Spin Polarization and Magnetic Properties of $V_{\text{Ga}}O_N$ and $V_{\text{Ga}}O_N\text{In}_{\text{Ga}}$ in GaN: GGA+$U$ Approach

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Abstract

Electronic structure of a defect center containing the gallium vacancy and substitutional oxygen atom at nitrogen site ($V_{\text{Ga}}O_N$) in zinc blende and wurtzite GaN was analyzed within GGA+$U$ approach. The +$U$ term was applied to $d$(Ga), $p$(N), $p$(O), and $d$(In). Neutral $V_{\text{Ga}}O_N$ is in the stable high spin state with spin $S = 1$. The defect structure is strongly dependent on geometry of the defect and the charge state. Two spin structures, which arise due to two different configurations in $V_{\text{Ga}}O_N$, with ON either along the $c$-axis or in one of three equivalent tetrahedral positions in wurtzite structure were analyzed. The weak ferromagnetic coupling between centers was found. The strength of magnetic coupling is increased when there is a complex containing $V_{\text{Ga}}O_N$ with additional substitutional indium atom at the second neighbor to vacancy gallium site ($V_{\text{Ga}}O_N\text{In}_{\text{Ga}}$). Magnetic coupling between $V_{\text{Ga}}O_N\text{In}_{\text{Ga}}$ is ferromagnetic due to strong spin polarization of $p$ electrons of the nearest and distant nitrogen atoms.

Keywords GaN · In doping · Complex $V_{\text{Ga}}O_N$ · Magnetic coupling · LDA+$U$ calculations

III-nitride materials such as GaN and InGaN have found their applications in advanced solid-state lighting technologies [1–3] and optoelectronic devices including diodes or solar cells [4–6]. Moreover, the ferromagnetism (FM) in GaN or InN without doping by transition metal atoms was recently observed [7–11]. This FM was ascribed to the formation of native defects, such as cation vacancies or their complexes. For example in Ref. [9], analysis of the characteristics of hysteresis curves in irradiated GaN showed that the coercive field increases in line with the increase of concentration of gallium vacancy ($V_{\text{Ga}}$). The stable FM was discovered in $n$-type GaN, and it can be due to the presence of non-intentional donors, such as oxygen.

Measurements using positron annihilation spectroscopy of $V_{\text{Ga}}$ and its complexes containing the gallium vacancy and substitutional oxygen atom at nitrogen site $V_{\text{Ga}}O_N$ have found them to be dominating defects in as-grown $n$-GaN [12–21]. Both defects were intensively studied in experiments for their optical properties [12, 13, 16, 17, 19, 21] indicating defects as possible sources of the green (GL) and yellow (YL) luminescence in GaN. In Ref. [12], Son et al. detect spin-polarized $V_{\text{Ga}}O_N$ by electron paramagnetic resonance and suggest that YL and GL bands can be explained by $0^−$ and $\sim 2 V_{\text{Ga}}O_N$–optical transition levels. Moreover, in Ref. [13], two electronic structures were observed, which arise due to two different configurations of $V_{\text{Ga}}O_N$ in wurtzite (w) GaN, one with ON either along the $c$-axis (axial configuration, referred below as $a$) and other one in one of three equivalent tetrahedral positions (basal configuration, referred below as $b$).

However, there is not unequivocal agreement on the structure of $V_{\text{Ga}}$ and its complexes—both experimental [12, 13, 16, 17, 19, 21, 22] and theoretical [14–18, 20–32] results are divergent and sometimes contradictory. For example, the prediction of multiband luminescence due to the difference in geometries of $V_{\text{Ga}}O_N$ was demonstrated in Ref. [32]. Nevertheless, the magnetic properties for both $a$- and $b$-configuration of $V_{\text{Ga}}O_N$ were not discussed there. Moreover, various aspects of magnetism in GaN are not yet fully elucidated. There are few works which analyze magnetic coupling of $V_{\text{Ga}}$–$V_{\text{Ga}}$ [24, 25, 27, 31] or $V_{\text{Ga}}O_N$–$V_{\text{Ga}}O_N$ [26].

In fact, the calculated defect structure strongly depends upon the used exchange-correlation functional [14–18, 20–34]. $V_{\text{Ga}}O_N$ in III-N nitrides were theoretically investigated by local density approximation (LDA)/generalized gradient approximation (GGA) [14, 15, 20] or the hybrid functional (HyF) calculations [16–18, 21, 26, 32, 33]. Finally, GGA+$U$...
approach with the $+U$ term imposed on $p(N)$ in an examination of $V_{Ga}$ was also tried [30, 31]. Both the GGA+$U$ [30, 31] and the HyF [16–18, 21, 22, 26–28, 32–34] methods enhance spin polarization $V_{Ga}$ or its complexes and push defect levels deeper into the band gap in comparison with LDA/GGA [14, 15, 20, 29]. According GGA+$U$ calculations, $V_{Ga}$ demonstrates the $U$-induced splitting of partly occupied multiplets, and “negative-$U_{eff}$” behavior. The results for $U(N) = 5$ eV [30, 31] reproduce well the HyF calculations [17, 18, 23, 27, 28, 32, 33]. Both approaches give rise to stronger localization of the wave functions. This, in turn, alters the character of magnetic coupling between defects [31]. In fact LDA/GGA calculations are erroneous as they lead to overly long-ranged magnetic interactions [24, 25, 27, 31].

The increasing localization of the defect wave function has the opposite effect on the stability of the local magnetic moment and on the collective magnetization: in the former case, the increased localization stabilizes the high-spin (HS) state, while the coupling through the overlap of the wave functions of the neighboring defects is decreased [27, 30, 31, 35]. For both the $V_{Ga}$ and the $V_{Ga}O_N$ complexes, strong localization can lead to stable local spin moments [26, 27, 35] but it does not guarantee automatically a stable interaction between them [31, 35]. Partial delocalization of defect-induced bands may reduce the stability of HS state defect but also be responsible for the long-range magnetic interactions. This stabilization can be due to $p$-$d$ exchange interaction of impurity like Mn [36] or the spin polarization of $p$ electrons in low In-content InGaN.

