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The *E*1–*E*2 center in gallium arsenide is the divacancy

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Abstract

Based on defect energy levels computed from first-principles calculations, it is shown the E1-E2 center in irradiated GaAs cannot be due to an isolated arsenic vacancy. The only simple intrinsic defect with levels compatible with E1 and E2 is the divacancy. The arsenic monovacancy is reassigned to the E3 center in irradiated GaAs. These new assignments are shown to reconcile a number of seemingly contradictory experimental observations.

Keywords: gallium arsenide, defects, density functional theory (Some figures may appear in colour only in the online journal)

1. Introduction

Performance of semiconductor electronic devices is dictated ultimately by the presence and behavior of point defects. A detailed knowledge of the properties and chemical evolution of those defects is crucial to the engineering of robust devices that operate effectively in normal operation conditions and under irradiation. In silicon, a vast, detailed understanding of defect behavior [1] has been developed, an result of extensive study over decades, and this has been of inestimable value in designing robust modern electronics. After silicon, gallium arsenide is perhaps the most studied semiconductor, with uses in a variety of devices. Despite early optimism [2] and notwithstanding intense effort over decades, definitive knowledge concerning even simple intrinsic defects in GaAs remains scanty. With electron paramagnetic resonance studies impaired in GaAs as a sensitive probe for the chemical structure of a defect [2,3], theory proved invaluable in identifying the arsenic antisite As_{Ga}—an arsenic replacing a gallium atom in the lattice—as the defect responsible for the EL2 [4, 5], the dominant center in as-grown GaAs.

Electron irradiation is an effective approach to study primary defects in materials. The initial products of electron irradiation are simple intrinsic defects: vacancies and interstitials resulting from primary lattice displacements. The

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EL2, only observed in as-grown GaAs, has been identified as As_{Ga}. The identities of defect centers observed in irradiated GaAs have proven more inscrutable. In n-type GaAs, a series of levels, designated E1 through E5, have been observed in e-irradiated GaAs above mid-gap [2]. The E1 level that appears at 32–45 meV below the conduction band (CB) and the E2 level at 0.13–0.18 eV [3, 6–8] are demonstrated to be two distinct transitions of the same defect [3], and are usually attributed to v_{As} [2]. The E3, E4, and E5 are transitions due to different defects, the E3 being the best characterized of these and is most commonly attributed to a v_{As} – As_i pair.

Identification of defects beyond the EL2 has been hampered by the lack of an experimental tool to unambiguously determine and characterize the chemical structure of a defect associated with an energy level. Conversely, first-principles calculations, particularly well suited for determining defect structure, have been unable to predict defect levels with quantitative accuracy sufficient to identify defects. In this work, I apply a robust method for computing accurate defect energy levels [9] to simple intrinsic defects in GaAs and determine that long-standing assignments of defect energy levels in GaAs need to be reassessed. The radiationinduced E1-E2 center, formerly associated with the arsenic monovacancy v_{As} , cannot be v_{As} and, instead, must be due to the divacancy vv. The v_{As} , no longer conscripted to account for the E1-E2 center, proves to give an apt account of the properties of the E3 center, once a site-shifting bistability is incorporated into a comprehensive description of v_{As} . These new assignments correlate with observed

behavior and reconcile seemingly contradictory experimental observations.

2. Methods

All density functional theory (DFT) calculations are performed using the Gaussian-basis pseudopotential code SeqQuest [10]. Exercising a new method for the accurate evaluation of energy levels of defects within a supercell approximation [9], we obtained all levels for all simple intrinsic defects of GaAs: both monovacancies and the divacancy, both antisites and the double antisite, and both single interstitials [11]. For silicon defects, this method predicted defect energy levels with an average accuracy of 0.1 eV and maximum error of \sim 0.2 eV. In GaAs, the computed span of defect levels, $\gtrsim 1.5 \,\mathrm{eV}$ [12], almost exactly matches the experimental band gap of 1.52 eV [13]. Furthermore, As_{Ga} calculations quantitatively match EL2 properties, e.g. the computed (0/1+) level is mid gap, 0.73 eV above the valence band, the (1 + /2 +) 0.24 eV below that [11]. This accuracy is comparable to the silicon defect study and validates using this computational model for accurate defect simulations in GaAs.

