Structures and Local Electronic States of Dislocation Loop in 4H-SiC via a Linear-Scaling Tight-Binding Study

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The atomic- and electronic-level structures of a dislocation loop and a stacking fault in 4H-SiC crystal are investigated by using large-scale tight-binding (TB) molecular-dynamics simulation. We employ a linear-scaling TB method implemented on a parallel computer in order to accelerate the 9,600-atoms calculation which is required for such a nanoscale simulation. We find that the initial configuration that involves unstable C-C networks around the dislocation loop is relaxed to a structure having six-membered rings, and that the distribution of electron populations is inhomogeneous on the loop. The local electronic density of states shows two peaks in the bulk band gap, where one of these peaks may correspond to the defect state observed in EBIC and CL experiments.

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1. Introduction

Silicon carbide (SiC) is a wide band gap semiconductor suited for applications in high-power, high-temperature, and high-frequency electronics. Among various polytypes of SiC, 4H-SiC has higher bulk mobility and smaller anisotropy, and thus is very promising for the applications unreached by silicon technology. Recent advances in 4H-SiC homoepitaxy have made it possible to eliminate harmful micropipes and to reduce dislocations. For further miniaturization of the devices, however, microscopic properties of defects have to be known prior to designing fabrication processes.

In the device applications, formation and subsequent development of stacking faults (SFs) in the active region of semiconductors are known to be one of the reasons for the degradation phenomenon in the I-V characteristics. The formation of SFs by dissociating basal plane dislocations (BPDs) is often observed under forward bias stressing. Recently, Electron-beam-induced current (EBIC) and cathodoluminescence (CL) study on 4H-SiC have shown that BPDs are easily dissociated into two partial dislocations with a SF between them under the electron-beam irradiation.

In this paper, we report a study on the atomic- and electronic-structure of a dislocation loop and a stacking fault in 4H-SiC. We focus on the local atomic structure similar to that reported in the aforementioned experiment and investigate by using large-scale tight-binding (TB) molecular-dynamics simulation. Using this computer simulation, we analyze the atomic structure, Mulliken’s local electronic populations, and the electronic density of states (DOS). The interplay between these characteristics and experimental observations is also discussed.

2. Computational Method

Molecular-dynamics (MD) simulations were performed using a velocity-Verlet algorithm. For the calculations of the electronic parts of the energy and the forces, we employed an \( O(N) \) tight-binding method with a paralle-
calculations of $E_{\text{tot}}$ and the electronic forces. Also, the truncation at the physical distance between atomic sites can make the data structures localized, hence a scalable parallelization can easily be achieved in the present algorithm. We employed a spatial domain-decomposition technique for calculating matrix-matrix multiplication. In this algorithm, data transmission between neighboring nodes is performed only for near-boundary atoms and matrix elements because of the spatially localized nature of the density matrix. Internode data transmission is performed using the standard MPI library. In the present study, we have performed the parallel TBMD simulations on our 32-cores parallel PC cluster. The criterion for convergence of the Chebyshev expansion was set to be $1.0 \times 10^{-12}$ in relative value of each element of the $F$-matrix and the pseudo-electronic temperature was chosen to be 0.1 eV.

Parameterization of matrix elements in the semi-empirical TB Hamiltonian is an essential ingredient for TBMD simulations. Among a number of parameterizations of the TB Hamiltonian for SiC systems, we have chosen Mercer’s set of the parameters which is based on an $sp^3$ orthogonal basis set. It also includes environment-dependent contributions to the onsite energies through intra-atomic terms. These terms give important contributions to variation of the electron transfer between Si and C atoms, especially for inhomogeneous systems, although it does not include explicitly the Coulomb interaction between ions. The original parameterization by Mercer, however, leads to some discrepancies with experimental data such as the lattice constant.

### 3. Structure of Dislocation Loop and Stacking Fault in 4H-SiC

4H-SiC crystal has the periodic structure containing four (ABCB) layers per unit cell, as shown in Fig. 1(a). In the present study, the system in a simulation cell is constructed by duplicating the above structure, and contains 9,600 atoms. The simulation cell has dimensions of 3.95 nm long, 4.30 nm wide and 5.06 nm depth. The periodic boundary condition is applied to the system simulated. In the defect structure proposed from the EBIC and CL experiments, a basal dislocation of the Burger’s vector $\mathbf{b} = 1/3[11\bar{2}0]$ is dissociated into two partials with $\mathbf{b}_1 = 1/3[10\bar{1}0]$ and $\mathbf{b}_2 = 1/3[01\bar{1}0]$, accompanying an extension of a SF regions between these partials. To model the defect structure, some atoms within a rhombohedral region surrounded by four (1000) and two (0001) planes are shifted along $\mathbf{b}_1$. This creates partial-dislocation loop on (0001) planes with two symmetric Burger’s vectors, $\mathbf{b}_1$ and $-\mathbf{b}_1$, as depicted in Fig. 1(b). We adopt this model, instead of creating both $\mathbf{b}_1$ and $\mathbf{b}_2$, because having the full dislocation in the simulation cell requires extra (or missing) atomic layers in the loop regions. The existence of such layers induces the large strain in entire region of the cell, which is turned out to be fatal in evaluating the net effect of the partials and SFs individually on the electronic structure (e.g., the local DOS outside the loop region may be shifted significantly due to the strain).