Both In concentration and microscopic In distribution strongly influence the electronic structure and physical properties of InGaN [37]. The localization of the valence band maximum (VBM) states and the domination of the light emission of InGaN with low In content were observed [37]. Similar to GaN, $V_{Ga}$-complexes were also suggested to be the important non-radiative defects in InGaN quantum well [38, 39]. Strong effects in electronic structure of $V_{Ga}O_N$-hydrogen complex were found in $p$-type InGaN for high In content [33].

To check the hypothesis that magnetic coupling between $V_{Ga}O_N$-complexes can be more stable in InGaN, in the present paper, we study $V_{Ga}O_N$-structure in zb (zinc blende)- and w-GaN and InGaN using GGA+$U$ approach. After presenting the details of calculations in Sect. 1, the justification of the chosen approach is underlined. Next, we present results of calculations of the formation energies of defects (Sect. 1.3). Sections 2.1–2.3 contain the results of calculations of electronic structure of $V_{Ga}O_N$ in GaN and transition levels (Sect. 2.2) and electronic structure of InGaN (Sect. 2.3). The magnetic coupling between defects is analyzed in Sect. 2.4. Next, the structural relaxation is discussed in Sect. 3. Finally, Sect. 4 summarises the results.

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### 1 Method of Calculations

#### 1.1 Details of Calculations

Calculations based on the density-functional theory were performed using the ultrasoft pseudopotentials [40], the Perdew-Burke-Ernzerhof GGA exchange-correlation potential [41], including the $+U$ term implemented in the QUANTUM-ESPRESSO code [42] along the theoretical framework developed in Ref. [43]. We employed ultrasoft atomic pseudopotentials and chose $3d$, $4s$, and $4p$ orbitals for Ga; $4d$, $5s$, and $5p$ for In; $2s$ and $2p$ for N; and O as valence orbitals. The plane wave basis with the kinetic energy cutoff $(E_{cut})$ of 40 Ry provided a convergent description of the analyzed properties. The Brillouin zone summations were performed using the Monkhorst-Pack scheme with a $2 \times 2 \times 2$ k-point mesh [44]. Methfessel-Paxton smearing method with the smearing width of 0.136 eV was employed for obtaining partial occupancies. The zb 216- and 512-atoms and w 128-, 192-atoms supercells were considered, and ionic positions were optimized until the forces acting on ions were smaller than 0.02 eV/Å. The spin-orbit coupling was neglected. Formation energies were calculated according to Ref. [45] for N-rich conditions [30]. The $+U$ corrections were imposed on $d$(Ga) and $p(N)$ [30, 31]; $d$(Ga), $p$(N), and $p(0)$; and $d$(Ga), $p$(N), $p$(O), and $d$(In) for pure GaN, GaN:$V_{Ga}O_N$, and InGaN:$V_{Ga}O_N$, respectively.

#### 1.2 Justification for the Chosen Method

##### 1.2.1 Impact $+U$ Correct on Band and Crystal Structure of GaN

The band gaps $E_{gap}$ calculated within LDA/GGA are shrunk and amount only to about 1.4 and 1.6 eV for the zb- and w-crystals, respectively (see Fig. 1 and Refs. [30, 31]). Moreover, both LDA and GGA give too high calculated...
energy positions of \( d(Ga)\)-derived bands [46]. Inclusion of large \( U(Ga) = 10 \) eV term solves the problem with the correct position of \( d(Ga)\) [46–48], but \( E_{\text{gap}}\) is still about 2.5 eV (see Fig. 1). Our previous works showed that increasing \( U(N)\) from 0 to 5 eV opens the band gap of w-GaN from 1.6 to 3.0 eV [30, 31], but with \( d(Ga)\) centered about 13 eV below the VBM, similar to LDA/GGA results [30, 46]. Here, we systematically analyze how the \( U(Ga)\) and \( U(N)\) terms affect the electronic structure in both the zb- and w-crystals. On-site +\( U\) parameter, varying from 0 to 10 eV, was applied separately on \( d(Ga)\) and \( p(N)\), and then together on both \( d(Ga)\) and \( p(N)\) orbitals, where \( U(N)\) = 5 eV. The energy \( E_{\text{gap}}\) calculated as a function of \( U\) is shown in Fig. 1.

We found that \( U(Ga) = 3.0 \) eV along with \( U(N) = 5 \) eV reproduces the experimental \( E_{\text{gap}}\) of 3.2 and 3.4 eV for both zb- and w-GaN [49] (Fig. 1, (3, 3')) and the binding energy of Ga 3\( d\) level centered about 15.5 eV below the VBM—in agreement with Ref. [50]. These values are also in agreement with HyF results [51]. Such an underestimation of the band gap and sensitivity of \( E_{\text{AFM}}\) result with Ref. [50]. These values are very close to the experimental data of \( \mu_{\text{bulk}}\) (GaN) centered about 13 eV below the VBM, and (\( \mu_{\text{bulk}}\)) for zb- and w-structure, respectively.

1.2.2 Impact of \( +U\) Correction on Electronic Structure of GaN with Defect

The \( +U\) corrections were imposed on \( d(Ga)\), \( p(N)\), \( p(O)\), and \( d(In)\). \( U(O) = U(N) = 5 \) eV, and \( U(In) = U(Ga) = 3 \) eV for GaN. However, including \( U(O)\) and \( U(In)\) terms have a small effect on \( V_{\text{GaOn}}\) structure due to small contributions of \( p(O)\) and \( d(In)\) orbitals to the defect states (see Sects 2.1 and 2.2 for the detailed discussion of electronic structure of both GaN: \( V_{\text{GaOn}}\) and GaN: \( V_{\text{GaOn}}\)In\( _{\text{Ga}}\)).

