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First principles modeling of defects in the Al₂O₃/In_{0.53}Ga_{0.47}As system

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Density functional theory paired with a first order many-body perturbation theory correction is applied to determine formation energies and charge transition energies for point defects in bulk In_{0.53}Ga_{0.47}As and for models of the In_{0.53}Ga_{0.47}As surface saturated with a monolayer of Al₂O₃. The results are consistent with previous computational studies that As_{Ga} antisites are candidates for defects observed in capacitance voltage measurements on metal-oxide-semiconductor capacitors, as the As_{Ga} antisite introduces energy states near the valence band maximum and near the middle of the energy bandgap. However, substantial broadening in the distribution of the Ga_{As} charge transition levels due to the variation in the local chemical environment resulting from alloying on the cation (In/Ga) sublattice is found, whereas this effect is absent for As_{Ga} antisites. Also, charge transition energy levels are found to vary based on proximity to the semiconductor/oxide interfacial layer. The combined effects of alloy- and proximity-shift on the GaAs antisite charge transition energies are consistent with the distribution of interface defect levels between the valence band edge and midgap as extracted from electrical characterization data. Hence, kinetic growth conditions leading to a high density of either GaAs or AsGa antisites near the In_{0.53}Ga_{0.47}As/Al₂O₃ interface are both consistent with defect energy levels at or below midgap. Published by AIP Publishing. [http://dx.doi.org/10.1063/1.4975033]

I. INTRODUCTION

As the number of transistors on a single integrated circuit has increased to the point of exceeding tens of billions for logic circuits and hundreds of billions for random access memories, critical dimensions of 10 nm or below are required for production technologies. The negative impact that electrically active defects have on device performance for larger device scales is exacerbated for nanoscale transistors; conventional problems such as gate screening due to interface and oxide charges occur to an even higher degree, and larger device to device variations are introduced that must be understood and eliminated, or at least controlled. 1-3 Nanoelectronics design finds itself in a regime of power-constrained scaling in which power density cannot be significantly increased in silicon technologies without overcoming problems related to further device scaling.^{4,5} Due to the larger electron mobility compared to silicon, III-V semiconductors such as In_{0.53}Ga_{0.47}As remain candidates for overcoming barriers to scaling into the sub 10 nm device technology nodes⁵ and for high performance applications. It should be noted that the bulk mobility for devices with channel lengths less than the electron mean free path display ballistic transport and the bulk mobility becomes less of a criterion. For ballistic transport, the current drive is determined by the source density of states and the electron injection velocity. Due to a low electronic density of states (DoS) at the conduction band edge for typical III-V materials considered for electronics, the advantage of a "high mobility" material is no longer a decisive factor for the scaled devices. The low density of states limiting the current drive in ultra-scaled devices is referred to as the DoS bottleneck.

A large defect concentration at III-V/oxide interfaces has traditionally been another significant obstacle to the integration of III-V materials into mainstream complimentary metal-oxide-semiconductor (CMOS) technologies, as the electrostatic control of the semiconducting channel's charge density at the semiconductor surface by the gate electrode becomes screened by charged defect states. Determination of the atomic structure of electrically active defects aids developing means to either eliminate the formation of the defects, or to devise schemes to passivate the defects subsequent to their formation. Recent electrical studies on metal-oxide-semiconductor (MOS) capacitors consisting of In_{0.53}Ga_{0.47}As/Al₂O₃ and In_{0.53}Ga_{0.47}As/HfO₂ allow for interface density of states (D_{it}) in the energy gap of In_{0.53}Ga_{0.47}As near the semiconductor/oxide interface to be extracted.^{6–9} Key findings include the following. The dominant interface defects are electrically active for the range of gate voltages typical for device operation; the defects are believed to be associated with the semiconductor and largely independent of the specific gate oxide material; the highest defect density within the bandgap occurs between the valence band maximum (VBM) and midgap, ⁶ with a donor-like character. Acceptor-like defect states, which can also be detrimental to device operation, are also observed in the conduction band. The electrical techniques applied to extract the defect states in the bandgap cannot directly identify the atomic structure of the defects, nor provide insight into avoiding the formation of the defects, nor guide strategies for defect passivation. Hence, the purpose of the calculations presented in this study is to narrow the possible set of atomistic configurations giving rise to defects levels in the bandgap, and thereby motivate the development of growth conditions and processing steps that either passivate the defects or avoid their formation.

Recent density functional theory (DFT) investigations of defects in GaAs and InGaAs alloys have been reported. These studies summarized in Ref. 10 provide predictions for the stability and amphoteric nature of native defects in bulk and interface models using hybrid exchange-correlation (XC) functionals, with the hybrid functionals chosen to overcome the DFT bandgap problem. In such approaches, a fraction a of Hartree-Fock exchange is mixed into the XC functional (giving rise to a "hybrid"), and the value of a is usually adjusted to reproduce the experimental bandgap. The authors 10,11 conclude that the position of the midgap charge transition levels (CTLs) of the As_{Ga} antisite, combined with a predicted lower formation energy relative to other commonly studied point defects, suggests that this antisite is responsible for the midgap D_{it} states observed in the capacitance-voltage (CV) response of In_{0.53}Ga_{0.47}As/high-k oxide MOS capacitors. In addition, As dangling bonds have also been suggested to give rise to D_{it} distributions below midgap, and cation dangling bond defects have been proposed for D_{it} features in the conduction band. ¹² In other works, studies of bonding mechanisms at the III-V/oxide interfaces GaAs/Al₂O₃ and GaAs/HfO₂ have been presented, 13 and predictions of charge transition levels (CTLs) of As and P vacancies at (110) oriented GaAs and InP surfaces are reported in Ref. 14. The latter utilizes many body perturbation theory (MBPT) to avoid the need to empirically parameterize the XC functional. The approach to determining CTLs presented in Ref. 14 is applied in the present

To explore the influence of the position of the defect with respect to the semiconductor/oxide interface, a comparison is performed using a 64 atom simulation cell with periodic boundary conditions to model bulk In_{0.53}Ga_{0.47}As with 17 In and 15 Ga atoms randomly distributed on the cation sublattice; further details are given in Section II. This structure is chosen as a reference configuration for the defects forming in the "bulk." Three point defects are studied with the bulk simulation cell: the antisites Ga_{As} (Ga on an As site) and As_{Ga} (As on a Ga site), and a Ga vacancy, the latter denoted as V_{Ga}. Only point defects in the semiconducting region are chosen due to the experimental indication that the measured density of interface defects is largely independent of the oxide. 8,15,16 This new set of calculations allows for a comparison between the use of hybrid functionals and the DFT + GW approach described in Section II. The DFT + GW approach allows for calculations free of the empirical parameterization introduced to calibrate the theoretical bandgap to the experimental value as required with hybrid functional approaches. Additionally, the explicit influence of alloying on the cation sublattice on the GaAs CTL is investigated. To explore the effects of the chemical environment on the antisites GaAs and AsGa that form within the vicinity of a semiconductor/oxide interface, an interface model is introduced with Al₂O₃ passivation of a (100) In_{0.53}Ga_{0.47}As surface. The CTLs of these point defects are re-examined for varying positions in the semiconducting region relative to the interfacial monolayer. A surface cleaved at an arsenic layer of atoms is used in building the oxide terminated surface model to reflect that for growth of indium gallium arsenide by metal organic chemical vapor deposition (MOCVD) that the samples are typically cooled in the presence of an AsH₃ flux. While previous calculations have suggested that the interfaces may involve Ga-O bonding, ^{17,18} As terminated surfaces are used in this work as a means to describe the As-rich conditions reported in experiments.

Defect formation energies have been studied as a function of chemical potential to assess the stability of defects in anion-rich, cation-rich, or varying stoichiometric conditions 10,19,20 for defects in a bulk-like environment. CTLs calculated by evaluating changes to formation energies have been studied for bulk defects¹⁰ and for the arsenic dimer (As₂), which can only be constructed from an explicit interface model.¹¹ Key results from these calculations are summarized in Fig. 1. This work focuses on the evaluation of CTLs and extends previous studies by considering defects in a bulk-like region and in regions only a few atomic distances from an explicit alumina terminated semiconductor interface model, as well as including the effects of alloying on the cation sublattice. DFT + GW is employed to aid in reducing errors when determining electron affinities and ionization potentials required to add or remove charges to the defect sites. In our calculations, the following broadening mechanisms are taken into account: the broadening of energy levels due to proximity to the interfacial layer, the local environment due to alloying, and the effects of thermal broadening on the distribution of defect states. These considerations allow for an analysis that can be directly compared to

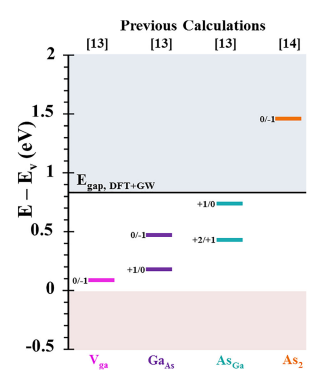


FIG. 1. A summary of previous calculations of charge transition levels employing hybrid density functional theory for bulk defects 10 and the interface defect As₂. 11

experimental capacitance-voltage measurements. This will be shown to lead to conclusions consistent with previous studies and also points to additional defects that can give rise to experimentally determined interface defect states.

