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DOI
10.1016/j.esci.2022.09.001

Publication date
2022

Document Version
Final published version

Published in
eScience

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Research Paper

Highly aligned lithiophilic electrospun nanofiber membrane for the multiscale suppression of Li dendrite growth

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HIGHLIGHTS

- A highly aligned lithiophilic TiO2/SiO2 (A-TS) electrospun nanofiber membrane is developed.
- A custom-made electrospinning system is introduced to control the orientation of the fibers in the membrane.
- Suppressing dendrite growth at both membrane and fiber scales to achieve excellent cycling performance in various batteries.

GRAPHICAL ABSTRACT

ARTICLE INFO

Keywords:
- Aligned structure
- Protective layer
- Li dendrite
- Lithium sulfur battery
- Li metal battery

ABSTRACT

Using inorganic fibrous membranes as protective layers has yielded success in suppressing dendrite growth. However, conventional fibrous membranes usually have large voids and low affinity for Li, promoting inhomogeneous charge distribution and allowing some dendrites to grow. Herein, we introduce a highly aligned TiO2/SiO2 (A-TS) electrospun nanofiber membrane as a protective layer for the Li metal anode. The A-TS membrane is fabricated by a custom-made electrospinning system with an automatic fiber alignment collector that allows control of the fibers’ orientation. At the scale of the individual fibers, their high binding energies with Li can attract more “dead” Li by reacting with the SiO2 component of the composite, avoiding uncontrollable deposition on the metal anode. At the membrane scale, these highly ordered structures achieve homogeneous contact and charge distribution on the Li metal surface, leaving no vulnerable areas to nucleate dendrite formation. Additionally, the excellent mechanical and thermal stability properties of the A-TS membrane prevent any potential puncturing by dendrites or thermal runaway in a battery. Hence, an A-TS@Li anode exhibits stable cycling performance when used in both Li–S and Li–NCM811 batteries, highlighting significant reference values for the future design and development of high-energy-density metal-based battery systems.

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https://doi.org/10.1016/j.esci.2022.09.001
Received 18 May 2022; Received in revised form 23 August 2022; Accepted 2 September 2022
Available online 11 September 2022
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1. Introduction

In recent years, there has been an increasing demand for safe high-energy Li batteries for electric grids, portable electronics, and electric vehicles [1,2,3,4]. Li metal is regarded as the most attractive anode material in Li-based batteries due to its ultrahigh theoretical specific capacity of 3680 mAh g⁻¹ (compared with 372 mAh g⁻¹ for a graphite anode) and low redox potential (−3.04 V versus a standard hydrogen electrode) [5,6,7,8]. In addition, Li metal anodes can be matched with various cathodes (e.g., sulfur, oxygen, LiCoO₂, LiFePO₄, LiNiₓMnₓCo₁₋ₓO₂) in some promising high-energy batteries, such as lithium–sulfur and lithium–air.

Despite the advantages of the Li metal anode, there are still serious challenges that need to be overcome before it can achieve its full potential. The uncontrollable growth of Li dendrites resulting from inhomogeneous charge distribution on the anode surface is at the top of the challenge list. Li dendrites can break the solid electrolyte interphase (SEI) layer and accelerate the consumption of organic electrolytes, decreasing the Coulombic efficiency and cycle life [9,10,11]. In addition, dendrites can penetrate the separator and consequently short-circuit the cell, potentially creating safety hazards [12,13,14].

Several concepts have been suggested to tackle the dendrite growth problem, including deploying a three-dimensional (3D) conductive skeleton to accommodate Li protrusion, designing a functional separator facing the Li side, preforming a protective SEI layer on the Li metal, and others [15,16,17,18]. Among these, constructing an artificial interlayer between the separator and the Li anode is reported to stabilize the SEI layer and regulate Li⁺ deposition [19,20,21,22]. In this regard, a 3D porous structure based on continuous nanofibers or electrospun nanofiber membranes demonstrated great success when used as an artificial protective layer [23,24,25,26,27]. The abundant volume available in the porous structure, combined with its high mechanical flexibility, allows it to accommodate the significant volumetric change that the Li metal anode undergoes during repeated deposition and stripping [28,29]. Further, the large specific surface area facilitates the processes at the electrolyte/Li interface, extending the Sand’s time and ensuring a stable and homogeneous Li⁺ flux. It is also possible to improve the separator and anode wettability and other physicochemical properties by inducing some functionality on the nanofiber membranes, further suppressing the growth of Li dendrites. In addition, the electrospinning technique is cheap and industrially mature, enabling large-scale fabrication and commercialization [29].