1.2.3 GGA+\( U\) vs HSE Results

It was noted above that HyF [16–18, 21, 22, 26–28, 32, 33] calculated \( V_{\text{GaOn}}\)-structure is in agreement with GGA+\( U\) results for \( U(N) = 5 \) eV [30, 31]. Here, in order to verify the agreement further, we perform calculations for Heyd, Scuseria, and Ernzerhof functional, based on the PBE functional where parameter \( \alpha\) is a fraction of the exchange that is replaced by Hatree-Fock exchange [54]. Calculations were done for isolated \( V_{\text{GaOn}}\) and \( V_{\text{GaOn}}\)In\( _{\text{Ga}}\) (3NNs axial configuration in notation of Sect. 2.4) for w-GaN. Screening parameter \( \alpha = 0.25\) was set to reproduce band gap of ~ 3.0 eV. Nevertheless, the results indicate that energies of spin polarization (\( \Delta E_{\text{AFM}}\) – FM) (defined in Sect. 2.1) agree to within 0.05 eV or less, and energies of magnetization (\( \Delta E_{\text{AFM}}\) – FM) (defined in Sect. 2.2) agree to within 0.005 or less. That shows the good agreement for calculations of magnetic properties within these two approaches.

We note that the problem of choosing the \( \alpha\) parameter for getting accurate defect levels is still an open issue [55, 56], as well choosing the \( U\) parameter in GGA+\( U\) approach [30].

1.3 Formation Energy of Defects

Formation energy of charged \( V_{\text{GaOn}}\) was calculated. One geometry of \( a-V_{\text{GaOn}}\) was considered for cubic GaN and two \( a\)- and \( b-V_{\text{GaOn}}\) configurations were analyzed for w-GaN. Because the aim was to understand the influence of \( In\) doping on spin-polarized properties, the number of configurations of complex \( V_{\text{GaOn}}\) was chosen: \( \text{In}_{\text{Ga}}\) as second nearest neighbor to \( \text{Ga}_{\text{Ga}}\) when forming \( \text{O}_{\text{O}}\)In\( _{\text{Ga}}\) (referred below as \( \alpha\) or \( \alpha\)-\( \text{In}_{\text{Ga}}\)) (referred later as \( n\) ) chains where these \( \text{O}_{\text{O}}\) and \( \text{N}_{\text{N}}\) are the nearest possible positions to \( \text{Ga}_{\text{Ga}}\) neighbors. Hence, the two geometries as referred as \( a-o\)- and \( a-n-V_{\text{GaOn}}\)In\( _{\text{Ga}}\) were considered for zb-crystal. And four geometry configurations referred as \( a-o\), \( a-n\), \( b-o\), and \( b-n-V_{\text{GaOn}}\)In\( _{\text{Ga}}\) were analyzed for w-GaN. In this work, the formation enthalpy of \( \text{In}_{\text{Ga}}\) was calculated and the binding energy, \( E_{\text{bind}}\) defined as a difference in the total energies of compounds that contain \( V_{\text{GaOn}}\)In\( _{\text{Ga}}\) or \( V_{\text{GaOn}}\)In\( _{\text{Ga}}\) were taken into consideration.

Formation energy \( E_{\text{form}}\) of a defect was calculated by formula taken from Ref. [45]

\[
E_{\text{form}} = E_{\text{tot}}(\text{GaN} : D) - E_{\text{tot}}(\text{GaN}) + (-)\Sigma n_i\mu_i + +q(\epsilon_F + E_{\text{VBM}}) + E_{\text{correct}} \tag{1}
\]

The first two terms on the right-hand side are the total energies of the supercell with and without the complex, respectively. \( n_i\) is number with the \(+(-)\) sign corresponding to the removal (addition) of atoms. \( E_{\text{VBM}}\) is the energy of the VBM of bulk GaN, and \( \epsilon_F\) is the Fermi energy referenced to this \( E_{\text{VBM}}\). The energy \( E_{\text{VBM}}\) is determined from the total energy difference between the pure crystal with and without a hole at the VBM in the dilute limit by algorithm from Ref. [45]. \( \mu_i\) are the variable chemical potentials of atoms in the solid, which in general are different from the chemical potentials \( \mu_i(\text{bulk})\) of the ground state of elements (Ga \( \text{bulk}\), In \( \text{bulk}\), and \( \text{N}_2, \text{O}_2\)). Chemical potentials of the components in the standard phase are given by total energies per atom of the elemental solids: \( \mu(\text{Ga \( \text{bulk}\)) = E_{\text{tot}}(\text{Ga \( \text{bulk}\))}, \mu(\text{In \( \text{bulk}\)) = E_{\text{tot}}(\text{In \( \text{bulk}\)), while } \mu(\text{N \( \text{bulk}\)) = E_{\text{tot}}(\text{N}_2)/2\) and \( \mu(\text{O \( \text{bulk}\)) = E_{\text{tot}}(\text{O}_2)/2\) (\( \Delta H_F(\text{NO})\) was neglected. In N-rich condition,
\[ \mu(Ga) = E_{\text{tot}} \text{(Ga bulk)} + \Delta H_f \text{(GaN)} \] and \[ \mu(In) = E_{\text{tot}} \text{(In bulk)} + \Delta H_f \text{(InN)} \] are taken, where \( \Delta H_f \) is the enthalpy of formation per formula unit, and it is negative for stable compounds. \( \Delta H_f \) at \( T = 0 \) K is obtained by considering the reaction to form or decompose a crystalline GaN and InN from its components and dependent on an cohesive energy, \( E_{\text{coh}} \), of Ga, In, N, and O. The obtained results for \( E_{\text{coh}} \) of Ga, N, and O were shown in Refs. [30, 57]. Calculated \( E_{\text{coh}} \) (In) and \( \Delta H_f \) (zb-GaN), \( \Delta H_f \) (InN) are 2.56 (2.5 [58]) and \(-1.24 \text{ (} -1.27\text{),} -0.36 \text{ (} -0.32\text{) eV [59]}\), (experimental values presented in brackets).

