On the nucleus composition during isothermal alloy solidification

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Accurate determination of the nucleus composition during isothermal alloy solidification still represents a great challenge nowadays. In this paper, a kinetic scheme was added to the Hillert-Rettenmayr thermodynamic analysis of the deviation from local equilibrium at migrating phase interfaces. A so-called interface permeability was introduced to account for the unambiguous determination of the energy dissipation of the solute rearrangement at the liquid-solid interface and the driving force for interface movement, from which the nucleus composition can be then evaluated. After benchmark test, a pragmatic nucleation model for solidification was also proposed, and applied in three-dimensional phase-field simulations of nucleation and subsequent dendritic growth during isothermal solidification process in one hypothetic Al-Si alloy. Moreover, the influence of the interface permeability on nucleation rate was fully discussed by exploring its effect on the initial nucleus components and the corresponding nucleation driving force.

When alloy melt cools down continuously or cools rapidly to a certain temperature with certain undercooling, nucleus of primary solid phase may start to form in the melt, and thus trigger the evolution of microstructure during the solidification process. Nucleation is the prior stage during the solidification process, and draws numerous attentions in the field of materials¹,². In order to gain a comprehensive and quantitative view of nucleation process in alloys, accurate determination of the composition for nucleus, especially for the first nucleus, during solidification is prerequisite, which still remains a challenge in experimental and theoretical investigations nowadays³.

Baker and Cahn⁴ firstly explored the thermodynamically possible composition range for nucleus, as demonstrated in Fig. 1.

Assuming in a fictitious binary A-B alloy during isothermal solidification, \( x_{\text{local}}^\alpha \) and \( x_L^\alpha \) are the equilibrium compositions while \( x_L^\alpha \) is the initial composition of alloy melt. Baker and Cahn⁴ concluded that it would be thermodynamically possible to nucleate \( \alpha \) with any composition between \( x_{\text{min}}^\alpha \) and \( x_{\text{max}}^\alpha \) from the melt. In order to further determine the specific composition for the nucleus, two hypotheses are generally used. One is the well-known local equilibrium hypothesis (i.e., the common tangent construction)⁵, as schematically described in Fig. 1. Under the local equilibrium hypothesis, the chemical potentials of the solid and liquid phases should be equal, and thus the composition of \( \alpha \) nucleus from melt is exactly the equilibrium composition \( x_{\text{local}}^\alpha \). While the other is the so-called quasi equilibrium hypothesis (i.e., the parallel tangent condition)⁶, as also demonstrated in Fig. 1. Under the quasi equilibrium hypothesis, the diffusion potentials of the solid and liquid phases are required to be equal, and thus the composition of \( \alpha \) nucleus from melt is exactly the equilibrium composition \( x_{\text{quasi}}^\alpha \). While the above two hypotheses are widely used in the field of materials, the nucleation process is simply treated to be either equilibrium or near equilibrium state. While in practice, the nucleation process should be non-equilibrium and even strongly non-equilibrium in some cases.

On purpose of precisely determining the nucleus composition, one approach is in great need for describing the non-equilibrium process by suspending the above-mentioned thermodynamic hypotheses, which is the major target of this work. Recently, Hillert and Rettenmayr⁷ gave a thermodynamic view of deviation from local equilibrium at migrating phase interfaces. Their constructions recapitulated that a part of the driving force is consumed via the exchange of solute atoms between two phases over the interface, resulting in a deviation from the local equilibrium. With the Hillert-Rettenmayr analysis, one may analyze the driving force for the non-equilibrium nucleation process, which is the necessity for the later quantitative description of microstructure evolution during solidification process. However, before the unambiguous determination of driving forces for non-equilibrium nucleation process, the accurate nucleus composition should be given. However, the thermodynamic alone cannot predict the actual composition of nucleus.

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Received: 5 January 2018
Accepted: 6 March 2018
Published online: 28 March 2018
Consequently, a kinetic scheme will be added on the Hillert-Rettenmayr thermodynamic analysis in this paper, from which the nucleus composition for alloys during isothermal solidification can be unambiguously determined. Furthermore, based on the thermodynamic and kinetic analysis, a new but pragmatic nucleation model for isothermal solidification is thus to be proposed, and further applied to simulate the nucleation and dendritic growth process in a binary Al-Si alloy using the phase-field model with finite interface dissipation.

