Study of induced half-metallic ferromagnetic property in CdO monolayer with non-metallic doping

Lalmuanchhana¹, Lalrinthara Pachuau², Dibya Prakash Rai², Ramesh Chandra Tiwari¹, Lalhriatzuala²

¹Department of Physics, School of Physical Sciences, Mizoram University, Aizawl 796004
²Department of Physics, Mizoram University, Pachhunga University College, Aizawl 796001

Doping is known to be an efficient way of altering the properties of materials. We have studied the structural, electronic and magnetic properties of non-metallic element (B, C and N)-doped CdO monolayer based on spin-polarized density function theory within the generalized gradient approximation (GGA) as revised for solids by Perdew, Burke and Ernzerhof (PBE-sol). Substituting B, C and N in O site with one atom in CdO monolayer induces spin polarized localized states in the gap or near the valence band and exhibit local magnetic moments ~ 1 μΒ, ~ 2 μΒ and ~ 1 μΒ respectively. It is found that the magnetic states in these doped systems are associated with the difference between the electronegativity of the dopant and the anion in the host. Our study shows that non-metal doped CdO is a promising candidate for novel nano-materials for future applications.

Keywords: Doping, CdO monolayer, non-metallic element, spin.

Introduction

Advancement in the modern technology ignites surge for the miniaturization of electronic devices. Series of 2D nano-materials with unique physical and chemical properties have drawn much attention since graphene was successfully prepared in 2004.¹ These 2D materials exhibit intriguing and abundant physical properties, ranging from metallic conductor to semiconductor, which guarantee them promising wide applications in electronic and optoelectronic devices in nano-scale.²-⁴ Many low dimensional materials such as silicone,⁵,⁶ transition metals dichalcogenide,⁷-⁹ and hexagonal boron nitride¹⁰ have been successfully prepared in experiments. Theoretical-based DFT calculations are also employed to predict new 2D material,¹¹-¹⁷ specifically among the honeycomb monolayers of II-VI group which exhibit tuneable properties.¹⁸-²⁰ Among II-VI group compounds, CdO single layer is found to exhibit a good dynamic stability using first principles calculations.²¹ With its tuneable properties, this result motivated further studies of the doped CdO single layer physical properties. But the issue of doping with non-transitional metal is that when electronegativity of the dopant is weaker than the replaced atom, it leads to ferromagnetism and if the electronegativity of the dopant is stronger than the replaced atom, it does not induce spin-polarisation.²² These results indicate that the electronegativity difference between the dopant and the anion of the host semiconductor plays an
important role for the magnetism of such doped semiconductors. However, the tuneable properties of these materials are capable of enhancing the electrical, optical, and structural properties of a semiconductor and open a new way in the advancement of miniature technology.

Cadmium oxide (CdO) is an n-type oxide belonging to the II-VI group having direct and indirect band gaps in the range of 2.2-2.5 eV and 1.36-1.98 eV respectively.\(^23\)\(^-\)\(^25\) Due to its ionic nature and low electrical resistivity and high optical transmission in the visible region, nano-scaled CdO are suitable candidate for several applications in various fields such as optical coatings, photovoltaic cells, phototransistors, IR reflectors, transparent electrodes, and gas sensor.\(^26\)\(^-\)\(^33\)

Various CdO nanostructures have been, in fact, studied in recent years.\(^34\)\(^-\)\(^40\) Zheng et al.\(^21\) confirmed the structural stability using density functional theory (DFT), Khan et al.\(^40\) studied the electronic and optical properties of CdO bulk and monolayer and provided a relative comparison. Xiaoxuan et al.\(^38\) explored electronic properties of CdO monolayer and nanotube. The strain effect on the optoelectronic properties of CdO monolayer has been investigated by Chaurasiya et al.\(^34\) and results indicate that the band gap decreases when the lattice distortion is applied and the direct-indirect gap transition can be achieved at tensile strain of 8%. Chaurasiya et al.\(^36\) also presented the magnetism induced upon point defects and doping with non-transitional metal with CdO monolayer.

