

First Principles Study on Li-doped and Li,O-codoped AlN

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This paper focuses on the detailed investigation of the structural and electronic properties of wurtzite AlN crystals doped by Li with and without oxygen with the first principles calculation. All the calculations have exhibited significant structural distortions. Compared with the monoclinic doping, the oxygen codoping has improved the structure deformation and lowered the formation energy of Li dopants. The calculated electronic density of states (DOS) reveals that all doping configurations still preserve semiconductor characteristics. The states around the valence band maximum cross the Fermi level, which implies p-type doping. The induced extra levels are extremely localized and flat in Li-doped AlN while much more delocalized in oxygen codoped models. The mono-doping of Li is in general energetically unfavorable while the codoping improves the formation and makes the intercalation of Li more stable in AlN. According to the results, the codoping configuration of Li with O in AlN has provided a useful way of modifying the corresponding properties.

Keywords: semiconductor, AlN, codoping, first principles.

1. INTRODUCTION

As a wide bandgap semiconductor, AlN exhibits a direct bandgap of 6.02 eV (at room temperature), good dielectric strength and high thermal conductivity. These excellent properties make AlN a promising material for ultraviolet (UV) optoelectronic devices such as LEDs, acoustic wave devices, Schottky diodes and UV detectors. [1, 2] Doping is an efficient way to modify the electronic and optical performances of AlN for such applications. It has been proved experimentally that AlN doped with different impurities has shown unusual properties [3–8].

The alkali metals have attracted much attention as dopants due to the simplicity of their free-electron-like, delocalized valence state [9, 10]. For example, Li with low electronegativity and small ionic radius facilitates high reactivity and diverse chemical bonding in the solid state [11, 12]. Inserting Li into the Al site in host AlN crystals will induce holes leading to mixed-valent systems, which are beneficial for realization of p-type semiconductors. The properties of Li accommodated within the doped structure are very important and interesting. It is well known that oxygen being the most common impurity in AlN crystals is very hard to be removed in the growth process. Giving that the codoping method is a possible way for p-type doping of AlN [2, 13, 14], we think the theoretical study of codoping with Li and O atoms in AlN will be reasonable for future experiments.

First principles calculations are widely used for investigation of the ground-state properties of numerous materials. In this paper, first principles calculations of the structural and electronic properties in Li-doped and Li,O-codoped AlN supercells were performed by the generalized

gradient approximation (GGA) together with plane-wave basis sets.

2. METHOD OF CALCULATION

With the WIEN2K code [15, 16], we predict the structural and electronic properties of doped wurtzite AlN by first principle method based on density functional theory (DFT) in the GGA. The full-potential linearized augmented plane wave (FP-LAPW) method was used to describe the interaction between ion cores and valence electrons. In this method, the unit cell is divided into non-overlapping muffin-tin atomic spheres centered on the atoms and interstitial spaces. There are two kinds of electrons, which are core and valence electrons, respectively. The Al ($3s^2 3p^1$), O ($2s^2 2p^4$), N ($2s^2 2p^3$) states and outermost shell of Li are treated as valence electrons. Different basis sets are used for these two types of regions. Core electrons are described by atomic wave functions while the valence electrons are expanded into spherical harmonic functions in the atomic sphere and plane waves in the interstitial space. The electronic wave functions are obtained by solving Kohn-Sham equations self consistently.

The model of AlN unit cell including two Al atoms and two N atoms are building. According to the experimental results, the lattice parameters of a and c are chosen to 0.311 nm and 0.498 nm, respectively. The supercell ($2 \times 2 \times 2$) is a model consisting of 32 atoms. The Li dopants are placed on the Al sites while oxygen atoms replace N atoms. The muffin-tin radii (R_{MT}) for Al, N, Li and O atoms are assumed as large as possible in a way that the spheres do not overlap. The plane wave expansion of the electronic eigenfunctions in the interstitial region is cutoff with $R_{MT} \times K_{max} = 7.0$, where R_{MT} is the minimum radius of the muffin-tin spheres and K_{max} is the magnitude of the largest K vector in the plane wave. The

