Ab initio Monte Carlo simulations for finite-temperature properties: Application to lithium clusters and bulk liquid lithium

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Ab initio Monte Carlo simulations have been performed to determine the equilibrium properties of liquid lithium and lithium clusters at different temperatures. First-principles density-functional methods were employed to calculate the potential-energy change for each proposed change of configuration, which was then accepted or rejected according to the Metropolis Monte Carlo scheme. The resulting structural properties are compared to data from experimental measurements and ab initio molecular dynamics simulations. It is shown that accurate structural information can be obtained with ab initio Monte Carlo simulations at computational costs comparable to ab initio molecular dynamics methods. We demonstrate that ab initio Monte Carlo simulations for the properties of fairly large condensed-matter systems at nonzero temperatures are feasible.

Keywords: Ab initio calculations; Monte Carlo simulations; Density-functional theory; Lithium; Clusters; Liquid; Pair correction functions

I. INTRODUCTION

Molecular dynamics (MD) and Monte Carlo (MC) simulations have been the major techniques for calculating finite-temperature properties of condensed-matter systems. In particular, ab initio MD simulations, which combine classical molecular dynamics with first-principles quantum theory, determine the atomic and electronic structures of a system simultaneously and accurately, as well as the temperature-dependent properties and time evolution of the system. Ab initio MD methods have been widely used to provide reliable and accurate information for various systems since Car and Parrinello first introduced their work in 1985.

On the other hand, ab initio Monte Carlo simulations, which employ first-principles quantum theory to calculate the potential energy of a system at each classical MC step, have so far been used only to a very limited extent. The majority of MC simulations for condensed-matter systems have been performed with empirical or semiempirical atom-atom interaction potentials, which are fitted to available experimental data or ab initio calculations. While the atom-atom potentials may work well in some cases, they do not provide reliable results in other instances. Since the quantum many-body effects and the detailed electronic structures cannot be included in empirical or semi-empirical potentials, first-principles theory would obviously be desirable for providing reliable energetics. The major concern with ab initio Monte Carlo simulations is the high computational demand. In particular, Monte Carlo simulations with empirical potentials often generate millions of sampling configurations. It is a formidable task to determine the potential energies of such a huge number of configurations with ab initio total-energy calculations. In recent years, however, several groups have performed ab initio MC simulations for very small clusters, and they have tried to reduce the number of sampling configurations (to typically $10^4$) [3,4,5,6,7,8,9,10,11,12,13,14]. Ishikawa et al. [4,5,7,10] applied ab initio calculations to the Monte Carlo simulated annealing algorithm [15]. They performed simulations for the Li$_6$ cluster, small hydrogenated lithium clusters (Li$_5$H, Li$_4$H$_2$, and Li$_7$H), and HCl(H$_2$O)$_n$ ($n = 3, 4$) clusters [3,12,14]. Other groups [3,5,7,10] used the ab initio molecular orbital (MO) method in classical MC simulations to study the interactions of ions and molecules with small water clusters. Structural and thermal properties of water dimers [11] and lithium clusters (Li$_4$, Li$_6^+$, and Li$_8$) [8,9,13] were also investigated with ab initio MC simulations.

We emphasize that all the previous ab initio MC simulations were limited to very small clusters. Since experimental data for the structures of the clusters at nonzero temperatures were not available, direct comparisons of the above simulations with experiments were not possible. Therefore, the accuracy and efficiency of ab initio MC simulations remain to be determined. Furthermore, the feasibility of the extension of ab initio MC simulations to bulk systems, which require models containing many more atoms than the small clusters, has yet to be tested. As a first application of ab initio MC simulations to bulk systems, we have chosen the liquid phase of bulk lithium. We have also investigated the structural properties of a cluster containing 16 lithium atoms at different temperatures. The simulated results for the lithium cluster are physically reasonable, and the structure of liquid lithium obtained from our ab initio MC simulations is in good agreement with the available experimental measurements. We also demonstrate that the computational demands of simulating liquid lithium with our ab initio
MC algorithms are comparable to those of \textit{ab initio} MD simulations.

The remainder of this paper is organized as follows. Section II describes our \textit{ab initio} MC algorithm. Section III describes the simulation results for a 16-atom lithium cluster (Sec. III A) and for liquid bulk lithium (Sec. III B), with a discussion of general aspects and possible improvements of our \textit{ab initio} MC algorithm in Sec. III C. Sec. IV contains our conclusion.