The last term, \( E_{\text{corr}} \), includes two corrections. The first one, \( \Delta E_{\text{PA}} \), is the potential alignment correction of the VBM. The VBM in the ideal supercell and in the supercell with a (charged) defect differs by the electrostatic potential and is obtained by comparing the potential at two reference points far from the defect in the respective supercells with \( (P[0]) \) and without \( (P[0]) \) the defect, \( \Delta E_{\text{PA}} = q[P[D^\dagger] - P[0]]. \) Second correction is an image charge correction as expressed by 2-order Makov-Payne form: \[ E_{\text{MP}} = \frac{\alpha_{\text{die}}}{} \text{, where } \alpha_{\text{die}} \text{ is the lattice-dependent Madelung constant, which for hexagonal structure is 3.5, W is the supercell volume, and } \varepsilon \text{ is the static dielectric constant.} \] \( E_{\text{MP}} \) was calculated to be 0.2, 0.4 eV for charged defects \( (q = 1, -1) \). Results of calculations are presented in Sect. 2.2.

## 2 Results

This section summarizes the obtained results for the defect structure and formation energy and discusses magnetic interaction between defects.

### 2.1 Electronic Structure and Spin Polarization of \( V_{GaO}_N \)

The defect states of \( V_{GaO}_N \) stem from the result of the interaction between the vacancy orbitals and the O-impurity states. Local atomic configuration of this defect has the \( C_{3v} \) point symmetry. In both the zb- and the w-structures, the vacancy is tetrahedrally surrounded by three N and one O atoms, and the respective defect states are localized on the resultant broken \( sp^3 \) bonds (Fig. 2) that split into a singlet \( a_g \) and a higher in energy quasiletur “\( t_2 \)”. Energies of \( a_g \) and “\( t_2 \)”, calculated below as relative to the VBM, depend on the crystal structure, geometry of defect and the charge state. \( a_g \) is a resonance state with the valence bands. “\( t_2 \)”- \( V_{GaO}_N \) is located in the band gap and splits into a doublet \( e_g \) and a singlet \( a_1 \) with a splitting energy of about 0.2, 0.5, and 0.7 eV for the zb-GaN, \( a \)- and \( b \)-geometries of \( V_{GaO}_N \) in w-crystals, respectively (see Fig. 3, the green lines). The energy splitting contains the contributions of a weak perturbation by the w-symmetry in w-GaN and the \( U \)-induced so-called quasi-Jahn-Teller (JT) effect [30].

In the case of non-vanishing spin polarization, the exchange coupling splits \( e_g \) into spin-up \( e_g \) and spin-down \( e_g \) states by the splitting exchange energy defined as \( \Delta \varepsilon_{\text{ex}} = \varepsilon(e_{\uparrow}) - \varepsilon(e_{\downarrow}) \), where \( \varepsilon \) is the energy of the defect level, and \( a_1 \) into \( a_{1\uparrow} \) and \( a_{1\downarrow} \) (Fig. 3, the blue lines). The \( \Delta \varepsilon_{\text{ex}} \), in general, depends on the symmetry of defect, the charge state and \( U \). The \( e_{\downarrow} \) and \( a_{1\uparrow} \) of neutral \( V_{GaO}_N \) in zb-GaN are localized in the band gap at 2.6 and 1.9 eV above the VBM, respectively. The \( e_{\downarrow} \) and \( a_{1\downarrow} \) are resonances with the VBM (Fig. 3a). According to this point of view \( V_{GaO}_N \) is a deep acceptor containing two holes.

As presented above in Section 2.3, in w-crystal, the defect can exists in two different geometries tagged as \( a \)- and \( b \)- \( V_{GaO}_N \). According to GGA+U calculations, single-electron level representations of electronic structures are different for \( a \)- and \( b \)- \( V_{GaO}_N \) (Fig. 3 b, c, left panels). Structure of \( a \)- \( V_{GaO}_N \) in w-GaN is similar to the one of zb-GaN, \( e_{\downarrow} \) and \( a_{1\downarrow} \) are 2.45 and 1.65 eV with the respect to the VBM, and \( e_{\downarrow} \) and \( a_{1\downarrow} \) are hybridized with the valence bands (Fig. 3b). Introducing O atom into basal plane of defect leads to strong symmetry perturbation, \( e_{\downarrow} \) is split by 0.5 eV into two \( a_{2\downarrow}(1) \) and \( a_{2\downarrow}(2) \), singlet states. Thus, “\( t_2 \)” in this case is a composite band containing \( a_{1\uparrow}, a_{2\downarrow}(1), \) and \( a_{2\downarrow}(2) \) levels located about 1.4, 2.3, and 2.8 eV above the VBM, respectively (Fig. 3c).

The energies of \( V_{GaO}_N \) levels strongly depend on the charge state \( q \). Single-electron energy levels of \( V_{GaO}_N \) for \( q = 0, -1, -2 \) with their respective charge states are shown in Fig. 3. The physics behind the calculated electronic structure of charged defects is determined by the following counteracting effects [30]: (i) the intracenter Coulomb repulsion is dominant in the non-spin-polarized calculations. Without spin polarization, the energy of \( e_{\downarrow} \), \( a_1 \) increases by \(-0.5 \text{ to } -0.6 \text{ eV with the } q \text{ changing from } 0 \text{ to } -2 \text{ (the levels are shown in green color in Fig. 3}); (ii) The effect of the value exchange splitting, for example, in zb-GaN with \( q \) changing from 0 to 2, \( \Delta \varepsilon_{\text{ex}} \) decreases from 2.6 to 0 eV; (iii) the \( U \)-induced potential which is attractive (repulsive) for occupied (unoccupied) orbitals [43]. This effect is clearly seen for \( e_{\downarrow} \) that decreases with \( q \) changing from 0 to 1 (Figs. 3a, b). These results are in agreement with HYF calculations from Ref. [17] where negative-\( U_{\text{eff}} \) behavior of \( V_{GaO}_N \) was observed.

“\( t_2 \)” defect state is occupied by 4, 5, and 6 electrons for \( q = 0, -1, -2 \), respectively. As shown in Figs. 2, 3 unpaired spins of the electrons localized on the dangling bond states lead to the local magnetic moment of \( V_{GaO}_N \). Unlike \( V_{GaO}_N \), which is a closed-shell system, both \( V_{GaO}_N \) and \( V_{GaO}_N \) are spin polarized. The energy gain \( \Delta \varepsilon_{\text{PM-FM}} \) due to spin polarization, defined as the difference in total energy of
the spin-nonpolarized (PM) and spin-polarized (FM) calculations, is shown in Table 1. Generally, \( \Delta E_{PM} - FM \) assumes the maximal value when \( "t_2" \) is occupied with 4 electrons with the spin \( S = 1 \), and it vanishes when \( "t_2" \) is fully occupied. Moreover, the energy of antiferromagnetic state (AFM) of single defect was considered also in the analysis [23].