The approximation considered to the correlation-exchange functional in our DFT calculation is the generalized gradient approximation (GGA) in the form of Perdew, Burke and Ernzerhof for solids (PBE-sol).\(^41\) Although popular Generalized Gradient Approximation (GGA)\(^2\) is widely used, it is biased on account of free atom energies.\(^41\) PBE-sol is the restoration of the first principles gradient expansion for exchange over a wide range of density gradients. GGA (PBE-sol) is found to be more appropriate than GGA (PBE) for structural properties in solids and are

### Table 1: Structural parameters of pure CdO ML, B-doped CdO ML, C-doped CdO ML and N-doped CdO ML. D represent the dopant atom (B/C/N). Ef is the formation energy of the doped CdO ML.

<table>
<thead>
<tr>
<th>System</th>
<th>Ground state energy (eV)</th>
<th>Bond length (Å)</th>
<th>Bond angle (°)</th>
<th>Ef (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure CdO ML</td>
<td>-121.970603</td>
<td>2.027</td>
<td>-</td>
<td>120</td>
</tr>
<tr>
<td>B-doped CdO ML</td>
<td>-117.486384</td>
<td>2.024</td>
<td>2.09</td>
<td>121.88</td>
</tr>
<tr>
<td>C-doped CdO ML</td>
<td>-119.682749</td>
<td>2.024</td>
<td>2.03</td>
<td>122.17</td>
</tr>
<tr>
<td>N-doped CdO ML</td>
<td>-120.969983</td>
<td>2.024</td>
<td>2.03</td>
<td>121.29</td>
</tr>
</tbody>
</table>

* Results obtained by Chaurasiya et al.\(^36\) using GGA-PBE.

### Table 2: Band gap of pure CdO ML, B-doped CdO ML, C-doped CdO ML and N-doped CdO ML, local magnetic moments of dopants (B/C/N) and magnetic moments per supercell of the doped structures.

<table>
<thead>
<tr>
<th>System</th>
<th>Band gap (eV)</th>
<th>Local magnetic moment of dopant(μΒ)</th>
<th>Magnetic moment per supercell (μΒ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pristine CdO</td>
<td>0.69</td>
<td>0.115</td>
<td>0.99</td>
</tr>
<tr>
<td>B-doped CdO</td>
<td>0.35</td>
<td>0.497</td>
<td>1.99</td>
</tr>
<tr>
<td>C-doped CdO</td>
<td>0.64</td>
<td>0.94*</td>
<td>0.99*</td>
</tr>
<tr>
<td>N-doped CdO</td>
<td>0.69</td>
<td>0.94*</td>
<td>0.99*</td>
</tr>
</tbody>
</table>

* Results obtained by Chaurasiya et al.\(^36\) using GGA-PBE.
in good agreement with experimental values.\textsuperscript{43} The GGA (PBE-sol) thus eliminates the biasing towards the description of free atom energies and improved equilibrium properties of densely packed solids and their surfaces.\textsuperscript{41, 44}

In this work, we present a comprehensive and comparative study of the structural, electronic and magnetic properties of B, C and N doped CdO ML obtained using spin-polarized density functional theory calculations. The structural stability of the monolayer is confirmed by the ground state energy. It is observed that the doped structures exhibit half-metallic behaviour due to the spin polarized 2p orbitals of the dopant, which are localized within the energy gap of the host lattice. The analysis of electronic structures showed that the d-orbitals of Cd play a crucial role in the electronic characteristics such as defining its valence band maxima and accommodation of most charges of its host atom.

Computational Details

In this study, a 4x4 supercell of CdO monolayer with 16 atoms each of Cd and O is taken as the undoped structure. A vacuum of 15Å is inserted along the z-direction to avoid the interactions between the monolayers during periodic calculations. Carbon, boron and nitrogen doping are done in the structure by substituting one of the oxygen atoms with a single dopant atom, thus making a doped structure with one dopant (B/C/N) atom in every 15 oxygen atoms.

