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Divacancy in 3C- and 4H-SiC: An extremely stable defect

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Using first-principles calculations for divacancy defects in 3C- and 4H-SiC, we determine their formation energies and stability, their ionization levels, and relaxed geometries (symmetry point groups) for neutral as well as for charged states. For 4H-SiC all four possible nearest-neighbor divacancy configurations are considered. We find not only a remarkable high binding energy of about 4 eV, but also a strong site dependence (cubic or hexagonal lattice sites) of the formation energies. Applying a Madelung-type correction to deal with the electrostatic interactions between charged supercells, our results indicate a negative-U behavior at $E_V + 0.7$ eV between the charge states $1^+/-1^-$—only for nearest-neighbor divacancies on different lattice sites (mixed cubic and hexagonal) in 4H-SiC, but not for all the other cases (pure cubic or pure hexagonal) in 4H—or for the cubic divacancy in 3C-SiC.

I. INTRODUCTION

SiC is a very promising material for semiconductor device applications which have to work under extreme conditions. Due to its good thermal conductivity, its high radiation resistance, and high breakdown voltage, it is well suited for demanding applications in harsh environments.

Even though progress in crystal growth during the past years has been able to reduce imperfection in SiC to a great extent, many properties of grown-in defects or those produced by irradiation damage are not well understood. However, new defects are introduced during doping wafers by ion implantation. To get rid of unwanted damage and to bring the dopant atoms on lattice sites—to activate them—one has to employ certain annealing steps, where unnecessarily high temperatures have to be avoided, since this would cause the dopants to diffuse and the sharp ion implantation profile would be lost. During annealing, irradiation-induced defects may or may not anneal out. In the “hard” SiC most primary defects are stable at room temperature and, hence, are easily accessible experimentally.

Since defects of the crystal lattice influence the electrical properties of the material, there is a great need to identify and to understand them. In the past there were difficulties in understanding the electric properties of GaAs, due to the electrically active point defect EL2, existing even in as-grown crystal in high concentrations. Now—after more than 20 years of studies—consensus has been reached on EL2 being the isolated As antisite in GaAs. To improve the reliability of defect identification, it is necessary to compare experimental data on defect symmetries, formation energies, and ionization levels to first-principles calculations. Defect identification is the prerequisite to avoid or—if unwanted—to deactivate them.

A characteristic property of SiC is its polymorphism—an uncommon feature in nature. Polymorphism means that SiC is able to crystallize in different modifications, so-called polytypes. The polytypes differ in the stacking sequence of hexagonally close-packed double layers of Si and C atoms. Depending on the next-nearest-neighbor atom arrangement, the atom sites in SiC crystal are classified to be cubic or hexagonal ones. In 4H-SiC the divacancy $V_{Si}V_{C}$—formed by nearest-neighbor silicon ($V_{Si}$) and carbon ($V_{C}$) monovacancies—has four possible configurations, since both Si and C atoms can be located either in cubic (“cub”) or hexagonal (“hex”) lattice sites. The four possible configurations are $V_{Si-cub}V_{C-cub}$, $V_{Si-cub}V_{C-hex}$, $V_{Si-hex}V_{C-cub}$, and $V_{Si-hex}V_{C-hex}$. In 3C-SiC, on the other hand, there is only one kind of lattice site, the cubic one, and thus only one kind of divacancy $V_{Si-cub}V_{C-cub}$.

Already identified intrinsic point defects in SiC such as the silicon and carbon vacancies ($V_{Si}, V_{C}$) or the silicon antisite $Si\text{C}$ cannot explain the high thermal stability of a number of commonly detected photoluminescence (PL) centers in SiC. Since the presence of simple point defects (vacancies, interstitials, antisites) can be excluded, one has to look for more extended defects—the simplest examples are the divacancy or vacancy-antisite pairs.

Although the divacancy $V_{Si}V_{C}$ has experimentally not been conclusively identified in SiC, it has in the past often been proposed to be an important defect center. Hence, there is a must to study the divacancy in 3C- and 4H-SiC by first-principles calculations. The $Z_1$ and $Z_2$ defect centers observed by various capacitance transient techniques and photoinduced electron spin resonance (ESR) have been proposed to be associated with $V_{Si}V_{C}$. Also for one of the most common and stable defect centers in SiC labeled D1—widely detected in PL studies—the divacancy has been discussed in the past as a possible candidate. However, no full consensus prevails regarding the origin of D1. In the experimental literature, divacancy-related defect centers are discussed also in several positron annihilation studies (cf., e.g., Refs. 15–17).

