Atomistic Simulation Protocol for Improved Design of Si–O–C Hybrid Nanostructures as Li-Ion Battery Anodes: ReaxFF Reactive Force Field

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Supporting Information

ABSTRACT: Recently, silicon–oxygen–carbon hybrid nanostructures have received great attention as a promising anode material for Li-ion batteries, for which their diverse structures can be synthesized. Here, using molecular dynamics (MD) simulations with a reactive force field (ReaxFF), we studied the atomistic lithiation behaviors of sp² carbon-coated Si and SiOₓ nanostructures, such as nanowires (NWs) and nanoparticles (NPs), in which various kinds and sizes of carbonaceous coating layers were explored. The introduction of an sp² carbonaceous coating layer to Si-based anodes makes Li diffusion more facile, which leads to improved battery performances such as faster charge/discharge rates. Moreover, the carbonaceous coating layer can also provide a buffer effect to volume changes during lithiation along with the well-known functions of preventing the loss of electrical continuity and increasing electrical conductivity of Si-based anodes. However, a thick carbonaceous coating layer can strongly suppress the volume expansion behavior of Si-based nanostructures and thus prevent Li penetration into the nanostructures, leading to a very low Li capacity. According to our ReaxFF-MD simulations, the critical size of the carbonaceous coating layer that can act as a buffer layer is approximately C/Si = 2.4, which is the circumference ratio of the carbonaceous coating layer over the Si NWs. For a coating layer that has a higher ratio, Li cannot penetrate into the Si NWs; instead, they exist only on and in the sp² coating layers including in the spaces between two graphene layers. Moreover, the shape of the Si nanostructures (e.g., NW and NP) does little to affect the anode properties, such as Li capacity and volume change, although Si NP confined in a carbon nanotube shows anisotropic volume expansion behavior during lithiation. We expect that the ReaxFF will provide a useful protocol for designing Si–O–C hybrid anodes to obtain better performing Li-ion batteries.

1. INTRODUCTION

Silicon (Si) is one of the most promising and widely investigated candidates as a next generation anode material for Li-ion batteries (LIBs) because of its high Li capacity and abundance.1–4 However, the large volume fluctuation during the lithiation/delithiation process leads to fracturing of the material followed by rapid capacity fading due to the loss of electrical continuity.5,6 For strategies to overcome the drawbacks of Si, several approaches, such as nanosized structures,7–13 silicon suboxides (SiOₓ),14–32 and silicon–carbon composites,33–36 have been investigated. In particular, a combination of Si with carbonaceous materials such as graphite,33 graphene,35,36 and carbon nanotube (CNT),37 has improved the cyclic stability in which carbon provides an enhanced buffering effect and electrical conductivity. Moreover, Si@SiOₓ@C nanocomposites with a core–shell structure have demonstrated significantly improved battery performances, such as high reversible capacity, excellent cyclability, and high rate capability.37,38 Thus, the development of novel Si–O–C hybrid nanostructures is of great significance for LIBs.

In developing such anode materials, understanding of the atomistic lithiation mechanism using computer simulations can provide remarkably valuable information. Of the various computational methods, a reactive force field (ReaxFF) can act as a sweet spot for this type of study because chemical reactions between atoms in real nanosized systems can be predicted with reasonable accuracy.39–41 Indeed, we previously clarified atomistic lithiation mechanisms of pristine Si and SiOₓ nanowires (NWs) with diameters of approximately 5 nm and lengths of approximately 10 nm using molecular dynamics simulations.
In particular, a ReaxFF-MD simulation study showed that the anisotropic volume expansion behavior of Si NWs during lithiation relies on the surface structures of Si NWs, which was supported by an experiment. However, the volumes of fully lithiated Si NWs are almost identical irrespective of the surface structures. Moreover, in the ReaxFF-MD simulation, we observed the transient formation of silicene-like structures in the Si NWs. Independently, several research groups also reported ReaxFF studies on Si-based anodes for LIBs.

In this regard, ReaxFF is definitely useful in predicting anode properties (Li capacity, volume change, Li diffusion, etc.) of the Si−O−C hybrid nanostructures and for unraveling their lithiation mechanisms. In this work, using the ReaxFF method that we recently developed via first-principles calculations, we report the atomistic lithiation behaviors of carbon-coated Si and SiO$_x$ nanostructures, such as NWs and nanoparticles (NPs), in which various kinds and sizes of the carbon coating layers were considered.

