Structural, electronic and magnetic properties of the ordered binary FePt, MnPt, and CrPt3 alloys

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A B S T R A C T

We perform ab initio simulations to investigate the structural, electronic and magnetic properties of the ordered binary FePt, MnPt, and CrPt3 alloys. In particular, equilibrium structural lattice parameters, electronic properties such as density of states (DOS), partial density of states (PDOS) and electronic band structure of each binary alloys are investigated and interpreted. Moreover, the magneto-crystalline anisotropy energies (MAE) are calculated. We found MAE values of FePt, MnPt and CrPt3 ordered alloys to be 2.66, 0.46 and 0.42 meV/f.u., respectively, corresponding to magneto-crystalline anisotropy constant K of $7.6 \times 10^7$, $1.3 \times 10^7$ and $1.1 \times 10^7$ erg/cm$^3$, respectively. The large MAE and K values reveal that FePt, MnPt and CrPt3 binary alloys are eligible to be key components in magneto-optical and opto-electronic devices. In addition, we estimated the Curie temperatures of the three ordered alloys from exchange energy. We found the $T_C$ of L10-FePt, L10-MnPt and L12 CrPt3 to be 955 K, 989 K and 762 K, respectively. The high Curie temperatures obtained enable the ordered alloys to serve as write assist in Heat-Assisted Magnetic Recording (HAMR). We believe that our findings would pave the way to fabricate bulk and thin films based on the ordered binary FePt, MnPt, and CrPt3 ordered alloys that have attractive electronic and magnetic properties.

1. Introduction

In the past two decades, the interest in hybrid materials formed by coupling transition metals with platinum (Pt) has attracted much attention both experimentally and theoretically due to their excellent structural and magnetic properties. In particular, high chemical and thermal stabilities, good display performance and a large magneto-crystalline anisotropy (MAE) of the ordered binary FePt MnPt, and CrPt3 alloys made them superior to other binary alloys used in magneto-optical recording devices [1, 2, 3, 4] and communication devices [5], optical filters [6], biomedical applications [7] and spintronic applications [8, 9]. The FePt and MnPt binary alloys are considered as potential candidates for state-of-art spintronic applications like magnetic random access memory (MRAM) [10, 11] and drivers in micro- and nano-electromechanical systems (MEMS/NEMS) [12,13,14,15,16]. It has been reported that high-purity L10 FePt NPs could be obtained by controlling the proportion of iron and platinum only in the vicinity of the equiatomic composition [17]. Several works have focused on the investigation of FePt thin films and related materials [18, 19, 20, 21]. First-principles based calculations addressing the magnetic and electronic properties of L10 FePt and MnPt alloys have presented interesting results [22]. The out-of-plane ground state Magnetocrystalline anisotropy energy (MAE) of MnPt was found to be more than an order of magnitude (~0.1 meV/f.u.) less than that of FePt (~2.9 meV/f.u.) [23, 24]. The synthesis and characterization of a well-organized magnetic selection of such materials can promote the design of magnetic media sufficiently proficient of recording densities beyond 1 Tb/in$^2$ [25,26].

The MAE is defined as the change in the total energy associated with a change of the orientation of the spin moment along different crystallographic axes of the crystal. The contributions to MAE are two-fold: The first is the volume shape anisotropy and the second is magnetocrystalline anisotropy. The shape anisotropy is a long-range effect depends on the shape of the crystal arise from the dipole-dipole interactions that favor always the in-plane orientation of the magnetization axis [17]. The MAE arises mainly from the spin-orbit interaction which is a short-range effect localized around atomic cores [18,20,21]. Moreover, MAE indicates the degree of magnetic stability and is related to the value of the spin magnetic moment of certain ferromagnetic materials [19]. In this work, we investigate the structural, electronic and magnetic properties of the ordered binary FePt, MnPt, and CrPt3 alloys that crystallize in L10 (also

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known as AuCu and L12 (known as AuCu3) as illustrated schematically in Figures 1 and 2.

