Impact of local composition on the energetics of E-centres in Si$_{1-x}$Ge$_x$ alloys

Stavros-Richard G. Christopoulos$^1$, Navaratnarajah Kuganathan$^{1,2}$ & Alexander Chroneos$^{1,2}$

The energetics of the defect chemistry and processes in semiconducting alloys is both technologically and theoretically significant. This is because defects in semiconductors are critical to tune their electronic properties. These processes are less well understood in random semiconductor alloys such as silicon germanium as compared to elementary semiconductors (for example silicon). To model the random silicon germanium alloy we have employed density functional theory calculations in conjunction with the special quasirandom structures model for different compositions. Here we show that, the energetics of substitutional phosphorous-vacancy pairs (E-centres) in Si$_{1-x}$Ge$_x$ alloys vary greatly with respect to the local Ge concentration and the composition of the alloy. The most energetically favourable E-centres have a Ge atom as a nearest neighbour, whereas the dependence of the binding energy of the E-centres with respect to alloy composition is non-linear.

The substitution of the silicon native oxide with high-k dielectrics has enabled the replacement of silicon (Si) with materials such as germanium (Ge) or silicon germanium (Si$_{1-x}$Ge$_x$), which have better materials properties (e.g. higher mobilities)$^{1-10}$. Si$_{1-x}$Ge$_x$ is a random alloy in the sense that two atom species can occupy one lattice site. This in turn can lead to Si-rich and Ge-rich regions that can impact the energetics of point defects. This has been previously demonstrated in Si$_{1-x}$Ge$_x$ alloys where there is a compositional dependence of self-diffusion (refer to$^{1,11}$ and references therein) but also oxides where local compositional variation impacts diffusion$^{12,13}$. Recently, Saltas et al.$^{14}$ studied self-diffusion in Si$_{1-x}$Ge$_x$ alloys as a function of temperature and Ge concentration within the cBO2 thermodynamic model$^{14-17}$. In this study considerable deviations from linearity of the activation energies with respect to compositions were observed$^{14}$. This non-linear behaviour was attributed to the diverse behaviour of the bulk properties of Si and Ge and is consistent with the experimental data.

The theoretical investigation of random alloys using advanced techniques such as density functional theory (DFT) is not trivial as it requires a very extensive number of calculations in large supercells. To reduce the supercell size and the number of calculations methods such as the special quasirandom structures (SQS) have been previously proposed$^{18}$ and implemented in technologically important random alloys (more details are given in what follows)$^{19-23}$. Considering group IV random alloys previous work has investigated the defect processes in Si$_{1-x}$Ge$_x$, Sn$_{1-x}$Ge$_x$ and Si$_{1-x}$Ge$_x$Sn$_x$ alloys$^{24-28}$.

The main aim of the present study is to use extended SQS cells in conjunction with systematic DFT calculations to investigate the energetics of substitutional phosphorous-vacancy pairs (E-centres) in Si$_{1-x}$Ge$_x$ alloys ($x = 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875$). Particular emphasis will be placed on the effect of the composition of the alloy as well as the nearest neighbour environment to the defect.

Results and Discussion

Si$_{1-x}$Ge$_x$ structure and E-centres. Typical DFT calculations involve the construction of a supercell with periodic boundary conditions and as such they are immediately applicable for perfectly ordered structures. The situation is not straightforward, however, for disordered random alloys. Brute force methods (i.e. construct a very large supercell and randomly inserting the host atoms) is practically unfeasible for DFT as it necessitates very large supercells to reproduce the statistics of random alloys. Conversely, the SQS approach efficiently mimics the statistics of random alloys using small supercells and therefore it is a method that is compatible with DFT particularly when numerous defect calculations are required$^{19-23}$. The qualitative difference of the SQS is that they are designed small-unit-cell periodic structures that mimic the most relevant near neighbour pair and multisite...
correlation functions of the random substitutional alloys. Their atomistic nature ensures that there is a distribution of distinct local environments that also exist in real random alloys. For the Si$_{1-x}$Ge$_x$ alloys considered the Si or Ge atoms can be surrounded by various Si$_n$Ge$_{1-n}$ coordination shells (n ranging from 0 to 4). This forms a distribution of local environments, which in turn influence the formation and energetics of dopant-defect interactions such as the E-centre$^{29}$. The efficacy of the SQS technique to model random alloys has been demonstrated previously in numerous systems including group IV alloys$^{5,20,28}$, oxides$^{21,30}$, and III-V alloys$^{22,31}$.

