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Structure design and mechanism analysis of silicon anode for lithium-ion batteries

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ABSTRACT Silicon-based material is one of the most promising substitutes of widely used graphite anodes for the next generation Li-ion batteries due to its high theoretical capacity, low working potential, environmental friendliness, and abundant natural resource. However, the huge volume expansion and serious interfacial side reactions during lithiation and delithiation progresses of the silicon anode are the key issues which impede their further practical applications. Rational designs of silicon nanostructures are effective ways to address these problems. In this progress report, we firstly highlight the fundamental scientific problems, and then focus on recent progresses in design, preparation, *in-situ* characterization methods and failure mechanism of nanostructured silicon anode for high capacity lithium battery. We also summarize the key lessons from the successes so far and offer perspectives and future challenges to promote the applications of silicon anode in practical lithium batteries.

Keywords: silicon anode, Li-ion batteries, structure design, *in situ* characterization, mechanism analysis

INTRODUCTION

Rechargeable lithium-ion batteries (LIBs), as one of the energy storage technologies, are the key part of the contemporary energy system which is closely coupled with renewable energy generation, transmission and utilization [1–4]. Meanwhile, due to the high energy density, long lifespan and environmental friendliness, LIBs have been highly developed over the past decades and widely used in our daily life including portable electronics, electrical transportation, and even grid energy storage [5–7]. However, the current most mature LIBs cannot ever meet the ever-increasing demands [8–9]. Thus, a lot of new electrode materials with different storage mechanisms

have been explored to improve the battery performance and reduce the cost [10–13]. Among them, alloy-type Si anode is one of the outstanding examples demonstrating great promise and broad research interest because of its high capacity (4200 mA h g^{-1}), low operation potential (0.2–0.3 V vs. Li^+/Li), environmental friendliness, and high abundance [14]. Therefore, Si appears to be a very promising anode material for the next generation battery, especially in electric vehicles which require to be powered by LIBs with higher energy density and less safety issues [15–17].

However, the alloy reaction mechanisms of silicon are significantly different from those of conventional layered electrodes. There are many challenges that emerge during lithiation/delithiation progress [18–20] (Fig. 1): complete destruction of crystal structure producing huge volume expansion (ca. 420%) at the electrode and cell level, chemical bond breaking/reformation with serious interfacial side reactions, and relatively low intrinsic electronic conductivity (ca. $10^{-3} \text{ S cm}^{-1}$) of silicon, which will limit the cycling stability, Coulombic efficiency (CE) and rate capability, and thus severely impede their further practical applications. In the past several years, significant breakthroughs have been achieved by combining advanced nanomaterial preparation method with *in-situ* characterization technologies to improve the electrochemical performance and gain a better understanding of the mechanism under the real working condition of Si anode [21]. Although there have been some excellent reviews in this topic [22–23], the rapid development and newly emerging research results are achieved in the last six to seven years, and more than 700 papers are published every year (Fig. 2). It is essential to summarize the developing trend and direction of silicon anode in recent

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years to provide some revelation for the further applications.

In this review, we first briefly discuss the fundamental scientific challenges and solutions as well as the development history of silicon anodes, and then focus on the recent progress of rational structure designing; specific examples are provided to illustrate these fundamental design principles. We also highlight *in-situ* techniques utilized for in-depth understanding the silicon anode performance and fading mechanisms, including atomic force microscopy (AFM), transmission electron microscopy (TEM), nuclear magnetic resonance (NMR), and X-Ray powder diffraction (XRD). Finally, the major challenges and opportunities on silicon based anode materials are presented.

BASIC INTRODUCTION OF SILICON ANODE

Fundamental scientific problems

Silicon (Si) commonly exists in silicates and silica, which are widely found in rocks, gravel, and dust, but it rarely appears in the form of elemental silicon in nature. Silicon atom locates in IV main group of the periodic table with an atomic number of 14. The four valence electrons in the outermost layer make silicon atoms in a metastable structure, which plays a dominant role in conductivity and other properties. The crystalline silicon is three-dimensional (3D) diamond structure in cubic $Fd-3m$ space group with lattice constants of 5.431 \AA , as shown in Fig. 3a. Silicon is bonded to four equivalent Si atoms to form corner-sharing SiSi_4 tetrahedra. Chemical properties of silicon are relatively stable. It is difficult to react with other substances (except the hydrogen fluoride and lye) at room temperature. Commercial silicon is usually divided into metallic silicon and semiconductor silicon. Metallic silicon is mainly used to make compounds of poly-

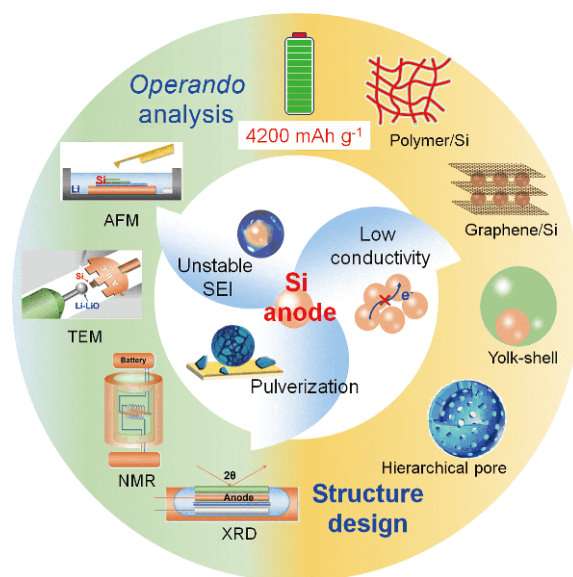


Figure 1 Overview of silicon-based anode for LIBs: fundamental challenges, recent structure design strategies toward commercialization, and failure mechanism analysis technologies.

crystalline silicon, monocrystalline silicon, silicon-aluminum alloy and silicon-steel alloy. Semiconductor silicon is used for producing semiconductor devices. Silicon-based materials have been widely used in aerospace, electronics, electrical, construction, transportation, energy, chemical, textile, food, light industry, medical, agriculture and other industries [24].

When Si is applied as anode materials for LIBs, it is important to know the properties of Li. Lithium is the lightest metal element with an atomic number of 3, two electrons in the K layer and one in the L layer. Since lithium has a high charge density and a stable helium double shell, it is easy to polarize other molecules or ions, but not easy to polarize itself. Lithium metal has a typical body centered cubic structure as shown in Fig. 3b. When

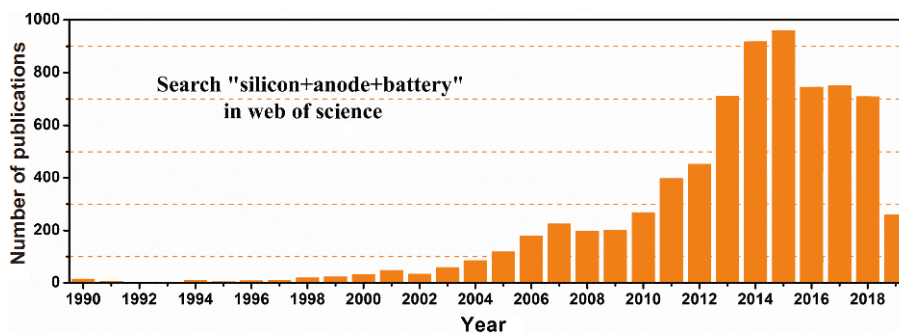


Figure 2 Statistical analysis of the publications on silicon anode in last three decades. The data was up to May 1, 2019.

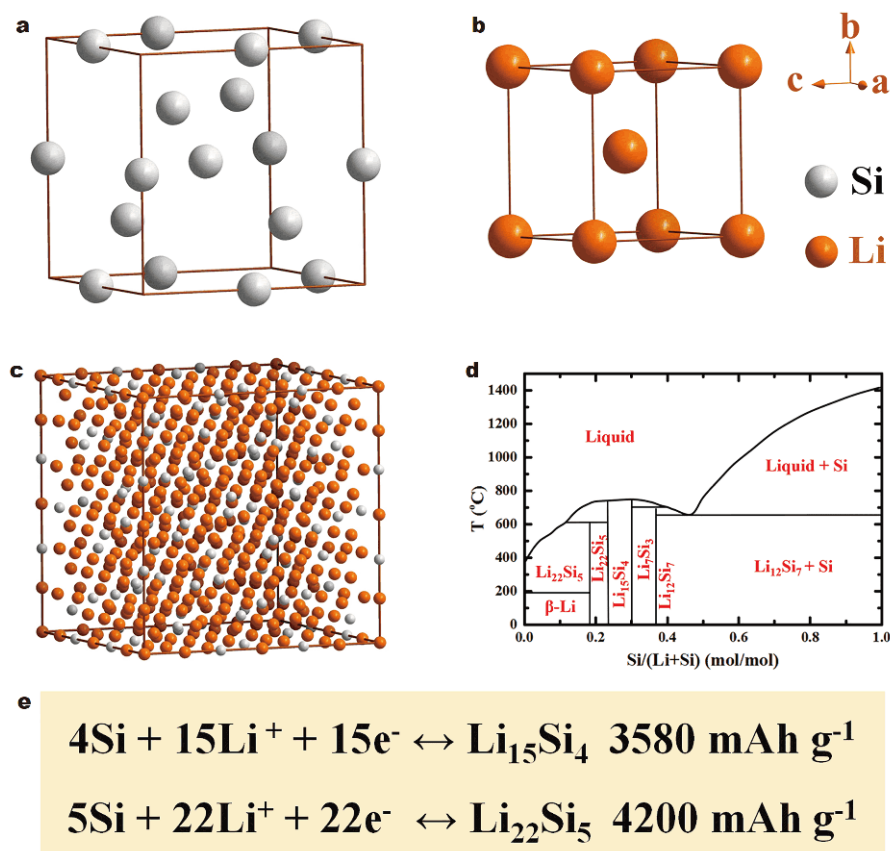


Figure 3 The crystal structures of silicon (a), lithium (b) and $\text{Li}_{22}\text{Si}_5$ (c). Redraw phase diagram of Li-Si system (d). Reprinted with permission from Ref. [24], Copyright 2015, Elsevier. Electrode reactions and theoretical capacity of different alloy products (e).

the temperature drops to -201°C , it starts turning into a face-centered cubic structure—the lower the temperature is, the greater the change will be, but not completely change. Lithium has the most negative electrode potential, making it the most active metal. Silicon is an alloy type anode, and each silicon atom can host 4.4 Li atoms at most to form $\text{Li}_{22}\text{Si}_5$ (Fig. 3c, e), reaching the maximum theoretical capacity. This mechanism of lithiation is associated with massive volume changes due to the fact that the unit cell volume of cubic Si is 40.88 \AA^3 , and 1617 \AA^3 for cubic $\text{Li}_{22}\text{Si}_5$ [25]. According to the Si-Li system phase diagram as shown in Fig. 3d, a series of Si-Li phases exist during the alloying process, such as LiSi, $\text{Li}_{12}\text{Si}_7$, $\text{Li}_{15}\text{Si}_4$, and $\text{Li}_{22}\text{Si}_5$.

