Atomic structure of dislocation kinks in silicon

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We investigate the physics of the core reconstruction and associated structural excitations (reconstruction defects and kinks) of dislocations in silicon, using a linear-scaling density-matrix technique. The two predominant dislocations (the 90° and 30° partials) are examined, focusing for the 90° case on the single-period core reconstruction. In both cases, we observe strongly reconstructed bonds at the dislocation cores, as suggested in previous studies. As a consequence, relatively low formation energies and high migration barriers are generally associated with reconstructed (dangling-bond-free) kinks. Complexes formed of a kink plus a reconstruction defect are found to be strongly bound in the 30° partial, while the opposite is true in the case of 90° partial, where such complexes are found to be only marginally stable at zero temperature with very low dissociation barriers. For the 30° partial, our calculated formation energies and migration barriers of kinks are seen to compare favorably with experiment. Our results for the kink energies on the 90° partial are consistent with a recently proposed alternative double-period structure for the core of this dislocation. [S0163-1829(98)10617-3]

I. INTRODUCTION

Dislocations are of fundamental importance in the physics of semiconductors, both from a mechanical and from an electronic point of view. They are the carriers of plasticity in crystals and act as trapping and scattering centers for electronic carriers. A wealth of experimental information is available about the properties of dislocations in tetrahedrally bonded semiconductors.1–7 In Si, the predominant slip systems are the 60° and the screw dislocations oriented along [110] directions in a {111} slip plane. Both are known to occur in the glide configuration and to dissociate into pairs of partial dislocations bounding a ribbon of intrinsic stacking fault.1–3 Dissociation lowers the strain energy and is made energetically favorable by the low energy of the stacking fault in this material. (Evidence indicates that the above is also true in the case of germanium and for III-V and II-VI semiconductors.1–3) The resulting 90° and 30° partials are believed to undergo core reconstruction, which eliminates the unsaturated bonds, thus restoring the fourfold coordination of the atoms at the cores. This picture is consistent with the low density of dangling bonds, as suggested by EPR measurements.1,2

Dislocation motion occurs by nucleation and propagation of kinks along the dislocation line. Due to thermal fluctuations or the action of an applied stress, double kinks can be nucleated along the dislocation line. When these reach a critical separation, dissociation occurs and the individual kinks propagate in opposite directions, thus generating a displacement of a dislocation segment.8 A detailed understanding of the atomic-scale structure of the kinks and the barriers associated with their motion is thus of the greatest importance.

In semiconductors, according to the theoretical model proposed by Hirth and Lothe (HL),8 double-kink nucleation and the high Peierls potential barrier to kink motion control the rate of dislocation propagation. This is to be contrasted with the case of metals, where kinks experience a very low barrier to motion, and the rate is controlled by nucleation only. The HL model is often used in the interpretation of dislocation mobility experiments, although the occurrence of such high Peierls barriers has been questioned by some authors.9 Furthermore, an alternative theoretical model has been proposed in which dislocation motion is controlled by the pinning of kinks by obstacles distributed along the dislocation line.10,11 Recent experimental evidence suggests that the barriers are indeed high, but experiments cannot clearly decide between these two theoretical models.4–7 A complete microscopic picture is still lacking. Related issues, such as the dependence of dislocation mobility on doping and the photoplastic effect in semiconductors,1 would also benefit from a better understanding of dislocation structure at the atomic level.

On the computational side, large-scale problems of this nature have been mostly studied by using classical interatomic potentials. Such studies are not always reliable, since these potentials are often unable to reproduce effects of intrinsic quantum-mechanical nature such as bond reconstruction and Peierls or Jahn-Teller symmetry breaking. For example, while the Stillinger-Weber12 potential has been used to study the core reconstruction and kinks of the 30° partial,13 it fails to reproduce the spontaneous symmetry-breaking core reconstruction of the 90° partial from the symmetric “quasi-fivefold” reconstruction.14,15 A proper quantum-mechanical description of the electronic structure is clearly needed. One is thus led to consider tight-binding (TB) and ab initio methods.

