Synthesis of interconnected graphene framework with two-dimensional protective layers for stable lithium metal anodes

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1. Introduction

Lithium ion batteries (LIBs) have been successfully used in portable electronics and electric vehicles for many years [1–3]. Although the demand for high-energy-density keeps increasing, the existing commercial LIB systems are getting close to their theoretical capacity limits. In order to push the energy density to an even higher level, Li commercial LIB systems are getting close to their theoretical capacity demand for high-energy-density keeps increasing, the existing commercial LIB systems are getting close to their theoretical capacity limits. In order to push the energy density to an even higher level, Li commercial LIB systems are getting close to their theoretical capacity limits. In order to push the energy density to an even higher level, Li commercial LIB systems are getting close to their theoretical capacity limits. In order to push the energy density to an even higher level, Li commercial LIB systems are getting close to their theoretical capacity limits. 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can no longer be utilized, leading to low Coulombic efficiency.

The current collector is regarded as a key component in Li metal anodes, and can make significant impact on the deposited Li metal morphology as well as the electrochemical performance. Therefore, a lot of effort has been made to modify current collectors for stable Li metal deposition. Because one main factor that leads to Li dendrite formation is inhomogeneous Li-ion flux distribution, it is straightforward to design a three-dimensional (3D) current collector that can increase the surface area and reduce the effective current density on the electrode. Following this strategy, 3D porous metal foil [23–26], porous graphene framework [27–29] and carbon nanofiber/nanotube network [30–36] are demonstrated to suppress dendrite growth and improve Coulombic efficiency effectively. However, this strategy can only reduce the growth rate of Li dendrites to some extent. After long cycles, mossy surface morphology can still be formed. In addition, another approach is to use a chemically stable and mechanically strong thin layer, such as interconnected hollow carbon nanospheres [37] and two-dimensional (2D) materials (including graphene [38], hexagonal boron nitride [38] and molybdenum disulfide [39]), as interfacial layer. During cycling, the interfacial layer can constrain Li deposition underneath it, leading to flat surface morphology. Nevertheless, one single thin protection layer may not be sufficient when the Li metal loading is high. Additionally, if the ion conductivity of the protection layer is low, it’s hard to achieve good cycling performance at high current densities.

Inspired by the previous reports, we rationally designed an interconnected graphene (IG) framework consisting of both 3D graphene hollow spheres and 2D graphene flakes. The 3D graphene hollow spheres were used to increase the surface area and provide enough ion transport channels. The 2D graphene flakes worked as interfacial layers to flatten the surface morphology and suppress dendrite growth. Using the IG current collector, we demonstrated highly reversible Li plating/stripping with an average Coulombic efficiency of 97.5% over 100 cycles at a current density of 1 mA cm⁻² for a total of 1 mAh cm⁻² of Li. In addition, the symmetric cell presented stable voltage profiles with small hysteresis up to 300 h, indicating the formation of stable SEI. Based on detailed characterizations, we believe that the innovative structure of the IG current collector plays an important role in the improvement of the performance, which brings about a promising route to tackle the intrinsic issues of Li metal anodes.

2. Results and discussion

We used a rather scalable method to prepare the 3D IG framework. Fig. 1 illustrates the schematic of the preparation process and corresponding SEM images after each step during the synthesis. Fig. 1a shows the general preparation process. The vacuum filtration was utilized to get the composite film consisting of multiple layers of graphene oxide (GO) and abundant nickel (Ni) microspheres, whose diameter is in the range of 0.5–1 μm. The surface of the film was covered by large GO flakes (as shown in Fig. S1a) and the Ni spheres were dispersed between two GO layers (as shown in Fig. 1b). The thickness of the composite film is about 25 μm according to the side-view SEM images (Fig. S2a). After vacuum filtration, the composite film was punched to small circular disks with an area of 1.5 cm². Then, the small circular disks were loaded into a furnace for graphene chemical vapor deposition (CVD) [40–43], using methane (CH₄) as carbon source and argon (Ar) as carrier gas, and graphene was grown on the surface of Ni microspheres (Fig. 1c and Fig. S1b). The thickness of the film increased to about 30 μm according to the side-view SEM image shown in Fig. S2b. Afterwards, the obtained samples were immersed in Ni etchants to etch the Ni microspheres away. As a result, massive hollow graphene spheres were distributed in between each two graphene layers. From the side-view SEM image in Fig. 1d, the edges of the 2D graphene flakes can be clearly observed; however, we can only see the top surface of the 2D graphene flakes in the top-view SEM image in Fig. S1c. In this way, a freestanding and binder-free IG framework was produced, consisting of vertically stacked 2D graphene flakes together with sufficient void space. From the side view we can see that the thickness of the as-prepared sample is still about 30 μm (Fig. S2c), and most of the graphene hollow spheres are porous, which is good for Li ion transport during cycling.