It is well known that the value of MAE can be tuned over broad range by varying c/a ratio [24]. Experimental investigations have reported c/a ratio to be 0.96, 0.92 and 1.00 for FePt, MnPt, and CrPt3 alloys, respectively [25, 26, 27]. Several theoretical calculations and experimental measurements have been performed to elucidate the values of MAE of L10-like FePt1-x structures. C.J. Aas [28] performed first-principles calculations of the MAE of the L10-like FePt1-x samples. Several theoretical studies have used the coherent potential approximation (CPA) [18, 29, 30] to determine the MAE of partially ordered alloys. Furthermore, a pioneering experimental study by Barmak et. al [31] reported an epitaxial growth of FePt films with nominal thicknesses of 42 or 50 nm by sputtering onto single crystal MgO(001) substrate. The calculated MAE was found to be in the range 0.453–0.775 meV/f.u. at Fe concentration 46%-66%. Rie Y. Umetsu [32] presented theoretical calculations of MAE of the L10-type MnPt alloy using local spin-density approximation (LSDA) and implementing the Linear Muffin-Tin Orbital (LMTO) method with atomic sphere approximation including the spin-orbit interaction [3]. The magnitude of the MAE of L10-type was estimated to be about 0.51 meV/f.u. This corresponds to a magnetocrystalline anisotropy constant Kc/a of approximately 1.39 × 107 erg/cm3. A pioneering theoretical study reveals that the CrPt3 is the most stable phase of the CrxPt1-x system and it exhibits an abnormally large MAE [2, 25, 33]. This inconsistent with the fact that the most stable phase of CrPt3 is the cubic AuCu structure, in which symmetrical considerations would discard a large MAE value [34, 35, 36]. The large MAE value of CrPt3 systems has attracted a great deal of interest since such materials can be used in the fabrication of novel magneto-optical devices. P.M. Oppeneer [2] examines computationally the origin of the unique magnetism observed in crystalline CrPt3 films and found that CrPt3 has a large orbital moment of 0.15 μB develops on Cr parallel to the spin moment of 0.72 μB. The MAE in the cubic 3d transition metals is a very small quantity of only a few μeV/f.u. [36, 37, 38]. In light of the above, we are motivated to calculate MAE of the three ordered alloys with high degree of accuracy and to interpret the physical origin of such large values.

2. Computation details

We have done self-consistent electronic structure calculations that are predominantly carried out within the spin-polarized DFT approach implemented in the Vienna ab initio simulation package (VASP) [37, 38, 39, 40]. The Projector Augmented Wave (PAW) pseudopotentials [41, 42, 43] are used to describe electron-ion interactions. The electronic configuration of the metals involved in the three ordered binary alloys are as follow: Fe: 3d7 4s1, Mn: 3d6 4s2 Cr: 3d5 4s1 and Pt: 5d9 6s1, respectively. The exchange-correlation functional [44] is accounted for by using the spin-polarized generalized gradient approximation (GGA) as parameterized by Perdew–Burke–Ernzerhof (PBE) [45, 46, 47].

Previous studies have emphasized the importance of k-sampling in the full Brillouin zone to get stable and reliable results of MAE of FePt alloys [48]. Consequently, we used large supercell to ensure high accuracy in determination of MAE values. We did carry out sets of calculations of all binary alloys with a mesh-grid of 24 × 24 × 24 k-point corresponds to a slightly less than 14,000 k-points. We use an energy cut-off of 300 eV. The spin-orbit coupling was taken into account only for the calculation of MAE using the force theorem [49, 50]. MAE is calculated in terms of the difference in band energies of the two magnetization directions with spin-orbit coupling included in the Kohn-Sham equation. Two sets of self-consistent calculations were performed, one with magnetic moments aligned parallel “E⊥” and the other oriented perpendicular “E⊥ “to the easy axis (c-axis). According to this approach:

\[
MAE = E_\perp - E_\parallel.
\]

We employed a model based on simultaneous use of force theorem and LSDA to compute the MAE of all the binary structures investigated in this work. The justification of implementing this approach is vindicated by the vanishing charge- and spin-density variations confirming that the spin-orbit coupling is correctly computed [44]. The LSDA method is very simple and promising for first principles determinations of MAE. In this method, on-site Coulomb potential depends on spin only. More precisely, LSDA is the best for transition-metal compounds where the spin-orbit coupling generally lowers the symmetry of a magnetic system compared to its crystalline symmetry. The local spin density + Hubbard U approximation (LSDA + U) was shown to provide a reliable representation of the magnetic ground state features when Coulomb correlations among atomic species are included for both 3d and 5d elements in FePt. [51]. The electron-electron interaction plays a crucial role in the determination of MAE in d- [52] and f-electron [53] magnetic materials. The pioneering study implemented the LSDA + U method and full potential linearized augmented plane wave method [53, 54] to describe the Coulomb repulsion U among different atomic species. The main idea of LSDA + U method is that “+U” potential has both onsite spin- and orbital-contributions. The charge/spin densities are converged better than 5 × 10−5 electron/(a.u.)2 in order to accomplish accurate total energy convergence. In calculating the ionic relaxation (relaxed geometry), we performed a self-consistent (SC) calculation using a conjugate gradient algorithm (CG) [55, 56, 57]. We constructed the unit cell of each of the investigated alloys by starting up with experimental lattice parameters (a = 3.86, c/a = 0.96, a = 4.00, c/a = 0.92 and a = 3.87, c/a = 1) for FePt, MnPt and CrPt3 alloys, respectively [3, 25, 27]. All the structures are then fully relaxed as described above. For FePt and MnPt, a 9 × 9 × 9 Monkhorst–Pack k-point mesh [58, 59] and an energy-cutoff of 300 eV were sufficient for energy convergence while for the CrPt3, we used a 6 × 6 × 6 Monkhorst–Pack k-point mesh. The Methfessel Paxton method [60, 61] is implemented for all structures to speed up the
integration over the Brillouin zone. The relaxed atomic positions were followed by minimizing the total energy (i.e., as small as 10⁻⁶ eV), and the Hellmann–Feynman forces. These forces were as small as 0.0003 eV/Å at convergence in the unit cell of all different electronic structure relaxations that has been performed. Furthermore, to ensure high accuracy of calculating the DOS of each alloy, we used 18² centered Monkhorst Pack k-point mesh with an energy cut-off of 400 eV.