The E-centre in Si or Ge is a substitutional donor atom (for example P, As, Sb) bound with a nearest neighbor vacancy. The structure and energetics of E-centres in Si or Ge have been extensively investigated using both experimental and theoretical techniques$^{32–36}$. Here as a criterion for the formation of the E-centre we consider the binding energies. A defect pair such as the PV is bound if the energy difference between the pair and the isolated defects (i.e. P and V being at distances long enough that they do not associate) is negative. The more negative the binding energies the more likely it would be for the PV pair to form.

Modelling E-Centres. Figure 1 is a schematic representation of the 32-atom SQS cells for the Si$_{1-x}$Ge$_x$ compositions considered. These have been reported in previous work$^{29}$, however, they have not been employed to systematically investigate the defect processes in Si$_{1-x}$Ge$_x$ alloys. Here to study the binding of the phosphorous substitutional-vacancy defect (PV pairs or E-centres) we have calculated the energies of all the different nearest neighbour PV combinations in the seven Si$_{1-x}$Ge$_x$ alloys considered. To limit defect-image errors typically introduced in computational modelling small supercells we have constructed supercells consisting of two SQS cells each (i.e. 64 atomic sites). Therefore, the total number of PV defects considered were 896, whereas more than 1350 DFT calculations were performed to predict the binding energies. The present approach allows the thorough study of defects for a range of compositions and importantly local environment (i.e. Si-rich or Ge-rich). It is an aim of this investigation to derive trends on the impact of composition and environment around the defect on the energetics of E-centres.

Figure 2 considers the effect of the nearest neighbour host atoms to the P substitutional atom. For Si$_{1-x}$Ge$_x$ cells with up to 62.5% Si content we calculate that there are also positive PV binding energies. This implies that PV pairs in Si$_{1-x}$Ge$_x$ ($x \leq 0.625$) may not form in some areas of the random alloy. This is not the case for Si$_{1-x}$Ge$_x$ ($x > 0.625$) were the PV binding energy is always negative (refer to Fig. 2). This is not of course implying that it

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Figure 1. Schematic representation of the 32-atom SQS cells for the Si$_{1-x}$Ge$_x$ ($x = 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875$).
would be equally likely for the PV to form at any sites in Si$_{1-x}$Ge$_x$ (x > 0.625). The more negative PV pairs will be more likely to form and more stable against dissociation.

Figure 3 examines the impact of the first nearest neighbour host atoms to the lattice vacancy. The trend observed is that the $E$-centre will have a more negative binding energy if it forms in the vicinity of Ge atoms, irrespective of the Si$_{1-x}$Ge$_x$ random alloy composition. The physical basis of this can be traced to the lattice relaxation of the Ge host atoms (which are larger than Si atoms) in the vacant space.

Figure 4 attempts to sum up the impact of nearest neighbours around the $E$-centre. This is to gain an understanding of whether the $E$-centres in Si$_{1-x}$Ge$_x$ prefer to form in Si-rich or Ge-rich regions of the alloy. What can be
deduced from Fig. 4 is that $E$-centres form in the vicinity of one or more Ge atoms, irrespective of the alloy composition. For all the alloys considered Ge is strongly represented in the nearest neighbours around the minimum energy $PV$. The most energetically favourable $PV$ defects and their nearest neighbour atoms are schematically represented (refer to Fig. 5) for every alloy considered here. The present findings are consistent with previous experimental and theoretical work where it was shown that vacancies preferentially form in the vicinity of Ge atoms$^{25,37}$.