Recent ab initio and TB theoretical works have concentrated on such issues as the core reconstruction of the 90° partial14,16 and the elastic interaction between dislocations of

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a dipole in the shuffle \cite{17} and glide sets\cite{16}. Using a relatively small supercell, one first-principles study has determined a kink mobility barrier in the 30° partial,\cite{18} but only one kink species was studied, out of a very rich variety characteristic of this system. As will become clear from the conclusions of the present work and from Ref. 13, the formation and migration energies of other kinks are needed for a proper comparison with the experimental results. An important recent development is our prediction,\cite{19} on the basis of classical-potential, tight-binding, and \textit{ab initio} calculations, that the reconstruction of the 90° partial in Si is not the above-mentioned symmetry-breaking structure, as had generally been accepted in the theoretical literature.\cite{14-16,20-28} Instead, we proposed a double-period (DP) reconstruction whose core structure is reminiscent of that of a double kink of minimal length that would form on the core of the original single-period (SP) structure.\cite{19} Cluster calculations on kinks and solitons\cite{29} in the SP reconstruction of the 90° partial have also been reported.\cite{23,25,27,28} These calculations have identified many of the basic types of defects in this system, but must be taken at a semiquantitative level, since they suffer from the lack of coupling of the defect local strain fields with the lattice elastic fields.

To address properly the issues related to dislocation mobility, a comprehensive study of dislocation kink structure and dynamics would require the use of very large supercells, for which the application of \textit{ab initio} techniques is still computationally prohibitive. In view of this, the natural choice is the application of more efficient quantum-mechanics based methods to study the electronic and structural excitations in the dislocation cores. In this work, we employ the tight-binding-total-energy (TBTE) Hamiltonian of Kwon and collaborators\cite{30} to carry out a detailed atomistic study of the atomic structure of both the 30° and the 90° partial dislocations in Si. To make these calculations tractable, we use the linear-scaling or \textquotedblleft \textit{O}(N)\textquotedblright \ method of Li, Nunes, and Vanderbilt\cite{31} to solve for the electronic-structure contribution to energies and forces, enabling us to treat system sizes up to 10^5 atoms easily on a workstation platform. Our work addresses some of the fundamental issues associated with these two systems. More specifically, we address the ground-state structural properties of the dislocation cores and of defects in the core, such as kinks and reconstruction defects (RD), as well as energy barriers to motion of the various defects.

In this work, when considering the 90° partial, we will discuss only the SP reconstruction. Despite the fact that this is not the correct ground state for this dislocation in Si, we hope that understanding this somewhat simpler system will help us in the study of the myriad of defects in the more complicated ground-state DP reconstruction, to which the former is related.\cite{19} Moreover, we should keep in mind that the 90° partial is equally important in other materials, such as germanium (Ge), diamond (C), and the III-V and II-VI semiconductors.\cite{1-3} Preliminary calculations,\cite{19} using a Keating model,\cite{32,33} indicate that in C the SP reconstruction is more likely to be lower in energy, while Ge, like Si, would prefer the DP structure. More accurate calculations are needed to reveal which of the two reconstructions would be favored in each case. Therefore, the study of the SP structure is still important from a theoretical point of view.

The paper is organized as follows. The next section contains the technical details of the calculations we performed. In Sec. III, we discuss our results for the core reconstruction and related defects in the 30° partial dislocation. Our main results for the SP reconstruction of the 90° partial are described in Sec. IV. Finally, in Sec. V we summarize the main conclusions and results, and compare our kink energies and barriers with the available experimental results. In particular, we will argue that our results appear to be consistent with the HL theory of dislocation glide.

II. TECHNICAL DETAILS OF THE CALCULATIONS

We use the TBTE parameters of Kwon \textit{et al.},\cite{30} which describe well the acoustic-phonon modes and elastic constants of Si, thus being adequate to describe the strain fields associated with the dislocation cores and related defects. Owing to its good transferability between different crystal structures, ranging from diamond to fcc, this Hamiltonian is also expected to give a good description of the coordination defects in the present study. The \textit{O}(N) method of Li \textit{et al.}\cite{31} is used to solve for the electronic structure contribution to the energies and forces. For the density matrix, we initially work at a real-space cutoff $R_c = 6.2$ Å on the range of the density matrix used in the tests presented in Ref. 31 for the \textit{O}(N) method. In a second stage, we improve the convergence of our results by further relaxing the ionic positions and the electronic structure with a larger cutoff value of $R_c = 7.3$ Å. The numerical minimization of the \textit{O}(N) functional was carried out by the conjugate-gradient algorithm, with the internal line minimization performed exactly. To obtain the right number of electrons, the chemical potential is adjusted iteratively, in each case. Usually, this procedure has to be repeated only at the initial steps of the structural relaxation procedure, after which the chemical potential converges to the adequate value and remains constant. Ground-state structures were computed by allowing all atomic coordinates to relax fully (average forces less than 0.1 meV/Å).