These crystalline phases possess lower formation energy than the corresponding amorphous phases, which tend to be more kinetically stable. However, the amorphous phases Li_xSi (with $x=0-3.75$) are favorably formed during the electrochemical lithiation. It has been found that when x reaches 3.75, the amorphous phase will transform to crystalline phase. These amorphous phases and con-

tinuous unstable solid electrolyte interphase (SEI) production result in large volume change. The large volume expansion/contraction of Si usually causes cracking and pulverization of Si, resulting in unstable SEI and loss of electrical contact of the current collector [26], which finally lead to low CE and rapid capacity fading. In addition, the Li^+ insertion-induced large deformation and elastic strain lead to fracture or change in morphology of electrodes, which will significantly slow down the kinetics of lithiation process. When silicon cannot withstand the resulting stress, the cracks will occur due to the brittleness and the material will peel off from the electrode, lose electrical contact, convert to dead weight and thereby decrease the specific capacity of the electrode [27].

History of silicon anode research

With the passage of time and development of technology, substantial efforts have been made in recent decades to solve the problems of the silicon anode. A timeline of selected important breakthroughs in the silicon anode materials is shown in Fig. 4. Due to the fast capacity

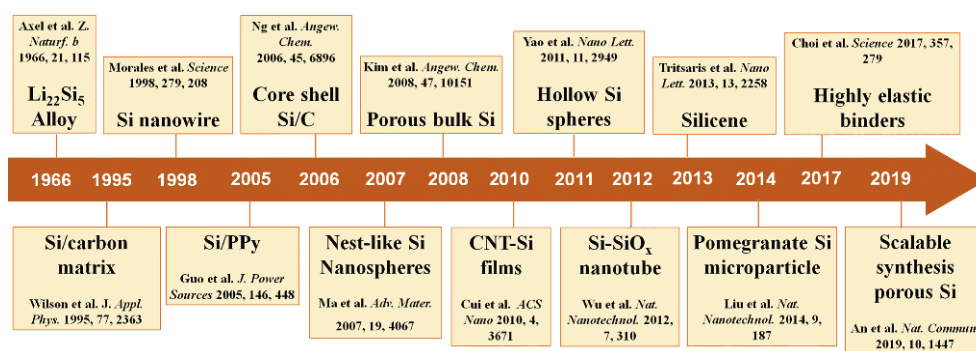


Figure 4 Timeline of selected important breakthroughs in the silicon-based anode.

degradation for limited cycles, silicon anode has been overlooked for a long time since the first discovery of Li₂₂Si₅ alloy in 1966 [28]. In 1971, Dey [29] found that lithium could electrochemically alloy with Sn, Pb, Al, Au, Pt, Zn, Cd, Ag and Mg at room temperature. Then Seefurth and Sharma [30] reported the formation of Li–Si alloys at high temperature cells. But the reversibility of normal Si powder was very poor due to the volume expansion. Thus, developing effective strategies to resolve the problem of volume expansion has been identified as a central issue of the research on Si-based LIB anodes. In 1995, with the development of nanotechnology, nano-sized Si composite had been regarded as the promising method to improve the electrochemical performance. Wilson *et al.* [31] dispersed nano-sized Si into carbon matrix and obtained a capacity of 600 mA h g⁻¹, which was about twice that of the commercial graphite anode. Li *et al.* [32] prepared Si/C composite by manually grinding the mixture of nano-sized Si and carbon black under ambient condition. The composite exhibited a capacity of 1700 mA h g⁻¹ even after ten cycles. It was found that nano-sized Si particles played an important role in the improvement of the cyclic stability due to their good plasticity and deformability. In 1998, Lieber's group [33] successfully synthesized Si nanowires which were considered to be superior to nanoparticles. Benefiting from the continuous electronic pathway, nanowires can efficiently facilitate charge transport. The Si nanowires could realize the theoretical charge capacity of silicon, but the capacity also degraded rapidly after a few cycles. With the development of conductive polymers in the 21st Century, conductive polymers such as polypyrrole (PPy) [34] were used to enhance the charge carrier transportation and inhibit the huge volumetric change. In order to increase the surface area, Si nanotubes were synthesized [35]. The first discharge and charge capacities of the Si nanotubes were 3648 and 3247 mA h g⁻¹ at the current density of

600 mA g⁻¹ (0.2 C). However, the Si nanotubes had very low mass loading density, which limited the practical applications. Therefore, constructing 3D Si network incorporated with pores became an attractive strategy to resolve the problem of the Si nanotubes. Kim *et al.* [36] prepared 3D porous bulk Si particles. The 3D porous structure effectively accommodated stress without pulverization after 100 cycles, and maintained charge capacity of more than 2800 mA h g⁻¹ at 400 mA g⁻¹. Our group also prepared nest-like Si particles composed of hollow Si spheres through solvothermal reaction in 2007, which displayed an initial specific capacity of 3052 mA h g⁻¹ at the current density of 2000 mA g⁻¹, and retained 1095 mA h g⁻¹ after cycling up to 48 cycles [37]. Because the commonly used heavy metal copper current collector for traditional Si film anode was much heavier than the Si active material itself, it unavoidably increased the total mass of the LIBs. To resolve this problem, a novel anode structure free of heavy metal current collector was designed by integrating a flexible, conductive carbon nanotube (CNT) network into a Si anode [38]. The CNT-Si films showed a high specific charge storage capacity and a good cycling life. Besides the conventional porous structures, nanostructured hollow Si was also designed and synthesized. Cui's group [39] fabricated hollow Si nanospheres which exhibited an initial discharge capacity of 2725 mA g⁻¹ with less than 8% capacity degradation every hundred cycles for total 700 cycles at 220 mA g⁻¹. The well-designed nanoparticles with a free volume in the hollow particle interior and the porosity in the shell effectively cushioned the volume change and consequently improved the electrochemical performance.

In order to address the SEI stability issue, Cui's group [40] designed a novel double-walled Si-SiO_x nanotube anode, in which the inner wall was active Si and the outer wall was the confined SiO_x, which still allowed Li ions to pass through. The outer surface of the Si nanotube was

prevented from expansion by the oxide shell. The expanding inner surface was not exposed to the electrolyte. This specific structure design helped to stabilize the SEI. The Si nanotube anode exhibited a high specific charge capacity (1780 mA h g^{-1} at 0.2 C). In 2013, Wang's group [41] employed the first-principles calculations to investigate the interaction of Li with Si in model electrodes of free-standing single-layer and double-layer silicone. Inspired by the structure of pomegranate, Cui's group [42] proposed a hierarchical structured silicon anode where single silicon nanoparticles were encapsulated by a conductive carbon layer that left enough room for expansion and contraction following lithiation and delithiation. As a result of this hierarchical arrangement, the SEI remained stable and spatially confined, resulting in superior cyclability (97% capacity retention after 1000 cycles). Up to now, scalable synthesis of silicon anodes with good cyclability and low electrode swelling remains a significant challenge.

Tremendous strategies have been proposed to deal with the issues listed above and improve the LIBs performance of silicon-based anode, such as downsizing silicon with various dimensions and microstructures [43–48], compositing with carbon [49–55], doping with metal [56–59], metal oxides [60–63] and conductive polymer modification [64–66]. In the following section, recent progress in these strategies will be discussed in detail.

RATIONAL DESIGNED SILICON ANODES

Nanostructured Si anode

Engineering of nanostructure has been proven to be one of the effective ways to buffer the significant volume change of silicon in the charge/discharge process due to their native features: large specific surface area promoting the infiltration between active materials and electrolyte, smaller particle size shortening the diffusion distance of electrons and lithium ions, as well as reducing the inhomogeneous lithium diffusion-induced stress and strain [14,67]. Thus, different nanostructures have been designed to improve the electrochemical performance of silicon anodes. In this part, various kinds of silicon nanostructures reported lately are briefly discussed, covering 0 dimensional (0D) nanoparticles, 1D nanowires, 2D nanosheets and silicene, 3D hierarchical nanostructure, hollow and porous structures. Turning the bulk silicon to nanoscale demonstrates one of the efficient strategies to overcome the large expansion/contraction problem by relieving the internal strain. In this regard, Kim *et al.* [68] reported well-dispersed 0D Si nanoparticles with various

particle sizes by a reverse micelle method for LIB anode. The capacity can still be maintained at 2700 mA h g^{-1} after 40 cycles at 0.2 C. The highest charge capacity was obtained with a critical size of 10 nm. Another example of such a particle-size-dependent conclusion was confirmed by Liu *et al.* [69]. By using *in-situ* TEM technology, the morphological changes of silicon particles with different sizes after the first lithium insertion were compared. It was found that with the decrease of size, silicon exhibited stronger resistance to mechanical strain and its cracking degree decreased. Researchers proposed that 150 nm was the critical point. Recently, consistent with these conclusions, Lin *et al.* [70] prepared crystalline Si nanoparticles through a reduction of micro-sized silicon zeolite by metallic Al in molten AlCl_3 at a low temperature of 200°C , as shown in Fig. 5a. The prepared Si used as an anode for LIBs reached reversible capacities of 2663 mA h g^{-1} at 0.5 A g^{-1} after 50 cycles and 870 mA h g^{-1} at 3 A g^{-1} after 1000 cycles.

In addition to 0D silicon nanoparticles, 1D silicon nanowires can also effectively improve the electrochemical performance. In 2007, Cui's group [71] firstly used Si nanowires vertically aligned on stainless steel substrate with a diameter of 90 nm as anodes of LIBs by the chemical vapor deposition (CVD) method. In their study, the improved LIB performances were ascribed to the advanced architecture of the Si nanowires electrodes, which provided sufficient space between the nanowires to release the strain from large volume change, and enhanced the electrical contact between Si nanowires and the substrate. Very recently, Wang *et al.* [72] used a magnetron sputtering deposition with metal-catalyzed electroless etching technology to produce silicon nanowire film (Fig. 5b), which allowed for achieving the high capacity and excellent cyclability. The prepared anode delivered an initial reversible discharge capacity of 3158 mA h g^{-1} and maintained a capacity of 2840 mA h g^{-1} after 100 cycles when cycled at a current density of 200 mA g^{-1} .

Additionally, 2D Si nanostructure possesses a smaller specific variable SEI surface area than that of other dimensional Si nanostructures. From this aspect, Chen *et al.* [73] synthesized ultrathin mesoporous 2D Si nanosheets by a soft template and subsequent magnesiothermic reduction method. Owing to the ultrathin 2D characteristics and mesoporous structure, it could simultaneously buffer the structural stress, mitigate the volume expansion, accelerate the electrolyte permeation and shorten the lithium diffusion pathway, giving rise to a superior cycle performance. Moreover, Zhang *et al.* [74] developed