Despite the enhancement of anode stability when a nanofibrous protective layer is used, there are still issues that need to be addressed before practical applications can be considered. The random spatial orientation of conventional electrospun nanofibers can lead to large voids within the membrane that allow the growth of Li dendrites [30]. Most previous studies in the literature have used a carbon-based nanofibrous protective layer as the membrane skeleton, owing to the high electrical conductivity and promising Li storage ability of carbon materials (e.g., hard carbon and graphite) [26,31,32,33,34]. However, the Li affinity of some carbon materials is lower than that of elementary metal substances (e.g., Ag, Si, and Mg) and metal oxides (e.g., Al₂O₃, TiO₂, ZnO, and SiO₂), and they have a low Li nucleation energy barrier [33,35,36], originating from a lattice mismatch between the body-centered cubic (bcc) lithium structure and the hexagonal structure [37]. Although some “lithiophilic” metal sulfides (MoS₂, ZnS, and VS₂) were recently used as an artificial interlayer to improve Li nucleation via electrochemical conversion between the metal sulfides and Li [38,39,40], their large-scale production is hindered by synthetic complexity, environmental instability, and high cost. Therefore, the design of a protective layer with high chemical stability, uniform pore size distribution, and lithiophilic materials may bring a breath of fresh air to the design of high-performance Li anodes.

Herein, we propose a highly aligned TiO₂/SiO₂ (A-TS) electrospun nanofiber membrane to serve as a protective layer for the Li metal anode. We also introduce a custom-made electrospinning system with an automatic fiber alignment collection device to prepare the inorganic membranes. At the membrane scale, the well-arranged nanofibrous array structure of the A-TS membrane guarantees homogeneous charge distribution and fast Li⁺ diffusion. The membrane also provides room to accommodate any dendrites formed and, together with the mechanically strong fibers, significantly minimizes the chance of short-circuiting. At the fiber scale, the “lithiophilic” inorganic ceramics (TiO₂ and SiO₂) chemically bind/react with “dead” Li to further suppress dendrite growth and achieve the efficient deposition and stripping of Li. As a result, the designed A-TS membrane supports the Li metal anode to achieve outstanding cycling and rate capability performance that outshines most Li metal anodes in the literature, including those using nanofiber membranes.

2. Results and discussion

Fig. 1 shows schematic illustrations of Li metal batteries with bare Li foil, C-TS@Li, and A-TS@Li anodes. Compared with bare Li, where dendrite growth is fast and uncontrollable (Fig. 1a) [41,42], the lithiophilic TS electrospun nanofiber membrane inhibits uncontrolled dendrite growth and provides physical space to accommodate any formed dendrites. However, due to the inherent bending instability effect of the jet in conventional electrospinning systems, the C-TS membrane is inevitably composed of randomly arranged nanofibers (Fig. 1b) [30,43]. As a result, the as-spun C-TS membrane retains plenty of large voids, and the charge diffusion is slightly inhomogeneous.

Highly aligned A-TS electrospun nanofiber membranes can regulate the charge distribution and further improve the ability to inhibit Li dendrites. The A-TS membrane is prepared using a custom-designed electrospinning system consisting of two parallel electrodes and one additional assistant electrode (Fig. 1b). An electric reciprocating mechanical arm driven by digital pulse power is also added into this electrospinning configuration, aiming to realize the fully automated and large-scale production of highly aligned fibers. More details and photos of the working mechanism of the custom-designed electrospinning configuration are shown in Fig. S1. The densely and highly ordered fiber-stacked structure of the A-TS membrane, combined with the excellent physical/chemical properties of the TiO₂/SiO₂ fibers, promotes more effective suppression of Li dendrites and enhances the cycling performance of Li metal batteries.

