A novel theoretical study of elastic and electronic properties of Os$_2$YAl, (Y=Sc, Ti, V) Heusler Alloys

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In this study, we have investigated the structural, electronic, and elastic properties of a new series of Os$_2$YAl, (Y=Sc, Ti, V) alloys called “Full Heusler”, based on the Wien2k code using the functional density theory (DFT). The exchange and correlation energy are evaluated as part of the LDA approximation. The results showed that Os$_2$VAI was more stable and harder than Os$_2$ScAl, and Os$_2$TiAl. The electronic band structures and density of states (DOS) of the compounds indicate that they are metallic because there is no bandgap in these three materials these results have been shown by three approaches (LDA, TB-mBJ, and SOC). Near the Fermi level, the energy is mainly occupied by the Os-5d and Sc, Ti, V-3d electrons. According to the results of the second-order elastic constants, these compounds met Born’s criteria for mechanical stability. The elastic properties indicate that our compounds are ductile, anisotropic, and rigid. All the calculations and the data were compared with the results obtained with different methods in terms of its mechanical and electronic behavior, Os$_2$VAl was found to have better physical properties than Os$_2$ScAl, and Os$_2$TiAl.

Keywords: Full-Heusler alloy; metallic; Ab-Initio calculations.

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1. Introduction

Heusler alloys were discovered in 1903 by Fritz Heusler, they are gaining more and more attention from researchers [1]. Full and Half-Heusler alloys have a broad family of multifunctional materials for spintronic compound [2-8], shape memory alloys [9], superconducting ground state [10], and thermoelectric materials [11]. Heusler alloys have SlaterPauling behavior [12], higher curie temperature [13-15], Heusler’s alloys are important materials in terms of their interesting properties such as electronic localisation, itinerant magnetism, antiferromagnetism, helimagnetism, Pauli paramagnetism or the behavior of heavy fermions [16]. Many researchers have discovered some full-Heusler alloys are not half-metals, but they present a metallic character because of the metallic nature of their spin-polarized electronic structures [17-20]. Among full-Heusler alloys, the Rh$_2$CrGe [17] and Rh$_2$MnTi alloys [18] which showed the metallic character, due to overlap between valence bands and conduction bands at the Fermi level with the two spins channels. While other researchers have shown that the metallic character of certain compounds products from the contribution of Rh, and (Cr,Mn)-3d states around the Fermi level [17,18]. The regular Heusler compounds X$_2$YZ, with 2:1:1 stoichiometry, crystallize in the cubic L$_{21}$ structure (Fm3m, space group 225) with Cu$_2$MnAl as the prototype [21,22]. Generally, the X and Y are transition metals and Z atoms are from III, IV, and V group elements [23]. If the atomic number Z(Y) > Z(X), then the inverse Heusler structure XXYZ (prototype Cu$_2$HgTi, F43m, space group 216) is formed [21], or in the particular case when the Y element is more electronegative than X, the crystal structure observed for full-Heusler compounds, has Cu$_2$HgTi prototype and F43m space group [22]. The discovery of the giant magnetoresistance effect (GMR) in magnetic multilayer has revolutionized the field of information technology. Today, we are in contact with spintronics in our daily lives. Giant magnetoresistance (GMR) are used in magnetic hard drives, position detection on pneumatic cylinders, Position detection in robotics, speed measurement and position of ball bearings, wheel speed measurement, detection of banknotes by measurement of the magnetic ink, detection of electrical short circuits (e.g. in batteries), vehicle detection (road traffic count), and the Earth’s magnetic field detection. The objective of this study is to predict the structural parameters, electronic structure, magnetic properties, and elastic properties of Os$_2$YAl, (Y=Sc, Ti, V) Heusler alloys by using the first-principles calculations of the full-potential (linearized) augmented plane wave (FP-(L)APW) method. This document is organized as follows: computational details are described in Sec. 2, the results are discussed in Sec. 3, and finally, the conclusions are presented in Sec. 4.

