transport processes and mechanics in bulk electrolytes to predict morphological transitions at a metal anode. This work showed that marginal stability is achieved at any current density in an electrolyte with mechanical modulus about twice that of the metal electrode. One study using block copolymer electrolytes based on hard polystyrene and ion conductive poly(ethylene oxide) segments to systematically adjust the electrolyte modulus provided early support for this prediction (16–19). This finding, however, stands in contrast to a large and growing body of recent experimental reports which show that rough deposition of metals can be inhibited in gel-like electrolytes (20, 21), cross-linked polymer electrolytes (18, 22, 23), and in ion-conducting polymer electrolytes (24–26), all with mechanical moduli three or more orders of magnitude below that of the metal electrode. Recently, Zhao et al. (27) showed that a composite polymer electrolyte with immobilized ions and ceramic fillers can indeed inhibit dendrites at ambient conditions. Even more dramatic are reports which show that dendrite-induced short circuits in lithium and sodium batteries can be delayed or even eliminated at low and moderate current densities in liquid electrolytes in which a fixed fraction of anions is maintained at the interface (28–33), or in liquid electrolytes in which halide ions are present at the electrode–electrolyte interface (11, 34–37). Essentially nothing is known about the working mechanisms by which such low-modulus materials prevent metal dendrite proliferation at either low or high current densities, relative to the diffusion-limiting current J∞, and a variety of qualitative arguments have been proposed to explain their working mechanism.

The present study is motivated by recent theoretical work by Tikekar et al. (7, 38), which extended the MN analysis to consider the effect of local stress–transport coupling with ion migration profiles and the impact such coupling has on stability of metal electrodeposition. This analysis shows that at current densities below J∞, stable electrodeposition of metals can be achieved under a broad range of conditions, including in electrolytes with shear moduli as low as 0.1 MPa (i.e., more than four orders of magnitude lower than that of the metal electrode). Another conclusion from the analysis is that electrolytes created by hosting simple liquids in porous materials should be able to produce large increases in gravimetric energy density for energy storage in batteries. To facilitate stable, long-term operation of such cells a variety of structured electrolytes have been designed in different physical forms, ranging from soft polymer gels to hard ceramics, including nanoporous versions of these ceramics that host a liquid or molten polymer in their pores. In almost every case, the electrolytes are reported to be substantially more effective than anticipated by early theories in improving uniformity of deposition and lifetime of the metal anode. These observations have been speculated to reflect the effect of electrolyte structure in regulating ion transport to the metal electrolyte interface, thereby stabilizing metal electrodeposition processes at the anode. Here we create and study model structured electrolytes composed of covalently linked polymer grafted nanoparticles that host a liquid electrolyte in the pores. The electrolytes exist as freestanding membranes with effective pore size that can be systematically manipulated through straightforward control of the volume fraction of the nanoparticles. By means of physical analysis and direct visualization experiments we report that at current densities approaching the diffusion limit, there is a clear transition from unstable to stable electrodeposition at Li metal electrodes in membranes with average pore sizes below 500 nm. We show that this transition is consistent with expectations from a recent theoretical analysis that takes into account local coupling between stress and ion transport at metal–electrolyte interfaces.

Significance

Electrochemical cells based on lithium and sodium metal anodes are considered among the most versatile platforms for high-energy electrical energy storage. Unfortunately, unstable dendritic metal deposition at currents below the diffusion limit lead to premature cell failure by internal short circuits. In a volatile electrolyte, the omnic heat generated by these short poses significant safety risks. The article reports that electrodeposition of metals can be stabilized by confining ion transport to length scales below a few hundred nanometers. It is also shown that dendrite growth can be arrested in electrolytes with mechanical moduli well below that of the metal. This finding contradicts current orthodoxy, which holds that solid-state electrolytes with moduli higher than the metal are required for preventing dendrite growth.


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prevent rough electrodeposition of any metal when the average pore size of the host falls below a certain critical electrolyte-dependent value. This prediction is important because it means that solid-state electrolytes are not required to stabilize electrodeposition at metal anodes in batteries; it is also testable. Here, we design structured electrolytes based on cross-linked nanoparticle–polymer hybrid membranes in which the effective pore size can be facilely manipulated by changing the volume fraction of particles in the precursor material. We also use these electrolytes to systematically investigate the effect of electrolyte network structure and mechanical properties on the stability of metal deposition at currents close to the critical current density, \( J^* \). The study takes advantage of an optical visualization technique that allows time-dependent changes in morphology of the metal–electrolyte interface to be directly imaged. Used in combination with the analysis of Tikekar et al. (38), these efforts are shown to lead to a comprehensive understanding of how and why structured electrolytes that do not meet the MN modulus criteria are able to prevent metal dendrite proliferation.