We first investigated the optimum conditions to obtain a robust, free-standing electrospun nanofiber membrane of C-TS and A-TS (as shown in Figs. S2–4). Pure TiO₂ electrospun nanofibers are too fragile to form free-standing membranes. TiO₂/SiO₂ electrospun composite membranes, on the other hand, can achieve excellent mechanical strength/flexibility with optimal material properties when calcined at 500 °C. Thermogravimetric analysis (TGA) demonstrated that the organic compounds from the precursors were entirely removed at 500 °C to form inorganic C-TS and A-TS nanofibers (Fig. S5). Hence, the C-TS and A-TS membranes were fabricated and then calcined at 500 °C to serve as a protective layer for the Li metal anode. It should be mentioned here that the average diameters of the C-TS and A-TS nanofibers were almost the same, 303.2 nm and 306.6 nm, respectively, despite changes in the electrospinning configuration (Figs. 2a–c and Figs. S2 and S6).

The formation of homogeneous TS nanofibers in both C-TS and A-TS was proven by EDS elemental mapping and XPS analysis, as can be seen in Figs. 2g–i and Fig. S7. According to the TEM and XRD analyses shown in Figs. 2d–h and Figs. S6e–i, the C-TS and A-TS nanofibers possessed lower crystallinity than the C-TS nanofibers calcined under the same conditions. Unlike the clear crystalline structure of the C-T nanofibers, as presented in their SAED and XRD patterns, only one weak diffraction ring and a shallow peak corresponding to the (101) plane of anatase TiO₂ (JCPDS No. 21–1272) were observed in the traces of C-TS and A-TS.
These results suggest the amorphous SiO2 phase introduced in the TS nanofibers retarded the long-order growth of the original TiO2 crystals [44]. The low crystallinity of TiO2 and SiO2 improved the affinity of the fibers toward Li and thus guided uniform Li nucleation [35]. The low crystallinity of TiO2 and SiO2 improved the affinity of the fibers toward Li and thus guided uniform Li nucleation [35]. The addition of SiO2 to the TiO2 seems to have given the protective layer another advantage with respect to enhancing the Li anode performance. As can be seen from Fig. 2j and Fig. S8, the N2 adsorption–desorption isotherms suggest an over four-fold increase in the BET specific surface area in the composite fibers compared to C-T, without influencing the intrinsic pore size distribution; this is mainly attributed to the introduced amorphous SiO2 phase, which increased the surface area of the original TiO2 nanofibers [45,46]. The higher surface area of the protective layer guaranteed more active sites and room to react with and deposit “dead” Li, thereby alleviating the issue of dendrite penetration [47,48].

The protective layer’s mechanical and thermal stability properties are important for increasing battery life and ensuring safe operation. As shown in Fig. 3, the composite TS nanofibers and the highly aligned fiber-stacked structure demonstrated high mechanical strength when attached to the separator. A PP separator with an A-TS membrane exhibited the highest puncture resistance compared with both a pure PP separator and a separator reinforced with a C-TS membrane, implying the protected separator’s stronger ability to physically resist being punctured by the Li dendrites (Figs. 3a and b and Fig. S9). The high mechanical stability of TiO2 and the fiber-stacked structure of the A-TS membrane also minimized potential volume change in the SiO2 and Si after lithiation [49,50]. The thermal stability and flame retardancy of the inorganic TiO2/SiO2 membrane was much higher than that of the commercial PP separator. While the PP separator instantly caught fire upon exposure to flame, the TS membranes showed significant resistance and maintained their original shape after over a minute of exposure to direct flame (Fig. 3c and d). The nonflammability and higher thermal resistance of this protective layer thus will increase battery safety by impeding heat diffusion if the PP separator shrinks and thermal runaway happens.

The binding energies of anatase TiO2, amorphous SiO2, and graphene towards Li atoms were further calculated using density functional theory (DFT) to evaluate their lithiophilic features. As shown in Figs. 3e, the binding energies of a Li atom on the TiO2 (101) surface and on amorphous SiO2 can reach up to –2.07 and –1.86 eV, respectively, much stronger than on graphene (–0.91 eV), proving the high chemical affinity of both anatase TiO2 and amorphous SiO2 toward Li. Low Li affinity will lead to inhomogeneous Li⁺ flux and easily give rise to uneven Li nucleation, resulting in a loose Li interphase and lower cyclability for the Li anode, especially under high current densities [37,51]. In contrast, as the nucleation and deposition of Li preferentially happen on the surface of polar metals or metal oxides, TiO2 and SiO2 have stronger binding energies with Li and show better lithiophilicity [52,53].