II. METHODS

This section describes the methods for calculation of the formation energies and CTLs of defects in the bulk and at interfaces using models developed for In_{0.53}Ga_{0.47}As bulk and an Al₂O₃ terminated In_{0.53}Ga_{0.47}As surface. Formation energies are given as the energy required to form a defect relative to the energy of a defect-free supercell achieved by adding and/or removing atoms to and/or from an ideal atom reservoir. All energies are for configurations relaxed to minimize the total cell energy. CTLs occur at the value of the Fermi level for which the charge state of a defect changes formally by one electronic charge. Charged state formation energies are decomposed into two contributions following the approach described in Ref. 14: a structural relaxation energy term Δ and an electron addition (removal) term A (I), given by the vertical electron affinities and ionization potentials, respectively. A DFT approximation is retained for the relaxation term Δ which is found as the energy difference between two supercells with the same number of electrons thereby avoiding the difficulties commonly associated with approximate XC functionals when treating systems with varying numbers of electrons. ^{21,22} The GW approximation as a quasiparticle theory is well equipped to treat addition or removal of charges into a system. The more computationally demanding GW approximation is reserved for the calculation of electron affinities A and ionization potentials I.

The DFT calculations are performed using normconserving pseudo-potentials and the Perdew-Zunger form of the local density approximation (LDA) to the XC functional²³ to determine total energies for the relaxed supercell configurations. The relaxations of the atomic positions within a supercell and atomic reference energies are calculated with Quantum Espresso.²⁴ Relaxation of the lattice vectors for the defect free cell circumvents any possible errors due to non-equilibrium cell volumes or lack of convergence with respect to energies as discussed in Ref. 19. Following the structural relaxation of the pristine cell, point defects are inserted and the atomic positions within the supercell are allowed to relax to a new minimum energy configuration. This procedure is also repeated for each charge state of a defect with a compensating uniform background charge to prevent divergences of the Coulomb interactions between periodic image charges. Regarding the numerical details of the DFT calculations and subsequent GW corrections, the following parameters were applied to pristine and defect simulation cells: a 60 Rydberg kinetic energy cutoff, 2×2 \times 2 (2 \times 2 \times 1) k-point meshes for geometry optimization of the bulk (surface) defects. For the bulk cell, 972 unoccupied bands are used to calculate the self-energy required for the GW correction. For the bulk cell simulation cells, 972 unoccupied bands are used to calculate the self-energy term in the GW correction. For the surface models, 1772 unoccupied bands along with a value of $\sqrt{N} = 5041$ are used, where N is the number of elements in the dielectric matrix. The value of N was chosen based on the convergence of the bandgap of the pristine cells. For the bulk cell, this leads to a highly converged bandgap of 0.84 eV from the *GW* calculation, in good agreement with the experimental low temperature bandgap of 0.82 Ev. ²⁵ For the slab models, a convergence error of less than 10 meV in the bandgap is achieved.

For the case of bulk defects, a 64 atom supercell with 17 In and 15 Ga atoms randomly distributed on the cation sublattice, which hosts a representative distribution for arsenic sites with a varying number of In/Ga nearest neighbors, is selected; the distribution of nearest neighbors about the arsenic sites is shown in Fig. 2. The slightly asymmetric distribution about $N_{As-In}=2$ is consistent with the stoichiometry of the cation sublattice in $In_{0.53}Ga_{0.47}As$. Defects are introduced into these cells, and formation energies and structural relaxation energies Δ are calculated. The A addition and removal I energies are calculated from the GW approximation using the relaxed geometries obtained from DFT.

For the surface model, passivation Al₂O₃ is chosen to compare with recent experimental data.8 The building of interfacial models to represent oxides on III-V surfaces requires more advanced considerations to achieve passivation relative to Si/SiO_x interfaces.²⁶ The interface model proposed by Robertson and Lin for Al₂O₃ passivation¹³ is used; this model satisfies the electron-counting rules and results in a stable, intrinsic (bandgap is free of defect states and the Fermi level lies at midgap) semiconducting region. The semiconducting region bonding directly to the oxide consists of an As layer. The bottom of the slab is an Asterminated surface passivated by pseudo-hydrogens with charges q = 0.75. For the structural relaxation of the point defects in Al₂O₃ passivated (100)-surface In_{0.53}Ga_{0.47}As models, 304 atoms (208 semiconductor atoms, 40 oxide atoms, 24 H, and 32 pseudo-H) supercells are used resulting in the periodic images for the defects being separated by approximately 16 Å (the supercell size is doubled in the x and y directions compared to that shown in Fig. 3). Defects are introduced into the model and formation energies

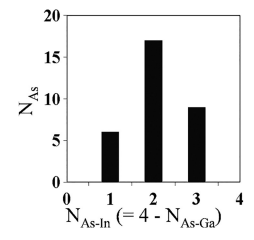


FIG. 2. Histogram associated with the number of cations bonding directly to each As atom in the 64 atom bulk cell. N_{As} is the number of As atoms with $N_{As\text{-}In}$ bonds to indium atoms.

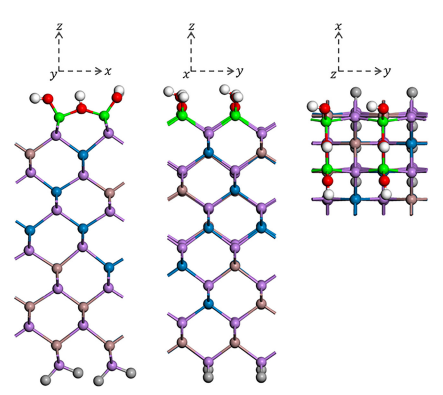


FIG. 3. 1 \times 1 unit cells of (100) InGaAs passivated with Al₂O₃. x, y, and z supercell directions are shown at the top. White, green, and red spheres indicate H, O, and Al, respectively. Pink, blue, and brown spheres indicate As, In, and Ga, respectively. Grey spheres on the bottom are for pseudohydrogens with valence = 0.75.

calculated. Periodic images of the slabs in the direction normal to the interfacial layer are separated by $\sim 18 \,\text{Å}$ of vacuum. These distances minimize the interaction between periodic images of defects while maintaining a reasonable computational effort. For the distribution of In and Ga atoms in the surface model, we investigate the use of special quasirandom structures (SOS)^{27,28} to define the cation sublattice. While the SQS described in Ref. 27 is constructed to mimic the multi-site correlation functions of a bulk fcc A_{0.5}B_{0.5} alloy, application of the SQS8 configuration to the 304 atom (100)-surface model results in a composition of In_{0.53}Ga_{0.47} for the cation sublattice due to the truncation of the structure in the slab model; the resulting cation configuration is representative of a random In_xGa_{1-x} alloy with a slight excess of In content as reflected in the histogram of As-In/Ga bonds shown in Fig. 4. As in the 64-atom bulk case, the small asymmetry about $N_{As-In} = 2$ is consistent with a small excess

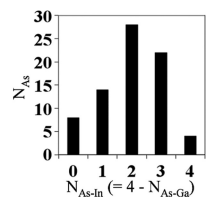


FIG. 4. Histogram associated with the number of cations bonding directly to each As atom which is 4-fold coordinated to the native cation sub-lattice in the 304 atom surface model. N_{As} is the number of As atoms with N_{As-In} bonds to indium atoms.

of In content on the cation sublattice. Thus, both the surface model and bulk model used in this work adopt cation configurations that are both representative of random alloys and maintain compositions consistent with each other, as well as with recent experimental studies.⁸

Due to the computational demand, smaller 152 atom supercells were used to compute the GW correction; smaller supercells for the electron addition energy term has been used previously when employing the GW correction for both neutral and charged cells.¹⁴ The justification for the use of smaller cells hinges on the lack of electrostatic (Hartree) contributions to the self-energy correction Σ , i.e., the latter only involves exchange and correlation terms leading to the supercell dimensions having a less pronounced effect on Σ .

