An optimized interatomic potential for silicon and its application to thermal stability of silicene

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(Dated: March 28, 2017)

Abstract

An optimized interatomic potential has been constructed for silicon using a modified Tersoff model. The potential reproduces a wide range of properties of Si and improves over existing potentials with respect to point defect structures and energies, surface energies and reconstructions, thermal expansion, melting temperature and other properties. The proposed potential is compared with three other potentials from the literature. The potentials demonstrate reasonable agreement with first-principles binding energies of small Si clusters as well as single-layer and bilayer silicenes. The four potentials are used to evaluate the thermal stability of free-standing silicenes in the form of nano-ribbons, nano-flakes and nano-tubes. While single-layer silicene is mechanically stable at zero Kelvin, it is predicted to become unstable and collapse at room temperature. By contrast, the bilayer silicene demonstrates a larger bending rigidity and remains stable at and even above room temperature. The results suggest that bilayer silicene might exist in a free-standing form at ambient conditions.

PACS numbers:

Keywords: Atomistic modeling, interatomic potential, silicon, silicene, thermal stability.
I. INTRODUCTION

Silicon is one of the most important functional materials widely used in electronic, optical, energy conversion and many other applications. Not surprisingly, Si has been the subject of many classical molecular dynamics (MD) and other large-scale atomistic computer studies for almost three decades. Although classical atomistic simulations cannot access electronic or magnetic properties, they are indispensable for gaining a better understanding of the atomic structures, thermal and mechanical properties of the crystalline, liquid and amorphous Si and various nano-scale objects such as nano-wires and nano-dots. Atomistic simulations rely on semi-empirical interatomic potentials. The accuracy of the results delivered by atomistic simulations depends critically on the reliability of interatomic potentials.

Several dozen semi-empirical potentials have been developed for Si. Although none of them reproduces all properties accurately, there is a trend towards a gradual improvement in their reliability as more sophisticated potential generation methods are developed and larger experimental and first-principles datasets become available for the optimization and testing. The most popular Si potentials were proposed by Stillinger and Weber (SW)\cite{1} and Tersoff\cite{2,3,4}. The original Tersoff potentials were modified by several authors by slightly changing the analytical functions and improving the optimization\cite{5,6,7,8,9}. Other Si potential formats include the environment-dependent interatomic potential\cite{11}, the modified embedded atom method (MEAM) potentials\cite{12,13,14,15,16,17,18}, and bond-order potentials\cite{19,20}.

One of the most significant drawbacks of the existing Si potentials is the overestimation of the melting temperature $T_m$, in many cases by hundreds of degrees. Other typical problems include underestimated vacancy and surface energies and positive Cauchy pressure ($c_{12} - c_{44}$), which in reality is negative ($c_{ij}$ being elastic constants). Kumagai et al.\cite{7} constructed a significantly improved Tersoff potential that predicts $T_m = 1681$ K in close agreement with the experimental value of 1687 K, gives the correct Cauchy pressure, and is accurate with respect to many other properties. This potential, usually referred to as MOD\cite{7}, is probably the most advanced Tersoff-type potential for Si available today. However, it still suffers from a low vacancy formation energy, low surface energies, and overestimated thermal expansion at high temperatures and the volume effect of melting.

The goal of this work was twofold. The first goal was to further improve on the MOD potential\cite{7} by addressing its shortcomings with a minimal impact on other properties. This was achieved by slightly modifying the potential format and performing a deeper optimization. When testing the new potential, we compare it not only with MOD but also with the popular SW potential\cite{1}. We further include the MEAM potential developed by Ryu et al.\cite{14} to represent a different potential format. To our knowledge, this is the only MEAM potential whose melting point is close to experimental.

The second goal was to test the four potentials for their ability to predict the energies
of low-dimensional structures, such as small Si clusters and single- and double-layer forms of silicene (2D allotrope of Si). Si potentials are traditionally considered to be incapable of reproducing low-dimensional structures. This view is largely based on testing the SW potential. The MOD and MEAM potentials have not been tested for the properties of clusters or silicenes in any systematic manner. Such tests were conducted in this work using all four potentials. The results suggest that the present potential, MOD and MEAM do capture the main trends and in many cases agree with first-principles density functional theory (DFT) calculations. As such, they can be suitable for exploratory studies of thermal and mechanical stability of Si clusters and 2D structural forms of Si. In this work we apply them to evaluate the stability of free-standing single-layer and bilayer silicenes at room temperature.

II. POTENTIAL GENERATION PROCEDURES

The total energy of a collection of atoms is represented in the form
\[ E = \frac{1}{2} \sum_{i \neq j} \phi_{ij}(r_{ij}), \]
where \( r_{ij} \) is distance between atoms \( i \) and \( j \) and the bond energy \( \phi_{ij} \) is taken as
\[ \phi_{ij} = f_c(r_{ij})[A \exp(-\lambda_1 r_{ij}) - b_{ij} B \exp(-\lambda_2 r_{ij}) + c_0]. \]  
Here, the bond order \( b_{ij} \) is given by
\[ b_{ij} = (1 + \xi_{ij})^{-\delta}, \]
where
\[ \xi_{ij} = \sum_{k \neq i,j} f_c(r_{ij})g(\theta_{ijk}) \exp[\alpha(r_{ij} - r_{ik})^\beta]. \]
The term \( (1 + \xi_{ij}) \) represent an effective coordination number of atom \( i \) and \( f_c(r_{ij}) \) is a cutoff function. The latter has the form
\[ f_c(r) = \begin{cases} 
1, & r \leq R_1 \\
\frac{1}{2} + \frac{9}{16} \cos\left(\pi \frac{r - R_1}{R_2 - R_1}\right) - \frac{1}{16} \cos\left(3\pi \frac{r - R_1}{R_2 - R_1}\right), & R_1 < r < R_2 \\
0, & r \geq R_2,
\end{cases} \]
where \( R_1 \) and \( R_2 \) are cutoff radii. The outer cutoff \( R_2 \) is chosen between the first and second coordination shells of the diamond cubic structure. The angular function \( g(\theta_{ijk}) \) has the generalized form
\[ g(\theta) = c_1 + \frac{c_2 (h - \cos \theta)^2}{c_3 + (h - \cos \theta)^2} \left\{ 1 + c_4 \exp \left[ -c_5 (h - \cos \theta)^2 \right] \right\}, \]
where $\theta_{ijk}$ is the angle between the bonds $ij$ and $ik$. These functional forms are the same as for the MOD potential\cite{7} except for the new coefficient $c_0$ that was added to better control the attractive part of the potential.

The adjustable parameters of the potential are $A$, $B$, $\alpha$, $h$, $\eta$, $\lambda_1$, $\lambda_2$, $R_1$, $R_2$, $\delta$, $c_0$, $c_1$, $c_2$, $c_3$, $c_4$ and $c_5$. The power $\beta$ is a fixed odd integer. In the original Tersoff potential\cite{2,4} $\beta = 3$, whereas Kumagai et al.\cite{7} chose $\beta = 1$. We tried both numbers and found that $\beta = 3$ gives a better potential.

The free parameters of the potential were trained to reproduce basic physical properties of the diamond cubic (A4) structure and the energies of several alternate structures. Specifically, the fitting database included the experimental lattice parameter $a$, cohesive energy $E_c$, elastic constants $c_{ij}$, and the vacancy formation energy $E_{fv}$. The alternate structures were: simple cubic (SC), $\beta$-Sn (A5), face-centered cubic (FCC), hexagonal closed pack (HCP), body-centered cubic (BCC), simple hexagonal (HEX), wurtzite (B4), BC8, ST12, and clathrate (cP46). Their energies obtained by DFT calculations are available from open-access databases such as Materials Project\cite{21}, OQMD\cite{22} and AFLOW\cite{23,24}. Some of these structures were found experimentally as Si polymorphs under high pressure, others were only generated in the computer for testing purposes. The parameter optimization process utilized a simulated annealing algorithm. The objective function was the sum of weighted squares of deviations of properties from their target values. Numerous optimization runs were conducted using the weights as a tool to achieve the most meaningful distribution of the errors over different properties. Several versions of the potential were generated and the version deemed to be most reasonable was selected as final.