2. COMPUTATIONAL DETAILS

To simulate the lithiation behavior of the Si−O−C hybrid nanostructures (NW and NPs) at the atomic level, we used an MD simulation employing ReaxFF. Very recently, we developed ReaxFF for the Si−Li to Si−Li−O systems, with which we successfully simulated the formation of the solid-electrolyte interphase on Si-based anodes. In the work, while developing the ReaxFF parameters for Si−C and C−Li pairs on the basis of our previous Si−Li−O ReaxFF, we considered various relevant trainsets: equation of states (EOSs) of various Si−C crystals, bond dissociations of C−Si single and double bonds in relevant clusters (H$_2$C−SiH$_3$ and H$_2$C=SiH$_2$), bond dissociations of the C−Li σ bond in C$_6$H$_5$Li, and dispersion interactions between Li and graphene. Thus, the ReaxFF for the Si−Li−O−C−H−F system is applicable for lithiation in the Si−O−C hybrid nanostructures studied in this work.

The MD simulations were performed using LAMMPS software on the iBat simulation platform (http://battery.vfab.org) for LIBs, which was recently developed by our research group. To integrate Newton’s equations of motion during the MD simulations, a Verlet algorithm was used with a time step of 0.5 fs (fs). All calculations were performed in a canonical NVT ensemble at 300 K, in which the temperature was maintained using a Nosé−Hoover thermostat with a damping parameter of 0.01 fs$^{-1}$.

As the Si−O−C hybrid anodes in this work, two nanostructure-type (NWs and NPs) Si-based anodes (Si and SiO$_x$) were considered along with CNT or graphene-coated layers. Here, the Si NWs have a diameter of 3 nm and a length of 10 nm, and the NPs have a diameter of 3 nm. For the SiO$_x$ NWs and NPs, we obtained the SiO$_x$ materials through chemical reactions between O$_2$ molecules and pristine Si materials that were simulated using the ReaxFF-MD simulations, which are similar to our previous simulation. The obtained SiO$_x$ has a composition of SiO$_{0.43}$ with a Si@SiO$_2$ core@shell structure.

For the lithiation simulation of the Si−O−C hybrid nanostructures, a method was applied that is similar to that used in our previous works to investigate the lithiation of Si and SiO$_{0.43}$ NWs. The Si−O−C nanostructures were immersed in a simulation cell with dimensions of 100 Å × 100 Å × 97.8 Å that was filled with Li atoms, and any Li atoms that overlapped the nanostructures were removed, as represented in Figure S1 of the Supporting Information (SI). The total number of atoms in the system was approximately 45 000. Additionally, periodic boundary conditions in the x-, y-, and z-directions were applied during the MD simulations, in which the NW axis is in the z-direction, leading to lithiation of the NWs along the x- and y-directions, although lithiation of the NPs allows all directions.

For penetration of a hexagon ring of carbon in CNTs or graphene, an extremely high energy barrier (10.2 eV) is required, which hardly allows Li penetration through the hexagon ring. However, the penetration can be facilitated by carbon vacancies. According to our density functional theory (DFT) calculations (Figure S2), the mono-, di-, tetra-, and hexavacancies significantly lower the penetration barrier to 2.30, 3.01, 0.02, and 0.01 eV, respectively. Here, the energy barriers for mono- and divacancies are similar to the reported DFT results. Although the ReaxFF slightly overestimates the Li penetration barriers in comparison with the DFT-D2, it reasonably predicts the tendency. Therefore, in this work, we introduced hexavacancies in the CNT or graphene coating layers to induce effective penetration of the coating layers. In addition, in building the Si or SiO$_{0.43}$ materials in CNTs, the optimum distances between the CNT and the Si-based material were determined by the ReaxFF, as represented in Figure S3.

3. RESULTS AND DISCUSSION

3.1. Prediction of the Lithiation Behaviors of Various Si−O−C Hybrid Nanostructures.

The lithiation behaviors of various Si−O−C NWs that were obtained from the ReaxFF-MD simulations are shown in Figure 1, along with their Li capacities and volume changes. Here, the Si and SiO$_{0.43}$ NWs
with CNT and graphene coating layers are considered, and the behavior of a pristine Si NW with no carbonaceous layer is also presented for comparison. The carbonaceous coating layers include carbon vacancies of 5% for efficient layer penetration of Li. Additionally, for the graphene layers, we considered two different sizes: One has exposed Si atoms that are not completely covered by the graphene layer (Figure 1d), and the second has a Li diffusion channel between two graphene layers (Figure 1e), although all Si atoms are covered by the graphene layer. The carbon edges of the graphene covalently bond to Si, as observed in a prior experiment.6 Here, the Si–O–C hybrid nanocomposites considered in this work could indeed be synthesized.27,35,36