Defect formation energies are calculated as

$$E_{form}(D) = E(D) - E(ref) + \sum_{a} n_a E(a), \qquad (1)$$

where E(D) is the total energy of a neutral relaxed simulation cell containing a single defect, E(ref) is the energy of a neutral relaxed simulation cell of the pristine host, E(a) is the energy of an isolated atom, and $n_a = +1$ (-1) if atomic species a is removed (added). The latter term corresponds to the chemical potentials of added or removed species, i.e., the energy relative to an ideal atomic reservoir. This is analogous to the standard procedure for calculating interfacial energies, in which grand canonical thermodynamics are used to evaluate energies of surfaces and interfaces relative to the dissociated components. 29,30

The thermodynamic CTL $\varepsilon_{q/q'}$ is defined as the value of the Fermi level, where energetically competing charge states have equal charge formation energies. The energy for forming a charged defect state q, relative to charge state q' is given by

$$E_{form}(q/q') = E(q, R_q) - E(q', R_{q'}) + (q - q')\Delta\varepsilon_F,$$
 (2)

where $E(q,R_q)$ is the total energy of a simulation cell containing a single defect in charge state q with the geometry relaxed to the configuration of the qth charge state denoted by R_q . The final term $\Delta q \Delta \varepsilon_F$ is the Fermi level with respect to the valence band maximum (VBM) and accounts for the transfer of electrons to and from a charge reservoir. The slope of $E_{form}(q/q')$ is given by the coefficient of the $\Delta \varepsilon_F$ term, 31 i.e., the final change in the formal charge state of a defect after an addition/removal process as the Fermi level is swept from the VBM to the conduction band minimum (CBM). This dependence of E_{form} on the Fermi level can be used to extract the CTL $\varepsilon_{q/q'}$. The energy E_{form} is plotted as a function of $\Delta \varepsilon_F$ for differing charge states, the Fermi level at which differing E_{form} intersect corresponds to a CTL. 19,32

As the name suggests, charge transition levels involve calculations with different numbers of electrons at or near a defect site. Thus, the well-known deficiencies of DFT rooted in the lack of a derivative discontinuity in the XC functional with respect to particle number^{21,22} exacerbate the computation of CTLs within a DFT framework. For example, the underestimation of the fundamental gap which arises from the above-mentioned weaknesses can lead to a qualitatively wrong picture in which CTLs resonate with the host bands, whereas experiment and more rigorous theoretical approaches would indicate that they lie within a semiconductor's bandgap.³³ To overcome the shortcomings of DFT, the formation of charged defects can be decomposed into two contributions: the energetic cost of adding or removing an electron, and the energy change arising from the relaxation of surrounding atoms upon addition of the electron or hole. This is achieved by rewriting Eq. (2). Let $E(q, R_{q'})$ be the energy of a simulation cell with charge state q but with atomic positions optimized for charge state q'. Adding and subtracting $E(q, R_{q'})$ and grouping all terms appropriately

$$E_{form}(q/q') = E(q, R_{q'}) - E(q', R_{q'}) + E(q, R_q) - E(q, R_{q'}) + (q - q')\Delta\varepsilon_F = A(q, q', R_{q'}) + \Delta(R_q, R_{q'}, q) + (q - q')\Delta\varepsilon_F.$$
(3)

The first term on the right-hand-side, A, represents the energy to add an electron neglecting the rearrangement of surrounding atoms $E(q,R_{q'})-E(q',R_{q'})$, known as the vertical electron affinity. The second term Δ arises from the relaxation of atoms in the presence of the extra charge. The decomposition is such that the charge states relating to the energy terms in Δ do not differ. As Δ can be computed without changing the number of electrons, the problem of a discontinuous XC functional with respect to electron number does not apply to the relaxation term, 14,33 and approximate DFT can be used. However, by necessity, the charge state changes for the calculation of the electron affinity A, and the GW approximation is a suitable method to determine this term. For electron addition $(q' \rightarrow q' - 1)$, the vertical transition corresponds to the electron affinity of the defect; in this

case, the electron is absorbed from the surrounding reservoir into the defect level. It is also possible to form charge states by removal of electrons. In this case, it is assumed that the vertical electron ionization energy (I) of q to q' equals the negative of the vertical addition energy (-A) of q' to q. For electron removal $(q \rightarrow q+1)$, a vertical transition would correspond to the ionization energy of the defect, where the removed electron is transferred *from* the defect level *to* the electron reservoir. The charge state formation energy can, thus, be re-expressed in terms of the energy to remove an electron plus the energy associated with the subsequent relaxation of atoms due to the hole that is created

$$E_{form}(q'/q) = E(q', R_q) - E(q, R_q) + E(q', R_{q'}) - E(q', R_q) + (q' - q)\Delta\varepsilon_F$$

$$= I(q', q, R_q) + \Delta(R_{q'}, R_q, q') + (q' - q)\Delta\varepsilon_F.$$
(4)

It has previously been shown that the difference between a defect level position calculated using A or I, in other words the CTL obtained from forming q from q' by adding an electron (A calculated level) to charge state q' compared to the CTL obtained by removing an electron from q (I calculated level), can vary by about 200 meV. 34 A method to overcome this error is to take the average value between the highest occupied state of charge state q and lowest unoccupied state of charge state q', since this is equivalent to the energy of the highest occupied state at half occupation under the assumption that the total energy can be written as a continuous function of the number of electrons. 34,35 In fact, the discrepancy between either A or I calculated levels and energy at half occupation is of the order of 100 meV.³⁴ The error is the same order of the error intrinsic to experimental studies; we take the more computationally convenient approach and neglect the calculation of both pathways to obtain the average. The CTLs of neutral to negative transitions are then calculated from vertical electron addition energies (A), while transitions involving positive charge states utilize the downward electron removal energies. The GW method as implemented in the YAMBO code³⁶ is applied to calculate the vertical charge addition and removal energies. In this approach, a first order perturbation correction to the Kohn-Sham eigenvalues ε_{KS} are obtained to yield the quasiparticle levels

$$\varepsilon_{qp} = \varepsilon_{KS} + \langle \varphi_{KS} | (\Sigma - V_{xc}) | \varphi_{KS} \rangle,$$
 (5)

where φ_{KS} are the Kohn-Sham orbitals assumed to be sufficiently similar to the quasiparticle wavefunctions, 37 Σ is the electronic self-energy, and V_{xc} is the XC potential (subtracted to avoid double counting between the Kohn-Sham eigenvalue and the self-energy expectation value). The separation of the highest occupied and lowest unoccupied levels at the Γ -point is taken as A. In order to calculate this correction, the self-energy is required, which in turn requires a convolution of Green's function G and the screened Coulomb interaction G. The former is constructed using wave functions and the band energies acquired from a self-consistent DFT calculation, 36,37 while the latter depends on the inverse

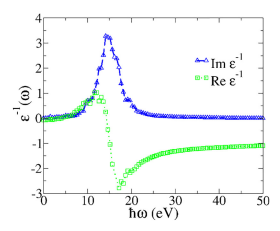


FIG. 5. Real and Imaginary parts at the Γ -point component (G = G' = 0) of the inverse of the dynamic dielectric matrix, obtained from the Al_2O_3 passivated (100) $In_{0.53}Ga_{0.47}As$ surface model, plotted as a function of energy.

dielectric response function ε^{-1} that takes into account the local field effects and the dynamics of the screened interaction. In the reciprocal space, the relation between the screened Coulomb interaction and ϵ^{-1} can be written as

$$W_{\mathbf{G},\mathbf{G}'}(\mathbf{q},\,\omega) = \epsilon_{\mathbf{G},\mathbf{G}'}^{-1}(\mathbf{q},\,\omega)\nu(\mathbf{q}+\mathbf{G}'). \tag{6}$$

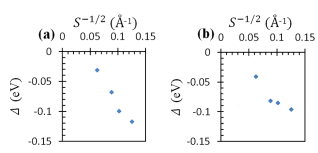
Here, \mathbf{q} is an arbitrary wave vector, while \mathbf{G} is a reciprocal lattice vector, and $v(\mathbf{q}+\mathbf{G}')$ is the bare Coulomb interaction. The local fields arise from the off-diagonal $\mathbf{G} \neq \mathbf{G}'$ elements. The connection back to DFT-calculated quantities is made from the relation between the static dielectric function and the non-interacting polarizability P obtained from the Kohn-Sham wavefunctions and eigenvalues. 37

The plasmon pole approximation (PPA)^{36,38} to describe the frequency dependence of ϵ^{-1} is made. The PPA assumes the spectral function for the screened interaction to be a single narrow peak in the plasmon energy $E (= \hbar \omega)$, where ω is the plasmon frequency).³⁸ If this condition does not hold,³⁹ the dielectric function must be explicitly computed throughout the full frequency axis leading to large increases in computational time and memory requirements. A single peak in the dielectric function is observed for bulk In_{0.53}Ga_{0.47}As, and the surface model exhibits a single peak in the imaginary part of the inverse dielectric response function as shown in Fig. 5. This allows for the application of the GW method to supercells containing >150 atoms.