In computational studies for defect properties in SiC, the electronic structure of the divacancy in SiC has received little attention up to now. While Talwar and Feng used a semiempirical tight-binding (TB) approach for a divacancy in 3C-SiC, Wang et al. studied the electronic structure of the divacancy with ab initio methods, where the calculations were performed in 3C-SiC using a 32-atom supercell without full ionic relaxations. The TB study by Talwar and
Feng revealed that the divacancy has localized impurity states in the semiconductor gap. The symmetry point group was determined to be $C_{3v}$ for the neutral divacancy, whereas in the \textit{ab initio} study the formation energy was evaluated to be 8.1 eV.\textsuperscript{18}

Even though for many point defects in SiC (antisites, monovacancies, and antistructure pairs) accurate pseudopotential calculations, employing full ionic relaxations\textsuperscript{19-22} and large (128 or 216 atom) supercells, have appeared during past years, the electronic structure of the divacancy in SiC has not been studied recently using accurate state-of-the-art first-principles methods. In this paper, the electronic structure of all nearest-neighbor divacancies in $3C$- and $4H$-SiC polytypes will be studied. While the electronic structure of the divacancy in GaAs and Si has been successfully studied using plane-wave pseudopotential calculations,\textsuperscript{23,24} to the knowledge of the authors, no works have been reported on divacancies in SiC.

II. METHODS

Our calculations are based on the density-functional theory (DFT) with the electron exchange correlation treated in the local density approximation (LDA) and local spin density approximation (LSDA). We have used a Car-Parrinello-like pseudopotential approach.\textsuperscript{27} While for the C ion the Vanderbilt-type ultrasoft pseudopotential\textsuperscript{28} has been employed in order to reduce the number of plane waves needed to describe the electronic wave functions, a standard norm-conserving Bachelet-Hamann-Schütz pseudopotential\textsuperscript{29} has been used for the Si ion. Good convergence with respect to the basis set size is obtained at a 20 Ry kinetic energy cutoff,\textsuperscript{30} which has been used throughout this work. In the defect calculations, the initial atomic configurations have been randomized slightly from the ideal structure to remove any spurious symmetries. All the ions in the supercell have been allowed to relax without any symmetry constraints using the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm.\textsuperscript{31} For the electronic structure minimization, we employed damped second-order dynamics\textsuperscript{32} combined with the Williams-Soler algorithm.\textsuperscript{33} All calculations have been performed in a massively parallel CRAY-T3E system using the carefully optimized FINGER (finnish general electron relaxator) code.\textsuperscript{34} While in $3C$-SiC the calculations are performed for a 128-atom-site fcc supercell, for the hexagonal structure of $4H$-SiC the supercell is elongated in the $c$ direction due to the ratio $c/a = 1.63$. Hence, to keep the distance between the defect and its periodic replica in the superlattice as large as possible, we choose—after carefully testing other alternatives—a rectangular supercell having a shape close to cubic (the hexagonal lattice of $2H$, $4H$, and $6H$-SiC can be transferred to cubic coordinates resulting in rectangular supercell). In all calculations a rectangular 128-atom supercell has been employed. Considering a proper $k$-point set,\textsuperscript{35} use of the $\Gamma$ point provides sufficiently well-converged results as well as the required efficiency in computing.\textsuperscript{19,30} For spin-unpaired charge states ($V_2^{1-}$ and $V_2^{1+}$), we performed a spin-polarized calculation employing the LSDA scheme.\textsuperscript{26}

A. Formation energies for defects

The formation energy of defects $E_F(q)$ is calculated using the standard formalism by Zhang and Northrup.\textsuperscript{36} For all positions of the electron chemical potential and the possibly appearing stoichiometries between C- and Si-rich compounds, the formation energy $E_F(q)$ is defined as follows:

$$E_F(q) = E_{\text{tot}}(q) + q (E_V + \mu_e) - n_Si \mu_{Si} - n_C \mu_{C}$$

$$= E_{\text{tot}}(q) - \frac{1}{2} (n_{Si} + n_C) \mu_{Si}^{\text{bulk}} - \frac{1}{2} (n_{Si} - n_C) (\mu_{Si}^{\text{bulk}} - \mu_{C}^{\text{bulk}}) + q (E_V + \mu_e) - \frac{1}{2} (n_{Si} - n_C) \Delta \mu,$$ \hspace{1cm} (1)