Fig. 1A shows the synthetic route used to create cross-linked hairy nanoparticle-based electrolytes. Briefly, silica nanoparticles (diameter \( \sim 10 \) nm) are first grafted in aqueous solution with an alkoxysilane terminated oligomeric polyethylene glycol (PEO-OH, \( M_w = 500 \) g/mol). As shown in our previous work (18, 39, 40), the approach produces \( \text{SiO}_2 \) nanoparticles densely grafted (\( \sim 1 \) chain per nm\(^2\); from thermal gravimetric analysis) with PEO chains bearing a reactive terminal hydroxyl group. The resultant \( \text{SiO}_2–\text{PEO}–\text{OH} \) nanoparticles are subsequently used as multifunctional node points for cross-linking poly(propylene oxide) (PPO, \( M_w = 2,000 \) g/mol) chains functionalized with isocyanate groups at both ends. A straightforward calculation shows that on average there are \( \sim 300 \) PEO–OH molecules anchored to a single \( \text{SiO}_2 \) nanoparticle, meaning that materials with a high degree of cross-linking are achieved. Additionally, because the PEO chains are short and the PPO linkers are amorphous at ambient temperatures, the cross-linked materials exist as free-standing membranes with high flexibility and toughness for silica content of 6.5% (by weight) (Fig. 1B). The large number of grafting sites available on each \( \text{SiO}_2 \) nanoparticle makes it possible to introduce different functionalities, including ionic species (41) or flame retardants such as phosphates (42) in the materials. Cross-linking these nanostructures provides a facile route to nanoporous membranes in which the surface chemistry of the pores can be manipulated for regulating the ion transport characteristics or for preventing thermal runaway in lithium batteries. Furthermore, by varying the volume fraction of particles it is possible to systematically change the configuration of the particle-tethered PEO and PPO chains in the cross-linked materials to create membranes with vastly different pore sizes and mechanical properties.

It is possible to monitor the progress of the cross-linking reaction by measuring the time-dependent development of elasticity in the materials by means of dynamic shear rheological measurements (Fig. 1C). Specifically, we employed oscillatory shear to continuously measure the storage and loss modulus of the initial slurry of PEO-grafted nanoparticles and PPO-disocyanate chains at \( 60^\circ \text{C} \) using a small-amplitude shear strain of 1% and a fixed oscillatory shear frequency of 10 Hz. It is observed that the storage modulus increases over time indicative of the cross-linking process, which reaches its maximum value in about 24 h. It is important to note that although the chemical reaction between an isocyanate and hydroxyl group is very fast, the overall cross-linking reaction kinetics are under diffusion control as it becomes progressively harder for PPO chains tethered to one particle to bond to a neighboring site on another particle to increase network elasticity. It can also be seen from Fig. 1C that the loss modulus decreases and rolls over to very low values at long times, consistent with the gradual loss of dissipation processes as the network becomes more fully cross-linked. These effects are most dramatically demonstrated by comparison of the storage and loss moduli for the unlinked PPO polymer. It is seen that whereas the loss modulus dominates storage for the unlinked, liquid-like material, the storage modulus dominates as cross-linking nears completion. It is remarkable that at just 6.8 wt % of \( \text{SiO}_2 \) in the materials there is increment of over five orders of magnitude in modulus.

