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Defect energetics in an high-entropy alloy fcc CoCrFeMnNi[†]

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High-entropy alloys (HEAs) are promising candidate materials in nuclear applications owing to their excellent mechanical properties and improved resistance to radiation damage. Defect formation energy in HEAs is directly dependent on the choice of chemical potential, and different chemical potentials can contribute to the discrepancies in the statistical spread. Herein, first-principles calculations were performed to investigate the defect energetics in a fcc CoCrFeMnNi HEA based on chemical potentials, which are back-derived in a self-consistent manner. Chemical potentials are more accurate and precise without additional computational cost and associated uncertainty. The computational results show that defect formation energy is strongly dependent on the local atomic environment and weakly dependent on the chemical composition. Moreover, vacancies prefer the Cr- and Mn-deficient as well as Ni-rich local atomic environment. There is a distinct relationship between the interstitial-specific formation energy and the number of X atoms in 1-NN, while, irrespective of interstitial species, there is no obvious trend in the dependence of the interstitial formation energy and the local atomic environment. The average vacancy migration energies follow the order: Cr < Fe < Mn < Ni < Co, which is lower than that in pure Ni, and the Fe and Cr atoms in 1-NN around vacancies prefer to exchange with vacancies to help vacancy migration. Furthermore, the local atomic environment of the Mn and Ni atoms of the chemical species that exchanges with vacancies suppresses vacancy migration and that of the Co and Fe atoms facilitates vacancy migration. The results of this work can contribute to understanding the enhanced irradiation resistance of HEAs and may provide a new paradigm for the design of advanced radiation-tolerant HEAs.

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1. Introduction

The development and deployment of new-generation fission and fusion reactors is directly dependent on the evolution of advanced high-performance structural materials that can tolerate the extreme environments produced by intense radiation fields, elevated temperatures, and chemical reactive conditions.^{1–3} As we all know, in these environments, high-dose, high-energy neutrons collide with lattice atoms, creating numerous point defects in the material, such as interstitials and vacancies. Further evolution of these irradiation-induced defects leads to the formation of extended defect structures such as dislocation loops, stacking fault tetrahedra, and voids during continuous irradiation.^{2,4} These defects can degrade the physical and mechanical properties of the

material, thus shortening the lifetime of nuclear reactor components and affecting the safe operation of nuclear reactors.^{1,2} Therefore, the development of high-performance materials that are radiation-resistant is critical for advanced nuclear reactor systems.

High-entropy alloys (HEAs),^{5–8} usually a single phase solid solution with multiple principal elements in equiatomic or near equiatomic composition, make them promising candidate materials in nuclear applications⁹ owing to their excellent mechanical properties^{10–12} and improved resistance to radiation damage.^{13,14} It is well established that the production and migration of point defects under irradiation directly affect the number, distribution and agglomeration of surviving defects, and consequently, the microstructure and properties of the materials.⁴ Therefore, it is important to develop a quantitative understanding of the properties of point defects in HEAs. Moreover, reliable data for defect energetics from comprehensive density functional theory (DFT) calculations can provide insights into the key factors on the early stage of irradiation-induced defect production and also provide fundamental parameters in multi-scale simulations for defect evolution, which is highly crucial in understanding and even predicting

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the microstructural evolution of materials under irradiation environments. In addition, the formation and migration energies of point defects are important for the development and testing of semi-empirical interatomic potentials for studying radiation damage behaviors by molecular dynamics (MD) simulation in HEAs.

Theoretical calculations have been made recently to unveil the defect energies in a series of CoCrFeMnNi HEA subsystems, including CoNi,¹⁵ FeNi,^{15–17} CrNi,¹⁵ CoCrNi,¹⁸ CrFeNi,¹⁷ NiFeMnCr¹⁹ and CoCrFeNi.^{18,20–22} However, in HEAs, the extreme chemical disorder induced by the random arrangement of different elements can lead to complex energy landscapes. Consequently, the formation and migration energies of vacancies and interstitials exhibit a distribution of values, which can significantly affect the defect production and evolution. Therefore, the theoretical determination of the defect energetics of various point defects is rather challenging. Due to the computational demands and complexity, few studies on the defect energetics of the high-entropy alloys have been performed. The vacancy formation and migration energies in CrMnFeCoNi have been investigated,²³ and the average vacancy formation energy of each element is almost the same at approximately 2.0 eV, while the average vacancy migration energy of each element increases with increasing atomic number. The effects of the chemical environment and magnetic moment on point defect formations in CoCrNi, CoCrNiFe, and CoCrNiFeMn have been systemically characterized.²¹ It was found that vacancies prefer Ni-rich and Cr-deficient environments, and interstitial defects are primarily dominated by Co and Mn. Furthermore, the point defect formation energies are found to be negatively correlated with the anti-magnetic moment changes in 1-NN atoms.

Different studies on defect energetics in the same HEAs can produce different results. There are two key factors in the first-principles calculations: (1) supercell size; and (2) chemical potential. In general, studies based on smaller supercells produce a wider statistical spread of defect energies than those based on large supercells. Moreover, the defect formation energy is directly dependent upon the choice of chemical potential in HEAs, and different chemical potentials used in these studies can also contribute to the discrepancies in the statistical spread. In the studies on the defect energetics in HEAs, there are different methods to compute the chemical potentials of constituent elements in HEAs. For example, the chemical potential of the defect species was approximated by the total energy of their bulk counterpart with the most stable phase,^{20,22} but it was not precise and suitable for the defect formation energy in the HEAs. Furthermore, the chemical potentials of all chemical components can be calculated using the Widom-type substitution technique.^{15,19,21} Usually, sufficient separate calculations with extra computational costs are needed to obtain an accurate average value and a level of uncertainty. Therefore, a new computational method of chemical potentials of constituent elements in HEA will be proposed without extra computational costs and uncertainty.