The optimized potential parameters are listed in Table I. The potential has been incorporated in the molecular dynamics package LAMMPS (Large-scale Atomic/Molecular Massively Parallel Simulator)\cite{25} as the pair style `tersoff/mod/c`.

The transferability of the new potential was evaluated by computing a number of physical properties that were not included in the training database and comparing the results with experimental data and/or DFT calculations available in the literature. The same comparison was made for the MOD, MEAM and SW potentials to demonstrate their strengths and weaknesses relative to the new potential. We utilized the MOD and SW potential files from the LAMMPS potential library. The MEAM potential file was obtained from the developers.\cite{14} The potential testing results are reported in the next Section.

### III. PROPERTIES OF SOLID SI

Table II summarizes some of the properties of crystalline Si predicted by the four potentials. All properties have been computed in this work unless otherwise is indicted by citations. The defect energies are reported after full atomic relaxation.
A. Lattice properties

The present potential, MOD and MEAM accurately reproduce the elastic constants. The SW potential gives less accurate elastic constants and a positive Cauchy pressure contrary to experiment. The phonon density of states (DOS) and phonon dispersion relations were computed by the method developed by Kong and implemented in LAMMPS. The MD simulation was performed at 300 K utilizing a primitive $16 \times 16 \times 16$ supercell with 8192 atoms. The DOS plots are shown in Fig. 1(a) and the respective zone-center optical frequencies $\nu_{\text{max}}$ are indicated in Table II. The present potential, MOD and SW predict surprisingly similar $\nu_{\text{max}}$ values that underestimate the experimental frequency by about 2 THz. The MEAM potential overshoots $\nu_{\text{max}}$ by about 10 THz and the entire DOS is stretched by a factor of 1.63. Note that none of the four potentials reproduces the sharp peak at about 5 THz arising from the acoustic zone-boundary phonons.

Fig. 1(b) displays the phonon dispersion curves predicted by the present potential. While general agreement with experiment is evident and the longitudinal acoustic branches are reproduced accurately, the potential overestimates the transverse acoustic zone-boundary frequencies and the optical frequencies.

The cubic lattice parameter $a$ was computed as a function of temperature by zero-pressure MD simulations. The linear thermal expansion coefficient $(a - a_0)/a_0$ relative to room temperature ($a_0$ at 295 K) is compared with experimental data in Fig. 2. The SW potential demonstrates exceptionally good agreement with experiment. The present potential slightly overestimates the experiment at temperatures below 1300 K and underestimates at higher temperatures. The negative slope at high temperatures is unphysical, but the overall agreement with experiment is reasonable. The MOD potential gives a similar thermal expansion at low temperatures but over-predicts it at high temperatures. The MEAM potential grossly overestimates the thermal expansion. Given also the poor agreement for phonons, care should be exercised when using this potential for thermodynamic calculations of crystalline Si. Note that neither phonon properties nor thermal expansion were not included in the fitting databases of the potentials.

B. Lattice defects

According to DFT calculations, a Si vacancy can exist in several metastable structures. In the lowest-energy structure, the four neighbor atoms slightly move towards the vacant site preserving the tetrahedral ($T_d$) symmetry and leaving four dangling bonds. A slightly less favorable structure is obtained when one of the four atoms moves towards the other three and forms six identical bonds. This configuration has a hexagonal ($D_{3d}$) symmetry and is referred to as the “dimerized” or “split” vacancy. This vacancy reconstruction eliminates the dangling bonds but increases the elastic strain in the surrounding lattice.
The present potential and MEAM correctly predict the split vacancy to be less stable than the $T_d$ vacancy. The latter has the formation energy within the range of DFT calculations and consistent with the experimental value of 3.6 eV\textsuperscript{33} (It should be noted, though, that the experiments are performed at high temperatures at which the vacancy structure is unknown.) The MOD and SW potentials significantly under-predict the formation energy of the $T_d$ vacancy. In addition, with the MOD potential the split vacancy spontaneously transforms to a $D_{2d}$ structure with the energy of 3.41 eV (the DFT value is 3.46 eV)\textsuperscript{39}, whereas the SW potential predicts the split vacancy to be mechanically unstable and spontaneously transform to the $T_d$ structure.

Self-interstitials can exist in four distinct configurations: hexagonal (hex), tetrahedral ($T_d$), bond center (B) and $\langle 110 \rangle$ split (Table II). Given the large scatter of the DFT formation energies, all four potentials perform almost equally well. There is one exception: the MEAM and SW potentials predict the hexagonal interstitial to be mechanically unstable and spontaneously transform to the tetrahedral configuration. Both potentials overestimate the B-interstitial energy.

Surface energies were computed for the low-index orientations \{100\}, \{110\} and \{111\}. Experiments have shown that these surfaces can undergo reconstructions to several different structures\textsuperscript{39-42} Reconsrtuction of the \{110\} and \{111\} surfaces are accompanied by a modest energy reduction of about 0.3-0.4 J/m$^2$. In this paper, these surfaces were tested in unreconstructed states. By contrast, the dimer reconstruction of the \{100\} surface to the more stable $2 \times 1$ structure reduces the surface energy by almost 1 J/m$^2$. In this case, both reconstructed and unreconstructed structures were compared with DFT calculations. Table II shows that the SW potential does an excellent job reproducing the DFT surface energies. The MOD potential is the least accurate: it systematically underestimates the surfaces energies for all orientations. The present potential demonstrates a substantial improvement over MOD: all energies are higher and closer to the DFT data. The MEAM potential is equally good for all surfaces except for the unreconstructed \{100\} structure. The latter is mechanically unstable with this potential and reconstructs to the $2 \times 1$ structure spontaneously during static relaxation at 0 K. This instability was not observed in the DFT calculations.\textsuperscript{33} The surface energy of 1.74 J/m$^2$ shown in the table was obtained by constrained relaxation of this surface, in which the atoms were only allowed to move in the direction normal to the surface to prevent the dimerization. With the potential proposed in this work, the unreconstructed \{100\} surface is stable at 0 K and forms symmetrical rows of dimers corresponding to the $2 \times 1$ reconstruction upon heating to 1000 K and slowly cooling down to 0 K.

As another test of the potentials, unstable stacking fault energies $\gamma_{us}$ were calculated for the \{111\} and \{100\} crystal planes. Such faults are important for the description of dislocation core structures. In silicon, dislocations glide predominantly on \{111\} planes. The spacing between \{111\} planes alternates between wide and narrow. In the former case
the chemical bonds are normal to the planes while in the latter they are at 19.47° angles.

A generalized stacking fault is obtained by translation of one half-crystal relative to the other in a chosen direction parallel to a \{111\} plane. Depending on whether the cutting plane passes between widely spaced or narrowly spaced atomic layers, the stacking fault is called shuffle type or glide type, respectively. After each increment of crystal translation, the atoms are allowed to minimize the total energy by local displacements normal (but no parallel) to the fault plane. The excess energy per unit surface area plotted as a function of the translation vector is called the gamma-surface. If the dislocation Burgers vector is parallel to a crystallographic direction \langle hkl \rangle, then its core structure is dictated by the \{111\} \langle hkl \rangle cross-sections of the gamma-surface. The unstable stacking fault energy \( \gamma_{us} \) is the maximum energy in this cross-section.

Figure 5 displays three cross-sections of the \{111\} gamma surface computed with the four potentials in comparison with DFT calculations. The figure additionally includes the \{100\} \langle 110 \rangle cross-section for which DFT data is available. The respective \( \gamma_{us} \) values are summarized in Table V. While none of the potentials reproduces the DFT curves well, the SW potential tends to be the least accurate. For some of the cross-sections, the Tersoff-type potentials “chop off” the tip of the curve due to the short range of atomic interactions and a relatively sharp cutoff. It should also be noted that the potentials do not reproduce the stable stacking fault predicted by DFT calculations [Fig. 5(c)]. This fault arises due to long-range interactions and is not captured by these potentials.