Every considered geometry of the neutral \( V_{GaON} \) is the magnetic centrum in HS state with the local magnetic moment \( \mu_{tot} = 2 \mu_B \). For example, \( \Delta E_{PM} - FM(V_{GaON}) \) are 1.7, 1.62, and 1.86 eV for defects in zb-GaN, and \( a \)- and \( b \)-geometry in w-GaN, respectively (Table 1). \( V_{GaON}^{-1} \) is the paramagnetic centrum with \( \mu_{tot} = 1 \mu_B \). \( \Delta E_{PM} - FM(V_{GaON}^{-1}) \) are 0.38 and 0.3, 0.4 eV for defects in zb-GaN, and \( a \)- and \( b \)-geometry in w-GaN, respectively (Table 1). Finally, \( V_{GaON}^{2-} \) is the non-spin-polarized centrum (in this case \( \Delta E_{PM} - FM = 0 \)).

According to our calculations, \( b \)-configuration stabilizes the
HS state in neutral $V_{Ga}O_N$ by about 0.24 eV in comparison with $a$-geometry. This value is close to the energy gain of formation energies in $b$-$V_{Ga}O_N$ and $a$-$V_{Ga}O_N$ (see Sect. 2.2).

The geometry of the defect affects also the localization of the $V_{Ga}O_N$ states. The effect is displayed in the plots of the density of spin polarization, Fig. 2a,e–f. Figure 2 indicates that the $V_{Ga}O_N$ states are dominated in all cases by the localized and spin-polarized contribution of the three $sp^3$ orbitals of the N nearest neighbors because O-atom is more electronegative than N-atom and two electrons with the opposite spins are located on $sp^3$ oxygen orbital. In contrast to GGA method in which electrons that occupy for example, “$t_{2g}$” are spread over four $p$ orbitals of the nearest neighbors [31, 60], GGA+U calculations showed that the partial occupancy is avoided [31, 60]. Moreover, in the case of $b$-$V_{Ga}O_N$, one can observe the anisotropy for three $sp^3$ of N orbitals (Fig. 2d, f). The $U$-induced symmetry breaking of $e_g$ level, i.e., the quasi-Jahn–Teller effect, was observed in $b$-$V_{Ga}O_N$. In $a$-$V_{Ga}O_N$ contributions of the three N neighbors to the $V_{Ga}O_N$ states of the vacancy, wave function are almost equal, whereas for $b$-$V_{Ga}O_N$ the wave function is dominated by the two basal N ions located in the $(x, y)$ plane, and the contribution of the remaining N ion is strongly reduced, see Fig. 2d–f.

### 2.2 Formation Energy and Transition Levels

The calculated $E_{form}$ of $V_{Ga}O_N$ and assumed $\varepsilon_F$ at the VBM are given in Table 2. $E_{form}$ in all configurations is the same. The same trend was observed also in Ref. [32]. The obtained values 1.64, 3.54, and 5.73 eV in w-GaN are close to results of HSE (1.9, −, and 5.3 eV [17]) and HyF (B97-2- functional) (1.5, 4.0, and 6.2 eV) [32]. Our calculations demonstrate that $E_{form}$ is similar in all geometries.

The change in the defect charge state is determined by the transition level $\varepsilon(q_1/q_2)$, defined as the Fermi energy relative to the VBM at which formation energies of the $q_1$ and $q_2$ charge states are equal. We find $\varepsilon(0/−)$ are 1.84 and 1.9, 2.1 eV for defects in zb-GaN, and $a$-, $b$-geometry in w-GaN, respectively, and $\varepsilon(−/−)$ are 2.1 and 2.2, 2.4 eV for defects in zb-GaN, and $a$-, $b$-geometry in w-GaN, respectively, which is consistent with $V_{Ga}O_N$ energies shown in Fig. 3. Comparison of the calculated energies with the results for other exchange functionals is shown in Table 3. Table 3 contains dataset for only $a$-$V_{Ga}O_N$. Comparable values of $\varepsilon(0/−)$ and $\varepsilon(−/−)$ were obtained with GGA+U and different HyF exchange-correlation functionals. For example, our results are close to the obtained by HSE06 approach (with 20% exact exchange) [17, 18].

### 2.3 Spin Polarization of $V_{Ga}O_N In_Ga$

In Sect. 1.3 the geometrical configurations of $V_{Ga}O_N In_Ga$ were discussed. Although our calculations show that binding energy in such a complex is low, ~0.1–0.3 eV, the formation energy is 1.9 eV which is a little higher than in the case of $V_{Ga}O_N$ and considerably lower than in the case of $V_{Ga}$ [30, 34]. Because the formation energy for different geometries ($a$- or $b$-) is similar, both $V_{Ga}O_N$ and $V_{Ga}O_N In_Ga$ can exist in non-equivalent atomic configurations.

In this section, the results for calculated energy of spin polarization, local magnetic moment, spin density, and density of charge of $V_{Ga}O_N In_Ga$ are presented. Moreover, the effect of addition of In impurity on the spin-polarized properties is analyzed by comparing these results with similar results for $V_{Ga}O_N$. Next, in order to get a clearer picture of the influence of In on electronic structure, the complex $V_{Ga}O_N In_Ga$ was investigated in zb-GaN.

The defect states of $V_{Ga}O_N In_Ga$ stem from the result of the interaction between the vacancy orbitals and the O- and In-impurity states.

