Ab initio study of defects in CdMnTe: Electronic structure and related properties

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Abstract: Band structure, density of states, total energy and magnetic moment are calculated for ideal and defective supercell CdTe and CdMnTe by ab initio method. The optimization of crystal structure and atom relaxation has been carried out. The band gap, local levels in the band gap and magnetic moments are defined for various defective supercell CdTe and CdMnTe in ferromagnetic and antiferromagnetic states. It has been defined that as Mn atoms, the vacancy, interstitial atom and Frenkel pair in the crystal structure form magnetic moment.

Keywords: Semimagnetic Semiconductor, Electron Structure, Defect, Vacancy, Interstitial Atom, Frenkel Pair, Ab Initio, Magnetic Moment, Density of States

1. Introduction

Semimagnetic semiconductors (SMS) are the new class materials, including the properties of both ordinary and magnetic semiconductors. The main qualitative difference between SMS and ordinary semiconductors is the exchange interaction between magnetic ions and band charge carriers, which start to appear only within the external magnetic field [1]. The injection of manganese ions to the crystal lattice of A²⁺B⁶⁺ compounds and formation of solid solutions A⁺⁺Mn⁶⁺ lead to increase of band-gap width. Thus it is possible to manage these materials’ photosensitivity by Mn ions concentration in SMS. These features cause strengthening of a number of effects, as well as Faraday effect in these materials [2, 3]. On the other hand, there are unusual changes of electron structure of these materials in magnetic field, which allows managing properties of these materials with magnetic field and temperature.

The energy spectrum and wave functions of electrons for arbitrary wave vector as well as Faraday effect for Cd₁₋ₓMnₓTe thin films were obtained in our previous works [3]. We calculated interband Faraday effect in Cd₁₋ₓMnₓTe in the framework of two band model with taking into account the exchange interaction in a nonquantizing magnetic field. Cd₁₋ₓMnₓTe SMS are promising materials for γ- and x-ray detectors, solar cells, optic insulator etc. [4]. To obtain high-sensitive and radiation-resistant materials, as well as creation of devices based on them, it is necessary to know the influence of defects on physical properties of Cd₁₋ₓMnₓTe. Defects in semiconductors not only influence on electrical and optic properties of these materials, but also display their interesting physical properties.

Ab initio calculations of electron structure and magnetic properties of Cd₁₋ₓMnₓTe were carried out in some works [5-8]. Unlike these works we carried out first principle calculations based on density functional theory (DFT) by using Atomistix ToolKit (ATK) programme for study of defects on structural and magnetic properties of supercell CdTe and Cd₁₋ₓMnₓTe. In this work we have theoretically investigated electronic band structure (EBS), density of states (DOS) and magnetic moment (MM) for various ideal and defective supercells CdTe and Cd₁₋ₓMnₓTe.

2. Electronic Structure of Ideal and Defective Supercell CdTe

The first-principle calculations of electronic structure of ideal and defective supercells CdTe based on spin- polarized
DFT are performed in the local spin density approximation with regard to Hubbard-U correction (LSDA+U) on double zeta double polarized (DZDP) basis set using licensed program software ATK [9].

In this section it was given the results of calculations of EBS, DOS and MM of ideal and defective supercell Cd32Te32.

2.1. Ideal Supercell Cd32Te32

The electronic structures of ideal supercell Cd32Te32 is calculated. It was defined the band gap of ideal supercell Cd32Te32 $E_g = 1.64$ eV, the lower level of conductivity band $E_{c1} = 0.25$ eV and the upper level of valence band $E_{v1} = 1.39$ eV. The axes indicate to the spin states. The calculated value of the energy gap corresponds to experimentally measured one. In so doing, we have been able to reproduce the experimental value of the energy gap only if Hubbard-U corrections were accounted for. It has been taken the value of Hubbard-U parameter for 5p-states of Te, $U_{t_e} = 3.7$ eV [10].

DOS analysis shows that upper levels of valence band located in the range of [-5; 0] eV, mainly comes from 5p-states of Te atoms, and lower levels of conductivity band - from 5s-states of Cd atoms (Fig.1). The total energy of the ideal supercell Cd32Te32 is $E_{cell} = -68285.96$ eV.

