Density Functional Study of Electronic, Magnetic and Chemical Bonding Properties of Spinel CdCr$_2$O$_4$

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Abstract

This paper is presented a theoretical study of electronic, magnetic and chemical bonding properties of spinel CdCr$_2$O$_4$ with a general formula of AB$_2$X$_4$ (A, B=transition- metal, X=oxides, chalcogenides) using density functional calculation method combined with spin-polarized theory within generalized gradient approximation. Density functional calculation is performed to observe the effects of magnetic ordering on the electronic and chemical bonding properties of spinel CdCr$_2$O$_4$ with both cubic and tetragonal structure from a pyrochlore lattice, using Quantum ESPRESSO package. Consequently, in order to investigate the magnetic properties in paramagnetic, ferromagnetic and antiferromagnetic orderings of spinel CdCr$_2$O$_4$, a first-principles study of the electronic structure as well as chemical bonding properties of spinel CdCr$_2$O$_4$ compound in two different structural form is performed: the cubic structure in order to investigate the desired properties in paramagnetic and ferromagnetic orderings, and tetragonal structure to calculate the same properties in antiferromagnetic ordering.

Keywords: Density functional theory; Electronic; Magnetic and chemical bonding properties; Spinels; Spin-polarized theory

Introduction

Spinels are the magnesium aluminum oxide member of this large group of materials with the following formula, Mg$^{2+}$Al$^{3+}$O$_2$. It gives its name to the family of compounds that are identified by two cation sites: an octahedral site and a tetrahedral site [1]. Any material that have the general formula of A$^{2+}$[B$^{3+}$]$_2$[X$^-$]$_4$, which crystallizes in the face-centered cubic crystal system and are described by the space group $Fd\bar{3}m$ (No. 227) are labeled as a spinel. Spinel is an attracting class of materials that demonstrate rich complex behaviors at ground states. Spinel-type compounds with the general formula of AB$_2$X$_4$ (A, B=transition-metal, X=oxides, chalcogenides) have attracted extensive interest not only due to their diverse properties but also wide applications in electronics, catalysis, magnetism and electrochemical technologies [2-5]. Spinel oxides AB$_2$O$_4$ with magnetic B cations have received special attention due to their identification by three-dimensional geometrical frustration. Spinel oxides with Cr$^{3+}$ ions on the B sites are good examples to study the frustration. Spinel CdCr$_2$O$_4$ has been chosen as a model system to study because it has a well-defined magnetic order [6,7]. Spinel CdCr$_2$O$_4$ is a magnetic compound that forms into a cubic spinel structure, and the magnetic properties stem from the three-dimensional network of Cr$^{3+}$ magnetic ions, which is indicated as a corner-sharing tetrahedral [8,9]. In this paper, density functional calculations are performed to study the effects of magnetic ordering on the electronic and chemical bonding properties of spinel CdCr$_2$O$_4$, a pyrochlore lattice with magnetic Cr cations and non-magnetic Cd cations. The electronic, and chemical bonding properties of spinel CdCr$_2$O$_4$ with cubic $Fd\bar{3}m$ and tetragonal ($I4_1/amd$) structures are calculated using density functional calculation method combined with the spin-polarized theory, and compared the results in both cubic and tetragonal structures for different magnetic orderings. Density functional theory applied with the ground-state theory recovered in the zero temperature limits.

To search for the most stable structure of spinel CdCr$_2$O$_4$ in term of magnetic ordering, the lowest energy in each structure with different magnetic ordering is calculated. Fallowing that, the effect of magnetism is obtained and analyzed on the basis of total density of states, projected density of states, and charge density distribution within paramagnetic, ferromagnetic and antiferromagnetic orderings using density functional calculations and understanding of the principles of Quantum ESPRESSO in magnetic materials. In continue, to complete the findings of the electronic density of states for spinel CdCr$_2$O$_4$, the density of states for each atom is calculated, in order to analyze the band gap in each state, separately. Finally, to clarify and compare the chemical bonding properties of spinel CdCr$_2$O$_4$ in different magnetic orderings, the electronic charge density distribution are obtained for both cubic and tetragonal structures, in the (1 1 0) crystallographic planes.