3. Results

3.1. Crystal structures and total energy minimization approach

Generally, L₁₀ MPt [M = Mn, Fe] structure consists of two types of atoms as shown in Figure 1. Mainly, Mn or Fe atoms occupy the [0 0 0] and [½ ½ 0] lattice sites and Pt atoms occupy the corresponding face center with [½ 0 ½], and [0 ½ ½] site positions [24, 62, 63], respectively. As shown in Figure 1, both FePt and MnPt exhibit a distorted tetragonal structure labeled by L₁₀ with space group (P4/mmm). Total energy minimization plots of L₁₀ phase of FePt and MnPt ordered alloys are shown in Figures 3 and 4. The obtained lattice parameters were found to be \(a = 3.88\ \text{Å},\ \ c = 3.73\ \text{Å},\ \ c/a = 0.961\) and volume \(= 56.1\ \text{Å}³\) for FePt ordered alloy and, \(a = 4.03\ \text{Å},\ \ c = 3.69\ \text{Å},\ \ c/a = 0.916\) and volume \(= 60\ \text{Å}³\) for MnPt ordered alloy, respectively. As illustrated in Figure 2, CrPt₃ was found to exhibit the cubic L₁₂ phase with space group (Pm̅₃m). In L₁₂ structure Cr and Pt atoms are located at the corner and face center, respectively with [0 0 0] for Cr and [½ ½ ½], [½ ½ 0], and [0 ½ ½] lattice sites for Pt [64, 65]. Total energy minimization plot for L₁₂ structure is shown in Figure 5. The calculated lattice parameters of the L₁₂ structure was found to be, \(a = c = 3.91\ \text{Å}\) and volume \(= 60.2\ \text{Å}³\).

The Antiferromagnetic (AFM) of MnPt and Ferromagnetic (FM) FePt phases have been intensively studied in past two decades due to their inherited outstanding properties. They were used for quantum computing applications and high-density magnetic recording. Both MnPt and FePt binaries crystallize in tetrahedral distorted L₁₀ (CuAu) atomic structure. Total energy minimization approach is used to calculate the equilibrium lattice constants of both binaries, as well as, (CuAu₃) CrPt₃ ordered alloys. The calculated equilibrium lattice constants of the three ordered binary alloys are listed in Table 1.

The details of the crystal structures and total energy minimization approach implemented for obtaining the optimized lattice parameters of the three ordered magnetic alloys are provided in the supplementary of this manuscript.

3.2. Density of state and band structure (BS) of FePt, MnPt, and CrPt₃ binaries

To obtain a deeper insight into the electronic properties of FePt, MnPt, and CrPt₃ ordered alloys, an accurate investigation of electronic DOS and BS are important. Figures 6, 7, 8, 9, 10, and 11 show our results of the total density of state (TDOS), partial density of state (PDOS) and band structure of the FePt compound. Figure 9 shows the TDOS where the Fermi level is set to 0 eV. The contributions to TDOS of FePt come mainly from the hybridization of Fe-3d and Pt-5d electronic states while the s and p electronic states contributions are found to be negligibly small as shown in Figures 7 and 8. As indicated by figures, the highest DOS of the majority spin "spin up" valence bands, are characterized by energy values located at ~ -3.1 eV, ~ -1.9 eV, and ~ -0.9 eV whereas, the highest DOS of the minority spin "spin down." conduction bands, are found at ~ 0.46 eV, ~ 0.78 eV, and ~1.44 eV. We found that the width of the majority band ~6 eV is narrower than the corresponding energy width of the minority band ~ 8eV which means that the DOS is larger in the minority bands than in the majority bands. The behavior of TDOS near \(E_f\) depends largely on the contribution of Fe-3d states and exhibits a low value near the Fermi energy where the lower energy part of the Fe-3d states are almost empty and the Pt-3d states are almost completely empty.
occupied. Therefore, the contribution of one-spin channel of the Fe-3d states electrons to the hybridization with electrons of neighboring atoms is extremely small. This weaker hybridization leads usually to the narrowing of the bandwidths and enhancing the exchange splitting. Furthermore, we present the projected DOS for the Fe, Pt atoms and FePt bulk as shown in Figures 6, 7, and 8. The d bands of both Fe and Pt significantly overlap below Ef in the majority spin state indicating that both d bands in the majority spin state strongly hybridize with each other. Therefore, ferromagnetic spin configuration of FePt originates from the d-d hybridization between Fe and Pt. However, the minority spin state, d bands of Pt and Fe form bonding and anti-bonding states respectively; this is where hybridization comes into place.

