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# First-principles study for quasi-static growth model in FeAl intermetallic based on Wulff cluster model

 Lin Song,<sup>a</sup> Anchen Shao,<sup>a</sup> Dong Li,<sup>d</sup> Xuelei Tian,<sup>b</sup> Zhuhui Qiao,<sup>id</sup>\*<sup>ac</sup> Huaguo Tang\*<sup>ac</sup> and Xiaohang Lin<sup>id</sup>\*<sup>b</sup>

In order to investigate the structure of FeAl mesoscopic crystals segregating in liquid state alloys, we have determined their equilibrium structures (Wulff shape) based on the Wulff cluster model. For non-stoichiometric surface terminations, the chemical environment is taken into account through the chemical potential of the constituents. In this case, different cluster shapes change as a function of the chemical environment. In order to model the growth process in more detail, we propose a quasi-static growth model based on the sequential addition of (sub-)monolayers in the most favorable surface directions. Thus, a sequence of different Wulff shapes results in the growth process, as illustrated for the FeAl intermetallic compound. This model is proved preliminarily by calculating the concentration trend of Al/Fe atoms on both Al-terminated and Fe-terminated surfaces, and by simulating the most stable layer adsorbed on these two surfaces. This model might be helpful in analyzing the growth processes including nucleation barriers during nucleation processes theoretically.

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## 1. Introduction

Solidification processes are necessary and important for most alloy manufacturing. The solidification processes, especially the nucleation stage, have a significant influence on the properties of the alloys including mechanical properties, electrochemical properties, catalytic properties, *etc.*<sup>1–3</sup> In short, nucleation is the process in which the nanoparticles grow gradually and reach the critical nucleation radius under certain supercooling conditions in alloy melts. Obviously, the structure and growth process of nanoparticles in melts, as an origin, will directly affect the nucleation process.

The ways to research the melts' properties could mainly be divided into experimental and theoretical methods. For experimental ones, high temperature X-ray diffraction (HTXRD)<sup>4,5</sup> and neutron diffraction<sup>6</sup> were adopted to explore the structures of metallic melts.<sup>4–18</sup> The theoretical methods, including molecular dynamics (MD),<sup>14–18</sup> phase fields, cellular automata (CA) and first principle simulation are used to study the crystallization and growth process after nucleation.<sup>19–22</sup> Unfortunately, due to the limitation of high temperature, the behavior

of nanoparticles in melts at the atomic level, especially the growth process, is still unclear.

In previous research, we proposed a model (Wulff cluster model) to investigate the ordering structures in metallic melts. It uses the most probable structure as the equivalent structure to describe the short-range ordering distribution of metallic melts under the thermal equilibrium state. In this model, the equivalent structure is determined by the Wulff construction with the crystal structures inside. The simulated XRD results of the model show good agreement with the high-temperature X-ray diffraction (HTXRD) experiments, which indicates that it is a reasonable equivalent model to describe pure metals, binary homogeneous alloys, eutectic alloys, and intermetallic compound melt systems.<sup>23–26</sup> This model gives us a chance to further investigate the growth processes, especially the processes before nucleation.

The FeAl intermetallic shows many attractive properties like high strength, excellent corrosion, low density at high temperature and economical price.<sup>27–30</sup> Exactly these fantastic characteristics make the FeAl intermetallic a potential candidate for high temperature alloys. Moreover, FeAl which has a wide range of phase diagram components is one of the simplest intermetallic compounds. Therefore, it was selected to explore the structure of nanoparticles in the melt before nucleation and the effect of external environment on the structure growth mechanism. In addition, this paper attempted to explain the mechanism/processes for the growth order of the FeAl compound before nucleation.

Recently, first principle calculations based on density functional theory was widely used to investigate the structures and

<sup>a</sup>Shandong Laboratory of Advanced Materials and Green Manufacturing at Yantai, Yantai, 264006, People's Republic of China

<sup>b</sup>Key Laboratory for Liquid-Solid Structural Evolution and Processing of Materials, Ministry of Education, Shandong University, Jinan 250061, People's Republic of China. E-mail: lxh12345@sdu.edu.cn

<sup>c</sup>State Key Laboratory of Solid Lubrication, Lanzhou Institute of Chemical Physics, Chinese Academy of Sciences, Lanzhou, 730000, People's Republic of China. E-mail: laotang@licp.cas.cn; zhqiao@licp.cas.cn

<sup>d</sup>Shandong Ludian Line Equipment Co. Ltd, China



other properties of nano-particles at the atomic level.<sup>31,32</sup> In this work, DFT calculations were applied to study the equilibrium shape as a function of the chemical environment. A quasi-static model based on the chemical potential correction was proposed to determine the evolution of the growth morphology of nano-particles. The non-equilibrium growth process is approximated to a series of equilibrium nodes to better understand the growth process. Finally, the mechanism/processes for the FeAl inter-metallic growth order is given by calculation.