Calculations with charged cells have a spurious electrostatic interaction due to the images created by the periodic boundary conditions. For the calculation of the relaxation energies of charged defects, energies are obtained as the difference between energies for simulation cells with the same charge state but in different geometries. Thus, errors associated with electrostatic interactions may, to a large degree, systematically cancel using the energy decomposition in Eq. (3), 19 assuming that the localization of the charge state is not significantly affected by atomic relaxation, as indicated by the relatively small changes in total energy between geometries. To explore the effect of the error due to interaction between periodic image charges, the Ga_{As} defect in a q = -1 charge state is taken as a test case. Fig. 6 shows the relaxation energies Δ versus the reciprocal of the square root of the surface area $1/\sqrt{S}$ where $S = a \times b$ in which a and b are the cell parameters parallel to the plane of the surface in each supercell. Four distinct (100)-surface orientation In_{0.53}Ga_{0.47}As/Al₂O₃ models are constructed consisting of 72, 114, 152, and 304 atoms. Figs. 6(a)–6(c) show the variation in the total energy differences with increasing cell size for relaxations from the neutral to negatively charged defect geometry $E(q_{-1}, R_{-1}) - E(q_{-1}, R_0)$ in (a), negative to neutral defect geometry $E(q_0, R_0) - E(q_0, R_{-1})$ in (b), and positive to neutral defect geometry $E(q_0, R_0) - E(q_0, R_{+1})$ in (c). Comparison between the three cases reveals a similar variation of the relaxation energy as a function of surface area for all the three cells. Comparison of the charged defect relaxation (a) to the change in the relaxation energy in the neutral cells reveals that the elastic defect-defect interactions dominate the relaxation energies and that the error due to the Coulomb interactions between the charged defect images is substantially less. These results are consistent with the typical estimates for a computational error of $\sim 100\,\text{meV}$ in defect formation energies from DFT, 11,19,32,33 and comparable to similar errors quoted from experimental studies^{8,40} of defect levels. Therefore, as in previous works, the electrostatic correction term in the relaxation energies is omitted due to the favorable cancellation that occurs when taking differences to obtain the formation energies. 19 The resulting estimate of the error in the formation energies due to the finite cell sizes used in these calculations are estimated to be of the order of tens of meV.

III. RESULTS

The structures for the gallium vacancy V_{Ga} , the gallium antisite Ga_{As} , and the arsenic As_{Ga} in bulk-like environments following relaxation of the atomic positions and cell parameters in the simulation cells are considered. Similar results for Ga_{As} and As_{Ga} antisites are then presented in Section III B for the surface models in the proximity of the $In_{0.53}Ga_{0.47}As/Al_2O_3$ surface, and for the Ga_{As} antisite when formed



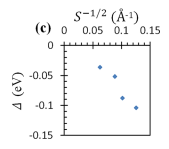


FIG. 6. Energy difference $\Delta=E(q_{-1},R_{-1})-E(q_{-1},R_0)$ for (a), $\Delta=E(q_0,R_0)-E(q_0,R_{-1})$ for (b), and $\Delta=E(q_0,R_0)-E(q_0,R_{+1})$ for (c), all versus $S^{-1/2}$. Variation in Δ from the largest to smallest supercell is 85 meV for (a), 68 meV for (c), and 55 meV for (b).

directly at the semiconductor/oxide interfacial layer are presented.

A. Bulk defects

1. Structural properties

Displacement of the atom's neighboring neutral defects is measured relative to a pristine (defect-free) cell, while displacements of the neighboring atoms for the charged defects are measured relative to the relaxed neutral defects. Focusing on the Ga vacancy (V_{Ga}), removal of a Ga atom results in the four surrounding As atoms relaxing inward towards the vacancy site on average by 0.5 Å. The tendency of the surrounding anions forming weak bonds across a cation vacancy has been noted in previous computational studies on III-Vs materials. 41 The structural effect of charging the vacancy is slight compared to the structural changes associated with the formation of the vacancy, leading to small inward displacements on the order of $\sim 0.01 \,\text{Å}$ for the four neighboring anions when an electron is added. Removing an electron from the neutral state and allowing the defect site to relax lead to a positively charged vacancy with a similar small degree of relative change in the defect configuration but with the four neighboring As atoms becoming displaced outwards away from the vacancy site.

The geometry of the Ga_{As} antisite is studied for the configuration, where the antisite bonds with two Ga and two In atoms. Relative to the unrelaxed substitutional site, the bonds between the neutral defect and the surrounding Ga atoms shorten by 0.08 A, whereas the bonds between the defect and the nearest-neighbor In atoms elongate by 0.04 A. Thus, the GaAs remains in a four-fold coordinated arrangement but moves slightly away from the tetrahedral symmetry of the non-defective As site. Charging the Ga_{As} site to a positive state pushes the defect center toward the two nearest neighbor Ga atoms by 0.04 Å compared to the position of the neutral defect. Charging this antisite to a negative charge state results in a slight movement of the defect center by 0.03 Å compared to the neutral position but in the opposing direction to the positive state relaxation, and toward the two In nearest neighbors.

For the neutral As_{Ga} antisite, the surrounding bonds exhibit larger changes relative to the bond length changes upon formation of Ga_{As} : an outward relaxation of 0.16 Å relative to the bonds of As bonding to Ga. Charging to a positive state causes the As- As_{Ga} bonds to contract by 0.05 Å relative to the neutral case, with tetrahedral symmetry preserved and the defect center remains in the same position relative to the relaxed neutral charge state. This displacement is larger for a doubly charged state of the As_{Ga} antisite; an average of 0.06 Å contraction of the anion antisite bond lengths relative to the +1 charge state is found.

For all defects studied, the bond lengths located along atomic planes which bisect the spacing between periodic images of the defect center undergo displacements of less than 0.01 Å on average, a measure relative to the corresponding pristine cell that indicates the defect-defect interactions are limited for the selected cell size. This indicates that spurious elastic effects due to periodic boundary conditions are

reduced to acceptable levels in our bulk model and is consistent with the finding of Van de Walle and Neugebauer that 64-atom cells are sufficient to estimate the relaxation energy associated with a point defect in bulk semiconductor calculations. Furthermore, as will be seen, the relatively small relaxations of the neighboring atoms upon changing charge state relative to the relaxed neutral defect geometry suggest that the relaxation energies will tend to be small compared to the vertical charge addition and removal energies.

2. CTLs for bulk defects

Bulk CTLs for As and Ga antisites (As_{Ga}, Ga_{As}) and the As and Ga vacancies (V_{As}, V_{Ga}) calculated with hybrid DFT have been reported. 10 For the V_{Ga}, a neutral to negative transition denoted as $\varepsilon_{0/-1}$ is found to occur at an energy $0.08\,eV$ above the VBM. A $\epsilon_{+1/0}$ transition for the As_{Ga} antisite occurs at 0.74 eV above the VBM, while the $\varepsilon_{+2/+1}$ transition for the As_{Ga} defect lies close to midgap at 0.42 eV above the VBM. The latter is found to be 50 meV lower than the $\varepsilon_{0/-1}$ transition for Ga_{As} , while the $\varepsilon_{+1/0}$ Ga_{As} transition lies slightly above the $\varepsilon_{0/-1}$ V_{Ga} transition found from hybrid DFT calculations. In the work reported in Ref. 10, a study of the defect formation energies as a function of growth conditions approximated by varying the chemical potentials of the added/removed species is also performed. Taking this study together with the proximity of the As_{Ga} transition to the midgap D_{it} feature measured at InGaAs/ oxide interfaces by electrical spectroscopy, 6-8 those authors assign the As_{Ga} antisite with the transition level $\varepsilon_{+2/+1}$ as a strong candidate for the defect responsible for experimentally extracted midgap defect densities. The V_{As} vacancy is also considered in Ref. 10; it is shown that this defect exhibits a positive to neutral CTL very close to the CBM, and no midgap CTLs. This finding is coupled with the fact of a high cost of formation of V_{As} for substrates grown under As-rich conditions⁹ to rule out V_{As} as a candidate for midgap D_{it} . In addition, under (approximated) Ga-rich growth conditions, the Ga_{As} antisite is found to be more stable than V_{As} , ¹⁰ which also suggests the relatively small contribution of V_{As} to midgap D_{it} , even for growth conditions corresponding to low As concentration. It should also be noted that V_{As} is situated on an anion site, and hence, can be subjected to alloying effects due to changes in the cation sublattice. The influence of such effects on the energy of V_{As} has already been studied in detail.⁴² Murphy et al. found a large variation (~500 meV) of the V_{As} formation energy as a function of variations in the alloy. However, the authors of Ref. 42 did not consider anion-situated antisites in their work nor did they report calculations of CTLs as a function of the local alloy. We also investigate the role of local alloying on an anion situated defect; however, due to the aforementioned findings regarding the midgap D_{it} candidacy of V_{As} , combined with a lack of studies of the effect of alloying on CTLs of antisites, we exclude V_{As} from this work, and instead focus on the effects of changes to the local cation alloy on the Ga_{As} antisite.