### Table 1

| Defect | $V_{Ga}O_N$ | $V_{Ga}O_N In_Ga$ |
|--------|-------------|-------------------|
| $q$    | $\Delta E^{PM-FM}$ | $\Delta E^{PM-FM}$ |
| 0      | 1.70$^a$    | 1.78$^{a-o}$     |
| 2      | 0.38$^a$    | 0.50$^{a-o}$     |
| 2      | 0.62$^a$    | 1.69$^{a-o}$     |
| 2      | 0.33$^a$    | 0.50$^{a-o}$     |

### Table 2

| $q$ | $a-V_{Ga}O_N$, zb-GaN | $a-V_{Ga}O_N$, w-GaN | $b-V_{Ga}O_N$, w-GaN |
|-----|----------------------|---------------------|---------------------|
| 0   | 1.35                 | 1.64$^a$            | 1.4$^b$            |
| 2   | 3.1                  | 3.54$^a$            | 3.5$^b$            |
| 5.2 | 5.73$^a$             | 5.9$^b$             | 5.9$^b$             |
According to GGA+U calculations, in zb-GaN, the electronic structure of \( V_{GaON} \) is similar to \( V_{GaON} \). There is only a difference in the defect energy levels of about 0.1–0.2 eV. With the increasing number of In content from 1 to 4, the level of \( e_2 \) increases no more than 0.5 eV. It demonstrates the stability of defect symmetry in zb-structure.

Strong effect on the electron structure was observed for wurtzite crystal (see Fig. 3b, c, right panels). As with O-atom in basal plane, also in this case for both \( a \)- and \( b \)-\( V_{GaON} \), “\( t \)” is split into three singlet states \( a_{11}, a_{21}(t), \) and \( a_{21}(c) \) due to strong tetragonal perturbation generated by In atom in the crystal structure. \( a_{21}(c) \) of \( b-n-V_{GaON} \) is higher in energy, and it is a resonance state with the conduction bands (Fig. 3c).

Calculated \( \Delta E_{PM-FM} \) values for different configurations of \( V_{GaON} \) are given in Table 1. According to our calculations, the inclusion of \( In_{Ga} \) stabilizes HS state of the defect by 0.1–0.16 eV in comparison to \( V_{GaON} \) (see Table 1). For example, in zb-GaN, the differences in the spin polarization energies of \( V_{GaON} \) and \( V_{GaON} \) are 0.08 and 0.14 eV, for \( q = 0 \) and \( -1 \), respectively. The same energy gain takes place in w-structure, for example, the differences between \( \Delta E_{PM-FM} \) (\( b-n-V_{GaON} \) and \( b-V_{GaON} \)) are 0.16 and 0.08 eV, in cases \( q = 0 \) and \( -1 \), respectively. Generally, the increase of \( \Delta E_{PM-FM} \) is in agreement with the changes of electron structure. Shifting up the single-electron levels (\( a_{21}(c) \) in particular) to the CBM leads to the increase in the energy of exchange splitting, and therefore in the energy of \( \Delta E_{PM-FM} \). The above results obtained for \( V_{GaON} \) are typical: the stability of the HS state predicted for \( V_{GaON} \) is 10–12% higher than that predicted for \( V_{GaON} \).

The increase in discrepancy of \( \Delta E_{PM-FM} \) was observed with the rising of In content. \( \Delta E_{PM-FM} \) is 1.7, 1.78, 1.88, 1.9, and 1.8 for \( n = 0, 1, 2, 3, \) and 4. For \( n = 4 \), \( \Delta E_{PM-FM} \) is smaller than for \( n = 2 \) and \( n = 3 \).

The spin (Fig. 2b, c, h-j) and electron (Fig. 2k, l) contour densities of \( V_{GaON} \) are shown in Fig. 2. Like in the case of \( V_{GaON} \), \( V_{GaON} \) states are localized mainly on the \( p \) orbitals of the three N atoms (Figs. 2b, c, h-j). Spin density of \( V_{GaON} \) is more delocalized, since they comprise long-range tails that involve \( p \) orbitals of distant N ions or even \( p \) orbital of oxygen (refer to \( V_{GaON} \) In \( Ga \), see Fig. 2c). These orbitals constitute the VBM (Fig. 4) of GaN. \( V_{GaON} \)-spin-up states are resonances degenerate with the valence bands, i.e., both \( a_{11} \) and \( e_{21} \) hybridize with the upper part of the VBM, which forces also a partial delocalization of their wave functions. It is evident from Fig. 2 that enhancing correlation effects by addition of In atoms leads to the decrease of the localization of the spin density on the broken bonds and to the increase of its axial anisotropy.

The contour (Fig. 2k, l) is plotted in the (100) plane and shows that a large contribution to the electron density comes from the nitrogen atoms and strong ionic bonds resulting from \( sp^3 \) hybridization. The spherical symmetry around anions is observed, which indicates that the bonds in \( GaN \) are dominated by the ionic component.

By calculating the contributions of individual atoms projected onto relevant atomic orbitals to the total DOS (Fig. 4a), one finds that the main contribution comes from the defects states \( p(N) \) of the N nearest neighbors of \( V \) (Fig. 4b)—in agreement with Fig. 2. Both \( p(N) \) and \( p(O) \) orbitals also build the VBM of GaN (see Fig. 4b–d). The contribution of the \( d(In) \) orbitals to the spin density is non-negligible due to the substantial contribution of \( d(In) \) to the VBM and defect states (see Figs. 2 and 4f).

### 2.4 Magnetic Interaction of \( V_{GaON} \) and \( V_{GaON} \)

In the present section, we study \( V_{GaON} \) and \( V_{GaON} \) defect pairs and analyze the impact of crystal distortion on their properties by comparing the results of magnetic interaction calculations. The magnetic coupling between vacancy complexes is discussed as the possible origin of the experimentally observed FM in GaN. Electronic structure of a defect pair is determined by three factors: (i) the distance between vacancy complex, (ii) the relative orientation of complexes with respect to each other and to the crystal axes, and (iii) the charge state. \( V_{GaON} \) and \( V_{GaON} \) configurations were considered, in which the defects are the third nearest neighbors (3NNs) and fourth nearest neighbors (4NNs) with respective spatial separation of about ~5.2 and ~6.5 Å (it is a distance between \( V \) in the relaxed structure). In w-GaN, the defects can be located either in the same \( (x,y) \) basal plane perpendicular to the \( c \)-axis, which is referred to as the \( xy \)-case, or they can be oriented along the \( c \)-axis, which is denoted here as the \( c \)-case. Finally, we mention that the defect pair in such configurations has eight nearest neighbor atoms (two O and six N atoms). Because the goal of work was to analyze the FM in GaN, we do not consider...
here 1NNs configurations with the seven nearest neighbor atoms.