As an application, we have chosen the well-studied prototype equiatomic fcc CoCrFeMnNi HEA, which was the first proposed

HEA, also known as the Cantor alloy.⁸ In addition, most work has focused on the fcc CoCrFeMnNi HEA as a model system to study the irradiation resistance of HEAs.¹⁴ Herein, first-principles calculations have been performed to study the defect energetics in fcc CoCrFeMnNi HEA to obtain the formation energies of vacancies and interstitials, as well as the migration energies of the vacancies based on the special quasi-random structures (SQS) approach. First, the chemical potentials of constituent elements were calculated based on a statistical approach to give the distribution of vacancies and interstitial formation energy. Then, the defect formation energies of vacancies and interstitial were calculated, and the effect of local atomic environment on the defect formation energy was also analyzed. Finally, the migration energies of the representative vacancies were also presented. Exploiting the defect energetics in high-entropy alloys can contribute to understanding the enhanced irradiation resistance of HEAs, and may provide a new paradigm for designing structural materials with more improved radiation resistance for the future nuclear applications.

2. Computational methods

The first-principles calculations were performed in the framework of density functional theory (DFT), as implemented in the Vienna *ab initio* simulation package (VASP) code.^{24,25} The interaction between ions and electrons was treated by the projector augmented wave (PAW) method,^{26,27} and the exchange–correlation functional was described by the generalized gradient approximations (GGA) with the Perdew–Burke–Ernzerhof formalism.²⁸ The cut-off energy for the plane-wave basis set was set to 350 eV for all calculations. The Brillouin zone was sampled with a $3 \times 3 \times 3$ Γ -centered k -point mesh for the supercell. The first-order Methfessel–Paxton smearing method of the Fermi surface was used with a smearing width of 0.1 eV to account for orbital partial occupancies of the HEA systems. Furthermore, all calculations were carried out with spin polarization to take into account the magnetic properties, and according to previous ref. 29 and 30 a ferrimagnetic structure was initialized with Cr and Mn spins being aligned anti-parallel to the Fe, Co and Ni atomic spins.

To model the full disorder with the best possible small periodic supercell for the fcc CoCrFeMnNi HEA, a 125-atom supercell ($5 \times 5 \times 5$ primitive cell) of special quasi-random structures (SQS)³¹ was generated by mcsqs code in the alloy theoretic automated toolkit (ATAT) package.³² The equilibrium volume was determined by computing the total energies for at least nine volumes around the equilibrium volume. For each structure, the internal coordinates of all atoms were first relaxed at a given volume by conjugate gradient method with a convergence criterion of 10^{-4} eV and 10^{-2} eV \AA^{-1} for the energy and force, respectively. The optimal lattice parameter was obtained by fitting the calculated energy–volume relation to the Birch–Murnaghan equation of state. Afterwards, all of the defect properties calculations were carried out under this optimal lattice by relaxing only the atomic coordinates with the same convergence criteria.



The defect migration energies were calculated using the climbing-image nudged elastic band (CI-NEB) method,³³ as implemented in VASP. A number of intermediate images are optimized along the migration path with the same convergence criteria as the defect properties calculations. To acquire the minimum energy paths (MEPs) of defects, twelve migration energies for each defect towards its first nearest neighbors (1-NN) were all calculated.

3. Results and discussion

3.1. Chemical potentials

In the HEAs, vacancies and self-interstitials were introduced by removing atoms from their regular lattice positions and placing atoms into their interstitial sites, respectively. Consequently, the defect formation energy can be calculated by comparing the difference between the energy of the relaxed defective supercell and that of the reference state, as follows:^{15,21}

$$E_f = E_d - E_0 \pm \mu_x \quad (1)$$

where E_d and E_0 are the energies of defective and defect-free supercells, respectively; μ_x is the chemical potential of the defect species X (Co, Cr, Fe, Mn and Ni) that was added to (–) or removed (+) from the perfect supercell for interstitials and vacancies, respectively. Accordingly, it can be found that the defect formation energy is sensitively dependent on the choice of chemical potential of the element being removed or added in the HEAs. Since the chemical potential of species is defined as the change of the total free energy per atom of that species and it is highly dependent upon the metallic bonding around that species, it can be expected that the chemical potentials in HEAs are substantially different from those of elements in their pure reference states. Thus, calculations of the chemical potentials of the species are required for the calculations of the defect formation energies in HEAs.

In this study, the chemical potentials of each component were calculated based on a statistical approach to give the distribution of defect formation energy. During the calculation of vacancy formation energy, the chemical potential of the defect species X can be calculated from:³⁴

$$\mu_X = E_0 + \langle E_V^f \rangle - \frac{1}{N_X} \sum_{k=1}^{N_X} E_{d,X,k} \quad (2)$$

$$\langle E_V^f \rangle = \frac{1}{N} \sum_{n=1}^N E_{d,n} - \frac{N-1}{N} E_0 \quad (3)$$

where $\langle E_V^f \rangle$ is the average vacancy formation energy in HEAs; E_0 is the energy of the perfect supercell; $E_{d,x,k}$ is the energy of the defective supercell in which the atom of element X is removed from atomic sites with index k ; N_X is the total number of atoms of element X present in the HEAs; $E_{d,n}$ is the energy of the defective supercell irrespective of the element type removed from the atomic site with index n ; and N is the number of atoms present in the HEAs.

Following the same idea,³⁴ in the course of the calculation of the X-interstitial formation energy with the X–X' dumbbell, the chemical potential of type X with the X–X' dumbbell can be calculated from,

$$\mu_{X-X'} = \frac{1}{N_{X-X'}} \sum_{k=1}^{N_{X-X'}} E_{d,X-X',k} - E_0 - \langle E_I^f \rangle \quad (4)$$

$$\langle E_I^f \rangle = \frac{1}{N} \sum_{n=1}^N E_{d,n} - \frac{N+1}{N} E_0 \quad (5)$$

where $\langle E_I^f \rangle$ is the average interstitial formation energy in HEAs; E_0 is the energy of the perfect supercell; $E_{d,x-x',k}$ is the energy of the defective supercell in which the atom of element X is added to the HEAs to form the X–X' dumbbell with index k ; $N_{X-X'}$ is the total number of atoms of X elements present in the HEAs to form the X–X' dumbbell; $E_{d,n}$ is the energy of the defective supercell irrespective of the element type added to the HEAs to form the X–X' dumbbell with index n ; and N is the number of atoms present in the HEAs.