where $E_{\text{tot}}(q)$ is the total energy of the defect supercell in question and $q$ the charge state. $n_{Si}$ and $n_{C}$ are the numbers of Si and C atoms in the supercell, respectively. $\mu_e$ is the electron chemical potential measured relative to the valence-band maximum $E_V$. $\mu_{Si}^{\text{bulk}}$ and $\mu_{C}^{\text{bulk}}$ are the chemical potentials of the Si and C atoms in the bulk Si and C lattices, respectively. $\mu_{Si}^{\text{bulk}}$ is the chemical potential of the SiC pair in the bulk SiC compound. Actually it is possible to know only

$$\mu_{Si}^{\text{bulk}} + \mu_{C}^{\text{bulk}},$$ \hspace{1cm} (2)

not the chemical potentials $\mu_{Si}$ and $\mu_{C}$ of the single atom separately. This fact is circumvented in Eq. (1) by introducing the chemical potential difference

$$\Delta \mu = (\mu_{Si} - \mu_{C}) - (\mu_{Si}^{\text{bulk}} - \mu_{C}^{\text{bulk}}).$$ \hspace{1cm} (3)

The allowed range of $-\Delta H < \Delta \mu < \Delta H$ is determined from the calculated heat of formation for SiC defined as

$$\Delta H = \mu_{Si}^{\text{bulk}} + \mu_{C}^{\text{bulk}} - \mu_{SiC}^{\text{bulk}}.$$ \hspace{1cm} (4)

The atomic chemical potential difference $\Delta \mu$ determines the deviation from the ideal stoichiometry. It can vary from $-\Delta H$ (C rich) to $\Delta H$ (Si rich).

We plot the formation energies as functions of the electron chemical potential from the valence-band maximum (VBM) $E_V$ to the value of the band gap ($E_{\text{gap}}$). The appropriate value for the chemical potential difference $\Delta \mu$ depends on the defect and the defect formation process.

The ionization level ($q/q'$) of a given defect is the position of $\mu_e$ in the band gap where the most stable charge state changes from $q$ to $q'$. Ionization levels can be obtained by solving the following equation for the value of the electron chemical potential $\mu_e$:

$$E_{\text{tot}}(q) + q (E_V + \mu_e) = E_{\text{tot}}(q') + q' (E_V + \mu_e).$$ \hspace{1cm} (5)

The computational DFT-LSDA method has some unresolved problems, which can have a significant influence on the ionization levels. The most important is the gap problem: in the electronic structure calculations the band gap is extracted from Kohn-Sham single-particle energy levels. For-
nally, these levels do not describe any physical quantity—only the total energy is well defined. Although Kohn-Sham eigenvalues sometimes agree quite nicely with the true quasiparticle band structure, the width of the band gap is always severely underestimated. While the corresponding experimental values for 3C- and 4H-SiC are $E_g = 2.39$ and 3.27 eV, respectively, we obtain for the energy gap of 3C- and 4H-SiC $E_g = 1.24$ and 2.6 eV, respectively. Since defect states appear in the band gap, there can be problems in interpreting the actual positions of the ionization levels according to Eq. (5). In this work, the calculation of the ionization levels is based on the total energies. We will consistently report ionization levels measured from the valence-band maximum. However, when plotting the formation energies, we indicate also the position of the LDA gap and leave it to the reader to measure the distance of an ionization level with respect to the experimental conduction-band minimum (CBM) or with respect to the Kohn-Sham band edge.

Neglecting the phonon contribution to the entropy, the formation energy $E$ is given by a formula

$$E = E_0 - q^2 \alpha - \frac{2 \pi q Q}{3L} + 0(L^2),$$

where $E_0$ is the total energy of the supercell obtained from the calculations, $Q$ is the quadrupole moment of the charge distribution in the supercell, $\alpha = 1.41865$ (rectangular 128-atom cell of size 19.89 a.u.) the Madelung constant, $e = 6.7$ the experimental dielectric constant, and $L$ the size of used supercell. The contribution arising from the second term of the Madelung correction is only one-third of that coming from the first correction term. Since this may be beyond the accuracy of the method, it will not be considered here. For 4H-SiC, we obtain a correction for $q = \pm 1$ around 0.3 eV and for $q = \pm 2$ about 1.2 eV.

III. RESULTS

A. Formation energies

In Table I, we present our results for the formation energies of mono- and divacancies in 3C- and 4H-SiC under stoichiometric conditions ($\Delta \mu = 0$). The binding energies for the neutral divacancy defects in 3C- and 4H-SiC are given for all lattice site configurations in Table I as well.