After 1000 picoseconds (ps) of lithiation, the pristine Si NWs had a volume expansion of 252% with a Li capacity of 1760 mAh/g (Li1.38Si). However, the Si NWs with the CNT coating (SiNW@CNT) had a volume expansion of 111% and a Li capacity of 1374 mAh/g (Li1.26Si), and the SiO0.43 NWs with the CNT coating (SiO0.43NW@CNT) had a volume expansion of 105% and a Li capacity of 1345 mAh/g (Li1.88Si). In Figure 1h, it is apparent that both the Si and SiO0.43 NWs coated with CNTs exhibit a smaller volume expansion behavior than that of pristine Si NWs at a given Li concentration, which clearly shows that the CNT coating layer provides a buffer effect on volume expansion of the Si or SiO0.43 NWs during lithiation.

For Si NWs coated with graphene (SiNW@graphene), we explored the effects of the graphene size on the lithiation behavior in which the two circumference ratios of graphene over the Si NW (C/Si = 0.94 and 2.09) were considered (Figure 1d,e). The SiNW@graphene composite with a ratio of C/Si = 0.94 shows a similar trend (volume expansion of 233% and a Li capacity of 1694 mAh/g (Li2.28Si) at 1000 ps) as the Si NW with no graphene layer, although the graphene provides an effect as a buffer layer on volume expansion during lithiation in the early stage of lithiation (Figure 1h). However, the Si NW@graphene with C/Si = 2.09 shows a smaller amount of lithiation, followed by a smaller volume expansion (volume expansion of 62% for a Li capacity of 1147 mAh/g (Li1.84Si) after 1000 ps of lithiation). From these results, the thick carbonaceous coating prevents the efficient lithiation of Si, implying that an appropriate thickness or size of the carbonaceous layer for a given Si nanostructure would exist, which plays a role in the buffer layer for volume expansion and does not simultaneously prevent the lithiation behavior.

In Figure 1h, compared to the pristine Si NW, the SiNW@CNT shows a smaller volume change at a given Li capacity than the pristine Si NW in the early stage of lithiation, although the SiNW@CNT and pristine Si NWs eventually show a similar Li capacity after lithiation of 1000 ps. This indicates that the early stage of lithiation occurs mainly on the CNT layer rather than in the Si bulk. The SiNW@graphene shows a trend similar to that of the SiNW@CNT. However, the SiNW@graphene with a C/Si = 2.09 shows a longer incubation time for the volume expansion and a smaller Li capacity than the composite with C/Si = 0.94. Because the SiNW@graphene with C/Si = 2.09 is a longer graphene layer, it can store relatively more Li on and in the graphene layer than the composite with C/Si = 0.94 and, in particular, between two graphene sheets. Moreover, after lithiation of 1000 ps, the SiO0.43NW@CNT shows a smaller volume expansion than the SiNW@CNT because the surface silicon oxide layer can act as a buffer for the volume expansion on the lithiation along with the CNT layer. However, in the early lithiation, the SiO0.43NW@CNT has a higher volume expansion because Li preferentially interacts with SiO0.43 NW rather than with the CNT layer, in contrast to the case of SiNW@CNT. The detailed lithiation mechanism will be discussed in Section 3.3.

In addition, we investigated the effects of the vacancy concentrations in the CNT layer on the lithiation behaviors of the Si nanostructures. As shown in Figure S4, a higher vacancy concentration leads to a faster lithiation; however, the slopes for volume change versus Li concentration are almost similar, irrespective of the vacancy concentration.

### 3.2. Effects of Sizes of the Carbonaceous Layers on the Lithiation Behaviors

Intrinsically, Si anodes own the volume expansion characteristics during the interaction with Li, which reveals that a strong suppression of the carbonaceous layers for the volume expansion leads to a low lithiation capacity. Therefore, it is important to find an appropriate size for the layer to buffer the significant volume change of the Si anodes without leading to a serious sacrifice of the Li capacity.