The Raman data in Fig. 2a exhibits more information about samples after vacuum filtration, graphene CVD, and Ni etching. In the spectrum of the sample after vacuum filtration, the D-band (at about 1300 cm⁻¹) and G-band (at about 1600 cm⁻¹) were broad, and the 2D band (at 2700 cm⁻¹) did not exist, which is typical for Raman spectra of GO [44]. After CVD, the D-band and G-band became narrower, and the 2D-band appeared, indicating the successful synthesis of a graphene framework after the CVD process [45]. Also, the ratio of intensity of the G-band and 2D-band suggests the existence of few-layer graphene. During the CVD process, few-layer graphene was grown on Ni spheres and the 2D GO flakes were reduced to 2D graphene flakes at the same time, which is consistent with the Raman spectrum. The presence of D-band indicates that the GO flakes were not completely reduced and there were still some oxygen containing functional groups left. After Ni was etched way, the overall Raman spectrum almost did not change, but the D-band intensity increased a little bit, probably due to some defects created during the etching process. The X-ray photoelectron spectroscopy (XPS) measurement was also performed to study the change of the surface chemistry after each step during the preparation process. The XPS spectrum of the sample after vacuum filtration, which is depicted in Fig. S3a, can be separated into three peaks (284.8 eV, 286.8 eV and 288.0 eV). These three peaks can be assigned to C-C, C-O and O-C=O, respectively [46]. Samples after CVD (Fig. S3b) and after etching (Fig. S3c) have similar XPS spectra and they can be divided into two peaks (284.8 eV and 285.1 eV), representing the bonding of C-C and C=O, respectively. It also confirms the conclusion from the Raman spectroscopy.

In order to gain further insight into the morphology of graphene hollow spheres grown on Ni, we used transmission electron microscope (TEM) to characterize samples after CVD and after etching (shown in Fig. 2 and Fig. S4). Fig. S4a shows the low-magnification TEM image of the sample after CVD. One can clearly see that the structure of “graphene oxide layer-Ni spheres-graphene oxide layer” is still maintained after graphene CVD. While in high magnification, the graphene layer whose thickness is ~10 nm can be observed on the surface of the Ni spheres (Fig. S4b and Fig. 2b). After Ni etching, graphene hollow spheres can be clearly observed and the thickness of the graphene shell kept similar to the thickness of samples before etching (shown in Figs. S4c and 2c). Fig. 2d to Fig. 2f present the scanning transmission electron microscopy (STEM) images and corresponding energy-dispersive X-ray spectroscopy (EDX) mapping data of the samples after Ni etching, which gives us information about element distribution. It is clear that the major element of the sphere is C, and Ni is only in a very small amount, which is consistent with the data shown in Fig. S5 given by SEM EDX measurement. The atomic ratio of Ni was 35.2% before etching and it decreased to less than 1% after the sample was immersed in Ni etchant for 40 min. The remaining trace amount of Ni may serve as Li nucleation seeds for stable Li deposition [47,48].