We find the lowest formation energies for divacancies in 4H-SiC when both $V_{Si}$ and $V_{C}$ are along the $c$ axis (both vacancies either only on cubic or only on hexagonal lattice sites). The by-far lowest formation energy is obtained for the pure cubic lattice site $V_{Si}^c V_{C}^c$. The values are 7.22 eV for 3C-SiC and 7.74 eV for 4H-SiC, while for a purely hexagonal divacancy in 4H-SiC, $V_{Si}^h V_{C}^h$, we find a somewhat higher formation energy of 8.0 eV. By far the highest formation energies are those for the mixed cases $V_{Si}^c V_{C}^h$ and $V_{Si}^h V_{C}^c$ (divacancies not along the $c$ axis in 4H-SiC), where the formation energy $E_F$ is 0.6 eV larger than the pure cubic case (see Table I).

It has to be pointed out that the formation energy of divacancies on pure cubic or pure hexagonal lattice sites is in fact lower than the formation energy of isolated silicon monovacancies on corresponding lattice sites (see Table I for stoichiometric material). For the pure cubic divacancy this energy difference is as high as 0.6 eV, while for the pure hexagonal divacancy it is about 0.3 eV. Due to this signifi-

![Table I. Monovacancies and divacancies in 4H- and 3C-SiC: shown are the calculated formation energies $E_F$ in electron volts [eV]. In 4H-SiC, $E_F$ is evaluated for all possible, cubic (cub) or hexagonal (hex), lattice sites. The values of the formation energy are given for stoichiometric ($\Delta \mu = 0$) SiC. A remarkable high binding energy $E_b$ is obtained for all divacancies.](image-url)
ciant difference, one may expect to find a higher concentra-
tion of divacancies than the corresponding one of isolated
silicon monovacancies under equilibrium conditions accord-
ing to Eq. (6).

Since the formation energy for divacancies is low com-
pared to monovacancies, we also find an extraordinarily
large binding energy (3.8–4.4 eV) between vacancies on
different sublattices in 4H-SiC (see Table I). The pure cubic
or the pure hexagonal cases have the highest binding energy.
They are larger by about 0.6 eV compared to the mixed
ones. The cases where the silicon vacancy on a cubic lattice
site is involved—either pure or mixed—are favored by about
0.1 eV. Additionally, the pure cubic case is favored com-
pared to the pure hexagonal one by a difference in the for-
amation energy of 0.26 eV. Again we see a strong site depen-
dence.

For 3C-SiC the binding energy found is significantly
smaller: about 3.3 eV. This is in qualitative agreement with
previous calculations by Wang et al., employing only a 32-
atom supercell, where the binding energy was evaluated to
be 4.6 eV in 3C-SiC.18

B. Ionization levels

In Fig. 1 the formation energies for all possible nearest-
neighbor (NN) divacancies in 4H-SiC are given as a func-
tion of the electron chemical potential without any correc-
tion. The ionization level (1+/2+) is located just above the
VBM for 4H-SiC while it is slightly higher for 3C-SiC. The
ionization level (2−/1−) is situated slightly below midgap.
A negative-\(U\) behavior between the (1−/1+) charge states
is found generically for 4H-SiC at about 0.7 eV above the
TABLE II. Divacancies in 4H- and 3C-SiC: ionization levels for the relaxed defects. The results from LSDA calculations are given in eV above the valence-band maximum. The upper part is without any correction, while in the lower part the Madelung correction is applied. In 4H, the results are given for all four nearest-neighbor divacancy combinations. The unphysical states are given in brackets. Depending on the type of the divacancy, there is a minor deviation in the position of the negative-U transition.

<table>
<thead>
<tr>
<th>Ionization levels [eV] above the VBM</th>
<th>2+/1+</th>
<th>1+/0</th>
<th>1+/-1</th>
<th>0/-1</th>
<th>1-/2-</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Without correction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4H: $V_{\text{cub}}^{\text{cub}}$</td>
<td>0.14</td>
<td>(0.71)</td>
<td>0.68</td>
<td>(0.65)</td>
<td>1.20</td>
</tr>
<tr>
<td>4H: $V_{\text{cub}}^{\text{hex}}$</td>
<td>0.11</td>
<td>(1.02)</td>
<td>0.69</td>
<td>(0.36)</td>
<td>1.24</td>
</tr>
<tr>
<td>4H: $V_{\text{hex}}^{\text{cub}}$</td>
<td>0.06</td>
<td>(1.17)</td>
<td>0.69</td>
<td>(0.22)</td>
<td>1.28</td>
</tr>
<tr>
<td>4H: $V_{\text{hex}}^{\text{hex}}$</td>
<td>0.23</td>
<td>(0.80)</td>
<td>0.76</td>
<td>(0.71)</td>
<td>1.25</td>
</tr>
<tr>
<td>3C: $V_{\text{Si}}V_{\text{C}}$</td>
<td>0.42</td>
<td>(1.00)</td>
<td>0.85</td>
<td>(0.70)</td>
<td>1.16</td>
</tr>
</tbody>
</table>