For the calculations of the defect formation energies, the chemical potentials of constituent elements for vacancies and interstitials with the X–X' dumbbell are presented in Table 1, in comparison with those of pure metals and the results of references in the HEA. For the same constituent element, the chemical potentials for the vacancy and interstitials with different X–X' dumbbells are different, while those of pure metals and the references' results^{21,23} in the HEA are all the same single values. The chemical potentials depend on the vacancy and dumbbell composition. In this sense, our result should be more accurate and precise. Interestingly, the chemical potentials of Co, Fe and Ni in HEA fluctuate around those of the pure metals, while the chemical potential of Cr in the HEA is larger than that of pure Cr and the chemical potential of Mn in the HEA is smaller than that of pure Mn, which can be attributed to the ferromagnetism of Co, Fe and Ni and the anti-ferromagnetism of Cr and Mn. Moreover, our chemical potentials are comparable with the results of references^{21,23} in the HEA, demonstrating that our result is reasonable without additional computational cost and the associated uncertainty.

3.2. Vacancy formation energy

Due to the chemical disorder caused by the random arrangement of various elements, the defect formation energy is

Table 1 Chemical potentials (in eV) of the constituent elements in the fcc CoCrFeMnNi HEA and pure metals

		μ_{Co}	μ_{Cr}	μ_{Fe}	μ_{Mn}	μ_{Ni}
Vacancy		–7.00	–9.27	–8.15	–8.98	–5.51
Interstitials	Co–X	–6.98	–9.41	–8.26	–9.01	–5.26
	Cr–X	–7.09	–9.13	–8.36	–8.91	–5.43
	Fe–X	–6.98	–9.29	–8.24	–9.03	–5.37
	Mn–X	–7.14	–9.09	–8.31	–8.95	–5.42
	Ni–X	–7.01	–9.43	–8.21	–8.98	–5.29
Ref. 23		–7.03	–9.35	–8.21	–9.00	–5.54
Ref. 21		–7.10	–9.76	–8.39	–8.96	–5.56
Pure metals		–7.04	–9.53	–8.25	–8.86	–5.48



dependent on the local atomic environment in HEAs. As a result, there are large variations in defect energies. The defect formation energies take on a distribution, rather than a single value as the pure metal and dilute alloy.³⁵ Moreover, in HEAs, the defect formation energy is usually element-specific.²² In other words, the defect formation energies are also dependent on the type of element or composition,²² *i.e.*, vacancies produced by removing the atoms of different elements or interstitials formed by adding the atoms of different compositions. In this study, the defect formation energies of all atomic sites in the SQS supercell were calculated to analyze their statistical distribution.

For the convenience of description, the vacancies are denoted by the element that is removed for vacancy production. The distribution of the vacancy formation energy is plotted in Fig. 1. The vacancy formation energy for each component in HEAs varies in a relatively wide range. Due to the different local chemical atomic environment, the vacancy formation energy varies from 1.54 eV to 2.33 eV for Co, 1.47 eV to 2.36 eV for Cr, 1.66 eV to 2.33 eV for Fe, 1.68 eV to 2.21 eV for Mn, and 1.52 eV to 2.32 eV, as shown in Fig. 1 and Table 2. Interestingly, the range of vacancy formation energy for Mn is very small in comparison with those for other components, which can be attributed to the complicated magnetism of Mn. In fact, Mn exhibits a very complex magnetic structure in its stable elemental phase,^{19,36} and cannot be simply described as either ferromagnetic or anti-ferromagnetic. As a result, the magnetic behavior of Mn is very different from that of the other elemental constituents in CoCrFeMnNi HEA. The complex magnetic structure of Mn reduces the fluctuation of the vacancy formation energy of Mn in CoCrFeMnNi HEA, and hence the smaller spread of the vacancy formation energy of Mn. The distribution of the vacancy formation energy is different for each chemical species of the vacancy (*i.e.*, Co, Cr, Fe, Mn or Ni), indicating that the vacancy formation energy is dependent on not only the local atomic environment, but also the chemical composition.

In order to quantify the mean and fluctuations (standard deviations) effect, all the data and the element-specific data can be fitted to the Gaussian function, as follows,

$$y = y_0 + A \times \exp\left(-\frac{1}{2} \times \left(\frac{x - \mu}{\sigma}\right)^2\right)$$

Table 2 The mean value (μ) and standard deviation (σ) (in eV) of Gaussian function fit from the distribution of vacancy formation energy and the range (in eV) of the vacancy formation energy in fcc CoCrFeMnNi HEA

Species	μ	σ	Range
V	1.88	0.11	1.47–2.36
V _{Co}	1.88	0.11	1.54–2.33
V _{Cr}	1.87	0.10	1.47–2.36
V _{Fe}	1.90	0.04	1.66–2.33
V _{Mn}	1.77	0.02	1.68–2.21
V _{Ni}	1.89	0.06	1.52–2.32
V in Ni	1.43	—	—

where μ and σ are the mean value and standard deviation, respectively. It has been shown that the Gaussian function is suitable for describing the distribution of the defect formation energy in HEAs, following the literature.¹⁹ From the Gaussian fit, irrespective of the element type removed from the atomic site in the fcc CoCrFeMnNi HEA, the mean vacancy formation energy is 1.88 eV, which is larger than the experimental value of 1.69 ± 0.13 eV or 1.72 ± 0.18 eV,³⁷ and smaller than the other theoretical value of 2.0 eV.²³ For the element-resolved data, the mean vacancy formation energy is 1.88 eV, 1.87 eV, 1.90 eV, 1.77 eV and 1.89 eV for Co, Cr, Fe, Mn and Ni, respectively, as shown in Table 2, demonstrating that the difference in the mean vacancy formation for different vacancies is very small except for the Mn vacancy. It is worth noting that the mean vacancy formation energy for Mn is the smallest among those in the HEA for the complicated magnetic structure. Furthermore, the fluctuation of the vacancy formation energy for Mn is negligible among those in the HEA, consistent with its small range. The mean vacancy formation energy in the HEA is also significantly larger than that in pure Ni of 1.43 eV. The vacancy formation energy of a material is directly related to its irradiation-resistant ability at the early stage of irradiation.^{20,21} Thus, it is more difficult to generate a vacancy in the HEA than pure Ni, and hence more resistant to radiation for the HEA compared with pure Ni. It can be found that the difference of the mean vacancy formation energy for different chemical species is very small, suggesting no preference of vacancy formation for a specific element in the HEA. As a matter of fact, there is no pre-