Our supercells are chosen with the dislocation direction (corresponding to a [1 0 0] crystalline direction) lying on the $y$ axis. The glide plane (which contains a stacking fault) is normal to the [1 1 1] direction and coincides with the $xy$ plane of the cell. (Figure 1 shows the glide plane of the 30° partial dislocation, with the crystalline directions indicated.) The $z$ direction of the cell is thus parallel to the [1 1 1] direction. Each supercell contains two dislocations having opposite Burgers vectors (a dislocation dipole), which allows us to use periodic boundary conditions. Supercell vectors are chosen such as to array the dislocations in a quadrupole configuration, as suggested in Ref. 14, to avoid the spurious shear strains associated with the minimal dipole cell.

To ensure the convergence of our calculations with respect to supercell size, we used three different cells, containing 216, 264, and 308 atoms, respectively, for the simulation of the reconstructed core of the 30° partial dislocation. Each cell corresponds to a slab of atoms at a 60° angle with respect to the dislocation direction, including twice the lattice period in that direction, to allow for the period-doubling reconstruction of the 30° partial. The two parameters characterizing the geometry of each cell are the separation between the two dislocations in the glide plane (i.e., the width of the stacking fault) within a given unit cell, and the distance be-
which produced a change of only 

\[ R_c \] computed using a larger cutoff (\( c \) for the density matrix, kink and core energies were 

Therefore, we consider our calculations to be converged 

kink separation 

defect interaction, this kink was studied with a larger kink-

308-atom slab was only 

\[ \frac{1}{2} \]

direction. The change in defect energy with respect to the 

z direction. In 

our calculations, these distances are, respectively, 15.0 and 
18.8 Å for the 216-atom cell, 18.3 and 18.8 Å for the 264-
atom cell, and 18.3 and 21.9 Å for the 308-atom cell.

The supercells for the computation of defect energies are 
obtained by repeating the core slabs several times along the 
dislocation direction. The defect energies we quote are re-
ferral to the corresponding supercell containing defect-free 
(but reconstructed and fully relaxed) dislocations. For the 
kinks in the 30° partial, each of the core slabs were repeated 
times (two and a half times for the RD, and three and a 
half times for the kink-RD complexes) along the dislocation 
direction, so that the defect-defect separation along the line 
was 19.2 Å or larger, depending on the type of defect. Be-
low, we describe the procedure we used for the computation 
of defect energy barriers. Because of the higher computa-
tional demands involved, in this case we employed only the 
smaller cells (three times the 216-atom slab for kinks and 
two and a half times the same slab for the RD).

Table I in the next section illustrates the convergence of 
our results with respect to dislocation separation. As a further 
check, we also computed the energies of the core and of one 
the kinks [the left kink (LK), as described below] with an 
even larger slab, consisting of 600 atoms for the recon-
structed core (1800 atoms for the defect). In this case, dislo-
cation distances are 24.9 Å in the \( xy \) plane and 31.4 Å in the 
z direction. The change in defect energy with respect to the 
308-atom slab was only \( \sim 0.02 \) eV. To test the effect of 
defect interaction, this kink was studied with a larger kink-
kink separation (with the smallest slab repeated eight times), 
which produced a change of only \( \sim 0.01 \) eV in the energy. 
Therefore, we consider our calculations to be converged within 
0.03 eV with respect to core-core and defect-defect 
interactions. To estimate the error involved in our choice of 
cutoff for the density matrix, kink and core energies were 
computed using a larger cutoff (\( R_c = 8.1 \) Å). The kink for-
mation energy changed by only \( \sim 0.06 \) eV, which justifies 
our choice of cutoff. From these results we can also estimate 
that our defect energy barriers are converged within 0.3 eV.

The supercells used for the study of the 90° partial, are as 
described in Refs. 15,19. In this case, despite the fact that we 
are only interested in the qualitative nature of our results, our 
values are well converged, with dislocation distances on the 
order of 26.6 Å, and defect-defect separations of at least 
\( \sim 13.4 \) Å. (As in the case of the 30° partial, barriers are 
computed using smaller cell sizes, corresponding to a dislo-
cation separation of 13.3 Å.)