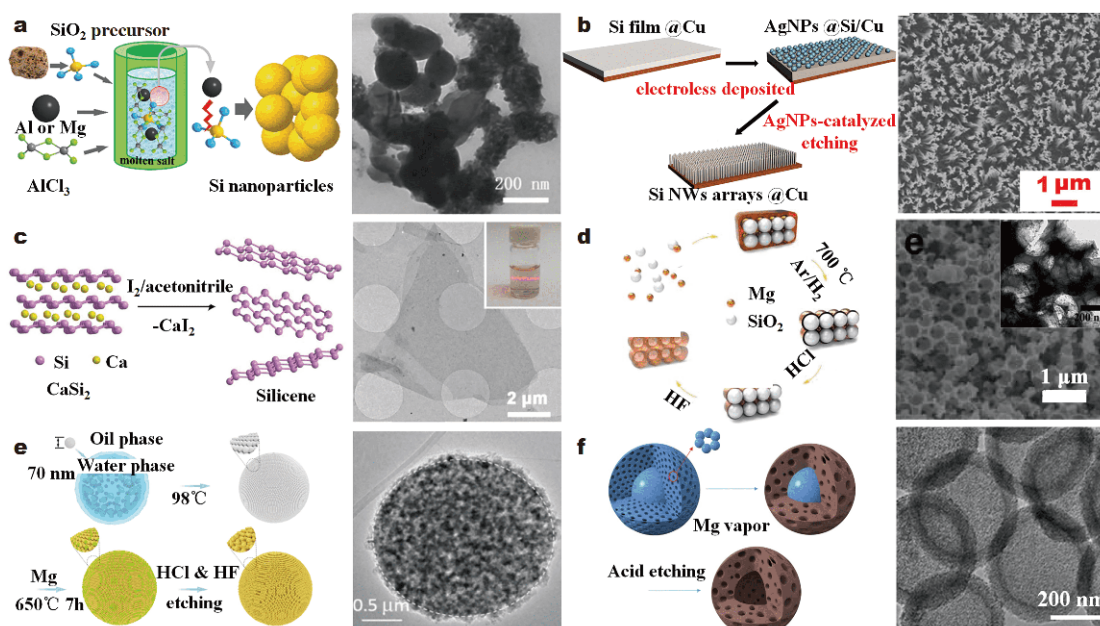


Figure 5 Schematic illustration of the synthesis process of different nanostructured silicons for LIBs: (a) 0D nanoparticles. Reproduced with permission from Ref. [70]. Copyright 2015, the Royal Society of Chemistry. (b) 1D nanowires. Reprinted with permission from Ref. [72]. Copyright 2017, Springer-Verlag GmbH Germany. (c) 2D nanosheets. Reprinted with permission from Ref. [75]. Copyright 2018, Wiley-VCH Verlag GmbH & Co. (d) 3D hierarchical structures. Reprinted with permission from Ref. [76]. Copyright 2016, American Chemical Society. (e) Porous structure. Reprinted with permission from Ref. [77]. Copyright 2018, Elsevier Ltd. (f) Hollow structures. Reprinted with permission from Ref. [78]. Copyright 2015, Nature Publishing Group.

a dual stabilized silicon building block, comprised of inherently interconnecting, spatially orientated silicene nanoplates to address the structure and interface stability issues of silicon. Lately, a scalable preparation of free-standing high-quality silicene *via* liquid oxidation and exfoliation of CaSi_2 was reported by Liu *et al.* [75] (Fig. 5c). The obtained silicene with monolayer or few-layer thickness was further used for LIB's anode, exhibiting a reversible capacity of 721 mA h g^{-1} at 0.1 A g^{-1} and an extraordinary cycling stability with no capacity decay after 1800 cycles.

Apart from the downsizing silicon with regulable morphology strategy, creating macro/mesopores silicon structure is another effective way to accommodate distinct volume change that accompanies with the lithiation process. Furthermore, the interconnected pores provide the channels for quick lithium ions transport and full infiltration of electrolyte, which guarantee an excellent rate capability and cycle stability. Well-crystallized 3D hierarchical macro-/mesoporous silicon structure (Fig. 5d) was synthesized through the magnesiothermic reduction process using 0D silicon particles as self-template [76]. The macro-/mesoporous silicon exhibited significantly improved cyclic and rate performance. A

reversible capacity of 959 mA h g^{-1} was retained after 300 cycles at 0.2 A g^{-1} with a high mass loading density of 1.4 mg cm^{-2} . Consistent with this strategy, Jia *et al.* [77] synthesized micrometer-sized silicon-based anodes with a unique porous structure (Fig. 5e) demonstrating good electrochemical performance. The void space derived from mesopores by the removal of the MgO phase could effectively accommodate the volume change of silicon and thus alleviate the breakdown of the SEI layer, all of which led to a high reversible capacity of 1467 mA h g^{-1} at 2.6 A g^{-1} , good rate capacity, and superior cycling performance up to 370 cycles.

Introduction of space into the nanostructured silicon to deliver an extra room is another efficient way to accommodate the unavoidable volume expansion and buffer the Li-ion diffusion induced mechanical strain. Xiao *et al.* [78] developed a combination of magnesiothermic reduction and acid etching method for the synthesis of hollow porous Si nanospheres (Fig. 5f). *In situ* TEM and chemo-mechanical modelling characterization demonstrated that both the mesoporous shell and hollow internal void were responsible for the inward expansion, contraction and stable SEI formation during lithiation and delithiation, which further enhanced the capacity

retention and cycling performance.

Si-carbon composites

Combining nanoscale silicon with various morphologies and structures has been proven to be an efficient strategy to improve the LIB performance. Although nanosized silicon can address the pulverization, the volume expansion of silicon still exists, which means that the SEI films will be destroyed and continuously produced. This will consume a large number of lithium ions, seriously reducing the CE and the reversible specific capacity of batteries. In addition, the conductivity of nano silicon is still an issue to improve the rate performance.

To date, carbon material is considered as one of the best candidates to address the above issues of silicon electrode because of its outstanding electrical conductivity and mechanical strength [79]. Conventional Si-C composites were commonly produced by mechanical mixing method, leading to an inhomogeneous disperse and poor connectivity of each component. In this regard, carbon uniformly coated on silicon with diverse structures becomes a promising approach to improve the

electrochemical performance [54,55].

Encapsulation of polybenzimidazole (PBI) derived from pyrrolic N-enriched carbon onto microsized silicon spheres [80] was achieved by an aerosol-assisted assembly combined with physisorption process (Fig. 6a). Thanks to the intrinsic high electronic conductivity, abundant pyrrolic nitrogen and structure robustness, the mesoporous Si-PBI carbon composite can avoid direct contact between silicon and electrolyte, prevent uncontrolled SEI film formation and create numerous extrinsic defects and active sites for extra lithium storage, which further enable the Si-PBI carbon composite with superior electrochemical performance in terms of high reversible specific capacity (2172 mA h g^{-1}), superior rate capability (1186 mA h g^{-1} at 5 A g^{-1}) and prolonged cycling life.

In order to buffer the volume expansion effect of silicon material more effectively, introducing voids between silicon and carbon layer is a more effective way [42,81]. In addition, the carbon shell prevents the inner silicon cores from aggregation and assists the formation of a stable SEI layer on the shell surface. Guo *et al.* [82] fabricated porous silicon nanoparticles (p-SiNPs) loaded in con-

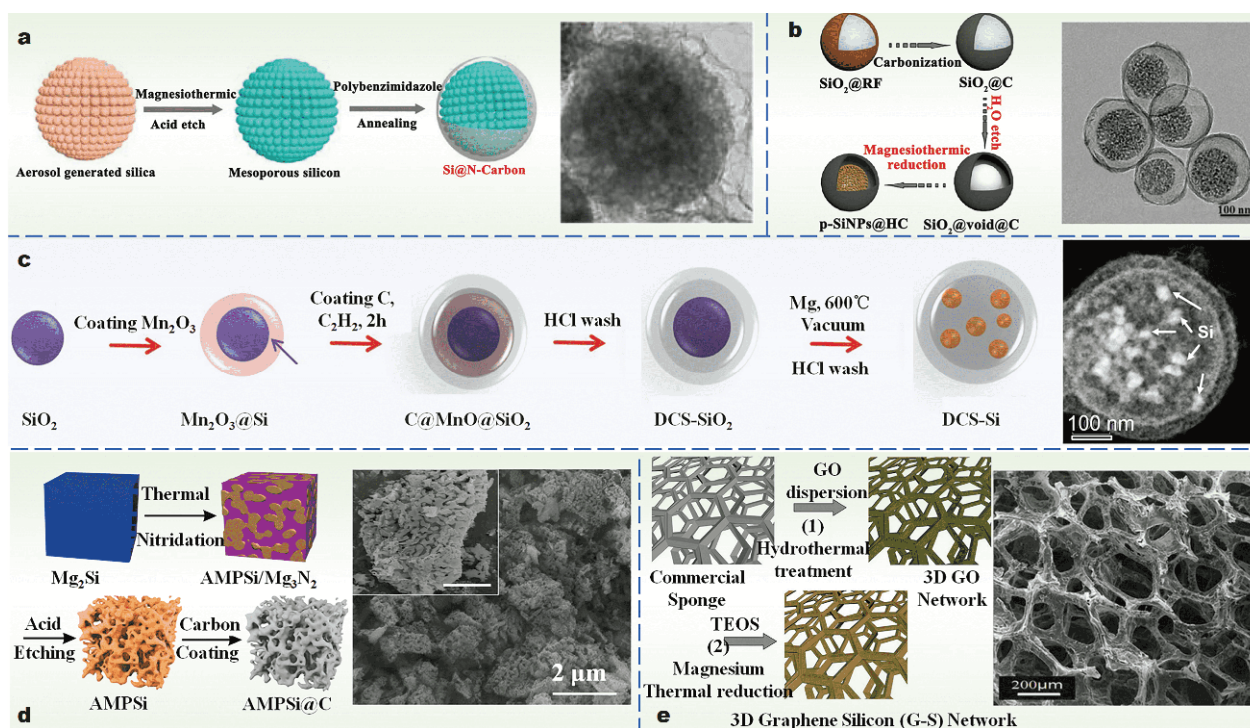


Figure 6 Schematic illustration of the synthesis process of Si-C composites and the corresponding morphologies or microstructures for LIBs: (a) Core-shell Si@N-C. Reproduced with permission from Ref. [80]. Copyright 2017, American Chemical Society. (b) Yolk-shell p-SiNPs@HC. Reproduced with permission from Ref. [82]. Copyright 2017, American Chemical Society. (c) SiNPs@double carbon shells. Reproduced with permission from Ref. [83]. Copyright 2017 Wiley-VCH Verlag GmbH & Co. (d) Ant-nest-like porous Si-C. Reprinted with permission from Ref. [84]. Copyright 2019, Nature Publishing Group. (e) 3D graphene-Si network. Reproduced with permission from Ref. [85]. Copyright 2015 WILEY-VCH Verlag GmbH & Co.

trollable void carbon spheres with yolk-shell structure by combining hydrothermal water etching and magnesiothermic reduction strategy, as shown in Fig. 6b. The optimized porous silicon nanoparticles@hollow carbon (p-SiNPs@HC) nanohybrids showed excellent LIB performance, delivering a capacity of 1400 mA h g^{-1} after 100 cycles at 0.2 A g^{-1} and 720 mA h g^{-1} at a high current density of 4 A g^{-1} . Moreover, Chen *et al.* [83] proposed a “double-shell” concept (DCS-Si) to confine the Si volume changes and stabilize the SEI layer using double carbon shells with certain inter spaces by CVD and magnesiothermic reduction method (Fig. 6c). The inner carbon shell provided certain voids to withstand volume changes inside the inner carbon shell and the outer shell stabilized the SEI layer in case of any cracks. Compared with bare Si and single carbon layer coated Si electrodes, the DCS-Si electrode showed a higher specific capacity of 1802 mA h g^{-1} at a current rate of 0.2 C, superior rate capability and good cycling performance up to 1000 cycles.

Porous structure designs to improve the performance of Si-C composite are similar to those of hollow structures. The space derived from porous structure can endure the volume expansion and alleviate the stresses. In addition, large surface area and uniformly distributed channels shorten the diffusion path for Li ions and increase reactivity of the composites, thus leading to enhanced rate capability. Lately, An *et al.* [84] reported a scalable top-down technique by thermal nitridation of the Mg-Si alloy and acid etching method to produce ant-nest-like porous silicon (shown in Fig. 6d). Synchrotron radiation tomographic reconstruction images and *in-situ* TEM characterization reveal that the ant-nest-like porous silicon consisting of 3D interconnected silicon nanoligaments and bicontinuous nanopores could prevent pulverization and accommodate volume expansion during cycling. The carbon-coated porous silicon anode delivered a high capacity of 1271 mA h g^{-1} at 2100 mA g^{-1} with 90% capacity retention after 1000 cycles and had a low electrode swelling of 17.8% at a high areal capacity of 5.1 mA h cm^{-2} .