2. Calculation details

Electronic structure calculations of our full-Heusler compounds were performed using the first-principle calculations of density functional theory [24,25] based on the FP(L)APW method is implemented in the Wien2k code [26-29]. For
exchange-correlation potential, we used the local density approximation (LDA) proposed by Perdew and Wang [30], and the GGA-PBE approximation proposed by J.P. Perdew, K. Burke, and M. Ernzerhof [31]. It calculates the self-consistent solution of the equations of Kohn and Sham [25]. In these calculations, we have chosen the values of the radii of the atoms of Os, Ti, Sc, and Al of 2.2, 2, 2, 2.1, and 2 u.a., respectively, so that there is no overlap Muffin-Tin spheres. The electronic configurations of the sets of the system studied are: Os (6s²4f⁴5d⁹), Ti (3d² 4s²), Sc (3d¹ 4s²), V (3d⁴ 4s²), and Al (3s² 3p¹). We have used a 1500K-point Monkhorst-Pack mesh [32,33] in the Brillouin zone for all compounds. We chose the $R_{MT} \times K_{\text{max}} = 7$ (where $R_{MT}$ is the mean radius of muffin-tin spheres). The energy cutoff was chosen as $-6$ Ry, used for separation between the valence and the core states. The basic functions and potentials are extended in combinations of spherical harmonics around atomic sites, that is to say, the atomic spheres with a cutoff of $l_{\text{max}} = 10$, and in Fourier series in the interstitial. The self-consistent convergence of the total energy was set at 0.1 mRy.

### 3. Results and discussion

#### 3.1. Structural properties

The compounds Os₂YAl, (Y=Sc, Ti, and V), as most “Full Heusler” alloys crystallize in type structure “regular” (Cu₂MnAl, L₂₁ prototype) with space group Fm³m N°225, the atoms are located at the Wyckoff coordinates: with Os atoms are positioned at 8c (0.25,0.25,0.25), Y=Sc, Ti, V atoms at 4b (0.5,0.5,0.5), and Al 4a (0,0,0) [34]. The CrySDen [35] package has been used to plot the crystal structure of Os₂YAl, (Y=Ti, Sc, V), as shown in Fig. 1. We used Michael Gillisen’s parameter with a magnetic moment equal to zero [36], and the data available [34,36,37] so we performed detailed structural optimizations by minimizing all energies. Figure 2 presents the total energy as a function of the volume for the three compounds. We have plotted the evolution of the total energy as a function of the volume according to the Murnaghan equation [38] given by the following relation,

\[
E = E_0V + \frac{BV}{B(B-1)} \left( B \left( 1 - \frac{V_0}{V} \right) + \left( \frac{V_0}{V} \right)^B \right),
\]

where $E_0$ is the minimum energy at $T = 0$ K, B is the bulk modulus, B' is the bulk modulus derivative and $V_0$ is the equilibrium volume, all the points are effectuated by using the LDA approximation [30]. The optimized lattice parameter is almost the same as what has already been reported [34,36,37]. The results concerning the optimization of three materials Os₂YAl, (Y=Sc, Ti, V), (lattice parameters, the compressibility modulus, and their derivatives, and the equilibrium energies, optimized with the LDA and PBE), are grouped in Table I. The calculated lattice equilibrium constant values ($a_0$) of the Os₂YAl compounds (y = Sc, Ti, and V) are compared to the lattice parameters available in the literature which are very close to each other [34,36,37]. The lattice constants for Os₂ScAl, Os₂TiAl, and Os₂VAl differs by a percentage of approximately 1.83%, 1.8%, and 1.53%, respectively using LDA approximation, for PBE approach the results are improved. No experimental data were found for the lattice parameters of the compounds Os₂TiAl and Os₂VAl for comparison. The mass modulus calculated for the material Os₂ScAl is closer to the available results [34]. For the mass modulus of the compounds Os₂TiAl and Os₂VAl, no experimental value was found for comparison.

We can say that the Os₂VAl compound has a higher negative energy and harder than Os₂ScAl, and Os₂TiAl compounds due to the large value of the compressibility module. We note that when we go to an increased Sc-Ti-V and fixing X = Os and Z = Al, the mass modulus of $X_2YZ$ increases.