### IV. MELTING TEMPERATURE AND LIQUID PROPERTIES OF SI

The melting temperature was computed by the interface velocity method. A periodic simulation block containing a (111) solid-liquid interface was subject to a series of isothermal MD simulations in the NPT ensemble (zero pressures in all directions) at several different temperatures. The interface migrated towards one phase or the other, depending on whether the temperature was above or below the melting point. The total energy of the system was monitored in this process and was found to be a nearly linear function of time. The slope of this function gives the rate of the energy change due to the phase transformation. A plot of this energy rate as a function of temperature was used to find the melting point by linear interpolation to the zero rate (Fig. 3). For the present potential, the melting temperature obtained was found to be \( T_m = 1687 \pm 4 \) K (the error bar is the standard deviation of the linear fit). This temperature is in excellent agreement with the experimental melting point of 1687 K, even though it was not included in the fitting procedure.

To verify our methodology, similar calculations were performed for the MOD potential. The result was \( T_m = 1682 \pm 4 \) K, which matches 1681 K reported by the potential developers. For the SW potential, the same method gives \( T_m = 1677 \pm 4 \) K. This number is consistent (within the error bars) with \( T_m = 1691 \pm 20 \) K obtained by thermodynamic
calculated. The energy rate versus temperature plots for the MOD and SW potentials can be found in the Supplemental Material to this paper.

Table II summarizes the predictions of the four potentials for the latent heat of melting \( L \) and the volume effect of melting \( \Delta V_m \) relative to the volume of the solid \( V_{\text{solid}} \). None of the potentials reproduces these properties well. The present potential gives the most accurate volume effect \( \Delta V_m/V_{\text{solid}} \) but the least accurate latent heat \( L \). The MOD potential predicts a better value of \( L \) but overestimated the volume effect a factor of two.

Prediction of structural properties of liquid Si presents a significant challenge to interatomic potentials. The nature of atomic bonding in Si changes from covalent to metallic upon melting, causing an increase in density. In this work, the structure of liquid Si was characterized by the pair correlation function \( g(r) \) and the bond-angle distribution function \( g(\theta, r) \). These functions were averaged over 300 uncorrelated snapshots from NPT MD simulations under zero pressure at 1750 K using a simulation block containing 6912 atoms. The angular distribution \( g(\theta, r) \) was computed for bonds within the radius \( r_m \) of the first minimum of \( g(r) \) and normalized by unit area under the curve.

The results are shown in Fig. 4. The present potential turns out to be the least accurate for the liquid properties. The first maximum of \( g(r) \) is too high and the first minimum too deep in comparison with experiment. The other potentials perform better but still show significant departures from the experiment. For the bond-angle distribution, the results computed with the four potentials are very different and none agrees with the DFT simulations. The DFT simulations (ab initio MD) yield a broader distribution with two peaks of comparable height centered at 60° and 90°. The present potential strongly underestimates the 60° peak, overestimates the peak at 90°, and creates another peak at the tetrahedral angle of 109.47°. Using the other potentials, the position of the large peak varies between 90° and 109.47°. Overall, our potential overestimates the degree of structural order in the liquid phase. This seems somewhat surprising given that this potential predicts the most accurate volume effect of melting.

V. ALTERNATE CRYSTAL STRUCTURES OF SI

Tables III and IV show the equilibrium energies of several crystal structures of Si relative to the diamond cubic structure and the respective equilibrium atomic volumes. All these structures were included in the potential fitting procedure except for two. The h-Si\(_6\) structure was recently found by DFT calculations as a new mechanically stable polymorph of Si attractive for optoelectric applications due to its direct band gap of 0.61 eV and interesting transport and optical properties. The h-Si\(_6\) structure is composed of Si triangles forming a hexagonal unit cell with the P6\(_3\)/mmc space group. Si\(_{24}\) is another mechanically stable polymorph that has recently been synthesized by removing Na from the Na\(_4\)Si\(_{24}\) precursor. The orthorhombic Cmcm structure of Si\(_{24}\) contains open channels composed of 6 and 8-
member rings. This polymorph has a quasi-direct 1.3 eV band gap and demonstrates unique electronic and optical properties making it a promising candidate for photovoltaic and other applications. The h-Si₆ and Si₂₄ structures were used for testing purposes to evaluate the transferability of the potentials. All structures were equilibrated by isotropic volume relaxation without local displacements of atoms. For the HCP and Wurtzite structures, the c/a ratios were fixed at the ideal values. For the simple hexagonal, β-Sn and h-Si₆ structures, c/a was fixed at the DFT values of 0.94, 0.552 and 0.562, respectively. It is worth mentioning that the present potential and MOD predict the wurtzite phase to be mechanically unstable at 0 K, which appears to be a generic feature of Tersoff-type potentials.

In Tables III and IV, we compare the predictions of the four potentials with DFT calculations available in the literature. Since the tables are overloaded with numerical data, we found it instructive to recast this information in a graphical format. In Figs. 6 and 7 we plot the energies (volumes) predicted by each potential against the respective DFT energies (volumes) computed by different authors. The bisecting line is the line of perfect correlation. The first thing to notice is the large scatter of the DFT data reported by different sources, which makes a comparison with potentials somewhat ambiguous. For each potential, the agreement was quantified by the root-mean-square (RMS) deviation of the data points from the bisecting line. The RMS deviations obtained are shown in the last row of Tables III and IV. It should emphasized that these RMS deviations reflect not only the differences between the potentials and the DFT calculations but also the scatter of the DFT points themselves. Thus, only comparison of relative values of the RMS deviations makes sense. It should also be noted that the energy deviations are strongly dominated by high-energy structures, such as the close-packed FCC and HCP phases. With this in mind, it is evident that the present potential is the least successful in reproducing the structural energies, whereas the MOD potential is the most successful. For the atomic volumes, however, the present potential and MOD are equally accurate, while the SW and MEAM potentials show significantly larger deviations.

It is interesting to note that the present potential gives the most accurate predictions for the energy and volume of the novel h-Si₆ and Si₂₄ structures that were not included in the fitting database. The MOD potential comes close second, whereas the MEAM and SW potentials are significantly less accurate. The energy-volume plots for several selected structures can be found in the Supplemental Material to this article.

VI. SILICON CLUSTERS

Structure and properties of small Si clusters offer a stringent test of interatomic potentials. Potentials are usually optimized for bulk properties, whereas the clusters display very different and much more open environments in which the coordination number and the type of bounding may change very significantly from one structure to another. Si poten-
tials are traditionally considered to be incapable of reproducing cluster properties, unless such properties are specifically included in the fitting process as in the case of the Boulding and Andersen potential. It was thus interesting to compare the predictions of the four potentials with first-principles calculations.

Figs. 8 and 9 show the structures of the Siₙ (n = 2 – 8) clusters tested in this work. Several different structures are included for each cluster size n whenever first-principles data is available. Such structures are labeled by index m in the Siₙ.m format in the order of increasing cohesive (binding) energy according to the DFT calculations. Thus, the structure labeled Siₙ.1 represents the DFT-predicted ground state for each cluster size n (except for the dimer Si₂ that has a single structure). In addition to the DFT calculations, we included the results of quantum-chemical (QC) calculations on the Hartree-Fock level. Such calculations are more accurate but the energy scale is not fully compatible with that of the DFT calculations. To enable comparison, we followed the proposal that the QC energies be scaled by a factor of 1.2 to ensure agreement with experiment for the dimer energy.