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RK_{max} parameter controls the size of the basis set in the FP-LAPW method. The largest vector in charge density Fourier expansion (G_{max}) is set to 12.0 bohr⁻¹. The wave functions inside the muffin-tin spheres, which are expanded into spherical harmonics, are taken up to $l_{max} = 10$. The Monkhorst-Pack k -mesh ($5 \times 5 \times 3$) is performed to sample the supercell Brillouin zone (BZ) in all our calculations [15], special k -points are obtained and the tetrahedron method [17] is used for integrations in the irreducible Brillouin zone (IBZ). The total energy was converged to less than 0.1 mRy (1.36 meV) by self-consistent calculations.

3. RESULTS AND DISCUSSION

The structural improvement was realized through a full relaxation of the atomic positions and internal parameters with volume optimizations of the total energy for all the calculated systems in order to reach the stable structure configuration. During the process of geometry optimization, all atoms were fully relaxed until residual forces on individual atoms become smaller than 1 mRy/bohr. The volumes were optimized with the minimum energy succeeding the atomic positions relaxations. Internal structural parameters were modified using the calculated interatomic forces to determine the equilibrium atomic positions of Al, N, Li and O. Due to the absence of experimental data concerning Li-doped AlN, the parameters of pristine AlN as initial conditions are used.

As seen in Table 1, Li-N moves outward after the doping. The expansion in basal plane becomes relatively smaller as the substitution takes place in the c -axis direction. The heterogeneously distributed bonds of tetrahedrons in pristine AlN remain anisotropy after the doping which is important for spontaneous polarizations. The bond of Li-N in c -axis direction shares a very approximate distance in Li and Li-O doped AlN. At the same time, the bond length along the direction perpendicular to the c -axis is less affected as the codoping has extended the bonding.

In order to analyze the relative stability, we study the formation energy of a separated Li and a Li-O complex in AlN supercells. The formation energy with alkali metal impurity (X) of charge state zero in AlN is defined as [2, 13]:

$$E^f(X) = E_{tot}[AlN, X] - E_{tot}[AlN, bulk] - \sum_i n_i \mu_i \quad (1)$$

where $E_{tot}[AlN, X]$ is the total energy derived from doped AlN supercells, and $E_{tot}[AlN, bulk]$ is the total energy of pristine AlN with the equivalent supercell size. n_i denotes the number of atoms of type i (host or impurity atoms) that have been added to ($n_i > 0$) or removed from ($n_i < 0$) the

supercell when impurities are intercalated. μ_i is the corresponding chemical potentials of these atoms. Lower formation energies imply that the atoms are doped simpler and more stable in AlN.

The results clearly show that the substitution of an individual alkali metal on the Al site in wurtzite AlN is energetically unfavorable with only one exception. The formation energies of a separated Li on Al site are -0.82 eV for the N-rich condition and 3.29 eV for Al-rich condition. Thus Li doped AlN is stable in N-rich growth ambient. In contrast, corresponding values of Li-O complex in AlN are -4.03 eV and -0.51 eV respectively. By codoping with O, the insertion of Li and O in tetrahedral positions of AlN becomes energetically favorable regardless if they take place in Al-rich or N-rich growth conditions. A very high formation energy is generated through the Li doping due to the fact that two extra impurity states are induced leading to substantial repulsive interaction with the neighboring N ions. The defect levels and weakened repulsion were modified through the O incorporation. The main advantage of the codoping method is reducing the formation energy by forming various complexes. In III-IV group semiconductors, O incorporation has always produced a number of complexes with vacancies or dopants. [2, 18] The results of this research differ from previous works [19], more theoretical and experimental investigations are expected as the calculation of formation energy is supercell-size-dependent [20].

Fig. 1 a and b show the calculated band structure of Li-doped AlN and Li,O-codoped AlN, respectively. It can be clearly observed that both of them have the direct band gaps at the Γ point. Furthermore, the calculated electronic density of states (DOS) of AlN crystals with Li doping and Li,O-codoping were obtained (presented as Fig. 2 and Fig. 3 respectively.). It is demonstrated that the distribution of states was affected by the type of doped impurities. After the doping, the electron density changes drastically, new states are produced at the top of the valence band forcing it to shift across the Fermi level.