#### 3.2. Formation and cohesive energy

To confirm the structural stability we calculated the cohesion energy and the energies of the individual atoms by increasing...
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Table I. Calculated equilibrium lattice constants $a$ (Å), bulk modulus $B_0$ (GPa), and its first derivative $B'$, Equilibrium Energy, and the valence electron concentration (val-el), of Os$_2$YA$_L$ (Y=Sc, Ti and V)

<table>
<thead>
<tr>
<th></th>
<th>$a$ (Å)</th>
<th>$B_0$ (GPa)</th>
<th>$B'$</th>
<th>$E_{Ch}$ (eV/atom)</th>
<th>$E_{Fermi}$</th>
<th>$E$ (Ry)</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Os$_2$ScAl</td>
<td>6.13</td>
<td>222.57</td>
<td>4.31</td>
<td>-2.605</td>
<td>0.88</td>
<td>-71097.499</td>
<td>LDA</td>
</tr>
<tr>
<td>Previous [36]</td>
<td>6.244</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6.21</td>
<td>193.31</td>
<td>4.44</td>
<td></td>
<td></td>
<td></td>
<td>PBE</td>
</tr>
<tr>
<td>Os$_2$TiAl</td>
<td>6.01</td>
<td>273</td>
<td>4.24</td>
<td>-2.614</td>
<td>0.92</td>
<td>-71276.567</td>
<td>LDA</td>
</tr>
<tr>
<td>Previous [36]</td>
<td>6.12</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Os$_2$VAl</td>
<td>5.94</td>
<td>307.4</td>
<td>4.23</td>
<td>-2.931</td>
<td>0.99</td>
<td>-71467.353</td>
<td>LDA</td>
</tr>
<tr>
<td>Previous [36]</td>
<td>6.032</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The unit cell of a face-centered cubic structure [39] up to 30 Bohr (about 16 Å) for the three compounds. The cohesion energy $E_{coh}^{Os_2YA_L}$ of Os$_2$YA$_L$ (Y=Si, Ti, V) is known as the overall energy of the constituent atoms minus the total energy of the compound is given by [39,40]

$$E_{coh}^{Os_2YA_L} = E_{Tot}^{Os_2YA_L} - \left[ xE_{iso}^{Os} + yE_{iso}^{Ti} + zE_{iso}^{Al} \right] / (x + y + z), \quad (2)$$

Y = Sc, Ti, and V with $E_{iso}^{Os}$, $E_{iso}^{Ti}$, $E_{iso}^{Al}$, refer to the energies of the isolated atoms, and $E_{Tot}^{Os_2TiAl}$ is the total energy of the unit cell used in the present calculation, $x$, $y$, and $z$ are the numbers of atoms Os, Ti, Al in the unit cell respectively.

The cohesive energies are also indicated in Table I. It is found that the cohesion energy of the Os$_2$ScAl, Os$_2$TiAl and Os$_2$VAl compounds are -2.605 eV / atom, -2.614 eV/atom, and -2.931 eV/atom, respectively. From these results, we can say that the compound Os$_2$VA has high stability compared to the compounds Os$_2$ScAl and Os$_2$TiAl, and the Os$_2$TiAl compound is more stable than the Os$_2$ScAl compound. These results are comparable to the results of the optimization part.

### 3.3. Electronic properties

#### 3.3.1. Band structures

Figures 3, 4 and 5, show the band structure of the studied systems Os$_2$YA$_L$ (Y=Sc, Ti, V) with LDA, TB-mBJ [41-44], and LDA+SOC, respectively, calculated at their equilibrium lattice constants at different high points of symmetry in the Brillouin zone. The valence bands pass through the Fermi level and enter the conduction band for all structures, but do not overlap except Os$_2$VAl for the three approximations, and the absence of a forbidden band which clearly indicates the metallic character. The dispersed upper bands are due to the strong hybridization of the (d) states of (Os, and Ti, Sc, and V). These results are similar to other results of the same family [34].