The energetics for V_{Ga} , As_{Ga} , and Ga_{As} defects determined within the DFT + GW approach are computed and

compared to the hybrid DFT results. The transition to a neutral V_{Ga} defect is obtained by adding an electron to the relaxed geometry of the positively charged defect and subsequently relaxing the geometry of the simulation cell of the neutral defect. The energy difference between the neutral cell in its relaxed geometry and the neutral cell with the atomic positions fixed to those of the +1 charge state results in a relaxation energy of $\Delta = -0.01 \,\text{eV}$. The DFT + GW approach yields a vertical electron affinity of 0.07 eV, which results in a charge state formation energy of $E_{form}(q_0/q_{+1})$ $= 0.06 \, eV$ for the neutral V_{Ga} defect. The -1 charge state is formed by adding an electron to the neutral defect. This yields $E_{form}(q_{-1}/q_0)$ of 0.21 eV for the V_{Ga} defect. As anticipated from the analysis of the charged defect geometries, both charge state relaxation energies for the vacancy are relatively low. The charge formation energies vary as the Fermi level ε_F is varied and a CTL occurs as the energies cross and a new charge state becomes more stable,

$$\varepsilon_{q/q'} = \frac{E_{form}(q') - E_{form}(q)}{q - q'} , \qquad (7)$$

for V_{Ga} q=0 and q'=-1, the CTL $\varepsilon_{0/-1}$ occurs at $0.15\,\mathrm{eV}$ above the VBM.

For the +1 charge state of the Ga_{As} antisite in the relaxed configuration, an electron affinity of $0.11\,\mathrm{eV}$ is obtained for the DFT + GW approximation. Together with the structural relaxation contribution, this yields a formation energy of $0.06\,\mathrm{eV}$ for the neutral charge state of Ga_{As} . Forming the negative and positive charge states by adding and removing an electron to and from the neutral state, the CTLs $\varepsilon_{+1/0}$ and $\varepsilon_{0/-1}$ are found to lie at $0.19\,\mathrm{eV}$ and $0.31\,\mathrm{eV}$ above the VBM, respectively.

For the As_{Ga} antisite, the +1 and +2 states are formed by successive removal of electrons starting from q=0. The GW-corrected charge transition energies for As_{Ga} are found slightly below midgap for the +2/+1 transition and close to the experimental CBM for the +1/0 transition. In all the cases, the magnitude of the relaxation energy Δ reflects the degree of structural rearrangement between the differing charge states; the Δ term averages $-97\,\text{meV}$ for As_{Ga} , whereas for Ga_{As} the average is $-49\,\text{meV}$.

Comparing the values for $\varepsilon_{q/q'}$ obtained by the DFT + GW approach with those reported in Ref. 10 determined from a hybrid DFT approach, reasonable agreement is found for the CTLs occurring near the VBM and the CBM. CTLs occurring deeper in the gap tend to be slightly lower in energy compared to the hybrid DFT results. In general, the bulk defect transition levels calculated by the DFT + GW approximation agree with hybrid DFT results typically within 100 meV for shallow defect levels ($\varepsilon_{+1/0}$ for V_{Ga} , Ga_{As} , and As_{Ga}) and within 200 meV for deeper levels ($\varepsilon_{0/-1}$ and $\varepsilon_{+1/0}$ for Ga_{As} and As_{Ga} , respectively).

Having focused on the As_{Ga} antisite as a likely candidate for the midgap defect states, previous studies have not reported specific values of CTLs of the Ga_{As} antisite as a function of alloying on the cation sublattice. ^{10,42} To examine the influence of local cation disorder, the atoms bonding directly to the Ga_{As} antisite are replaced with either four Ga

nearest neighbors or with four In nearest neighbors, recalling that the initial calculation consisted of a configuration with two Ga and two In nearest neighbors denoted as In_{0.5}Ga_{0.5}-NN. When the defect is surrounded by four nearest neighbor Ga atoms, denoted $In_{0.0}Ga_{1.0}$ -NN, the CTL $\varepsilon_{0/-1}(Ga_{As})$ increases by $60\,\text{meV}$ to $0.37\,\text{eV}$. The $\varepsilon_{+1/0}(Ga_{As})$ CTL moves to within 20 meV of the VBM for the In_{0.0}Ga_{1.0}-NN configuration. These changes, relative to the case of In_{0.5}Ga_{0.5}-NN, occur mainly through a decrease in the magnitude of the electron addition energies in the positive charge states $A(q = +1, R_{q=0})$ and $A(q = +1, R_{q=+1})$, on average by 76 meV, while the electron addition energy of the neutral charge state $A(q = 0, R_{q=0})$ decreases by 32 meV. All the relaxation energies (Δ) maintain their values within 10 meV compared to the values obtained for the In_{0.5}Ga_{0.5}-NN configuration revealing that the largest change to the electron affinities and ionization potentials are due to the change in the electronegativity of the defect due to local changes in the chemical environment. When the GaAs antisite is bonded to four In atoms In_{1.0}Ga_{0.0}-NN, the $\varepsilon_{0/-1}(Ga_{As})$ CTL increases by approximately 55 meV compared to the In_{0.5}Ga_{0.5}-NN case resulting in a level at 0.37 eV relative to the VBM. The +1/0 CTL $\epsilon_{+1/0}(Ga_{As})$ = 0.16 eV for In_{1.0}Ga_{0.0}-NN configuration and is approximately 30 meV lower than for the In_{0.5}Ga_{0.5}-NN case. While the 0/-1 CTL is approximately equal for the two cases In_{0.0}Ga_{1.0} NN and In_{1.0}Ga_{0.0} NN, the latter exhibits a slightly increased value of $A(q = 0, R_{q=0})$ of 11 meV compared to $In_{0.5}Ga_{0.5}$ NN configuration. The value of $A(q = +1, R_{q=+1})$ decreases by 58 meV relative to the In_{0.5}Ga_{0.5}-NN bonding configuration and $A(q = +1, R_{q=0})$ increases by 11 meV. Again, all relaxation energies for In_{1.0}Ga_{0.0} NN change by less than 10 meV relative to the In_{0.5}Ga_{0.5}-NN configuration. The largest change in the CTLs for the Ga_{As} antisite as a function of local cation disorder occurs for the +1/0 transition level which lies 0.02 eV above the VBM when bonding to four Ga atoms, compared to 0.19 eV above the VBM when bonding to two In and two Ga atoms. Local cation disorder appears to have a greater effect for donor-like transitions, while acceptor-like transitions change by less than 60 meV.

The effect of changes in the cation sublattice alloy configuration has consequences for the donor-like feature of the experimentally observed D_{it} distribution.⁸ Variations in the local cation distribution can shift donor-like transition states by up to 200 meV, while the CTLs remain within the lower half of the bandgap. Therefore, one could conclude that the D_{it} feature occurring below midgap and extending lower to the valence band may have broadened contributions from the variable bonding arrangements due to the random cation alloy; this point will be examined when comparing the experimental broadening of the CTLs.

B. Surface defects

1. Structural properties

The influence of the oxide terminated semiconductor surface $In_{0.53}Ga_{0.47}As/Al_2O_3$ on the energetics of defects formed near the interfacial layer is examined for V_{Ga} , Ga_{As} ,

and As_{Ga} . The effect on defects directly bonded with atoms forming the oxide layer is also considered. The energetics of these surface defects are calculated with the effects of oxide terminating layer included.