Barrier energies were calculated by identifying the 
2\( N \)-dimensional configuration-space vector \( R_{12} = R_2 - R_1 \) 
pointing from one equilibrium position \( R_1 \) of the defect to a 
neighboring position \( R_2 \), and defining a reaction coordinate 
\( Q = R - R_{12} \) measuring the progress from \( R_1 \) to \( R_2 \). Then, for 
a series of values of this coordinate, we computed the energy 
with this coordinate fixed and all others fully relaxed. This 
approach is efficient in simple cases, but we find that it often 
fails to converge to the saddle-point configuration when the 
reaction path\(^ {35} \) makes sharp angles with respect to \( R_{12} \). 
In these cases, we can usually find two configurations, \( R_{1i} \) and 
\( R_{1i}' \) near the saddle point, with nearly the same value of \( Q \) 
but with opposite forces along \( R_{12} \). By exploring the space 
spanned by \( R_{12} \) and (\( R_{1i} - R_1 \)) while allowing all other coor-
dinates to relax, we were able to determine the energy bar-
riers with good accuracy for all cases studied (average forces 
less than 1.0 meV/Å).

### III. THE 30° PARTIAL DISLOCATION

#### A. Core reconstruction

In Fig. 1(a), a top view of the atomic structure of the 
unreconstructed 30° partial in the glide plane is shown. The 
shaded area represents the stacking fault, and the dislocation 
line is indicated by the boundary between shaded and un-
shaded areas. The crystalline directions are also displayed. 
Atoms shown as white (black) are above (below) the glide 
plane; each atom is bonded to another either above or below 
it, and these are not shown in the picture. Thus, fourfold 
coordinate atoms have three of their bonds in the plane of the

| Table I. Formation energy of defects in the 30° partial dislo-
ca tion, in eV. Defect energies are referred to a defect-free dislo-
cation core. TB results for three supercell sizes are shown. For the 
PSD, supercells contain 5/6 of the number of atoms shown. For the 
LK, in the third column we indicate in parenthesis the formation 
energy computed with a 1800-atom cell. The fourth and fifth col-
umns contain Keating energies for TB-relaxed structures, computed 
for the largest cell, with the QC (Ref. 38) and the KT (Ref. 32) 
parameter sets. |
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FIG. 1. (a) Unreconstructed core of the 30° partial dislocation, 
viewed from above the (111) slip plane. Shaded region indicates 
stacking fault. Black (white) atoms lie below (above) the slip plane. 
(b) Same view of the double-period reconstructed structure. Cryst-
talline directions are also shown.
figure. The atoms at the core of the defect are threefold coordinated, with a dangling bond lying nearly parallel to the dislocation line. In Fig. 1(b) we show a reconstruction in which the fourfold coordination of the atoms at the core is restored by atoms bonding in pairs along the line, leading to a doubling of the period in that direction. This reconstruction is well accepted as being the ground state of the 30° partial, and has been discussed theoretically by other authors. In Ref. 13, it was found to be 0.21 eV/Å lower in energy than the unreconstructed structure, using a Stillinger-Weber potential. A Tersoff potential calculation obtained a value of 0.14 eV/Å, while we find a higher value of 0.36 eV/Å for the reconstruction energy.

A look at the distribution of bond lengths for this structure shows that the reconstruction is indeed strong, with maximum bond-length deviations of only 3.0% (maximum and minimum bond lengths are 2.42 and 2.31 Å, respectively) with respect to Si bulk values (2.35 Å). The core energy is mostly due to the strain associated with bond-angle distortions at the core of the defect, with bond angles ranging between ~90° and ~126° (109.5° is the bulk value). No mid-gap levels are expected for this structure, in accordance with the EPR evidence.

A rich variety of core defects is associated with this reconstruction, including kinks and RD's, and complexes of these basic types. A very extensive study of these defects is found in Ref. 13, including structural features and energetics under a Stillinger-Weber potential. To a large extent, our study of this specific dislocation relies on this previous study, adding to it the benefits of a quantum-mechanical treatment of the electronic structure. More specifically, the defects considered in this work are the ones identified in Ref. 13. As we proceed, it will be seen that some of our results differ qualitatively from those in Ref. 13, and also that we find a better agreement with the experimental results.

### B. Phase switching defect (PSD)

Figure 2(a) shows a RD associated with the core of the 30° partial. We shall refer to this defect as a phase switching defect (PSD). The existence of such defects has been hinted at since the realization that the core of the partials might undergo reconstruction. We computed the energy of a fully relaxed PSD by repeating the atom slabs five times along the (110) direction, and introducing one PSD in each dislocation line. Our value for the PSD energy formation is 1.32 eV (see Table I), which is somewhat higher than the value of 0.81 eV obtained in Ref. 13. We believe our result to be more reliable, given the quantum-mechanical nature of our approach, in particular for a defect containing a dangling bond. A previous study based on a combined Keating linear combination of atomic orbitals (LCAO) approach has estimated the PSD energy to be a few tenths of an eV less than 2 eV, in agreement with our value. To a first approximation, this defect can be understood as a p dangling-bond defect, which indicates that a formation energy on the order of 1 eV (roughly the bond-breaking energy in bulk Si) is to be expected. The exact value is determined by the relaxation of the atoms surrounding the defect.