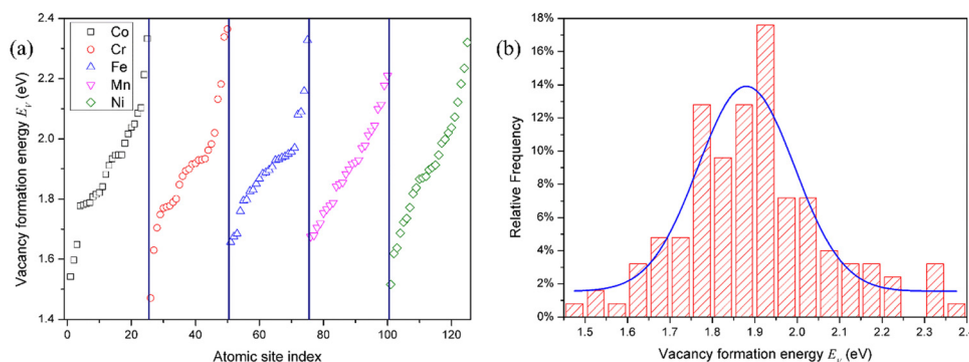


Fig. 1 (a) Vacancy formation energy for each chemical species in fcc CoCrFeMnNi HEA. There are 25 different calculations for the vacancies removed for each species, and these data are compiled in ascending order. (b) Total distribution of vacancy formation energies of all 125 samples in fcc CoCrFeMnNi HEA. The blue line represents Gaussian function fit.



existing atom at the vacancy site in actual materials, and no atom needs to be removed to form a vacancy. Accordingly, the dependence of composition for the vacancy formation energy is artificial and does not exist. The wide distribution of vacancy formation energy for each chemical species and the small difference of the mean value in the vacancy formation energy for different chemical species show that the vacancy formation energy is weakly dependent on the chemical species, while strongly dependent on the local atomic environment.

To investigate the effect of the local atomic environment on the vacancy formation energy in the HEA, the dependence of the vacancy formation energy on the number of atoms in the first nearest neighbor (1-NN) around the defect site was systematically analyzed, as shown in Fig. 2. It can be found that there is no obvious dependence of the vacancy formation energy on the number of Co and Fe atoms in the 1-NN around the vacancy, in agreement with the reference results.²¹ On the other hand, the vacancy formation energy increases with the number of Cr and Mn atoms in the 1-NN, while the vacancy formation energy decreases with the number of Ni atoms in the 1-NN, consistent with the trend from ref. 21 and 23. Thus, the number of Cr, Mn and Ni atoms in the 1-NN is the key factor affecting the vacancy formation energy, and vacancies will more easily form in a Cr- and Mn-deficient, as well as Ni-rich environment in fcc CoCrFeMnNi HEA.

It should be noted that the tendency of the vacancy formation energy with the number of Mn atoms is opposite with the previous results,²¹ which can be attributed to the complicated magnetism of the Mn element. Mn exhibits a very complicated magnetic behavior in its stable ground state.^{19,36} Hence, it cannot be simply described as either a ferromagnetic or anti-ferromagnetic alignment in the fcc CoCrFeMnNi HEA. Importantly, for ferromagnetic elements, there is no significant

dependence between the vacancy formation energy and the numbers of Co or Fe atoms in the 1-NN, while the vacancy formation energy is seen to be negatively correlated with the number of Ni atoms in the 1-NN, which can be attributed to the Ni atoms having the smallest radius among all the elements in the fcc CoCrFeMnNi HEA. During vacancy formation, an increase in the number of Ni atoms in the 1-NN results in a decrease in the atomic perturbation induced by the smallest radius. Moreover, as the radius of the Cr atom is the largest among the constituent elements, the interaction between Cr atoms in the 1-NN is stronger than that for other elements, and increasing the number of Cr atoms in the 1-NN can help suppress the vacancy formation. It can be speculated that the atomic radius of elements can affect the vacancy formation energy in the HEA. Furthermore, the 3d orbitals of the Cr atoms are more delocalized than those of the other elements in the HEA.^{18,23} Thus, the degree of orbital overlap with neighboring atoms increases with the increase of Cr atoms. On contrary, the shrinking of the 3d orbitals is most significant for Ni atoms,^{18,23} and leads to the decrease in the vacancy formation energy with the number of Ni atoms in the 1-NN. Statically, vacancies occur more easily for those with lower formation energy, and the vacancies with mean formation energy occurs more frequently. In fact, the mean vacancy formation energy is only an artificial value, which is generated by removing the atom from the constructed supercells in the HEA. It should be noted that thermodynamically, vacancies occur more easily and frequently for those with the lowest vacancy formation energies.

3.3. Interstitial formation energy

In pure fcc Ni, among six possible interstitial configurations: octahedral, tetrahedral, and crowdion, as well as [100], [110]