Figure 2 shows the lithiation behaviors of SiNW@graphene composites, where various sizes of graphene layers (i.e., C/Si ratio = 0.94, 1.10, 1.35, 1.74, and 2.09) were considered. The Li anodes without leading to a serious sacrifice of the Li capacity.
The Journal of Physical Chemistry C

(\text{Li}_{x,y}\text{Si}) for 2.09, and their volume expansion is 233% for C/Si = 0.94, 201% for 1.10, 128% for 1.35, 88% for 1.74, and 62% for 2.09. Using these values, we also investigated the correlations for the volume change versus the graphene size (C/Si ratio), which is shown in Figure 2g. Using the extrapolation of the correlation, we found that when the C/Si ratio is 2.4 the volume change is zero. Thus, for a larger C/Si ratio, Li atoms cannot penetrate into the Si NW. Instead, they exist only on and in graphene layers including the spaces between two graphene layers.

To further explore the constraining effect of the graphene layer, we calculated the von Mises stress distributions of the SiNW\@graphene composites in Figure 3, where atomic structures at the same Li capacity (600 mAh/g) were considered. The von Mises stress implies one that is required to start yielding or deformation of a material. Additionally, the von Mises stress \( (\sigma_v) \) is calculated according to eq 1:

\[
\sigma_v = \frac{1}{2} \left( \sigma_{xx} + \sigma_{yy} + \sigma_{zz} + 2 \sqrt{ \sigma_{xx}^2 + \sigma_{yy}^2 + \sigma_{zz}^2 - 2 \sigma_{xx} \sigma_{yy} - 2 \sigma_{yy} \sigma_{zz} - 2 \sigma_{zz} \sigma_{xx} + 2 \sigma_{xy} \sigma_{yx} + 2 \sigma_{yz} \sigma_{zy} + 2 \sigma_{zx} \sigma_{xz} } \right)
\]

(1)

where the \( \sigma_{xx}, \sigma_{yy}, \text{ and } \sigma_{zz} \) are the normal stresses, and the \( \sigma_{xy}, \sigma_{yx}, \text{ and } \sigma_{zx} \) are the shear stresses in an \( x-y-z \) coordinate system. Here, the calculated von Mises stress values are similar to the reported experimental lithiation-induced stress values in nanostructured Si anodes by \textit{in situ} measurements.\(^{50,61}\)

For the pristine Si NW, the Si surface exhibited smaller von Mises stress values than the interior, which indicates that the Si surface is easier to deform due to lithiation. However, for the SiNW\@graphene composites, the Si surface near the graphene coating layer has higher von Mises stress values than the core region of the Si NWs. This indicates that the graphene coating leads to a constraint effect for the deformation of the Si surface region, which prevents Li penetration into the Si NW.

We additionally calculated the residual stress distributions of the graphene-coated Si NW (SiNW\@graphene) in the radial direction \( (\sigma_r) \) and then compared the results with those for the pristine Si NW, as shown in Figure S22. Here, the residual stress was obtained by averaging the atomic stresses of Si and Li atoms in the concentric shell between \( r \) and \( r + \delta r \), where \( r \) is the radial distance from the center of the NW. The radial stress \( (\sigma_r) \) is calculated using eq 2.\(^{33}\)

\[
\sigma_r = \sigma_{xx} \cos^2 \theta + 2 \sigma_{xy} \sin \theta \cos \theta + \sigma_{yy} \sin^2 \theta
\]

(2)

Here, the \( \sigma_{xx} \) and \( \sigma_{yy} \) are the normal and shear stresses in an \( x-y-z \) coordinate system, respectively, and the \( \theta \) is an angle.

During the lithiation of a pristine Si NW, the lithiated phase (Li\(_x\)Si) has tensile (positive) stresses, which serve as a driving force for the volume expansion of the NW.\(^{53}\) On the other hand, in the initial SiNW\@graphene composite, the Si core is subjected to compressive stresses that are induced by the tensile stresses of the Si–graphene interface. Although the lithiation proceeds, the interface has still tensile stresses and acts as a buffer layer alleviating abrupt changes in stresses. This induces the lithiated Si phase (Li\(_y\)Si) to be subjected to compressive stresses, which is in contrast to the pristine Si NW case. Such compressive stresses exerted on the Li\(_y\)Si phase induce a retardation effect on the volume expansion of the NW during lithiation. Moreover, the interface also induces compressive stresses to the lithiated graphene layer that is an outmost layer of the NW.