The analysis here (except where noted) focuses on the results of local density approximation (LDA) [14] simulations using Ga pseudopotentials with 3d electrons in the core, in 216-site cubic supercells with a 2³ k-sampling grid offset from the Γ -point. Computed energy levels for all intrinsic defects were also extrapolated from 512-site, and also selected 1000site (both these with 2^3 k-sampling) supercell calculations, to demonstrate that the computed levels were converged to the asymptotic dilute limit. The results extrapolated from 512-site cells matched to within $\sim 0.05 \, \text{eV}$ of the same levels extrapolated from 216-site supercells, demonstrating robust control of finite size effects, k-point sampling, and boundary conditions. For the defects discussed here, the conclusions are insensitive to use of a generalized gradient approximation such as PBE [15] rather than the LDA, or including the Ga 3d electrons as valence electrons.

3. Results and analysis

The presence of the divacancy in GaAs has historically been dismissed, due to early calculations predicting a high vvformation energy that precluded any significant population in equilibrium conditions [16], and positron annihilation calculations that yielded predictions of lifetimes in vvincompatible with any experimental observation in irradiated GaAs [17]. In the current calculations, vv takes charge states from (2+) to (4-). The formation energy of vv(0), 4.19 eV, only slightly exceeds $v_{As}(0)$, 3.55 eV [11]. previously unanticipated (4–) and (3–) charge states of the divacancy prove pivotal in explaining the radiation-induced E1-E2 centers. The v_{As} takes charge states from (3+) to (3-). A unified treatment of a site-shift bistability in the description of the arsenic vacancy proves crucial in accounting for its observed behavior. Structural changes for the negative charge states and all computed levels for these defects is illustrated in figure 1.

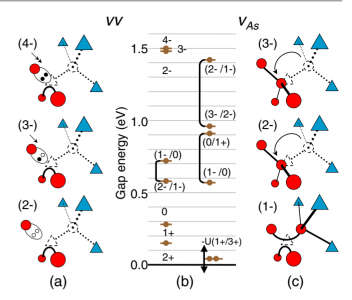


Figure 1. Schematic illustration of crucial features of the vv and $v_{\rm As}$ model for the E1-E3 centers in irradiated GaAs. (a) Bonding in the highly negative vv charge states that result in the levels near the CB. The (blue) triangles depict arsenic atoms and (red) circles depict gallium atoms. The open shapes depict vacated sites. (b) Defect level diagram for vv and $v_{\rm As}$. The solid lines connect the levels of -U systems. (c) Bonding in the $v_{\rm As}$ charge states responsible for levels near the CB. The global site-shift bistability occurs between $v_{\rm As}(1-)$ and $v_{\rm As}(2-)$, as a Ga adjacent to the simple As vacancy hops into the vacant site.

The vv can be described usefully as a $v_{\rm Ga}-v_{\rm As}$ pair. The vv(2-) has three full As lone pairs on the $v_{\rm Ga}$ side. The two electrons on the $v_{\rm As}$ side form a stretched Ga–Ga bond pair, illustrated in figure 1(a). The remaining trivalent Ga retreats from the vacant site. Adding an electron occupies the dangling bond on this Ga, causing it to pucker inwards by $0.6\,\text{Å}$ in the vv(3-). The vv accommodates yet another electron in the vv(4-), puckering this Ga another $0.2\,\text{Å}$ inwards. The resulting vv(4-/3-) and vv(3-/2-) levels are closer to the CB than any level from any other simple intrinsic defect. The next lower vv level is too distant from the conduction band (CB) to be of interest here; the vv(2-/0) is a negative-U transition near mid-gap, the inversion of charge transition levels (the -U) triggered by the dissociation of the Ga–Ga bond pair across the $v_{\rm As}$ side of the divacancy.

The arsenic vacancy proves globally bistable, as illustrated in figure 1(c), between a simple $v_{\rm As}'$ and a nearest-neighbor $v_{\rm Ga}$ –Ga_{As} pair obtained by an adjacent Ga atom shifting from its lattice site to the vacant site, a structure designated $v_{\rm As}^*$. The four valence electrons of $v_{\rm As}'(1-)$ couple into two Ga–Ga bond pairs in the ground state of $v_{\rm As}(1-)$, with D_{2d} symmetry. The ground state of $v_{\rm As}(2-)$ and $v_{\rm As}(3-)$ is the site-shifted $v_{\rm As}^*$. The large lattice relaxation (the site shift) reverses the standard order of the transitions, forming a -U system: $v_{\rm As}(3-/2-)$ is more strongly bound than $v_{\rm As}(2-/1-)$.