The V_{Ga} model is created by removing a single Ga atom located on a monolayer below the oxide, this is the cation layer closest to Al_2O_3 but not directly bonding to the oxide. All the remaining atoms are allowed to relax about the vacancy site. Similar to the vacancy model in the bulk, removal of the Ga atom results in an inward movement of the atoms nearest the vacancy by an average of 0.43 Å relative to their positions prior to creation of the vacancy site. Relaxation in the positive charge state results in a slight outward movement of the atoms surrounding the vacancy by 0.01 Å, while relaxation in the negative charge state results in an inward movement of atoms towards the vacancy shown in Fig. 8(a) by 0.03 Å. The degrees of atomic displacements are again in proportion to the magnitudes of charge state relaxation.

The Ga_{As} defect in Fig. 8(b) bonds directly to the oxide layer. For the neutral defect, the defect site is displaced towards the oxide relative to the antisite position prior to performing a geometry relaxation. The defect moves away from Al atoms in the oxide and towards the nearest hydroxyl group -OH for the relaxed configuration; the relaxed GaAs-OH and Ga_{As}-Al separations are 2.12 Å and 2.59 Å, respectively, compared to the corresponding As-OH and As-Al distances prior to creating the antisite defect, which average 3.42 Å and 2.42 Å, respectively. In analogy to the bulk Ga_{As} antisite, the surface GaAs antisite moves towards the nearest Ga neighbor and away from the nearest In neighbor relative to the pristine simulation cell in the absence of the antisite. Relative to the bulk defect, the relaxed surface Ga_{As}-Ga bond length is 0.07 Å less than the corresponding surface As-Ga bond length with a 0.08 Å reduction for the bulk Ga_{As} case. The relaxed surface GaAs-In distance increases by 0.16 Å relative to the corresponding surface As-In separation compared to only a $0.04\,\text{Å}$ increase of the same bond length for the bulk case. Charging to q_{+1} results in a further contraction of the Ga_{As} -OH distance to $2.09\,\text{Å}$, while the Ga_{As} -Ga and Ga_{As} -In separations both increase by an average of $0.02\,\text{Å}$ relative to the neutral defect. Adding an electron to the neutral state to bring the charge state to q_{-1} results in a Ga_{As} -OH separation of $2.13\,\text{Å}$, a slight increase relative to q_0 state. The Ga_{As} -Ga and Ga_{As} -In separations both decrease by an average of $0.02\,\text{Å}$ relative to q_0 geometry; i.e., the q_{-1} relaxation is almost the same magnitude but opposite in direction compared to q_{+1} . These relaxations occur mainly through a movement of the defect center towards a -OH and away from the neighboring cations for q_{+1} , and away from -OH and towards the cations for q_{-1} .

The As_{Ga} antisite at the surface is depicted in Fig. 8(c) and is not bonding directly to the oxide as the surface of the semiconductor layer is cleaved at a plane of As atoms. Hence as the nearest neighboring bonds are similar to the bulk As_{Ga} defect in terms of structural rearrangement. The As_{Ga} -As bonds are on average 0.12 Å greater than the Ga-As bonds in the simulation cell in the absence of the defect. Removing an electron and relaxing in the positive charge state results in 0.02 Å reduction in the As_{Ga} -As bond lengths relative to q_0 , and removing another electron to bring the charge state to q_{+2} yields a further 0.02 Å shortening of As_{Ga} -As bond lengths relative to q_{+1} .

2. CTLs for surface defects

The arsenic dimer has been calculated for the neutral to negative charge transition, to provide a reference point against previous calculations. In good agreement with Ref. 11, we also find that the CTL is positioned into the conduction band, see Fig. 7.

For the Ga_{As} antisite bonding directly to Al_2O_3 , CTLs of $\varepsilon_{0/-1}=0.61\,\text{eV}$ and $\varepsilon_{+1/0}=0.22\,\text{eV}$ above the CBM are obtained. Thus, as in the case of the dimer which is likewise

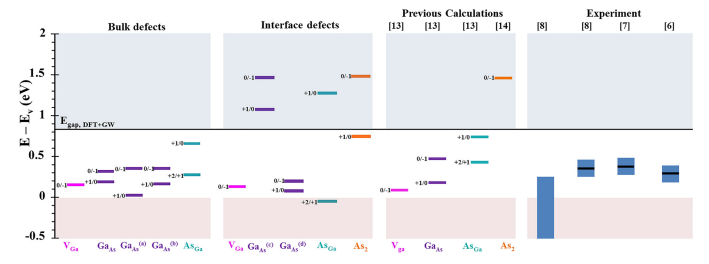


FIG. 7. A comparison of CTLs from this work for bulk (first panel) and interfacial defects (second panel), previous theoretical work 10,11 (third panel and as also depicted in Fig. 1), and experimentally extracted defect states $^{6-8}$ (fourth panel, levels are shown as dark lines and the broadening/distributions are approximated by the blue rectangles). A bandgap of $0.84\,\text{eV}$ has been calculated for pristine bulk $In_{0.53}Ga_{0.47}As$ within the DFT +GW approach using the converged parameters discussed in Section II. $Ga_{As}^{(a)}$ and $Ga_{As}^{(b)}$ refer to the Ga antisite bonding to 4 Ga (a), and bonding to 4 In atoms (b). $Ga_{As}^{(c)}$ is the Ga antisite calculated in the interface model and bonding directly to the oxide, while $Ga_{As}^{(d)}$ is the Ga antisite located 2 monolayers away from the oxide. In the third panel V_{Ga} , Ga_{As} , As_{Ga} refer to bulk defects, whereas the As_2 defect level is for the interface model.

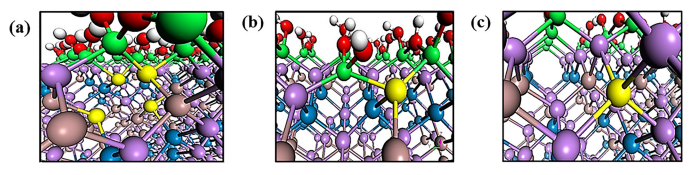


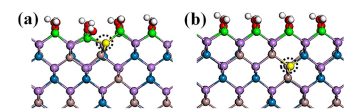
FIG. 8. Relaxed structures of neutral defects ((a) V_{Ga} , (b) Ga_{As} , (c) As_{Ga}) in (100) InGaAs:Al₂O₃. Blue, brown and, pink spheres represent In, Ga, and As, respectively. Red, white, and green spheres represent O, H, and Al, respectively. Defect atoms are highlighted in yellow in each image.

occurring directly at the $In_{0.53}Ga_{0.47}As/Al_2O_3$ interface, this antisite defect exhibits CTLs residing well above the conduction band energy of $In_{0.53}Ga_{0.47}As$, and hence, is not expected to play a role in the experimental midgap D_{ii} . For the defects studied to date, defects bonding directly with the oxide tend to have the CTLs lying much higher in energy due to the distinctly different local chemical bonding.

A V_{Ga} defect created in the Ga layer separated from the oxide layer by the single plane of surface As atoms, which terminate the semiconductor slab, yields CTLs which are relatively close to the values obtained for the same defect in a bulk environment, but with a slightly lower energy level with $\epsilon_{0/-1}=0.14\,\mathrm{eV}$ relative to the VBM, significantly lower than the experimentally reported midgap states but certainly possibly contributing to the high density of defects found near the VBM.

Turning now to the As_{Ga} antisite reveals the same qualitative picture as the Ga_{As} antisite and the As₂ dimer; defects in close proximity to the oxide, either bonding directly to or a monolayer away from the oxide, do not exhibit charge transition levels which match the energies corresponding to the distribution of midgap interface states extracted from CV measurements. In the case of the As_{Ga} antisite in close proximity to the oxide, the defect does not bond directly to Al₂O₃ in the As-terminated (100) In_{0.53}Ga_{0.47}As surface model as can be seen in Fig. 8(c). For this configuration, the CTLs are found to be $\varepsilon_{+2/+1} = -0.06$ eV relative to the VBM and $\varepsilon_{+1/0} = 0.43 \,\mathrm{eV}$ relative the CBM, or in other words, within the valence and conduction bands, respectively, and do not give rise to states in the semiconductor bandgap. This result for the $\varepsilon_{+1/0}$ level is in contrast to previous calculations of the As_{Ga} antisite near the oxide in a model of the GaAs/ Al₂O₃ interface⁴³ calculated within a hybrid DFT framework, which exhibited little difference to the corresponding CTLs of the bulk AsGa antisite. The discrepancy may be ascribed to a combination of factors including that the wide GaAs bandgap may result in significantly less hybridization between the band edges and the respective charged defect states compared to the narrower bandgap In_{0.53}Ga_{0.47}As material. Significantly, the models of the oxide bonding to the semiconductor substrate differ substantially. Interfacial bonding with O atoms bonding directly to a Ga in turn bonded to an As atom about the antisite is studied in Ref. 43, whereas the metal (Al) atoms of the oxide are bonding directly to As atoms at the semiconductor surface in the