More in-depth knowledge of the configuration and dynamics of the linker polymer chains is possible from dielectric relaxation measurements. It is known that polymers such as PPO and cis-1,4-polyisoprene possess type-A dipole that facilitate alignment of the polymer chain end-to-end vector in the direction of an imposed electric field; thus one can quantify the end-to-end chain relaxation from the dielectric loss spectrum (43, 44). Additionally, from the magnitude of the loss peak, one can calculate the dielectric strength, which for flexible, amorphous chains can be used to estimate the relative degree of chain stretch (45). SI Appendix, Fig. S1 A and B shows the dielectric loss from the cross-linked PPO at various frequencies and temperatures. The experimental data were fitted using the Havriliak–Negami (H-N) equation: \( e'(\omega) = e_\infty + \Delta e/(1 + i\omega\tau_\beta)^\alpha \), where \( \alpha \) and \( \beta \) are associated with broadening dielectric peaks, \( \tau \) represents the H–N relaxation time, \( \omega \) is frequency, \( e_\infty \) is dielectric constant (\( e' \)) at the high-frequency limit, and \( \Delta \) is the dielectric strength. The relaxation times at various temperature obtained from the H-N fits are shown in SI Appendix, Fig. S1C. It can be seen that the relaxation time for all temperatures follow the Vogel–Fulcher–Tammann (VFT) model, represented by the continuous lines in the figure. The VFT model is given as \( \tau = A \exp(-B/(T - T_h)) \), where \( A \) is the prefactor, \( B \) is the activation energy, and \( T_h \) is reference temperature, shown in SI Appendix, Table S1. It can be seen that the activation energy of the relaxation dynamics of PPO polymer increases dramatically from 9.8 kJ/mole to 20 kJ/mole as a result of cross-linking, which is consistent with expectations based on the molecular weight of the polymers and high levels of cross-linking achieved (18). Comparison of the dielectric strength for the untethered and tethered PPO indicates that the linkers are also highly stretched (shown in SI Appendix, Fig. S1D). For random coil type-A dielectric polymer chains, the dielectric strength \( \Delta \) is proportional to the mean squared end-to-end vector of the polymer chain. The results in SI Appendix, Fig. S1D show that \( \Delta \) measured using the cross-linked membranes is \( \sim 250 \) times larger than the corresponding value for the linear unlinked polymer. Considering that the molecular weight of the PPO chains is just
2 kDa, the large increases in Δε also imply that dipoles on the particle tethered chains are also highly correlated.

The cross-linked hairy nanoparticles provide an opportunity to understand the underlying physics of metal electrodeposition in heterogeneous environment, because these materials serve as model nanoporous structures such that the silica nanoparticles act as nonconductive physical barriers and the polymeric chains in the interparticle gaps as pathways for ion migration. In this context, the volume fraction of silica nanoparticles in the entire composite was varied and the corresponding interparticle distance was estimated by assuming random close packing (r.c.p.) arrangement of spherical nanoparticles: \( d_{p-p} = d \left( (0.63/\Phi) \right)^{1/3} \), as reported in SI Appendix, Table S2 for different weight fraction of silica content. We utilized transmission electron microscope (TEM) to analyze the spatial arrangement of the nanoparticles in the polymer composite as shown in SI Appendix, Fig. S24 for r.c.p. pore sizes of 20, 100, and 500 nm. The interparticle distance of the silica nanoparticles was further measured from the TEM images and the corresponding variation was fitted to a normal distribution function obtained from the respective mean and SDs and plotted as an inset of the images. The average interparticle distances measured from TEM are observed to be similar to r.c.p. pore sizes (Fig. 2A).

However, the variance of the distribution is seen to increase with decreasing silica volume fractions (SI Appendix, Fig. S3). We further performed frequency sweep measurements at a constant amplitude of 5% for different cross-linked PPO as well as neat PPO polymer. The storage and loss modulus of obtained from the measurements are plotted in SI Appendix, Fig. S2 and C, respectively. The results show that the composite materials at all cross-linking densities have higher G’ than G", indicating elastic behavior that is unseen at such low core volume fractions in colloidal suspensions. The elastic modulus is seen to progressively increase with increasing core volume fraction; the associated cage volume (\( kT/G’ \)) is reported in Fig. 2A. Membranes with r.c.p. pore size 20 nm show the highest shear modulus of \( \sim 1 \) MPa, while in the uncross-linked state the materials are simple Newtonian liquids. The higher variance of spatial distribution of silica nanoparticles and reminiscence of elasticity at low core fractions indicate that the particles form hierarchical string-like mass fractals due to the fixed length of the cross-linker (PPO), as shown in Fig. 2B, also confirmed using small-angle X-ray scattering in our previous work (18).