Figure 9 displays the electronic band structure of FePt system. It is obvious that the FePt system is metallic in nature. By careful inspection of Figures 10 and 11, we can distinguish three regions of interest. Mainly, the region dominated by Pt-5d electronic states in the (-7 to -4 eV) range below Ef level. The second region characterized by the strong hybridization of Pt-5d states and Fe-3d states in the energy range (-4 to -1.5 eV). The third region above Ef level is dominated by minority spin states of Fe-3d occupying the empty levels of the conduction band. Moreover, the Fe-3d and Pt-5d states hybridization for majority spin becomes stronger than that for minority spin. One can see clearly that the 3d states of Fe atoms lie on the higher-energy side of the 5d states of Pt atoms because these states lie slightly on the lower energy side of Fe-3d states as the difference between the Fe and Pt is two electrons per atom.

The DOS and PDOS plots of MnPt ordered system are shown in Figures 12, 13, and 14. The figures illustrate the majority and minority spin states, respectively. Inspection of Figure 12 demonstrates clearly that the MnPt has an extremely low TDOS to the right of Ef level. As seen before for FePt, the contributions to TDOS of MnPt come mainly from Mn-3d and Pt-5d electronic states and it depends largely on the contribution of Mn-3d states. The contributions of s and p electronic states are negligible as illustrated in Figures 13 and 14. The width of the majority band is ~6 eV narrower than the corresponding energy width of the minority band (~8.5 eV) indicating a larger DOS in the majority bands than the corresponding DOS in the minority bands.

The d bands of both Mn and Pt significantly overlap below the Ef in the majority spin state as shown in Figure 12. This means that both d bands in the majority spin state are strongly hybridized indicating that the ferromagnetic spin configuration of MnPt originates from the d-d hybridization between d states of Mn and Pt, while in the minority spin state, the d bands of Pt and Mn form bonding and anti-bonding states respectively, this is where hybridization comes into place analogous to what we found for FePt alloy.

As seen before for FePt, Figure 15 shows that MnPt system has zero bandgap indicating that it is a pure metallic compound. Figures 16 and 17 indicate that the Mn-3d and Pt-5d states hybridization for majority spin is considerably stronger than the corresponding hybridization for minority spin states.

Careful inspection of Figures 6, 7, 8, 10, 11, 12, 13, 14, and 16 that display L10 FePt and MnPt ordered binaries (TDOS and PDOS). Our results indicate that Fermi level of ordered MnPt alloy is slightly higher than that of ordered FePt alloy. For both ordered FePt and MnPt alloys, a

| Material | a(Å) | c(Å) | a ex(Å) | c ex(Å) | a th(Å) | c th(Å) | a/c | a ex/c ex | c/a th/c th |
|----------|------|------|---------|--------|---------|--------|-----|-----------|------------|
| FePt     | 3.88 | 3.73 | 3.86    | 3.71   | 3.87    | 3.73   | 0.96| 0.96      | 0.96       |
| MnPt     | 4.03 | 3.69 | 4.00    | 3.71   | 3.99    | 3.70   | 0.92| 0.92      | 0.93       |
| CrPt3    | 3.91 | 3.91 | 3.87    | 3.92   | 3.92    | 3.92   | 1   | 1         | 1          |

Figure 6. Density of state of FePt binary. The dashed lines represent partial density of states (PDOS) of Fe and Pt elements. The vertical line corresponds to the Fermi energy.

Table 1. Equilibrium lattice constants of L10 (CuAu) FePt and MnPt, as well as L12 (CuAu3) CrPt3 ordered alloys.

a Ref. [27].
b Ref. [62].
c Ref. [3].
d Ref. [24].
e Ref. [25].
f Ref. [63].

distinctive characteristic of spin up is that it has an extremely deep TDOS minimum. The d states of Pt exhibit lower energy than d states of Fe. This could be interpreted in terms of the difference in number of electrons per atom (2 electrons/atom). The Pt, Mn and Fe atoms exhibit magnetic moments of zero, 4 and ~3, respectively. The d states of Mn and Fe atoms are found to mainly exhibit degenerate local majority states. Furthermore, the AFM of Mn is a result of local minority spin channel of Mn d states that have no adjacent similar state of the same energy for hybridization to take place. The energy interfaces between the Mn-local minority d states the adjacent states of same energy in the same spin channel are positive. Consequently, $E_{\text{Mn}_1}^d$ d states are relatively higher in energy.