Methodology

The key starting point in the existing methodology is finding a stable crystal structure of the spinel CdCr$_2$O$_4$ for both cubic and tetragonal structures as a plane wave basis, followed by obtaining the electronic, magnetic and chemical bonding properties of the compound. Therefore, the structural stability, electronic properties as well as chemical bonding properties of the spinel CdCr$_2$O$_4$ is calculated within density functional calculation method, using Quantum ESPRESSO package. All pseudopotentials are obtained from the plane wave self-consistent field (PWscf) pseudopotentials online references within generalized gradient approximation parameterized by Perdew-Burke-Ernzerhof exchange correlation calculation [10].

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pseudo potentials are applied for both cubic and tetragonal structures. In the case of generalized gradient approximation, the following valence electrons are considered in this calculation:

- Cd\(^{4+}\) (group 12) 1s\(^2\) 2 (s\(^2\)p\(^6\)) 3 (s\(^2\)p\(^4\)d\(^3\)) 4 (s\(^2\)p\(^4\)d\(^3\)) 5 (transition metal): 4d\(^3\), 5s\(^2\)
- Cr\(^{3+}\) (group 6) 1s\(^2\) 2 (s\(^2\)p\(^3\)) 3 (s\(^2\)p\(^4\)d\(^4\)) 4s\(^2\) (transition metal): 3d\(^3\), 4s\(^2\)
- 1s\(^2\) 2 (s\(^2\)p\(^3\)) (nonmetal): 2s\(^2\), 2p\(^4\)

The crystal structures of spinel CdCr\(_2\)O\(_4\) is optimized with paramagnetic, ferromagnetic and antiferromagnetic orderings. The crystal structure of spinel CdCr\(_2\)O\(_4\) is cubic with the space group \(F\bar{4}3m\) (No. 227), and the lattice constant \((a=b=c=8.667 \text{ Å}, \ a=\beta=\gamma=90°)\) in the paramagnetic and ferromagnetic ordering, whereas for the antiferromagnetic ordering is tetragonal with the space group \(I4/\text{amd}\) and lattice constant \((a=8.634 \text{ Å and } c=8.694 \text{ Å}, \ a=\beta=\gamma=90°)\). The calculated data of lattice constant is in a very good match with the experimental results in reference 6. In this reference, the results are reported \(a=8.588 \text{ Å for the cubic phase and } (c-a)/a=0.5 \times 10^{-3} (c=8.592 \text{ Å})\) for the tetragonal phase from elastic and inelastic neutron scattering measurements on CdCr\(_2\)O\(_4\) [6].

The lattice parameters of the spinel CdCr\(_2\)O\(_4\) are demonstrated for both cubic and tetragonal structures, with the Cr magnetic ions and the Cd non-magnetic. Crystal Structure is studied using XcrysDen visualization software. The density functional theory together with the general gradient approximation is used to find the electronic ground state properties of spinel CdCr\(_2\)O\(_4\). It is necessary to know that choosing the suitable crystal structure with suitable symmetry and exact lattice parameters for different magnetic orderings is the first step of the Quantum ESPRESSO calculation, and also by carrying out the self-consistent convergence test, the key of Quantum ESPRESSO calculation is finding the actual energy cut-off and k-point mesh in order to have the better and correct results. In order to calculate the total energy, the exchange and correlation functional is described within the generalized gradient approximation based on exchange-correlation energy optimization. The optimization of atomic positions and cell parameters is approved through the minimization of energy using Hellman-Feynman forces acting on atoms with the Broyden-Fletcher-Goldfarb-Shanno scheme and to get the actual relaxed atomic positions and cell parameters for each element the PWscf (VC-relax) input code is applied.

Results

In order to search for the most stable structure of spinel CdCr\(_2\)O\(_4\) in the case of magnetic orderings, the total energy in each structure is calculated.