Next, we systematically study the morphology of the lithium–electrolyte interface in the cross-linked hairy nanoparticles soaked with liquid electrolyte using direct visualization of the lithium metal anode during electrodeposition in an optical cell. The cell comprises a lithium metal and stainless-steel electrodes separated by a chamber containing electrolyte of 1 M LiPF6/EC/ DMC (10). For the conditions used in the experiments, the diffusion-limited current density for the different membranes were calculated from the measured conductivity and transference number: \( J^* = 2zc_FD_{app}t_c(L/t_c)^{-1} \). Here, \( z \) is the charge number of cation, \( F \) is the Faraday constant, \( c_F \) is salt concentration, \( t_c \) is the anion transference number, and \( L \) is the interelectrode spacing (1 mm) and \( D_{app} \) is the diffusion coefficient. Using the Bruce–Vincent method (33) the transference number (SI Appendix, Fig. S4) was determined to be 0.38 for the cross-linked nanoparticles (r.c.p. = 20 nm). The variation of conductivity and critical current density (\( J^* \)) for the cross-linked nanoparticles at different pore sizes are reported in SI Appendix, Fig. S5. It is seen that \( J^* \) varies in the range \( \sim 5 \times \) m A/cm² for the materials studied. Using the previously reported values of Li⁺ transference number (0.36) and lithium ion diffusivity (3 \( \times \) \( 10^{-6} \) cm²/s) for 1 M EC/DMC-LiPF6 electrolyte, the diffusion-limited current density can be estimated to be \( 9 \times \) m A/cm² for the same interelectrode spacing. (46, 47).

Fig. 3A reports snapshots of the negative electrode at different stages of electrodeposition for the cases without any separator, with a commercial glass-fiber separator and cross-linked hairy nanoparticles (r.c.p. pore size = 20 nm) at a constant current density of 8 mA/cm², which is 0.9 times the \( J^* \) calculated for liquid electrolyte. Through visual inspection, it is obvious that Li electrodeposition in the absence of a separator is uneven, mossy, and blistered, which is indicative of the side reactions between the electrolyte and electrolyte as well as the porous nature of the deposited layers. At first appearance, the deposits are already quite large, at least 100 μm in diameter, which is much larger than typical Li dendrite nucleate sizes assumed in the literature (6). Deposition studies based on a commercial glass-fiber separator show that deposition is relatively compact until a capacity of around 2 mAh/cm² (i.e., ~10 μm/cm² Li is transferred), where after dendritic structures develop and proliferate in time, ultimately piercing the separator and continue their growth through the separator. This sequence is associated with the short-circuiting phenomena described in the Introduction and can lead to catastrophic cell failure by fire, explosion, or both. In comparison with the cases with and without separator, the deposition pattern with the cross-linked hairy nanoparticles display uniform deposition for entire 1 h of the charging process. The accelerated movies of electrodeposition are provided for all of the three cases. Movies S1–S3 are 10-s movies of 1,500-s-long electrodeposition for neat electrolyte, with glass-fiber separator and cross-linked nanoparticles (r.c.p. = 20 nm), respectively.

By tracking the advancing front of the Li electrode over time, it is possible to quantify the dendrite growth as a function of time (Fig. 3B). The apparent growth rate obtained from the initial time through the assumption of linearity is shown in the inset. The larger-sized error bars for the neat electrolyte case is a reflection of the unevenness of electrodeposition. The apparent growth rate of dendrite in the commercial glass-fiber separator is
calculated to be \( \approx 0.13 \mu m/s \), while for the cross-linked hairy nanoparticles it is estimated to be \( 0.03 \mu m/s \), which is modestly larger than the theoretical rate \( (0.011 \mu m/s) \) for perfectly smooth electrodeposition at \( 8 \text{ mA/cm}^2 \).

This idea that one could control the stability of electrodeposition by manipulating the nanostructure of an electrolyte (SI Appendix, Fig. S6) was previously discussed in the theoretical analysis by Tikekar et al. (7, 38) and has also been adapted in an ad hoc manner to explain previous experimental results (48, 49). In this analysis, deposition in membranes with pores larger than the critical nucleate diameter at which electrodeposition is unstable is thought to lead to unchecked growth of dendrites that penetrate through these pores, as in the case of the glass-fiber separator. In contrast, in the cross-linked hairy nanoparticles the pore size of 20 nm is evidently much smaller than the nucleate size, leading to a significant retardation of dendrite growth. To determine the effect of solid electrolyte interphase formation and chemical instability on the topological features observed during the electrodeposition process, we performed postmortem analysis of the lithium electrode after repeated charge and discharge cycling in a coin cell. Specifically, we utilized a symmetric lithium battery with lithium electrode after repeated charge and discharge cycling in a coin cell. Specifically, we utilized a symmetric lithium battery with glass-fiber separator. In contrast, in the cross-linked hairy nanoparticles the pore size of 20 nm is evidently much smaller than the nucleate size, leading to a significant retardation of dendrite growth.