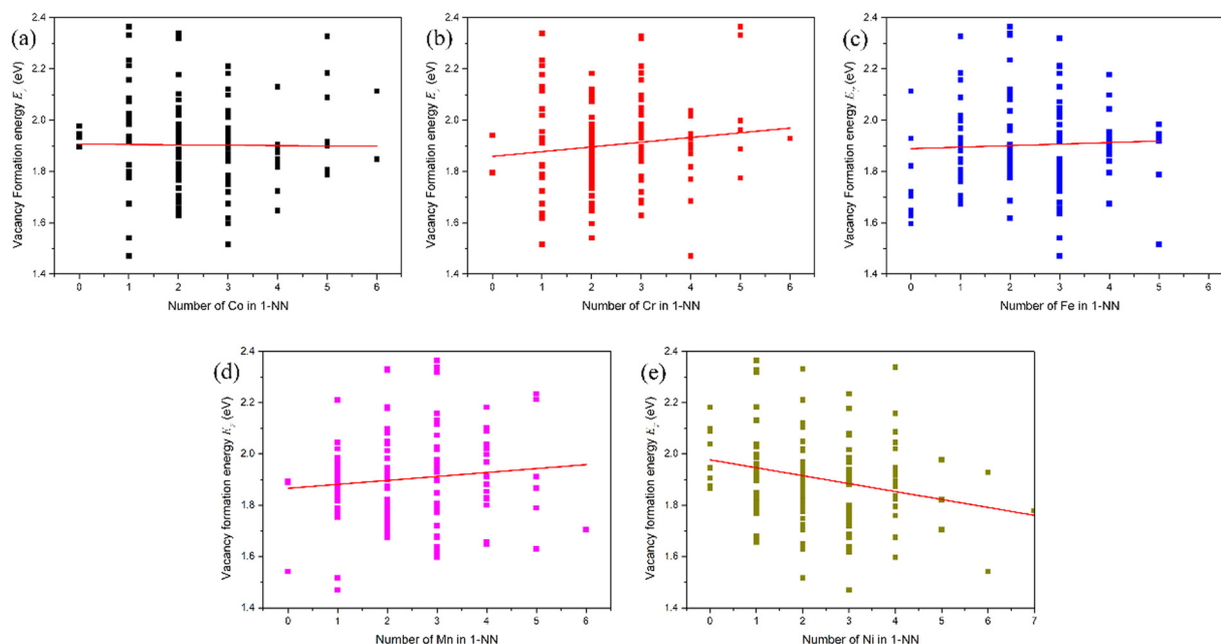


Fig. 2 Evolution of vacancy formation energy with the number of X atoms in the first nearest neighbor (1NN) in fcc CoCrFeMnNi HEA: (a) Co, (b) Cr, (c) Fe, (d) Mn, and (e) Ni. The line is fitted from the data.



and [111] dumbbells, it can be found that the [100] dumbbell has the lowest formation energy. Thus, the [100] dumbbell is the most stable interstitial defect configuration.¹⁵ Therefore, the interstitial formation energy in the fcc CoCrFeMnNi HEA was also calculated in the form of the [100] dumbbell. All the possible configurations in the form of the [100] dumbbell for the Co, Cr, Fe, Mn and Ni interstitials were considered, *i.e.*, Co-Co, Co-Cr, Co-Fe, Co-Mn and Co-Ni [100] dumbbells for the Co interstitial, Cr-Co, Cr-Cr, Cr-Fe, Cr-Mn and Cr-Ni [100] dumbbells for the Cr interstitial, Fe-Co, Fe-Cr, Fe-Fe, Fe-Mn and Fe-Ni [100] dumbbells for the Fe interstitial, Mn-Co, Mn-Cr, Mn-Fe, Mn-Mn and Mn-Ni [100] dumbbells for the Mn interstitial, as well as Ni-Co, Ni-Cr, Ni-Fe, Ni-Mn and Ni-Ni [100] dumbbells for the Ni interstitial, respectively, in fcc CoCrFeMnNi HEA.

For clarity, the Ni interstitial formation energy for the Ni-Co, Ni-Cr, Ni-Fe, Ni-Mn and Ni-Ni [100] dumbbells is presented in Fig. 3, while the other X (Co, Cr, Fe and Mn) interstitial formation energies for the X-Co, X-Cr, X-Fe, X-Mn and X-Ni [100] dumbbells in fcc CoCrFeMnNi HEA are presented in Fig. S1–S4 in the (ESI[†]). The range of Ni interstitial formation energies for the Ni-Co, Ni-Cr, Ni-Fe, Ni-Mn and Ni-Ni [100] dumbbells is from 2.84 to 3.66 eV, 3.01 to 3.88 eV, 2.54 to 3.54 eV, 2.87 to 3.68 eV and 3.01 to 4.18 eV, respectively, as shown in Table 3, and those of other X interstitial formations are also presented in Table 3. It can be seen that different [100] dumbbells produce different ranges of interstitial formation energy for different X (Co, Cr, Fe, Mn and Ni) interstitials. The distribution of interstitial formation energy mainly originates from the local atomic environment around the interstitial defect. Moreover, the range of interstitial formation energy is larger than that of the vacancy formation energy. This is due to the fact that the variation of distortion induced by the interstitial is larger than that induced by vacancy.

From Gaussian fit, irrespective of the [100] dumbbell type, the mean interstitial formation energies are 3.31 eV, 3.30 eV, 3.26 eV, 3.26 eV and 3.30 eV for the Co, Cr, Fe, Mn and Ni interstitials, respectively, as shown in Table 4, which are smaller than that in pure Ni of 4.38 eV. That the difference in

Table 3 The range (in eV) of the interstitial formation energy in the fcc CoCrFeMnNi HEA

Interstitial	X-Co	X-Cr	X-Fe	X-Mn	X-Ni
Co	2.87–3.79	2.95–3.79	2.62–3.62	2.83–3.52	2.99–3.81
Cr	2.73–3.66	2.68–3.82	2.48–3.99	2.69–3.94	2.74–4.05
Fe	2.86–3.79	2.95–3.81	2.65–3.83	2.76–3.84	2.99–3.72
Mn	2.78–3.84	2.81–4.14	2.70–3.78	2.82–3.88	2.92–3.85
Ni	2.84–3.66	3.01–3.88	2.54–3.54	2.87–3.68	3.01–4.18

the mean interstitial formation energy is very small shows that there is no obvious relation between the interstitial formation energy and interstitial species added by different elements. Moreover, the interstitial formation energies for many dumbbell-specific species do not follow a Gaussian distribution. In summary, the interstitial formation energy in fcc CoCrFeMnNi HEA is smaller than its counterpart in pure Ni, while the vacancy formation energy in fcc CoCrFeMnNi HEA is larger than that in pure Ni. Accordingly, it is easier for interstitial formation in fcc CoCrFeMnNi HEA than that in pure Ni, while it is the opposite for vacancy formation. Since the production of a vacancy is easier than that of an interstitial in the same material, it is more difficult to produce point defects in fcc CoCrFeMnNi HEA than pure Ni. Hence, fcc CoCrFeMnNi is more resistant to radiation damage.