model used in this study, which are in turn bonding to the As atom corresponding to the antisite. The local chemical environment in Ref. 43, in this sense, is more "bulk-like" and the differences highlight how significantly differing electrical properties arise from differences in the local chemical environment of a defect. The shift in the CTLs between the bulkand the As_{Ga} configuration occurring near the interfacial layer is found to be significantly smaller than the CTL shifts for the bulk- and surface-model of the GaAs antisite bonding directly to the oxide; the latter sites have a different local chemistry due to the bonding mechanism to oxide atoms, and hence, the larger associated differences in the CTLs relative to bulk defects are not unexpected. If a Ga_{As} anti-site is introduced into the arsenic layer nearest the interface but not bonding to the oxide as can be seen in Fig. 9, the CTLs are much nearer to the corresponding values occurring in the bulk with a value of $\varepsilon_{0/-1} = 0.2 \,\text{eV}$ relative to the VBM, although still lower in energy by 0.11 eV relative to the same transition in the bulk. A similar shift towards the VBM is also found for the positive to neutral CTL of the Ga_{As} antisite in proximity to the interfacial layer with a value of $\varepsilon_{+1/0} = 0.08 \, eV$ for this case. While these values are much closer to the corresponding values in the bulk case, these states are significantly lower in energy than the midgap D_{it} states extracted from measurement. However, clearly these states can be associated with the large defect density seen near the VBM in most experiments. The calculations also indicate that bonding to the oxide plays a significant role in the position of defects states and that it is unlikely, for the defects considered, the bonding directly to the oxide has a significant role in introducing defect levels within the bandgap of In_{0.53}Ga_{0.47}As.



a. Defect formation energies—bulk versus surface. A comparison of the formation of surface defects with the bulk counterparts is discussed below. Their formation energies are tabulated in Table I.

An observation for antisite stability is that bonding directly to the oxide, or bonding to the As layer that bonds to the oxide, results in less energy required to form a given defect than the energy required to form it in the bulk. However, if the antisite is two monolayers or more from the oxide, its formation energy is already within 50 meV of the bulk counterpart. Regarding the vacancy, bulk V_{Ga} defects (relaxed structure shown in Fig. 8(a)) are significantly less stable than the antisites, a trend which has been observed in the previous works 10,19,20,44 and continues to hold for defects formed close to the surface. The Ga vacancies near the surface are less stable than their bulk counterparts. The consistent trend of higher formation energies for V_{Ga} compared to other defects implies the reduced concentration of this defect for both bulk and surface variants of this defect center—although this comment is predicated upon an equilibrium argument. For the non-equilibrium conditions that occur during growth, the formation energies are only suggestive of the probability at which various defects can be formed. For the Ga_{As} anti-site bonding directly to the oxide, the formation energy is 0.12 eV lower than the Ga_{As} anti-site in the bulk, and the latter has a formation energy that is 0.97 eV higher compared to the bulk As_{Ga} antisite.

An increase of $0.17\,\mathrm{eV}$ in $E_{form}(\mathrm{Ga_{As}})$ is calculated when this antisite is moved two monolayers away from the oxide terminated surface; refer to Fig. 9 for the configuration. This results in an overall decrease in the stability of only $50\,\mathrm{meV}$ relative to the formation energy in the bulk. Hence, even when situated within a few atomic distances from the oxide, the stability of the defect effectively resembles that of the bulk defect. However, again it is noted that the formation energetics do not accurately reflect the energies of the surface during growth and the growth kinetics can alter the picture suggested by the formation energies for defects calculated relative to an ideal interface. Comparing $E_{form}(\mathrm{As_{Ga}})$ between the bulk and oxide-terminated surface cases, the $\mathrm{As_{Ga}}$ antisite exhibits a $0.18\,\mathrm{eV}$ increase in the formation energy when the defect is moved from the surface towards the bulk chemical environment.

IV. ALLOY BROADENING

The shift in CTLs with respect to a large range of defect formation conditions has been considered. These include the

TABLE I. Formation energies of bulk and surface defects calculated using total energy differences within DFT-LDA (see Eq. (1)). GaAs - Al_2O_3 refers to the Ga_{As} antisite bonding directly to Al_2O_3 . The formation energy of the Ga_{As} antisite calculated within the surface model but located two monolayers away (see Fig. 9) from the oxide is within 50 meV of the bulk counterpart. For V_{Ga} and As_{Ga} calculated in the surface model, these defects bond to the As layer that bonds to Al_2O_3 . All energies in electron volts (eV).

Defect	E_{form} , $bulk$	$E_{\it form}$, surface
V_{Ga}	6.94	7.09
Ga_{As}	2.86	2.91
$Ga_{As} - Al_2O_3$	N/A	2.74
As_{Ga}	1.89	1.71

cases in which the defect is formed at the semiconductor surface, near the surface, or in a more bulk like environment. Excluding the large changes in CTLs due to a defect directly bonding to the oxide, the shift in energies due to "proximity" broadening is found to be on the order of 100 meV (see Fig. 7 and compare Ga_{As} with $Ga_{As}^{(d)}$, the +/0 and 0/- transitions each differ by 110 meV). In addition, for defects formed on the anion sublattice and bonding to the random alloy of In and Ga atoms on the cation sublattice, there is a different local chemical environment based on the specific local distribution of group III atoms. The effect of the random nature of the cation alloy results in an additional broadening for some defects referred to as alloy broadening. To a first approximation, the alloy broadening is ascribed to the nearest neighbor bonds between an anion antisite or vacancy to the cation sublattice. Hence, the effect of alloy broadening is anticipated to be small for the As_{Ga} antisite, as there are no nearest neighbor bonds to group III atoms. Conversely, for an anion situated defect, there is a different local chemical environment due to bonding to the cation sublattice. Here, the effect of alloy broadening on the Ga_{As} antisite is considered in detail.

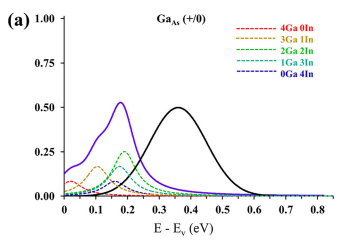
When extracting interface state density profiles from measured CV or conductance-voltage (GV) characteristics, there is an inherent thermal broadening of the actual interface state density distribution due to the occupation of states by the Fermi-Dirac distribution and the method of extraction. This is best illustrated by considering the case of a monoenergetic defect level in an MOS system for a CV measurement and an interface state density profile extraction at room temperature. Due to the finite temperature, a percentage of the interface states are occupied when the Fermi level is below the interface state energy, and as a consequence of the Fermi-Dirac distribution, the monoenergetic defect level results in a broad feature on the CV or GV response. When extracting the interface state density profile at each gate voltage (and corresponding surface potential), the difference in the high and low frequency CV characteristics, or the "stretch out" of the high frequency characteristics, is attributed to an interface state concentration in units of cm⁻²eV⁻¹ at that voltage (or surface potential). As a consequence, a monoenergetic level is extracted as a broad feature in the energy gap, and it can easily be shown that the energy distribution of the extracted interface state density profile resulting from a monoenergetic level is precisely the derivative of the Fermi Dirac distribution with respect to energy, at the temperature of the measurement and the interface state density extraction process. From this effect, the minimum thermal broadening for a CTL for measurements made at T = 300 K yields a minimum peak width of 91 meV for full width at half maximum (FWHM). Hence, each of the CTLs present in Fig. 7 would have a minimum broadening equal to the theoretical lower limit for the thermal broadening of 91 meV. For Ga_{As}, the CTLs are re-calculated with either 4 In nearest neighbors, 2 In and 2 Ga nearest neighbors, or 4 Ga nearest neighbors. The energetics for the sites with either 3 In and 1 Ga nearest neighbors or 1 In and 3 Ga nearest neighbors are interpolated. The $Ga_{As}(+/0)$ and $Ga_{As}(0/-)$ antisite CTLs for each local alloy composition are broadened

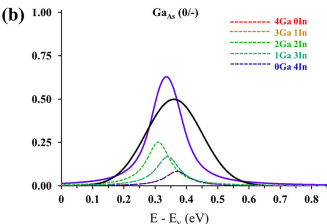
by the minimum theoretical thermal peak width, and each peak is weighted by the distribution of a random alloy on the cation sublattice. The different CTLs are then summed to give the net effect of the thermal and alloy broadening on the Ga_{As} antisite CTLs. The resulting prediction for the minimum broadening, excluding the proximity broadening, is shown in Fig. 10. The net result of accounting for both charge transitions in all 5 local alloy compositions with thermal broadening is shown in Fig. 10(c). It is interesting to note the position of the valley between the two main peaks in Fig. 10(c) (denoted by the dashed arrow). The D_{it} profile reported in Ref. 8 exhibits a valley between the midgap D_{it} peak and the broad D_{it} feature extending into the valence band at a similar position in the bandgap—approximately 250 meV above the VBM. The similarity of the broadened defect level profile of the GaAs antisite compared to the experimental D_{it} profile suggests that the Ga_{As} should not be ruled out as a midgap D_{it} candidate.