| **Madelung correction applied**      |       |      |       |      |       |
| 4H: $V_{\text{cub}}^{\text{cub}}$   | —     | 0.52 | —     | 0.85 | 1.76  |
| 4H: $V_{\text{cub}}^{\text{hex}}$   | —     | (0.98)| 0.69  | (0.41)| 1.84  |
| 4H: $V_{\text{hex}}^{\text{cub}}$   | —     | (0.83)| 0.69  | (0.55)| 1.80  |
| 4H: $V_{\text{hex}}^{\text{hex}}$   | —     | 0.61 | —     | 0.90 | 1.81  |
| 3C: $V_{\text{Si}}V_{\text{C}}$     | —     | 0.81 | —     | 0.89 | 1.72  |

VBM for all cases [see Figs. 1(a)–1(d)], while for 3C-SiC it is about 0.85 eV above the VBM [see Fig. 3(a)]. For details see Table II.

If the Madelung correction is applied, the picture changes in some important aspects. As we can see from Figs. 2(a)–2(d), the 2+ charge state is not stable anymore; i.e., it vanishes to the valence band. In turn the stability region of the single positive-charge state of the divacancy 1+ becomes wider and extends from the VBM to about $E_{\text{v}} + 0.5–0.7$ eV. Due to the correction, the ionization level (2+/1−) has moved higher up in the band gap and is now found at about $E_{\text{v}} + 1.8–2.0$ eV above the VBM for both 4H- and 3C-SiC. For 3C-SiC this means that the (2+/1−) level is now found above the Kohn-Sham band edge but still below the experimental CBM. Hence, the charge state 1− extends over a wide region around midgap—from $E_{\text{v}} + 0.7$ eV up to $E_{\text{v}} + 1.8$ eV for 4H-SiC and from $E_{\text{v}} + 0.9$ eV up to $E_{\text{v}} + 1.7$ eV for 3C-SiC. For 4H-SiC our results indicate that the charge state 2− is stable even with the applied Madelung correction, provided that the value of the electron chemical potential is high (n-type material). For details see Table II.

Concerning the negative-U behavior in 4H-SiC, it is still found for the mixed case (one of the vacancies is on a cubic site and the other on a hexagonal site—$V_{\text{Si}}^{\text{hex}}V_{\text{C}}^{\text{cub}}$ or $V_{\text{Si}}^{\text{cub}}V_{\text{C}}^{\text{hex}}$). For the pure cubic or pure hexagonal case (divacancies formed along the c axis: $V_{\text{Si}}^{\text{cub}}V_{\text{C}}^{\text{cub}}$, $V_{\text{Si}}^{\text{hex}}V_{\text{C}}^{\text{hex}}$), on the other hand, by applying the Madelung correction the neutral state stabilizes over a wide range of the electron chemical potential and the negative-U behavior is removed. The reason for this different behavior can be found in the lower formation energy of the charge-neutral divacancy in the latter cases and, hence, the neutral-charge state stabilizes in the range $E_{\text{v}} + (0.5–0.9)$ eV. This means that all charge states from 1+ to 2− may stabilize for certain values of the electron chemical potential.

For the nearest-neighbor divacancy in 3C-SiC the negative-U behavior between the 1+ and 1− charge state [see Fig. 3(a)] is removed when the Madelung correction is applied and the neutral divacancy may stabilize in a small range of the electron chemical potential [see Fig. 3(b)].

C. Relaxations

The relaxations of the atoms surrounding the different divacancies are listed in Table IV below. The relaxations of the three Si atoms sitting around one end (carbon vacancy) are denoted by the bond lengths between them, $d_{12}, d_{13}, d_{23}$. Respectively, the bonds between the C atoms surrounding the other end of the divacancy (around the missing Si atom) are denoted by $d_{45}, d_{46}, d_{56}$. For 4H-SiC the symmetry groups are given considering only the NN arrangements, i.e., provided that there is locally a mirror plane or a rotational symmetry.

Generally we find a strong outward relaxation of the C atoms around the missing Si atom ($V_{\text{Si}}$ in $V_{\text{Si}}V_{\text{C}}$), while the Si atoms around the missing C atom ($V_{\text{C}}$ in $V_{\text{Si}}V_{\text{C}}$) are only slightly moved—after the relaxation they are found nearly in their ideal places. In detail, we can describe the relaxation around the divacancy in the charge states 2+ or 2− by a breathing-mode type or weak pairing type according to the obtained symmetry group $C_{3v}$. For singly negative- or positive-charge states the relaxation is found to be mostly of the pairing type (symmetry group $C_{1h}$). The neutral-charge state—according to our results—is not stable in all cases. It tends to have the resonant-bond-type relaxation (symmetry group $C_{1v}$).