Figure 6 summarizes the binding energies with respect to alloy composition. The fitted line indicates that binding energies are not linear as a function of alloy composition (i.e. Vegard’s law is invalid in this case). The
deviation from linearity of the binding energies of $\text{Si}_{1-x}\text{Ge}_x$ ($E_b^{\text{SiGe}}$) as a function of composition can be described via the following relation:

$$E_b^{\text{SiGe}} = (1-x)E_b^{\text{Si}} + xE_b^{\text{Ge}} + x(1-x)\theta$$

Where $\theta$ is called the bowing parameter and is calculated to be $-2.0$ eV. Although the trend is very similar, this bowing parameter is considerably higher as compared to the one calculated in previous work. This is due to the more substantial SQS cells (32 atom SQS as compared to 16 atom SQS in ref.5) and the more $\text{Si}_{1-x}\text{Ge}_x$ compositions used here that allow far more $E$-centre configurations to be considered.

Figure 4. The dependence of the binding energy of PV defects in $\text{Si}_{1-x}\text{Ge}_x$ ($x = 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875$) with respect to the total nearest neighbour atoms to the P and vacancy.
Figure 5. A schematic representation of the lowest binding energy PV defects and their nearest neighbour atoms in Si<sub>1-x</sub>Ge<sub>x</sub> (x = 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875).

Figure 6. The dependence of the binding energy of PV defects with respect to concentration in Si<sub>1-x</sub>Ge<sub>x</sub> (x = 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875).
The non-linear dependence of the binding energies with respect to composition is consistent with previous work, however, the present study is far more detailed given more compositions were explored, larger SQS cells and more E-centres with richer nearest neighbor environments. What is the nature of this deviation from linearity and is it determined in other defect processes in Si$_{1-x}$Ge$_x$ alloys? The Ge content dependence of the most strongly bound P-V defects is in agreement with previous experimental studies for donor atom (As and Sb) diffusion in Si$_{1-x}$Ge$_x$ alloys. In particular these experimental investigations established that the activation enthalpies of diffusion of As and Sb are not linearly dependent as a function of the Ge content of Si$_{1-x}$Ge$_x$ alloy. More recently, Kube et al. determined the self-diffusion of Si and Ge in Si$_{1-x}$Ge$_x$ over a wide range of Ge concentration ($x=0.0, 0.05, 0.25, 0.45$ and $0.70$) and temperatures (963 K-1543 K). These investigations showed a non-linear dependence of the activation enthalpy of self-diffusion with Ge concentration with a bowing that is consistent with the present study. Thereafter, Saltas et al. analysed the experimental results within the cBΩ thermodynamic model. In this study, Saltas et al. used the thermoelastic properties (bulk modulus, mean atomic volume and thermal expansion coefficient) of Si and Ge to study the composition and temperature dependence of self-diffusion in Si$_{1-x}$Ge$_x$. The resulting deviations from Vegard’s law were attributed to the diversification of the bulk properties of Si and Ge. This is anticipated to be the reason for the bowing in the binding energies of the PV defects with respect to composition calculated in the present study, however, further thermodynamic analysis will need to be performed in future work.

Apart from relaxation and thermodynamic issues the electronic properties of the defects including charge transfer can impact the properties of Si$_{1-x}$Ge$_x$. To consider this we have performed spin polarized DFT calculations. These calculations enabled us to calculate the amount of charges on the P and its nearest neighbor atoms and plot charge densities localized on the P atoms in each configurations. The relaxed configurations of P interacting vacancies in Si$_{1-x}$Ge$_x$ are shown in Fig. 7. In each configurations, there is a substantial interaction between