As demonstrated by Figures 18 and 19, the contribution of s and p electrons to the density of states of Cr and Pt is negligibly small. Therefore, the electrons of Cr-3d and Pt-5d are mainly responsible for the structural properties of the CrPt$_3$ compound. A careful inspection of TDOS of CrPt$_3$ reveals a large peak in the DOS of minority spin state that formed from the Cr-3d states above Fermi level. This minority peak originates mainly from Cr-3d bands with much smaller contributions from Pt-5d derived states as illustrated in Figures 17 and 19. The major peaks of this compound come mostly from the electronic states of 5d-Pt atoms. We found that the width of the majority band ~7 eV is narrower than the corresponding energy width of the minority band ~ 9eV indicating that the DOS is larger in the majority bands than in the minority bands. Analogous to the discussion of the band width of DOS of FePt and MnPt, the orbitals are strongly localized and the effective mass is higher leading to a large number of states over a small energy range.

Figure 7. Density of state of FePt binary. The dashed lines represent partial density of states (PDOS) of Fe. The vertical line corresponds to the Fermi energy.

Figure 8. Density of state of FePt binary. The dashed lines represent partial density of states (PDOS) of Pt. The vertical line corresponds to the Fermi energy.
As shown in Figure 18, the 3d states of the Cr atoms located near the top of the Pt-5d band form relatively narrow 3d bands. The hybridization of Cr-3d and Pt-5d states can be interpreted quantitatively by calculating the PDOS. By inspecting Figure 19 near the Fermi energy, we notice that Pt-d densities follow distinctly the Cr-d densities, a clear sign of strong hybridization (see Figure 20).

The electronic band structure of CrPt$_3$ system is displayed in Figure 21 demonstrating that the CrPt$_3$ system has zero bandgap. As
shown in Figures 22 and 23, the Pt 5d states lie below the 3d states of the Cr atoms on the lower energy side. The Cr-3d bands with spin-up and spin-down shift to the lower- and higher energy sides, respectively. The hybridization between the Cr-3d and Pt-5d states with spin-up becomes stronger than that with down spin. This refers to the number of d holes on Pt in the spin-up bands that are greater than in the spin-down bands.

Having interpreted the TDOS and PDOS of the three ordered systems, we then focus our attention on the interplay between electronic properties and magnetic properties in the following section of the paper.

3.3. Magnetocrystalline anisotropy energy

In order to study the MAE of both L1_0 of FePt and MnPt, we include spin-orbit coupling. For both binaries, our calculations predict that, from energy considerations, the magnetic moments of MnPt and FePt ordered alloys oriented preferentially in the 001 plane. Magnetic moment orientation shifts from a perpendicular plane to another parallel plane have been reported for specific composition range of L1_0 stoichiometry. Such experimental abrupt response of MAE to slight variations of composition above 1:1 equiatomic composition is the main reason of the small MAE value of MnPt ordered binary found to be 0.46 meV/f.u. for
MnPt system. Our results indicate that MAE is very sensitive to the details of first principles calculations. For instance, MAE can be greatly altered for small variations of lattice constants and axial ratio in the supercell. We employed accurately calculated lattice parameters for both systems to ensure a high degree of accuracy in calculating MAE values.

In crystals, physical properties are generally anisotropic as they exhibit different values and behaviors along different crystallographic directions. The magnetic moment in a single crystal is principally aligned toward the easy direction. The easy and hard directions arise from the interaction of the spin magnetic moment with the crystal lattice (spin–orbit coupling). Accurate calculation of magnetic anisotropy in crystals is very important as its value is considered to be vital in current and future technological applications of spintronic and magneto-optical devices. In this study, we report MAE values of FePt, MnPt and CrPt$_3$ binaries calculated as described in the computational details section. Our results agree well with the previous experimental and theoretical studies within numerical precision.

As indicated by Table 2, the L1$_0$ phase of FePt exhibits larger MAE and K values than the corresponding values of L1$_0$ MnPt. However, the total magnetic moment of MnPt is larger than that of L1$_0$ phase of FePt.

Figure 13. Density of state of MnPt binary. The dashed lines represent partial density of states (PDOS) of Mn. The vertical line corresponds to the Fermi energy.

Figure 14. Density of state of MnPt. The dashed lines represent partial density of states (PDOS) of Pt. The vertical line corresponds to the Fermi energy.
Furthermore, we found that CrPt3 exhibit an abnormally large MAE and K values for a typical compound adopts a cubic phase. It also exhibits a total magnetic moment of 2.51 $\mu_B$. The smaller values of MAE and K in MnPt and CrPt3 binaries compared to L10 FePt binary are due to a relatively weaker exchange splitting. In general, total magnetic moment in the investigated binary systems depends on the filling of electrons in 3d orbital of (Fe, Mn, and Cr) atoms \{Fe \((3d^7 \ 4s^1)\), Mn \((3d^6 \ 4s^2)\) and Cr \((3d^5 \ 4s^1)\)\].

Figure 15. Electronic band structure of the MnPt system. Band gap is missing, and bandgap energy is identically zero.