Comparing to the relaxation patterns obtained for monovacancies in 3C- and 4H-SiC we note the following: The C atoms around the silicon monovacancy $V_{\text{Si}}$ relax outward $\sim 10$% in $T_{d}$ or $C_{3v}$ symmetry for all polytypes. The relaxations around the missing silicon atom in $V_{\text{Si}}V_{\text{C}}$ are more distorted, and the outward movement is not as large as in the case of the silicon monovacancy. For the carbon monovacancy $V_{\text{C}}$ a Jahn-Teller-distorted, pronounced inward relaxation is found, whereas—as already noted—for $V_{\text{Si}}$ in $V_{\text{Si}}V_{\text{C}}$ the Si atoms stay nearly at their ideal lattice positions.

IV. DISCUSSION

A. Formation energies

Due to the wide band gap of SiC (experimental values are $E_{g} = 2.4$ eV for 3C-SiC and 3.3 eV for 4H-SiC), the for-
mation energies of defects depend heavily on the position of the electron chemical potential [see Eq. (1)]. In hexagonal polytypes, such as $4H$-$SiC$, the band gap is about 1 eV wider than in cubic $3C$-$SiC$, which favors negative-charge states (theoretical values are $E_g = 1.24$ eV for $3C$-$SiC$ and 2.6 eV for $4H$-$SiC$).

Considering the site dependence on the formation energies for monovacancies, it turns out that the energy differences are smaller than 0.1 eV (see Table I and Ref. 30). But for divacancies the story is different: there we find a strong dependence on the lattice sites (cubic or hexagonal) from which the atoms have been removed. The most expensive divacancies are for mixed cases (one vacancy from a cubic site and the other from a hexagonal site). If, on the other hand, both silicon and carbon atoms are removed from hexagonal sites, the formation energy is smaller by about 0.35 eV, while for atoms removed both from cubic sites the energy gain is another 0.25 eV. The reason is an energy gain by larger relaxations due to Jahn-Teller distortions in the latter cases. Calculations done for the unrelaxed divacancies show that the total energy differs by less than 0.1 eV between different cases (see Table III). On the other hand, the energy lowering by Jahn-Teller distortion shows that the most favored case is “cubcub” followed by “hexhex” (see Table III).

Comparing the formation energies for divacancies, calculated here, to those for monovacancies we find the remarkable fact that divacancies formed from pure cubic or pure hexagonal lattice sites, i.e., situated along the c axis, are energetically more favorable by up to 0.6 eV than the corresponding Si monovacancies under equilibrium conditions for stoichiometric or C-rich material (see Table I).

Comparing to others, our results for $4H$-$SiC$ for the isolated carbon and silicon monovacancies in are very good agreement with the pseudopotential calculations by Zywietz et al., since in those calculations the same rectangular su-

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{$V_{Si}V_{C}$: divacancy in $4H$-$SiC$ in all possible site configurations for stoichiometric material ($\Delta \mu = 0$). Just like the figure above only the Madelung correction is applied. Note that the negative-$U$ effect is removed for the pure cubic or pure hexagonal case.}
\end{figure}
percell (128 atom) was used. However, Zywietz et al.\textsuperscript{22} calculated the formation energy for the monovacancies in 3C-SiC in a 216-atom supercell. Thus in the case of the 3C structure their results should be more accurate due to a smaller defect-defect interaction.\textsuperscript{39}

For 3C-SiC, we do not find such a good agreement for the formation energies of monovacancies with Wang et al.,\textsuperscript{18} since they used only a 32-atom supercell, and no full ionic relaxations were allowed for. Hence, the values for the formation energies differ by up to 2 eV.

### B. Ionization levels

The ionization levels for the divacancy in different configurations in 4H-SiC and in 3C-SiC are presented in Fig. 4(a) without the Madelung correction and in Fig. 4(b) with the Madelung correction applied. One observes that the negative-$U$ behavior is removed in some cases (4H, cubcub, and hexhex, 3C) when the Madelung correction is applied.