Figure 1: 2D Charge density distribution contour of CdCr\(_2\)O\(_4\) with tetragonal structure using XcrysDen software.
CdCr₂O₄ is calculated, initially without considering the magnetization (Starting-magnetization=0.0), and then, by considering the parameter of magnetization. The spin-polarized total density of states and projected density of states for spinel CdCr₂O₄ is obtained, using 8×8×8 Monkhorst-Pack grid [12]. Figure 3 shows the total density of states of spinel CdCr₂O₄ in three different magnetic configurations. By considering the charge density plot and also the total and absolute magnetization values obtained from density of states output files (Total magnetization=0.0 and Absolute magnetization=0.0), it can be found that in the paramagnetic configuration, the two spin contributions are exactly the same, as its ground state is non-magnetic. In another word, the valence bands (corresponding to bonding states) are all doubly occupied and the total magnetization is zero. Also, by applying the initial magnetization for Cr³⁺(3d³) magnetic atoms, the same density of states calculation is calculated for ferromagnetic and antiferromagnetic orderings, as well. In the case of ferromagnetic ordering, when the spins are oriented parallel to each other, the magnetic energy is minimized. For that reason, the material may turn into a strong magnet in the presence of an externally applied magnetic field. Also, in the case of antiferromagnetic ordering, the magnetic energy is minimized when pairs of nearby spins are oriented antiparallel to each other. This state proposes that the signs of the magnetic interactions in an antiferromagnet are opposite those in a ferromagnet [13]. From Figure 3, it can be seen that in ferromagnetic ordering (Total magnetization=12.00 (Bohr magneton/cell) and Absolute magnetization=12.45 (Bohr magneton/cell)), one spin population is larger than the other and the density of states of the two spin are not aligned anymore. Although, in the antiferromagnetic ordering (Total magnetization=0.0 and Absolute magnetization=25.21 (Bohr magneton/cell)), half of the atoms have a magnetization that is opposite to the magnetization of the other half, therefore, the total magnetization is zero and the two spin contributions are the same. Also from Figure 3, it can be found that spinel CdCr₂O₄ is metallic in paramagnetic ordering as well as antiferromagnetic ordering, as the density of states can be considered to start at the bottom of the valance band and fill up the Fermi level. Since the conduction band and valance band are overlapped, the Fermi level is in conduction band and there are plenty of electrons available for conducting. While in ferromagnetic ordering, it is semiconductor as in a semiconductor, there is a small gap between the conduction and valence bands and the Fermi level is surrounded by a small band gap (1~3 eV).

In order to analyze the band gap in each state, the density of states of spinel CdCr₂O₄ is calculated for each atom, separately. Figure 4 shows the projected density of states of CdCr₂O₄ in paramagnetic ordering. It can be seen from the plot that in Cadmium, from approximately -10 to -8 eV, d orbital derived bands, in Chromium, d orbitals split into two rather sharp peaks in the interval from approximately -2 to 4 eV, and in Oxygen, from approximately -21 to -19 eV, S orbital derived bands, these bands are somewhat isolated and far away from the other bands.
that we will pay little attention to them, also from approximately -10 to -3 eV, p orbital derived bands. Therefore, according to the LSDA calculations, the band gap seems to be placed between the two crystal field split Cr d manifolds, and the O p states are deeper. On the other word, by clarifying the band for all orbitals (Cd s p d), (Cr s p d) and (O s p), it is clear that the contribution of s and p orbitals in charge density for both Cd and Cr atoms, can be neglected compare to the contribution of orbital d. In oxygen atom orbital s can be neglected because these bands are isolated and far away from the other bands.