The Cd atoms in bulk configuration (Fig.1a) is indicated by yellow color, Te atoms by red color.

The contribution to MM of Cd vacancy ($V_{Cd}$) in the supercell Cd32Te32 was defined. 4 anion atoms which have 4 broken bonds near the Cd vacancy, acquires MM $4.864\mu_B$. The MM of all 31 Cd atoms is $-9.271\mu_B$, and all 32 Te atoms is $11.276\mu_B$. The total MM of all atoms is $2.005\mu_B$.

Thus, the main contribution to the MM gives 5p orbitals of Te atoms in the upper of the valence band. The contribution from $p_x, p_y, p_z$ orbitals are the same. Calculations were performed with an accuracy of $0.001\mu_B$.

The electronic structure of supercell Cd32Te32 having $V_{Cd}$ is calculated (Fig.2). The band gap is changed from $E_g = 1.64$ eV to $E_g = 1.22$ eV. It is formed no local level in the band gap. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{c1} = 1.47$ eV, $E_{v1} = 0.25$ eV. The total energy of the supercell Cd32Te32 with $V_{Cd}$ is $E_{cell} = -66730.36$ eV.

The contribution to MM of Te vacancy ($V_{Te}$) in the supercell Cd32Te32 is insignificant. 4 cation atoms which have 4 broken bonds near the Te vacancy, acquires insignificant value of MM $0.004\mu_B$. The MM of all 31 Te atoms is $0.004\mu_B$, and all 32 Cd atoms is $0.011\mu_B$. The total MM of all atoms is $0.015\mu_B$.

The band gap is changed from $E_g = 1.64$ eV to $E_g = 1.97$ eV. It is formed 1 local level in the band gap: $E_l = 0.21$ eV. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{c1} = 1.53$ eV, $E_{v1} = -0.44$ eV (Fig.3). The total energy of the supercell Cd32Te32 having $V_{Te}$ is $E_{cell} = -67692.97$ eV. The total amount of electrons is 568 and spins of 284 of them directed upward, 284 downward.
Thus, $V_{\text{Cd}}$ decrease the band gap in the supercell Cd32Te32 but $V_{\text{Te}}$ increase it and forms 1 local level in the band gap.

### 2.2.2. Interstitial Atom in the Supercell Cd32Te32

The electronic structure of supercell Cd32Te32 having interstitial Cd atom ($I_{\text{Cd}}$) has been calculated. The band gap doesn’t changed. It is formed no local level in the band gap. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{v\uparrow}=E_{c\downarrow}=0.58eV$, $E_{v\uparrow}=E_{c\downarrow}=-1.05eV$ (Fig.4). The total energy of the supercell Cd32Te32 having $V_{\text{Cd}}$ is $E_t=-69844.74eV$.

$I_{\text{Cd}}$ atom has 3 bonds: 2 bonds with Cd atoms and 1 bond with Te atom. $I_{\text{Cd}}$ leads to forming MM. The main contribution to the MM give interstitial Cd atom 0.489µB, two Cd atoms each of them is 0.399µB and Te atom 1.516µB having bonds with interstitial Te atom. The MM of all 33 Cd atoms is -1.469µB and all 32 Te atoms is 1.464 µB. Finally the total MM of all atoms is -0.005µB.

**Fig. 4. Supercell Cd32Te32 with $I_{\text{Cd}}$: a) Bulk configuration b) EBS**

The electronic structure of supercell Cd32Te32 having interstitial Te atom $I_{\text{Te}}$ has been calculated. The band gap doesn’t changed. It is formed 1 local level in the band gap: $E_{v\uparrow}=0.29eV$, $E_{c\downarrow}=0.44eV$ (Fig.5). The total energy of the supercell Cd32Te32 with $V_{\text{Cd}}$ is $E_t=-68890.2eV$.

$I_{\text{Te}}$ atom has 5 bonds: 2 bonds with Te atoms and 3 bonds with Cd atoms. $I_{\text{Te}}$ leads to forming large MM. The main contribution to the MM give interstitial Te atom 5.467µB, two Te atoms each of them 2.521µB and 3 Cd atoms each of them 2.9µB having bonds with interstitial Te atom. The MM of all 32 Cd atoms is -15.264µB and all 33 Te atoms is 19.891 µB. Total MM of all atoms is 4.627 µB.