## 2. Theoretical calculations

### 2.1 Numerical simulation details

All DFT<sup>33,34</sup> total energy calculations in this paper have been performed using the Vienna *Ab initio* Simulation Package (VASP) within the generalized gradient approximation (GGA) to describe the exchange–correlation effects, employing the Perdew, Burke and Ernzerhof (PBE) exchange–correlation functional.<sup>35–38</sup> The ionic cores are represented by projector augmented wave (PAW) potentials<sup>39</sup> as constructed by Kresse and Joubert.<sup>40</sup> It has already been proved to give reasonable results of Fe–Al system with enough accuracy. The structures were optimized until atomic forces were converged to 0.01 eV Å<sup>-1</sup>. A 7 × 7 × 1 *k*-point sampling is chosen for all calculations based on careful convergence tests.<sup>41</sup> The plane wave cut-off energy was set to 400 eV. Moreover, due to magnetic of iron atoms, the spin polarization has been taken into considered.

### 2.2 Surface models

In this work, different types of low-index surfaces have been built, including FeAl(100), FeAl(110), FeAl(111), FeAl(210), FeAl(211) and FeAl(221) surfaces (Fig. 1). The calculation method of surface energy is a typical slab model. The thickness of vacuum layer is more than 20 Å to ensure the accuracy of surface calculation results. Both Fe-terminated and Al-terminated of FeAl are considered.

For elemental materials, surface energies ( $E_{\text{surf}}$ ) are defined according to

$$E_{\text{surf}} = \frac{1}{2A}(E_{\text{tot}} - NE_{\text{bulk}}) \quad (1)$$

where  $E_{\text{tot}}$  and  $E_{\text{bulk}}$  are the total energies of surface system and bulk, respectively.  $N$  is the number of bulk unit cells in the slab.

It can be seen from Fig. 1, the Fe and Al atoms in some stable slab structures are located on different layers, which results in different stoichiometric ratios in calculation. These surfaces with such characteristics appear on FeAl(100), FeAl(111), FeAl(210), FeAl(221) surfaces. In other faces, like FeAl(110) and FeAl(211) surfaces, each face has the same amount of Al and Fe atoms and the same layer structure. As a consequence, the values of chemical potential of these surfaces are invariant constants, which also correspond to the calculation results below.

Compared with elemental materials, surfaces of compounds are more complex as the stoichiometry at the surfaces might be different from the one in the bulk. Therefore, generally

speaking, then the bulk energy can no longer be used as the only reference for the surface energy, in addition the chemical potential of the constituents has to be considered. Consequently, the most stable structure of a FeAl compound surface is given by the minimum of surface energy

$$\gamma = \frac{1}{2A}[E_{\text{surf}} - \mu_{\text{FeAl}}N_{\text{Al}} - \mu_{\text{Fe}}(N_{\text{Fe}} - N_{\text{Al}})] \quad (2)$$

Here  $A$  is the area of the FeAl surface,  $E_{\text{surf}}$  is the total energy of the FeAl surface per surface unit cell.  $\mu_{\text{FeAl}}$  and  $\mu_{\text{Fe}}$  are the chemical potentials of the FeAl bulk and Fe bulk, respectively. And  $N_{\text{Fe}}$  and  $N_{\text{Al}}$  are the numbers of Fe and Al atoms in the slab per surface unit cell.

In addition, the chemical potential is related to the elemental bulk chemical potentials through the heat of formation  $\Delta H_{\text{FeAl}}$ , which is given by

$$\Delta H_{\text{FeAl}} = E_{\text{tot}}(\text{FeAl}) - E_{\text{bulk}}(\text{Fe}) - E_{\text{bulk}}(\text{Al}) \quad (3)$$

Here  $E_{\text{bulk}}(\text{Fe})$  and  $E_{\text{bulk}}(\text{Al})$  are the bulk energies of Fe and Al, and the  $E_{\text{tot}}(\text{FeAl})$  is the total energy of FeAl bulk per formula unit.