In continue, to observe the effect of magnetization in projected density of states, the projected density of states is plotted for three different magnetic orderings of spinel CdCr₂O₄, considering the densities of Cd 4d, Cr 3d, and O 2p states with the occupied bands in the energy range down to -10 eV below the Fermi level (Figure 5). Figure 5 shows that the density of states of Cd and O atoms in cubic structure of CdCr₂O₄ are remaining the same for both paramagnetic and ferromagnetic orderings and only the density of Cr d states is changing from paramagnetic to ferromagnetic. However, the density of states of the three atoms is changed in tetragonal structure of CdCr₂O₄, but the difference in densities is manly for Cr atoms. Figure 6 shows the projected density of states of CdCr₂O₄ for both spin up and down in ferromagnetic ordering. It can be seen that the charge density in cadmium and oxygen remains the same and only in chromium atom, the population of spin up and spin down is changed as the initial magnetization is applied only on chromium atom. Figure 7 shows the projected density of states calculated for paramagnetic spinel CdCr₂O₄. As it is obvious from Figure 7, Cd d states in CdCr₂O₄ form a narrow group of bands below the bottom of O p states, which results in the appearance of a sharp peak of d DOS between -8.5 and -9.5 eV. Therefore, Cd d states have no magnetic effects, as they are completely occupied and lie well below E₁. O p-derived bands in CdCr₂O₄ spread over the energy range from -8 to -3 eV and are separated by a gap of ~1.5 eV from Cr 3d states, which give prevailing contribution to the bands crossing the Fermi level and a prominent DOS peak at E₁. Cr d states also are split into a triplet t_{2g} and a doublet e_{g} with two sharp peaks in the range from -2 to 4 eV. As it is mentioned earlier, in the transition metal oxides, the d level is fivefold degenerate. The degeneracy of d level is split into the lower energy t_{2g} level and higher energy e_{g} level by the crystal field splitting in an octahedral field which in this system chromium atom octahedrally coordinated by oxygen.

In order to explain the chemical bonding properties of spinel CdCr₂O₄ the electronic charge density is calculated in different
magnetic orderings. Figure 8 shows the electronic charge density maps along the (110) crystallographic plane. The bonding has a significant covalent character due to sharing of charge between O atoms. Moreover, the charge transfer occurs mainly from Cr atoms towards O atom. It can be found from Figure 8 that in cubic structure of CdCr₂O₄, there is no significant change in electronic charge density contour in both paramagnetic and ferromagnetic orderings, while there is a major transformation in charge density distribution as shifting from cubic to tetragonal structure. Also, in paramagnetic and ferromagnetic orderings of CdCr₂O₄, Cd and O atoms shows the ionic nature though the charge density contours around the cadmium is not completely circular but it shows the ionic bonding. Ionic bonding occurs between atoms that have opposite needs for electrons and results in a transfer of electrons. However, covalent bonding occurs between atoms that have similar needs for electrons and results in a sharing of electrons, so, due to sharing of charge between Cr and O atoms, Cd-O makes the covalent bonding. Finally, in the cubic structure of CdCr₂O₄, Cd atom shows a very week charge density but there is the stronger charge density as moving from cubic to tetragonal structure, and also the Cd-O makes partial covalent bonding in tetragonal structure. It is another reason to accept that tetragonal structure is the most stable and satisfied structure of spinel CdCr₂O₄, knowing that most of the transition metal oxides have strong covalent characteristics.

**Conclusion**

In the present work, a computational study of the electronic, magnetic and chemical bonding properties of spinel CdCr₂O₄ is reported using density functional theory (DFT) based on first principle performance within Quantum ESPRESSO method, with GGA exchange correlation functional. All the calculations perform in the basis of DFT by applying a plane wave basis set joined with spin-polarized theory. Furthermore, since chromium spinel belongs to transition metals with partially filled d-subshell, for electronic structure calculations, spin polarization had to be taken into account. In addition, the first-principles calculations are used to examine the effects of magnetic orderings on the ground state properties of this compound. It has been found from the outcomes that the most stable structure is the antiferromagnetic spinel CdCr₂O₄, with tetragonal structure compares to the other magnetic configurations in terms of energy and charge density distribution, and
also the structures would have electronic ground state ranging from metallic to semiconductor electronic character in different magnetic orderings.

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