Table VI summarizes the predictions of the four potentials in comparison with DFT calculations and unscaled QC energies. In addition to the clusters, we included an infinitely long linear chain for the sake of comparison. To aid visual comparison, Fig. 10 shows the cluster energies grouped by the cluster size (same-size clusters are connected by straight lines). The QC energies are plotted in the scaled format. Note that the scaling does indeed bring the QC and DFT energies to general agreement with each other. Despite the significant scatter of the individual energies on the level of 0.2-0.4 eV/atom, both calculation methods predict the same ground state for trimers, tetramers and pentamers. None of the potentials predicts the correct ordering for all DFT/QC energies. The present potential and MOD show about the same level of accuracy, but the present potential makes less mistakes in the ordering. Both potentials tend to slightly under-bind the clusters. The MEAM potential is the most successful in reproducing the cluster energies, except for the dimer energy for which it is least accurate. There are mistakes in the ordering, but overall the deviations from the first-principles calculations are about the same as the difference between the two first-principles methods. The SW potential performs poorly: for some of the clusters, the binding energy is underestimated by more than 1 eV/atom. For the infinite atomic chain, the present potential and MOD are in closest agreement with the DFT/QC energies (Table VI).

This comparison leads to the conclusion that, at least for the cluster structures tested here, the present potential, MOD and MEAM are quite capable of predicting the general trends of the cluster energies with a reasonable accuracy without fitting.
VII. 2D SILICON STRUCTURES

A. Single-layer silicenes

Silicenes are 2D allotropes of Si that have recently attracted much attention due to their interesting physical properties and potential device applications.57–61 By contrast to carbon, the sp³ hybridized Si would seem to be an unlikely candidate for a 2D material. Nevertheless, epitaxial honeycomb Si layers have been found on metallic substrates such as (111)Ag.57–59,62–69 Unlike in graphene, some of the 2D forms of Si can have a band gap and could be incorporated in Si-based microelectronics. In particular, electric field applied to the buckled honeycomb structure of silicene, which is normally semi-metallic, can open a band gap whose magnitude increases with the field. It was predicted,70 and recently demonstrated,71 that single-layer silicene can work as a field-effect transistor.70 Experimentally, it has not been possible so far to isolate free-standing silicenes. They are presently considered hypothetic 2D materials and have only been studied by DFT calculations. Such calculations predict that single-layer silicene can possess remarkable electric, optical and magnetic properties,72–75 in addition to ultra-low thermal conductivity.76

The planar (graphene-like) silicene [Fig. 11(a)] is mechanically unstable and spontaneously transforms to the more stable buckled structure [Fig. 11(b,c)].77–79 The latter has a split width Δ of about 0.45-0.49 Å and a first-neighbor distance $r_{1}$ slightly different from that in the planar structure.70,77,79–82 Furthermore, adsorption of Si ad-atoms on the buckled silicene creates a series of periodic dumbbell structures that are even more stable.76,78,81 An ad-atom pushes a nearby Si atom out of its regular position and the two atoms form a dumbbell aligned perpendicular to the silicene plane. The dumbbell atoms have a fourfold coordination (counting the dumbbell bond itself) consistent with the sp³ bonding. One of the best studied dumbbell silicenes has the $\sqrt{3} \times \sqrt{3}$ structure shown in Fig. 11(d,e,f) (the dumbbell atoms are shown in blue and green). The dumbbells distort the hexagonal structural units and create three slightly different nearest-neighbor distances: $r_{\text{I},\text{II}}$, $r_{\text{II},\text{III}}$ and $\Delta_{\text{III},\text{III}}$ [Fig. 11(f)].

The energies and geometric characteristics of the three silicene structures predicted by the four potentials are listed in Table VII. The results of DFT calculations reported in the literature are included for comparison. The agreement with the DFT data is reasonable, especially considering that the 2D structures were not included in the fitting datasets of the potentials. The present potential, MOD and MEAM demonstrate about the same agreement with the DFT calculations. The SW potential tends to be less accurate. For the planar structure, the MOD potential is the most accurate, followed by the present potential, MEAM and then SW. All four potentials correctly predict that the planar structure is mechanically unstable and transforms to the buckled structure. The present potential, MEAM and SW correctly predict that the $\sqrt{3} \times \sqrt{3}$ dumbbell structure has a lower energy.
than the buckled structure. By contrast, the MOD potential predicts that the $\sqrt{3} \times \sqrt{3}$ dumbbell structure has a higher energy, which is contrary to the DFT calculations. All four potentials overestimate the split width $\Delta$ in the buckled structure and the distance $\Delta_{\text{III,III}}$ between the dumbbell atoms in the $\sqrt{3} \times \sqrt{3}$ structure, the present potential being closest to the DFT data.

Thermal stability of single-layer silicenes has been evaluated by MD simulations. The simulated systems were subject to periodic boundary conditions at zero pressure. Fig. 12 demonstrates that a nano-ribbon of buckled silicene is unstable at finite temperatures and quickly collapses to a cluster before temperature reaches 300 K. Likewise, a free-standing sheet (flake) of buckled silicene (Fig. 13) collapses into a cluster with the shape of a bowl when temperature reaches 300 K. The nano-ribbon and nano-flake made of the $\sqrt{3} \times \sqrt{3}$ dimerized silicene collapse as well.

A single-wall nano-tube was also tested for thermal stability. The latter was obtained by wrapping a layer planar silicene into a tube 49 Å in diameter (Fig. 14). The period along the tube axis was 122 Å. As soon as temperature began to increase starting from 0 K, the wall of the tube transformed to the buckled structure and then collapsed before the temperature reached 300 K. Qualitatively the same behavior of the single-layer silicene structures was found with all four potentials. In all cases, the single layer silicene easily developed waves due to thermal fluctuations until neighboring surface regions came close enough to each other to form covalent bonds. Once this happened, the bond-forming process quickly spread over the entire surface and the structure collapsed. This chemical reactivity and the lack of bending rigidity are the main factors that cause the instability of free-standing single-layer silicenes at room temperature.

B. Bilayer silicenes

Another interesting 2D form of silicon is the bilayer silicene. Like the single-layer silicene discussed above, the bilayer silicene was found experimentally on top of metallic surfaces such as Ag(111). By contrast to bilayer graphene, the interlayer bonds in bilayer silicene are covalent sp$^3$ type. As a result, the formation of a bilayer is accompanied by a significant energy release. It can be expected, therefore, that bilayer silicene should be more stable than two single layers.

Several structural forms of the bilayer silicene have been found in experiments and studied by DFT calculations, depending on the type of stacking of the two layers and whether they are planar or buckled. Three of the structures, referred to as AA$_p$, AA$'$ and AB, are shown in Fig. 15. The AA$_p$ structure is obtained by stacking two planar silicene layers (A) on top of each other and connecting them by vertical covalent bonds [Fig. 15(a)]. This structure is characterized by the geometric parameters $b$ (side of the rhombic structural unit) and the interlayer spacing $h$. The bond length between Si atoms is $d_1 = b/\sqrt{3}$ within
each layer and \( h \) between the layers. In the AA' structure, both layers are buckled, and the buckling of one layer (A') is inverted with respect to the buckling of the other layer (A) [Fig. 15(b)]. As a result, half of the interlayer distances are short, leading to the formation of covalent bonds, and the other half of the distances are longer and covalent bonds do not form. The geometric parameters of the structure are \( b \) (defined above), the in-layer bond length \( d_1 \), the interlayer bond length \( d_2 \), and the split width of each layer \( \Delta \). The distance between the layers is \( h = d_2 + \Delta \). Finally, in the AB structure, two buckled silicene layers A and B are stacked together so that half of the atoms of one layer project into the centers of the hexagonal units of the other layer [Fig. 15(c)]. The remaining half of the atoms project onto each other and form vertical covalent bonds. As with the single-layer silicenes, it has not been possible so far to isolate free-standing bilayer silicene experimentally.

The cohesive energies \( E_c \) and geometric parameters of three bilayer silicenes computed with four interatomic potentials are compared with DFT data in Table VIII. The Table also shows the energies \( \Delta E \) of the buckled bilayers AA' and AB relative to the planar bilayer AA\(_p\). None of the potentials matches the DFT calculations accurately. However, the present potential displays the closest agreement. The MOD potential incorrectly predicts that the buckled structures AA' and AB are more stable than AA\(_p\) (negative \( \Delta E \) values), which is contrary to the DFT calculations. It should be noted that all four potentials predict virtually identical properties of the AA' and AB silicenes. This is not surprising: considering only nearest neighbor bonds, the local atomic environments in the two structures are identical. Their DFT lattice parameters \( b \) are indeed the same (3.84 Å), but the DFT energies are different (0.33 and 0.17 eV/atom, respectively; our potential gives \( \Delta E = 0.12 \) eV/atom for both). This discrepancy apparently reflects a common feature of all short-range Si potentials.