Li *et al.* [85] reported an efficient approach to facile and large-area fabrication of 3D graphene-silicon (G-Si) networks that possessed porous structure, good flexibility, ultrathin hybrid walls and high electrical conductivity, as displayed in Fig. 6e. All these unique properties are beneficial to minimizing the volume change of silicon and facilitating the diffusions of both lithium and electron. As a consequence, the resulting 3D G-Si network exhibited a superior rate capability, cyclability and high reversible

capacity over 2050 mA h g^{-1} after 200 cycles.

Si-metal composites

The electrochemical properties of silicon can also be improved by compounding silicon with other metal materials especially at extremely high rates. On the one hand, metal doping can effectively improve the conductivity of silicon and promote the charge transfer process. On the other hand, the proportion of silicon in materials can be reduced by replacing part of silicon, and thus the volume expansion degree can be reduced. Moreover, the introduced metal can buffer the volume expansion of silicon to improve the electrode structural stability by using its good ductility. And some metals can form alloying products with silicon under heat treatment conditions, which are closely bound to silicon and further act as buffer layer to enhance the structural stability of materials when the volume of silicon expands. Besides, compounding with Li-active metals is also attractive, because they have different potentials for lithiation and delithiation. When Li is inserted into one component, the other will alleviate the volume change as a buffer. Song *et al.* [86] delivered a template-free synthesis of highly connected hollow Si-Cu nanotubes for high-rate and durable LIB binder-free anode. CuO was firstly grown on the copper foam by a thermal oxidation treatment, as illustrated in Fig. 7a. Then, coating amorphous silicon on CuO nanowires was achieved *via* a plasma-enhanced CVD technology. Finally, hollow Si-Cu nanotubes were obtained by an *in situ* H_2 annealing. The resulting hollow Si-Cu nanotubes anode possessed a high specific capacity of 780 mA h g^{-1} after 1000 cycles at 20 A g^{-1} , with a capacity retention rate of 88%.

Recently, 3D nanoporous SiGe alloy with tunable morphology and porosity [87] was synthesized as a high-performance LIB anode using a dealloying method with a ternary AlSiGe ribbon serving as the precursor, as shown in Fig. 7b. By adjusting the Al content in the precursor, the morphology and porosity could be controlled. With an Al content of 80%, the 3D-NP SiGe with continuous ligaments and hierarchical micropores and mesopores coral-like structure delivered a high reversible capacity of 1158 mA h g^{-1} after 150 cycles at a current density of 1000 mA g^{-1} with excellent rate capacity.

In addition, according to different preparation conditions and design methods, silicon/metal composites with different structures can be obtained. Zhang *et al.* [88] developed a composite anode of Cu/Si/Ge nanowire arrays, where each nanowire consisted of a core of Cu segments and a Si/Ge bilayer shell, as illustrated in Fig. 7c.

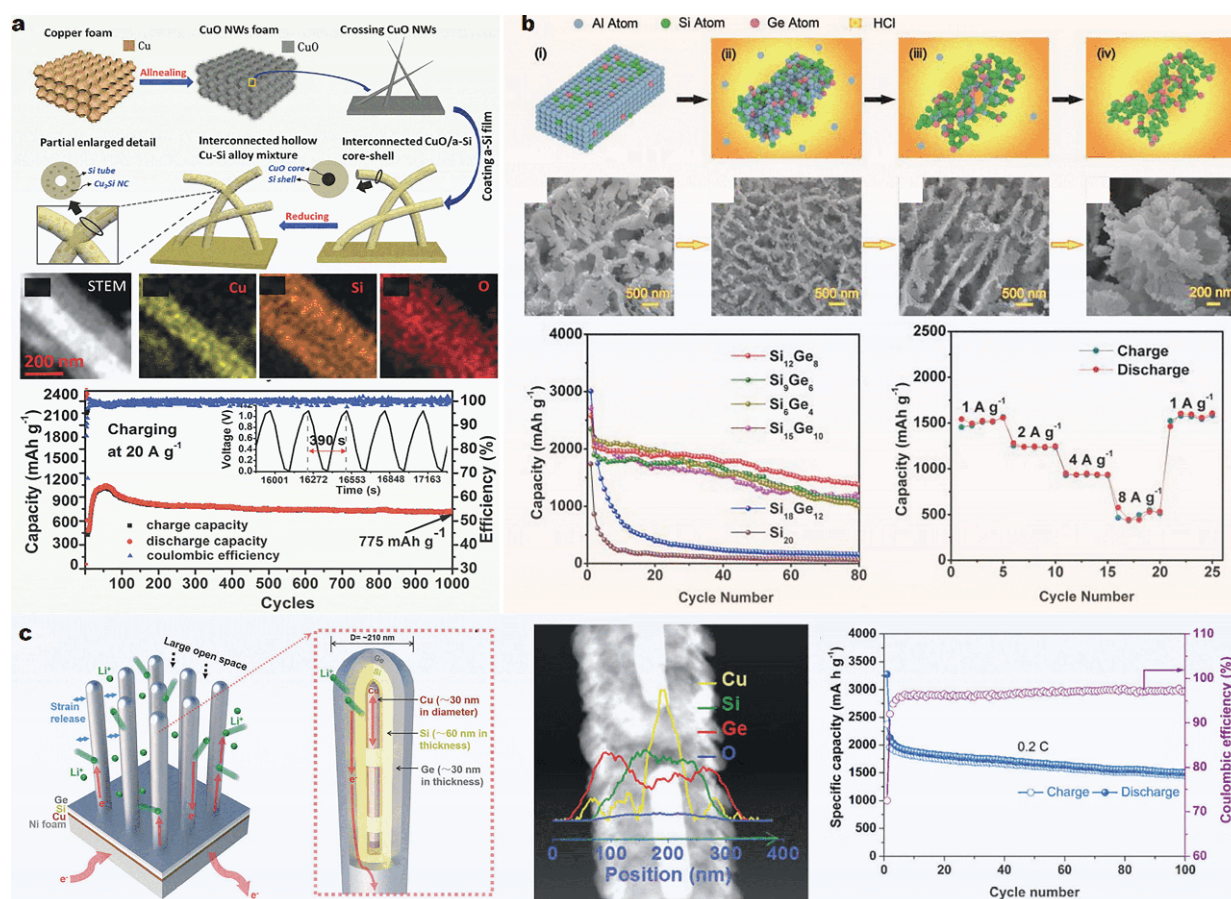


Figure 7 Si-metal composites. (a) Schematic illustration of the synthesis process of Cu-Si alloy structures, scanning transmission electron microscopy (STEM), energy-dispersive X-ray spectroscopy (EDS) mapping of CuO/amorphous-Si core-shell structure and the long term cycling performance at 20 A g⁻¹. Reprinted with permission from Ref. [86]. Copyright 2015, WILEY-VCH Verlag GmbH & Co. (b) Schematic illustration and SEM of the 3D-NP SiGe structure evolution *via* chemical dealloying. Cycle performance of the 3D-NP SiGe anodes and Si₂₀ anode at 100 mA g⁻¹ and rate capability of the Si₁₂Ge₈ anode at different current densities from 1 to 8 A g⁻¹. Reprinted with permission from Ref. [87]. Copyright 2018, American Chemical Society. (c) Schematic showing of Cu/Si/Ge array grown on a Ni foam substrate, STEM image of a Cu/Si/Ge nanowire with corresponding EDS spectra and capacity performance at a low rate of 0.2 C. Reprinted with permission from Ref. [88]. Copyright 2018, the Royal Society of Chemistry.

This unique electrode architecture exhibited several favorable properties: the nanoscale nanowires enabled facile strain relaxation; the free space between nanowires and the hollow space between Cu segments accommodated volume expansion; the core Cu segments enhanced electron transport. Moreover, the outer Ge shell also served as an active high-capacity coating. By using *in-situ* TEM and electrochemical testing, a novel co-lithiation/co-delithiation reaction mechanism was proposed, which effectively alleviated the electrochemically induced mechanical degradation and thus greatly enhanced the long-cycle stability of the electrode.

Si-metal oxide composites

Unlike alloys, Si/metal oxide composites with a core-shell

structure share a similar working mechanism with Si/C core-shell composites. The metal oxide coating layer on Si can separate silicon from electrolyte and stabilize SEI film. To add on this concept, Fang *et al.* [89] reported a rational design of core-shell Si@TiO₂ with Si NPs encapsulated in TiO₂ hollow spheres by combination of hydrolysis with magnesiothermic reduction method, as shown in Fig. 8a. In this architecture, the void space inside provided enough space for accommodation of the volume expansion, and the robust TiO₂ shell could improve the electrical conductivity and prevent the direct contact of Si with the electrolyte. As a result, the Si@TiO₂ hollow nanospheres maintained a specific capacity of 804 mA h g⁻¹ after 100 cycles at 0.1 C.

