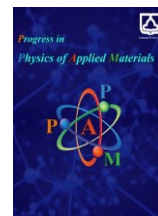




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## Structural, Electronic, and Magnetic Properties of $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$ ( $x=0.0-1.0$ ) Alloys

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### ABSTRACT

The structural, electronic, and magnetic properties of the novel  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  ( $x=0.0-1.0$ ) alloys were surveyed by using the density functional theory. The formation and cohesive energy results confirm that all members of these series are thermodynamically stable, but the  $\text{Hg}_2\text{CuTi}$ -type structure has lower stability compared to the  $\text{Cu}_2\text{MnAl}$ -type structure. The results also reveal that with increment of Si content, the lattice constant decreases linearly from 6.01 to 5.88 Å, while the bulk modulus increases. The total magnetic moment decreases from 2.00  $\mu_B$  (for  $x = 0.0$ ) to 0.99  $\mu_B$  (for  $x = 1.0$ ). The results of electronic structure show that the alloys with  $x = 0.0, 0.25$ , and  $0.50$  have a half-metallic nature with a real gap in the down-spin band, and 100% spin polarization. For other alloys, the spin polarization decreased with increasing  $x$  from 0.75 to 1.0. Although 100% spin polarization was not found for all members of these series, it is quite high value which can be used in industries.

## 1. Introduction

In the last decades, the physical properties of the half-metallic (HM) materials have been investigated because of their diverse applications in the electronic devices [1-8]. In these materials, an energy gap around the Fermi level ( $E_F$ ) is observed for one spin band, while for the other band a metallic nature is exhibited. Therefore, they may have 100% spin-polarization ( $P$ ) [2, 9-12]. This feature makes them excellent candidates for spintronic applications [3, 9, 12, 13]. The spintronic is a swiftly advanced area in the field of modern technology. The main concept of this new technology is to use the spin freedom degree of electrons. Spintronics has the potential advantage of nonvolatility, speeding up of data processing, high circuit integration density, and reduction in power consumption [14]. However, it needs the adequate materials as a source of the spin-polarized charge carriers [15, 16]. For these purposes, the charge carriers injection into a semiconductor, the conventional ferromagnetic metals are not suitable because

of low spin polarization. But with the aid of magnetic semiconductors, a high spin polarization of the injected charge carriers has been shown. On the other hand, these require bias fields because they are not spontaneous magnets which make them unsuitable for a variety of applications. For polarized charge-carrier sources, half-metallic systems such as the Heusler alloys are an obvious choice [1, 13, 14, 16, 17]. These alloys are utilized as the magnetic electrodes for using in the giant magneto-resistance (GMR), magnetic sensors, and tunnel magneto-resistance (TMR) devices [5, 18-20]. Among HMs, the Heusler alloys, with high  $P$  and Curie temperature ( $T_C$ ) are favorable [21-23]. In some of these materials, a new ferromagnetic shape-memory behavior has been found which motivated to explore the new alloys from this category [5, 24]. The Heusler alloys can be manufactured using a variety of methods such as the mechanical alloying, arc melting, induction melting, sputtering, and electrodeposition. However, these methods have several challenges such as the contamination problem, high fabrication cost, chemical composition inhomogeneity,

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time consumption, and susceptibility to oxidation [1, 3, 19]. Accordingly, the theoretical study on Heusler alloys has become widely important due to the difficulties involved in synthesizing and producing these materials. In fact, theoretical investigation is essential for understanding the properties and feasibility of synthesis. The researches emphasize that predicting the physical, magnetic, and electronic properties of Heusler alloys theoretically can lead to the targeted design of compounds that are difficult to synthesize in practice or require the precise control of processing conditions [1, 3, 14, 25].

Among the Heusler alloys, the ferrimagnetic alloys are more favorable than the ferromagnetic ones because of their internal spin compensation and low magnetization, while  $T_c$  remains fairly high [9, 26]. Therefore, the external magnetic fields have less effect on these materials as they can be favorable for spintronic applications [3, 9, 27, 28]. Having high  $T_c$  guarantees the good thermal stability in these materials [9]. In the Heusler alloys,  $T_c$  is much higher than that in other HMs [26]. These alloys are classified as half (XYZ), full ( $X_2YZ$ ), and quaternary ( $XX'YZ$ ), where X, X', and Y are transition metals and Z is a sp element [16, 21, 22]. The  $X_2YZ$  alloys crystallize either in  $AlCu_2Mn$ -type (normal) or in  $CuHg_2Ti$ -type (inverse) structure [29] with atomic positions of  $A=(0,0,0)$ ,  $B=(0.25,0.25,0.25)$ ,  $C=(0.5,0.5,0.5)$ , and  $D=(0.75,0.75,0.75)$  [30]. In the normal structure, the position of X atom is A and C sites, Y atom is B, and Z atom is D. For inverse structure, X atom occupied A and B sites, while Y and Z atoms occupied C and D, respectively [31]. Slater and Pauling [32, 33] have observed that the magnetic moment of these alloys can be determined according to average valence electron number per atom. As will be discussed later, the Slater–Pauling (SP) rule relates the number of valence electrons to the spin magnetic moment [28, 34]. With substitution of different elements, the functional materials with desirable magnetic properties can be achieved. After the Co-based compounds,  $Mn_2YZ$  alloys are the second family that contains many half-metallics (HM). However, it is less studied compared with the first family [35]. The Mn-based alloys have considerable applications in the spintronic, magnetic refrigerators and energy technology [36].