Therefore, we obtain the possible range of Fe chemical potentials that is determined by the energy of bulk Fe and the heat of formation.

$$\mu_{\text{Fe}}^{\text{bulk}} - \Delta H_{\text{FeAl}} < \mu_{\text{Fe}} < \mu_{\text{Fe}}^{\text{bulk}} \quad (4)$$

## 3. Results and discussion

### 3.1 FeAl bulk calculation

FeAl bulk is a simple cubic structure which contains one Fe and Al atom in the unit cell. The calculated formation energy and lattice parameters of FeAl are shown in Table 1. From our calculation, the lattice constant is  $a = b = c = 2.872$  Å, which only 0.59% differs from the experimental data.<sup>43</sup> Furthermore, due to the presence of Fe element in the FeAl compounds, the results of magnetism have been considered. The ferromagnetism is the most stable state, which shows a good agreement with other results.<sup>44,45</sup> Therefore, the subsequent calculations are all based on ferromagnetism.

### 3.2 Surface energy

Based on Wulff cluster model mentioned above, the equivalent structure is determined by the Wulff construction with the crystal structures inside. According to Wulff theory,<sup>46</sup> the shape with the lowest surface energy can be determined by the following methods: draw the radius vector from the origin of the polar diagram of surface energy. At the intersection, a plane perpendicular to the corresponding radius vector is constructed, and then the plane with the lowest surface energy is connected to form a closed convex shape.

As the first step to derive the Wulff shape, we consider the surface energy of low-index surfaces. Recall that for non-stoichiometric surfaces the gas-phase environment