To confirm the constraint effect of carbonaceous coating layers, we performed nudged-elastic band calculations using DFT within the PBE-D\(^2\)\(^{62}\) framework to investigate the effects of the carbonaceous layer on the energy barriers for penetration of a Li atom into the Si surface (Figure 4). For a Si(111) surface with no graphene coating layer, the energy barrier is 1.00 eV, which is similar to the previous DFT calculation.\(^{53}\) However, the graphene coating increases the energy barrier due to the constraint effect of the coating layers. Indeed, for Si(111) surfaces with two graphene layers, the energy barrier is increased by 1.34 eV, which clearly shows the constraint effect of the graphene layers for Li penetration. According to the analysis (Table S1) of the DFT stress tensor components for the initial structures of Figure 4, \( \sigma_{xx} \) in the pristine Si is 0.57 kbar (tensile stress), while \( \sigma_{zz} \) in the two-layer graphene-coated Si is \(-5.04\) kbar (compressive stress). Here, we focus on the stress tensor component along the \( z \)-direction because the Li insertion into the Si bulk proceeds along the direction. Indeed, the carbon coating layer can induce a compressive strain effect, which leads to a higher Li penetration barrier. In addition, we need to mention that the current ReaxFF provides the Li penetration barrier similar to that of the DFT (DFT 1.34 eV vs ReaxFF 1.31 eV).

To validate prediction of atomic charges by the ReaxFF, we compare the ReaxFF values with the DFT ones, as shown in Figure S23a. For the structure shown in Figure 4b, the ReaxFF predicts very similar atomic charges for Si and C atoms to those of DFT, although the ReaxFF tends to underestimate atomic
charges of Li atoms. Therefore, the present ReaxFF can reasonably predict atomic charges in various carbon-coated Si hybrid nanostructures. We have also investigated effects of sizes of the carbon coating layers on atomic charge distribution in graphene-coated Si NWs in Figure S23b–f. Our ReaxFF simulations reveal that the effect of the graphene coating size is not significant, which implies that the NWs have similar driving forces for the lithiation process.

Thus far, we have focused on the lithiation behavior of Si or SiO\textsubscript{0.43} NWs with carbonaceous coating layers. As shown in Figure 5, we additionally investigated the lithiation behaviors of spherical Si NPs in CNTs, which can be experimentally synthesized.\textsuperscript{27} Spherical Si NPs without the CNT coating show uniform volume expansion during lithiation. However, the CNT coating leads to anisotropic volume expansion of the Si NP in which the volume expansion behavior along the axis direction of the CNT is more feasible because the Si NP is under constraint along the radial direction by the CNT layer. Moreover, we observed that the shape of Si nanostructures (e.g., NW and NP) does little to affect the anode properties, such as Li capacity and volume change (Figure 5d).

It was reported that an anisotropic volume expansion during the lithiation of Si NWs occurred due to the different levels of expansion along different crystallographic orientations.\textsuperscript{64} Here, largely incompatible strains are generated at the intersections of adjacent crystalline (110) facets near the outer surface, leading to surface crack nucleation at well-defined angular sites. Although we have observed such an anisotropic volume expansion in the CNT-coated Si NP, the surface crack nucleation induced by the phenomenon has not been found, which is probably due to the time limit of MD simulations.

**3.3. Lithiation Mechanisms of Si and SiO\textsubscript{0.43} Nanostructures Coated with Carbonaceous Layers.** To clarify the lithiation mechanism of Si or SiO\textsubscript{0.43} NWs coated with carbonaceous layers, we scrutinized the MD trajectories for Si and SiO\textsubscript{0.43} NWs coated with CNT layers, which is shown in Figure 6. For pristine Si NWs, Li atoms interact with Si surfaces, and amorphous Li\textsubscript{4}Si phases are generated in which lithiation proceeds uniformly (Figure 6a). Detailed information on the lithiation mechanism can be found in our previous work.\textsuperscript{43} However, in the SiNW@CNT composite (Figure 6b), Li atoms penetrating through carbon vacancies in the CNT layer tend to preferentially diffuse at the interface between the CNT layer and Si surface rather than penetrate into the Si surface. However, for the SiO\textsubscript{0.43}NW@CNT composite (Figure 6c), the Li atoms penetrating through the defects in the CNT interact with oxygen atoms in SiO\textsubscript{0.43}, leading to the evolution of Li\textsubscript{2}O and Li\textsubscript{4}SiO\textsubscript{4}, as reported in our previous study.\textsuperscript{43} Simultaneously, Li atoms that do not interact with the oxygen atom preferentially diffuse into the interface between the CNT layer and the SiO\textsubscript{0.43} surface rather than penetrate into the interior of the SiO\textsubscript{0.43} NW, as similarly observed in the SiNW@CNT.