The Mn-based alloys have low magnetic moment which is desirable for using in the spintronic and transfer torque-based devices [27, 36, 37]. The first ferromagnet HM of this family was  $Mn_2VAl$  [38]. Later, other Mn-based alloys such as  $Mn_2VZ$  ( $Z = Sn, Ge, Si, In, Ga, Al$ ) [26],  $Mn_2CrZ$  ( $Z = Sb, Ge, Si, Ga, Al$ ) [39],  $Mn_2CuZ$  ( $Z = Al, Ge$ ) [40, 41],  $Mn_2TiZ$  ( $Z = Sb, As, P, Sn, Ge, Si, In, Ga, Al$ ) [9, 15],  $Mn_2CuSi$  [42], and  $Mn_2ZnZ$  ( $Z = Ge, Si$ ) [18, 43] were investigated. Jum'h et al. [35] also reported the HF behavior in  $Mn_2VAl$  and  $Mn_2CrAl$  Heusler compounds. Based on these results, in the Mn-based Heusler alloys, the half-metallic ferrimagnet behavior can be observed [1, 17, 35].

However, more investigations are done on the 3d transition metals and fewer researches are concentrated on the 4d transition metal [5, 28]. In this regard, Abada et al. [5] predicted of new  $Mn_2$ -based full Heusler alloys  $Mn_2ZrSi$  and  $Mn_2ZrGe$  by first-principle calculations. Their study indicated that these alloys can be used as promising

magnetic materials in spintronic applications. In our previous works [20, 21, 36], we reported the HF characteristics in  $Mn_2ZrGa_{1-x}As_x$  (for  $x=0.5$  and  $0.75$ ) and  $Mn_2ZrGa_{1-x}Ge_x$  (for  $x=0.75$  and  $1.0$ ) systems by first-principle calculations. Based on the results reported in [22, 44, 45], the  $Mn_2RuZ$  ( $Z = Si, Ga, Ge$ ) and  $Mn_2RhZ$  ( $Z = Si, Ge, Sn$ ) alloys can also be potential candidates for the spintronic. Ghosh et al. [46] studied  $Mn_2YAl$  ( $Y=Zr, Nb, Mo$ ) by the Pseudo-potential method and concluded that only the  $Mn_2NbAl$  compound shows HF behavior. However, their research lacked the analysis of the band structure. In another research, Saber et al. [6] concluded that the Nb-d states can create or regulate the half-metallic characteristics in the  $Ru_2NbMn$  Heusler compounds. However, to our best knowledge, there are few papers that investigate the Mn-based full-Heusler compounds which contain Nb atoms. In this regard, Piyasin et al. [47] indicated that the Heusler compounds containing Mn and Nb can be suitable for thermoelectric and spintronic applications. Kervan et al. [48] observed the HM behavior in  $Mn_2NbAl$  alloy by using the first-principles calculations. Sofi et al. [49] studied newly Mn-based Heuslers  $Mn_2NbX$  ( $X=Al, Ga, In$ ) by ab-initio calculations within GGA and mBJ approximations. Their results indicated that the  $Mn_2NbX$  alloys are vital applicants for thermoelectric effects, spintronic, power generation, and green energy sources for future technologies. However, there is a shortcoming in their research, and that is the failure to consider the Hubbard parameter  $U$ . In fact, the transition metal d electrons are generally strongly correlated and cannot be adequately described within the GGA approximation, because of not including the on-site Coulomb interaction [2, 50]. This limitation of standard DFT can be taken care of by introducing the Hubbard parameter  $U$  (DFT+ $U$  scheme). The Hubbard parameter consists of the on-site Coulomb interaction parameter  $U$  and the exchange interaction parameter  $J$  [50]. Accordingly, in the present study, we used the DFT+ $U$  scheme. Moreover, the half-metallic ferrimagnet with low total magnetic moment would reduce the energy losses in spintronic devices because the devices would be less affected by the external magnetic field due to its low magnetic strain [12, 37].

The half-metallic ferromagnet with high total magnetic moment would also hinder the spintronic device performance since the external magnetic field would strongly affect the devices due to its large dipole field [51]. The conducted studies on these alloys indicate the energy gap ( $E_g$ ) usually increases with the lattice volume contraction [11, 34]. The Al substitution by Si in the  $Mn_2NbAl$  is expected to affect the magnetic and HM properties of parent alloy, because of the smaller atomic radius of Si and one more valence electron. Si is a semiconductor and its resistance is much higher than Al. Therefore, its substitution can reduce the Eddy currents and energy losses for use in the electronic devices. With these mindsets in this research, the effect of Si substitutions on the physical properties of  $Mn_2NbAl_{1-x}Si_x$  ( $x=0.0-1.0$ ) are theoretically investigated. In the following, the organization of our work is: Section 2 describes the method of calculations. Section 3 surveys the results and their Interpretation, and Section 4 states a summary of results.

## 2. Computational Method

The first-principles calculations for simulation of the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys were done by the density functional theory (DFT) via the WIEN2k code [52, 53]. Fig.1 illustrates the unit cell of the supercells which were used for the simulation of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys. The muffin-tin (MT) sphere radii for the Mn, Nb, Al, and Si atoms were respectively set to 2.25, 2.20, 2.10, and 2.10 a.u. The valence and core states were separated by cut-off energy of -6.5 Ry. These values were chosen such that, while preventing the overlap between the MT spheres, the charge leakage outside these spheres was minimized. The cut-off parameters  $R_{\text{MT}} \times K_{\text{max}}$ ,  $I_{\text{max}}$ , and  $G_{\text{max}}$  were set to 8, 10 and 14 respectively. Moreover, In the irreducible Brillion zone, 120  $k$ -points were used by the Monkhorst-Pack method [54]. These values were optimized separately using the energy convergence criterion. In this method, a graph of the changes in total energy of the lattice is plotted versus the changes of that quantity until further increases in the quantity result in no significant energy changes. The generalized gradient approximation with considering the Hubbard term (GGA+U) was selected as the exchange-correlation energy [50]. The value of the Hubbard parameter ( $U_{\text{eff}}$ ) of Mn-3d states in  $\text{Mn}_2\text{NbAl}$  alloy was calculated using the method presented by Madsen et al.[55] and was found to be of about 4.12 eV. The same amount was also used for other alloys in this series ( $x=0.25-1.0$ ). The spin-orbit coupling is neglected, but it may have strong influence on the spin polarization when the Z atom is a heavy element such as Sn or Sb [9, 11, 28]. The convergence criterion for the self-consistent cycles was  $10^{-5}$  Ry and  $10^{-4}$  electrons simultaneously.