We also computed the migration barrier for the propagation of the PSD along the dislocation direction. The relaxed structure of the barrier (saddle-point) configuration is shown in Fig. 2(b). In this case, the symmetry between adjacent positions of the defect along the line indicates that the saddle-point configuration is at the halfway position. It was somewhat surprising to find that even in this case, we had to resort to a two-dimensional reaction coordinate as described in Sec. II. Our saddle-point configuration, with an energy barrier of 0.3 eV, is very similar to that in Ref. 13. In can be seen that the atom at the center becomes fivefold coordinated, which leads to a smooth process of bond substitution as the PSD propagates to the right. This explains the low-energy barrier involved in this process.

### C. Kinks

The period doubling of the reconstructed core gives rise to a multiplicity of kinks in this system. Two distinct families of such defects appear, depending on whether the dislocation “kinks” to the left (Fig. 3) or to the right (Fig. 4). The period doubling of the core introduces a choice of phase of the core reconstruction both ahead of, and behind, the kink. Of the four configurations generated in this way, two of them (those necessarily containing a coordination defect) will be classified as PSD-kink complexes, and will be considered in Sec. III D. The remaining two configurations will be classified as “pure” kinks and are considered here. The two left kinks LK and LK’ are shown in Figs. 3(a) and 3(b), while the two right kinks RK and RK’ are shown in Figs. 4(a) and 4(b).

The energies for each type of kink were computed using the TBTE Hamiltonian, as well as with a classical Keating model, with two different sets of parameters, one being the set introduced by Qian and Chadi (QC), and the other, the original set proposed by Keating (KT). In Table I, we show the TBTE results for each of the three slabs described in Sec. II, along with the Keating results for the 924-atom cell. For one of the kinks, the energy computed with a 1800-atom slab
is also shown in parenthesis; note the convergence of these results with respect to cell size.

In the following discussion, we use the Keating model only as a tool to enable us to examine the local-strain contributions to the energy of each defect. We do not intend to draw any quantitative conclusions from our Keating results. For this reason, what we compute are Keating energies for the TB-relaxed structures. It is known that the KT set of parameters describes well long-wavelength elastic deformations, while overestimating short-wavelength deformations such as those that are present at the dislocation and kink cores.\textsuperscript{3,39} On the other hand, by reducing the ratio between the bond-bending and bond-stretching forces to one third its original value,\textsuperscript{3,39} short-wavelength deformations are well described, at the expense of underestimating the long-wavelength elastic modes.\textsuperscript{3,39} Thus, we should expect the QC set of parameters to give a more reliable description of the short- to medium-range strain of the kinks. As will become clear from our results, despite the model shortcomings at the quantitative level, the qualitative trend of kink energies is correctly described. Thus, the Keating model should suffice for our analysis of the strain fields associated with these defects.

1. Left kinks

The left kinks LK and LK’ are shown in Fig. 3, together with the saddle-point configuration for the LK→LK’ translational motion. The energies, as given in Table I, show that reconstruction produces low energy kinks in this case, as compared to the energy of the unreconstructed PSD defect. At first sight, the formation energy for these reconstructed defects is expected to be mostly associated with the local strain at the kink cores. The Keating-model results can give us a qualitative understanding of these local-strain effects. The LK is found to add little additional strain to that imposed by the core reconstruction itself, with a formation energy that varies from slightly negative to small and positive, depending on the parametrization. On the other hand, the LK’ is found to have a Keating formation energy of 0.44 and 1.27 eV, for the QC and KT parameters, respectively. Despite the strong dependence of the kink energies on the choice of parameters, we note that the trend in energies for the two left kinks is in qualitative agreement with the TB results. This is actually true for all four kink types, as can be seen by looking at the kink energies for the two sets of parameters in the fourth column of Table I.

For the LK saddle-point configuration in Fig. 3(c) we computed an energy barrier of 1.52 eV. This result is in very good agreement with experimental estimates. In our concluding section, we will discuss more extensively the significance of our results in light of the available experimental evidence. Here, we note that such a high barrier can be understood by the presence of severe bond-bending and stretching distortions at the core of the defect, along with the presence of malcoordinated atoms. Bond angles as small as 50.4° are found, as well as bonds stretching to 2.80 Å.