#### 3.3.2. Densities of states

The projected total and partial state densities (DOST) (DOSP), between -18.5 and 9 are illustrated, respectively, in Figs. 4a), b), c), and the Fermi level is taken as the origin of the energies. The analysis of the figures of the total and partial state density of the Os$_2$ScAl, Os$_2$TiAl, and Os$_2$VAl indicates a non-zero density at the Fermi level and the ab-

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ence of the forbidden band $E_g$ which makes it possible to deduce that these materials have a metallic nature, (since the DOS has a great value at the Fermi level, see Table I). At the Fermi level, the DOS is 47.6, 34, and 13.6 states per cell unit per eV, respectively for Os$_2$ScAl, Os$_2$TiAl, and Os$_2$VAl. Therefore, there is a downward disposition concluded which makes Os$_2$ScAl more conductive than Os$_2$TiAl and Os$_2$VAl. We find that the TDOS around the Fermi level come mainly from electrons (Ti-d), and (Os-d) the states (p) of the (Al) elements occupy the lowest part of the valence states and have a small contribution around the Fermi level. In addition, there are two atoms of (Os), and only one atom of (Sc, Ti, and V) but the contribution of the Sc, Ti and V atoms is less efficient compared to the Os atom. The hybridization between the states Os (d), Y (d), and Al (p) becomes stronger when the lattice parameter increases with the atomic number of the elements (Y = Sc, Ti, and V). We can also see that the partial DOS of Y=Sc, Ti, and V (3d) orbitals in three materials exhibit the same behavior.

### 3.4. Elastic properties and mechanical stability

Furthermore, we have also considered to mechanical properties of the Os$_2$YAl (Y=Sc, Ti, and V) full-Heusler alloy. The elastic properties describe the mechanical behavior of materials, their study is also important for the field of engineering. Knowledge of the elastic properties of materials is important for fundamental research, particularly for understanding the mechanisms of the bonds between atoms. The cubic structure materials have three independent elastic constants: $C_{11}$, $C_{12}$, and $C_{44}$. To obtain the elastic constants for these compounds, we used a first-principles numerical calculation using the method developed by Reshak and Morteza and integrated into the WIEN2k code [45]. From Table II, one can say that the Os$_2$TiAl, Os$_2$ScAl, and Os$_2$VAl compounds are mechanically stable because all these elastic constants are positive and meet the criterion of mechanical stability verify the relation in Eq. (3) [46-49].

<table>
<thead>
<tr>
<th>Material</th>
<th>$C_{11}$ (GPa)</th>
<th>$C_{12}$ (GPa)</th>
<th>$C_{44}$ (GPa)</th>
<th>A</th>
<th>$C_\rho = C_{12} - C_{44}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Os$_2$ScAl</td>
<td>308.73</td>
<td>181.67</td>
<td>98.73</td>
<td>1.55</td>
<td>82.94</td>
</tr>
<tr>
<td>Previous [34]</td>
<td>261.908</td>
<td>160.436</td>
<td>102.312</td>
<td>2.016</td>
<td>58.124</td>
</tr>
<tr>
<td>Os$_2$TiAl</td>
<td>401.16</td>
<td>205.15</td>
<td>158.1</td>
<td>1.61</td>
<td>47.05</td>
</tr>
<tr>
<td>Os$_2$VAl</td>
<td>516.9</td>
<td>202.98</td>
<td>163.6</td>
<td>1.04</td>
<td>39.38</td>
</tr>
</tbody>
</table>

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**Figure 4.** Band Structure of Os$_2$YAl (Y=Sc, Ti, and V) with TB-mBJ approximation.

**Figure 5.** Band Structure of Os$_2$YAl (Y=Sc, Ti, and V) with LDA+SOC approximation.
TABLE III. The bulk modulus $B$, shear modulus $G$, Young’s modulus $E$, Poisson’s ratio $\nu$, and the bulk-modulus-to-shear-modulus ratio, for the Os$_2$YAl (Y=Sc, Ti and V).