Fig. 3. Stability of electrodeposition by direct visualization of electrode. (A) Snapshots of the electrode and electrolyte every 15 min during charging at the rate of \( 8 \text{ mA/cm}^2 \). (B) Height of dendrite at various points of the electrode for the initial 1,500 s. (Inset) Comparison of the growth rate by assuming a linear growth for electrodeposition at \( 8 \text{ mA/cm}^2 \).

Fig. 4 summarizes the growth rate deduced from experiments as a function of average membrane pore size and elastic modulus, respectively, at \( 8 \text{ mA/cm}^2 \) as well as for varying current densities such that \( \frac{J}{J^*} \) remains constant at 0.9 (SI Appendix, Fig. S5). Optical snapshots of the initial deposition and the corresponding current densities (at \( \frac{J}{J^*} = 0.9 \)) are reported in SI Appendix, Fig. S9. The experimental results reveal a sudden transition from stable deposition to dendrite growth between 100–500-nm pore size in agreement with our hypothesis that electrodeposition stability is function of electrolyte length scale. Also, the shear modulus measured from rheology does not have a linear relation with growth rate, which can be rationalized by the fact that the cross-linked membrane comprises building volume fractions of silica nanoparticles corresponding to interparticle distance of order 500 nm, SI Appendix, Fig. S10 shows two regimes of dendrite growth, wherein the growth rate at the initial time is low, followed by an accelerated rate of dendrite propagation at longer time. The results in SI Appendix, Fig. S9 also show that for the membranes with 500-nm pores, dendrites with “needle-like” morphology are visible at 4 \( \text{mAh/cm}^2 \), in contrast to the mossy-blackened deposited lithium apparent when the pores are larger and in neat electrolytes. Furthermore, it is seen that in membranes with effective pore size of around 100 nm (Movie S4), the deposition is similarly restricted as for the 20-nm case, implying that there is a threshold pore size between 500 and 100 nm where the resistance to dendrite proliferation is large enough to prevent growth.
blocks having different inherent modulus (SiO₂ and polymer). As the deposition takes place in the interparticle gap, the growing “dendrites” only experience the modulus of the liquid electrolyte and polymers; thus, the length scale of the polymeric phase plays a greater role than the overall modulus of the material. We also analyze the growth rate of dendrites as a function of pore size obtained from TEM micrograph analysis as well as the pore volume calculated from plateau modulus as $kT/G'$. As shown in *SI Appendix*, Fig. S12, a similar nonlinear behavior is observed.

Fig. 5A plots the perturbation growth rate versus wavelength deduced from the analysis of Tikekar et al. (38). It is seen that the growth rate is negative for small wavelengths and goes through a maximum at a particular wavelength that decreases as the current density approaches critical current density, $J^*$. The first of these results can be shown to arise from the greater influence of surface tension in suppression growth of small dendrite nucleates, while the presence of the maximum reflects the effect of the physical (macroscopic) dimensions of the simulation cell in restricting nucleate size below certain limits. As the current density increases, the growth rate of the fastest (most unstable) mode increases and its wavelength decreases because the interfacial tension has to compete with progressively more aggressive electroconvection to stop the roughening. The fastest growing mode can therefore be thought of as representative of the typical size of dendrites that one would observe in an experiment under similar conditions. A porous material would restrict the size of the growing dendrite, since the deposition can only happen on length scales smaller than the pore size. In Fig. 5B, we showed the calculated growth rate for the fastest growing mode as a function of the critical wavelength. For very large pores, the pore size is predicted to have no effect on the dendrite growth, and the growth occurs more or less at the same rate as in the absence of the porous material. For smaller pores, however, the pore diameter restricts the sizes of the dendrites that can form in them, causing the growth rate of the fastest mode to plummet. Below a certain size, there can be no unstable deposition, as surface tension adequately competes with current-density-driven dendritic growth at all possible length scales. This critical pore size decreases with current density due to the requirement of a higher contribution from surface tension as seen in *SI Appendix*, Fig. S13. *SI Appendix*, Fig. S13 (Inset) plots the analogous growth rate deduced from experiments as a function of membrane pore size for a fixed current density, $Jf^* \sim 0.9$. The experimental results show a similar trend as calculated from the linear stability analysis, wherein there is sudden transition from stable deposition to dendritic growth. The transition point from the theoretical prediction corresponding to nonzero growth is \( \sim 1.6 \) μm, while the experiments indicate the cross-over between 100–500-nm pore size as previously mentioned in Fig. 4A. Considering the variety of sources of error in comparing the experiments and theory, including the tortuosity of the pores in the cross-linked materials, the polydispersity of the particle size, uncertainty in the value of the Li transference number under the conditions of the experiments (48), and the deviation from linearity in calculation of initial growth rate, the agreement between the theoretical and observed transition is fair.