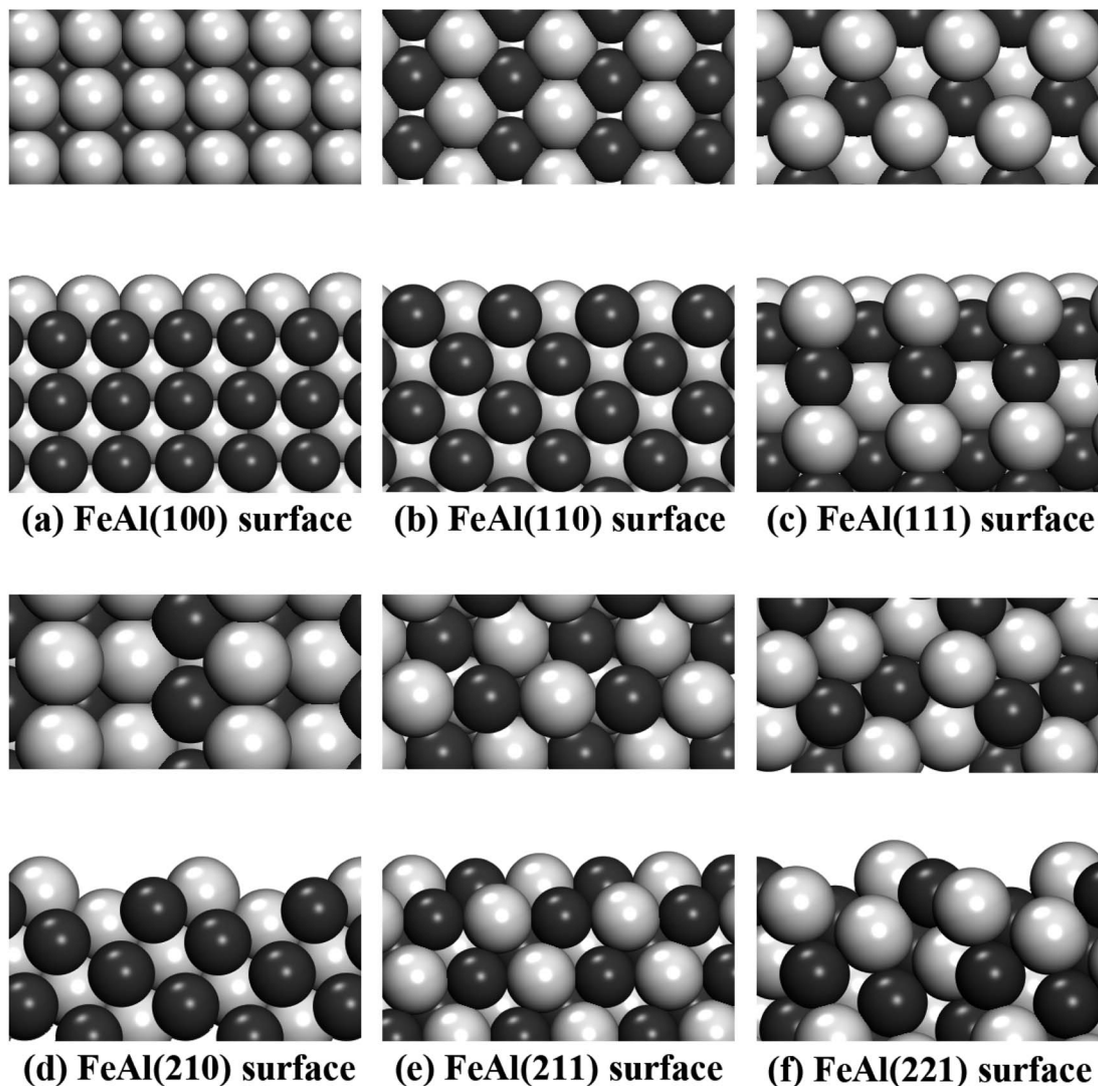


Fig. 1 (a)–(f) show the top and side views of the surface (100), (110), (111), (210), (211) and (221), respectively. The silver balls represent the Al atoms, and the black balls represent the Fe atoms.

Table 1 The calculated parameters ( $\text{\AA}$ ), the calculated  $E_{\text{for}}$  (eV per atom) of the FeAl intermetallic

Species	Space group	$a$ ( $\text{\AA}$ )	$b$ ( $\text{\AA}$ )	$c$ ( $\text{\AA}$ )	$E_{\text{for}}$ (eV per atom)
FeAl	 $Pm3m$	2.872	2.872	2.872	−6.359
		2.851 <sup>a</sup>	2.851 <sup>a</sup>	2.851 <sup>a</sup>	−6.947 <sup>a</sup>
		2.889 <sup>b</sup>	2.889 <sup>b</sup>	2.889 <sup>b</sup>	

<sup>a</sup> Cal. in ref. 42. <sup>b</sup> Exp. in ref. 43.

contributes to the stability of the surfaces through the corresponding chemical potentials (shown in eqn (2)). In fact, most of the FeAl surfaces that we consider are not stoichiometric. As we are using slab models to calculate surface energies, there are

always two surfaces present in our surface model, which are furthermore not necessarily equal. However, typically the different surface energies for asymmetric slabs can not be separated so that only an average surface energy can be derived. In the case of the FeAl surfaces, we have been able to always construct slabs with the same first layer of atoms, but the second layer can differ. Still, we take the average value which is reasonable considering that the first layer on both sides is the same.

To deal with the non-stoichiometric surfaces, it is more common to plot the surface energy as a function of the chemical potential  $\mu_{\text{Fe}}$  of the constituents according to eqn (2), as we have done in Fig. 2. There, the dashed lines indicate the range of possible Fe chemical potentials given by eqn (3) and (4) with  $\Delta H_{\text{FeAl}} = 0.63$  eV obtained from our calculation. Low Fe chemical potential is corresponding to the environment of Al-rich while the Fe bulk chemical potentials give the Fe-rich limit. It is obvious from Fig. 2 that the FeAl(100) surface of Al terminated



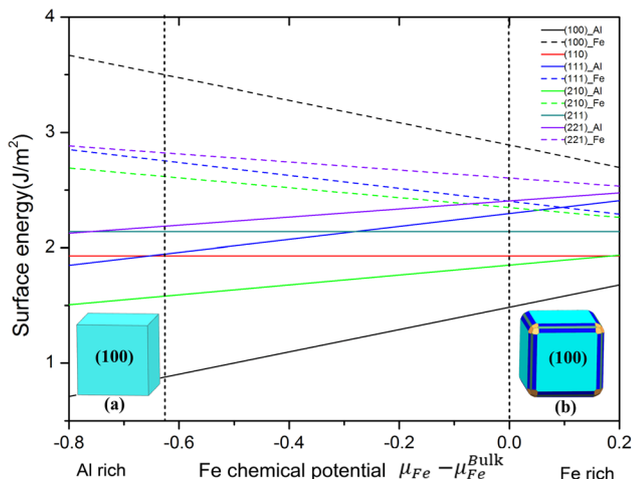


Fig. 2 Surfaces energies of different surface and Al-terminated FeAl including (100), (110), (111), (210), (211) and (221) in  $\text{J m}^{-2}$  as a function of the difference of the chemical potential of Fe with respect to bulk Fe. The area between two vertical short dashed lines represents the stable termination at a given growth condition. Here the different colors represent different surfaces. The shapes on both sides represent the Wulff shape at the position of vertical short dashed lines, respectively.