According to previous reports, the lithiation mechanism of TiO2 and SiO2 can be explained by the following steps [54,55]:

\[
\begin{align*}
\text{TiO}_2 + \text{Li}^+ + e^- &\rightarrow \text{LiTiO}_2 \\
5\text{SiO}_2 + 4\text{Li}^+ + 4e^- &\rightarrow 2\text{Li}_2\text{Si}_2\text{O}_3 + \text{Si} \\
2\text{SiO}_2 + 4\text{Li}^+ + 4e^- &\rightarrow \text{Li}_4\text{SiO}_4 + \text{Si} \\
4\text{Si} + 15\text{Li}^+ + 15e^- &\rightarrow \text{Li}_{15}\text{Si}_4
\end{align*}
\]

Based on steps (2), (3), and (4), the presence of amorphous SiO2 enables the activation of more “dead” Li, based on its specific alloying reaction mechanism and low energy barriers; in addition, it enhances the mechanical flexibility of the free-standing membrane. The formation of the intermediate Li silicate phases and the Li–Si alloy were proven by the subsequent XPS analysis presented in Fig. 6g [54,56]. Meanwhile, according to equation (1), the anatase TiO2 contained in

![Fig. 1.](image)

(a) Schematic illustrations of Li metal batteries with bare Li, C-TS@Li, and A-TS@Li anodes; (b) conventional electrospinning system for the fabrication of C-TS electrospun nanofiber membrane, and the self-designed modified electrospinning system used to fabricate the A-TS electrospun nanofiber membrane.
TS fibers can not only stabilize the volume change experienced by SiO2 and Si after lithiation but also provide intercalation space to accommodate and react with Li [53,55]. After the initial lithiation process, all the Li-containing compounds served as efficient Li ionic conductors to ensure a higher Li\(^+\) transfer rate. In addition, owing to its aligned porous structure and high electrolyte affinity, the A-TS membrane also exhibited much smaller contact angles with different electrolytes, suggesting faster Li\(^+\) diffusion would be achieved in various electrolyte systems (Fig. S10).

The interaction between deposited Li and TS nanofiber is illustrated in Fig. 3f. During battery cycling, the high chemical affinity between the TS nanofiber and Li drives a strong binding force to anchor nonactivated Li, slowing down the vertical growth of Li dendrites. More adsorbed Li\(^+\) reacts with the amorphous SiO2 to form lithiophilic Li-containing compounds based on the alloying reaction mechanism, and the Li-containing compounds further provide more sites to adsorb Li\(^+\) and facilitate Li\(^+\) diffusion [35,54]. Therefore, the deposition and plating behaviors of Li are stabilized under the protection of the TS membrane. This mechanism was proven experimentally using Li–TiO2 and Li–TiO2/SiO2 batteries as assembled with a TiO2 or TiO2/SiO2 cathode and a Li metal anode (details on the cell assembly are in the Experimental section in the Supporting Information). A sufficient Li source enabled Li ions to fully embed into the TiO2 or TS electrospun nanofibers in every cycle in the tested battery system. After 400 embedding/de-embedding cycles (Figs. 3g and h), the SEM images show that the pure TiO2 nanofibers still had a relatively smooth surface with no obvious anchoring effect; in contrast, the TS composite fibers were almost covered with anchored Li-containing nanoparticles. In addition, the TS nanofibers presented a nearly 1.5 times higher capacity during the Li\(^-\) embedding process than the pure TiO2 nanofibers at all cycles (Fig. 3i). These results confirmed the role of the amorphous SiO2 phase in the continuous anchoring and conversion of “dead” Li.

The membrane-scale advantages of the A-TS membrane for the suppression of Li dendrite growth are further investigated in Fig. 4. A bare Li anode without a protective layer had difficulty avoiding Li dendrite growth and the volumetric change Li metal undergoes during repeated deposition/stripping processes [16,57,58]. The unevenly distributed charges also induced the Li\(^+\) to aggregate near the dendrites, thereby accelerating the uncontrollable growth of dendrites, resulting in severe electrolyte consumption and the fragmentation of the SEI layer [9,16]. Hence, the surface of the bare Li anode after cycling was quite rough, with plenty of micron-sized dendrites growing out of control over the whole anode.