To assess thermal stability of bilayer silicenes, MD simulations were conducted for the same nano-ribbon, nano-flake and nano-tube configurations as discussed above. The most stable AA\(_p\) silicene was chosen for the tests. The samples were heated up to 300 K and annealed at this temperature for 10 ns. The systems developed significant capillary waves, especially the nano-ribbon, but none of them collapsed (Fig. 16). Although 10 ns is a short time in comparison with experimental times, these tests confirm that the bilayer silicene has a much greater bending rigidity and smaller reactivity in comparison with its single-layer counterpart. As such, it has a much better chance of survival in a free-standing form at room temperature.

In additional tests, the nano-flake was heated from 300 K to 1000 K in 6 ns followed by an isothermal anneal for 2 ns at 1000 K. The surface of the flake developed a set of thermally activated point defects, such as ad-atoms and locally buckled configurations, but the flake itself did not collapse. This again confirms the significant thermal stability of the bilayer silicene, possibly even at high temperatures. The same tests were conducted with all four potentials and the results were qualitatively similar. With the MOD potential,
the initial $AA_p$ silicene quickly transformed to the more stable buckled structure, but the system still did not collapse.

VIII. DISCUSSION AND CONCLUSIONS

Silicon is one of the most challenging elements for semi-empirical interatomic potentials. It has over a dozen polymorphs that are stable at different temperatures and pressures and exhibit different coordination numbers and types of bonding ranging from strongly covalent to metallic. The diamond cubic phase displays a rather complex behavior with several possible structures of point defects, a number of surface reconstructions, and an increase in density upon melting. It is not surprising that the existing Si potentials are not nearly as successful in describing this material as some of the embedded-atom potentials for metals.\textsuperscript{88–90} In this work, we developed a new Si potential with the goal of improving some of the properties that were not captured accurately by other potentials. For comparison, we selected three potentials from the literature that we consider most reliable\textsuperscript{7,14} or most popular.

Extensive tests have shown that the present potential does achieve the desired improvements, in particular with regard to the vacancy formation energies, surface formation energies and reconstructions, thermal expansion factors and a few other properties. The potential is more accurate, in comparison with other potentials, in reproducing the DFT data for the novel Si polymorphs $h$-$Si_6$ and $Si_{24}$ without including them in the fitting database. But the tests have also shown that each of the four potentials has its successes and failures. The present potential makes inaccurate predictions for the energies of high-lying Si polymorphs (although their atomic volumes are quite accurate), for the latent heat of melting, and for the short-range order in the liquid phase. The MOD potential\textsuperscript{7} has its own drawbacks mentioned in Section I. The MEAM potential\textsuperscript{14} grossly overestimates the phonon frequencies and thermal expansion factors, in addition to the incorrect $\{100\}$ surface reconstruction. The SW potential successfully reproduces the surface energies and thermal expansion factors but predicts a positive Cauchy pressure and systematically overestimates the atomic volumes of Si polymorphs (as does the MEAM potential).

The potentials were put through a very stringent test by computing the binding energies of small $Si_n$ clusters. Such clusters were not included in the potential fitting procedure and are traditionally considered to be out of reach of potentials unless specifically included in fitting database. Surprisingly, the present potential, the MOD potential\textsuperscript{7} and especially the MEAM potential\textsuperscript{14} reproduce the general trends of the cluster energies reasonably well (Fig. 10). In many cases, the ranking of the energies of different geometries for the same cluster size $n$ agrees with first-principles calculations. The SW potential is less accurate: it systematically under-binds the clusters and makes more mistakes in the energy ordering.

Encouraged by the reasonable performance for the clusters, we applied the potentials
to model single-layer and bilayer silicenes, which were not included in the potential fitting either. While none of the potentials reproduces all DFT calculations accurately, they generally perform reasonably well. One notable exception is the MOD potential, which under-binds the $\sqrt{3} \times \sqrt{3}$ dumbbell structure of the single-layer silicene and fails to reproduce the correct ground-state of the bilayer silicene. Furthermore, all four potentials predict identical energies of the AA' and AB bilayer silicenes, whereas the DFT energies are different. Other than this, the trends are captured quite well. The present potential demonstrates the best performance for the bilayer silicenes.

Experimentally, silicenes have only been found on metallic substrates. Whether they can exist in a free-standing form at room temperature remains an open question. Evaluation of their thermal stability requires MD simulations of relatively large systems for relatively long times that are not currently accessible by DFT methods. Although interatomic potentials are less reliable, they can be suitable for a preliminary assessment. The MD simulations performed in this work indicate that single-layer silicenes are unlikely to exist in a free-standing form. Their large bending compliance and chemical reactivity lead to the development of large shape fluctuations and eventually the formation of covalent bonds between neighboring surface regions at or below room temperature. By contrast, bilayer silicenes exhibit much greater bending rigidity and lower surface reactivity. Nanostructures such as nano-ribbons, nano-flakes and nano-tubes remain intact at and above room temperature, at least on a 10 ns timescale. The fact that this behavior was observed with all four potentials points to the generality of these observations and suggests that free-standing bilayer silicenes might be stable at room temperature. Of course, this tentative conclusion requires validation by more detailed and more accurate studies in the future.

The four potentials discussed in this work are likely to represent the limit of what can be achieved with short-range semi-empirical potentials. Further improvements can only be made by developing more sophisticated, longer-range, and thus significantly slower potentials. Analytical bond-order potentials offer one option. Recent years have seen a rising interest in machine-learning potentials. While even slower, they allow one to achieve an impressive accuracy of interpolation between DFT energies, in some cases up to a few meV/atom. However, the lack of transferability to configurations outside the training dataset remains an issue.

**Acknowledgements**

This work was supported by the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering, the Physical Behavior of Materials
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Table I: Optimized parameters of the new Si potential. Parameters of the MOD potential\textsuperscript{[7]} are listed for comparison.

| Parameter | Present        | MOD\textsuperscript{a} |
|-----------|----------------|------------------------|
| $A$ (eV)  | 3198.51383     | 3281.5905              |
| $B$ (eV)  | 117.780724     | 121.00047              |
| $\lambda_1$ (Å$^{-1}$) | 3.18011795  | 3.2300135              |
| $\lambda_2$ (Å$^{-1}$) | 1.39343356    | 1.3457970              |
| $\eta$   | 2.16152496     | 1.0000000              |
| $\eta \times \delta$ | 0.544097766 | 0.53298909 |
| $\alpha$ | 1.80536502     | 2.3890327              |
| $\beta$  | 3              | 1                      |
| $c_0$ (eV) | -0.0059204  | 0.0                    |
| $c_1$   | 0.201232428    | 0.20173476             |
| $c_2$   | 614230.043     | 730418.72              |
| $c_3$   | 9.96439.097    | 1000000.0              |
| $c_4$   | 3.33560562     | 1.0000000              |
| $c_5$   | 25.2096377     | 26.000000              |
| $h$     | -0.381360867   | -0.36500000            |
| $R_1$ (Å) | 2.54388270    | 2.7                    |
| $R_2$ (Å) | 3.20569403    | 3.3                    |

\textsuperscript{a}Ref.\textsuperscript{[7]}
Table II: Properties of diamond cubic Si computed with four interatomic potentials in comparison with experimental data and DFT calculations.