**Fig. 5. Supercell Cd32Te32 with $I_{\text{Te}}$: a) Bulk configuration b) EBS**

Thus, $I_{\text{Cd}}$ and $I_{\text{Te}}$ atoms in the supercell Cd32Te32 don’t change the band gap. It is formed large MM and 1 local level in the band gap in the supercell Cd32Te32 having $I_{\text{Te}}$.

### 2.2.3. Frenkel Pair in the Supercell Cd32Te32

The displaced Cd atom has 5 bonds with neighboring 2 Cd atoms and 3 Te atoms in the case of Frenkel pair $F_{\text{Cd}}$ in the supercell Cd32Te32. $F_{\text{Cd}}$ in the supercell Cd32Te32 leads to forming 2 local levels in the band gap: $E_{v\uparrow}=0.57eV$, $E_{c\downarrow}=0.43eV$. The band gap doesn’t changed. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{v\uparrow}=1.0eV$, $E_{c\downarrow}=-0.64eV$ (Fig.6). The total energy of the supercell having $F_{\text{Cd}}$ is $E_t=-68283.4eV$.

Frenkel pair $F_{\text{Cd}}$ leads to forming MM. The main contribution to MM give interstitial Cd atom -2.36µB, two Te atoms each of them 1.76µB and 2 Cd atoms each of them 0.75µB having bonds with interstitial Cd atom and neighboring Te atom 0.7451µB. The MM of all 32 Cd atoms is -4.351µB and all 32 Te atoms is 4.35 µB. Totally MM’s of all atoms compensate each other.

**Fig. 6. Supercell Cd32Te32 with $F_{\text{Cd}}$: a) Bulk configuration b) EBS**

In the case of Frenkel pair $F_{\text{Te}}$ in the supercell Cd32Te32 the displaced Te atom has 5 bonds with neighboring 2 Te atoms and 3 Cd atoms. $F_{\text{Te}}$ in the supercell Cd32Te32 leads to forming 4 local levels in the band gap: $E_{v\uparrow}=-0.05eV$, $E_{c\downarrow}=-0.3eV$, $E_{v\uparrow}=0.7eV$, $E_{c\downarrow}=-1.0eV$. The band gap doesn’t changed. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{v\uparrow}=0.25eV$, $E_{c\downarrow}=-1.35eV$ (Fig.7). The total energy of the supercell Cd32Te32 having $F_{\text{Te}}$ is $E_t=-68286.9eV$.

**Fig. 7. Supercell Cd32Te32 with $F_{\text{Te}}$: a) Bulk configuration b) EBS**

Frenkel pair $F_{\text{Te}}$ leads to forming MM. The main contribution to MM give interstitial Te atom 3.112µB, two Te atoms each of them is 1.531µB and 3 Cd atoms each of them is 0.751µB having bonds with interstitial Cd atom and neighboring Te atom 1.694µB. The MM’s of all 32 Cd atoms
3. Electronic Structure of Defective Supercell Cd$_{1-x}$Mn$_x$Te

The calculations are performed in ATK program within the spin-polarized DFT and LSDA+U on DZDP basis. We have used U$_{Mn}$ = 3.59 eV for 3d states for Mn atoms and U$_{Te}$ = 3.7 eV for 5p states of Te atoms [10, 11]. It has been given the results of calculations of EBS, DOS and MM of defective supercell Cd$_{30}$Mn$_2$Te$_{32}$ in both ferromagnetic and antiferromagnetic states.

3.1. Supercell Cd$_{30}$Mn$_2$Te$_{32}$

MM of supercells Cd$_3$MnTe$_4$ and Cd$_{15}$Mn$_2$Te$_{16}$ was calculated by ab initio method in the work [13].

We have investigated supercell Cd$_{30}$Mn$_2$Te$_{32}$ and have obtained that DOS of supercell Cd$_{30}$Mn$_2$Te$_{32}$ consist of three parts in the valence band: 1) the upper part of the valence band is mainly formed from p-states of Te atoms with some contribution of Cd s-states at bottom of the upper part 2) the middle part is formed from d-state of Mn atoms and from s-states of Cd atoms, which are below the valence band maximum on 7eV 3) the main peak at 10 eV below of the valence band maximum is formed mainly from s-states of Te atoms.