Lately, a facile sol-gel strategy was proposed to syn-

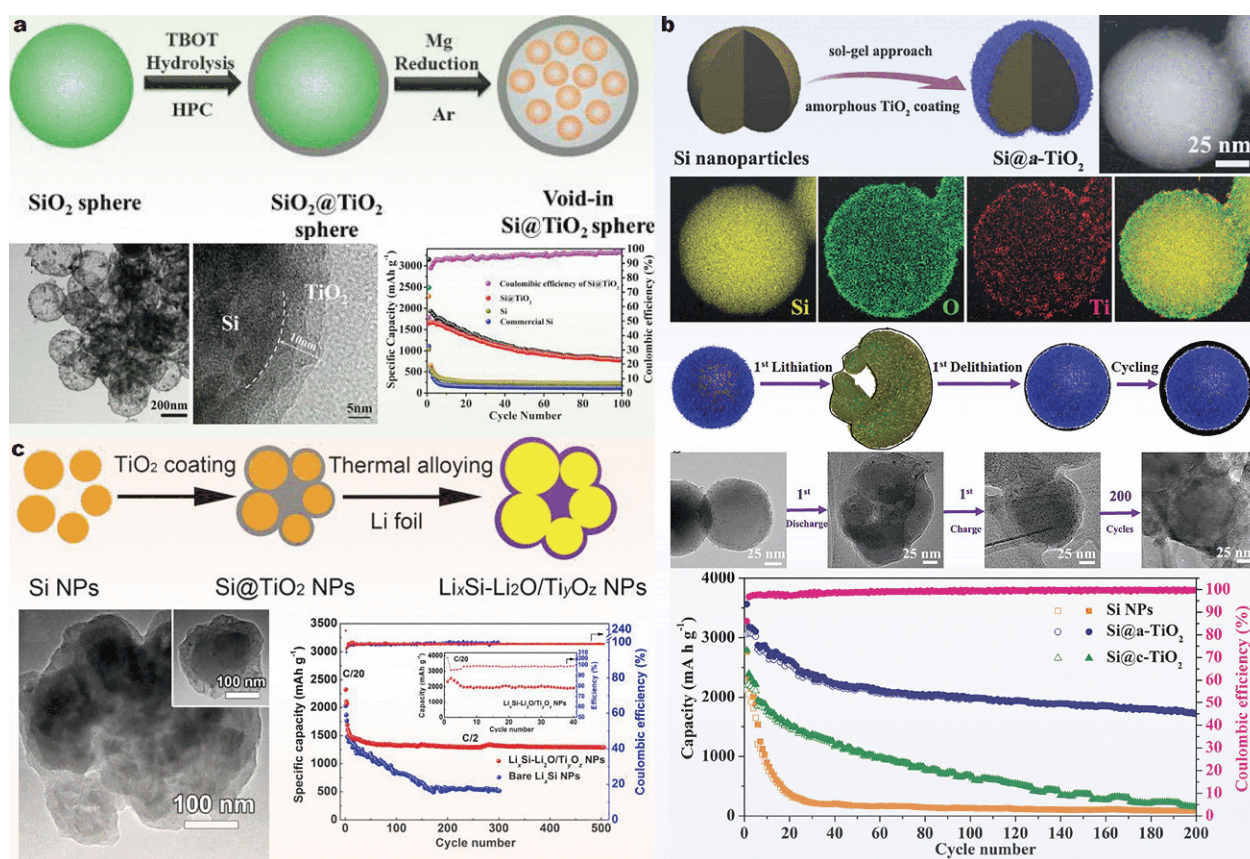


Figure 8 Metal oxides coating on silicon. (a) Schematic illustration for the fabrication process of the unique core-shell nanostructure of the Si@TiO₂ composite. TEM and high resolution TEM (HRTEM) images of the Si@TiO₂ composite. Cycling performance of the Si@TiO₂ composite and contrast samples at 0.1 C. Reproduced with permission from Ref. [89]. Copyright 2014, American Chemical Society. (b) Schematic of fabrication process of the amorphous-TiO₂-coated Si core-shell nanoparticles, STEM image and the corresponding EDS mapping images, schematic and TEM images of structural evolution of the amorphous-TiO₂-coated commercial Si nanoparticle electrode during electrochemical cycling, and cycling performance of the Si, Si@a-TiO₂, and Si@crystal-TiO₂ (c-TiO₂) nanoparticle electrodes. Reprinted with permission from Ref. [90]. Copyright 2017, WILEY-VCH Verlag GmbH & Co. (c) Schematic of the fabrication process for Li_xSi-Li₂O/Ti_yO_z core-shell NPs, TEM images of the resultant Li_xSi-Li₂O/Ti_yO_z NPs and cycling performances of Li_xSi-Li₂O/Ti_yO_z core-shell NPs and bare Li_xSi NPs. Reprinted with permission from Ref. [91]. Copyright 2018, American Chemical Society.

thesize core-shell Si@amorphous TiO₂ (~3 nm) by Yang *et al.* [90], as exhibited in Fig. 8b. STEM and the elemental mapping characterizations clearly verified that a thin TiO₂ nanograin layer was conformally coated on the Si nanoparticle core. *Ex situ* TEM measurements revealed the structural evolution of the core-shell Si@amorphous-TiO₂ (a-TiO₂) nanoparticles during the initial lithiation and delithiation processes. The a-TiO₂ shell showed elastic behavior during cycling, avoided the electrolyte impregnation within the encapsulated Si core and finally maintained the structural integrity. Further electrochemical investigation revealed that the a-TiO₂ shell offered superior buffering properties over the pristine Si nanoparticles and crystalline TiO₂ layers coated Si for unprecedented cycling stability. More recently, Wang *et*

al. [91] reported coating Si with a-TiO₂ layers and then thermally alloy lithiation approach to prepare lithiated-TiO₂ protected Li_xSi nanoparticles (Li_xSi-Li₂O/Ti_yO_z NPs) as a LIB anode (displayed in Fig. 8c). The robust lithiated-TiO₂ matrix not only improved the electrical conductivity but also spatially limited the direct SEI formation on Li_xSi/Si cores during cycling. More importantly, the coating layer protected the most inner Li_xSi alloys from corrosion, leading to high dry-air stability. As a result, the resulting Li_xSi-Li₂O/Ti_yO_z anode achieved a capacity of about 1300 mA h g⁻¹ after 500 cycles at a high current rate of 0.5 C.

Si-conductive polymer composites and binders

Besides the carbonaceous, metal and metal oxide mate-

rials, conducting polymers with unique features including excellent chemical stability, superior electronic conductivity and structural flexibility have also been introduced to accommodate the volume expansion of Si and to form conductive networks for enhanced reaction kinetics of the electrode. A strong interface bond between Si and polymer and a high electrical conductivity of polymer are the most critical parameters determining the electrochemical performance of Si/conducting polymer composite electrodes.

For instance, strong interfacial bond between polyaniline (PANi) and Si NPs was achieved through hydrogen bonding with the native oxide layer on Si particles by *in-situ* polymerization [92], as displayed in Fig. 9a. The resulting materials were in a well-connected electrically conductive 3D network consisting of Si NPs conformally coated by the conducting polymer, whose porous space was beneficial to volume expansion of Si particles. In

contrast to fast capacity fading for Si/PANi mixture, an electrode capacity of 550 mA h g^{-1} was still retained after 5000 cycles with over 90% capacity retention at current density of 6.0 A g^{-1} for the *in-situ* polymerized Si/PANi electrodes. Du *et al.* [93] reported the synthesis of PPy@porous silicon hollow spheres (PPy@PHSi) *via* the magnesiothermic reduction of mesoporous silica hollow nanospheres (MHSiO_2) and *in situ* chemical polymerization of PPy on the PHSi surface (shown in Fig. 9b). The synergetic effect of the porous hollow structure and the surface PPy coating is responsible for the highly electrochemical characteristics. The hollow structure and porous channels in the shell can not only buffer the huge volume change and reduce the diffusion-induced stress, but also facilitate the diffusion of Li^+ and electrolyte into the electrode. The surface PPy coating can significantly enhance the surface electronic conductivity and stabilize the whole structure. As a part of the electrode, polymer

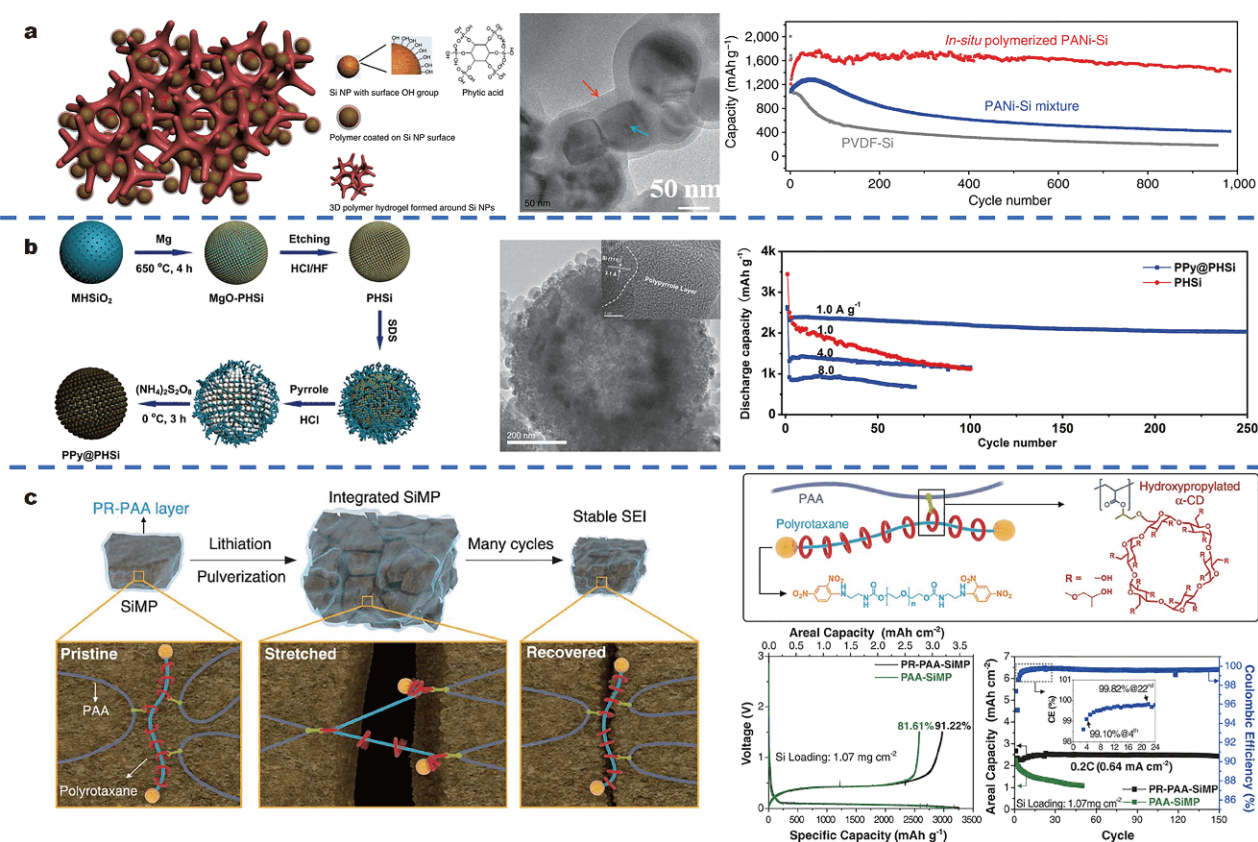


Figure 9 (a) Schematic illustration of the 3D porous PANi-Si electrodes, TEM of the SiNPs coated with PANi polymer layer and cycling performance of the PANi-Si composite electrode and contrast materials at current of 1.0 A g^{-1} . Reproduced with permission from Ref. [92]. Copyright 2013, Nature Publishing Group. (b) Schematic of the fabrication process of PPy@PHSi nanocomposite, TEM and cycling performance of the PHSi and PPy@PHSi nanocomposite at a variety of current densities. Reprinted with permission from Ref. [93]. Copyright 2014, WILEY-VCH Verlag GmbH & Co. (c) Graphical showing of the operation mechanism of PR-PAA during repeated volume changes of SiMPs, together with chemical structures of polyrotaxane and PAA and the corresponding electrochemical performance. Reproduced with permission from Ref. [95]. Copyright 2017, AAAS.

binders act as bonds between current collectors, active substances and conductive agents to ensure electrical contact and integrity of electrodes. Strong supramolecular interactions such as hydrogen bond, ion-dipole interaction and coordination bond play an important role in guiding the structural design of binders. Sodium alginate was firstly used as a binder for silica-based materials [94]. The polar functional groups in sodium alginate molecules could form hydrogen bonds with the silica layer on the silicon surface, which could effectively maintain the bonding between sodium alginate binder and silica materials and the integrity of the electrode structure. In addition, sodium alginate could maintain high strength even in electrolyte. Therefore, it could effectively keep a stable conductive network and buffer the volume change of silicon, thus improving the cycling performance of the electrode.

Recently, Choi *et al.* [95] combined the advantages of soft elastic material polyacrylic acid (PAA) and rigid plastic polyrotane (PR) for highly efficient binders (displayed in Fig. 9c). The soft elastic component can ensure the ionic and electronic conductance of active material and even keep the pulverized silicon coalesced without disintegration, enabling stable cycle life for silicon microparticle in the process of cycling. Noteworthy, the conducting polymers also could be substitution for traditional conductive additive or binder as a conductive binder which reduced the nonactive components [96]. For example, Zeng *et al.* [97] designed and prepared a novel polymer binder possessing high ion and electron conductivities by assembling ion-conductive polyethylene oxide (PEO) and polyethylenimine (PEI) onto the electron-conductive poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT:PSS) chains *via* chemical crosslinking, chemical reduction and electrostatic self-assembly. The obtained polymer binder possessed superior 14 and 90 times higher of lithium-ion and electron transport properties than those of carboxymethyl cellulose (CMC) (with acetylene black) binder system. What's more, the strong crosslinking and electrostatic interactions guaranteed the integrity of the Si anode during volume contraction. Consequently, the Si anode with the final conductive polymer binder showed enhanced delithiation rate capability, cycling stability, reversible capacity, and initial CE.