has the lowest surface energy in the whole range of chemical potential. In particular, Fig. 2 shows that the surface energies of Al-terminated surface structures are totally lower (about  $0.5\text{--}1 \text{ J m}^{-2}$ ) than those surface energies of Fe-terminated surface structures. In other words, the surface structure of Al-terminated is more stable than that Fe-terminated ones from the thermodynamic equilibrium, which agrees with other results.<sup>47,48</sup> These provide an important data foundation for the establishment of the subsequent nanoparticles Wulff shape.

The Wulff shape can be easily obtained from the surface energy corresponding to a certain chemical potential. Panels (a) and (b) in Fig. 2 illustrate the Wulff shape in the limit of Al rich and Fe rich condition. Upon increasing the chemical potential from the lower bound of the stability (shown in Fig. 2), the shape changes continuously. For Al rich conditions, we obtained the cubic Wulff shapes which are the FeAl(100) surface occupying the whole area. With the increase of Fe content in the chemical potential, the (210) and (110) surface exist on the edge; a (211) surface is formed on the vertex of the cubic shape. Note that although the (110) surface is the third stable surface, it only formed on the edge with the smallest area (area in blue in Fig. 2b). The results determine that FeAl(100) surface is the most stable surface of Wulff shape in FeAl intermetallics (the chemical potential changes from Al rich to Fe rich condition). Note that we have performed preliminary experiments of nucleation processes in Al/Cu melts at different temperatures. The results of first measurements using liquid X-ray diffraction (XRD) are in fact consistent with the predictions of our Wulff cluster model, giving credibility to the model presented here.<sup>23–26</sup>

### 3.3 The quasi-static growth model

In principle, the Wulff shape of FeAl mesoscopic crystals shows the equilibrium structure with the highest thermodynamic stability, which means that the mesoscopic FeAl crystals form such a shape in equilibrium. However, during the mesoscopic crystal growth to larger sizes in the ideal bulk structure, along one specific normal sub-monolayers of the involved species have to follow each other in the same order as in the bulk. This is the basis for our simple quasi-static growth model that will be explained in more detail below. We assume that always one type of facet grows completely by one (sub-)monolayer until the next surface will start to grow in another direction. Therefore, the growth process can be treated as a sequence of several quasi-static states, which can be described by a series of Wulff shapes.

To grow an ideal bulk crystal, in one particular direction different types of surface terminations corresponding to a sequence of sub-monolayers should follow in a specific order, according to the surface structures illustrated in Fig. 1. When such a new surface termination is completed, the Wulff shape is determined by this new surface structure and the unchanged surfaces in the other directions, whereas the old surface termination is not relevant anymore. The starting structure of the growth process should be the Wulff shape with the highest thermodynamic stability, which is formed by the most stable surfaces in every direction. Generally speaking, the growth processes prefer to follow the path with a relatively low Gibbs free energy. We propose a quasi-static growth model in which growth occurs sequentially by adding (sub-)monolayers along one surface normal that have the highest probability, *i.e.*, the Gibbs free energy per volume. Thus, we obtain a series of Wulff shapes corresponding to intermediate states which together describe the growth process from one most stable shape to the next larger one. In this model, we ignore the single deposition processes leading to the growth of a new (sub-)monolayer. Therefore, we call our approach a quasi-static model. Details of the growth of one particular facet are not considered, but still this model might give us a general idea about growth processes of mesoscopic crystals. This might be helpful in analyzing cluster shapes observed in cluster growth experiments.

To illustrate our model, in Fig. 3 we show the sequence of Wulff shapes of FeAl that result from our quasi-static approach. The chosen Fe chemical potential corresponds to the one of bulk Fe. The process starts with the most stable shape, a cubic-like structure with FeAl(210) and FeAl(211) surfaces at the edge. After every growth event, the volume of the cluster increases according to the addition of a (sub-)monolayer along the chosen directions given by the termination with the lowest surface energy. The structure of cluster almost no change until the third step occurs, only FeAl(100), FeAl(110) and FeAl(211) surfaces. Then, the structure changes from a simple cubic to a dodecahedron. After that, the cycle of growth finishes, the structure that corresponds to the starting configuration is formed.