The influence of the local atomic environment on the interstitial formation energy in fcc CoCrFeMnNi HEA was also analyzed following the analytic method for the vacancy formation energy. The evolution of the Ni interstitial formation energy with the number of X (Co, Cr, Fe, Mn and Ni) atoms in the 1-NN in fcc CoCrFeMnNi HEA is presented in Fig. 4. It can be found that there is no distinct relation between the Ni interstitial formation energy and the number of Co, Fe and Ni atoms in the 1-NN, while the Ni interstitial formation energy increases with the number of Cr atoms in the 1-NN and decreases with the number of Mn atoms in the 1-NN. For the Co, Cr, Fe and Mn interstitial, the dependence of the interstitial formation energy on the number of X (Co, Cr, Fe, Mn and Ni) atoms in the 1-NN was also studied, as shown in Fig. S5–S8 (ESI[†]). The dependence of the Co, Cr, Fe, Mn and Ni interstitial

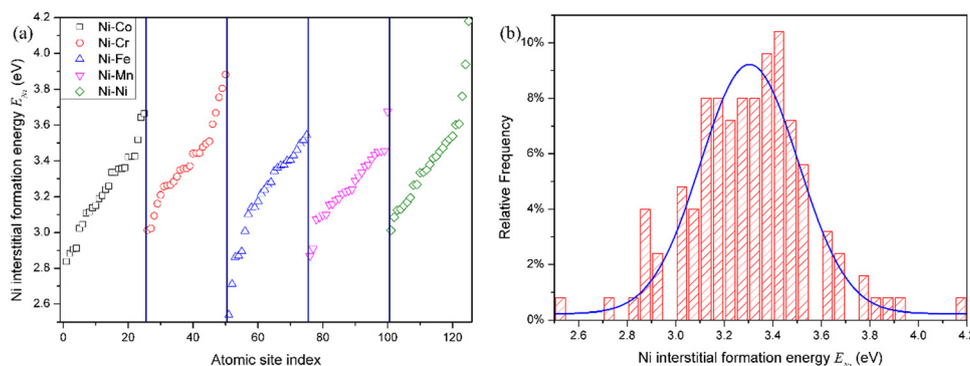


Fig. 3 (a) Ni interstitial formation energy for Ni-Co, Ni-Cr, Ni-Fe, Ni-Mn and Ni-Ni [100] dumbbells in the fcc CoCrFeMnNi HEA. There are 25 different calculations for each dumbbell, and these data are compiled in an ascending order. (b) Total distribution of Ni interstitial formation energies of all 125 samples in the fcc CoCrFeMnNi HEA. The blue line represents Gaussian function fit.



Table 4 Interstitial formation energy (in eV) from Gaussian function fit in the fcc CoCrFeMnNi HEA

Interstitial	X-interstitial	X-Co	X-Cr	X-Fe	X-Mn	X-Ni
Co	3.31 ± 0.21	—	3.44 ± 0.20	—	—	3.41 ± 0.14
Cr	3.30 ± 0.31	3.26 ± 0.18	—	3.22 ± 0.20	—	3.41 ± 0.33
Fe	3.26 ± 0.23	3.25 ± 0.12	—	3.12 ± 0.03	3.20 ± 0.09	—
Mn	3.26 ± 0.32	—	—	3.18 ± 0.08	—	3.42 ± 0.08
Ni	3.30 ± 0.20	—	3.35 ± 0.09	—	—	—
Int in Ni	4.38					

formation energy on the number of X (Co, Cr, Fe, Mn and Ni) atoms in the 1-NN is summarized in Table 5. It can be found there is a distinct relationship between the interstitial-specific formation energy and the number of X atoms in the 1-NN. Meanwhile, irrespective of the interstitial species, there is no obvious trend of the dependence of the interstitial formation energy on the local atomic environment due to the different pronounced lattice relaxations induced by the introduction of different X interstitials compared with the vacancy. The interstitial formation energy is also weakly dependent on the composition, and strongly dependent on the local atomic environment as vacancy formation energy.

3.4. Vacancy migration energy

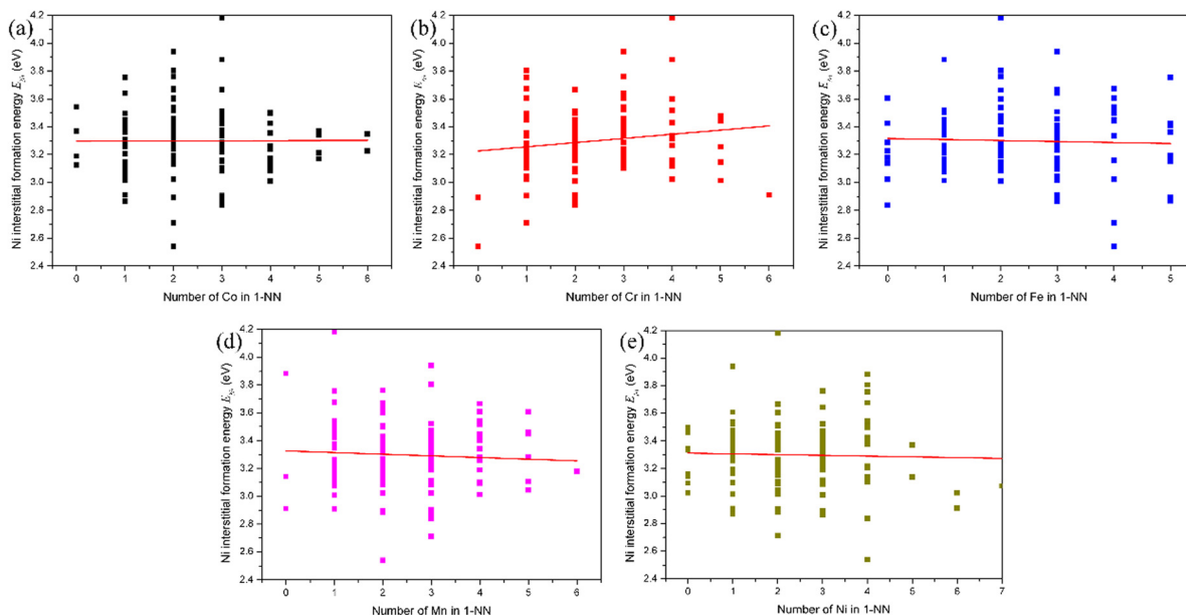
The migration barrier is an important parameter in atomic transport properties. Due to the complexity of the interstitial migration mechanism in the HEAs, it is difficult to calculate the interstitial migration energy in the fcc CoCrFeMnNi HEA. Accordingly, only the vacancy migration energy in the fcc CoCrFeMnNi HEA was studied in this work. The diffusion of vacancies is considered by directly exchanging a lattice atom between two first nearest neighbor sites. In the ref. 18 and 35, the vacancy migration barriers were calculated by directly optimizing the saddle-point structures located at the midpoints

Table 5 The relation between the X-interstitial formation energy and the number of X' atoms in the 1-NN

Interstitial	Co in 1-NN	Cr in 1-NN	Fe in 1-NN	Mn in 1-NN	Ni in 1-NN
Co	+	—	—	—	+
Cr	—	+	—	NA	—
Fe	+	+	+	—	NA
Mn	NA	+	NA	—	+
Ni	NA	+	NA	—	NA

Note: +: X-interstitial formation energy increases with the number of X' atoms in 1-NN; -: X-interstitial formation energy decreases with the number of X' atoms in 1-NN; NA: there is no obvious dependence of the X-interstitial formation energy on the number of X' atoms in 1-NN.

of the corresponding diffusion paths. However, due to the complicated local atomic environments around the vacancy site and the lattice atom exchange with vacancy, the saddle-point is not always located at the middle point of the diffusion path. Consequently, the vacancy migration energies were calculated using the CI-NEB method with one image, which was tested enough to get an accurate vacancy migration energy in pure Ni, as shown in Fig. S9 (ESI†). In addition, in the fcc NiFeMnCr HEA,¹⁹ the single-image NEB method provides the most efficient method to calculate the vacancy migration energy.