The doped structures are optimized until the forces on each atom have become less than 0.02 eV/Å and by setting the energy convergence criteria of $10^{-5}$ eV. All the computations are performed within the framework of spin-polarized plane-wave density functional theory (DFT) as implemented in the Vienna Ab-initio simulation package (VASP).\textsuperscript{45} The ionic potentials are represented by projector augmented wave potentials (PAW)\textsuperscript{46} and electron exchange correlation is represented by generalized gradient approximation (GGA) as revised for solids by Perdew, Burke and Ernzerhof (PBE-sol).\textsuperscript{41} An energy cut-off of 400 eV is used for the plane-wave expansion of the electronic wave function and the Brillouin zone integration is performed using 8 x 8 x 1 k-point mesh within the Monkhorst-Pack scheme.\textsuperscript{47} For electronic structure calculations, the Brillouin zone is sampled by 16 x 16 x 1 k-point mesh.

Results and Discussion

The schematic diagram for top view of a 4x4 CdO monolayer is shown in Fig 1(a). The lattice parameter of pristine CdO monolayer is optimized by calculating the total energy of monolayer. The total energy of CdO ML after structural optimization is -121.97 eV indicating that the structure is stable. The optimized bond length between Cd and O is 2.03 Å. Cd and O atoms lie in X-Y plane, forming a graphene like honeycomb structure, suggesting sp\textsuperscript{2} hybridization. The Cd-O-Cd/ O-Cd-O bond angle is 120º, showing no buckling height suggesting a planar stability. The calculated values are comparable with reported literatures.\textsuperscript{21, 36, 38}

The electronic properties of CdO monolayer are investigated by computing the band structure and density of states. The band structure is calculated using spin polarized GGA exchange-correlation functional along $\Gamma$-M-K-$\Gamma$ Brillouin zone. The Fermi energy is set at 0 eV, as shown in Fig 2(a). The band structure showed that valence band maxima and conduction band minima are located at $\Gamma$ point indicating a direct bandgap nature with a band gap value of about 0.691 eV. The computed bandgap value is in agreement with 0.66 eV as reported by Zheng \textit{et al.}\textsuperscript{21} However, the computed value is much lower than the experimentally reported value of 3.3 eV for thin films.\textsuperscript{33} This difference may be attributed to GGA exchange-correlation functions used for present calculations, which underestimates the bandgap of semiconducting materials. The calculated total and partial density of states are shown in Fig 2(b). The electronic energy states near Fermi energy in the valence band are contributed by cadmium 4d orbitals and oxygen 2p orbitals while cadmium 5s, cadmium 4p and oxygen 2s orbitals are
Figure 2 | (a) Bandstructure of pure CdO ML. (b) PDOS of pure CdO ML.

Figure 3 | (a) Bandstructure of B-doped CdO ML. (b) PDOS of B-doped CdO ML. (c) Bandstructure of C-doped CdO ML. (d) PDOS of C-doped CdO ML. (e) Bandstructure of N-doped CdO ML. (f) PDOS of N-doped CdO ML.
Figure 4 | ELF of (a) B-doped CdO ML; (b) C-doped CdO ML; (c) N-doped CdO ML.

Figure 5 | Magnetization density of (a) B-doped CdO ML, (b) C-doped CdO ML, (c) N-doped CdO ML
contributing to the deeper valence bands. The conduction band minima are mainly contributed by cadmium 5s orbitals. The symmetrical density of states (DOS) for spin up and spin down states confirm the non-magnetic behaviour of pristine CdO monolayer.

The atomic charge density is calculated to understand the charge sharing and bonding among atoms. The DFT calculations showed that oxygen atoms act as a charge accumulating site while cadmium atoms behave like charge depleting sites. The transfer of charge from Cd atoms to O atoms is confirmed from Mulliken charge analysis, where -1.2e excess charge each is observed on oxygen atoms. Electron localization function (ELF) plot in figure 1(b) shows that the electronic charges are more localized around the O atoms and therefore the chemical bonding in CdO monolayer is mostly ionic in nature. Atomic partial charges are also computed which shows that most of the charges, i.e. 9.757e out of the total 10.828e, are located on the d orbitals of Cd atoms. While in O atoms, majority of the charges are on p orbitals, i.e., 2.803e out of 4.057e total charges and there is also a significant amount of charges on the s orbitals, i.e., 1.249e out of 4.057e total charges. This indicates that the filled valence bands near Fermi level are mostly the bonding states of 4d orbitals of Cd atoms and 2p orbitals of O atoms.