In this paper, the areas where atoms reside on and near the partial dislocation cores, accompanied by left-pointing arrows in Fig. 1(b) are denoted as “high-density area”, and the area on and near the other dislocation cores are “low-density area”, respectively.

### 4. Relaxed Structure

Figure 2 shows the local structure in the high-density area before and after molecular-dynamics relaxation. While anomalous C-C bonds, which connect electronically having finite values of the bond order between them, are observed before the relaxation (Fig. 2(a)), those bonds disappear after the relaxation (Fig. 2(b)). Furthermore, the atomic structure around dislocation loops are composed of a six-membered ring similar to that of bulk structure, as shown in Fig. 2(c). At the low-density area, on the other hand, there are no significant changes of the atomic structure by the relaxation.

### 5. Mulliken’s Charge Analysis

Figure 3 shows effective charge transfers of atoms in the system after the relaxation. The effective charge transfer is calculated from the Mulliken’s method for the electron population at each atomic site. The color of each atom represents deviation of the charge from its bulk value. Compared with the result of bulk state, the silicon in the low-density area tend to lose electrons, while carbons in the area tend to attract them, indicated by small arrows in the left of Fig. 3, for example. On the other hand, the carbons in the high-density area lose electrons. In addition, the silicons on the [11\bar{2}0] dislocation line tend to lose electrons, while others...
get them, indicated by small arrows in the right of Fig. 3, for example. These results indicate that the electrons in the high-density area have tendency of being displaced along the direction of large arrows in Fig. 3.

6. Density of States

We have evaluated the electronic DOS of the 4H-SiC system which contains dislocation loops using a kernel polynomial method. Figure 4 shows a comparison between the DOS in the bulk crystal, in the system with the defects before the relaxation, and that after the relaxation. As shown in Fig. 4(a), the bulk gap $E_g$ obtained from this TB model is 3.2 eV, which is very close to the experimental value 3.3 eV. In the case of the system with dislocation loops, the gap state appears before the relaxation, as indicated by an arrow in Fig. 4(b). After the relaxation, however, that state disappears, as shown in Fig. 4(c). To find causes of this phenomenon, the local DOSs in four areas indicated in Fig. 5 are calculated.

Figure 6 shows the local DOSs of two areas in the high-density area before and after the relaxation. In these Figs., the gap state appears at 1.3 eV below the conduction band edge in the bulk gap region before the relaxation, and it disappears after the relaxation. It is assumed that this change is related to the change in bonding network induced by the relaxation as discussed in Sec. 4, that is, the peak in the gap region is related to the abnormal C-C bonds formed before the relaxation. Thus, this peak disappears after the relaxation with the disappearance of those abnormal bonds.

Similarly, Fig. 7 shows the local DOSs of two areas in the low-density area before and after the relaxation. As shown in Figs. 7(a) and (b), there are no peaks in the gap in the area 3 both before and after the relaxation. On the other hand, in the area 4, the gap state appears at shallow level in the gap both before and after the relaxation, as shown in Fig. 7(c) and (d). As in the case of the high-density area, this peak may be attributed to bonding states in the areas. There are dangling bonds in the area 4, whereas there are not in the area 3. In addition, those structures in the area 4 still exist even after the relaxation. This implies that the peaks observed in Fig. 7(c)
Fig. 6 Local DOSs of the high-density area. Inset is the close-up of highlighted area in the figure: (a) Area 1 before the relaxation; (b) area 1 after the relaxation; (c) area 2 before the relaxation; (d) area 2 after the relaxation.

Fig. 7 Local DOSs of the low-density area. Inset is the close-up of highlighted area in the figure: (a) Area 3 before the relaxation; (b) area 3 after the relaxation; (c) area 4 before the relaxation; (d) area 4 after the relaxation.
and (d) are induced by dangling bonds in the area 4. Furthermore, although the present model takes only the $\pm b_1$ partial dislocation loops into account, the area 4 can be considered as the part of the dislocation loop observed in EBIC and CL experiments. Therefore, the peak appeared 0.1 eV below the conduction band edge may correspond to the defect state observed 0.3 eV below the conduction band in EBIC and CL experiments.6)

7. Conclusions

Using a linear-scaling TB simulation, we investigated local structure and electronic states of partial dislocation loop and stacking faults in 4H-SiC. We find that the initial configuration that involves unstable C-C networks around the dislocation loop is relaxed to a structure having six-membered rings, and that the distribution of electron populations is inhomogeneous on the loop. The local electronic density of states shows two peaks in the bulk bandgap, where one of these peaks may correspond to the defect state observed in EBIC and CL experiments.

As discussed briefly in Sec. 3 and 6, the present model deals only one type of the partials proposed in the experiments. Also, we have observed a weak appearance of a gap state in the area containing no defects in the system. This may be attributed to the interaction between the defects in the simulation cell and those in the image cells. Analysis with larger simulation cell is therefore inevitable in order to investigate the local electronic states in the system with full types of dislocations and SFs more thoroughly. The simulations with several tens of thousands of atoms in a MD cell are in progress.

Acknowledgments

This work was supported by Grant-in-Aid for Scientific Research on Priority Areas “Nano Materials Science for Atomic Scale Modification 474” from MEXT of Japan.

REFERENCES