From the SiNW@CNT result, it can be expected that the interfacial diffusion of Li is more facile than bulk diffusion in the Si. We also calculated the mean-square displacements from the MD trajectories for the interfacial and bulk diffusion of Li atoms, as shown in Figure 7. Indeed, the interfacial diffusivity is approximately 5 times higher than the bulk diffusivity (4.5 × 10\textsuperscript{−8} m\textsuperscript{2}/s vs 0.9 × 10\textsuperscript{−8} m\textsuperscript{2}/s) because the sp\textsuperscript{2} carbon provides a slide effect of Li diffusion. A similar result was also reported in the previous first-principles MD work.\textsuperscript{65} This is also supported by a recent experimental measurement that showed that Li diffusion is extremely fast in bilayer graphene.\textsuperscript{66} In the c-Si bulk, a Li atom diffuses from one tetrahedral interstitial site into another interstitial site.\textsuperscript{67} At each interstitial site, one Li atom interacts with six Si atoms. On the other hand, in the case of the interfacial diffusion between Si surface and graphene, the Li...
atom interacts with three Si atoms. Of course, the Li atom also interacts with carbon atoms in graphene. Moreover, the Li diffusion channel size at the interface is bigger than that in the c-Si bulk. In Figure 4b, the Si–Si (covalent bonding) bond distance in the Si bulk and the Si–C (nonbonded interaction) distance at the interface along z-direction are 2.4 and 3.5 Å, respectively. The bigger diffusion channel at the interface results from delocalization stability of electron in the sp<sup>2</sup> carbon and then leads to facile diffusion of Li atoms. Using NEB calculations within the DFT framework, we additionally compared Li diffusion barriers in the c-Si bulk and at the interface, where the structure of Figure 4b was considered. Indeed, our DFT calculations reveal that the Li diffusion barrier in the c-Si bulk is higher (0.5 eV vs 0.2 eV), which supports our ReaxFF-MD simulation.

As a result, introduction of the sp<sup>2</sup> carbonaceous coating layer into Si-based anodes makes Li diffusion more facile, which leads to improved battery performances, such as faster charge/discharge rates. Of course, the carbon coating layer can also provide a buffer effect to volume changes during lithiation, along with preventing the loss of electrical continuity and increasing the electrical conductivity of the Si-based anodes.

4. SUMMARY AND CONCLUSION

Using MD simulations with ReaxFF, we can predict anode properties (e.g., Li capacities and volume changes) of various Si–O–C hybrid nanostructures for LIBs and understand the effects of carbonaceous coating layers on the anode properties. It has been typically believed that a significant volume expansion of Si anodes during lithiation can be prevented by the carbonaceous coating layer, leading to improved battery performance. Although the carbonaceous layer can provide a buffer effect on the volume expansion, the thick layer can also suppress Li penetration into the Si anode, which leads to a very low Li capacity. According to our ReaxFF-MD simulation, for a carbonaceous coating layer with a C/Si ratio of greater than 2.4, Li atoms cannot penetrate into the Si NW. Instead, they exist only on and in graphene layers including in spaces between two graphene layers.

In addition, a sp<sup>2</sup> carbonaceous coating layer can provide a facile Li diffusion channel along the interfaces between the carbon coating layer and the Si nanostructures, which leads to faster charge/discharge rates. Therefore, in CNT or graphene-coated Si-based anodes, Li atoms penetrating through the carbonaceous coating layer tend to diffuse preferentially on the interface between the layer and Si surface. After full coverage of Li on the interface, Li atoms begin to penetrate into the interior of the Si nanostructures. From these results, we expect that the ReaxFF will provide a useful protocol for designing Si–O–C hybrid anodes to obtain better performing LIBs.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b07095.

REFERENCES

(14) Kim, T.; Park, S.; Oh, S. M. Solid-State NMR and Electrochemical Dilatometry Study on Li<sup>+</sup> Uptake/Extraction...


63 Jung, S. C.; Han, Y.-K. Facet-Dependent Lithium Intercalation into Si Crystals: Si(100) vs. Si(111). *Phys. Chem. Chem. Phys.* 2011, 13, 21282−21287.