**Fig. 4** Evolution of the Ni interstitial formation energy with the number of X atoms in the first nearest neighbor (1NN) in fcc CoCrFeMnNi HEA: (a) Co, (b) Cr, (c) Fe, (d) Mn, and (e) Ni. The line is fitted from the data.

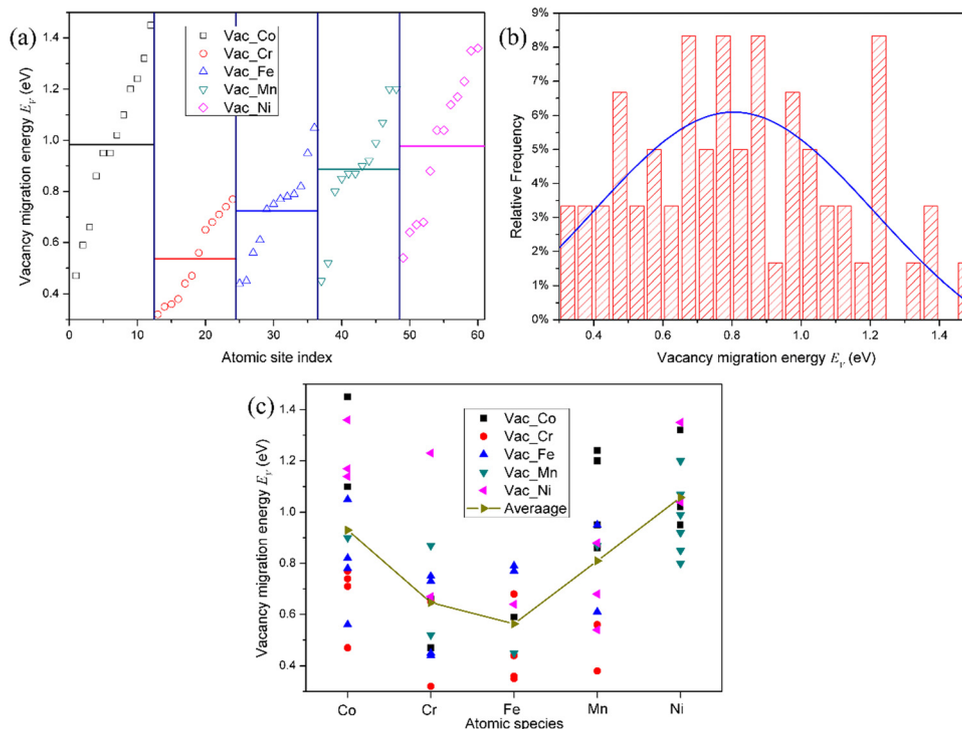


Fig. 5 (a) Vacancy migration energy for the representative of each chemical species in the fcc CoCrFeMnNi HEA. There are 12 different calculations for the representative vacancies removed for each species, and these data are compiled in ascending order. The horizontal line denotes the average migration energy for the vacancy removed for each species. (b) Total distribution of vacancy migration energies of all calculated 60 samples in the fcc CoCrFeMnNi HEA. The blue line represents Gaussian function fit. (c) Variation of vacancy migration energy with the chemical species that exchanges with vacancy in the fcc CoCrFeMnNi HEA.

Due to the limitation of computational resources, the migration energy of vacancy near the mean vacancy formation energy for the element-specific vacancy was calculated. The vacancy migration energy for the representative of each chemical species in fcc CoCrFeMnNi HEA is presented in Fig. 5. It can be found that due to chemical disorder, the vacancy migration barriers also exhibit distributions in the HEA. It can be seen that the average values of the vacancy migration energies in the HEA follow the order: Cr < Fe < Mn < Ni < Co, which are lower than that in pure Ni of 1.09 eV. This order corresponds to the experimentally observed order of atoms for individual constituent elements: Ni < Co < Fe < Cr < Mn,³⁸ *i.e.*, the lower the diffusivity, the larger the value of the migration energy. It should be noted that there is no obvious dependence of the average vacancy migration energy on the atomic number for each element, *i.e.*, it does not follow that the vacancy migration energy increases with the atomic number, as shown in the ref. 23. Actually, the vacancy migration energy decreases with the atomic radius. Furthermore, irrespective of the vacancy species, from the Gaussian fit, the mean vacancy migration energy is about 0.8 eV, which is also lower than that in pure Ni. Accordingly, the migration energy of the vacancy and interstitial will overlap with each other, which can enhance the recombination of defects in the HEA, thereby improving the irradiation resistance. It can be reasonably speculated that the lattice distortion from the chemical disorder facilitates the vacancy migration. Moreover, the order of chemical species

which exchanges with the vacancy is as follows: Co < Ni < Mn < Cr < Fe, as seen from Fig. 5(c), in excellent agreement with the atomic radius of Ni < Co < Mn < Fe < Cr. Therefore, Fe and Cr atoms prefer to exchange with the vacancy to help the vacancy migration. It is worth noting that the order of the chemical species for the vacancy exchange seems to not strictly correspond with the average values of the vacancy migration energies. However, from Fig. 5(c), the difference in the migration energy that the vacancy exchanges with Cr and Fe is very small. Therefore, we can consider that the order of the chemical species for the vacancy to exchange and the average values of the vacancy migration energies remain consistent.