When the complex is spin polarized, the orientation of the two spins can be ferromagnetic (FM), ferrimagnetic (FiM, when spins of the two complexes are different because the defects are in different charge states), and antiferromagnetic (AFM). In the latter case, the total spin vanishes but the spin polarization energy is finite. Energy of magnetization $\Delta E_{\text{AFM(FiM)}-\text{FM}}$ is defined as a difference between the total energies of AFM (FiM) and FM states and is positive when FM coupling is stable. In summary, two and four configurations were considered for $^{\text{V GaO}_N-^\text{V GaO}_N}$ and $^{\text{V GaO}_N\text{InGa}-^\text{V GaO}_N\text{InGa}}$ in zb-GaN, respectively, and four and eight geometries were considered for $^{\text{V GaO}_N-^\text{V GaO}_N}$ and $^{\text{V GaO}_N\text{InGa}-^\text{V GaO}_N\text{InGa}}$ both in w-GaN, respectively. $\Delta E_{\text{AFM(FiM)}-\text{FM}}$ are summarized in Table 4. In Table 4, we only briefly highlight the results.

zb-GaN:

(i) $^{\text{V GaO}_N-^\text{V GaO}_N}$ are spin-polarized and the coupling between defect spins is FM in both charge states. Magnetic moment is 4 and 2 $\mu_B$, for $q=0$ and $-2$, respectively. The observed FM is weak with $\Delta E_{\text{AFM-FM}}$ energy of about 0.0045 and 0.001 eV, for $q=0$ and $-2$, respectively (for 4NNs pair). The obtained values are a little larger than those obtained by HSE06 approach in Ref. [26] due to shorter spatial separation between defects in our work.

(ii) The inclusion of In impurity modifies the coupling. The absolute values of the coupling strength are few times higher than they are for 4NNs $^{\text{V GaO}_N}$ and amount to 0.01–0.014 and 0.006–0.008 eV, for $q=0$ and $-2$, respectively.

w-GaN:

(i) The dependence on geometry configuration was observed. For example, in $x\bar{y}$- basal plane, the ground state of $(^{\text{V GaO}_N-^\text{V GaO}_N})$ is FM with $\Delta E_{\text{AFM-FM}}$ of about 0.01, 0.006 eV (for 3NNs and 4NNs) and 0.09, 0.005 eV (for 3NNs and 4NNs) in $a$- and $b$-$^{\text{V GaO}_N}$ geometry, respectively.

(ii) Unlike zb-crystal, in wurtzite, the dependence of the coupling on the orientation in crystal of the pairs is rather strong: the total energy differences between the $x\bar{y}$- and $c$-orientations are larger than the corresponding values of $\Delta E_{\text{AFM-FM}}$. When defects are oriented along the $c$-axis, the magnetic interaction is AFM.

(iii) Similar to the results for zb-crystal, magnetic coupling of $^{\text{V GaO}_N\text{InGa}-^\text{V GaO}_N\text{InGa}}$ is larger than of $^{\text{V GaO}_N-^\text{V GaO}_N}$. Generally, for all configurations, In doping facilitates the stability of FM phase in GaN by order of magnitude (Table 4).

The dominant mechanism of magnetic interaction between $^{\text{V GaO}_N}$ is determined by the interplay between the counteraction of the bonding-antibonding (BA) and the exchange of spin-up–spin-down ($\Delta e_{\text{ex}}$) splittings.
Figure 5 shows spin densities calculated for several configurations of $V_{\text{GaON}}$ and $V_{\text{GaONInGa}}$. They reflect the shape and localization of the partially occupied defect states of the complexes. Generally, these states are localized on the $p$ orbitals of the six N atoms. In most cases, spin density of a given defect in w-structure is like in Fig. 5 where two directions, orthogonal ($xy$) and parallel to the c-axis, are presented. In the FM cases, they are close to the sum of spin densities of $V_{\text{GaON}}$ shown in Fig. 2. Spin densities in $b$-$V_{\text{GaON}}$ and $V_{\text{GaONInGa}}$ are more delocalized, since they are

| $q$ | Configuration | $\Delta E_{\text{AFM}}$ | $\mu$ | Configuration | $\Delta E_{\text{AFM}}$ | $M$ |
|-----|----------------|-----------------|------|----------------|-----------------|-----|
|     |                |                 |      |                |                 |     |
| 0   | $a-a$ (3NNs)   | 0.014           | 4 (FM)| $a-o-a$ (3NNs) | 0.1             | 4 (FM) |
|     | $a-a$ (4NNs)   | 0.0045          | 4 (FM)| $a-o-a$ (4NNs) | 0.01            | 4 (FM) |
|     |                |                 |      |                |                 |     |
| −2  | $a-a$ (4NNs)   | 0.001           | 2 (FM)| $a-o-a$ (4NNs) | 0.006           | 2 (FM) |
|     |                |                 |      |                |                 |     |
|     | $xy-a-a$ (3NNs) | 0.01            | 4 (FM)| $xy-a-n-a$ (3NNs) | 0.05           | 4 (FM) |
|     | $xy-a-a$ (4NNs) | 0.006           | 4 (FM)| $xy-a-n-a$ (4NNs) | 0.008           | 4 (FM) |
|     | $c-a-a$ (3NNs) | −0.016          | 0 (AFM)| $c-a-n-a-a$ (3NNs) | −0.1           | 0 (AFM) |
|     | $c-a-a$ (4NNs) | −0.002          | 0 (AFM)| $c-a-n-a-a$ (4NNs) | −0.007         | 0 (AFM) |
|     | $xy-b-b$ (3NNs) | 0.09            | 4 (FM)| $xy-b-n-b-n$ (3NNs) | 0.15        | 4 (FM) |
|     | $xy-b-b$ (4NNs) | 0.003           | 4 (FM)| $xy-b-n-b-n$ (4NNs) | 0.012         | 4 (FM) |
|     | $c-b-b$ (3NNs) | 0.008           | 4 (FM)| $c-b-n-b-n$ (3NNs) | 0.015         | 4 (FM) |
|     | $c-b-b$ (4NNs) | 0.00015         | 4 (FM)| $c-b-n-b-n$ (4NNs) | 0.009         | 4 (FM) |