Apart from the above direct interactions between the silicon and the binder that could enhance the electrochemical performance of silicon anode, an all-integrated electrode was achieved by Liu *et al.* [98] with indirect chemical interlinking between carbon shells of car-

bon@void@silica@silicon (CVSS) and the polymer binder. Based on density functional theory (DFT) calculations, the binding energy between the binder and the carbon layer of CVSS was higher than that of the direct interactions between the silicon and the binder, which led to a better electrode stability and higher electrochemical performance (a reversible capacity of 1640 mA h g⁻¹ after 100 cycles at a current density of 1 A g⁻¹, and a specific capacity of 720 mA h g⁻¹ can be maintained after 1000 cycles at 5 A g⁻¹).

IN SITU TECHNIQUES FOR FAILURE MECHANISM ANALYSIS

Promoting the LIB performance of silicon-based anode requires improved understanding of the internal electrochemical processes and the degradation mechanisms in the electrodes. Thus, various *in-situ* characterization tools have been playing indispensable roles in monitoring the material morphological change, phase evolution, SEI formation and evolution and mechanical property, etc., under real operating conditions. The obtained information is greatly beneficial to the optimization and design of more efficient and stable materials, which in turn improves the overall battery performance. Fundamental understanding and progress revealed by *in-situ* characterization techniques are discussed in the following section.

Atomic force microscopy

The lifetime of silicon-based electrodes is closely related to the formation and evolution of the SEI. These passivation films undergo substantial formation when the silicon electrode particles expand and crack during cycling. Thus, study of the formation mechanisms and structural evolution of SEI layers during cycles is of great importance. *In-situ* AFM has been an invaluable tool to probe the SEI, which can avoid unwanted damage in the sample preparation process such as washing and cleaning. Becker *et al.* [117] analyzed the shape evolution of a-Si pillars with 100 nm height and 100–1000 nm in diameter during cyclings by AFM, as exhibited in Fig. 10a. It was found that the pillars with diameters of 200 nm or larger reached the theoretical volume expansion of ~300% and the volume expansion of the 100 nm pillars was limited. Additionally, the volume effect and interface properties were correlated to the dimension of the materials. Liu *et al.* [118] investigated the morphology and Young's modulus of the individual Si nanowire during SEI evolution and the electrode volume changes during the first discharging/charging cycle *via in-situ* AFM (Fig. 10b).

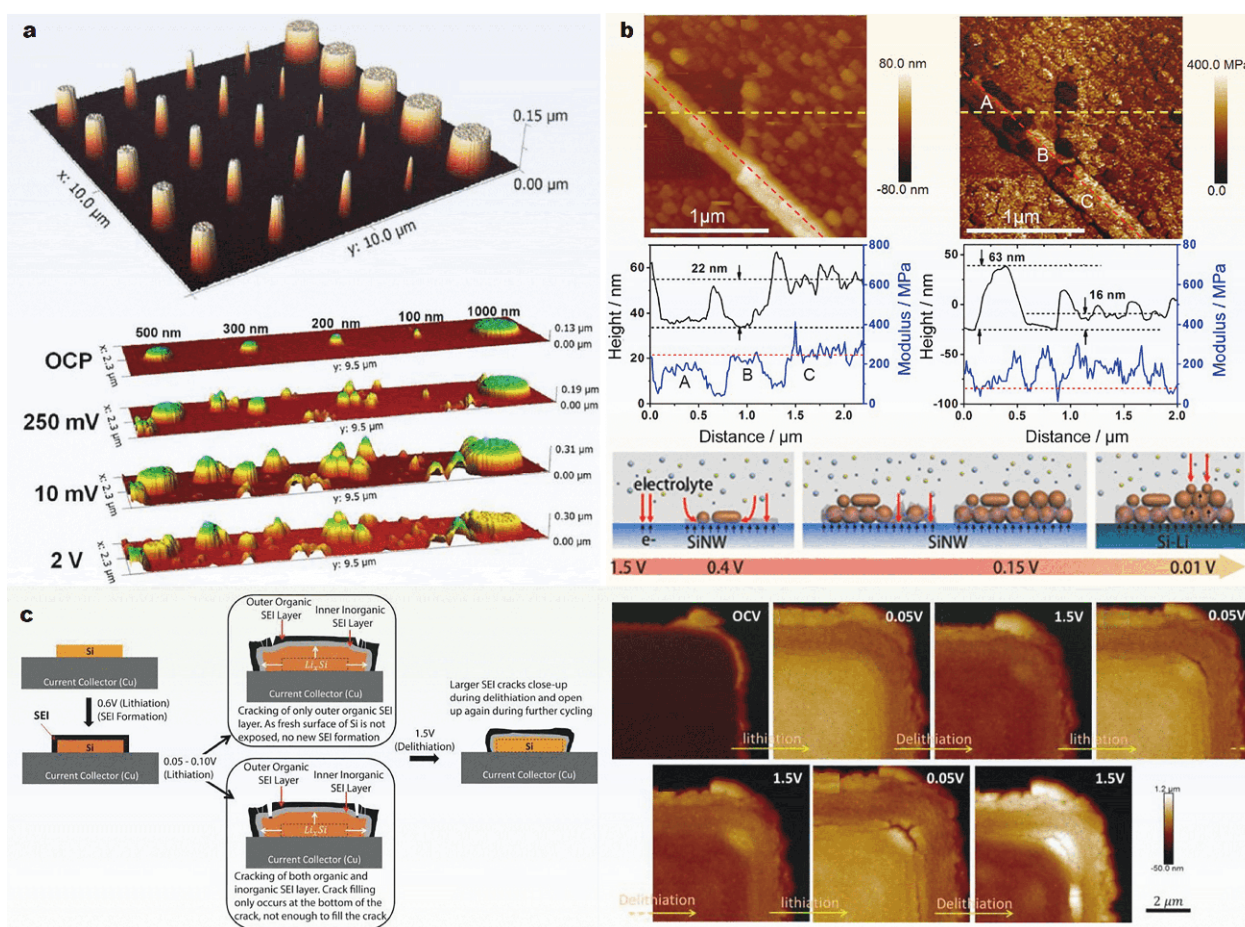


Figure 10 (a) AFM image of the as-fabricated array with a diameter of 1000, 500, 300, 200 and 100 nm pillars and *in situ* 3D AFM images during lithiation and delithiation of a-Si nanopillars at several electrochemical potentials in the first cycle. Reproduced with permission from Ref. [117]. Copyright 2013, American Chemical Society. (b) Images of height and modulus of the Si nanowire anode after being scraped. The line profiles of the topography (black) and modulus (blue) along the Si nanowire (marked with a dashed red line) and along the horizontal cross section (marked with a dashed yellow line). Reproduced with permission from Ref. [118]. Copyright 2014, American Chemical Society. (c) Schematic showing the structure evolution of patterned Si island during cycling and the resulting impact of volume changes on SEI formation and failure, *in-situ* AFM images showing opening and closing of SEI cracks that formed during the initial three cycles. Reproduced with permission from Ref. [119]. Copyright 2016, American Chemical Society.

Three distinct stages of the SEI formation were observed in the discharge process: thin primary film was slowly formed when voltage was above 0.4; thick and particle-like SEI grew rapidly when the voltage was in the 0.4–0.1 V range; the SEI continued growing with the voltage below 0.1 V finally with a thickness of 28 ± 10 nm. The Young's modulus of the Si nanowire interface slightly increased with the growth of SEI. And the final SEI showed a Young's modulus value within a range of 50–400 MPa, consistent with the composition nature of the SEI.

Kumar *et al.* [119] investigated the mechanical degradation of the extremely fragile SEI layer by inducing controlled strains through the unique Peak Force tapping

mode, as shown in Fig. 10c, which directly showed how the volume expansion and contraction of Si electrodes led to significant mechanical damage of the SEI layer. It was found that SEI film in the shear lag region underwent tensile strains during lithiation and the surface cracks did not fill up with new decomposition products in the further cycling.

Transmission electron microscopy

In-situ TEM is another powerful tool to monitor real-time silicon involved electrochemical reactions including phase evolution, anisotropic expansion and fracture behavior, atomic level details of the lithiation of crystalline Si, effects of coatings and mechanical properties of lithi-

ated Si [69,81,120–122].

He *et al.* [123] used *in-situ* TEM to probe the structural and chemical evolution of molecular layer deposition (MLD) alucone-coated silicon upon cyclic lithiation and delithiation. The native oxide layer (~2 nm in thickness) reacted with Li to form crystalline Li_2O (c- Li_2O) upon initial lithiation, which further insulated the particle during subsequent cycles. In comparison, the alucone MLD coated silicon showed no agglomeration tendency and Li_2O formation, and appeared to possess great flexibility, compatible stretching or shrinking ability along

with the expansion or shrinkage of the silicon nanoparticles during cycling (shown in Fig. 11a). Si nanoparticle (NP) confined in CNT was applied as a model to investigate the structural change of the Si NPs and the confinement effect of the CNT during the lithiation and delithiation processes by *in-situ* TEM [124]. It was found that the volume expansion (~180%) of the lithiated Si NPs was restricted by the wall of the CNTs and that the CNT could accommodate this volume expansion without breaking its tubular structure (displayed in Fig. 11b). The Si NP-filled CNTs showed a high reversible lithium sto-