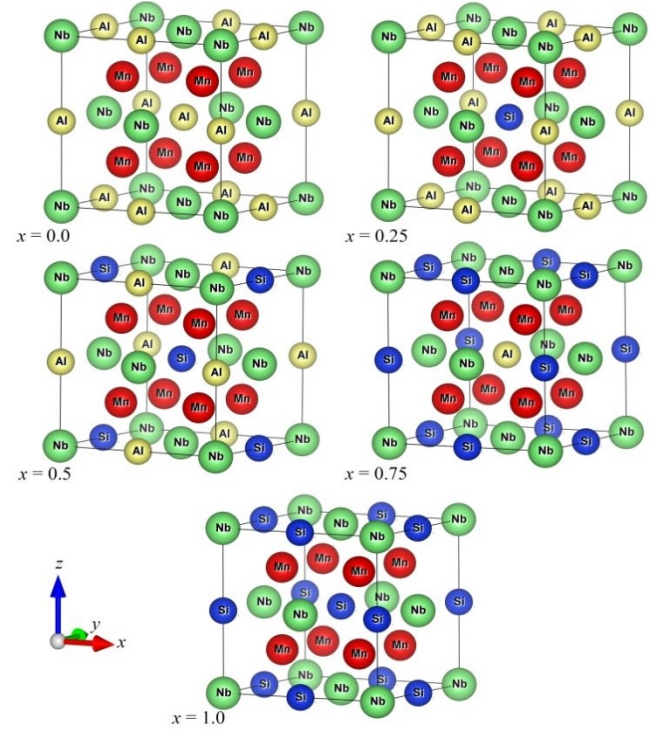
## 3. Results and Discussion

### 3.1. Structural Properties

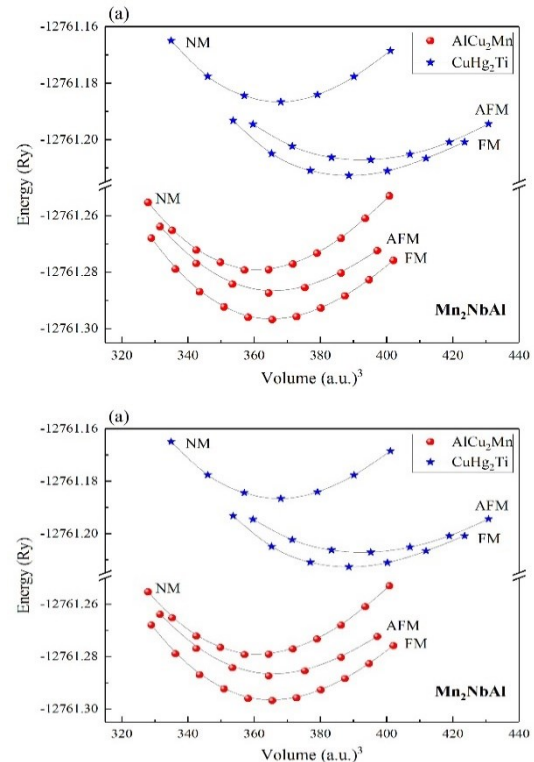
Fig.2 illustrates the calculated total energy  $E$  of the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys with  $x=0.0$  and  $1.0$  versus the cell volume  $V$ . The calculations were done for the anti-ferromagnetic (AFM), ferromagnetic (FM), and non-magnetic (NM) configurations in both type-structures, i.e.  $\text{CuHg}_2\text{Ti}$  (inverse) and  $\text{AlCu}_2\text{Mn}$  (normal). The obtained data then have fitted with Murnaghan equation of state [56] to obtain the equilibrium ground state parameters such as the lattice parameter, volume, bulk modulus and total energy (Table 1). As seen, the normal structure is thermodynamically more stable than the inverse structure.

Moreover, the total energy of FM states is more negative than that of AFM and NM states. Therefore, the FM states of normal structure of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloy can be considered as ground state. A similar trend was also observed for the alloys with  $x = 0.25, 0.5$ , and  $0.75$  which is not presented here. The following calculations for the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys are only done for the FM state of  $\text{AlCu}_2\text{Mn}$ -type structure. Table 1 presents some calculated structural

parameters of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys in FM equilibrium state for  $\text{AlCu}_2\text{Mn}$ -type structure. The accessible measured data by others are also presented for comparison. Moreover, the energy difference between NM and FM configurations ( $\Delta E = E_{\text{FM}} - E_{\text{NM}}$ ) is calculated for the  $\text{AlCu}_2\text{Mn}$ -type structure. As concluded, the calculated lattice constant for  $\text{Mn}_2\text{NbAl}$  ( $x=0.0$ ) is about 6.01 Å, which is in good agreement with the results of others [46, 48, 49].