Table 4 Energy of magnetic coupling $\Delta E_{\text{AFM-FM}}$ together with total magnetic moment (in $\mu_B$) of the complex in the charge state $q$ (it is a sum $q_1$ and $q_2$, where) obtained for zb- and w-GaN calculations. For simplicity, both $\Delta E_{\text{AFM-FM}}$ and $\Delta E_{\text{FiM-FM}}$ are denoted by $\Delta E_{\text{AFM-FM}}$ and the actual spin configurations and values of magnetic moment are given in the columns “$\mu$”.
comprised of long-range tails which involve $p$ orbitals of distant N ions (Fig. 5). Moreover, N neighbors of the complex are not equivalent, as displayed by the dissimilar contributions to the spin density. Indeed, the contribution of the axial N (or O) ions is almost vanishing (Fig. 5 a, c, e, g). Non-negligible spin density of In orbital demonstrates the important contribution of In atom into magnetic interaction between defects.

### 2.5 Crystal Structure zb- and w-GaN and Relaxation

All above results are demonstrated for atomic relaxed structures. The increase of FM stability when In content grows can be explained by the distortion effects in crystal structure due to the atomic displacement in the process of relaxation. As it can be seen in Figs. 2 and 5, significant and complex perturbations in the crystal structure of GaN are experienced when vacancy is introduced and it can be attributed to the fact that the radius of In is larger than the one for Ga and the radius for N is larger than the one for O. The interatomic distances between the neighbors of a complex defect are mainly determined by the lattice constants of the host crystal (but here, for low In content, it was neglected), and also by the atomic relaxations, i.e., displacements of the neighbors. In the studied compounds, the outward relaxation of the nearest neighbors elongates bonds by about 5–12%, i.e., 0.2 Å, and reduces the total energy by the relaxation energy $\Delta E_{\text{relax}}$ of 0.63, 0.86, 0.93, 0.98, and 0.99 for $V_{\text{Ga}}\text{O}_N\text{InGa}$, $V_{\text{Ga}}\text{O}_N\text{InGa}$, $V_{\text{Ga}}\text{O}_N\text{2InGa}$, $V_{\text{Ga}}\text{O}_N\text{3InGa}$, and $V_{\text{Ga}}\text{O}_N\text{4InGa}$, respectively. It can be noted that the

![Diagram of crystal structure](image)

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**Fig. 6** The bond lengths of Ga-N and In-N for (a) $V_{\text{Ga}}\text{O}_N$, (b) $V_{\text{Ga}}\text{O}_N\text{InGa}$ in w-GaN. Initial bond lengths were equal to the length of Ga-N bond, i.e., 1.944 Å.
structural relaxation in GaN:V_{GaON}In_{Ga} is about 0.23–0.33 eV larger than in the case of GaN:V_{GaON}In_{Ga}. From the results in Fig. 6, it follows that distortion (change in bond length) for the V_{GaON}In_{Ga} is larger than that for the V_{GaON}.

But we note that structural distortions are more complex. Although, the displacements of the second and third neighbors are an order of magnitude smaller, the effect of atomic relaxations around defects, involving not only the nearest but also more distant neighbors, cannot be neglected. The states of the defect complexes are determined by the overlap of the N and O dangling bonds given by the N-N (N-O) distances. In ideal structure (after relaxation without defect), N-N is equal to 3.18 Å. In V_{GaON}, the N-N and N-O dangling bonds are 3.55 and 3.48 Å, respectively. But in V_{GaON}In_{Ga}, these values are shorter; three N-N are 3.58, 3.52, and 3.48 Å, respectively, and N-O is 3.42 Å. It implies that the defect states are more localized, and the energy of spin polarization of such defects is higher.

3 Summary and Conclusions

In summary, spin states of V_{GaON} and V_{GaON}In_{Ga} complexes in both zb- and w-GaN, and magnetic coupling between them, were studied within GGA+U calculations. The U(Ga) = 3 eV and U(N) = 5 eV terms were imposed on d(Ga) and p(N) leading to the correct band gap of GaN.

Charge states q from 0 to −2 of V_{GaON} were considered. In both crystal structures for neutral V_{GaON} with S = 1, high-spin configuration is stable. Wave functions of V_{GaON} have a multi-orbital character, being composed of three p(N) and one p(O) orbitals of vacancy neighbors. But the main contribution to the spin density comes from sp^3 N orbitals.

Two different electronic structures, which arise due to two different geometry configurations of V_{GaON}, with O_N either along the c-axis and in one of three equivalent tetrahedral positions in w-GaN, were analyzed. The latter geometry configuration assumes stronger stability of HS state and more delocalization of defect state.

Introducing In_{Ga} as second neighbor to V_{Ga} on the one hand imposes changes to the electronic structure, on the other gives rise to the delocalized wave function of the defect as the crystal structure is perturbed, and finally contributes to long-tail spin density distribution. Magnetic moments originate mainly from sp^3 N orbitals, and the contribution of p orbitals of distant N, d(Ga), and d(In) states is about 20%.

Various relative orientations of the defects and several charge states (q = 0 and q = −2) were considered, and consequences regarding the observed FM in GaN were pointed out. Using a relation predicted from mean field model, T_c = 2\delta(S + 1)J/3k_B, the room temperature of FM implies that (assuming \delta = 6 neighbors and S = 1) the coupling constant J(V_{GaON}In_{Ga}) = 0.01 and J(V_{GaON}) = 0.0045 eV, i.e., 920 K and 415 K, respectively. These values have a limited reliability as the distribution of defects is random and the coupling depends on the distance between defects.

Comparing the obtained results with experiments, we note that, according to the results, the observed collective ferromagnetism in GaN systems [7–11] can originate from magnetic interaction between V_{GaON} defects. And in Ga-rich In_{Ga}N alloys, we predict even stronger FM.

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