VARIATION OF MAGNETISM OF Cr_{1-X}Ga_XN DUE TO THE PRESSURE: A DFT STUDY

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ABSTRACT

We report a firt-principles study of the pressure dependence of electronic and the magnetic properties of $Cr_{1-x}Ga_xN$ compounds (x= 0,25, 0,50 and 0,75) in wurtzite-derived structures. We use the full-potential linearized augmented plane wave method (FP-LAPW) within of the density functional theory framework. We found that, the lattice constant vary linearly with Ga-concentration. The magnetic moment changes for a critical pressure. At x = 0,75, a rather abrupt onset of the magnetic moment from 0 to 2,2 μ_B at P_{cr} = 22,65 GPa is observed. For x = 0,25 and 0,50 Ga concentrations, the magnetic moment increases gradually when the pressure decreases toward the equilibrium value. We study the transition pressure dependence to a ferromagnetic phase near the onset of magnetic moment for each $Cr_{1-x}Ga_xN$ compounds. The calculation of the density of states with Ga concentration is carried out considering two spin polarizations. The results reveal that for x = 0,75 the compound behaves as a conductor for the spin-up polarization and that the density of states for spin-down polarization is zero at the Fermi level. At this concentration the compound presents a half metallic behavior; therefore this material could be potentially useful as spin injector. At high pressures P > P_{cr} the compounds exhibit a metallic behavior.

Keywords: FP-LAPW, Magnetic semiconductors, Pressure dependence.

VARIACION DEL MAGNETISMO EN EL COMPUESTO Cr_{1-X}Ga_XN DEBIDO A LA PRESIÓN: UN ESTUDIO POR DFT

RESUMEN

Reportamos un estudio de primeros principios para estudiar los efectos de la presión sobre las propiedades electrónicas y magnéticas del compuesto $Cr_{1-x}Ga_xN(0,25, 0,50 \text{ y } 0,75)$ en la estructura wurtzita. Usamos el método Ondas Plana Aumentadas y Linealizadas Potencial Completo (FP-LAPW) en el marco de la Teoría del Funcional de la Densidad DFT. Hallamos que la constante de red del compuesto aumenta linealmente con la concentración de Ga. El momento magnético cambia para una presión crítica. Para x = 0,75, un cambio bastante abrupto comienza en el momento magnético de 0 a 2,2 μ_B a la presión P_{cr} = 22,650 GPa. Para las concentraciones de Ga x = 0,25 y 0,50 el momento magnético aumenta gradualmente cuando la presión disminuye hacia su valor de equilibrio. Estudiamos la dependencia con la presión de transición hacia la fase ferromagnética cerca del inicio del momento magnético para cada compuesto $Cr_{1-x}Ga_xN$. Calculamos la densidad de estados con la concentración de Ga considerando las dos contribuciones de polarización de espín. Los resultados revelan que a x = 0,75 el compuesto se hace conductor para la polarización de espín arriba y que la densidad de estados de espín abajo es cero en el nivel de Fermi. A esta concentración en el compuesto presenta un comportamiento half-metallic, por consiguiente este material podría ser potencialmente usado como inyector de espín. A presiones *P* > *P_{cr}* el compuesto exhibe un comportamiento metálico.

Palabras Claves: FP-LAPW, Semiconductores magnéticos, dependencia con la presión.

1. INTRODUCTION

At present, various high performance devices fabricated from III-nitride have generated great interest. Amongst the III semiconductor, Gallium nitride (GaN), whose more stable phase is hexagonal (wurtzite) [1]. Due to its wide direct band gap, gallium nitride is a promising candidate in semiconductor technology and has a broad range of potential applications for optoelectronic and high power electronic devices. Intensive activities over the recent years have made of the short wave length blue, violet light emitting diodes (LEDs) a commercial reality [2]. Due to its high chemical stability and high thermal conductivity, is also suitable for the applications in the harsh environments, such as, in high-temperature/highpower electronic devices [3, 4], as metalsemiconductor field effect transistors (MESFETs), high electron mobility transistors (HEMTs) and heterojunction bipolar transistors (HBTs) [5, 6]. Large piezoelectric constants of GaN point out possible applications of GaN-based materials in piezoelectric sensors [1]. Additionally, theoretically [7, 8] and experimentally [9], high Curie temperatures and room-temperature ferromagnetism have been found in GaN-doped with transitionmetal (MT) elements, which in principle opens the door for potentials use of this room-temperature ferromagnetic material for spintronic devices [7 -10].

The magnetic properties of transitions metal (MT) in GaN regained prominence due to potential application for Dilute Magnetic Semiconductors (DMS) [11, 12]. In particular, the 3d-MT elements can be expected that substitute Ga-atoms during crystal growth. The knowledge of their associated deep defects is very important to development a new kind of devices such as: electro-optic switches, ultra sensitive magnetic field sensors and quantum-mechanism-based logic for high speed computation [13].