**Fig. 1.** The unit cells of the calculated supercells for simulation of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys



**Fig. 2.** The total energy variations versus the volume of (a)  $\text{Mn}_2\text{NbAl}$  ( $x=0$ ) and (b)  $\text{Mn}_2\text{NbSi}$  ( $x=1.0$ ) for AFM, FM, and NM states of the  $\text{CuHg}_2\text{Ti}$  and  $\text{AlCu}_2\text{Mn}$ -type structures

The percentage error for the calculation of this parameter relative to the other measured values is 0.17. For other  $x$ -values, no other research results are available. However, by comparing the lattice constant of closely related alloys (Table1), it can be concluded that the obtained values can be reasonable. Moreover, as seen from Fig.3, the lattice constant shows a systematic behavior with changes in  $x$ , as with increment of  $x$ , it decreases linearly in agreement with Vegard’s law. This behavior can be due to the larger atomic radius of Al atom than Si atom as was also observed in our previous work [11] for  $\text{Co}_2\text{TiAl}_{1-x}\text{Si}_x$  Heusler alloys. Since the equilibrium lattice constants of the understudied alloys are close to that of zinc blende semiconductors such as  $\text{InAs}$  (6.058 Å),  $\text{CdSe}$  (6.077 Å), and  $\text{ZnTe}$  (6.088 Å)[5], it is suggested to experimentally realize these Heusler alloys in the form of thin films on appropriate substrates and to use them as new candidates for applications in the spintronic field.

However, the bulk modulus  $B$  increases as  $x$  increases. In fact,  $B$  is a measure of the substance’s resistance to uniform compression and represents the resistance to fracture [35, 57]. This means that the  $\text{Mn}_2\text{NbAl}$  is more compressible compared to  $\text{Mn}_2\text{NbSi}$ . This behavior is in agreement with the well-known relationship between  $B$  and the lattice constant:  $B \propto \frac{1}{V}$  ( $V=a^3$  is the unit-cell volume) as  $V$  decreases with increasing  $x$  [20]. The increase in  $B$  can also be attributed to the increase in strength of the atomic bonds due to the lattice contraction [7, 8].

**Table 1.** Comparison of the structural parameters and some physical findings of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys with the reported values

| Alloy                                               | $a$ (Å)          | $B$ (GPa) | $E_f$ (eV)   | $E_c$ (eV) | $\Delta E$ (eV) | Ref.                   |
|-----------------------------------------------------|------------------|-----------|--------------|------------|-----------------|------------------------|
| $\text{Mn}_2\text{NbAl}$                            | 6.01             | 196       | -1.31        | -19.18     | 0.24            | This work              |
| $\text{Mn}_2\text{NbAl}$                            | 6.00, 6.01, 6.00 | 203, 220  | -1.37, -1.56 | -29.59     |                 | [49], [48], [46]       |
| $\text{Mn}_2\text{NbAl}_{0.75}\text{Si}_{0.25}$     | 5.97             | 202       | -1.39        | -19.53     | 0.17            | This work              |
| $\text{Mn}_2\text{NbAl}_{0.5}\text{Si}_{0.5}$       | 5.94             | 211       | -1.48        | -19.89     | 0.12            | This work              |
| $\text{Mn}_2\text{NbAl}_{0.25}\text{Si}_{0.75}$     | 5.91             | 216       | -1.60        | -20.27     | 0.07            | This work              |
| $\text{Mn}_2\text{NbSi}$                            | 5.88             | 227       | -1.73        | -20.67     | 0.04            | This work              |
| $\text{Mn}_2\text{VAl}$                             | 5.81             | 188       |              | -17.72     |                 | [35]                   |
| $\text{Mn}_2\text{MoAl}$                            | 5.90             |           | -1.23        |            |                 | [46]                   |
| $\text{Mn}_2\text{ZrAl}$                            | 6.15             |           | -2.01        |            |                 | [46]                   |
| $\text{Mn}_2\text{ZrGa}$                            | 6.15             | 150       | -0.92        |            |                 | Our previous work [21] |
| $\text{Mn}_2\text{ZrGe}$                            | 6.10             | 176       | -0.97        |            |                 | Our previous work [36] |
| $\text{Mn}_2\text{ZrAs}$                            | 6.06             | 185       | -2.88        |            |                 | Our previous work [20] |
| $\text{Mn}_2\text{TiAl}$ , $\text{Mn}_2\text{TiSi}$ | 5.96, 5.82*      |           |              |            |                 | [9], [15]              |
| $\text{Mn}_2\text{RuGa}$ , $\text{Mn}_2\text{RuGe}$ | 6.00*, 5.92*     |           |              |            |                 | [44]                   |

\* Denotes the accessible experimental data

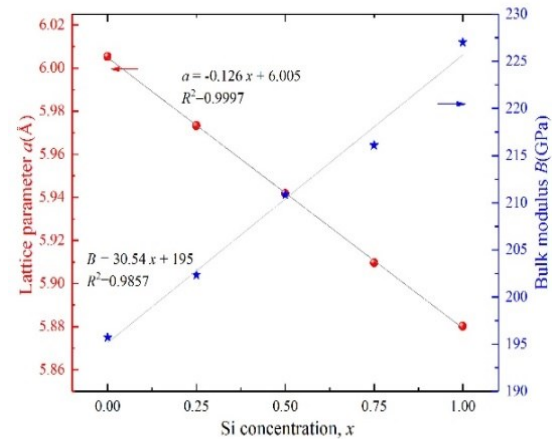
This is consistent with the increase in the magnitude of the calculated cohesive energy as  $x$  increases (Table1). Table1 presents the formation ( $E_f$ ) and cohesive ( $E_c$ ) energies of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys defined by the following equations:

$$E_f = E_{\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x}^{\text{tot}} - [2E_{\text{Mn}}^{\text{bulk}} + E_{\text{Nb}}^{\text{bulk}} + (1-x)E_{\text{Al}}^{\text{bulk}} + xE_{\text{Si}}^{\text{bulk}}] \tag{1}$$

$$E_c = E_{\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x}^{\text{tot}} - [2E_{\text{Mn}}^{\text{iso}} + E_{\text{Nb}}^{\text{iso}} + (1-x)E_{\text{Al}}^{\text{iso}} + xE_{\text{Si}}^{\text{iso}}] \tag{2}$$

where  $E_{\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x}^{\text{tot}}$  is the total energy of the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloy for formula unit,  $E_{\text{Mn}}^{\text{bulk}}$ ,  $E_{\text{Nb}}^{\text{bulk}}$ ,  $E_{\text{Al}}^{\text{bulk}}$ , and  $E_{\text{Si}}^{\text{bulk}}$  are the total energies for one atom of Mn, Nb, Al, and Si respectively. Similarly,  $E_{\text{Mn}}^{\text{iso}}$ ,  $E_{\text{Nb}}^{\text{iso}}$ ,  $E_{\text{Al}}^{\text{iso}}$ , and  $E_{\text{Si}}^{\text{iso}}$  correspond to the energies of the isolated Mn, Nb, Al, and Si atoms. The calculated negative values for  $E_f$  and  $E_c$  indicate that understudy alloys are stable from the thermodynamic point of view. Moreover, with increasing the Si content, the

magnitude of  $E_f$  and  $E_c$  increases which indicates that the Si-rich alloys have more stable structures. As  $x$  increases, the energy difference  $\Delta E$  of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys decreases possibly due to a reduction in the magnetic energy contribution as the total magnetic moment decreases (see section **Error! Reference source not found.**).





**Fig. 3.** The variation of the lattice constant  $a$ , and the bulk modulus  $B$  of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  versus Si concentration

### 3.2. Electronic Properties

The electronic structure is a key factor in predicting the HM nature and their magnetic properties of the materials [11, 43, 58]. Accordingly, the electronic structure calculations can be useful.

Fig.4 shows the total (TDOS) contribution of density of states  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys at their equilibrium structure (AlCu<sub>2</sub>Mn-type) and the partial density of states (PDOS) contribution of their constituent atoms. The zero energy values were devoted to  $E_F$ . As seen, for  $x=0, 0.25$  and  $0.5$  samples, a metallic characteristic is seen for all up-spin bands, while the down-spin bands have a semiconducting gap.

From the contribution of TDOS for the spin-up and spin-down at  $E_F$ , i.e.  $\rho_{\uparrow}(E_F)$  and  $\rho_{\downarrow}(E_F)$ , the spin-polarization  $P$  was calculated from [17]:

$$P = \frac{|\rho_{\uparrow}(E_F) - \rho_{\downarrow}(E_F)|}{\rho_{\uparrow}(E_F) + \rho_{\downarrow}(E_F)} \quad (3)$$

The calculated values for  $P$  are 100% for  $x=0, 0.25$ , and  $0.5$ , and 98% and 95% for  $x=0.75$  and  $1.0$  respectively. These calculated values are high enough to be useful for practical applications of this alloy in spintronic devices [44].

The high spin polarization is one of the main design criteria for materials and equipment in this emerging technology and influences the efficiency, speed, and stability of these devices [59]. In spin sensors or filters with fully spin-polarized current, the output current is only spin-up or spin-down electrons and hence the output signal and switching efficiency are improved. High spin polarization also reduces the probability of spin scattering and reversal during the transmission path, providing greater stability to spin information [59-61]. Accordingly, the samples with  $x=0.0, 0.25$ , and  $0.50$  can be more suitable for practical applications due to 100% spin polarization. For both spin channels, the TDOS around  $E_F$  is mainly originated from the Mn-d and Nb-d states which extend from occupied and unoccupied states. These widely spread d states hybridize with each other as the energy gap around  $E_F$  originates predominantly from this hybridization [43].

As seen, in the energy interval of about  $-6$  to  $0$  eV, d states of Mn and Nb and then the p states of Al and Si have dominant contributions in TDOS. These states are mixed and hybridized with each other, as the strength of the hybridization determines the energy gap width [27]. These mixed states are separated from the next bands which are highly localized within  $-6$  to  $-10$  eV originated from Si-s and Al-s states. In TDOS, the majority and minority spin states are spin-up and spin-down electrons. Therefore, the magnetic moment of these alloys is predicted to be positive (see section **Error! Reference source not found.**).

On the other hand, since each Si atom has one valence electron more than the Al atom, with increasing Si content, the majority states shift slightly towards more negative energies as the accessible states of spin-up band can be occupied by these extra electrons.

To reveal the role of the type of crystal lattice structure, TDOS was also studied for the CuHg<sub>2</sub>Ti-type structure (inverse) of  $\text{Mn}_2\text{NbAl}$  and  $\text{Mn}_2\text{NbSi}$  as presented in Fig. 4f. As seen, for CuHg<sub>2</sub>Ti type-structure, the shape of TDOS is different compared to that of AlCu<sub>2</sub>Mn-type structure. The TDOS for the CuHg<sub>2</sub>Ti-type structure has metallic nature as  $E_F$  crosses the up and down-spin bands. For this case, the spin polarization of about 70% and 13% is calculated for  $\text{Mn}_2\text{NbAl}$  and  $\text{Mn}_2\text{NbSi}$  alloys respectively.

Similar behaviors were also observed for TDOS of CuHg<sub>2</sub>Ti-type structure of the alloys with  $x=0.25, 0.5$ , and  $0.75$ , which are not presented here. Therefore, a change in the atomic positions of the alloy can change its electronic structure.