<table>
<thead>
<tr>
<th></th>
<th>$E$ (GPa)</th>
<th>$B$ (GPa)</th>
<th>$G$ (GPa)</th>
<th>$\nu$</th>
<th>$B/G$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Os$_2$ScAl</td>
<td>220.99</td>
<td>224.02</td>
<td>82.735</td>
<td>0.32</td>
<td>2.70</td>
</tr>
<tr>
<td>Previous [34]</td>
<td>214.922</td>
<td>194.260</td>
<td>81.681</td>
<td>0.31</td>
<td>2.398</td>
</tr>
<tr>
<td>Os$_2$TiAl</td>
<td>337.280</td>
<td>270.48</td>
<td>130.51</td>
<td>0.29</td>
<td>2.07</td>
</tr>
<tr>
<td>Os$_2$VAl</td>
<td>411.04</td>
<td>307.61</td>
<td>160.9</td>
<td>0.27</td>
<td>1.90</td>
</tr>
</tbody>
</table>

FIGURE 6. Total a) and partial density of states of Os$_2$ScAl b), Os$_2$TiAl c), Os$_2$VAl d). The Fermi level is set to zero energy and marked by a vertical dashed line.
Our results show that the value of the $C_{11}$ constant is larger than the other constants for three compounds. So, we can say that the length change resistance is the most important in these compounds. The $C_{ij}$ modules have a heavy weight in the study of materials especially the $C_{44}$ module, from this module, can determine several properties such as fragility, among others. The $C_{44}$ module shows that Os$_2$YAl is harder than Os$_2$TiAl, and Os$_2$ScAl; this result is similar to the one found in the optimization part. The elastic constants obtained for the compound Os$_2$ScAl are in good agreement with the available results, whereas the values of $C_{11}$ are greater than those reported in Ref. [34]. The value of $C_{44}$ was lower than the search result [34]. For the compounds Os$_2$TiAl, and Os$_2$ScAl, no results were found for comparison. By comparing the $C_{11}$ values of the compounds studied, we verified that its values are of the increasing order according to the increase in the atomic number of the element Y = Sc, Ti, and V. According to Pettifor [50,51], the character of atomic bonding in metals and compounds, also relates to the brittle or ductile characteristics can be found by considering the Cauchy pressure (CP). A negative value of CP shows a strong covalent bond while a positive value of CP indicates a strong metallic bond. The Cauchy pressure values are 82.94 GPa, 47.05 GPa, and 39.38 GPa for Os$_2$ScAl, Os$_2$TiAl, and Os$_2$Va1, respectively. The obtained values for the pressure are all positive, showing a metallic character and ductile nature of the compounds. These results confirm the metallic bonding character of the three materials, and the most stable compound was Os$_2$YAl, in agreement with the optimization and cohesive energy calculations, as well as the electronic band structure analysis.

The knowledge of the elastic constants $C_{ij}$ allows us to calculate other mechanical (elastic) quantities such as the modulus of compressibility $B$, the Young and shear modulus ($Y$ and $G$), as well as the Poisson’s ratio ($\nu$) which are important parameters in technological applications and provide a fundamental description of the mechanical behavior of a material. The bulk modulus is used to measure the hardness of materials to volume variation by the applied hydrostatic pressure [47,48], while the shear modulus $G$ represents the resistance to plastic deformation and gives the behavior of a material under a uniform pressure [47-49]. For a cubic crystal the bulk modulus $B$, the Voight-Reuss-Hill averaged shear modulus $G$, Poisson’s ratio $\nu$, and Young’s modulus are expressed as follows [35,36,38,46-48,52-54].

$$B_V = B_R = \frac{(C_{11} + 2C_{12})}{3}, \quad (4)$$

$$G_V = \frac{(C_{11} - C_{12} + 3C_{44})}{5}. \quad (5)$$

The Reuss bounds [46,55] of the bulk and shear modulus are $B_V = B_R$ and

$$G_R = \frac{2C_{11}(C_{12} - 2C_{44})}{[4C_{44} + 3(C_{11} - 2C_{12})]. \quad (6)$$

According to the Hill approximation [46,56] method, arithmetic mean of the Voigt and Reuss’s shear moduli gives the elastic modulus expressed as following:

$$G = \frac{(G_V + G_R)}{2} = G_H, \quad (7)$$

where $G_V$ is the Voigt shear modulus and $G_R$ is the Reuss shear modulus.