The bottom of the conductivity band is formed from s states of Cd atoms and p states of Te atoms. The peak of 2 eV above the conductivity band minimum is formed mainly from d states of Mn atoms [12].

The calculated band gap in ferromagnetic supercell Cd$_{30}$Mn$_2$Te$_{32}$ is $E_g$=1.73eV, the lower level of conductivity band is $E_{c\uparrow}$=0.632eV and the upper level of valence band is $E_{v\uparrow}$=1.05eV (Fig.8). The total energy of the supercell Cd$_{30}$Mn$_2$Te$_{32}$ in ferromagnetic supercell Cd$_{30}$Mn$_2$Te$_{32}$ is $E_B$=-66403.14eV.

3.2. Defective Supercell Cd$_{30}$Mn$_2$Te$_{32}$

3.2.1. Vacancy in the Supercell Cd$_{30}$Mn$_2$Te$_{32}$

V$_{Cd}$ in the supercell Cd$_{30}$Mn$_2$Te$_{32}$ in antiferromagnetic state forms insignificant value of MM. The main contribution to MM give two Mn atoms, of which leads to formation of MM 5.113$\mu_B$. The MM of all 30 Cd atoms -0.02 $\mu_B$ and all 32 Te atoms 0.024 $\mu_B$. The total MM’s of all atoms practically compensate each other.

Thus, impurity of two Mn atoms into the supercell Cd$_{30}$Mn$_2$Te$_{32}$ leads to forming large MM. Comparison of total energies for ferromagnetic and antiferromagnetic states shows that the antiferromagnetic state is more stable.

3.3. Supercell Cd$_{30}$Mn$_2$Te$_{32}$ in Antiferromagnetic State

The calculated band gap of supercell Cd$_{30}$Mn$_2$Te$_{32}$ in antiferromagnetic state is $E_g$=1.73eV, the lower level of conductivity band is $E_{c\uparrow}$=0.31eV and the upper level of valence band is $E_{v\uparrow}$=-1.42eV (Fig.9). The total energy of supercell Cd$_{30}$Mn$_2$Te$_{32}$ in antiferromagnetic state is $E_B$=-66403.1eV.

For the supercell Cd$_{30}$Mn$_2$Te$_{32}$ in antiferromagnetic state the main contribution to the MM give two Mn atoms, each of them leads to formation of MM 5.113$\mu_B$. The MM of all 30 Cd atoms -0.02 $\mu_B$ and all 32 Te atoms 0.024 $\mu_B$. The total MM’s of all atoms practically compensate each other. The total amount of electrons is 556 and spins of 278 of them directed upward, 278 downward.
The total energy of supercell Cd30Mn2Te32 having V_{Cd} in antiferromagnetic state is $E_t=-64848.24$ eV.

$V_{Cd}$ in the supercell Cd30Mn2Te32 in ferromagnetic state forms large MM due to the two Mn atoms, each of them leads to formation of MM 5.065 $\mu_B$, and 4 broken Te atoms with dangling bonds near the Cd atom. The MM of all 29 Cd atoms is 6.309 $\mu_B$ and all 32 Te atoms is -4.463 $\mu_B$. Finally the total MM of all atoms is 11.997 $\mu_B$.

The band gap doesn’t changed $E_g=1.73$ eV. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{c\downarrow}=1.377$ eV, $E_{v\downarrow}=-0.35$ eV. It is formed 3 local levels in the band gap $E_r=0.3$ eV, $E_r=0.13$ eV, $E_r=-0.132$ eV (Fig.11). The total energy of the supercell Cd30Mn2Te32 in ferromagnetic state with $V_{Cd}$ is $E_t=-64847.4$ eV.

Thus, in the supercell Cd30Mn2Te32 having $V_{Cd}$ it is formed 3 local levels in the band gap. It is formed MM due to the Mn atoms and Te and Cd vacancies. Comparison of total energies for ferromagnetic and antiferromagnetic states shows that the ferromagnetic state is more stable.