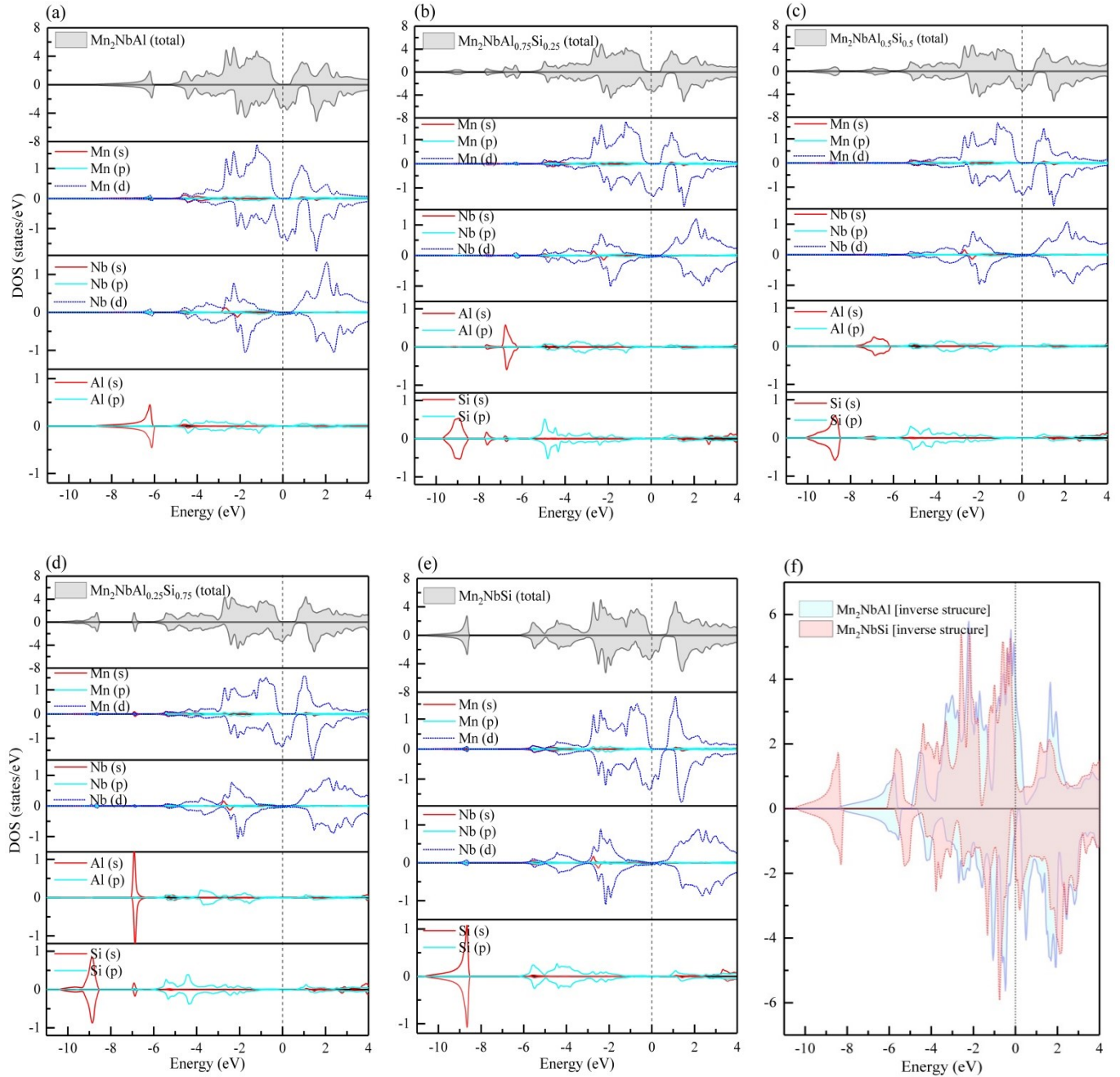
Fig. 5 displays the spin-up energy bands of AlCu<sub>2</sub>Mn-type structure for  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys at equilibrium geometry. As seen, for the samples  $x = 0.75$  and  $1.0$ , the spin-up bands have metallic characteristic, while for  $x = 0.0, 0.25$ , and  $0.5$  a real band gap about  $0.4$  eV is observed around  $E_F$ . This value is close to the reported values for the compounds  $\text{Mn}_2\text{NbAl}$  (GGA= $0.30$  eV, mBJ= $0.46$  eV) and  $\text{Mn}_2\text{VAl}$  (GGA= $0.40$  eV) [48, 49]. By increasing  $x$  from  $0.25$  to  $1.0$ , the highly localized states are observed in the energy range from  $-11$  to  $-8.5$  eV. From the PDOS plots (Fig. 4), one can conclude that these ones are related to Si-s states. These bands are well separated from the next bands that arise mostly from Al-s and Si-s states. As seen, with increasing  $x$ , the width of the bands located in energy range  $-6$  eV to  $E_F$  increases, possibly due to the decrease in lattice parameter and increase in hybridization of p-d orbitals.

The Mn-3d and Nb-3d electrons have dominant contribution for these states. For unoccupied states located above  $E_F$ , the dominant contributions belong to the d states of the Mn and Nb atoms. The spin-down bands for the whole series show the metallic natures which have not been demonstrated here. Moreover, it was found that band structure results of these alloys are well consistent with the calculated TDOS.