Moreover, the barrier of every growth model corresponding to the above structure in Fig. 3. We can clearly find that there is a small platform from (a) to (b). A key quantitative finding of this study is that a huge barrier ( $1.41 \text{ eV}$  per atom) is crossed



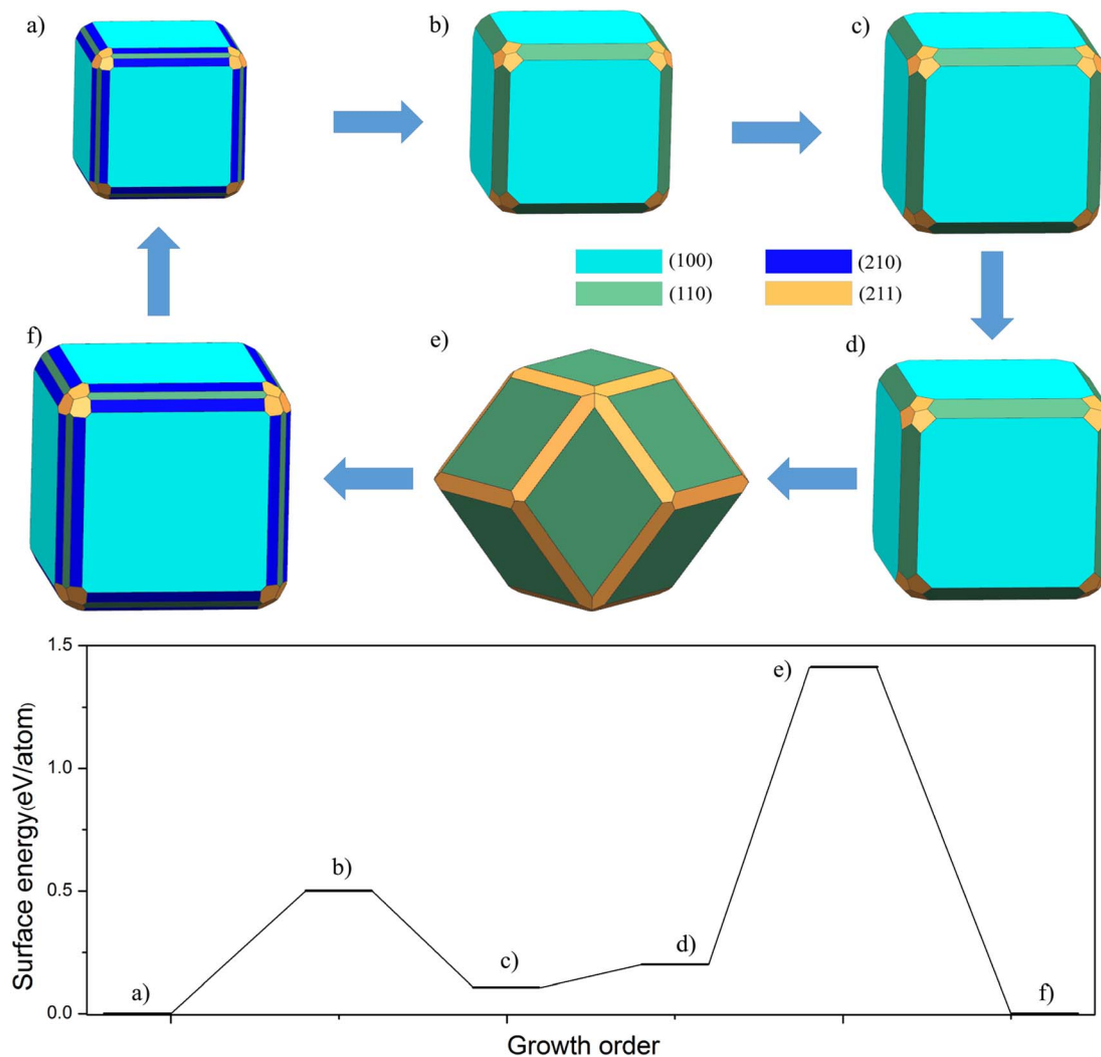


Fig. 3 Quasi-static cluster growth model of FeAl intermetallic by Wulff shape upon addition of sub-monolayers for  $\mu_{\text{Fe}} = \mu_{\text{Fe}}^{\text{bulk}}$ . The model (a)–(f) are the growth order of the clusters at the thermodynamically stable conditions. Different colors represent different surfaces. The barrier of every growth model corresponding to the above structure.