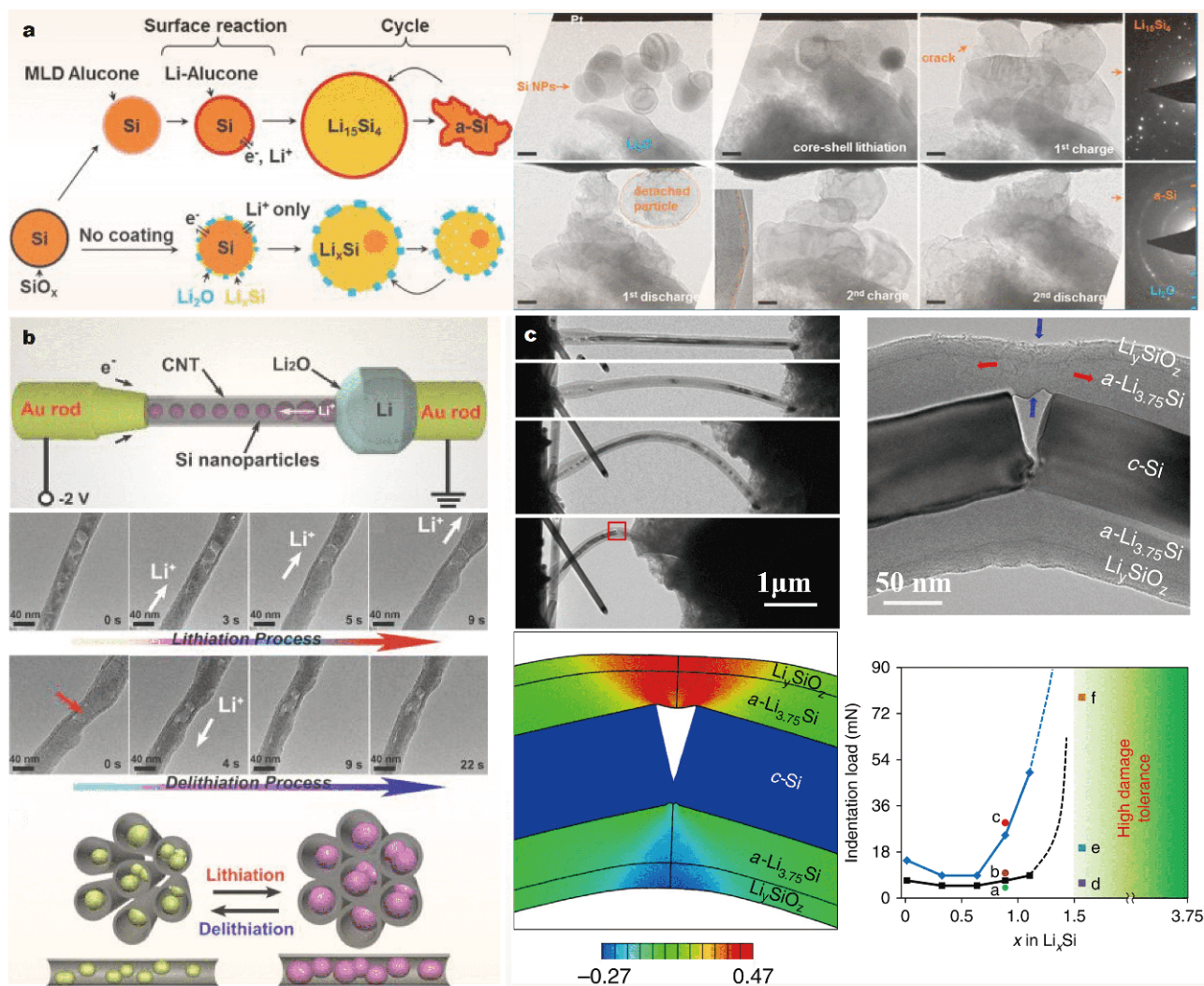


Figure 11 (a) Schematic of surface reactions and cycle behaviors of silicon nanoparticles with different coating conditions and captured *in situ* TEM images of the alucone-coated Si nanoparticles during the lithiation/delithiation behavior. Reproduced with permission from Ref. [123]. Copyright 2014, American Chemical Society. (b) Schematic of the construction of the electrochemical testing setup, *in-situ* captured images of a Si NP-filled CNT structure evolution during lithiation and delithiation. Reproduced with permission from Ref. [124]. Copyright 2015, American Chemical Society. (c) Sequential and zoom-in TEM images of the partially lithiated Si nanowire under axial compression, finite element result showing the simulated elastic-plastic deformation in the nanowire that agrees with the zoom-in TEM image and the indentation loads applied to the lithiated electrodes with different Li contents. Reproduced with permission from Ref. [125]. Copyright 2015, Nature Publishing Group.

rage capacity and desirable high rate capability, because the pulverization and exfoliation of the Si NPs confined in CNTs were efficiently prevented. The fracture resistance is a critical parameter to indicate the structural durability of Si electrode during cycles. Wang *et al.* [125] reported an *in-situ* TEM study on the damage tolerance of electrochemically lithiated silicon (exhibited in Fig. 11c). The results revealed a striking contrast of brittle fracture in pristine silicon *versus* ductile tensile deformation in fully lithiated silicon. The nano indentation testing of amorphous lithiated Si alloys indicated a drastic increase of fracture toughness as the Li to Si ratio was increased to above 1.5.

Other *in-situ* techniques

There are other *in-situ* techniques focusing on studying different aspects of Si-based electrode. For instance, Ogata *et al.* [126] used *operando* ^7Li NMR spectroscopy to study the kinetics of the electrochemical lithiation and delithiation reactions that occurred in nm-sized Si based anodes under realistic cycling conditions (with careful voltage/current controls) over multiple cycles. The ability

to control the voltage and current carefully (Fig. 12c) enabled an understanding of how the charge and discharge process in the NMR experiment were connected, for example, the inhomogeneous growth of crystalline $\text{Li}_{3.75}\text{Si}$ from the amorphous phase, the electrochemical and NMR signatures of the process involving the over-lithiation of $\text{c-Li}_{3.75}\text{Si}$ that occurred below 50 mV, and the formation of small clusters within the $\text{Li}_{3.75}\text{Si}$ phase on charge. These results, in combination with DFT calculations, showed that the phases formed during delithiation strongly depended on the rate of lithiation.

In-situ XRD technique has been applied to study the phase evolution of c-Si electrodes during lithiation/delithiation cycle (Fig. 13a, b). It was found that the formation of metastable $\text{c-Li}_{15}\text{Si}_4$ phase in Si NW electrodes could be avoided through devising several modifications to materials and processing parameters by limiting the cycles to a lower specific capacity cutoff or altering the SiNW growth temperature [127].

Recently, *Operando* Raman and synchrotron XRD were combined to probe the evolution of stress and elastic strain in SiNPs anodes during the first two cycles, under

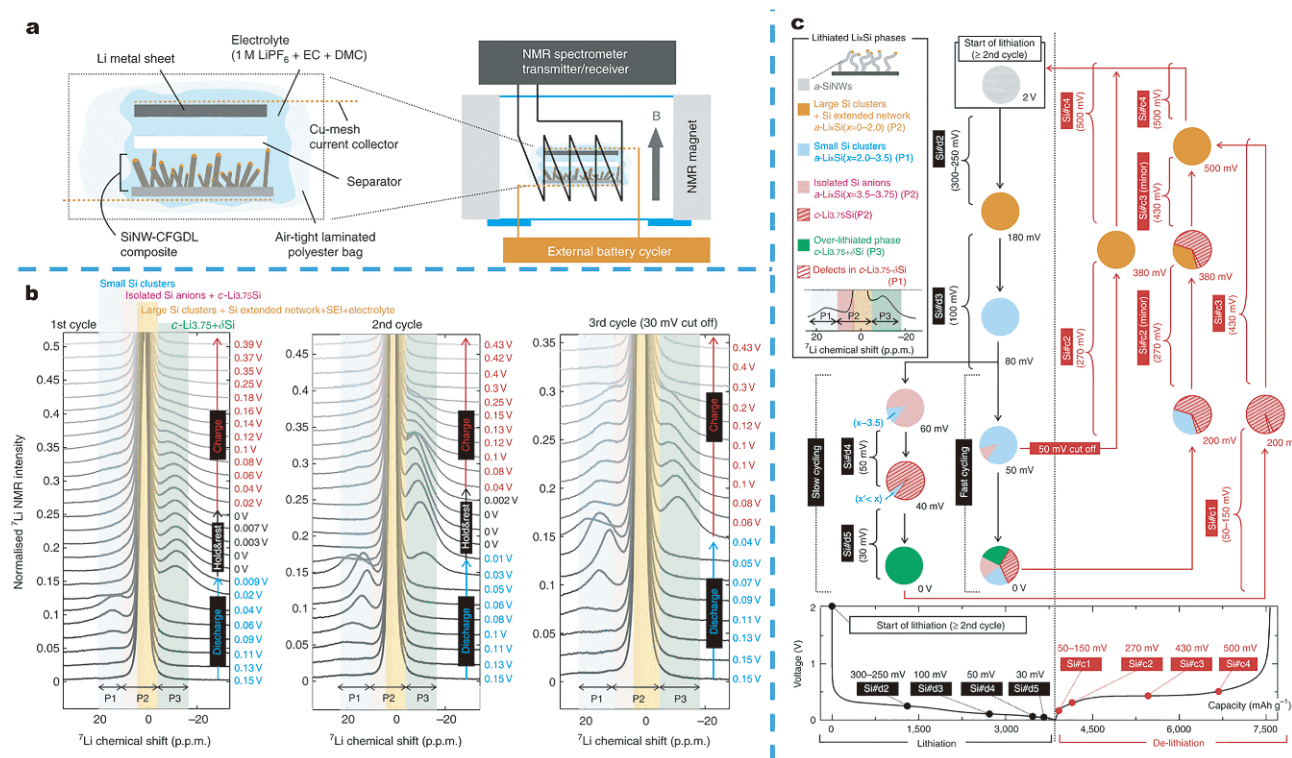


Figure 12 (a) Schematics of the Si nanowire-composite based cell for the *in situ* ^7Li NMR measurements. (b) Enlarged *in-situ* ^7Li NMR spectra of the Si nanowire-CFGDL composite obtained during the potentiostatic experiment in the first three cycles. (c) Phase transformation diagram for the amorphous silicon nanowires on lithiation and delithiation showing the dependence of the phase evolutions on the rate of cycling. Reproduced with permission from Ref. [126]. Copyright 2014, Nature Publishing Group.