### 3.3. Magnetic Properties

As mentioned in section 3.1, the FM states of AlCu<sub>2</sub>Mn-type structure for  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys were more stable than NM ones. Fig. 6 shows the changes in total magnetic moments per formula unit ( $M_{\text{tot}}$ ) of the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys versus the Si concentration. For a better analysis of this behavior, the contribution of the magnetic moment of the constituent atoms is also presented in Table 2. As seen, with the increasing of Si content,  $M_{\text{tot}}$  decreases in agreement with the SP rule [32, 33]. The SP rule states that for a Heusler alloy, the  $M_{\text{tot}}$  depends on the total number of valence electrons ( $N_t$ ):  $M_{\text{tot}} = 24 - N_t$ . Since  $N_t$  is 22 for the  $\text{Mn}_2\text{NbAl}$  alloy ( $N_t = (7 \times 2) + 5 + 3 = 22$ ),  $M_{\text{tot}} = 24 - 22 = 2.0 \mu_B$  is expected by this rule. Similarly,  $M_{\text{tot}}$  of  $1.75, 1.5, 1.25$ ,

and  $1.0 \mu_B$  is expected for the alloys with  $x = 0.25, 0.5, 0.75$ , and  $1.0$ , respectively.



**Fig.4.** The DOS of  $Mn_2NbAl_{1-x}Si_x$  alloys for (a)-(e)  $AlCu_2Mn$ -type structure, and (f)  $CuHg_2Ti$ -type structure of  $Mn_2NbAl$  and  $Mn_2NbSi$ . Positive and negative contributions are devoted to up and down-spin states

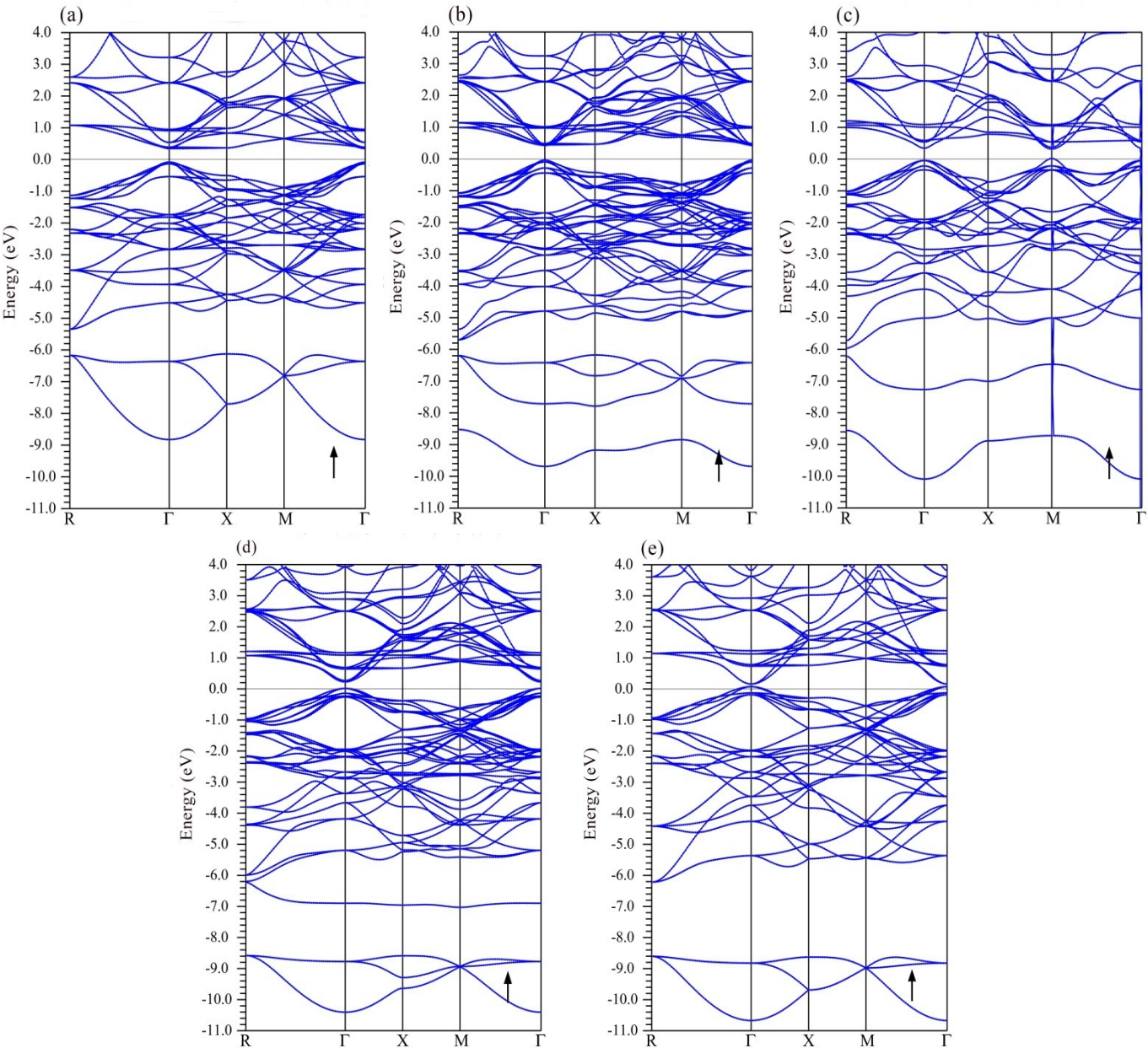
This result indicates that only the magnetic moments of the alloys with  $x = 0.0, 0.25$ , and  $0.50$  obey from SP rule. For  $x = 0.75$  and  $1.0$ , the total magnetic moment deviates from SP rule which arises from the existence of a pseudo-gap instead of a real gap around  $E_F$ , and the non-zero contribution of spin-up density of states in  $E_F$  [20, 62]. This also confirms the stability of HM behavior for  $x = 0.0, 0.25$ , and  $0.50$  as expected from the DOS plots (Fig.4). The Mn atoms have main contribution in  $M_{tot}$ , and the other constituent atoms have less influence. In fact, the hybridization of the p states of Al and Si atoms with Mn-3d