during the transition from cubic to dodecahedron. Although this growth barrier is not directly equivalent to the nucleation barrier, in our opinion, the barrier of this microstructure is closely related to the nucleation barrier under the undercooling condition. By comparing the nucleation and growth barriers of Fe containing intermetallics such as FeSi and FeAl, we believe that they are positively correlated. Thus, this model would be able to explain how clusters nucleated in a melt could keep their Wulff shape upon further growth.

Note that we have performed some preliminary experiments addressing the nucleation rate of undercooled Al/Cu melts, as mentioned above. The results of these experiments with respect to the structure of the nucleated clusters are consistent with our quasi-static growth model which will be the subject of a forthcoming publication.

Here, we'd like to do some preliminary proof by simulation. Generally speaking, there are two completely different models of cluster growth including layer by layer growth and free atom

adsorption in the melt. The layer–layer growth and free atom adsorption growth has been considered separately in the proof part. For the layer–layer growth, we use an approximation that a layer-by-layer growth happens following the order of compounds crystal structures to describe the growth of intermetallic compounds. This assumption is based on the fact that the intermetallic compounds' crystal structures do exist after the solidification of metallic melts. Could this model describe the growth processes of nanoparticle before nucleation correctly? In other words, we are wondering if it is really the favorable path to follow the order of crystal structure. We calculated the cohesive energy of Al layer (Fig. 4a)/Fe layer (Fig. 4b) adsorbed on Al-terminated FeAl(100) surface, which is defined by the following formula:

$$E_{\text{coh}} = E_{\text{tot}} - E_{\text{surf}} - N\mu_{\text{add}} \quad (5)$$



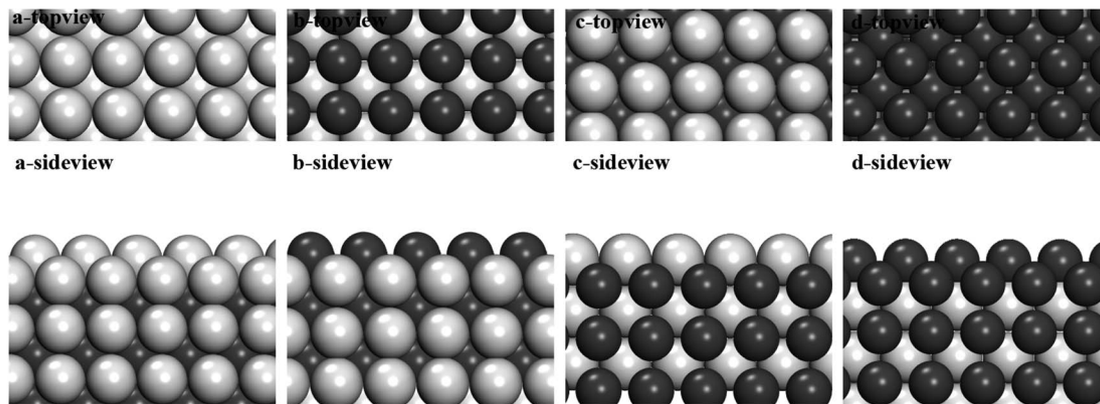


Fig. 4 (a)–(d) represent the top view and side view of different layers (Al layer and Fe layer) grown on Al-terminated and Fe-terminated of (100) planes, and their cohesive energies were shown below. The silver and black balls represent the Al and Fe atoms, respectively.

where  $E_{\text{surf}}$  is the total energy of Al/Fe terminated FeAl surface,  $N$  is the number of additional atoms.

Obviously, on the Al-terminated FeAl (100) surface, the cohesive energy of a Fe layer is much smaller than of a Al layer ( $1.6 \text{ eV } \text{\AA}^{-1}$  smaller). It is more favorable to grow a Fe layer on this surface, which is exactly the order of FeAl crystal. For the Al layer (Fig. 4c)/Fe layer (Fig. 4d) adsorbed on Fe-terminated FeAl (100) surface, Al layer is more stable (about  $0.5 \text{ eV } \text{\AA}^{-1}$  smaller), which also follows the FeAl crystal structure.