FIG. 3. Core structure of the left kinks in the 30° partial, and associated transition state. Kink notation is explained in the text. (a) LK. (b) LK’. (c) Transition state for the LK→LK’ transformation.

FIG. 4. Core structure and transition state of right kinks in the 30° partial. Kink notation is explained in the text. (a) RK. (b) RK’.
(c) Transition state for the RK→RK’ transformation.
where the Keating energy $E_i$ is the sum over all atoms within a distance $R$ from the dislocation core. Corresponding core energy is subtracted to yield defect energies.

### 2. Right kinks

Shown in Fig. 4 are the two kinks of the right family, RK and RK', together with the saddle-point corresponding to the RK$\rightarrow$RK' reaction. Despite the fact that both kinks are fully reconstructed, the formation energies of 1.24 eV for RK and 1.85 eV for RK' are surprisingly high. No single structural feature of the right kinks could be traced in order to explain the unexpected formation energies. The minimum and maximum distortions of bond lengths and angles do not vary drastically among the four kink types.

To help us better understand these results, we observe that the Keating energies can be decomposed in an atom-by-atom basis. Bond-bending energies, associated with changes in the angle between two bonds, are assigned to the vertex atom, and half of the bond-stretching energy of a given bond is assigned to each of the two participating atoms. To examine the nature of the strain fields associated with each kink type, using our largest cells (924 atoms), we looked at these atomic energies integrated over shells of atoms defined by their distance from the core of the defect. (Since our supercells contain two cores and thus two defects, we always choose the shortest distance to a defect.) The integrated energies are then defined by

$$E_i^d(R) = \sum_{R_i=R} E_i^d(R_i) - E_i^c(R_i),$$

where the Keating energy $E_i^d(R_i)$ of each atom in a correspondent kink-free supercell (containing only the dislocation dipole) is subtracted, and we sum over all atoms within a distance $R$ from the kink. The results are shown in Fig. 5, for the QC parameters. We see that the kink energies are determined by the medium-range behavior of the associated strains. At short range ($R<3.0$ Å) the LK is actually the highest in energy. As we advance away from the core of the kinks, the energies only approach their final relative values at a distance of about $R=10.0$ Å. Almost exactly the same qualitative pattern is obtained for the KT set of parameters (not shown), despite the fact that the final energies are different.

As in the case of the left kinks, a look at the RK saddle-point configuration shows that the rather high migration barrier of 2.03 eV for the right kinks is associated with the drastic bond distortions and malcoordination of atoms at the core. Note that this barrier is substantially higher than the 1.52 value we obtained for the left kinks, leading to a physical picture of “fast” and “slow” plasticity carriers for the 30° partial dislocation. In our concluding section, we discuss this point further.

Here, it is worth pausing to compare our results with those in Ref. 13. Individual kink energies are not obtained in their work, since in all their calculations the supercells contained a double kink (two kinks, one of each family). Therefore, we cannot compare our kink energies directly with their results. In their procedure, what is computed are the relative energies of kinks within each kink family, assuming the LK and RK to have the same energy. A first aspect to be pointed out is that the above assumption of degeneracy between the LK and RK is in sharp disagreement with our findings. In agreement with our work, they find an energy difference of $\sim 0.4$ eV between the two left kinks. On the other hand, while our results indicate that the two right kinks also differ by $\sim 0.4$ eV, they find these two kinks to be almost degenerate, with energies differing by 0.07 eV only. Our kink migration barriers are substantially higher, despite the fact that the associated saddle-point configurations seem to be very similar with those identified in their work. Below, our results will be seen to compare more favorably with experimental estimates of the kink barriers.

### D. PSD-kink complexes

Kinks and PSD’s can be considered as the fundamental types of excitations in the dislocation cores. Important structural features and modes of dislocation dynamics can also be associated with the complexes formed by these basic defect types. Moreover, since RD’s such as the PSD are malcoordinated (thus acting as weak links in the reconstructed core), they may act as preferential sites for the nucleation of double kinks, as suggested by Heggie and Jones.1,25 Possibly, a PSD-kink complex could result from such a nucleation process, as the double kink expands and eventually dissociates into single kinks. Therefore, it is important to understand the structure and energetics of these complexes.