II. \textit{AB INITIO} MONTE CARLO ALGORITHM

To simulate the cluster of 16 lithium atoms, the atoms were put in a periodically repeated cubic supercell. The size of the supercell ($10 \times 10 \times 10 \text{ Å}^3$) was much larger than the concentrated cluster (see Fig. 1.). Bulk lithium was represented by a cubic box (supercell) with periodic boundary conditions containing 54 Li atoms. The size of the supercell ($10.698 \times 10.698 \times 10.698 \text{ Å}^3$ and $10.874 \times 10.874 \times 10.874 \text{ Å}^3$ for simulations at $T = 523 \text{ K}$ and $T = 725 \text{ K}$, respectively) was determined to match the experimental density of liquid lithium at the given temperature.

The simulations were performed in canonical ensembles. The standard Metropolis algorithm, \cite{17,18}, which provides an efficient approach for simulating the equilibrium properties of an atomic system at a given temperature, was used in our \textit{ab initio} MC simulations. Starting from a pre-selected initial configuration, the simulations repeated the following steps. (1) Given the current configuration $I$ of the system, a new configuration $J$ is generated by making a small random displacement of one randomly chosen particle (suitable for small systems) or random displacements of all the particles simultaneously (more efficient for systems containing a large number of particles). (2) Once the total energies ($E_I$ and $E_J$) of these two configurations are calculated, the acceptance probability of the new configuration $J$ is then determined as

$$P(J|I) = \min\left[1, \exp\left(\frac{(E_J - E_I)}{k_BT}\right)\right], \quad (1)$$

where $T$ is the temperature and $k_B$ is Boltzmann’s constant. (3) If the configuration $J$ is accepted, it serves as the current configuration of the next MC step, and $E_I$ is set equal to $E_J$. If the configuration $J$ is not accepted, the configuration $I$ and its energy $E_I$ are retained and used to start the next step. In this way, the system will eventually reach equilibrium and evolve toward a Boltzmann distribution. \cite{17,18} The resulting sequence of sampling configurations is then used to obtain the equilibrium properties of the system.

To calculate the total energy at each MC step, we employed an \textit{ab initio} pseudopotential density-functional-theory (DFT) method with a plane-wave basis set. All the \textit{ab initio} total-energy calculations were carried out using the Vienna \textit{ab-initio} simulation package (VASP). \cite{19,20,21} The electronic exchange-correlation effects were treated with the generalized gradient-corrected functionals given by Perdew and Wang. \cite{22,23} We adopted the Vanderbilt ultrasoft pseudopotential to replace the core electron- \cite{24,25} and used the conjugate-gradient technique for performing electronic relaxations.\cite{26} The total-energy calculations were conducted with 4 \textbf{k}-points in the three-dimensional Brillouin zone with a plane-wave energy cutoff of 120 eV. The convergence of the total energy differences between different configurations was checked for selected configurations with higher cutoff energies (up to 200 eV) and more \textbf{k}-points (up to 16 \textbf{k}-points), and the convergence was found to be within a few meV. Because we used a pseudopotential approach, the total-energy calculations for the configurations with too small distances between atoms, which result in core-core overlap, would give wrong total energies. We therefore defined a hard-core distance of 2.0 Å between any two Li atoms. Trial configurations with any interatomic distance smaller than the cutoff distance were always rejected in the MC simulations. This is physically reasonable because such configurations would have very high potential energies due to the strong core-core repulsion.

III. RESULTS AND DISCUSSION

A. Small cluster

Starting from the configuration with the atoms at ordered bcc positions (shown in Fig. 1), we first performed our \textit{ab initio} MC simulations for the cluster of 16 lithium atoms at temperatures of 1000 K and 5000 K. The MC moves in the simulations for the cluster consisted of random displacement of one randomly selected lithium atom at each MC step, with the maximum displacement adjusted to give an overall acceptance rate of about 50%. The chosen atom was displaced from its old position with equal probability to any position inside a sphere surrounding the old position, with the radius of the sphere being the maximum displacement. We obtained maximum displacements of 0.75 Å and 1.2 Å for the simulations at the temperatures of 1000 K and 5000 K, respectively. About 10000 MC steps were needed to reach the equilibrium state. Averages for calculating the structural properties were taken over the next 10000 sampling configurations after equilibration. It took approximately 1 minute of CPU time to obtain the total energy of each configuration with a single 375 MHz processor on an IBM SP3 supercomputer.