For free atom adsorption growth, we calculate one single Al/Fe atom adsorbed on different FeAl (100) surfaces, in order to get the adsorption tendency of atomic type (Fig. 5). It is obvious that Fe atom ( $-3.32 \text{ eV}$ ) prefer to adsorb on Al-terminated surface compared with Al atom ( $-2.84 \text{ eV}$ ). This means that Fe free atom concentrated around Al-terminated surface in melts, which give the same results with the calculation above. For Fe-terminated surface, Al atom is more favorable.

All the results show that the intermetallic compounds formed when the nanoparticle grown in melts. The growth process following the order of crystal has the lowest Gibbs free energy, in other words, is the most favorable path. The quasi-

static growth model is reasonable to describe the growth processes of intermetallic compounds, and make it possible to further investigate the nucleation barriers theoretically.

## 4. Conclusion

In this paper, we proposed a quasi-static growth model of compound clusters based on a sequence of stable surface terminations created by the addition of (sub-)monolayers. Based on Wulff cluster model, the corresponding stable cluster shapes are derived within the Wulff construction scheme. We illustrated this scheme using FeAl intermetallic compound cluster. The surface energies have been derived from first-principles electronic structure calculations. For non-stoichiometric surface termination, the surfaces energies depend on the chemical environment as reflected in the chemical potentials of the elements. In this case, the shape of the mesoscopic crystals' changes with respect to the chemical potential. Using quasi-static growth model to describe the growth processes, we visualize the resulting sequence of FeAl cluster shapes which can be used to analyze the cluster shapes

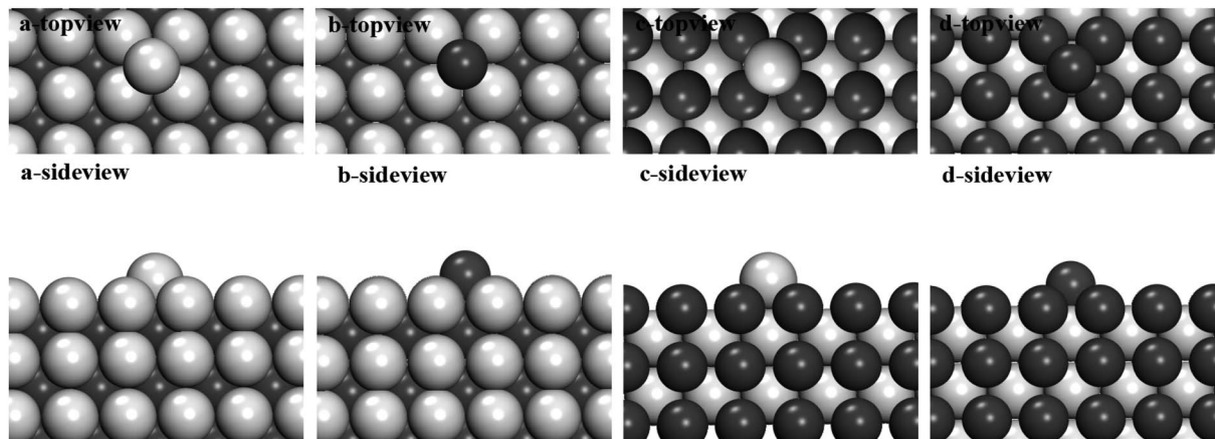


Fig. 5 (a)–(d) represent the top view and side view of different atoms (Al atom and Fe atom) adsorption on Al-terminated and Fe-terminated of (100) planes, and their adsorption energies were shown below. The silver and black balls represent the Al and Fe atoms, respectively.



observed in experiments addressing cluster growth processes. In addition, the growing barriers in different surfaces have also been determined. The barrier is applied to describe the growth cycle of microstructure, and it is likely to be closely related to the nucleation ratio under a supercooling condition. The model is proved preliminarily by calculating the adsorption trend of Al/Fe atom on both Al-terminated and Fe-terminated surfaces, and by simulating the most stable layer adsorbed on these two surfaces. The quasi-static growth model is reasonable to describe the growth processes of intermetallic compounds, and make it possible to further investigate the nucleation barriers theoretically.

## Conflicts of interest

There are no conflicts to declare.

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