$V_{Te}$ in the supercell Cd30Mn2Te32 in ferromagnetic state doesn’t form large MM. The main contribution to MM ferromagnetic supercell Cd30Mn2Te32 give two Mn atoms each of them is 5.095 $\mu_B$. The MM of all 30 Cd atoms is -0.248 $\mu_B$ and all 31 Te atoms is 0.06 $\mu_B$. The total MM of all atoms is 10.002 $\mu_B$.

The band gap is changed from $E_g=1.73$ eV to $E_g=2.08$ eV. It is formed 1 local level in the band gap below the Fermi level $E_{\uparrow\downarrow}=-0.73$ eV. The top of valence band and the bottom of conductivity band are displaced to the energy levels $E_{c\uparrow}=0.3$ eV, $E_{v\downarrow}=-1.78$ eV (Fig.13). The total energy of the supercell Cd30Mn2Te32 in ferromagnetic state having $V_{Te}$ is $E_t=-65817.15$ eV.

Thus, in the supercell Cd30Mn2Te32 having $V_{Te}$ it is formed 1 local level in the band gap and band gap is increased. Contribution of $V_{Te}$ to MM is insignificant. The main contribution to MM gives two Mn atoms. Comparison of total energies for ferromagnetic and antiferromagnetic states shows that the ferromagnetic state is more stable.

### 3.2.2. Interstitial Atom in the Supercell Cd30Mn2Te32

Interstitial Cd (I_{Cd}) atom in the supercell Cd30Mn2Te32 in antiferromagnetic state leads to formation of 1 local level in the band gap: $E_{\uparrow\downarrow}=-0.68$ eV. The band gap is changed from $E_g=1.73$ eV to $E_g=2.03$ eV. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{c\uparrow}=0.2$ eV, $E_{v\downarrow}=-1.83$ eV (Fig.14). The total energy of the supercell Cd30Mn2Te32 in antiferromagnetic state having $I_{Cd}$ is $E_t=-67953.46$ eV.

Main contribution to the MM give two Mn atoms, each of them is 5,114 $\mu_B$. The contribution of $I_{Cd}$ to MM insignificant.
The MM of all 31 Cd atoms is -0.023 \( \mu_B \) and all 32, Te atoms is 0.079 \( \mu_B \). The total MM of all atoms is 0.006\( \mu_B \).

\( I_{\text{Cd}} \) atom in the supercell Cd30Mn2Te32 in ferromagnetic state forms 1 local levels in the band gap: \( E_g = -1.1 \text{eV} \). The band gap doesn’t changed from. The top of valence band and the bottom of conductivity band are shifted towards the energy levels \( E_{\text{c} \uparrow} = -2.31 \text{eV}, E_{\text{c} \downarrow} = -0.58 \text{eV} \) (Fig. 15). The total energy of the supercell Cd30Mn2Te32 in ferromagnetic state having \( I_{\text{Cd}} \) is \( E_i = -67953.26 \text{eV} \).

\( I_{\text{Cd}} \) atom has 6 bonds: 3 bonds with Te atoms and 3 bonds with Cd atoms. The main contribution to the MM give two Mn atoms each of them is 5.153\( \mu_B \), interstitial Cd atom -2.143\( \mu_B \) and 3 Te atoms having bonds with interstitial Cd atom, each of them 2.049\( \mu_B \). The MM of all 31 Cd atoms is -0.023 \( \mu_B \) and all 32 Te atoms is 0.079 \( \mu_B \). The total MM of all atoms is 10.005\( \mu_B \).

Main contribution to the MM give two Mn atoms each of them is 5.64 \( \mu_B \). \( I_{\text{Tc}} \) doesn’t lead to forming large MM. The MM of all 30 Cd atoms is 0.107 \( \mu_B \) and all 33 Te atoms is -0.491 \( \mu_B \). The total MM of all atoms is -0.006\( \mu_B \).