Figure 16. Band structure and spin up PDOS of MnPt binaries.
We used LSDA $+$ U method to calculate the spin ($M_s$), orbital ($M_l$) magnetic moments of 3d and 5d states, as well as, total magnetic moment ($M_{tot}$) per formula unit (µB) of FePt, MnPt and CrPt$_3$ ordered binary alloys. The obtained results are presented in Table 3. The calculated MAE (meV/f.u.) values of ordered L1$_0$ FePt, L1$_0$ MnPt and L1$_2$ CrPt$_3$ are found to be 2.66, 0.46 and 0.42 respectively. Our results of MAE of the three investigated ordered alloys are found to be in good agreement with previous works [2, 32, 48]. The origin of the abnormally large MAE perceived for crystalline CrPt$_3$ is due to the large orbital moment of 0.149 µB created on Cr parallel to the spin moment of 2.563 µB. This abnormal property arises from strong Cr 3d–Pt 5d hybridization as can be clearly seen from the DOS and PDOS plots (Figures 22 and 23). This unusually strong hybridization leads to a relatively large computed MAE of 0.48 meV/f.u. that is reasonably larger than the room-temperature experimental value of 0.3 meV/f.u. In addition, we estimated the Curie temperatures of the three ordered alloys from the exchange energy data extracted. The high $T_c$ of L1$_0$ FePt, MnPt and L1$_2$ CrPt$_3$ enforces severe requirements on the media and optical system for Heat-Assisted Magnetic Recording [67, 68, 69].

From the exchange integral of the short range ordered neighboring atoms, we could estimate Curie temperature ($T_c$) of FePt, MnPt, and CrPt$_3$ alloys. The exchange integral $J$ and $T_c$ can be related to each other using Heisenberg model and the mean field model [70, 71],

$$T_c = \frac{2}{3k_B} \left( s + 1 \right) \left( z_1 J_1 + z_2 J_2 \right)$$

where $k_B$ is the Boltzmann constant, $J_1$, $J_2$ are the first, second nearest neighbor exchange constants and $z_1$, $z_2$ are the number of the first and second nearest neighbor atoms, respectively.

Figure 17. Band structure and spin down PDOS of MnPt binaries.

Figure 18. Density of state of CrPt$_3$. The dashed lines represent partial density of states (PDOS) of CrPt$_3$. The vertical dashed line corresponds to the Fermi energy.
As listed in Table 4, the calculated values of $T_C$ are in good agreement with previous theoretical values for L1$_0$-FePt, L1$_0$-MnPt and L1$_2$-CrPt$_3$ ordered binary alloys. However, we found a considerable discrepancy between the reported theoretical and experimental values of $T_C$. Therefore, it would be very interesting to compare the different theoretical and experimental results of $T_C$ of the three investigated ordered alloys presented in Table 4. The differences between theoretical and experimental $T_C$ values are still significant and they may be attributed to thermal spin fluctuations, since there is an experimental evidence of such spin fluctuations. Current development in the storage density of magnetic recording needs media containing extremely small magnetic grains featured by thermally stable magnetization with extremely high anisotropies. The Heat-Assisted Magnetic Recording (HAMR) technology has attracted the attention of researchers investigating optimum thermal conditions for high efficient FePt magnetic grains in scaled magnetic recording. (HAMR) is a swiftly emerging technology, aimed to

![Figure 19. Density of state of CrPt$_3$ binary. The dashed lines represent partial density of states (PDOS) of Cr. The vertical line corresponds to the Fermi energy.](image)

![Figure 20. Density of state of CrPt$_3$ binary. The dashed lines represent partial density of states (PDOS) of Pt. The vertical line corresponds to the Fermi energy.](image)
revolutionize magnetic recording. The high Curie temperatures $T_c$ obtained for the three investigated magnetic systems make them appropriate potential candidates to serve as write assist in HAMR magnetic media schemes. Furthermore, we investigate the behavior of magnetization of FePt ordered alloys in the vicinity of $T_c$. Due to the distinct features of L10 FePt phase, such as its low bulk $T_c$ and large anisotropy constant $K$ presently regarded as the best contender for HAMR magnetic media. Actual application of HAMR involves understanding the high-temperature phase transition behavior of FePt, including critical exponents ($\beta$) and $T_c$ diffusions as the self-motivated HAMR media scheme entails. The L10 FePt phase has been reported to exhibit huge uniaxial anisotropy. This is a critical feature mandatory to boost thermal stability of FePt magnetic grains. Throughout the operation of HAMR unit based on FePt, the high anisotropy of FePt is overwhelmed during magnetic recording by energetic laser that causes the heating of the magnetic grains close to their $T_c$. A deeper understanding of the dependence of $T_c$ on the size of FePt magnetic grains in granulated films is crucial for evolving HAMR technology. $\beta$ describes quantitatively the divergent performance of thermal behavior in L10 FePt near $T_c$. It describes the behavior of physical quantities near $T_c$. It is used in a diversity of applications of high-temperature magnetism.