The pair correlation function $g(r)$ can be defined as

$$g(r) = \frac{2V}{N(N - 1)} \frac{1}{4\pi r^2} \left(\sum_{i,j<i} \delta(r-r_{ij})\right), \quad (2)$$

where $V$ is the system volume, $N$ is the number of particles, $r_{ij}$ is the distance between a pair of particles $i$ and $j$, and $\langle \cdot \rangle$ signifies averaging over the sequence of simulated
FIG. 1: The initial configuration of the cluster of 16 lithium atoms for the \textit{ab initio} MC simulations. The distance between two nearest-neighbor atoms is 2.96 Å, the same as that determined by our \textit{ab initio} total-energy calculations for the crystalline bulk Li. The size of the supercell is $10 \times 10 \times 10$ Å$^3$.

configurations. The pair correlation function is proportional to the probability of finding a pair of particles a distance $r$ apart, normalized by the square of the particle density such that it approaches unity in the limit of a uniform random distribution at the same density. It hence provides a measure of the local spatial ordering of the system. As described in Ref. 26, we determined $g(r)$ by dividing the range of $r$ ([2 Å, 10 Å]) into 400 intervals of length 0.02 Å ($\Delta r$), calculating every $r_{ij}$ for a given configuration, and recording in narrow bins $[r, r + \Delta r)$ the frequency of occurrence of the different particle separations. In this way a histogram was built up and finally normalized at the end of the scan over the configurations.

Figure 2 shows the pair correlation functions for the cluster at temperatures of 1000 K and 5000 K, respectively. It can be observed that there are two peaks in the pair correlation function at $T = 1000$ K. This indicates the existence of two atomic shells for the structure, and therefore the system locally has a liquid-like structure. At a temperature of 5000 K, however, there is no obvious second peak in the pair correlation function, and thus the system has a gas-like structure. Given that the boiling point of bulk Li is about 1600 K, the results from our simulations for the cluster of lithium atoms are physically reasonable. Note that the pair correlation function $g(r)$ at $T = 5000$ K does not approach zero when $r$ is close to 2.0 Å, indicating that the configurations with very small separations (smaller than 2.0 Å) between atoms would contribute to the ensemble averages. This reflects the gas-like structure of the system at $T = 5000$ K because the probability of occurrence of configurations with very small separations between particles is expected to be higher for a gas-like structure than for a liquid-like structure.

B. Bulk system

Following the simulations for the lithium cluster, we carried out \textit{ab initio} MC simulations for the liquid phase of bulk lithium. In contrast to the simulations for the lithium cluster, the Monte Carlo moves in the simulations for liquid lithium consisted of random displacements of all the atoms in the supercell simultaneously. The other aspects of the MC moves were the same as in the simulations for the lithium cluster. In order to achieve an overall acceptance rate of roughly 50%, we found that the maximum displacements at temperatures of 523 K and 725 K were 0.05 Å and 0.06 Å, respectively. We started our simulations at $T = 725$ K from an initial configuration with all the Li atoms at their bulk bcc positions. Roughly 8000 MC steps were required to reach equilibrium. Since the initial configuration was a perfect crystalline structure, far away from the equilibrium structure of the liquid phase of bulk lithium, it required relatively long MC runs to reach equilibrium. In contrast, the initial configuration for the simulations at $T = 523$ K was chosen as one of the equilibrium sampling configurations obtained in the simulations at $T = 725$ K. A reduced number of MC steps (5000 MC steps) was then required for the equilibration period. After equilibration at either temperature, we generated 5000 sampling configurations for averages.

Our simulations were carried out in parallel on four 375-MHz processors of an IBM SP3 supercomputer. The total-energy calculation at each MC step required approximately 150 seconds of CPU time. On average, the electronic relaxation converged within seven self-

FIG. 2: Equilibrium pair correlation functions $g(r)$ for a cluster of 16 lithium atoms at temperatures of 1000 K and 5000 K, obtained from our \textit{ab initio} Monte Carlo simulations.
FIG. 3: Equilibrium pair pair correlation function $g(r)$ for liquid lithium at a temperature of 523 K, calculated from ab initio Monte Carlo and ab initio molecular dynamics simulations (histograms). Also shown are experimental data measured by X-ray diffraction (circles).

The software (VASP) that we used for the total-energy calculations contains computations for the forces after the electronic relaxation converges. Such computations are not needed in ab initio MC simulations. Should VASP be modified to exclude the force calculations, the computational costs would therefore be decreased by approximately 10% – 15%.