states induces a small spin polarization on the Al or Si, and hence they get a low magnetic moment. The spin polarization formed in the indirect exchange interaction depends directly on the hybridization between magnetic ions and the delocalized electrons. In other words, the spin of one ion influences the spin of another atom through the hybridization with these electrons [62, 63]. Similarly, Nb atom gets a low magnetic moment due to the hybridization of the d states of Mn and Nb atoms. Since the direction of the magnetic moments of the Mn atoms is antiparallel to Nb, Al, and Si atoms, the  $AlCu_2Mn$ -type structure of  $Mn_2NbAl_{1-x}Si_x$

alloys is a typical ferrimagnet. But indeed, the magnetic moments of about 2.00, 1.75, 1.50, 1.15, and 0.99  $\mu_B$  were calculated for the alloys with  $x = 0.0, 0.25, 0.5, 0.75$ , and 1.0 respectively.

**Table 2.** The total and partial magnetic moments ( $\mu_B$ ) of  $Mn_2NbAl_{1-x}Si_x$  alloys. The reported values for closest compounds are also presented for comparison. The accessible experimental data are indicated by \*

| Alloy ( $X_2YZ$ )          | Total        | Mn (X)           | Nb (Y)              | Al (Z)         | Si (Z) | Ref.                   |
|----------------------------|--------------|------------------|---------------------|----------------|--------|------------------------|
| $Mn_2NbAl$                 | 2.00         | 1.20             | -0.30               | -0.03          |        | This work              |
| $Mn_2NbAl$                 | 2.00         | 1.19, 1.21, 1.21 | -0.39, -0.30, -0.36 | -0.03          |        | [49], [48], [46]       |
| $Mn_2NbAl_{0.75}Si_{0.25}$ | 1.75         | 1.04             | -0.26               | -0.02          | -0.03  | This work              |
| $Mn_2NbAl_{0.5}Si_{0.5}$   | 1.50         | 0.92             | -0.20               | -0.02          | -0.03  | This work              |
| $Mn_2NbAl_{0.25}Si_{0.75}$ | 1.15         | 0.74             | -0.18               | -0.02          | -0.02  | This work              |
| $Mn_2NbSi$                 | 0.99         | 0.59             | -0.14               |                | -0.02  | This work              |
| $Mn_2VAl$                  | 1.99         | 1.40             | -0.741 (for V)      | -0.0147        |        | [35]                   |
| $Mn_2ZrAl$                 | 3.00         | 1.73             | -0.34 (for Zr)      | -0.05 (for Al) |        | [46]                   |
| $Mn_2ZrGa$                 | 2.99         | 1.74             | -0.27 (for Zr)      | -0.05 (for Ga) |        | Our previous work [21] |
| $Mn_2ZrGe$                 | 1.99         | 1.14             | -0.18 (for Zr)      | -0.04 (for Ge) |        | Our previous work [36] |
| $Mn_2ZrAs$                 | 0.99         | 0.59             | -0.10 (for Zr)      | -0.02 (for As) |        | Our previous work[20]  |
| $Mn_2MoAl$                 | 1.04         | 0.68             | -0.31 (for Mo)      | -0.01 (for Al) |        | [46]                   |
| $Mn_2RuGa, Mn_2RuGe$       | 0.29*, 1.55* |                  |                     |                |        | [44]                   |



**Fig.5.** The band structure of spin-up states for (a)  $x=0$ , (b)  $x=0.25$ , (c)  $x=0.5$ , (d)  $x=0.75$ , and (e)  $x=1.0$  in  $Mn_2NbAl_{1-x}Si_x$  alloys



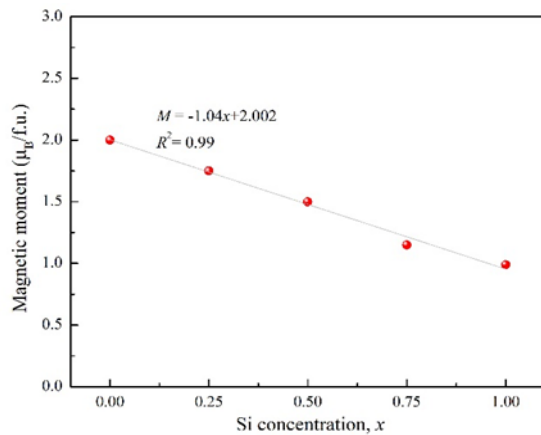


Fig. 6. The variation of total magnetic moment of the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  versus Si concentration

#### 4. Conclusions

The structural, electronic, and magnetic properties of  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  ( $x=0.0-1.0$ ) alloys were theoretically investigated. For the whole series of this alloy, from  $x = 0.0$  to  $1.0$ , the normal structures are more stable than the inverse ones. Moreover, the complete series  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  are found to be ferrimagnetic with total magnetic moment which decreases from  $2.0 \mu\text{B}$  to  $0.99 \mu\text{B}$  as  $x$  increases. The results of the density of states and band structure show that the alloys with  $x = 0.0, 0.25$ , and  $0.50$  have the HM nature with the spin polarization  $P=100\%$  and a real energy gap of about  $0.4 \text{ eV}$ . The results show that the  $\text{Mn}_2\text{NbAl}_{1-x}\text{Si}_x$  alloys can be promising materials for using in spintronic industries. The present results can provide some help for future research of  $\text{Mn}_2$ -based Heusler alloys.

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#### Conflicts of interest

The author declares no conflicts of interest.

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