In the vicinity of $T_c$ and near the Curie temperature, magnetization can be expressed as,

$$M(T) \propto (T_c - T)\beta.$$  \hfill (1)

Eq. (1) demonstrates the temperature dependence of magnetization very close to $T_c$, where $\beta$ stands for the critical exponent that depends on the type of the magnetic interaction. We consider the finite-size scaling analysis adopted in [73, 74, 75] to calculate $\beta$ of L10 FePt phase. To investigate the phase transition behavior of FePt, we consider the realistic effective spin Hamiltonian,

$$H = -\sum_{ij} J_{ij} S_i \cdot S_j + d^{(3)}_{ij} S_i^z S_j^z \cdot \sum_{i} d^{(3)}_{ii} (S_i^z)^3 + \mu_\text{Fe} H S_i^z$$ \hfill (2)

Eq. (2) accounts for all the possible interactions in high uniaxial anisotropic L10 FePt phase. In this equation, $\mu_\text{Fe} = 3.23 \mu_B$ is the magnetic moment of Fe atom. $\mu_\text{B} = 9.274 \times 10^{-24} \text{J/T}$ is the Bohr magneton and $H$ stands for the strength of the magnetic field directed along the easy $z$-axis. The effective exchange interaction $J_{ij}$ includes all possible interactions up to the second nearest-neighbors. To simulate the magnetic field and temperature dependent magnetization data of the model defined by Eq. (2), we use SLDA described in methodology section. We treated Fe magnetic grains as spherical particles with radii $R$ in the range

Figure 21. Electronic band structure of the CrPt$_3$ ordered alloys. Band gap is missing, and bandgap energy is identically zero.

Figure 22. Band structure and spin up PDOS of CrPt$_3$ alloys.
of 1–8 nm. The magnetic field strengths are specifically selected in the range 0–3 meV (0–16 T) and the temperature T ranges from 300 to 1000 K. We calculated $\beta$ to be 0.365 consistent with the value reported by [76, 77] within numerical accuracy.

4. Conclusions

In summary, First-principles structural stability investigations show that both FePt and MnPt exhibit the face centered tetragonal L10 (CuAu) structure labeled as L10 phase, whereas the most stable phase of CrPt3 is the cubic L12 (CuAu3). The interplay among electronic band structure, total spin density of states (TDOS), spin partial density of states (PDOS) and magnetic properties is examined and interpreted. We found that L10...
FePt, L10 MnPt and L12 CrPt exhibit MAE (K) values of 2.66 meV/f.u. (1.7 × 10⁻¹⁰ erg/cm²), 0.46 meV/f.u. (1.3 × 10⁻¹⁰ erg/cm³) and 0.42 meV/f.u. (1.1 × 10⁻¹⁰ erg/cm³), respectively. To clarify the influence of each atom to electronic and magnetic properties, we calculated the (TDOS) and (PDOS) of each alloy. We found that the major contribution to the (TDOS) comes from hybridization of M-3d and Pt-5d states (M = Fe, Mn). The preponderance of this kind of hybridization is predicted to boost both spin and orbital magnetic moments and thus enhances the magnetic properties. In L10 FePt, this hybridization is strong enough to enhance the energy data. The Tc values of L10 FePt, L10 MnPt and L12 CrPt3 are found attributed partially to the overestimation of the inter-sites interactions of each.