In conclusion, we utilized a cross-linked network by covalently linking polymer-grafted nanoparticles, where varying the volume fraction of the nanoparticles in the composite can conveniently change the effective pore size. It was shown that these cross-linked nanoparticles when used as gel electrolytes showed stable electrodeposition in contrast to mossy and dendritic morphology for cases without separator and with glass fibers, respectively. We performed visualization experiments for different pore sizes determined by volume fractions to demonstrate that below 500 nm, the deposition is smooth and compact, while it is dendritic for other cases. These experimental observations were rationalized using linear stability analysis of dendrite growth at different perturbation length scale representing pore diameter of the electrolytes. It was understood that below the limiting current density, the electrodeposition can be stabilized by implementing a separator with pore size lower than the length scale of the most unstable nucleate. Overall, this work will provide guidelines for designing solid-state electrolytes and separators for metal-based batteries.

**Materials and Methods**

**Materials.** Lithium rods, ethylene carbonate, dimethylene carbonate, lithium hexafluorophosphate, anhydrous copper chloride, dimethyl sulfoxide, glass fibers Ludox SM30 colloidal silica (d = 10 ± 2 nm), poly(propylene glycol)-toluene-2,4-diisocyanate, were all purchased from Sigma-Aldrich. Hydroxy terminated poly (ethylene oxide)-silane was obtained from Gelest. Copper foil was obtained from Alfa Aesar. All of the chemicals were used as received after rigorous drying in an \(~0\)-ppm water level and \(<5\)-ppm oxygen glove box.

**Linear Stability Analysis.** The details regarding the linear stability analysis were presented in a previous paper by Tikekar et al. (38)

**Cross-Linked Hairy Nanoparticles Synthesis.** The reaction procedure for cross-linking between the grafted silica nanoparticles and functionalized PPO polymer was given in a previous work (18). Different cross-linked nanoparticles with varying pore sizes were synthesized by adjusting the ratio between the silica nanoparticles and the PPO polymer as given in *SI Appendix*, Table S1. The ratio between the liquid electrolyte and the cross-linked composite was kept constant at 2:1 by weight.

**Dielectric Spectroscopy.** The dielectric spectroscopy measurements were done using a Novocontrol N40 Broadband Dielectric instrument. The cross-linked hairy nanoparticles as well as neat PPO (without liquid electrolyte) were casted at the center of a teflon o-ring. The measurements were done in a frequency range from \(10^{-3}\) to \(10^7\) Hz at various temperatures.

**TEM.** The TEM measurements were performed to understand the nanostructure of the cross-linked hairy nanoparticles using an FEI T12 Spirit TEM. A thin layer of the materials was casted onto the TEM grid using chloroform as the solvent and it was allowed to cross-link in the TEM grid. Before the measurements, the samples were dried overnight at 60 °C.

**SEM.** SEM analysis was done using the LE0155FESEM instrument. In this experiment, symmetric lithium coin cell was constructed using the cross-linked electrolyte (r.c.p. \(= 20\) nm). “Plate-strip” measurements were performed at a current density of 0.1 mAcm⁻² for 100 h, with each cycle comprising 3 h; thereafter, the lithium electrode was extracted and dried in the anechamber of a glove box overnight.

**Mechanical Properties.** The cross-linking process was studied mechanically by casting the precursor materials in the rheometer with parallel-plate geometry of diameter 25 mm using an ARES Rheometer. Specifically, for time-sweep measurements, a constant strain amplitude of 5% and fixed frequency of 10 Hz was applied until the storage modulus reached a fixed value. The frequency-sweep measurements were performed at a fixed strain amplitude of 5% between 1 and 100 Hz for all materials. All of the measurements were performed...
done without any addition of liquid electrolytes and at the same temperature value of 60 °C.

Direct Visualization Experiments. The visualization experiment was carried out for understanding the electrodeposition process with various forms of electrolytes mentioned in the text. In all of the measurements the ratio between the weight of the composite material and liquid electrolyte (1 M LiPF6 EC/DMC) was kept constant at 1:2. The details of the visualization cell for lithium deposition measurements were presented in a previous paper (10). The dendrite growth was done using a semi-automated analysis in MATLAB, where the sizes of the focused dendrites are measured over time.

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