The lattice distortion induced by the random arrangement of different species in the HEAs significantly modifies the diffusion kinetics due to different displacements at different atomic species sites. As a result, the diffusion barrier is dependent on the local atomic environment around the vacancy site and that around the chemical species which exchanges with vacancy. The evolution of the vacancy migration energy with the number of X atoms in the first nearest neighbor (1-NN) of the chemical species that exchanges with the vacancy in fcc CoCrFeMnNi HEA is presented in Fig. 6. It can be found that the vacancy migration energy increases with the number of Mn and Ni atoms, and decreases with the number of Co and Fe atoms in the 1-NN of the chemical species which exchanges with vacancy in fcc CoCrFeMnNi HEA. In other words, the local atomic environment of Mn and Ni atoms of the chemical



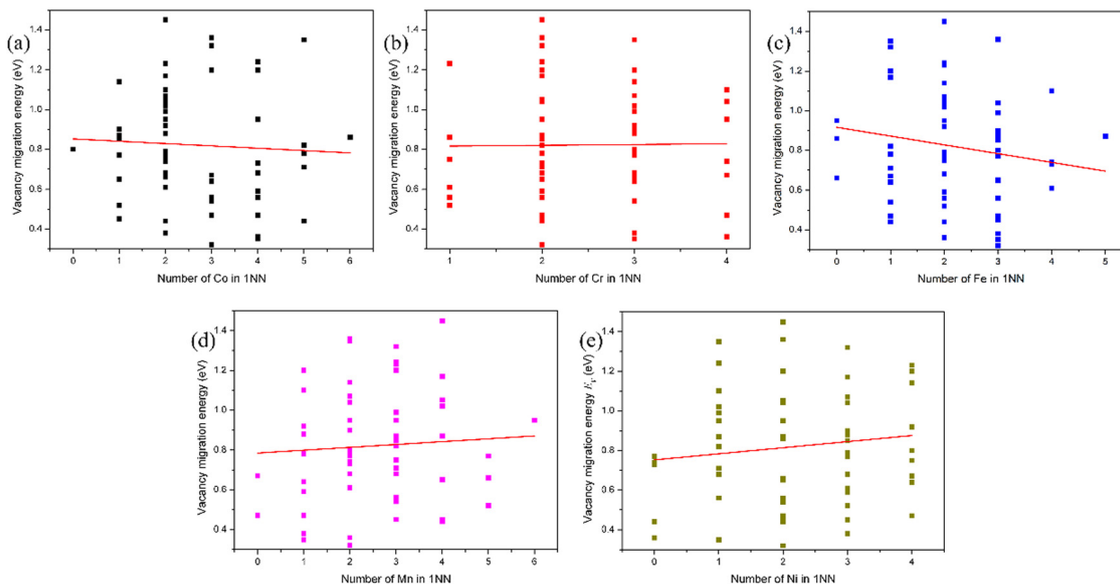


Fig. 6 Evolution of the vacancy migration energy with the number of X atoms in the first nearest neighbor (1-NN) of the chemical species that exchanges with vacancy in fcc CoCrFeMnNi HEA: (a) Co, (b) Cr, (c) Fe, (d) Mn, and (e) Ni. The line is fitted from the data.

species that exchanges with the vacancy suppresses the vacancy migration, and that of Co and Fe atoms facilitates vacancy migration. The vacancy migration energy is dependent on not only the vacancy species and its local atomic environment, but also on the chemical species which exchanges with the vacancy and its local atomic environment. The decrease of the vacancy migration energy will significantly enhance the recombination of vacancies and interstitials, and contribute to the defect annihilation, making the HEAs more irradiation-resistant. Due to the limitation of the local atomic environment, the number of data points is very small in the calculation. As a result, it seems that the fitted lines in Fig. 2, 4 and 6 do not align well with the data points. It should be noted that in fact, the lines in Fig. 2, 4 and 6 are fitted in the term of the least squares method.

4. Conclusion

First-principles calculations have been performed to investigate the defect energetics in fcc CoCrFeMnNi HEA to obtain the formation energies of vacancies and interstitials, as well as the migration energies of the vacancies based on the SQS approach. The chemical potentials of constituent elements in HEAs are back-derived in a self-consistent manner based on a statistical approach, and the chemical potentials are more accurate and precise without additional computational cost and the associated uncertainty.

The defect formation energy takes on a wide distribution due to chemical disorder induced by the random arrangement of constituent elements in fcc CoCrFeMnNi HEA. The defect formation energy is strongly dependent on the local atomic environment, and weakly dependent on the chemical composition. Moreover, vacancies prefer the Cr- and Mn-deficient, as well as Ni-rich local atomic environment. There is a distinct relationship between the interstitial-specific formation energy and the

number of X atoms in the 1-NN, while irrespective of interstitial species, there is no obvious dependence of the interstitial formation energy and local atomic environment. The interstitial formation energy in fcc CoCrFeMnNi HEA is smaller than that of its counterpart in pure Ni, while the vacancy formation energy in fcc CoCrFeMnNi HEA is larger than that in pure Ni. This demonstrates that it is more difficult to produce point defects in fcc CoCrFeMnNi HEA than pure Ni. Hence, fcc CoCrFeMnNi is more resistant to radiation damage. The vacancy migration energy also exhibits distributions in fcc CoCrFeMnNi HEA, and its average values follow the order: Cr < Fe < Mn < Ni < Co, which is lower than that in pure Ni. Moreover, Fe and Cr atoms in the 1-NN around vacancy prefer to exchange with the vacancy to help the vacancy migration. The local atomic environment of Mn and Ni atoms of the chemical species that exchanges with the vacancy suppresses the vacancy migration, and that of Co and Fe atom facilitates vacancy migration.

The results of this work can certainly improve our understanding of their enhanced irradiation resistance of HEAs, and provide guidance for the design of advanced radiation-tolerant HEAs. Recently, some ref. 39 and 40 pointed out that the defect formation energy is dependent on not only the 1-NN composition, but also equally the arrangement of 1-NN atoms around the defect in concentrated solid solution alloys. In the future, the effect of the arrangement of 1-NN atoms around the defect on the defect formation energy in fcc CoCrFeMnNi HEA will be studied using machine learning based on the dataset of the first-principles results of the defect formation energy.

Author contributions

Chan Gao – conceptualization, methodology, formal analysis, investigation, writing – original draft, writing – review & editing.



Shuyu Wang & Xiao Liu – validation, formal analysis, writing – review & editing. Chandra Veer Singh – software, resources, writing – review & editing.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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