The peaks around the Fermi level were split with decreased amplitudes from top to bottom in Fig. 2 and Fig. 3. Moreover, Fig. 4 depicts that the DOS profiles of the doped phases are similar with slight deference. The top of the VB consists predominantly of Li and N hybridization. The peaks around the Fermi level get narrower and split, which can be attributed to the weakening bonding of Li-N with doping induced structure deformation, as discussed in Table 1.

The bottom of CB constructs from Al and N ions, the affection of Li can be neglected. The p-like states of the doped Li atoms are fairly localized which also exist in Li doped AIP system [9].

Table 1. Bond lengths along and perpendicular to the c -axis in the pristine and doped AlN as well as the change ratio of bond in doped AlN systems compared with the undoped system

Structure	Bond along c -axis		Rate of change	Bond in basal plane		Rate of change
	type	length, a.u.		type	length, a.u.	
AlN	Al-N	3.5949	-	Al-N	3.5700	-
Li: AlN	Li-N	3.8067	5.9%	Li-N	3.8444	7.7%
Li-O: AlN	Li-O	3.8365	6.7%	Li-N	3.5803	0.3%

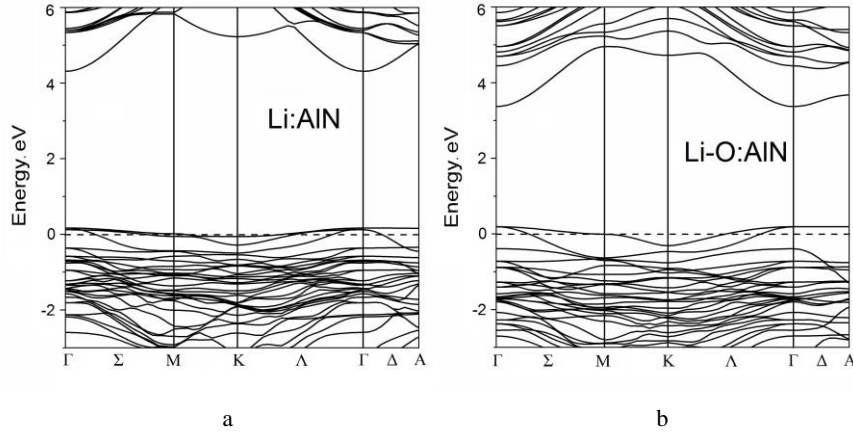


Fig. 1. Band structures for *a* Li-doped AlN and *b* Li,O-codoped AlN. The Fermi level is set as zero for reference

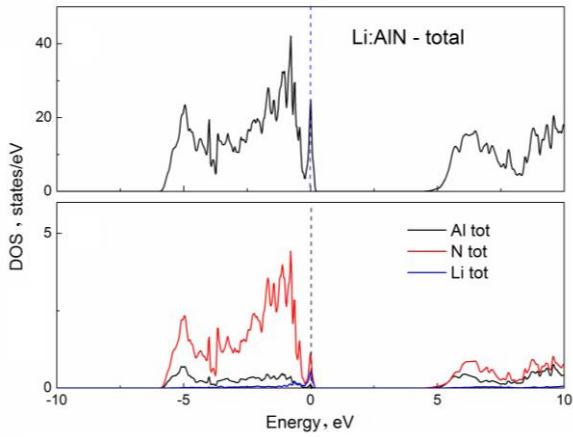


Fig. 2. DOS for Li-doped AlN: Total DOS and Partial DOS. The dashed line denotes the Fermi level