The unique structure of our 3D IG shows potential to be utilized for stable Li metal anodes. Fig. 3a and b illustrate our hypothesis of Li metal deposition behavior on planer Cu foils and IG. When Li metal is deposited onto Cu foils directly, Li dendrites will be formed. As the Li dendrites grow longer and longer after several cycles, some of them will lose electrical conduction and become dead Li. The dead Li cannot be stripped any longer, resulting in reduced Coulombic efficiencies. Furthermore, the Li dendrite can also puncture the separators, leading to short circuit and severe safety problems. On the other hand, when Li metal is deposited onto a 3D IG current collector, the effective current density would decrease due to the increased effective surface area, reducing the probability of Li dendrite formation. Even though some dendrites may still be formed occasionally, they would be suppressed.
by the planer 2D graphene layers at the early stage. Therefore, we are able to achieve more stable Li deposition with high Coulombic efficiencies and flat surface morphology. The electrochemical behavior of Li plating/stripping on 3D IG was examined in coin cells and the pristine Cu foil was tested as a control sample. Li foils were used as the counter electrodes, and 1 M lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) in 1,2-dimethoxyethane (DME) and 1,3-dioxolane (DOL) (volume ratio 1:1) with 0.2 M lithium nitrate (LiNO₃) as additive was used as the electrolyte. In each cycle, the batteries were first discharged for 1 h and then charged to 1 V at a constant current density of 1 mA/cm². Fig. 3c shows the cycling performance of a planar Cu current collector and a 3D IG current collector. The Coulombic efficiency of 3D IG electrode kept increasing in the first 5 cycles, which can be attributed to the formation of solid electrolyte interface (SEI), and then maintained at a high level up to 100 cycles. The overall average Coulombic efficiency was 97.5% and the average Coulombic efficiency after 5 cycles was as high as 98.5%. In comparison, the Coulombic efficiency of bare Cu electrode maintained above 90% in the early stage, but degraded dramatically after 30 cycles. After 90 cycles, the Coulombic efficiency was as low as about 20%. Obviously, the 3D IG showed much better electrochemical performance of Li plating/stripping, which is consistent with our hypothesis. The discharge/charge curves of the IG electrode are shown in Fig. 3d. In the first 10 cycles, the formation of SEI consumed part of plated Li, leading to the relatively low Coulombic efficiency in the discharge/charge curve. Afterwards, the Coulombic efficiency was close to 100% from 20 cycles to 100 cycles, indicating that almost all the plated Li metal can be stripped reversely after the formation of stable SEI. Similar behaviors were observed when the current density was increased to 2 mA cm⁻² (Fig. S6a), or when the Li loading was increased to 2 mA h cm⁻² (Fig. S6b). At higher current density or higher Li loading, the 3D IG electrodes can still facilitate stable Li deposition in the first 100 cycles, and performed much better than the bare Cu electrode. The specific capacities are calculated to be 2143 mA h g⁻¹ (with 1 mA cm⁻² Li loading) and 4286 mA h g⁻¹ (with 2 mA cm⁻² Li loading) based on the total mass of IG current collector, which is much higher than the theoretical gravimetric capacity of graphite (372 mA h g⁻¹) [49]. In order to further investigate the cycling stability of Li metal anodes with different current collectors, we assembled symmetric cells using IG current collector with deposited Li as the working electrodes, and Li foils as the counter electrodes. As control samples, we also assembled bare Cu current collectors with deposited Li as the working electrodes, and Li foils as the counter electrodes. The galvanostatic discharge/charge voltage profiles of symmetric cells with different current collectors are depicted in Fig. 3g and f. At 1 mA cm⁻², the Li metal anode with IG current collector exhibited excellent cycling stability with small overpotential and negligible voltage fluctuation up to 300 h, owing to the good electrode-electrolyte contact and uniform SEI layer. In contrast, the overpotential of the symmetric cell with a bare Cu current collector was much larger initially. After 75 h, the overpotential increased dramatically and the voltage profiles fluctuated a lot, which is related to non-uniform Li deposition and fractal dendrite formation [48]. Similar behaviors were observed when the current density was increased to 2 mA cm⁻². The Li metal anode with IG current collector showed stable voltage profiles up to 200 h; however, the overpotential of the Li metal anode with bare Cu current collector increased rapidly after 40 h, which is consistent with the symmetric cell tested at 1 mA cm⁻². In this way, we can confirm the advantageous electrochemical performance of Li metal anodes with IG current collector.