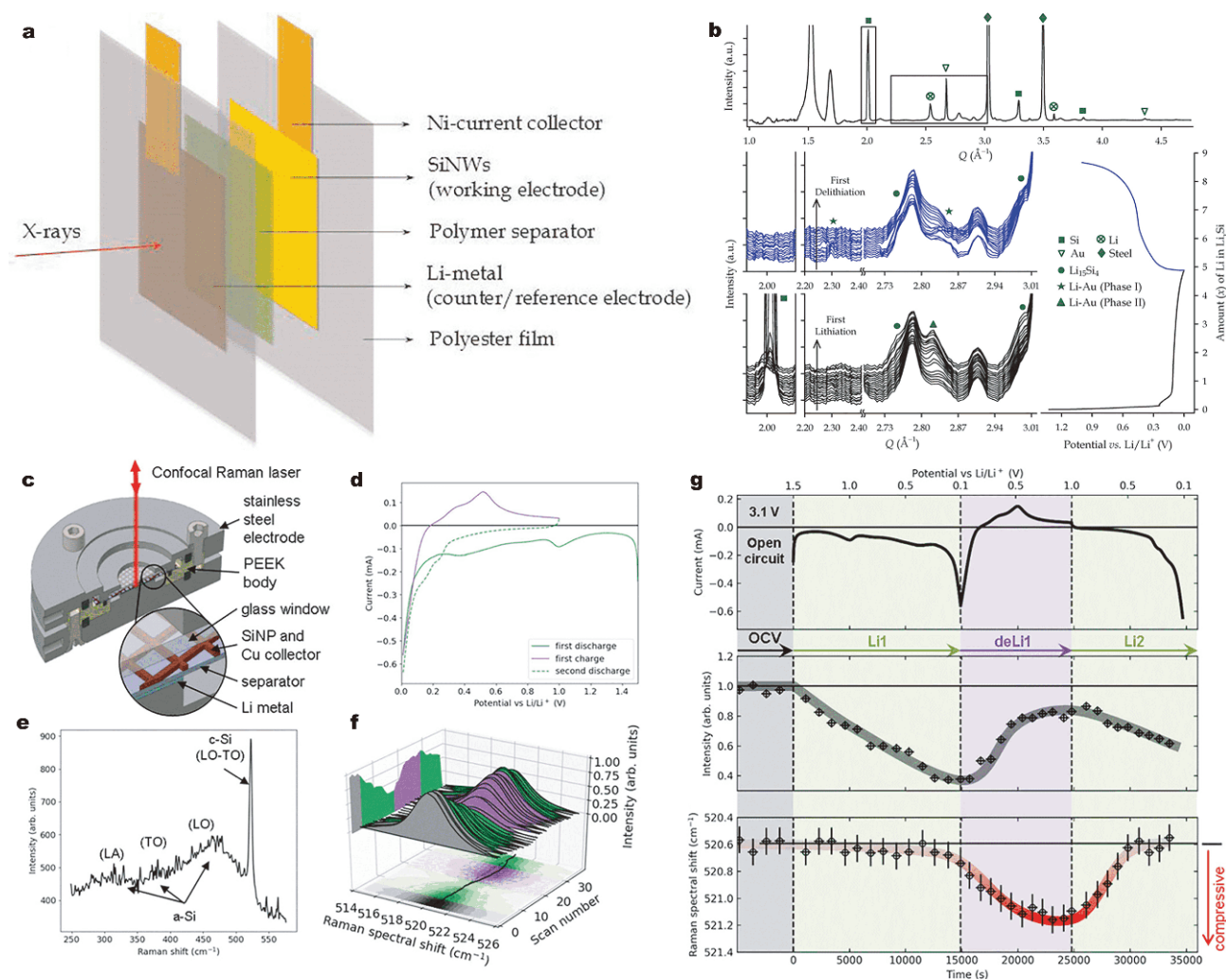


Figure 13 (a) Schematic figure of a cell for *in-situ* XRD test. (b) *In situ* XRD results for a stainless steel-mesh (SS-mesh) cell cycled at 0.2 C. Reproduced with permission from Ref. [127]. Copyright 2012, American Chemical Society. (c) *Operando* Raman setup and electrochemical cell. (d) Linear sweep cyclic voltammograms. (e) Typical Raman spectra including crystalline and amorphous silicon peaks. (f) Evolution of the c-Si transverse optical-longitudinal optical (TO-LO) peak as a function of the scan number during the open-circuit, discharge (green) and charge (purple) cycles. The shadow on the left panel shows the variation of the Raman peak intensity over cycling time and the intensity map on the lower panel indicates the shift of the peak position (highlighted by the black line). (g) Evolution of the current, Raman peak intensity and spectral position as a function of time and applied voltage. Reproduced with permission from Ref. [128]. Copyright 2017, American Chemical Society.

limited capacity cycling conditions [128]. Fig. 13g presents the detailed evolution of the current, Raman intensity and position with the applied voltage, which proved the crystalline SiNPs experienced little to no stress during the first lithiation and were compressed during the following delithiation. Moreover, by analyzing the variations of the Si XRD and Raman peaks, the pressure exerted from amorphized shell onto the continuously shrinking crystalline core was responsible for electrochemically driven variations of the internal stress. Lithiation/delithiation of the silicon under limited capacity conditions triggered the formation of “crystalline core-

amorphous shell” particles, which evidenced as a stepwise decrease in core size, as well as sequences of compressive/tensile strain due to the stress applied by the shell (Fig. 13c-f).

These conclusions from various *in-situ* technologies are much needed for the benchmarking of theoretical models and for the further rational design of SiNP-based electrodes.

THE GAP BETWEEN CURRENT STUDY AND REAL APPLICATION

After years of research and development, although great

improvements of the silicon-based anodes have been achieved in lab scale, the wide application of the Si-based anode in real batteries has not yet popularized. There are only a few reports about the commercialization of the Si-based LIB anodes. BTR New Energy Materials Inc. developed high capacity ($600\text{--}650\text{ mA h g}^{-1}$) Si based composite material with an initial CE of 89%–90%. Nexxon designed silicon based composite powder for hybrid or low loading anode electrodes (recommended use up to 10 wt.% loading) with capacities up to $400\text{--}450\text{ mA h g}^{-1}$. Further industrialization should focus on the energy density of full cell in practical application instead of half coin cells, and several critical concerns should be settled.

Firstly, the silicon volume expansion will become more serious when used in practical application that usually needs a higher current density. Mass loading of the Si-based electrode is a key issue to realize the decent energy output in real battery applications. Unfortunately, the real mass loading density is not high enough. Secondly, the CE is also crucial for practical battery applications. Normally, the initial CE of the Si-based anode is typically in the range of 50%–90% (Table 1), which still cannot meet the need for practical battery applications. Thirdly, the consistency and cost of the silicon anode materials are far from meeting scale industrial application requirements. The low yield and complicated fabrication process or harsh synthesis condition lead to the high price of silicon nanomaterials. Moreover, pairing with high capacity Ni-rich or Li-rich cathode is preferred to realize the commercialization of silicon-based anodes. However, the internal failure mechanism is more complex than that of half cells.

CONCLUSIONS AND PERSPECTIVES

Silicon is one of the most promising anode materials for next generation lithium batteries owing to its ultrahigh capacity, proper working potential, low-cost and high abundance. However, the inherent large volume expansion, unstable SEI layer and poor conductivity of Si have hindered its further practical application. In this review, we discussed and summarized the rational material design principles, the recent advances in design and synthesis of silicon anode structures and *in-situ* characterization techniques that facilitate the deep understanding of lithiation process and capacity degradation mechanism of silicon anode in LIBs.

Over the past decades, the above issues have been addressed through various methods. Especially, structural design is believed to be the most effective solution and it

can be summarized as follow: construction of silicon nanostructures (downsizing the Si to nanoscale with various morphologies, dimensions and porous structures), fabrication of silicon composite (compositing with carbon, metal, metal oxide and conductive polymer). Early studies have shown that nanosized silicon can effectively inhibit the volume expansion and pulverization phenomenon. However, the larger specific surface area results in much more formation of SEI and consumption of Li ion, which further leads to low CE. Compositing silicon with carbon, the conductivity of silicon-based materials can be effectively improved, which can further improve the rate performance. On the other hand, the introduced carbon materials can buffer the volume change effect of silicon and avoid direct contact between silicon and electrolyte, prevent uncontrolled SEI film formation at once, which significantly improve the electrochemical properties of silicon-based materials. However, the tap density and mass loading of silicon are still far from satisfaction to achieve higher energy density. Silicon/metal and conductive polymer composite almost share a similar working concept with silicon/carbon, while these materials are hindered by the high cost, limited resource and complicated preparation process for large scale application.

For now, coating or modifying Si with carbon has been highlighted and demonstrated the most common and effective way. Meantime, the Si/C has been partially commercialized. However, real breakthroughs are still in urgent need to come. Firstly, improving the tap density and mass loading of silicon to ensure the volume and specific energy density is the most important issue to meet the commercial need. Secondly, the synthesis of nanomaterials is in high cost, which is of key importance for widespread commercial applications. Thirdly, it is hard to ensure the consistency of materials when it comes to industrial production. So, it is crucial to explore facile and environmentally friendly routes to obtain low-cost, large-scale and high-quality silicon/carbon anode.

We have witnessed the fast development of several *in situ* electrochemistry techniques in the recent years, such as *in situ* TEM, SEM, AFM, XRD and NMR, for the evaluation of real-time energy storage behaviors of silicon-based batteries. The new insights contribute to a better understanding of device failure and monitoring on the electrode morphological change, phase evolution, SEI formation and evolution and mechanical property, guiding future material optimization and innovation. However, the present *in situ* cells are incapable of completely mimicking real battery-operation conditions, and

Table 1 Summary of synthesis method and electrochemical characteristics of selected Si-based anode nanostructure

Active materials	Synthesis method	Mass loading (mg cm ⁻²)	First cycle capacity (mA h g ⁻¹) (A g ⁻¹)	Initial Coulombic efficiency	No. cycles with 80% capacity retention
Mic-Si/graphene cage [81]	Ultrasonication	0.8	3300 (0.05)	93.2%	No decrease after 100 cycles
Hierarchically porous Si [78]	Dispersed in ethanol and cast	1	1850 (0.42)	87%	60
Si NPs [98]	Stirring for 6 h at 200°C	2.0–2.5	2500 (0.05)	85%	15
Si NPs [92]	Solution processes	0.3–0.4	1200 (1.0)	70%	180
SiMP [66]	Condensation reaction	0.5–0.7	3200 (0.5)	80%	80
Si/C [99]	CVD	1.8	2277 (0.42)	90%	16
p-Si/C [77]	Magnesiothermic reduction	1.1	2500 (2.6)	75%	10
Si nanorods [101]	Self-templating synthesis	1.1	2705 (0.25)	60.5%	No decrease after 500 cycles
PG-Si [102]	Chemical reduction or decompose	1	1464 (0.2)	58%	70
Si/C [103]	CVD	1–5	3207 (0.05)	79%	No decrease after 100 cycles
Mesoporous Si [104]	Pyrolysis	0.5	2660 (0.5)	72.8%	175
SiO/G/C [105]	Heat treatment	2	905 (1.0)	68.1%	No decrease after 200 cycles
SiO ₂ nanowires [106]	Pyrolysis	1.0	2215 (0.5)	88%	10
Si@C [107]	Pyrolysis	0.5	2936 (0.09)	85%	10
MHSiO ₂ @C [108]	Pyrolysis	1.73–1.91	880 (0.5)	68%	5
Si@TiO ₂ [109]	Sonication and pyrolysis	0.8	1562 (2.1)	65.8%	68
HSi@C [110]	Reduce HSiO ₂	0.7	1970 (2.0)	52.4	No decrease after 200 cycles
CNT-Si [111]	Etching by HF	1.3	2213 (0.42)	83.4%	50
Mesoporous Si [112]	Pyrolysis	0.7–1.2	2789 (0.36)	64.1%	70
Ferrosilicon [59]	High energy ball milling	0.5	1250 (0.4)	88%	No decrease after 100 cycles
Si/C [113]	Thermal decomposition	0.6–1	1992 (0.2)	60%	110
Si-SHP [114]	Ultrasonication	1.13	2850 (0.1)	80%	25
Si@GO [115]	3D tomography	1.2	2900 (0.42)	94%	70
Si/graphene [116]	Mechanical stirring	2	879 (0.05)	58.6%	6

detailed information about the electrochemical mechanisms, ion and electron kinetic transport at the electrode/electrolyte interface are still absent due to their complexity. On the other hand, one *in situ* characterization technique can obtain only single information, which may give rise to inconsistency of the observed information and lead to further disagreements on the conclusions. Thus, it is highly desirable to develop integrated technique that can get all the information at the same time so that a more complete picture can be drawn than one technique acting alone. This requires a fast data-collection rate, which is not possible using most of the present tools and expected to motivate future studies.

Despite some great achievements for enhancing battery performance, other challenges of Si anode such as the scalable synthesis and the compatibility to combine with novel components still remain for the commercialization of Si-based battery. Exploration of other interrelated

strategies, such as electrolytes, electrolyte additives, new current collector and binders, will also help to improve the electrochemical performance of lithium-based rechargeable batteries using nanostructured silicon electrode materials. With the fast development and extensive research interest of science and technology in this area, the obstacles that hinder the practical applications of Si anode material will be addressed successfully in the near future.

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锂离子电池中硅负极材料的设计及机理分析研究进展

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摘要 硅基负极具有理论容量高、工作电位低、环境友好、丰度高等优点, 被认为是下一代锂离子电池中最有希望替代石墨负极的材料之一。然而, 硅负极在脱嵌锂过程中体积膨胀大、界面反应剧烈是制约其进一步实际应用的关键问题。合理设计硅纳米结构有助于解决这些问题。本文首先介绍了大容量锂电池用纳米硅负极的基本科学问题, 然后重点介绍了其设计、制备、原位表征和失效机理等方面的最新进展, 总结到目前为止关键的经验以促进硅负极在实际锂电池中的应用。最后展望了硅负极未来的发展和挑战。