\( I_{\text{Tc}} \) in the supercell Cd30Mn2Te32 in ferromagnetic state leads to forming 2 local levels in the band gap: \( E_{\text{c} \uparrow} = 0.4 \text{eV}, E_{\text{c} \downarrow} = 0.54 \text{eV} \). The band gap doesn’t changed \( E_g = 1.73 \text{eV} \). The top of valence band and the bottom of conductivity band are shifted towards the energy levels \( E_{\text{c} \uparrow} = 0.783 \text{eV}, E_{\text{c} \downarrow} = 0.955 \text{eV} \) (Fig 17). The total energy of the supercell Cd30Mn2Te32 in ferromagnetic state with \( I_{\text{Tc}} \) is \( E_i = -66997.02 \text{eV} \).

To move Te atom from its position to the interstitial one it is have to break 4 bonds with Cd atoms. After the displacement the interstitial Te atom has 6 bonds - 3 bonds with Te atoms and 3 bonds with Cd atoms. Contribution to MM give these 6 atoms, each Cd atom -2.443\( \mu_B \) and each Te atom 2.823\( \mu_B \), interstitial Te atom itself 4.632\( \mu_B \) and two Mn atoms each of them 4.806\( \mu_B \). The MM of all 30 Cd atoms is -9.16 \( \mu_B \) and all 33 Te atoms is 13.52 \( \mu_B \). The total MM of all atoms is 12.994\( \mu_B \).

Thus, in the supercell Cd30Mn2Te32 having \( I_{\text{Cd}} \) it is formed 1 local level in the band gap. Band gap is increased in antiferromagnetic state and doesn’t changed in ferromagnetic state. The contribution of \( I_{\text{Cd}} \) atom to MM in antiferromagnetic state is insignificant. The main contribution to MM give two Mn atoms and \( I_{\text{Cd}} \) atom in ferromagnetic state. Comparison of total energies for ferromagnetic and antiferromagnetic states shows that the ferromagnetic state is more stable.

\( I_{\text{Tc}} \) in the supercell Cd30Mn2Te32 in antiferromagnetic state it is formed no local level in the band gap. The band gap doesn’t changed. The top of valence band and the bottom of conductivity band are shifted towards the energy levels \( E_{\text{c} \uparrow} = -0.1 \text{eV}, E_{\text{c} \downarrow} = -1.6 \text{eV} \). The total energy of the supercell Cd30Mn2Te32 in antiferromagnetic state having \( I_{\text{Tc}} \) is \( E_i = -66981.54 \text{eV} \) (Fig 16).

Main contribution to the MM give two Mn atoms each of them is 5.64 \( \mu_B \). \( I_{\text{Tc}} \) doesn’t lead to forming large MM. The MM of all 30 Cd atoms is 0.107 \( \mu_B \) and all 33 Te atoms is -0.491 \( \mu_B \). The total MM of all atoms is -0.006\( \mu_B \).

Thus, in the supercell Cd30Mn2Te32 having \( I_{\text{Cd}} \) it is formed no local level in the band gap in antiferromagnetic state and 2 local levels in ferromagnetic state. Contribution of \( I_{\text{Tc}} \) atom to MM in antiferromagnetic state is insignificant. The main contribution to MM give two Mn atoms and \( I_{\text{Tc}} \) atom in ferromagnetic state. Comparison of total energies for ferromagnetic and antiferromagnetic states shows that the antiferromagnetic state is more stable.

### 3.2.3. Frenkel Pair in the Supercell Cd30Mn2Te32

Frenkel pair \( F_{\text{Cd}} \) in the supercell Cd30Mn2Te32 in antiferromagnetic state leads to forming of 1 local level in the band gap: \( E_{\text{c} \uparrow} = 0.38 \text{eV} \). The band gap is changed from \( E_g = 1.73 \text{eV} \) to \( E_g = 1.9 \text{eV} \). The top of valence band and the bottom of conductivity band are shifted towards the energy levels \( E_{\text{c} \uparrow} = 1.4 \text{eV}, E_{\text{c} \downarrow} = -0.5 \text{eV} \) (Fig.18). The total energy of the supercell Cd30Mn2Te32 in antiferromagnetic state having \( I_{\text{Tc}} \) is \( E_i = -66400.55 \text{eV} \).