The calculated $v_{\rm As}$ and vv levels are presented in figure 1(b). The $v_{\rm As}$ does not have two distinct levels in the upper part of the gap, only a single -U transition. This -U(3-/1-) transition is much deeper in the gap than the E1 and E2 levels. The E1 and E2 were originally both

identified as acceptor levels [6], but negative charge states proved incompatible with the (1-), (0), and (1+) charge states required for a $v_{\rm As}$ model to be consistent with the results of positron experiments [18]. My computed $v_{\rm As}(1-/1+)$ charge transition is a single -U energy level near mid-gap. It is farther from the CB (and E1 and E2) than the maximum error seen in any LDA (or PBE) defect level calculation. The $v_{\rm As}$ cannot be responsible for E1 and E2.

The vv(4-/3-) and vv(3-/2-) transitions are the only viable candidates for E1 and E2. These transitions are near the CB, above all the computed $v_{\rm As}$ levels. No other simple intrinsic defect has computed levels within 0.5 eV of the CB edge. These vv levels are ideally located to correspond to E1 and E2, especially once spin polarization is included. Spin lowers the vv(3-) energy by 0.03 eV, increasing the splitting between vv(4-/3-) and vv(3-/2-) from 0.02 eV to 0.08 eV, good agreement with the observed 0.10 eV splitting between E1 and E2 observed in experiment, either $E_c-.032$ and $E_c-.129$ eV [8] or $E_c-.045$ and $E_c-.140$ eV [3]. To confirm the assignment of E1 and E2 to vv, a few other experimental observations must be explained.

One early experiment proposed the divacancy was responsible for the positron lifetimes associated with the E1 and E2 [19]. However, this assignment was rejected in later studies, replaced with a $v_{\rm As}$ model, partially based on analyses of positron studies. The positron annihilation lifetime calculated for vv by Puska and Corbel [17] was much longer than that observed for E1 and E2. This result was used to reject the vv model [2, 7]. However, those lifetime calculations used a vv structure where the atoms relaxed outwards, and did not consider (4–) and (3–) charge states. The atoms in vv(4-) and vv(3-) relax strongly inwards, and a smaller vacancy volume generally leads to a shorter positron lifetime.

The observed position trapping coefficient decreases after emitting one electron (E1), and disappears after emitting a second electron (E2) [18]. This observation motivated the reanalysis of the charge states in the $v_{\rm As}$ model: the neutral $v_{\rm As}(0)$ would bind a positron less well than the negative $v_{\rm As}(1-)$, positive charge in $v_{\rm As}(1+)$ would repel a positron, rendering the center invisible to positron annihilation spectroscopy. This progression of positron trapping behavior with charge state can be explained just as well in a vv assignment to this defect center, with transitions from vv(4-) to vv(3-) to vv(2-).

In the (4–), (3–), and (2–) charge states, the three As lone pairs on the $v_{\rm Ga}$ side of the vv are populated by six electrons. The electron charge changes occur on the $v_{\rm As}$ side, evidenced by the sequential structural rearrangements that occur there. In bulk GaAs, each As contributes $\frac{5}{4}$ electrons to each bond, Ga $\frac{3}{4}$ electrons. In a nominally neutral divacancy, therefore, each Ga dangling bond has $\frac{3}{4}$ electrons. In vv(4-), the $v_{\rm As}$ side distributes four valence electrons among three Ga dangling bonds in a Ga–Ga bond pair and two electrons in a pyramidal Ga lone pair, resulting in a nominal charge of $-\frac{7}{4}$. Going from vv(4-) to vv(3-) removes an electron from the Ga lone pair, causing that Ga atom to retreat slightly from the vacancy, reducing the charge on the $v_{\rm As}$ side to $-\frac{3}{4}$. The next electron, to form the vv(2-), also comes out

of this Ga lone pair, the now-trivalent Ga retreats strongly from the vacancy, toward a planar sp^2 -configuration, making the $v_{\rm As}$ side a slightly positive $+\frac{1}{4}$. This net positive charge makes it repulsive for positron capture. Furthermore, the $v_{\rm Ga}$ side of vv is highly negative, nominally $-\frac{9}{4}$, making the $v_{\rm Ga}$ side much more attractive for a positron than the nowpositive v_{As} side. Positrons trapped on the v_{Ga} side of vvare invisible to spectroscopy: the isolated $v_{Ga}(3-)$ has never been observed, suggesting that positron trapping at full As lone pairs leads to short positron lifetimes indistinguishable from annihilation in bulk GaAs. Hence, in positron annihilation experiments, vv mimics a bare v_{As} , with an offset in the total charge state. The vv model for E1 and E2 reconciles the original experiments that identify these as acceptors [6, 7] and positron experiments suggesting the lowest state was positively charged [18]. Positron lifetime calculations should be repeated with these new vv charge states and the correct inward-relaxed