![Fig. 2. Morphological and structural characterizations of various samples. Field emission scanning electron microscopy (FE-SEM) images of (a) C-TS and (b) A-TS membranes; (c) the diameter distribution of A-TS nanofibers measured from the membrane in Fig. 2b; (d–f) transmission electron microscopy (TEM) images of C-T, C-TS, and A-TS nanofibers; the inset is the selective area electron diffraction (SAED) pattern; (g) energy-dispersive spectrometer (EDS) mapping of the A-TS nanofiber; (h) X-ray diffraction (XRD) patterns of C-T, C-TS, and A-TS nanofibers; (i) X-ray photoelectron spectroscopy (XPS) fully scanned spectra of C-TS and A-TS nanofibers; (j) comparisons of Brunauer–Emmett–Teller (BET) specific surface area and BJH average pore size for C-T, C-TS, and A-TS nanofibers.](image-url)
surface (Figs. 4a and d). With the C-TS@Li anode, Li dendrite growth was alleviated to a certain extent. However, some small dendrites were still observed, mainly due to the uneven charge distribution and some large voids in the C-TS membrane (Figs. 4b and e). In contrast, the highly aligned fiber-stacked structure of the A-TS membrane not only enabled more overall physical protection against dendrite penetration and provided more exposed active sites for the anchoring and activation of “dead” Li [59], but also ensured an even electric field distribution for homogeneous SEI formation and Li\(^+\) diffusion; consequently, it exhibited a smooth surface with almost no dendrites (Figs. 4c and f).

To gain more insights into the dendrite suppression mechanism via different protective layers, COMSOL Multiphysics software was used to simulate the electric field distribution when Li dendrites appear on the surface of the Li metal anode (Figs. 4g–i). Without protection, Li ions are prone to gather around the formed dendrites due to the tip effect (Figs. 4g and j) [41,60]. In subsequent cycles, the charges focused around the dendrites will attract more Li ions, accelerating vertical dendrite growth until the battery short-circuits. Although the unevenly distributed electric field around Li dendrites can be dispersed by lithiophilic TS nanofibers, the tip effect still exists in the voids of crossed nanofibers (Figs. 4h and k). Owing to both the high lithiophilicity and the aligned fiber-stacked structure of the A-TS nanofibers, Li\(^+\) tends to adsorb uniformly on the A-TS membrane and, in turn, induces homogeneous charge distribution on the surface of the Li anode (Figs. 4i and l), thus reducing the local current density of the Li anode and extending the Sand’s time [61]. Moreover, homogeneous charge distribution can effectively prevent the tip effect and localized vertical growth of Li dendrites, contributing positively to smooth Li deposition and stripping during cycling.

The galvanostatic cycling performance of symmetric cells assembled with bare Li, C-TS@Li, and A-TS@Li anodes was assessed to further compare the stability of various anodes in repeated deposition/stripping processes (Fig. 5). At a current density of 0.5 mA cm\(^2\) with a stripping/plating capacity of 0.5 mAh cm\(^2\), the bare Li exhibited an initial overpotential of 90.9 mV, much higher than the C-TS@Li (59.5 mV) and A-TS@Li (48.9 mV) electrodes (Fig. 5a). The polarization of the bare Li became unstable after 50 h, and the overpotential further increased to ~220 mV after 300 h. This polarization increase can be attributed to the gradual accumulation of “dead” Li on the surface of the Li metal, leading to
an aggravated Li/electrolyte interface and fragmentation of the SEI layer [62,63]. Despite the similar initial overpotential recorded for the C-TS@Li and A-TS@Li anodes, the highly aligned A-TS@Li facilitated smoother Li stripping/plating than C-TS@Li, judging by the lower polarization after 200 h. The A-TS@Li also exhibited the most stable polarization among the three tested anodes at a higher current density and when switching between low and high current densities (Fig. 5b), indicating excellent rate capability. Furthermore, the A-TS membrane maintained a stable voltage profile during the long-term stripping/plating cycles, even at a higher current density of 5 mA cm$^{-2}$ ($\sim$73 mV after 300 h, in Figs. 5c and d). The same stability was observed for the A-TS@Li anode at a stripping/plating capacity as high as 5 mAh cm$^{-2}$ ($\sim$35.4 mV after 300 h, in Figs. 5e and f), more stable than that of the bare Li, shown in Fig. S14).