References

[1] J.M. MacLaren, R.R. Duplessis, R.A. Stern, S. Willoughby, First principles calculations of FePt, CoPt, Co/substrate 3/3, and Fe/substrate 3/3 alloys, IEEE Trans. Magn. 41 (2005) 4374–4379.
[2] P.M. Oppeneer, I. Galanakis, A. Grechnev, O. Eriksson, Unusual magnetism and magnetocrystalline anisotropy of CrPt3, J. Magn. Magn. Mater. 240 (2002) 377–383.
[3] K.M. Krishnan, C. Nelson, C. Eicher, R. Farrow, R. Marks, A. Kellock, Exchange biasing of permalloy films by Mn on Pt 1–x: role of composition and microstructure, J. Appl. Phys. 83 (1998) 6610–6812.
[4] M. Futamoto, M. Nakamura, M. Okabe, N. Inaba, T. Shimotsu, Growth of L10 ordered crystal in FePt and FePd thin films on MgO (001) substrate, AIP Adv. (2016), 085302.
[5] S.C. Tsang, C.H. Yu, H. Tang, H. He, V. Castelletto, I.W. Hamley, T. Narayanan, C.C. Lo, K. Tam, Assembly of centimeter long silica coated FePt colloidal crystals with tailored interstices by magnetic crystallization, Chem. Mater. 20 (2008) 4554–4556.
[6] R. Hao, R. Xing, Z. Xu, Y. Hou, S. Gao, S. Sun, Synthesis, functionalization, and biomedical applications of multifunctional magnetic nanoparticles, Adv. Mater. 22 (2010) 2729–2742.
[7] A. Alsaad et al. Heliyon 6 (2020) e03545
[8] T. Burkert, O. Eriksson, S.I. Simak, A.V. Ruban, B. Sanyal, L. Nordström, Neutron diffraction study of the magnetic anisotropy of L10 FePt and Fe1−xMnx alloys, Phys. Rev. B 81 (2010), 084705.
[9] A. Sakuma, First-principles study of the magnetocrystalline anisotropy energy of FePt and CoPt ordered alloys, J. Phys. Soc. Jpn. 81 (2012), 084705.
[10] D. Dai, H. Xiang, M.H. Whangbo, Effects of spin-orbit coupling on magnetic properties of discrete and extended magnetic systems, J. Comput. Chem. 29 (2008) 2187–2209.
[11] L. Xiaoyu, P. Sharma, Y. Zhang, A. Makino, H. Kato, Nano-imprinting potential of magnetic FeCo-based metallic glass, Nanotechnology 30 (2019) 305302.
[12] Z. Lu, K.-H. Müller, L. Schultz, O. Isnard, B.C. Sales, B. Saparov, M.A. McGuire, D.J. Singh, D.S. Parker, Ferromagnetism of Fe 3 Sn and alloys, Sci. Rep. 4 (2014) 7024.
[13] P. Bruno, Tight-binding approach to the orbital magnetic moment and magnetocrystalline anisotropy of transition-metal monolayers, Phys. Rev. B 39 (1989) 865.
[14] J.M. Wills, J.M. MacLaren, Magnetic anisotropy in density-functional theory, Phys. Rev. B 38 (1988) 8022.
[15] I. Galanakis, M. Alouani, H. Dreyss, Perpendicular magnetic anisotropy of binary alloys: a total-energy calculation, Phys. Rev. B 62 (2000) 6475.
[16] D. Li, Magnetocrystalline anisotropy of metallic nanotubes: tight-binding and first-principles studies, in: Université Pierre et Marie Curie-Paris VI, 2015.
[17] B.C. Sales, B. Saparov, M.A. McGuire, D.J. Singh, D.S. Parker, Ferromagnetism of Fe 3 Sn and alloys, Sci. Rep. 4 (2014) 7024.
[18] Y. Kota, A. Sakuma, Relationship between magnetocrystalline anisotropy and orbital magnetic moment in L10-type ordered and disordered alloys, J. Phys. Soc. Jpn. 81 (2012), 084705.
[19] A. Sakuma, First-principle calculation of the magnetocrystalline anisotropy energy of FePt and CoPt ordered alloys, J. Phys. Soc. Jpn. 63 (1994) 3053–3058.
[20] P. Bruno, Tight-binding approach to the orbital magnetic moment and magnetocrystalline anisotropy of transition-metal monolayers, Phys. Rev. B 39 (1989) 865.
[21] H. Jansen, Magnetic anisotropy in density-functional theory, Phys. Rev. B 38 (1988) 8022.
[22] I. Galanakis, M. Alouani, H. Dreyss, Perpendicular magnetic anisotropy of binary alloys: a total-energy calculation, Phys. Rev. B 62 (2000) 6475.
[23] D. Li, Magnetocrystalline anisotropy of metallic nanotubes: tight-binding and first-principles studies, in: Université Pierre et Marie Curie-Paris VI, 2015.
[24] B.C. Sales, B. Saparov, M.A. McGuire, D.J. Singh, D.S. Parker, Ferromagnetism of Fe 3 Sn and alloys, Sci. Rep. 4 (2014) 7024.
[25] D. Dai, H. Xiang, M.H. Whangbo, Effects of spin-orbit coupling on magnetic properties of discrete and extended magnetic systems, J. Comput. Chem. 29 (2008) 2187–2209.
[26] Y. Kota, A. Sakuma, Relationship between magnetocrystalline anisotropy and orbital magnetic moment in L10-type ordered and disordered alloys, J. Phys. Soc. Jpn. 81 (2012), 084705.
[27] A. Sakuma, First-principle calculation of the magnetocrystalline anisotropy energy of FePt and CoPt ordered alloys, J. Phys. Soc. Jpn. 63 (1994) 3053–3058.
[28] P. Bruno, Tight-binding approach to the orbital magnetic moment and magnetocrystalline anisotropy of transition-metal monolayers, Phys. Rev. B 39 (1989) 865.
[29] H. Jansen, Magnetic anisotropy in density-functional theory, Phys. Rev. B 38 (1988) 8022.
[30] I. Galanakis, M. Alouani, H. Dreyss, Perpendicular magnetic anisotropy of binary alloys: a total-energy calculation, Phys. Rev. B 62 (2000) 6475.
[31] D. Li, Magnetocrystalline anisotropy of metallic nanotubes: tight-binding and first-principles studies, in: Université Pierre et Marie Curie-Paris VI, 2015.