Here, we consider the energetics of the PSD-kink complexes. The important questions concern whether or not these complexes form bound states, as well as the associated binding energies and migration barriers. We considered each of the PSD-kink complexes in two configurations, as shown in Fig. 6. The left complex (LC=LK+PSD) is shown in the state of closest approach, LC(0), Fig. 6(a), in which the two constituents overlap and cannot be distinguished; and in an extended state, LC(1), Fig. 6(b), in which the PSD and the kink have been separated to adjacent positions. The corresponding right-complex cases RC(0) and RC(1) are shown in Figs. 6(c) and 6(d), respectively. In Table II we show our results for the energies of these four configurations, where it can be seen that the PSD binds strongly with both the left and the right kinks, in agreement with Ref. 13. Contrary to what is found in Ref. 13, our results indicate the LC to be more strongly bound than the RC. From the binding energies
and the energies of these more extended configurations, we obtain a lower bound of 0.80 eV (LC) and 0.49 (RC) for the dissociation barrier of these bound states. Below, these results will be shown to be in sharp contrast with those for kink-RD complexes in the SP reconstruction of the 90° partial dislocation. In this section, we summarize our main results for the core and related defects of the DP structure. In our calculations, the result is in good agreement with previous TB calculations of Ref. 16 (0.18 eV) and local-density approximation works, and also compares reasonably well with the value of 0.12 eV obtained in Ref. 26 (a Tersoff potential study). We find the SP core to be 0.18 eV/Å lower in energy than the QF one. This result is in good agreement with previous TB (Ref. 16) (0.18 eV) and local-density approximation works, and also compares reasonably well with the value of 0.12 eV obtained in Ref. 26 (a Tersoff potential study) and Ref. 37 (using a combined Keating-LCAO approach). In Ref. 14, it was found that symmetry breaking occurs spontaneously, a result that is confirmed by our model. In our calculations, the reconstructed bonds are stretched 3.0% with respect to the perfect crystal values (2.5% in Ref. 14), and the minimum and maximum bond angles are 97° and 135°, respectively (96° and 138° in Ref. 14). Core defects are considered next.

B. Direction switching defect (DSD)

Symmetry breaking in the SP core gives rise to a RD in which the direction of the bonds is switched, as shown in Fig. 7(d). We shall refer to this defect as a direction switching defect (DSD). Note that, like the 30°-partial PSD, this defect contains a dangling bond, which explains its formation energy of 1.45 eV. Our result is in reasonable agreement with the 1.2 eV value obtained in the cluster calculations of...
Ref. 27, being lower than the value of 2.0 eV obtained in Ref. 37. For the DSD motion, we computed an energy barrier of only 0.04 eV for the propagation between two adjacent equilibrium positions. Given such a small barrier, the DSD is expected to be extremely mobile even at low temperatures. As a test, we performed a molecular-dynamics simulation on a supercell having a pair of DSD defects, initially separated by 9.6 Å, on an otherwise defect-free partial dislocation. Remarkably, at a temperature of only 50 K, recombination of the pair took place after only 1.3 ps. Unlike PSD’s in the 30° partial, these complexes appear to be either unstable or marginally stable against the emission of a DSD, as discussed in Ref. 15. The dissociation barrier, if present, is basically the migration barrier of the SR core, this additional strain is more than compensated for by the attraction between the LR and RL kinks. We also computed an energy barrier of 1.62 eV for the motion of the LR and RL kinks. As is the case for reconstructed kinks in the 30° partial, such large energy barriers are associated with the existence of malcoordinated atoms and severe bond distortions at the core of the kink.

D. DSD-kink complexes

There are two additional kink-type defects associated with the SP reconstruction of the core. These are the RR and the LL defects, shown in Figs. 8(c) and 8(d). We prefer to regard these as complexes of a RR or a LL kink together with a DSD. Two LL complexes are possible (only one is shown in Fig. 8), and they share the same ‘quasisymmetry’ that the LR and RL kinks do, differing only by the position of the fivefold and dangling-bond-containing rings with respect to the stacking fault. In contrast with complexes in the 30° partial, these complexes appear to be either unstable or marginally stable against the emission of a DSD, as discussed in Ref. 15. The dissociation barrier, if present, is basically the DSD migration barrier, which indicates that these complexes should dissociate very easily at moderate temperatures. This was confirmed by a simulation performed at 300 K, with a supercell containing a pair of RR complexes in each dislocation, separated by a distance of 34.6 Å. On the time scale of 1 ps, one of the kink complexes undergoes the DSD-emission reaction RR → RL + DSD, with the DSD propagating rather easily towards the other RR complex, where a DSD + RR → LR process takes place. Overall, a dislocation containing a pair of RR complexes relaxes into one containing alternating RL and LR kinks, by means of DSD emission (absorption) and propagation.