In order to study the post-cycle surface morphology of Li metal anodes, SEM was performed for both IG and bare Cu current collectors after 50 cycles. The top-view SEM images of deposited Li on IG at different magnifications are presented in Fig. 4a and b. After the 50th plating process, 2D graphene flakes and 3D graphene spheres can still be observed on the flat surface. Although a small amount of Li can still be deposited on the top of the 2D graphene flakes, it would not grow to long dendrites and its effect can be ignored. In the high-magnification image shown in Fig. 4b, we can even clearly observe some Li dendrites grow underneath the 2D graphene flakes. Those dendrites were constrained by the 3D graphene top layer and cannot grow longer, which resolved the dendrite formation problem in the early stage. In comparison, the top-view SEM image of IG after the 50th stripping process is shown in Fig. S7. The morphology is similar to the morphology after plating due to the presence of SEI, while we can still observe more void space in the image after stripping. We believe that the stable SEI layers formed during cycling play an important role in the improved cycling performance. In order to study the composition of the SEI layers, XPS was conducted for IG electrodes before cycling, after 5 cycles and after 50 cycles, as shown in Fig. S8. From the C 1s spectrum, the O-C-O peak increased a lot during cycling, suggesting that COOLI is the main functional group in
From the F 1s spectrum, we can confirm the formation of LiF during cycling, which can work as a surface protection layer to suppress the Li dendrite growth [52, 53]. Fig. 4c and d shows the post-cycle cross-section SEM images of a Li deposited IG current collector with different magnifications. After Li plating, the thickness of the IG increased to about 50 μm, but the whole structure of the current collector can still be preserved thanks to the good flexibility of graphene. In the high-magnification image in Fig. 4d, the side edge of 2D graphene flakes can be clearly observed and deposited Li metal was sandwiched by two layers graphene. In addition, the deposited Li metal has particle-like morphology with a size of around 1 μm, which is similar to the size of the hollow graphene spheres. This phenomenon suggests that in the early stage of deposition, Li ions nucleate inside or around the hollow graphene spheres, as well as on the 2D graphene flakes. As more Li got deposited, the Li particles expanded and merged with each other. In this way, the presence of hollow graphene spheres can effectively increase the effective surface area. It is noticeable that the deposited Li metal is more likely to expand in the directions parallel to the 2D graphene flakes due to the physical confinement effect of the 2D flakes.

In contrast, dendritic growth of Li is observed when Li metal was deposited onto a bare Cu current collector, as shown in Fig. 4e. The Li dendrites can be as long as several tens micrometers and their diameters are usually less than 5 μm. These Li dendrites can either disconnect from the current collector and become dead Li, leading to reduced Coulombic efficiency, or easily puncture the separator, causing short circuit which may result in an explosion. In addition, the thickness of the deposited Li metal is as large as 120 μm, which is more than two times of the deposited Li on IG.
in thickness further confirms that a large amount of dead Li lost connection with the current collector during cycling, leading to continuous volume expansion.

In order to gain insight into the growth of Li metal, electrochemical impedance spectroscopy (EIS) analysis of samples using different current collectors was performed (shown in Fig. 5). The semicircle at...
high and intermediate frequencies corresponds to the interfacial resistance of the electrode and electrolyte. Before cycling, the two kinds of current collectors exhibit similar spectra. However, the cell using a bare Cu current collector shows much larger interfacial resistance than the cell using an IG current collector after 10 cycles, indicating the formation of a thicker and non-uniform SEI. The impedance spectra evolution of the cell with an IG current collector is depicted in Fig. 5b. An equivalent circuit (Fig. S9a) was used to fit the measured data and the results of interfacial resistance are shown in Fig. S9b. It’s worthy to note that there is almost no change in the
interfacial resistance (about 15 Ω) from the 5th cycle to the 50th cycle, confirming that a stable SEI was formed during cycling. We believe that the 2D graphene flakes play a key role in the formation of a stable SEI, which is very important to improve the Coulombic efficiency and avoid short circuit.