In this paper, we investigate mixed $C_{1-x}Ga_xN$ compounds. As will be shown, the dilution of CrN with Ga expands the lattice constant according to Vegard's law as expected. Also Ga incorporation produce the arise of magnetic moment formation. This makes these materials potentially useful as spin injector. In order to model the compounds, simple ordered model structures (Cr₃Ga₁N₄, Cr₁Ga₁N₂, Cr₁Ga₃N₄) based on supercells of the wurtzite

structure were investigated.

2. COMPUTATIONAL METHOD

The electronic structure calculations were performed employing the full-potential linearized augmented plane wave method (FPLAPW) as implemented in the WIEN2k package, which includes the LAPW[14], within of the spin density functional theory (DFT) framework [15, 16]. The exchange and correlations electronic energy. were calculated with Generalized Gradient Approximation (GGA) of Perdew et al. [17]. Separation energy between the valence and core states of -8,0 Ry, and the angular momenta up to lmax = 10, were used. The wave functions in the interstitial region were expanded in plane waves with a cutoff of Kmax = 8.0/RMT(where RMT is the smallest muffin-tin sphere radii inside the cell). For Ga_xCr_{1-x}N compounds muffintin radii of 1,90, 1,80 and 1,60 bohr for Ga, Cr and N atoms were selected respectively. In calculation of the electronic structure, a 144 k-points mesh were used in Brillouin irreducible zone generated according to the Monkhorst Pack scheme [18]. The iteration for self-consistence was continued until the convergence criterium of 1×10^{-4} Ry was reached.



Figure 1. Unit cell used in our calculation for $Cr_{1-x}Ga_xN$ compounds in wurtzite-like structure.

The $C_{1-x}Ga_xN$ compounds were modeled for x = 0,25, 0,50 and 0,75 compositions according to special quasirandom structures approach [19] and the disorder aspects were ignored. For $Cr_1Ga_1N_2$ (x = 0,50) an hexagonal unit cell with alternating [0001] layers of CrN and GaN in conventional wurtzite structure was employed. For $Cr_3Ga_1N_4$ (x = 0,25) and $Cr_1Ga_3N_4$ (x = 0,75) an hexagonal unit cell consisting of two wurtzite unit cells piled in the

c direction were used [20]. There are eight planes with one Ga (or Cr) or one N atom in a 1× 1configuration, as shown in Figure 1. In this structure, a Cr atom replaces to a Ga atom in the unit cell. This atomic substitution (Cr-Ga) has been experimentally observed by S. E. Park et al. [21] for Cr doped GaN has been successfully grown in the form of bulk single crystals in wurtzite phase using a sodium flux growth method [21] and also used metal organic chemical vapor deposition technique (MOCVD); additionally, Haider et al [22] grown Cr doped GaN by molecular beam epitaxy (MBE) on sapphire (0001) substrate. This has allowed to get, room temperature ferromagnetism of Cr-doped GaN single crystals has stirred further interest in the DMS systems [21].

The lattice parameters and cohesion energy were found by the fitting the total energy versus volume to the Murnaghan's state equation [23].

3. RESULTS AND DISCUSSION

The cohesion energy per unit cell as function of lattice constant for $Cr_{1-x}Ga_xN$ compounds are shown in Figure 2. As we showing figure, the minimum for each curve shifts to the right as it increases the concentration of Ga in the $Cr_{1-x}Ga_xN$ compound.



Figure 2. Cohesion energy as a function of the lattice constant for $Cr_{1-x}Ga_xN$ compound, the line is a Murnaghan equation of state fit.

From minimum point energy of Figure 2, the equilibrium lattice constant (square points) and the energy (circular points) per unit cell as a function of Ga concentration are presented in Figure 3. The increase of lattice constant with Ga concentration is due to that Ga-atom is bigger than Cr-atom. We found a linear dependence according to Vegard's law. Note that the energy also has a linear tendency with the Ga concentration as has been observed in other ternary nitrides [20, 24].



Figure 3. Equilibrium lattice constant and cohesion energy as function of Ga concentration in $Cr_{1-x}Ga_xN$.

Magnetic moment for $Cr_{1-x}Ga_xN$ compounds are shown in Figure 4. As we shown in this figure, for $Ga_3Cr_1N_4$ (square points) there is a rather abrupt onset of the magnetic moment from 0 to 2,2 μ_B at ~ 22,65 GPa (3.11 Å). The $Cr_1Ga_1N_2$ and $Cr_3Ga_1N_4$ compounds presented the same magnetic moment transition but with a slowly pressure dependence. From this figure, we observe the tendency towards magnetism increases as we further decrease the pressure. A similar behavior has been reported by González et al. $Ga_xV_{1-x}N$ compounds in wurtzita structures [25].