Observations showing that the $Z_1/Z_2$ centers exhibit negative-$U$ behavior in electron irradiated 4H- and 6H-SiC have been recently published.\textsuperscript{6,40} Like the negative-$U$ calculated for the divacancy, this negative-$U$ transition is found between the charge states $1^+$ and $1^-$, but at a different position in the band gap. These centers have been identified in as irradiated and at 900 °C annealed samples. Therefore, it is quite probable that the observed signal is related to divacancies, since a divacancylike signal has been detected also by positron annihilation after similar sample preparation.\textsuperscript{41,42}

In the tight-binding picture of $V_{Si}V_{C}$ by Talwar and Feng,\textsuperscript{14} the one-electron levels were calculated to be at 0.48 eV and 1.62 eV above the VBM, and the symmetry character for the neutral divacancy was determined to be $E(C_{3v})$. It is found that for this divacancy symmetry, the two triply degenerate $t_2$-type isolated-impurity levels split and produce two sets of doubly degenerate $E$- or $\pi$-type bonds and two nondegenerate $A_1$ or $\sigma$-type levels.\textsuperscript{14} Since the $\sigma$-type levels are derived from the mixing of $s$- and $p$-like orbitals of the two isolated defects, the $A_1$ states of the divacancy will be at energies significantly different from the two isolated vacancy states. In other words, the $a_1$-type resonance state of an isolated vacancy in the valence band can be pulled into the band gap by pairing it with an appropriate impurity. Hence, the earlier tight-binding calculations\textsuperscript{14} suggest for the nearest-neighbor divacancy defects having $C_{3v}$ symmetries, the possibility of localized impurity states to be detected in the gap. In this work, we find for the nearest-neighbor divacancy pair defects in the doubly positive- or negative-charge state a $C_{3v}$ symmetry. For other charge states we calculate lower symmetries.

### C. Relaxations

All studied divacancies in doubly positive- or negative-charge state exhibit a relaxation of the "breathing type" (symmetry...
relaxation towards a lower symmetry of the neutral divacancy is found to be stable, we find a pairing-type relaxation pattern in the mixed configuration. Hence, it is more expensive to form a divacancy in the mixed configuration. The lattice seems not to be able to change the bond length so much for neutral divacancies formed between cubic and hexagonal lattice sites. Hence, nearly no inward relaxation is observed. This is obviously the reason for the clearly lower formation energies by 0.35 eV and 0.6 eV found for the pure hexagonal and pure cubic case, respectively. The lattice seems not to be able to change the bond length so much for neutral divacancies formed between mixed cubic and hexagonal lattice sites. Hence, it is more expensive to form a divacancy in the mixed configuration.

V. CONCLUSIONS

We have performed first-principles calculations for the electronic and ionic structure of the divacancy in 3C- and 4H-SiC. To correct for the electrostatic self-interaction arising from charged supercells, we employed the Madelung correction, in order to better compare experimentally obtained and calculated ionization levels. Even though the correction is known to be overestimated slightly, the correction does not affect the physics of the final results here.

While in the 3C polytype only one configuration exists, there are four different ones in 4H-SiC due to atoms missing from a combination of cubic and hexagonal sites. In stoichiometric 4H-SiC the formation energies for isolated vacancies in the charge-neutral state are about 4 eV on the carbon site $V_C$ and about 8 eV on the silicon site $V_S$. Hence, it is remarkable that such a low formation energy of about 8 eV for the divacancy $V_S V_C$ in the charge-neutral state in 4H-SiC is calculated.

We observe a clear site dependence, i.e., variations in the formation energy of up to 0.62 eV depending on the lattice sites (cubic or hexagonal) in which the defect complex is situated. This is related to the different possible energy-lowering relaxations. We find for all cases extraordinarily large binding energies of about 4 eV, which, in turn, imply that for almost all divacancy configurations $V_S V_C$ their formation energy can even be up to 0.6 eV lower than the corresponding one for the isolated $V_S$ monovacancy. This means that the energy necessary for their dissociation is the binding energy plus the barrier for diffusion. Hence—once formed—they can be expected to be extremely stable defects.

The formation energy $E_F$ for divacancies and silicon monovacancies in bulk SiC is much higher than that for the carbon monovacancy and for the antisites. This indicates that the equilibrium divacancy concentration for stoichiometric semi-insulating material should be relatively low compared to antisites or carbon vacancies. Nevertheless, silicon monovacancies have been detected experimentally in as-grown material in different charge states. According to these facts one may speculate that in as-grown SiC divacancies may be the preferred defect species compared to isolated silicon monovacancies.