The displaced Cd atom has 3 bonds with neighboring Cd atoms. Frenkel pair \( F_{\text{Tc}} \) in the supercell Cd30Mn2Te32 in antiferromagnetic state leads to forming MM. The main contribution to MM give two Mn atoms each of them is 5.241 \( \mu_B \). The MM of all 30 Cd atoms is 0.221\( \mu_B \) and all 32 Te atoms is -0.707 \( \mu_B \). The total MM of all atoms is -0.619\( \mu_B \).
Displacement of Cd atom in the supercell Cd30Mn2Te32 in ferromagnetic state leads to forming F$_{Cd}$. Cd atom breaks 4 bonds with Te atoms and displaces to the interstitial position and forms 4 new bonds with 2 Cd atoms and 2 Te atoms. F$_{Cd}$ in the supercell Cd30Mn2Te32 in ferromagnetic state leads to formation of 2 local levels in the band gap: $E_v^\uparrow=0.51eV$, $E_v^\downarrow=0.62eV$. The band gap is changed from $E_g=1.73eV$ to $E_g=1.89eV$. The top of valence band and the bottom of the conductivity band are shifted towards the energy levels $E_{\uparrow\downarrow}^v=-0.67eV$, $E_{\downarrow\uparrow}^v=1.215eV$ (Fig.19). The total energy of the supercell Cd30Mn2Te32 in ferromagnetic state having $I_{Fe}$ is $E_{\uparrow}\uparrow=-66400.88eV$.

Frenkel pair F$_{Cd}$ in ferromagnetic state leads to forming MM. The main contribution to MM give two Mn atoms, each of them have $5.081\mu_B$, interstitial Cd atom with MM of $1.372\mu_B$ and two Te atoms with MM of $1.91\mu_B$ and $0.396\mu_B$. The MM of all 30 Cd atoms is $-2.663\mu_B$ and all 32 Te atoms is $2.623\mu_B$. The total MM of all atoms is $10.002\mu_B$.

Thus, in the supercell Cd30Mn2Te32 having F$_{Cd}$ it is formed 1 local level in the band gap in antiferromagnetic state and 2 local levels in ferromagnetic state. The band gap is increased. Contribution of F$_{Cd}$ atom to MM is insignificant. The main contribution to MM give two Mn atoms. Comparison of total energies for ferromagnetic and antiferromagnetic states shows that the antiferromagnetic state is more stable.

Displacement of Te atom leads to formation of F$_{Te}$ in the supercell Cd30Mn2Te32. Te atom breaks 4 bonds with Cd atoms displace to the interstitial position and forms 4 new bonds with 2 Cd atoms and 2 Te atoms.

Frenkel pair F$_{Te}$ in the supercell Cd30Mn2Te32 in antiferromagnetic state leads to formation of 4 local levels in the band gap: $E_v^\uparrow=0.8$, $E_v^\downarrow=-0.1eV$, $E_c^\downarrow=0.25eV$, $E_c^\uparrow=-0.3eV$. The band gap is changed from $E_g=1.73eV$ to $E_g=2.0eV$. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{\uparrow\downarrow}^v=-1.5eV$, $E_{\downarrow\uparrow}^v=1.5eV$. The band gap is changed from $E_g=1.73eV$ to $E_g=2.0eV$. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{\uparrow\downarrow}^v=-0.5eV$, $E_{\downarrow\uparrow}^v=-1.5eV$. The total energy of the supercell Cd30Mn2Te32 in ferromagnetic state having $I_{Fe}$ is $E_{\uparrow}\uparrow=-66405.26eV$.

Frenkel pair F$_{Te}$ in the supercell Cd30Mn2Te32 in ferromagnetic state leads to formation of 3 local levels in the band gap: $E_{t}\uparrow=0.1$, $E_{t}\downarrow=-0.3eV$, $E_{c}\downarrow=-0.5eV$. The band gap is changed from $E_g=1.73eV$ to $E_g=2.0eV$. The top of valence band and the bottom of conductivity band are shifted towards the energy levels $E_{t}\uparrow=0.5eV$, $E_{c}\downarrow=-1.5eV$. The total energy of the supercell Cd30Mn2Te32 in antiferromagnetic state having $I_{Fe}$ is $E_{\uparrow}\uparrow=-66404.26eV$.