To further demonstrate the significant role of the A-TS membrane in homogenizing Li deposition, it was used in a Li||A-TS@Cu cell. This cell delivered a high initial Coulombic efficiency of $\sim$97% and remained at $\sim$96% after 70 cycles at a current density of 1 mA cm$^{-2}$ with a capacity of 1 mAh cm$^{-2}$ (Fig. S15). All these results reflect that under the multiscale protection of the A-TS membrane, uniform deposition/stripping behaviors were well achieved for the Li metal anode.

Electrochemical impedance spectroscopic (EIS) and cyclic voltammetric (CV) analysis of Li–S full batteries provided more insights into the charge transfer mechanism of various anodes. Fig. 6a displays the EIS spectra of the bare Li, C-TS@Li, and A-TS@Li anodes at open-circuit voltage. The Nyquist plots of all three anodes constitute a high-frequency semicircle and a low-frequency sloped line, corresponding to the charge-transfer resistance ($R_{ct}$) and mass-diffusion process [64]. Based on the fitted equivalent electrical circuit model and the corresponding resistance data shown in the inset of Fig. 6a and in Table S1, the $R_{ct}$ value of A-TS@Li (80.6 $\Omega$) is lower than that of C-TS@Li (96.5 $\Omega$) and bare Li (114.9 $\Omega$). The lower charge-transfer resistance at A-TS@Li emphasizes the role of the lithiophilic membrane in easing charge...
movement across the Li/electrolyte interface and speeding up the Faradaic reaction kinetics. The CVs recorded for the different anodes (bare Li, C-TS@Li, and A-TS@Li) at various scan rates ranging from 0.1 to 0.9 mV s\(^{-1}\) are illustrated in Fig. S16.

The Li diffusion coefficient (\(D_{Li}^{+}\)) in the three cases is calculated using the Randles–Sevcik equation to describe the Li diffusion process [65,66]. As can be seen from the CVs, all the cathodic current peaks (\(I_{\text{Peak } A}\) and \(I_{\text{Peak } B}\)) and the anodic current peak (\(I_{\text{Peak } C}\)) show linear relationships with the square root of the scanning rate (\(\sqrt{\nu}\)) of the three anodes, indicative of their diffusion-limited process [66]. Based on the calculated \(D_{Li}^{+}\) results (Fig. 6b and Table S2), Li\(^+\) diffusion was faster on the A-TS@Li anode than on the bare Li and C-TS@Li anodes during both
reduction and oxidation. It should be mentioned here that for Li–S batteries, rapid Li\(^+\) diffusion facilitates the sulfur transformation chemistry at the cathode side and enables efficient utilization of active sulfur for enhanced capacity performance \([66, 67, 68]\).

The long-term cycling performance of various anodes in Li–S batteries is shown in Fig. 6c. The A-TS@Li anode delivered a high initial discharge capacity of 1218 mAh g\(^{-1}\) at 0.2C (1C = 1675 mAh g\(^{-1}\) in a Li–S battery), which remained stable at 869.5 mAh g\(^{-1}\) even after 400 cycles with a Coulombic efficiency of ~99% and a slow capacity attenuation (0.071% per cycle). Compared with a conventional Li–S battery, the weight increase in the whole cell was only 2.9% (Fig. S17). The galvanostatic charge/discharge profiles of the A-TS@Li anode also show stable discharge/charge characteristic voltage plateaus belonging to the Li–S battery from the 1st to the 400th cycle. In comparison, the capacity of the bare Li anode struggled with rapid capacity attenuation, reaching 689 mAh g\(^{-1}\) after 200 cycles. Although the C-TS@Li anode exhibited a high initial capacity of 1196.6 mAh g\(^{-1}\), similar to the A-TS@Li anode, it could not maintain a stable cycling performance after 200 cycles. Furthermore, the A-TS@Li anode presented a lower polarization (205 mV) compared with the C-TS@Li (240 mV) and bare Li (286 mV) anodes (Fig. 6d), suggesting a lower energy barrier for the charge transfer process \([64]\), in agreement with the EIS and diffusion coefficient results. The A-TS@Li anode also showed a stable rate performance, with the capacity decreasing to 756 mAh g\(^{-1}\) at 2C and recovering to 1066 mAh g\(^{-1}\) when the current density was reduced back to 0.1C (Fig. 6e and Fig. S19). This excellent performance by the A-TS@Li anode could be attributed to the lithiumophilicity and nano-fibrous array structure of the A-TS membrane, which helped achieve homogeneous charge transfer/diffusion and multiscale suppression of Li dendrite growth.