For comparison, we also performed ab initio MD simulations\textsuperscript{19,20,21} for liquid lithium. Canonical ensembles were generated with the algorithm of Nosé\textsuperscript{27} and the time step was set to 1 fs. The initial configuration for the ab initio MD simulations at both $T = 725$ K and $T = 523$ K was the same perfect bcc structure as that for the ab initio MC simulations at $T = 725$ K. The equilibration periods extended over 5000 and 3000 time steps at $T = 523$ K and $T = 725$ K, respectively. Ensemble averages were calculated from the sampling configurations of the next 5000 steps. Each step required about 400 seconds of CPU time on an IBM SP3 supercomputer (in parallel on 4 processors), and the corresponding wall-clock time was roughly 100 seconds.

The pair correlation functions for liquid lithium at temperatures of 523 K and 725 K, obtained from our ab initio MC and ab initio MD simulations are shown in Fig. 3 and Fig. 4, respectively. For comparison, the experimental data for liquid lithium at $T = 523$ K obtained by X-ray diffraction\textsuperscript{28} are also shown in Fig. 3. We observe that the agreement between ab initio MC, ab initio MD, and experiment is very good. In particular, the three peaks determined from our ab initio MC simulations are essentially located at the experimental positions. The intensities of the peaks obtained from the ab initio MC simulations are also basically in agreement with those from both the ab initio MD simulations and the X-ray data. The small differences of the intensities of the peaks between the theoretical results and the experimental data may be due to the small size of our supercells. The experimentally observed temperature effects\textsuperscript{29} that is, broadening of the peaks and slight shifts to higher positions of the second and third peaks at the higher temperature are also well reproduced by our ab initio MC simulations (see Fig. 3 and Fig. 4).

C. General discussion

Finally, we discuss some general aspects and possible improvements of our ab initio MC algorithm. The computational costs of ab initio MC simulations are dominated by the ab initio total-energy calculations. Since efficient iterative schemes for ab initio total-energy calculations with a plane-wave basis set have achieved order $N^2$ ($N$ is the number of the atoms in the supercell) scaling for systems containing up to 1000 electrons\textsuperscript{23} the cost of simulations with our ab initio MC algorithm would increase essentially as the square of the number of atoms.

While the computational costs are comparable, our ab initio MC algorithm is not any faster than ab initio MD simulations. In particular, a larger number of steps are required to reach equilibrium in our ab initio MC simulations for bulk lithium than in the corresponding ab initio MD simulations. For the lithium systems that we have investigated, our ab initio MC algorithm produces the required sampling configurations at a speed similar to the ab initio MD simulations. In this respect, our ab initio MC simulations are as efficient as ab initio MD simulations.

In some cases, like simulations for strongly covalent
systems and situations involving volume changes, however, our \textit{ab initio} MC algorithm may not be efficient due to the low acceptance rate of the traditional MC moves. In such situations, therefore, our \textit{ab initio} MC algorithm needs to be improved. A cluster MC move scheme proposed by Lee and Swendsen\cite{Lee79} may be used for this purpose. We have tested the cluster MC move scheme for the bulk lithium systems that we investigated with traditional MC moves as described in Sec. III. We found that the use of cluster MC moves in \textit{ab initio} MC simulations provided almost the same performance as the traditional MC moves, indicating that both kinds of MC moves might work equally well for metallic systems. It is also possible that the relative performance of different kinds of MC moves somewhat depend on the number of particles included in the supercell. It would be interesting to see if the cluster MC move scheme would be more efficient than the traditional MC approach in simulations for systems containing large numbers (for example, hundreds) of particles. In the cases where the boundary conditions or/and the volume of the simulated cells change, which result in a low acceptance rate with the traditional MC moves, the cluster MC move scheme has been shown to increase the acceptance rate significantly.\cite{Lee79} We also expect that the cluster MC move scheme would be more efficient than the traditional MC move method in simulations for systems containing strong covalent bonds. Another approach, a hybrid of MD and MC simulation in which trial MC moves are generated with classical MD calculations in classical Monte Carlo simulations provides structures of liquid lithium in agreement with experimental measurements and \textit{ab initio} MD simulations. Our work also demonstrates that \textit{ab initio} MC simulations for condensed-matter systems with up to several tens of atoms (and possibly with more than a hundred atoms) are feasible and may provide an alternative way to investigate finite-temperature properties of such systems.

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