No longer conscripted to explain E1 and E2, the v_{As} is now free to be assigned elsewhere. I propose that the E3 center in irradiated GaAs [3], asserted to be the same as (or very similar to) the EL6 center seen in semi-insulating as-grown GaAs, [20], is the v_{As} . The E3 center is known to be a primary defect of displacement damage—it must be an intrinsic defect—and is an important recombination center in GaAs [21]. The E3/EL6 energy level is reported to be 0.27– 0.38 eV below the conduction band edge [3, 20-25]. The only viable candidate in the computed survey of intrinsic defect levels is the $-U v_{As}(3-/1-)$ transition, at roughly -0.3 eVin LDA ($-0.35\,\text{eV}$ in PBE), where this defect will emit two electrons midway between the (3-/2-) and (2-/1-) energy levels. With the divacancy assigned to the E1 and E2, no other intrinsic defects have candidate levels this high in the band gap; all other computed intrinsic defect levels are more than 0.2 eV lower, and hence farther than the observed maximum error of 0.2 eV. The LDA (and PBE) calculations with these new assignments agree with the experimental observations for E1-E2 and E3 to within 0.1 eV.

An early LDA study using 216-site supercells [26] reports $v_{\rm As}$ charge states up to (3–), predicting that the (2 – /1–) and (3-/2-) defect levels are almost equal, nearly -U, high in the gap, which could be possibly consistent with an interpretation of the v_{As} highest two levels being associated with the E1-E2 levels. A more recent hybrid screened exchange (HSE) study using 64-site defect cells [27] reports their computed defect levels for the $v_{As}(3-/2-)$ and (2-/1-) transitions near the conduction band, and assert these correspond with the observed E1-E2 transitions, apparently confirming the classic association of the E1-E2 levels with the arsenic vacancy. The current results can reconcile these computational observations as well. The essence of the analysis is illustrated in the computed level diagrams of figure 2, In these, the computed level diagrams of v_{As} near the CB are depicted, for (a) v_{As} restricted to a paired- D_{2d} distortion (as in the HSE study [27]), then (b) switching to the ground state v'_{As} structures into the resonant- $D_{2d}(3-)$ ground state, and then (c) the full v_{As} energy level diagram that includes the v_{As}^* site shift.

Earlier studies [11,26] suggest that 216-site cells are necessary to get reliable results for defects, particularly

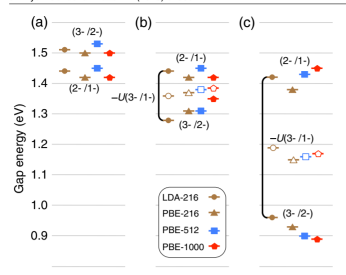


Figure 2. Computed level diagrams for the v_{As} in the upper half of the GaAs band gap: (a) restricted to paired- D_{2d} in simple vacancy v_{As}' ; (b) traversing the lower energy resonant- D_{2d} in $v_{As}'(3-)$; (c) the full thermodynamic energy levels for v_{As} , that traverses the site-shifted v_{As}^* ground state in the (2-) and (3-). The computed LDA energy levels extrapolated from 216-site supercells are given in (brown) circles. The PBE results in 216-site cells are marked with (brown) triangles, 512-site cells with (blue) squares, and 1000-site cells with (red) pentagons. Restricted to the paired- D_{2d} structure, the v_{As}' gives two levels nears the CB edge. Traversing the ground state structures instead causes inverted -U transitions, which would exhibit a single thermodynamic energy level in experiments, a double-electron emission midway between the computed charge transitions (open symbols).

for subtle relaxations around vacancies. The recent HSE results [27] reproduce very well our results with semilocal functionals, if one restricts the v_{As} vacancy to just the paired- D_{2d} structure. With either LDA or PBE, the $v'_{As}(3-/2-)$ level for the paired- D_{2d} is right at the CB edge, with the (2 - /1 -) slightly below, as seen in figure 2(a). However, earlier LDA results [11, 26], using larger 216-site cells with converged $2 \times 2 \times 2$ k-point sampling indicate that the resonant- D_{2d} structure is favored over the paired- D_{2d} for $v_{As}(3-)$, here, by 0.24(0.29) eV for LDA(PBE). The immediate consequence is that the $v_{As}(3-/1-)$ transition becomes -U, as depicted in figure 2(b), even before considering a site shift, already making it an unlikely match for the E1 and E2 levels. The results for a sequence of supercells of increasing size in figure 2 indicates that this result is converged to the bulk asymptotic limit with 216-site supercells.