For doping, one of the O atoms in the 4x4 CdO supercell is substituted by the dopant atom. By the same parameters used in the pure CdO system, the doped structure is then optimized using spin polarized DFT calculations. All the doped structures are perfectly planar without any buckling. However, due to the insertion of smaller atoms at the O site, the bond length and bond angle slightly change around the dopant atom. The bond length between Cd and B atoms is 2.09 Å and the bond angle around the B atom, i.e., Cd-B-Cd is 120.01°. For C doped sample, bond length between Cd and C atoms is 2.03 Å and the bond angle Cd-C-Cd is 119.99°. While for N doped sample, bond length between Cd and N atoms is 2.03 Å and the bond angle Cd-N-Cd is 119.99°. Total energy of the B, C and N doped structures are -117.49 eV, -119.68 eV and -120.96 eV respectively. These negative ground state energies indicate that these doped structures are stable even though O atom is substituted with dopant atom (B/C/N), which possesses smaller ionic radius and resulted in altering of its bond length and bond angle.

Formation energy of the doped structures is calculated to understand their relative stability using the relation:

\[ E_f = E_{\text{total}}(B/C/N \text{ doped CdO}) - E_{\text{tot}}(\text{pristine CdO}) + E_{O} - E_{\text{dopant}} \]

where \( E_{\text{total}}(B/C/N \text{ doped CdO}) \) and \( E_{\text{tot}}(\text{pristine CdO}) \) are the total energies per supercell of the relaxed B/C/N doped and pure CdO ML respectively. \( E_{O} \) and \( E_{\text{dopant}} \) represent the chemical potential of O and the dopant atoms (B/C/N) respectively. Formation energies of the B-doped and C-doped CdO ML structures are +5.116 eV and 2.959 eV respectively while N-doped CdO ML has +1.67 eV of formation energy. This indicates that N doped CdO ML is the most favourable stable structure.

The calculated electronic band structure of the B, C and N doped CdO ML are given in figure 3(a), 3(c) and 3(e) respectively. All the doped structures exhibit half-metallic behaviour with one of its spin states (spin down) showing metallic behaviour while the other spin state behaves as a semiconductor. The Fermi level also lies within the valence band for the spin down channel indicating the presence of unoccupied energy states within the valence band. In B-doped CdO ML, the energy gap in the spin up channel is 0.35 eV, which is about 49% smaller than the pristine CdO ML direct gap (0.691 eV). B atom having significantly smaller atomic radius occupy O site in the lattice, the chemical bonds with the neighbouring Cd atoms become stretched and creates a large distortion, which deforms the crystal structure around the dopant atom. This influences the electronic bandstructure and it results in reduced band gap of the host lattice in agreement with reported studies on the effect of strain on CdO ML.34 The C-doped CdO monolayer on the other hand shows energy gap of 0.64 eV which remains almost the same with the pristine structure. The separation between the valence band maximum and conduction band minimum for N-doped CdO ML is around 0.69 eV at the Γ point. Among the dopants, B atom significantly modifies the band gap while C and N dopants offer almost no change as compared to the pristine CdO ML. The flat band line observed in the bandstructure of these doped MLs could be due to the formation of defect energy states by the dopant atom. The two defect states in the case of B-doped CdO ML are noticed at 0.35 eV and 0.51 eV in the forbidden region near conduction band while in the case of C-doping, the defect states are observed around -0.58 eV in forbidden region near the valence band. In the case of N-doped CdO ML, the defect states are observed slightly above the Fermi level in the conduction band. The half-metallicity and observation of defect states is similar to the work of Chaursiya et al.36