| Property | Experiment | DFT | Present | MOD | MEAM$^w$ | SW$^u$ |
|----------|------------|-----|---------|-----|----------|--------|
| $E_c$ (eV/atom) | 4.63$^c$ | 4.84$^a$ | 4.630 | 4.630 | 4.630 | 4.337 |
| $a$ (Å) | 5.430$^a$ | 5.451$^a$ | 5.434 | 5.429 | 5.431 | 5.431 |
| $c_{11}$ (GPa) | 165$^a$; 167.40$^b$ | 172.6 | 166.4 | 163.8 | 151.4 |
| $c_{12}$ (GPa) | 64$^a$; 65.23$^b$ | 64.6 | 65.3 | 64.5 | 76.4 |
| $c_{44}$ (GPa) | 79.2$^a$; 79.57$^b$ | 81.3 | 77.1 | 76.5 | 56.4 |
| $\nu_{\text{max}}$ (THz) | 15.7$^o$ | 17.6 | 17.5 | 25.6 | 17.8 |

Vacancy:

- $E_f^V$ ($T_d$) (eV) 3.6$^j$ 3.17$^m$; 3.69$^t$ 3.54 2.82 3.57 2.64 3.29-4.3$^h$; 3.70-3.84$^a$
- $E_f^V$ ($D_{3d}$) (eV) 3.97$^c$; 4.29$^v$; 4.37$^n$ 3.67-3.70$^e$; 5.023$^i$

Interstitials:

- $E_f^I$ (hex) (eV) 3.31–5$^h$; 2.87-3.80$^s$ 3.51 4.13$^d$ – –
- $E_f^I$ ($T_d$) (eV) 3.43–6$^h$; 3.43-5.10$^s$ 3.01 3.27$^d$ 4.12 4.93
- $E_f^I$ (B) (eV) 4–5$^h$ 4.34 5.03$^d$ 6.78 5.61
- $E_f^I$ ($110$) (eV) 3.31–3.84$^h$; 2.87-3.84$^s$ 3.26 3.57$^d$ 3.91 4.41

Surface energy $\gamma_s$ (J/m$^2$):

- $\{111\}$ 1.24$^q$; 1.23$^p$ 1.57$^d$; 1.74$^f$ 1.11 0.89 1.2 1.36
- $\{100\}$ 2.14$^l$; 2.39$^f$; 2.36$^k$ 2.19 1.77 1.74$^e$ 2.36
- $\{100\}_{2\times1}$ 1.36$^p$ 1.71$^g$; 1.45$^f$; 1.51$^k$ 1.21 1.07 1.24 1.45
- $\{110\}$ 1.43$^p$ 1.75$k$ 1.36 1.08 1.41 1.67

Melting:

- $T_m$ (K) 1687 1687 1681$^d$; 1682 1687$^w$ 1691$^v$; 1677
- $\Delta V_m/V_{\text{solid}}$ (%) -5.1$^a$ -3.8 -12.5 -2.7 -7.2
- $L$ (kJ/mol) 50.6$^a$ 24.0 34.7 43.2 31.1

References:

- $^a$Ref. 26
- $^b$Ref. 27
- $^c$Ref. 98
- $^d$Ref. 7
- $^e$Constrained relaxation
- $^f$Ref. 43
- $^g$Ref. 99
- $^h$Ref. 100
- $^i$Ref. 17
- $^j$Ref. 34
- $^k$Ref. 37
- $^l$Ref. 29
- $^m$Ref. 101
- $^n$Ref. 36
- $^o$Ref. 38
- $^p$Ref. 35
- $^q$Ref. 102
- $^r$Ref. 103
- $^s$Ref. 104
- $^t$Ref. 33
- $^u$Ref. 1
- $^v$Ref. 14

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Table III: Energies (eV/atom) of alternate crystal structures of Si relative to the cubic diamond phase in comparison with first-principles calculations.

| Structure | Ab initio | Present MOD<sup>i</sup> MEAM<sup>j</sup> SW<sup>a</sup> |
|-----------|----------|-----------------|
| FCC       | 0.449<sup>c</sup>; 0.57<sup>f</sup>; 0.537<sup>k,m</sup> | 1.113 0.4473 0.8975 0.3963 |
|           | 0.6494<sup>n</sup>; 0.5536<sup>p</sup> |  |
| HCP       | 0.55<sup>f</sup>; 0.508<sup>m</sup>; 0.5946<sup>n</sup>; 0.5301<sup>p</sup> | 1.1019 0.4426 0.8909 0.3963 |
| BCC       | 0.43<sup>p</sup>; 0.435<sup>c</sup>; 0.46<sup>j</sup>; 0.53<sup>f</sup>; 0.523<sup>m</sup>; 0.6142<sup>p</sup> | 0.6945 0.4377 0.5354 0.2810 |
| HEX       | 0.293<sup>e</sup> | 0.7322 0.3901 0.5591 0.3876 |
| SC        | 0.276<sup>c</sup>; 0.35<sup>f</sup>; 0.38<sup>b</sup> | 0.2849 0.3076 0.4688 0.2745 |
| β-Sn      | 0.19<sup>d</sup>; 0.33<sup>d</sup>; 0.414<sup>d</sup>; 0.454<sup>d</sup> | 0.3725 0.3343 0.3671 0.2012 |
|           | 0.3264<sup>n</sup>; 0.27<sup>f</sup>; 0.32<sup>b</sup>; 0.290<sup>h</sup> |  |
|           | 0.2718<sup>p</sup>; 0.380<sup>f</sup>; 0.291<sup>m</sup> |  |
| BC8       | 0.13<sup>c</sup>; 0.159<sup>b</sup>; 0.126<sup>i</sup> | 0.2008 0.2127 0.2502 0.1880 |
|           | 0.110<sup>k</sup>; 0.166<sup>n</sup> |  |
| Wurtzite  | 0.011<sup>h,m</sup>; 0.016<sup>f</sup> | 0.0000 0.0000 0.00001 0.0000 |
| ST12      | 0.136<sup>i</sup>; 0.1181<sup>k</sup> | 0.3900 0.4470 0.6031 0.4857 |
| cP46      | 0.063<sup>h</sup>; 0.0637<sup>n</sup> | 0.0703 0.0581 0.0625 0.0502 |
| h-Si<sub>6</sub> | 0.35<sup>g</sup> | 0.5021 0.5863 0.6464 0.8417 |
| Si<sub>24</sub> | 0.09<sup>f</sup> | 0.1816 0.1864 0.2340 0.1949 |
| RMS error | 0.2883 0.1124 0.2138 0.1745 |

<sup>a</sup>Ref. 1<sup>i</sup> <sup>b</sup>Ref. 99 and references therein
<sup>c</sup>Ref. 7<sup>j</sup> <sup>d</sup>Ref. 105 and references therein
<sup>e</sup>Ref. 106 and references therein; <sup>f</sup>Ref. 103
<sup>g</sup>Ref. 50<sup>i</sup> <sup>h</sup>Ref. 21<sup>m</sup> <sup>i</sup>Ref. 7<sup>j</sup> <sup>j</sup>Ref. 107 <sup>k</sup>Ref. 108
<sup>l</sup>Ref. 14<sup>m</sup> <sup>l</sup>Ref. 22<sup>n</sup> <sup>n</sup>Ref. 109 <sup>q</sup>Refs. 2324
<sup>q</sup>Ref. 110 <sup>q</sup>Ref. 111 <sup>q</sup>Ref. 112 <sup>q</sup>Ref. 51
Table IV: Equilibrium volume per atom (Å³) of alternate crystal structures of Si in comparison with experiment and first-principles calculations.