Moreover, these calculations indicate that the site-shifted $v_{\rm As}^*(3-)$ and $v_{\rm As}^*(2-)$ are both favored over the simple arsenic vacancy $v_{\rm As}'$. For the (3-), the v^* is favored over the resonant- D_{2d} by $0.34(0.42)\,{\rm eV}$ for LDA(PBE); for the (2-), by 20–40 meV over the paired- D_{2d} (the resonant- $D_{2d}(2-)$ is higher yet by $\sim 0.15\,{\rm eV}$). This magnifies the -U effect for the (3-/1-) transition, and lowers the associated double-emission thermodynamic energy level to 0.30– $0.35\,{\rm eV}$ below the CB edge, as depicted in figure 2(c). It is impossible that this thermodynamic level can correspond to the E1 and E2 near the CB edge, and it is in extraordinarily good agreement with the observed E3 level. That both the LDA and PBE give nearly identical results lends further confidence to this prediction.

This unified defect level model creating this -U, incorporating this site-shift, is dependent upon this siteshift being thermally accessible within an energy level measurement; otherwise they would be observed as two separate defects. My computations indicate the site-shifting barriers are very small. As the site-shifted $v_{As}^*(3-)$ loses electrons, it becomes less favored with respect to v'_{As} . The $v_{As}^*(1-)$ starting with the $v_{As}^*(2-)$ atomic configuration collapses to the ground state $v'_{As}(1-)$ without a barrier, in both LDA and PBE, in all supercells, from 216-site through 1000-site. Conversely, postulating that a electron filling pulse starting with the $v'_{As}(1-)$ could conceivably become trapped in the simple vacancy in the (2-) or even the (3-) charge state (not finding a path to the more stable v^*), the barrier from the v' to v^* for these charge states is also relevant. Saddle point barrier calculations reveal that the simple v' has only a very small radius of stability against shifting into a v^* —a very small distortion of a neighbor toward the vacant site places the configuration on a downhill path to the lower energy siteshift—regardless of whether it is in the (2-) or (3-) charge state, with both LDA and PBE. The computed barrier for a site shift from $v'_{As}(2-)$ to $v^*_{As}(2-)$ is ~ 0.1 eV, the barrier from $v'_{\rm As}(3-)$ to $v^*_{\rm As}(3-)$ is less than 0.07 eV. The $v'_{\rm As}$ and $v^*_{\rm As}$ must both be included in a unified treatment of v_{As} defect levels.

Further experimental information also supports the notion of a larger -U transition. The EL6 emission exhibits a large Franck–Condon factor, $0.6\,\mathrm{eV}$, indicating this transition is associated with a large lattice relaxation [22]. It was later shown that the EL6 was but one of several emission peaks from a single defect: the intensities could be shifted between the EL5-EL6-EL7 family of levels by varying the filling pulse length in DLTS (deep-level transient spectroscopy), indicating a single DX-like center was responsible for these levels [25]. The predicted -U $v_{\mathrm{As}}(3-/1-)$ has a DX-like lattice relaxation, this transition involves a Ga atom hopping between two sites, from v'_{As} into v^*_{As} , as illustrated in figure 1(c). Therefore, v_{As} is compatible with known observations of the E3 and EL6 centers.

Identification of E3 with $v_{\rm As}$ vindicates treating the ${\rm Ga_{As}}-v_{\rm Ga}$ pair as a bistable form of the simple As vacancy, including it in computing levels for $v_{\rm As}$. The calculations demonstrated that $v_{\rm Ga}$ also exhibits a site-shift bistability [11], as originally proposed by Baraff and Schlüter [28]. These unified descriptions might be crucial to understand fermi-level pinning in GaAs: the site-shifting, charge-switching vacancies are the basis of an amphoteric native defect model for understanding many phenomena in GaAs [29, 30]. Though aspects of this amphoteric model are contested [31], our calculations confirm both monovacancies do exhibit the necessary features.