Partial density of states (PDOS) corresponding to the doped CdO ML bandstructure are also calculated to understand the atomic contribution in band structure. In B-doped CdO ML system, Fig 3(b), the valence band of the semiconducting spin up channel is contributed by Cd-5s, Cd-4p, Cd-4d, O-2p and B-2p orbitals. The flat band line observed in the bandstructure, as seen in Fig 3(a), arises due to the presence of B-2p, Cd-4d, Cd-4p, and O-2p orbitals. The PDOS in Fig 3(b), show that the spin down states...
behave as a metal due to the presence of B-2p orbital crossing Fermi level. The PDOS of C-doped CdO ML given in Fig 3(d), shows that the valence band of the spin up channel is mainly formed by the bonding states of Cd-4d, O-2p and C-2p orbitals. The plotted PDOS revealed that metallic behaviour for spin down state arises due to C-2p orbitals. Fig 3 (f) shows the calculated PDOS for N-doped CdO ML. The metallic behaviour in the spin down channel is mainly due to the energy states contributed from the N-2p orbitals and in addition to this, there are also contributions from Cd-4p, Cd-4d, and O-2p orbitals. Meanwhile the valence band of the spin up channel can be attributed to Cd-4d, O-2p and N-2p orbitals.

Electron Localization Function (ELF) of B-doped, C -doped and N-doped CdO ML are given in Fig 4(a), (b) and (c) respectively. In C-doped and N-doped structures, the electronic charges are more localized around the O atoms and the dopant C and N-atom in the doped monolayer. This charge localization is similar to the pure CdO ML and therefore the bonding in the case of C-doped and N-doped CdO ML remains ionic. Bader charge analysis also shows the charge transfer between the atoms. In C-doped CdO ML, the charge transfer in C atom is -0.6287e, and +0.895e to each of the neighbouring three Cd atoms. Meanwhile in N-doped structure, the charge transfer in N atom is -1.0048e and in the neighbouring three Cd atoms, it is +1.1097e each. The charge transfer in C atom is slightly less than that in O atom meanwhile the charge transfer in N is roughly the same as that of O atom. This is due to the lower electron affinity of the C atom in comparison to the O and N atoms. In the case of B-doped sample, the charge transfer on the B atom is only -0.1011e, and +0.7489e to each of its adjacent Cd atoms. The electronic charges are also localized around the B atom but the localization is much weaker than around the O atoms. Electronic charges around the B atom are localized almost near the middle of the chemical bonds with the adjacent three Cd atoms. Therefore, the chemical bond between Zn and B atoms is more of a covalent in nature.

From the spin-polarised calculation, pristine CdO ML resulted in zero magnetic moment in all the s, p and d orbitals which shows a non-magnetic behaviour. This is an indication that there are no half-filled atomic orbitals that could result in unpaired electrons responsible for magnetic moment. However, spin-polarized calculations on doped CdO ML show the presence of magnetic moments in the structures. The total magnetic moment of B doped CdO monolayer is 0.99 μ_B~ 1 μ_B and local magnetic moment of B atom is 0.115μ_B~ 0.12 μ_B. In C-doped system, the total magnetic moment is 1.99 μ_B~ 2 μ_B and local magnetic moment of C atom is 0.497μ_B~ 0.5μ_B. N-doped CdO ML shows a net magnetic moment of 0.991 μ_B~ 1 μ_B and N atom local magnetic moment is 0.4341 μ_B~ 0.4 μ_B. In the doped system, the magnetic moment arises mainly from the dopant atom and there are also small contributions from the neighbouring Cd and O atoms. Whatever magnetic moment observed in Cd and O atoms are therefore induced by the dopant atom (B/C/N), which decays gradually as we move further away from it. This variation is also observed in the case of charge transfer from Mulliken charge analysis. In all of the doped structures, the algebraic sum of all the magnetic moments associated with all the atoms in the system equals the net magnetic moment. Thus, it appears that all the magnetic moments associated with the dopant and the host atoms aligned along the same direction, which can happen only if there is a ferromagnetic coupling between the dopant atom (B/C/N) and the neighbouring Cd and O atoms.