V. COMPARISON WITH EXPERIMENTAL RESULTS

In Table III we summarize our results for the formation energies and migration barriers of kinks in the 90° and 30° partial dislocations. For the 30° partial, of the two equilib-
rium states of each kink \([\text{LK}, \text{LK}']\) and \([\text{RK}, \text{RK}']\), one is to regard as an intermediate metastable state in the propagation of the kink, given the substantial difference in formation energy between the two states. Only the state with the lower formation energy will determine the kink concentration in each case (this lower formation energy is the number included in Table III). For comparison, results from Ref. 13 are also included, as are the ranges of experimental results for both quantities, obtained from different techniques.4–7 We observe that, for the 30° partial, our values are in excellent agreement with the experimental ones.

The interpretation of these experiments is done according to the theory of Hirth and Lothe. In this theory, the dislocation velocity is given by

\[
v_d \approx 2 \times \exp \left[ -\frac{1}{kT} (U_i + W_m) \right],
\]

where \(U_i\) is the kink formation energy and \(W_m\) is the kink migration barrier. This equation is written under the assumption that the two kinks that result from the nucleation of a stable double kink (a kink-antikink pair) are equivalent. This assumption does not hold for the 30° partial, where the left and right kinks are intrinsically different. The more general form

\[
v_d \approx \exp \left[ -\frac{1}{2kT} (U_{\text{LK}} + U_{\text{RK}}) \right] \times \left[ \exp \left( -\frac{W_{\text{LK}}}{kT} \right) + \exp \left( -\frac{W_{\text{RK}}}{kT} \right) \right],
\]

must be used. We note that the quantity of interest in the first activated term is the average formation energy of the two kink species. The second term is derived from the kink velocities, and therefore the relative velocity appears in the generalized form. In the 30° partial this term is dominated by the velocity of the left kinks (fast carriers), given the much higher migration barrier of the right kinks (slow carriers).

VI. CONCLUSIONS

In this work, an extensive study of the core reconstruction and structural excitations in the cores of both the 30° and the 90° partial (in its SP reconstruction) in Si, was presented. For both partials, we find the core to undergo strong bond reconstruction, restoring the fourfold coordination of the core atoms. The reconstructed bonds are stretched by only \(~3\%\) with respect to bulk values, and the core energies are mostly associated with the bond-angle distortions present in the reconstructed cores.

In the case of the SP structure of the 90° partial, the RD (or DSD) is associated with a switch of direction of the reconstructed bonds, and is found to be highly mobile. Kink-DSD complexes are found to be only marginally stable against emission of a DSD, a reaction that is observed to proceed rather quickly in our simulations at room temperature. The LR and RL kinks have very low formation energy, indicating that they introduce little additional on the SP core, a result which is consistent with the lower energy of the DP core, as proposed in Ref. 19.

For the 30° partial, two kink species (RK and LK) are identified, and the RK’s are found to have higher formation energies that the LK ones. This is explained by the medium-range behavior of the associated strains. The RD (or PSD) is related to a phase switching of the core reconstruction, and binds strongly with kinks to form PSD-kink complexes. These are the more likely sites for unpaired electrons in the 30° partial core. The results for this particular dislocation can be directly compared with experiment, and we find good agreement between our calculated values for the kink formation energies and migration barriers and the experimental results.

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28. Reconstruction defects in the core of the 30° and 90° partials have commonly been termed as solitons, but variants such as antiphase defect have also been used. In this paper, we shall refer to these as phase-switching defect, in the case of the 30°, and direction-switching defect in the case of the 90° partial, as explained in the text.


32. The reaction path is the path joining the two minima in configuration space, along which the gradient is always parallel to the path. Physically, it is the minimal-action path that the system would follow to reach a minimum once released from the saddle-point configuration.


35. Several Keating and bond-charge model studies are summarized in Ref. 3. For the 30° partial core, the results range between 0.28 and 0.32 eV/Å, when for E_{db} (the energy of a dangling bond), we use our result for the PSD formation energy. For the 90° partial, these calculations give results ranging from 0.31 to 0.60 eV/Å, using our DSD energy as an estimate for E_{db}. The various estimates of kink and kink-pair energies listed in this reference are in very good agreement with our results.