Thus, in the supercell Cd30Mn2Te32 having F$_{Te}$ it is formed 4 local levels in the band gap in antiferromagnetic state and 3 local levels in ferromagnetic state. The band gap is increased. Contribution of F$_{Te}$ atom to MM in the supercell Cd30Mn2Te32 having $I_{Fe}$ is $5.113\mu_B$, 2.445 $\mu_B$, $3.495\mu_B$, and $-3.719\mu_B$. The MM of all 30 Cd atoms is $-0.221\mu_B$ and all 32 Te atoms is $0.707\mu_B$. The total MM of all atoms is $1.022\mu_B$.
ferromagnetic and antiferromagnetic states shows that the antiferromagnetic state is more stable.

4. Conclusion

It was calculated band structure, density of states and magnetic momentum for ideal and defective supercell CdTe and CdMnTe by ab initio method. The band gap, local levels in the band gap and magnetic momentum are defined for various defects in ferromagnetic and antiferromagnetic states. It has been defined, that as Mn atoms, the vacancy, interstitial atom and Frenkel pair in the crystal structure form magnetic moment.

It has been defined that \( V_{\text{Cd}} \) decrease the band gap in the supercell Cd32Te32 but \( V_{\text{Fe}} \) increase it and forms 1 local level in the band gap.

\( I_{\text{Cd}} \) and \( I_{\text{Fe}} \) atoms in the supercell Cd32Te32 don’t change the band gap. It is formed large MM and 1 local level in the band gap in the supercell Cd32Te32 having \( I_{\text{Fe}} \).

\( F_{\text{Cd}} \) and \( F_{\text{Fe}} \) atoms in the supercell Cd32Te32 don’t change the band gap. \( F_{\text{Cd}} \) forms 2 local levels and \( F_{\text{Fe}} \) forms 4 local levels in the band gap in the supercell Cd32Te32. It is formed MM in the supercell Cd32Te32 having \( F_{\text{Cd}} \) or \( F_{\text{Fe}} \) atom.

Impurity of two Mn atoms into the supercell Cd30Mn2Te32 leads to forming large MM. The antiferromagnetic phase is more stable in the supercell Cd30Mn2Te32.

In the supercell Cd30Mn2Te32 having \( V_{\text{Cd}} \) it is formed 3 local levels in the band gap. It is formed MM due to the Mn atoms, Te and Cd vacancies. The antiferromagnetic state in the supercell Cd30Mn2Te32 having \( V_{\text{Cd}} \) is more stable.

In the supercell Cd30Mn2Te32 having \( V_{\text{Te}} \) it is formed 1 local level in the band gap and band gap is increased. The contribution of \( V_{\text{Te}} \) to MM is insignificant. The main contribution to MM give two Mn atoms. The antiferromagnetic state is more stable in the supercell Cd30Mn2Te32.

In the supercell Cd30Mn2Te32 having **I** \( \text{Fe} \) it is formed no local level in the band gap in antiferromagnetic state and 2 local levels in ferromagnetic state. The contribution of \( I_{\text{Fe}} \) atom to MM is insignificant. The main contribution to MM give two Mn atoms and \( I_{\text{Fe}} \) atom in ferromagnetic state. The antiferromagnetic state is more stable.

In the supercell Cd30Mn2Te32 having \( F_{\text{Cd}} \) it is formed 1 local level in the band gap in antiferromagnetic state and 2 local levels in ferromagnetic state. The band gap is increased. The contribution of \( F_{\text{Cd}} \) atom to MM is insignificant. The main contribution to MM give two Mn atoms. The antiferromagnetic state is more stable in the supercell Cd30Mn2Te32 having \( F_{\text{Cd}} \).

In the supercell Cd30Mn2Te32 having \( F_{\text{Te}} \) it is formed 4 local levels in the band gap in antiferromagnetic state and 3 local levels in ferromagnetic state. The band gap is increased. The contribution of \( F_{\text{Te}} \) atom to MM in the supercell Cd30Mn2Te32 in antiferromagnetic state having \( F_{\text{Te}} \) is insignificant. The main contribution to MM give two Mn atoms and \( F_{\text{Te}} \) in ferromagnetic state. The antiferromagnetic state is more stable.

The supercell Cd30Mn2Te32 is more stable the supercell Cd30Mn2Te32 having \( F_{\text{Te}} \).

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