The calculated spin polarized magnetization density obtained for the doped structures are shown in Fig. 5. In both the structures magnetization density is mainly localized on the dopant B/C/N atom and distributed slightly over the three nearest neighbouring Cd atoms and the next nearest neighbouring O atoms. Atomic partial magnetic moments are also calculated, which indicates that the magnetic moment in the dopant atom (B/C/N) is mainly due to the p orbitals. In B-doped system, the p-orbitals of B-atom contribute 0.110μ_B out of the total 0.115 μ_B while the corresponding p-orbitals in N atom contributes 0.418 μ_B out of the total 0.434 μ_B magnetic moments in their respective structures. In the C-doped system, p-orbitals of C contribute 0.474 μ_B out of the total 0.497 μ_B local magnetic moment associated with the C atom. Fig 5(a) shows overlapping between magnetization densities from B -atom and the three neighboring Cd atoms in B- doped ML, while it is absent in the case of C-doped and N-doped structures. This may be attributed to the fact that B-doped CdO ML has shorter bond length between B and its neighboring Cd atoms. In all the doped structures, atomic partial magnetic moment in the neighbouring three Cd atoms is largest in the d-orbitals although there are small contributions from the s and p orbitals. The secondary neighbouring O atoms have its magnetic moment mainly from the p orbitals. This calculation is in agreement with the PDOS of the doped structures in which partially filled B-2p, C-2p, N-2p, Cd-4d and O-2p orbitals result in metallic behaviour in the spin down channel. Therefore, substitution of O atom with non-metallic dopant atom (B/C/N) having smaller atomic radius and weaker electron affinity results in formation of unpaired electrons transforming semiconducting CdO ML into half-metallic ferromagnets.

Conclusion

We have studied the impact of doping in CdO monolayer using spin-polarised density functional
theory calculations. Pure CdO monolayer is a thermodynamically stable direct bandgap (~ 0.7 eV) oxide monolayer system. Our calculations show that doping of CdO ML with 2p light non-metallic elements such as B, C, and N atoms can transform a wide band gap semiconducting CdO ML into a half-metallic ferromagnets. All the dopants; B, C and N atoms being an atom with weaker electronegativity, the charge transfer from the neighbouring Cd atoms are lesser and resulted in partially filled 2p orbitals of the dopant atom and 4d orbitals of the neighbouring Cd atoms, which subsequently influence the secondary neighbouring O atoms. This results in spin polarized localization of the dopant 2p orbitals and 4d orbitals of the neighbouring Cd atoms as well as 2p orbitals of the subsequent neighbouring O atoms within the energy gap of the host CdO ML, which is responsible for the half-metallic ferromagnetic behaviour. Among the doped system, C-doped CdO ML appears to be most favourable while N-doped system appears to be least stable. Atomic partial magnetic moment calculations show that the magnetic property of the doped CdO ML is mainly due to the partially filled 2p orbitals of the dopant atom (B/C/N). Ferromagnetic coupling is also observed between the dopant atom and the three neighbouring Cd atoms and the subsequent secondary neighbouring O atoms.

Comparison of our results with the results obtained by Chaurasiya et al. on this particular study shows there are noticeable difference in its properties especially its structural parameters of the proposed ML. These differences may be attributed to the approximation considered to the correlation-exchange functionals for the study. The previous work done by Chaurasiya et al. is based on GGA (PBE) whereas our calculation is based upon GGA (PBE-sol). GGA (PBE-sol), which is the restoration of the first principles gradient expansion for exchange over a wide range of density gradients, is found to be more appropriate than GGA (PBE) for structural properties in solids and are in good agreement with experimental values. Nonetheless, these studies shows that doping of CdO ML with non-metal is a promising way to tune its electronic and magnetic properties making it a suitable 2D system for numerous future nano-electronics and spintronic devices.

References