As shown in figure 1(b), both vv and v_{As} are computed to have donor levels near the valence band edge. Levels measured in p-type GaAs could be consistent with these. It has been hypothesized that the H0-H3 traps observed in irradiated GaAs were other transitions of the same defects responsible for E1-E3 [2, 3], and the results here suggest this hypothesis is likely correct. However, absent other discriminating evidence, definitive assignment of those observed hole traps to vv and v_{As} on the basis of defect level matching alone would be more

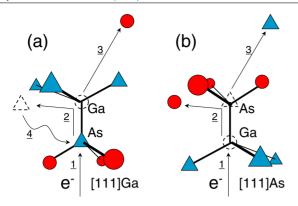


Figure 3. Schematic illustration of shadowing anisotropy leading to vv selectivity for a [1 1 1]As direction (b) as opposed to (a) [1 1 1]Ga-direction incident electron beam, when incorporating differential diffusivities of interstitials. For low energy displacements, the incident electron, $\underline{\mathbf{1}}$, causes the front atom to recoil, $\underline{\mathbf{2}}$, imparting most of its momentum to the knocked-on shadowed atom, $\underline{\mathbf{3}}$, ejecting it farther from the vacancy. If the front atom is Ga (b), the nearby Ga $_i$ is trapped, leaving behind vv, appearing to be a 'As sublattice' defect. If the front atom is As (a), the nearby As $_i$ is more likely to be recaptured, $\underline{\mathbf{4}}$, by the vv, forming v_{Ga} .

speculative, as other intrinsic defects are computed to also have candidate levels in the lower half of the band gap [11] and could, in principle, be responsible for those hole traps.

In electron radiation experiments, the E1-E3 and E3 traps were determined to be from defects with atoms displaced from the arsenic sublattice [2], based upon (1) an analysis of the expected anisotropy of defect production rates, a shadowing effect, from [111]-directed electron beams [32], and (2) these traps being annealed by pair-recombination in first-order kinetics, presumably through annihilation by mobile As_i . Identification on the As sublattice was a central motivation for originally assigning v_{As} to E1 and E2 [2], and appears inconsistent with a vv model for this center. Incorporating a differential diffusivity for ejected Ga and As interstitials into this simple billiard-ball analysis results in an anisotropy selective for vv on what appears to be the As sublattice.

The Ga_i diffuses with thermal barriers of $\sim 1 \text{ eV}$ [11, 33, 34]. Experiment inferred a 0.5 eV migration energy for As_i, confirmed in these DFT calculations [11]. In addition, the calculations indicate As_i will diffuse athermally, sequential capture of electron and holes driving the arsenic interstitial through the lattice [11]. Under e-irradiation, with copious carriers, athermal migration will be activated. A Ga_i ejected in a radiation-induced displacement is more likely to remain displaced. An As, displaced nearby has a greater probability of being recaptured by the (di)vacancy. Hence, a shadowing anisotropy results through the more or less distant ejection of As_i , as illustrated in figure 3. In one direction, the more distant As_i often escapes, resulting in more vv (and some v_{As}). In the reverse direction, the nearer As_i is often recaptured, resulting in more v_{Ga} (and only some vv). The key corollary is to note that, to this date, v_{Ga} has been invisible to electronic probes, so that it would not be detected if it were present, hence capture of a As_i by vv would cause it to vanish. The v_{Ga} has no levels above mid-gap, does not appear among the electron traps in n-type GaAs.

The threshold energy for displacement leading to generation of E1-E2 has been determined to be $\sim 10 \, \mathrm{eV}$ [35]. By analogy to irradiated silicon, this was taken to be a single displacement, and lack of a second threshold at twice this energy suggesting the double-displacement, the divacancy, was absent. The results here, compelling evidence supporting the identification of the vv with E1-E2, suggest that the assumptions that went into generalizing this threshold analysis from elemental silicon to a binary GaAs need to be reexamined, that the simple analogy to irradiated silicon is not quantitative.

4. Conclusions

In conclusion, the E1-E2 center in GaAs is determined to be due to the divacancy, on the basis of quantitative calculations of defect energy levels, and a detailed analysis of other experimental data. The standard model, v_{As} , cannot be responsible for E1 and E2 in GaAs. The v_{As} , instead, gives an apt description of the E3 (EL6) center. Previously unanticipated negative charge states for vv and a unified description of a site-shift bistable v_{As} are key insights needed to identify these traps. These assignments reconcile what had been seemingly conflicting experimental observations. Density functional theory supercell methods, using conventional semi-local functionals, are now accurate enough to make firm identifications of defects through defect level positions, adding v_{As} and vv to As_{Ga} (EL2) as intrinsic defects that are unambiguously identified in GaAs.

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