| Structure  | Experiment | Ab initio | Present MOD | MEAM † | SW ‡ |
|------------|------------|-----------|-------------|--------|-------|
| Diamond    | 20.024 †   | 20.264 c; 20.444 d; 20.439 h; 20.33 i; 19.59 j; 20.46 m; 19.03 p; 16.686 f; 20.385 i | 20.052 | 20.002 | 20.024 | 20.023 |
|            |            | 19.77 h; 20.42 b; 20.124 b; 20.21 b; 20.08 b |             |        |       |       |
| FCC        | 14.678 c; 14.484 d; 14.504 i; 14.810 j; 14.337 k | 14.448 | 14.262 | 17.312 | 17.824 |
| HCP        | 14.477 c; 14.313 i; 14.68 j | 14.439 | 14.257 | 17.279 | 17.824 |
| BCC        | 14.738 c; 14.2427 k | 14.483 | 14.045 | 15.592 | 17.082 |
| HEX        | 15.21 i; 14.56 i; 13.15 p | 15.423 | 14.992 | 17.457 | 18.230 |
| SC         | 16.179 c; 15.7653 k | 15.639 | 15.581 | 18.194 | 17.822 |
| β-Sn       | 14.0 f; 14.2 f | 15.479 c; 15.334 d; 16.0 f; 15.292 i | 15.016 | 15.085 | 16.560 | 17.275 |
|            |            | 14.92 b; 15.45 b; 15.25 b; 15.34 b |             |        |       |       |
|            |            | 15.31 b; 15.405 i; 15.35 m; 14.8859 k |             |        |       |       |
| BC8        | 18.13 f; 18.26 f | 17.724 f; 17.48 g; 18.44 j; 18.427 d | 18.112 | 18.079 | 19.374 | 17.902 |
|            |            | 18.2619 k; 18.082 n |             |        |       |       |
| Wurtzite   | 20.324 c; 20.440 d; 20.380 i; 19.7575 k | 20.052 | 20.002 | 20.024 | 20.023 |
| ST12       | 17.65 g; 17.57 g | 18.083 | 18.123 | 20.931 | 18.325 |
| cP46       | 23.256 d; 23.214 i; 23.128 j | 22.746 | 22.663 | 23.042 | 22.663 |
| h-Si6      | 27.188 g | 28.575 | 28.725 | 33.460 | 31.667 |
| Si24       | 21.52 r | 21.934 r | 21.861 | 21.809 | 23.189 | 22.083 |
| RMS error  |             | 0.6758 | 0.6609 | 1.9147 | 2.0452 |

*Ref. 11; †Ref. 105 and references therein; ‡Ref. 103; †Ref. 21
†Ref. 7; †Ref. 107 and references therein; ‡Ref. 108; †Ref. 22
†Ref. 109; †Ref. 20; †Ref. 113; †Ref. 111; †Ref. 114; †Ref. 106; †Ref. 50; †Ref. 51
Table V: Energies $\gamma_{us}$ (in Jm$^{-2}$) of unstable stacking faults computed with the present interatomic potential in comparison with other potentials and first-principles calculations.

| Property                  | Ab initio | Present MOD$^a$ | MEAM$^b$ | SW$^c$ |
|---------------------------|-----------|-----------------|----------|--------|
| (111)⟨110⟩ shuffle       | 1.81$^{d,e}$ | 1.09           | 1.04     | 1.40   | 0.87   |
| (111)⟨110⟩ glide         | 4.97$^f$   | 5.25           | 5.00     | 4.58   | 6.37   |
| (111)⟨211⟩ glide         | 2.02$^{d,e}$ | 2.39           | 2.05     | 2.86   | 3.09   |
| (100)⟨110⟩               | 2.15$^e$   | 2.44           | 1.77     | 2.19   | 1.61   |

$^a$Ref. 7; $^b$Ref. 14; $^c$Ref. 1
$^d$Ref. 115; $^e$Ref. 116; $^f$Digitized from Ref. 116
Table VI: Cohesive energies (eV/atom) of Si clusters relative to isolated atoms computed with four interatomic potentials in comparison with first-principles calculations. The asterisk marks mechanically unstable structures whose energies were obtained by anisotropic volume relaxation without local atomic displacements.

| Cluster | Experiment | *Ab initio* | Present MOD | MEAM | SW |
|---------|------------|-------------|-------------|------|----|
| Si₂     | 1.62<sup>d</sup> | 1.53<sup>d</sup>; 1.81<sup>e</sup> | 1.327 | 1.788 | 2.473 | 1.084 |
| Si₃.1   | 2.03<sup>d</sup>; 2.41<sup>e</sup> | 1.710 | 2.003 | 2.519 | 1.267 |
| Si₃.2   | 2.6<sup>d</sup> | 2.39<sup>d</sup>; 2.58<sup>e</sup> | 1.757 | 2.197 | 2.672 | 1.446 |
| Si₃.3   | 2.61<sup>e</sup> | 2.259 | 2.147 | 2.815 | 1.480 |
| Si₄.1   | 1.82<sup>d</sup>; 2.48<sup>e</sup> | 1.901 | 2.121 | 2.593 | 1.372 |
| Si₄.2   | 2.02<sup>d</sup>; 2.49<sup>e</sup> | 2.457 | 2.325* | 2.984 | 1.669* |
| Si₄.3   | 2.21<sup>d</sup>; 2.73<sup>e</sup> | 2.571 | 2.810 | 3.021 | 2.035 |
| Si₄.4   | 2.22<sup>d</sup> | 2.219 | 2.232 | 2.759 | 1.525 |
| Si₄.5   | 2.68<sup>d</sup>; 3.09<sup>e</sup> | 2.579 | 2.441* | 2.995 | 1.746* |
| Si₅.1   | 2.02<sup>d</sup>; 2.62<sup>e</sup> | 2.613 | 3.013 | 3.075 | 2.168 |
| Si₅.2   | 2.69<sup>d</sup>; 3.04<sup>e</sup> | 2.800 | 2.731 | 3.159 | 2.062 |
| Si₅.3   | 3.09<sup>f</sup> | 2.678 | 2.549* | 3.037 | 1.845* |
| Si₅.4   | 2.78<sup>d</sup>; 3.30<sup>e</sup> | 2.836 | 2.821 | 3.124 | 2.146 |
| Si₅.5   | 2.22<sup>d</sup> | 2.618 | 3.023 | 3.075 | 2.168 |
| Si₆.1   | 3.33<sup>e</sup> | 2.862 | 2.793 | 3.269 | 2.142 |
| Si₆.2   | 3.04<sup>d</sup>; 3.448<sup>e</sup> | 2.664* | 2.658* | 3.225 | 1.970* |
| Si₆.3   | 3.453<sup>e</sup> | 2.706* | 2.771* | 3.260 | 2.139* |
| Si₆.4   | 3.56<sup>e</sup> | 2.938 | 2.960 | 3.344 | 2.321* |
| Si₆.5   | 3.22<sup>e</sup> | 2.919 | 3.006* | 3.267 | 2.379* |
| Chain   | 2.260<sup>a</sup> | 2.477 | 2.475 | 2.771 | 1.680 |

<sup>a</sup>Ref. 7; <sup>b</sup>Ref. 14; <sup>c</sup>Ref. 11
<sup>d</sup>Ref. 54 and references therein; <sup>e</sup>Ref. 53
Table VII: Properties of single-layer silicenes computed with four interatomic potentials in comparison with DFT calculations.

| Property                  | Ab initio | Present MOD | MEAM | SW |
|---------------------------|-----------|-------------|------|----|
| Honeycomb planar:         |           |             |      |    |
| $E_c$ (eV/atom)           | 3.96f     | 3.6955      | 3.8280 | 3.6234 | 3.1450 |
| $b$ (Å)                   | 3.895f    | 4.042       | 4.019 | 4.306 | 4.104 |
| $r_1$ (Å)                 | 2.249f    | 2.332       | 2.321 | 2.486 | 2.369 |

| Honeycomb buckled:        |           |             |      |    |
| $\Delta E_c$\text{buckled–diamond} (eV/atom) | 0.76e     | 0.88        | 0.69  | 0.89 | 1.09 |
| $\Delta E_c$\text{buckled–}\sqrt{3}\times\sqrt{3}$ (eV/atom) | 0.048d    | 0.14        | -0.08 | 0.08 | 0.07 |
| $b$ (Å)                   | 3.88k; 3.87d; 3.83e | 3.870      | 3.820 | 3.944 | 3.840 |
| $r_1$ (Å)                 | 2.28d; 2.25e,i | 2.328       | 2.312 | 2.449 | 2.352 |
| $\Delta$ (Å)             | 0.44d,e; 0.45g,j | 0.655       | 0.694 | 0.901 | 0.784 |
|                           | 0.46i; 0.49f |            |      |      |      |

| $\sqrt{3} \times \sqrt{3}$ Dumbbell: |           |             |      |    |
| $b$ (Å)                 | 6.52d,h    | 6.475       | 6.471 | 6.312 | 6.604 |
| $r_{II,III}$ (Å)        | 2.40d,h    | 2.393       | 2.425 | 2.526 | 2.513 |
| $r_{I,II}$ (Å)          | 2.28d      | 2.333       | 2.425 | 2.456 | 2.359 |
| $\Delta_{III,III}$ (Å) | 2.76b      | 3.0564      | 3.111 | 3.160 | 3.261 |