The divacancy in the 4H polytype can exist at different charge states $[+, -, - -]$, where the neutral defect stabilizes only for some cases. Hence, we find a negative-U behavior between the charge states $1+$ and $1-$ for divacancies in 4H-SiC situated on mixed sites (cubhex or hexcub), where

FIG. 4. Energy level schemes for Si-C divacancies in 4H as well as in 3C-SiC, (a) without Madelung correction and (b) with Madelung correction. Results are given for all possible combinations of cubic and hexagonal lattice sites obtained from LSDA calculations employing a 128-atom supercell. In (b) highly positive-charged states are shifted by the Madelung correction into the valence band while highly negative-charge states move towards the conduction band.
the transitions occur at the same value of the electron chemical potential (0.7 eV above the VBM). Applying the Madelung correction for 3C-SiC the doubly negative-charge state moves above the Kohn-Sham band edge and, hence, it is doubtful if it can be observed. If the Madelung correction would slightly overestimate the shift in formation energies, there would be some room for the divacancy in 3C-SiC to exhibit a negative-$U$ behavior as well, which would then be slightly higher in the band gap at about 0.85 eV above the VBM. For 4H-SiC, on the other hand, divacancies formed

<table>
<thead>
<tr>
<th>Charge</th>
<th>Spin</th>
<th>Carbon end</th>
<th>Silicon end</th>
<th>Symm.</th>
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<tr>
<td></td>
<td></td>
<td>$d_{12}$</td>
<td>$d_{13}$</td>
<td>$d_{23}$</td>
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<tr>
<td>2+</td>
<td>0</td>
<td>0.1</td>
<td>0.0</td>
<td>0.2</td>
</tr>
<tr>
<td>1+</td>
<td>1/2</td>
<td>-0.6</td>
<td>+0.4</td>
<td>+0.6</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0.0</td>
<td>0.0</td>
<td>+0.2</td>
</tr>
<tr>
<td>1-</td>
<td>1/2</td>
<td>-2.3</td>
<td>+0.4</td>
<td>+0.6</td>
</tr>
<tr>
<td>2-</td>
<td>0</td>
<td>-2.4</td>
<td>-0.1</td>
<td>+0.1</td>
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</table>

**TABLE IV. Divacancies in 4H-SiC and 3C-SiC.** Relaxations around the divacancy depend on its lattice sites, charge, and spin state. They are given in % of the ideal tetrahedral distances between NN carbon (distances $d_{12}, d_{13}, d_{23}$) or silicon atoms (distances $d_{45}, d_{46}, d_{56}$). In the last column we give the symmetry groups for some of the defects.

<table>
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<th>Symm.</th>
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<td>$d_{12}$</td>
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<td>$d_{23}$</td>
</tr>
<tr>
<td>2+</td>
<td>0</td>
<td>+1.4</td>
<td>+1.4</td>
<td>+0.6</td>
</tr>
<tr>
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<td>1/2</td>
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<td>+0.9</td>
</tr>
<tr>
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<td>+1.1</td>
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<td>2-</td>
<td>0</td>
<td>-0.1</td>
<td>-0.2</td>
<td>-1.3</td>
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<th>Silicon end</th>
<th>Symm.</th>
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<td>$d_{23}$</td>
</tr>
<tr>
<td>2+</td>
<td>0</td>
<td>-0.1</td>
<td>-0.0</td>
<td>+0.1</td>
</tr>
<tr>
<td>1+</td>
<td>1/2</td>
<td>-0.3</td>
<td>+0.2</td>
<td>+0.2</td>
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<tr>
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<td>+0.9</td>
<td>-0.8</td>
</tr>
<tr>
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<td>1/2</td>
<td>-0.5</td>
<td>-0.2</td>
<td>-1.6</td>
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<tr>
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<td>-1.6</td>
<td>-1.5</td>
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<tr>
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<td>+0.9</td>
<td>+1.5</td>
</tr>
<tr>
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</tr>
<tr>
<td>1-</td>
<td>1/2</td>
<td>-0.7</td>
<td>+0.7</td>
<td>+0.8</td>
</tr>
<tr>
<td>2-</td>
<td>0</td>
<td>-0.4</td>
<td>-0.2</td>
<td>-0.2</td>
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</tbody>
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along the $c$ axis (cubcub or hexhex) have their neutral-charge state stabilized in a long range of the electron chemical potential and, hence, exhibit no negative-$U$ behavior.

The relaxation pattern found around the divacancy can be described as follows: The nearest-neighbor silicon atoms at the carbon end of the $V_{Si}V_{C}$ hardly move at all from their ideal positions. On the other hand, the nearest-neighbor carbon atoms around the silicon end of the $V_{Si}V_{C}$ move symmetrically outward by about 10%, forming more $sp^2$-like bonds. The slightly different relaxations found for the divacancy in the pure cases (cubcub or hexhex) compared to those in the mixed cases (cubhex or hexcub) do indeed explain the differences in the formation energies observed.

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