The mass modulus ($B$) is greater than the shear modulus ($G$) for all three compounds. Thus, these materials must be resistant to changes in their volumes under uniform pressure. The results of the modulus of mass $B$ obtained as a function of the elastic constants are of the same order as the results obtained by applying the Murnaghan equation [38], displaying that our estimated results of the elastic constants for Os$_2$ScAl, Os$_2$TiAl, and Os$_2$Va1 are exact and precise. The Young’s modulus $E$ and Poisson’s ratio $\nu$ for a cubic structure are related to the modulus of compressibility $B$ and shear $G$. Using the relation [36,46-48],

<table>
<thead>
<tr>
<th></th>
<th>Os$_2$ScAl</th>
<th>Os$_2$TiAl</th>
<th>Os$_2$Va1</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mu_{\text{Os}}$</td>
<td>0.00089</td>
<td>0.00086</td>
<td>0.00026</td>
</tr>
<tr>
<td>$\mu_{\text{Sc}}$</td>
<td>0.00145</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$\mu_{\text{Ti}}$</td>
<td>-</td>
<td>0.00141</td>
<td>-</td>
</tr>
<tr>
<td>$\mu_{\text{V}}$</td>
<td>-</td>
<td>-</td>
<td>0.00005</td>
</tr>
<tr>
<td>$\mu_{\text{Al}}$</td>
<td>0.00053</td>
<td>0.00001</td>
<td>0.00113</td>
</tr>
<tr>
<td>$\mu_{\text{inter}}$</td>
<td>0.00311</td>
<td>0.00060</td>
<td>0.00040</td>
</tr>
<tr>
<td>$\mu_{\text{Total}}$</td>
<td>0.00069, [36]</td>
<td>0.00028, [36]</td>
<td>0.00106, [36]</td>
</tr>
</tbody>
</table>

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The Young’s modulus is defined as the ratio between stress and strain and is used to provide a measure of the stiffness of the solid matter, i.e., the larger the value of $E$, entails a stiffer material [46,57]. We can see from the results mentioned in Table III, that the Young’s modulus value $E$ is in ascending order, and Os$_2$VAI compound is more rigid and harder than Os$_2$ScAl and, Os$_2$TiAl due to its higher value of Young’s modulus, and compressibility modulus ($B$). At the same time, the value of $E$ of compound Os$_2$ScAl is of the same order of results available [34]

With an increase in Poisson’s ratio, the plasticity of the crystal improves. The Poisson’s ratio has been suggested as 0.1 for covalent substances and 0.25 for ionic substances.

\[ E = \frac{9BG}{(3B + G)}, \quad (8) \]

\[ \nu = \frac{(3B - 2G)}{2(3B + G)}. \quad (9) \]

This ratio is associated with ductility (brittleness). The critical value separating the ductile and the brittle turned out to be 1.75. As shown in Table III, this ratio is greater than 1.75 for the three Os$_2$ScAl, Os$_2$TiAl, and Os$_2$VAI compounds, which are classified as ductile compounds, and this result confirmed previous data [34].

4. Conclusion

In summary, we have studied the structural, elastic, and electronic properties of the Heusler Os$_2$YAl$_3$(Y=Sc, Ti, V) alloy, with the space group Fm$ar{3}$m (# 225) in phase L$_{21}$, which were calculated using first-principle calculations of the full-potential linearized augmented plane wave (FP-LAPW) method, with local density approximation. After fitting the Murnaghan equation of state, the equilibrium lattice parameters obtained from our calculation agree well with the theoretical values available. We found that the compound Os$_2$VAI is more stable and harder than Os$_2$ScAl, and Os$_2$TiAl, a result that was confirmed by the calculation of the cohesion energy. The calculated state densities presented in this study identify the metallic behavior of Os$_2$YAl$_3$, (Y=Sc, Ti, and V), by three approaches (LAD, mBJ, SOC). The metal band property of these compounds was confirmed based on the Cauchy pressure values. The elastic constants of these compounds are calculated according to the method developed by Reshak and Morteza and the results obtained show the compounds studied are mechanically stable. The Poisson’s ratio values showed that the compounds had an ionic metallic character. Analysis of Young’s modulus and the $B/G$ ratio shows that the three materials were rigid and ductile.


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