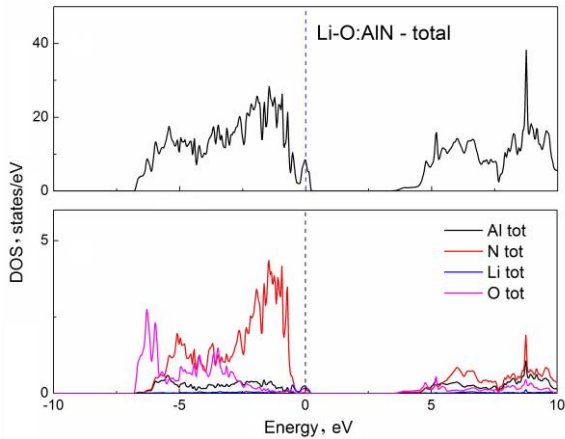


Fig. 3. DOS for Li,O-codoped AlN: Total DOS and Partial DOS. The dashed line denotes the Fermi level

It is also observed that a noticeable shift along the energy scale appears with extra O insertion, extra states are found in the lower energy levels induced. The change of patterns with given compositions can be mainly attributed to variations in the doping configuration. As mentioned above, charge redistribution effects made the cation and anion bonding in principle ionically polarized. The degree of s-p hybridization increases and the anisotropy becomes larger as the atomic number increases. As seen in the

density of states, in addition to distribution of the total states moved towards lower energies, the splitting of states around the Fermi level gradually becomes obvious going from Li to their heavier congeners doped AlN.

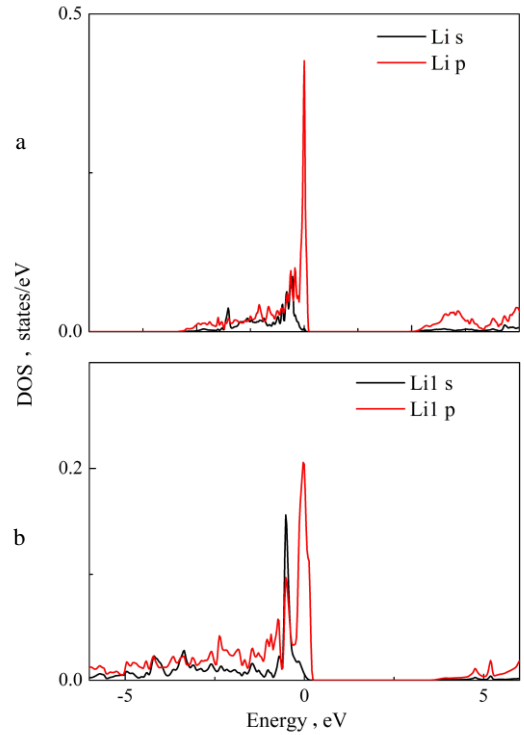


Fig. 4. DOS projected on Li atoms in *a* Li-doped AlN and *b* Li,O-codoped AlN

In Li doped AlN, Li atoms are positively charged. Strong overlap of the Li 2s, 2p states with the N electronic states will simulate an ionic-like charge transfer. The electrons compensating the holes induced by Li would be in anion (N, O) 2p bands. Due to the small radius, Li⁺ cations can move relatively independently [21] so that the inserted Li atoms are able to donate electron density towards the nearest ions. The electron donation from the doped Li mainly flows toward the O⁻ ions and partly to the N⁻ ions around in codoped supercells. Structural analysis highlights that the codoping leads to charge transfer and weakening of Al-N bonds along with smaller overlap of electronic states. The doped AlN compounds have turned into p-type semiconductors according to the calculated

band gaps. A negative relationship can be observed between the doped Li impurities and the band gap. The codoping of Li atom with extra O yields a significantly narrower band gap compared to the monoclinic doping.

4. CONCLUSIONS

The calculated electronic DOS of AlN crystals with Li-doping and Li,O-codoping demonstrated that the doped AlN still exhibits semiconductor nature with p-type character. The codoping has a general smooth effect on the profile. The top of the valence band is flat with large effective masses for localized states with anion p character in Li -doped AlN. It is possible to tune the bandgap of AlN by adjusting doping configuration. According to calculations of formation energy, codoping with O makes the intercalation of Li in tetrahedral positions of AlN energetically favorable regardless of Al-rich or N-rich growth conditions. Both Li mono-doping and codoping with O have provided an interesting way to manipulate the corresponding properties of wurtzite AlN crystals.

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