Motivated by the excellent performance of the A-TS@Li anode in Li–S batteries, we then tested the A-TS membrane in a Li metal battery system using an NCM811 (LiNi\(_{0.8}Co_{0.1}Mn_{0.1}O_2\)) cathode (Fig. 6f). The A-TS@Li
We further investigated the role of the A-TS membrane in suppressing dendrite growth on Li metal anodes via a series of post-mortem analyses after cycling in Li-S batteries. In addition to the surface morphological information from the cycled Li metal anodes, presented in Figs. 4d–f, the associated TEM images of the TS membrane are shown in Fig. S22, and the weight parameters of the Li anode and A-TS membrane before and after cycling are summarized in Table S5. After 200 cycles in Li-S batteries, the A-TS membrane enabled more effective anchoring of the “dead” Li than the C-TS membrane did. The high affinity of TS nanofibers toward Li and the more homogeneous Li⁺ diffusion during the deposition/plating processes effectively controlled the “dead” Li deposition on the Li metals and prevented the vertical growth of Li dendrites caused by the tip effect. The fiber-stacked structure of the A-TS membrane provided more active sites to accommodate the “dead” Li (Fig. S23). The high-resolution XPS spectra of Si 2p and Li 1s for the cycled A-TS membrane are shown in Fig. 6g. Compared with the original TiO₂/SiO₂ composite, the TS nanofibers after cycling showed the peaks belonging to Li-containing compounds. In the Si 2p orbit, in addition to the peaks at 103.0 and 102.1 eV from the inherent SiO₂ compound in TS nanofibers, a Li₂SiO₃ peak at 101.1 eV and a Li–Si peak at 98.1 eV were observed. Detection of Li–Si bonds confirmed that the electrochemical conversion and alloying reactions of SiO₂ played significant roles in binding with “dead” Li during cycling, as presented previously in Fig. 3 [69]. For the Li 1s orbit, the peak representing the Li–Si bond at 54.5 eV was again observed [69]. These results confirmed the effective electrochemical conversion of SiO₂ to Li₂SiO₃ and of Si to Li–Si alloy. In addition, the Li–F bond at 55.7 eV may have derived from the formed SEI layer or the interaction between the anchored “dead” Li and the electrolyte [70,71].

Based on the above analysis, the functional mechanism of the A-TS membrane in the multiscale suppression of Li dendrite growth is summarized in Fig. 6h. At the fiber scale, introducing SiO₂ greatly enhances the mechanical properties of the TiO₂ electrospun nanofibers, allowing the formation of free-standing TS membranes as protective layers for the Li metal anode. The high chemical affinity of the TS nanofibers for Li attracts Li to attach to the nanofibers in the form of “dead” Li and then continuously reacts with the TS nanofibers. Consuming the Li on the TS nanofibers removes the pressure on the Li anode surface and prevents the vertical growth of Li dendrites. The TS composite nanofibers, with their high mechanical strength and thermal stability, physically resist being punctured by any formed dendrites and prevent the battery from short-circuiting if thermal runaway occurs. At the membrane scale, the special array structure composed of the highly aligned electrospun nanofibers achieves even charge distribution as well as homogeneous and fast Li⁺ diffusion, removing the conditions that might favor dendrite seeding. The A-TS membrane also enables perfect physical protection to prevent rapid dendrite penetration through the membrane’s large voids, as well as ensuring more exposed active sites for anchoring “dead” Li. Hence, stable deposition and stripping of the Li metal anode can be achieved under the protection of the A-TS membrane. However, to better optimize the electrochemical performance of A-TS membranes for future commercialization of Li-S batteries, the interaction between polysulfides and the A-TS membrane needs to be further investigated to reveal the mechanism by which the “shuttle effect” is suppressed. The introduction of flexible additives with high ionic conductivity may also be an effective method to broaden the application scope of the A-TS membrane.