![Figure 7. Relaxed structures of seven different phosphorous substitutional-vacancy defect (PV pairs) configurations in Si$_{1-x}$Ge$_x$ alloys. A clear view of chemical environment showing bond distances between Si (Ge) and P and Bader charges on P and its nearest neighbour atoms are also reported.](image)
P and Si (or Ge) is observed. This is reflected in the negative Bader charge on P and positive Bader charge on Si (or Ge) (refer to Fig. 8). The amount of charge on P in each configuration is \(-3.00\). In each configurations, the P forms three-coordination with adjacent Si (or Ge) atoms and they donate \(1.00\) e each to P to form stable \(P^{3-}\) states. This is further evidenced by the charge density around the P atoms in each configurations (refer to Fig. 8). The Si–P bond distance is calculated to be \(2.30\) Å while the Ge–P bond distance is \(2.40\) Å. This is due to the larger atomic radius of Ge than that of Si.

**Summary.** In the present study, electronic structure calculations were used to investigate the defect process of the \(E\)-centre in \(Si_{1-x}Ge_x\) for a range of compositions. It is shown that the binding energies of the \(E\)-centres are strongly dependent upon the composition of the alloy, but also on the local environment around the defect. In particular, the more bound \(E\)-centres have at least one Ge atom as a nearest neighbour, whereas the dependence of the binding energy of the \(E\)-centres with respect to alloy composition is non-linear. It is shown that the more substantial 32 atom SQS can better describe the defect processes in \(Si_{1-x}Ge_x\) as compared to 16 atom SQS cells considered previously. The present study is a paradigm of the employment of the SQS method in conjunction with systematic DFT calculations to describe non-linear energetics in random semiconductor alloys. This approach can be extended to other technologically important systems such as random alloys for materials for photovoltaics and solid oxide fuel cells.

**Methods.** The binding nature of phosphorous with vacancy defects in \(Si_{1-x}Ge_x\) was examined by using the plane wave DFT code CASTEP\(^{39,40}\). The correlation and the exchange interactions are described using the

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**Figure 8.** Surface of the constant charge density associated with the interaction of phosphorous with vacancy in each of the seven different configurations in \(Si_{1-x}Ge_x\) alloys.
corrected density functional of Perdew, Burke, and Ernzerhof (PBE), the generalized gradient approximation (GGA), BFGS (Broyden-Fletcher-Goldfarb-Shanno) geometry optimisation algorithm in conjunction with the ultrasoft pseudopotentials. The calculations involved 64-atomic site supercell, the plane wave basis set by choosing the level of convergence of the atomic energies to 0.3 eV per atom, a $2 \times 2 \times 2$ Monkhorst-Pack (MP) k-point grid. We performed seven sets of calculations for different special quasirandom structures (SQS) configurations for the following concentrations of $\text{Si}_{1-x}\text{Ge}_x$ ($x = 0.875, 0.750, 0.625, 0.500, 0.375, 0.250, 0.125$).

To understand the electronic structures of substitutional phosphorous-vacancy pairs in $\text{Si}_{1-x}\text{Ge}_x$ alloys, spin-polarized DFT calculations were performed using VASP code, which uses plane wave basis sets. The exchange correlation term was modeled using generalized gradient approximation (GGA) parameterized by Perdew, Burke and Ernzerhof. A plane wave basis set with the cut-off energy of 500 eV and a $4 \times 4 \times 4$ Monkhorst-Pack k-point mesh which yields 36 k points. Both atom positions and simulation box were relaxed using a conjugate gradient algorithm. Forces on the atoms and stress tensors in all configurations were less than 0.001 eV/Å and 0.002 GPa respectively. Semi-empirical dispersion was included in all calculations as parameterized by Grimme et al. in the VASP code. Bader charge analysis was used to estimate the charges on the substitutional atom and its nearest neighbours.

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**Author Contributions**

S.-R.G.C. and N.K. performed the calculations. All authors analyzed and discussed the results and contributed to the writing of the paper.

**Additional Information**

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