3. Conclusion

In conclusion, we have demonstrated a novel design of 3D IG framework with vertically stacked 2D protective layers and its successful application for stable lithium metal anodes. The lithium metal anode with an IG current collector can achieve an average Coulombic efficiency of 97.5% in 100 cycles at current density of 1 mA cm⁻² and it can be cycled for over 300 h when assembled as symmetric cells with stable voltage profiles. This outstanding performance can be attributed to the large surface area of the IG network, sufficient ion channels provided by graphene hollow spheres, and excellent lithium dendrites suppression by the 2D graphene layers on the top. The morphological study also shows that after cycling the electrode surface is rather smooth and a stable SEI is formed, which is further confirmed by EIS measurement. These results indicate that the major challenge encountered in the development of lithium metal anodes, including safety risks and low Coulombic efficiency, can be resolved effectively. We believe this strategy can also be applied in other alkali metal anodes like Na and full cells with various cathodes, implying huge practical application potential in energy storage field.

4. Experimental section

4.1. Materials

Nickel spheres were purchased from US Research Nanomaterials Incorporation. The graphene oxide water dispersion was purchased from Graphenea Incorporation. The nickel etchant was purchased from Alfa Aesar. All chemicals were used as received.

4.2. Preparation of 3D IG network

In a typical process, 100 mg of 1 µm nickel microspheres was dispersed in 10 ml ethanol and was sonicated to form a uniform solution. 1 ml of graphene oxide water dispersion was then added into the solution and the mixed solution was stirred for 10 min. Vacuum filtration with the help of a vacuum pump was conducted immediately after stirring to form a flexible composite film. The film was punched into small circular disks with an area of 1.5 cm², and then loaded into a one-inch quartz tube furnace. The furnace was first evacuated and filled with a mixture gas of 80 scen Ar and 20 scen H₂. Then it was heated up to 850 °C in 15 min and was kept at this temperature for 10 min to anneal the films under 40 scen Ar and 10 scen H₂. The small circular films were then exposed to 40 scen Ar, 5 scen CH₄ and 5 scen H₂ for 10 min before they were cooled down to room temperature with a rate of 10 °C min⁻¹. The two-layer graphene was grown on the surface of the nickel microspheres during this process. The as-prepared films were immersed in nickel etchant for certain minutes, washed with deionized water twice, and dried in ambient environment to form the 3D IG frameworks. The total mass of an IG circular disk is about 0.7 mg.

4.3. Characterization

The morphology and the microstructure of the electrodes was characterized with a SEM (JEOL JSM-7001) operating at 10 kV and a TEM (JEOL 2100F). The element distribution was measured on the EDX attached to the TEM. Samples after cycling were washed with DOL/DME to remove the electrolyte residue and dried in an Ar-filled glovebox. The Raman spectroscopy measurement was carried out on a Renishaw inVia Raman microscope and the XPS data was measured using a Kratos Axis Ultra XPS microprobe system to analyze the surface chemistry.

4.4. Electrochemical measurements

For electrochemical testing, CR2032 coin cells were assembled in an Ar-filled glovebox with O₂ and H₂O concentration below 0.1 ppm. The Netware battery testing system and Gamry workstation were used to perform the measurements. The as-prepared 3D IG framework, a bare Cu foil, a Li metal foil were used as the modified electrode, the controlled electrode, and the counter electrode, respectively. Celgard 2400 was used as the separator and the electrolyte was the lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) in 1,2-dimethoxyethane (DME) and 1,3-dioxolane (DOL) (volume ratio 1:1) with 0.2 M lithium nitrate (LiNO₃) additive. The cycling test was carried out by discharging at 1.0 mA cm⁻² for 1 h and then charged to 1.0 V under the constant current density. For higher current rates, the cells were discharged at 2.0 mA cm⁻², and then charged to 1.0 V at the same current.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.ensm.2018.09.028.

References

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