We found that in the studied cases, $Cr_{1-x}Ga_xN$ has a nonzero magnetic moment at the zero pressure. Comparing the total energy of the ferromagnetic (FM) and antiferromagnetic (AFM) configurations for x = 0,25 and 0,50 compounds at the equilibrium volume, the FM is found to be lower in energy and is the predicted to be preferred state. Such as has been found experimentally R. K. Singh et al [26] and Sang Eon Park [27] for diluted concentrations of Cr atoms in the GaN semiconductor.



Figure 4. Magnetic moment as a function of the lattice constant for $Cr_{1-x}Ga_xN$. The line is guide to the eye.

In Table 1 we show a summary of the structural and magnetic results for $Cr_{1-x}Ga_xN$ compounds. We give the equilibrium lattice constant (a₀), critical pressure at which the magnetic moment appears (P_{cr}), saturation value of the magnetic moment per unit cell (μ) and the spin polarization P_F at the equilibrium lattice constant defined by:

$$P_F = \frac{N_{\uparrow}(E_F) - N_{\downarrow}(E_F)}{N_{\uparrow}(E_F) + N_{\downarrow}(E_F)} \tag{1}$$

With $N_S(E_F)$ being the DOS of spin $S (= \uparrow \text{ or } \downarrow)$ for the Fermi energy E_F .

In Table 1, we note that the calculated magnetic moment for each configuration (3 μ_B /Cr-atom) is due to the Cr³⁺ electronic configuration, because a Cr-atom provides two net electrons to the Cr_{1-x}Ga_xN compound [28] and the Cr atoms couple ferromagnetically when doped into GaN whether the host is a cluster or a crystal [10].

Table 1. Structural and magnetic properties of $Cr_{1-x}Ga_xN$ compounds.

Compound	a_0 (Å)	P_{cr} (GPa)	$\mu (\mu_B)$	$P_F(\%)$
$Cr_1Ga_3N_4$	3,235	22,65	3,0	100
$Cr_1Ga_1N_2$	3,200	24,55	3,0	78,8
$Cr_3Ga_1N_4$	3,160	26,60	9,0	31,8

We study the pressure dependence on the electronic

density of states (DOS) for the spin-up and spindown polarization in $Cr_{1-x}Ga_xN$ compounds. In Figure 5, we show the partial (PDOS) of d states of chromium calculated per atom, and the total DOS estimated per unit cell of the $Cr_1Ga_3N_4$ compound at the equilibrium lattice constant (P = 0).

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Figure 5. Partial DOS of d-states of Chromium calculated per atom and total DOS estimated per unit cell of the $Cr_1 Ga_3N_4$ compound at equilibrium pressure P = 0.

Near the Fermi level, the dispersion in the chromium impurity bands show that there is considerable interaction between neighboring Cr atoms in the unit cell. This compound behaves as conductor for spin-up polarization and as semiconductor for spin-down polarization. We found 100% spin polarization at the Fermi level which indicates that Cr₁Ga₃N₄ compound presents a half metallic behavior at P= 0. The figure 6 shows the density of states calculated at the high pressure P = 32 GPa over to the transition pressure P > Pcr. The Fermi level passes through the impurity and conduction bands in both spin polarization. The compound exhibits a metallic behavior. We can observe that the spin polarization splitting is dependent on the pressure and may be manipulated experimentally. Similar results we found for the other concentrations Cr₁Ga₁N₂ and Cr₃Ga₁N₄. On the basis of our present study, we believe that Cr_{1-x}Ga_xN compounds may be candidates for being ferromagnetic or half-metallic materials in semiconductor-based spintronic applications.



Figure 6. Partial DOS of d-states of Chromium calculated per atom and total DOS estimated per unit cell of the $Cr_1Ga_3N_4$ compound at over the transition pressure P > Pcr (P = 32 GPa). Energy is relative to Fermi energy.

4. CONCLUSIONS

An ab-initio study of pressure effects on the electronic and magnetic properties of Cr_{1-x}Ga_xN compounds (x = 0.25, 0.50 and 0.75) in wurtzita derived structures were presented. We used the fullpotential linearized augmented plane wave LAPW within of the spin density functional theory framework. The lattice constant is found to vary linearly with Ga-concentration. The magnetic moment changes for a critical pressure. At x = 0.75, a rather abrupt onset of the magnetic moment from to 2,2 μ_B at ~ 22,65 GPa, is observed. For x = 0,25 and 0.50 Ga concentrations the magnetic moment increased gradually when pressure is decreased toward the equilibrium volume. Also, we study the pressure dependence of transition to a ferromagnetic phase near the onset of magnetic moment for each $Cr_{1-r}Ga_{x}N$ compounds. Calculation of the density of states with Ga concentration was carried out considering two spin polarizations. Results reveal that when x = 0.75 the compound behaves as conductor for spin-up polarization and as semiconductor for spin-down polarization is zero at the Fermi level. At this concentration compound has a possible half metallic behavior; therefore this material could be potentially useful as spin injector. At high pressures P > Pcr the compounds exhibit a metallic behavior.

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