$^{a}$Ref. 7; $^{b}$Ref. 14; $^{c}$Ref. 1; $^{d}$Ref. 81; $^{e}$Ref. 77; $^{f}$Ref. 106

$^{g}$Ref. 82; $^{h}$Ref. 78; $^{i}$Ref. 70; $^{j}$Ref. 79; $^{k}$Ref. 58; $^{l}$Ref. 80
Table VIII: Properties of three structures of bilayer silicenes computed with interatomic potentials and DFT calculations.

| Property                  | Ab initio | Present | MOD\(^a\) MEAM\(^b\) | SW\(^c\) |
|---------------------------|-----------|---------|------------------------|----------|
| **Bilayer planar silicene AA\(_p\):** |           |         |                        |          |
| \(E_c\) (eV/atom)         | 4.16\(^d\); 4.27\(^d\) | 4.3067  | 4.2183  | 4.1739  | 3.8542  |
| \(b\) (Å)                 | 4.12\(^e\); 4.13\(^d\) | 4.3264; 3.9804 | 4.0913  | 4.2685  | 4.1497  |
| \(d_1\) (Å)               | 2.38\(^d,e\); 2.39\(^d\) | 2.3641; 2.3737 | 2.3621  | 2.4644  | 2.3958  |
| \(h\) (Å)                 | 2.41\(^d,e\) | 2.3916  | 2.4393  | 2.4869  | 2.4428  |
| **Bilayer buckled silicene AA\(_p\)′:** |           |         |                        |          |
| \(E_c\) (eV/atom)         |           | 4.1866  | 4.2776  | 4.1626  | 3.7945  |
| \(\Delta E_{\text{buckled-planar}}\) (eV/atom) | 0.33\(^e\) | 0.1201  | -0.0593 | 0.0113  | 0.0597  |
| \(b\) (Å)                 | 3.84\(^e\) | 3.8430  | 3.8245  | 3.9155  | 3.8402  |
| \(d_1\) (Å)               | 2.3405    | 2.3311  | 2.4081  | 2.3517  |
| \(d_2\) (Å)               | 2.3543    | 2.3515  | 2.3801  | 2.3517  |
| \(h\) (Å)                 | 3.0994    | 3.0990  | 3.2101  | 3.1356  |
| \(\Delta\) (Å)            | 0.7451    | 0.7475  | 0.8300  | 0.7839  |
| **Bilayer buckled silicene AB:** |           |         |                        |          |
| \(E_c\) (eV/atom)         | 4.10\(^d\); 4.25\(^d\) | 4.1866  | 4.2776  | 4.1626  | 3.7945  |
| \(\Delta E_{\text{buckled-planar}}\) (eV/atom) | 0.17\(^e\) | 0.1201  | -0.0593 | 0.0113  | 0.0597  |
| \(b\) (Å)                 | 3.84\(^d,e\); 3.86\(^d\) | 3.8429  | 3.8245  | 3.9155  | 3.8402  |
| \(d_1\) (Å)               | 2.32\(^d\) | 2.3405  | 2.3311  | 2.4082  | 2.3517  |
| \(d_2\) (Å)               | 2.51\(^d\); 2.54\(^d\) | 2.3543  | 2.3515  | 2.3801  | 2.3517  |
| \(h\) (Å)                 | 3.19\(^d\); 3.20\(^d\) | 3.0994  | 3.0990  | 3.2101  | 3.1359  |
| \(\Delta\) (Å)            | 0.66\(^d\); 0.68\(^d\) | 0.7451  | 0.7475  | 0.8300  | 0.7839  |

\(^a\)Ref. 7; \(^b\)Ref. 14; \(^c\)Ref. 1; \(^d\)Ref. 84; \(^e\)Ref. 86
Figure 1: Phonons properties of diamond cubic Si. (a) Density of states calculated with different interatomic potentials in comparison with experimental data. (b) Dispersion relations at room temperature computed with the present potential in comparison with experiment.
Figure 2: Linear thermal expansion of Si lattice, \((a - a_0)/a_0\), relative to room temperature \((a_0 \text{ at } 295 \text{ K})\) predicted by four interatomic potentials in comparison with experimental measurements.\(^{117,118}\)

Figure 3: Rate of energy charge as a function of temperature during melting and crystallization of Si modeled with the present potential. The line is the linear fit to determine the melting temperature.
Figure 4: Structure of liquid Si: (a) pair correlation function $g(r)$ and (b) bond-angle distribution $g(\theta, r)$ computed with the present interatomic potential at the temperature of 1750 K in comparison with first-principles calculation at 1767 K, experimental data at 1733 K, and the MOD, MEAM and SW potentials at 1767 K. The arrows indicate the angles of $60^\circ$, $90^\circ$ and $109.47^\circ$. 
Figure 5: Selected cross-sections of the \{111\} and \{100\} gamma surfaces predicted by the present potential in comparison with other potentials and DFT calculations.
Figure 6: DFT energies of crystal structures of Si versus the energies predicted by interatomic potentials: (a) present potential, (b) MOD potential, (c) MEAM potential, and (d) SW potential. The energies are counted per atom relative to the diamond cubic structure. The line of perfect correlation is indicated.
Figure 7: DFT atomic volumes of crystal structures of Si versus the atomic volumes predicted by interatomic potentials: (a) present potential, (b) MOD potential, (c) MEAM potential, and (d) SW potential. The line of perfect correlation is indicated.
Figure 8: Structures of dimer, trimer, tetramer and pentamer Si clusters tested in this work. The labels indicate the cluster notations.
Figure 9: Structures of hexamer, heptamer and octamer Si clusters tested in this work. The labels indicate the cluster notations.
Figure 10: Binding energies of Si clusters predicted by interatomic potentials: (a) present potential, (b) MOD potential, (c) MEAM potential, and (d) SW potential. First-principles energies computed by DFT and QC methods are shown for comparison. The clusters are divided into groups corresponding to the same number of atoms and are ordered with increasing binding energy. The cluster structures are shown in Figs. 8 and 9.
Figure 11: Silicene structures: (a) graphitic (planar) structure, (b,c) top and edge views of the buckled structure, and (d,e,f) top and edge views of the $\sqrt{3} \times \sqrt{3}$ dumbbell structure.
Figure 12: Snapshots of MD simulations of a non-ribbon of buckled silicene modeled with the present interatomic potential. The temperature increases with a constant rate from 0 K to 300 K over a 1 ns time period. The images show one repeat unit of the ribbon containing 1080 atoms. The time increases from (a) (initial state) to (d) (final state).
Figure 13: Snapshots of MD simulations of a 6120-atom free-standing nano-sheet (flake) of buckled silicene modeled with the present interatomic potential. The temperature increases with a constant rate from 0 K to 300 K over a 0.6 ns time period [snapshots (a), (b) and (c)] followed by an isothermal anneal at 300 K [snapshot (d) taken 0.2 ns into the anneal].
Figure 14: Snapshots of MD simulations of a single-wall nano-tube of planar silicene modeled with the present interatomic potential. The temperature increases with a constant rate from 0 K to 300 K over a 2 ns time period. The images show one period of the tube (diameter 49 Å, length 122 Å, 2160 atoms). The time increases from (a) (initial state) to (d) (final state).
Figure 15: Structures of bilayer silicenes: (a) AA$_p$, (b) AA', (c) AB.
Figure 16: Snapshots of MD simulations of the AA’ bilayer silicene after a 10 ns anneal at 300 K. (a) Nano-ribbon, (b) Free-standing nano-sheet (flake), (c) Nano-tube (the two layers are shown in different colors for clarity).