Taking into consideration the results of this analysis, the additional energy shifts to the CTLs due to defects being formed near the oxide terminated semiconductor surface, and the error inherent in the calculations, we infer the following. Although the $As_{Ga}(+2/+1)$ CTL has been identified as a candidate for the midgap states, the Ga_{As} antisite should not be ruled out as an additional strong candidate for generating the defect states at midgap in CV measurements. There have been two primary considerations that have previously focused attention on the As_{Ga} antisite. First is the experimental finding that the integrated charge across the energy gap is positive, and the second is that the calculated formation energy for the Ga_{As} antisite is on the order of 1 eV higher than the formation energy of the As_{Ga} antisite.

The first point does not rule out the Ga_{As} antisite as giving rise to the midgap states based on the experimental data. CV measurements indicate that the net charge due to the defect states integrated over the energy gap is positive. However, the midgap states form a relatively small peak and there is a much higher defect density in the gap but nearer to the valence band edge. The overall effect of a high density of positively charged defects giving rise to defect levels near the band edge maximum can compensate for the effect of a smaller population of negatively charged defects yielding midgap states. Hence, the overall charge contribution due to all defects does not *a priori* eliminate the possibility that the Ga_{As} antisite generates midgap levels.

The second point is based on the formation energies of the antisite defects. An analysis based upon formation energies relies on an equilibrium process for the formation of defects. The non-equilibrium processes occurring during the growth of a material such as In_{0.53}Ga_{0.47}As cannot be reduced to a set of processes described by equilibrium thermodynamics. For example, to provide reasonable growth conditions an As-rich supply of carrier gases must be provided to the growth chamber—the ratio of group V to group III precursors can exceed a factor of a hundred. However, this does not reflect the ratio of group III and V atoms available for growth at the surface of a substrate in the chamber. Hence, although the growth conditions are described as "As-rich," small variations in the growth conditions can lead to





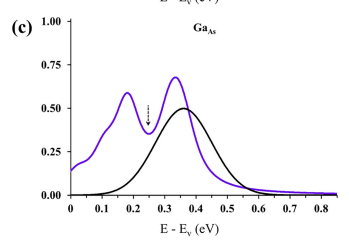


FIG. 10. Normalized distributions centered on defect levels showing the effect of thermal and alloy broadening on the (a) $Ga_{As}(+/0)$ and (b) $Ga_{As}(0/-)$ CTLs. The solid black curve in (a), (b), and (c) gives the experimentally extracted midgap defect peak from Ref. 8. The individual dashed peaks are for the CTLs calculated with a different distribution of cation nearest neighbors as identified in the legend within each figure. The solid purple curves in (a) and (b) are the weighted sum for the CTLs due to the alloy composition. The overall effect of this defect is depicted in (c), in which the solid purple curve is the sum of the purple curves from (a) and (b). The valley between the two main peaks is denoted by a dashed arrow, located at approximately 250 meV above the VBM.

either "As-rich" or "In/Ga-rich" conditions at the growth front, hence the possibility of forming either or both As_{Ga} and Ga_{As} antisites during the growth of $In_{0.57}Ga_{0.43}As$.

Therefore, our analysis leads us to the following conjecture. Some electrical characterizations see a peaked feature

in the midgap D_{it} , 8,15,16 whereas others do not. 45 Experiments suggest that the midgap states are largely independent of the oxide layer deposited on the In_{0.53}Ga_{0.47}As surface, 6-9 and thus, the defects are attributed to the semiconductor layer. Hence, our analysis suggests that a possible explanation is that there are many defect states providing positive charge with CTLs near the VBM maximum, and these states are observed in most CV measurements. However, if growth conditions are favorable to the formation of antisites, additional features at midgap are observed. If the charge associated with the midgap states increases the overall positive charge contribution of all states in the bandgap, then the defects should be identified with As_{Ga} antisites. If the net effect of the midgap states is to reduce the overall positive charge contribution of the states in the bandgap (but the integrated charge remains positive), then the midgap states can be associated with GaAs or related defects and the broad distribution of these midgap states can be associated with thermal alloying and proximity broadening.

V. CONCLUSIONS

Qualitative agreement for energies of CTLs for a range of defects native to In_{0.53}Ga_{0.47}As is found compared to previous computational studies based on hybrid DFT methods. Shallow defect levels agree well between the DFT + GWapproach and hybrid DFT. Deeper levels calculated by the DFT + GW approach are predicted to be 100 to 200 meV lower in energy compared to hybrid DFT results for defects occurring near and below the mid gap. 10 Good agreement between the methods is also found for the As₂ surface defect with both approaches predicting a transition level resonant with the conduction band. 11 Our findings are consistent, within the accepted theoretical and experimental accuracy, with the conclusions of Komsa and Pasquarello¹⁰ that the As_{Ga} antisite in a bulk-like local chemical environment is a candidate for giving rise to the electrical states below the midgap as observed at In_{0.53}Ga_{0.47}As/high-k oxide interfaces. However, a detailed investigation of the Ga_{As} antisite reveals that if this defect site is incorporated during kinetic growth conditions, then the CTLs associated with this defect and the effects of alloy broadening and proximity broadening suggest that this defect level is also compatible with the observation of the midgap defect states, and consistent with the broadening associated with this feature. The fact that the midgap states are only observed in some experimental CV measurements may be an indication that these defects are only present in a significant concentration under some growth conditions.

The stability of the neutral defect centers in various chemical environments is also evaluated. As is expected due to the loss of bonding energy upon the formation of a vacancy, a higher formation energy for the V_{Ga} compared to other defects in the bulk and oxide-terminated surface models is found, suggesting a lower concentration for this defect, and hence, a relatively small contribution to the D_{it} observed in III–V/high-k oxide MOS devices. The lower formation energy of the As_{Ga} antisite compared to other defects is observed; the Ga_{As} formation energy is approximately one

electron volt higher relative to the As_{Ga} antisite. During Asrich conditions or In/Ga-rich conditions at the surface during growth, it is not clear how to relate equilibrium formation energies to the population of the antisite densities. Hence, although the formation energies give some indication of relative stabilities, the actual relative population of the different defects will depend heavily upon growth conditions.

Comparing the CTLs for bulk-like defects and for defects formed near the oxide passivated semiconductor surface, summarized in Fig. 7, reveals a large range of energies for the various defect levels. The defect levels have been determined for bulk-like environments and these studies have been complemented by a quantitative analysis of the shift in the CTL energies as a function of their distance from the oxide. The calculations show that CTLs return to approximately 100 meV of the corresponding bulk values when situated even a few monolayers away from the semiconductor/ oxide interfacial layer. This is consistent with previous studies in which the defect levels of antisites located at various distances from a GaAs surface were found to return to bulklike values within a few monolayers from the surface. 46 In our simulations, the GaAs antisite exhibits CTLs which resonate with the In_{0.53}Ga_{0.47}As conduction band when directly bonded to the oxide at the semiconductor surface; however, the defect energetics become significantly more like their bulk counterparts even if situated two monolayers away from the semiconductor surface, being on the order of 0.1 eV of the corresponding bulk values. In addition, variable bonding arrangements due to local alloy disorder may account for the D_{it} feature extending into the valence band, 8,47 while defects bonding directly to the oxide have transition levels deep within the conduction band, potentially explaining the increase in D_{it} near and above the In_{0.53}Ga_{0.47}As conduction band edge observed in a number of experimental works. 8,15,47 This is also consistent with recent theoretical results. 11,48,49 III-V oxide interfaces have not received the amount of attention, particularly at the atomistic level, of the silicon-silicon oxide interfaces²⁶ or silicon-high k interfaces;³⁰ however, their increasing technological relevance is generating increased interest in these complex systems. Although there remain clear questions as to the specific nature of the atomic structures giving rise to the interface density of states with the bandgap of In_{0.53}Ga_{0.47}As/oxide interfaces, it appears that there are a variety of contributions and these contributions can potentially vary due to growth conditions. However, antisites seem to be the most likely candidates for generating midgap states and for generating a strong contribution to the high density of defect levels seen at the valence band edge.

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