**Results and Discussion**

**Hillert-Rettenmayr thermodynamic analysis.** During the formation of precipitation solid $\alpha$ from a supersaturated liquid $L$, a local decrease in Gibbs energy is yielding

$$\Delta G_m^{\text{total}} = (1 - x^\alpha)(\mu^A_L - \mu^A_L) + x^\alpha(\mu^B_L - \mu^B_L)$$

$$= (1 - x^L)(\mu^A_L - \mu^A_L) + x^L(\mu^B_L - \mu^B_L) + (x^L - x^\alpha)(\mu^B_L - \mu^B_L + \mu^B_L)$$

where $x^\alpha$ and $x^L$ are the local compositions at the interface of $\alpha$ and $L$, $\mu^A_L$ and $\mu^B_L$ are the chemical potentials of species A and B in liquid, while $\mu^A_\alpha$ and $\mu^B_\alpha$ are treated analogously in solid $\alpha$.

According to Hillert and Rettenmayr, the phase transformation during solidification can be divided into two detached processes. One is the transformation from liquid $L$ with composition $x^L$ to solid $\alpha$ with composition $x^\alpha$, while the other is the adjustment of the composition from $x^\alpha$ to $x^L$ by exchanging atoms between the two phases via diffusion over the interface, as shown in Fig. 2. Thus, one can separate the total driving force into the fraction $\Delta G_m$ which drives the phase transformation and the fraction $\Delta G_i$ which drives the redistribution of atoms between the phases.
Here, $\Delta = \Delta + \Delta$ indicates the diffusion potential difference of two phases. $\Delta = \Delta \cdot \Delta$ means the difference in concentrations between the two phases.

With the Hillert-Rettenmayr thermodynamic analysis, $x^\alpha$ is the initial alloy composition, and is known a priori. Accordingly, the chemical potentials $\mu^A_L$ and $\mu^B_L$ can also be derived. While in Eqs (3) and (4), $x$, $\mu^A_L$ and $\mu^B_L$ are unknown, but $\mu^A_L$ and $\mu^B_L$ depend on the nucleus composition $x^\alpha$. Thus, one can clearly separate $\Delta G_{m_{\text{total}}}$, $\Delta G_{m}$ and $\Delta G_{t}$ for specific alloy composition if the nucleus composition is known. But Eqs (3) and (4) are not enough to evaluate $x^\alpha$. Instead, only a corrected nucleus concentration range can be gained based on the constraint that both $\Delta G_{m}$ and $\Delta G_{t}$ cannot be less than zero. Thus, one needs more profound analysis to fix the freedom, from the perspective of kinetics.

**Kinetic analysis.** Taking a fictitious A-B binary system as an example, five different alloy compositions, i.e., a, b, c, d and e, at one constant temperature are chosen for demonstration, as presented in Fig. 3. Point a lies on left side of solidus line, point b lies between solidus line and T_0 line, point c lies on T_0 line, point d lies between T_0 line and liquidus line, while point e lies on the right side of liquidus line. The possible nucleus concentration ranges corresponding to different initial melt concentrations based on the Hillert-Rettenmayr thermodynamic analysis are obtained and illustrated schematically with the molar Gibbs energy diagrams in Fig. 3. It is well known that

$\Delta G_{m_{\text{total}}} = \Delta G_{t} + \Delta G_{m}$ (2)

$\Delta G_{m} = (\mu^A_L - \mu^A_L) - x^\alpha \cdot \Delta \mu$ (3)

$\Delta G_{t} = \Delta x \cdot \Delta \mu$ (4)
the vertical distances represent the driving forces for the precipitation of nucleus with various compositions from the melt, which are composed of $\Delta G_m$ and $\Delta G_t$ for each supersaturated melt concentration. As displayed in Fig. 3, the colored part characterizes $\Delta G_m$ and the blank part is $\Delta G_t$. For clarity, the distribution of driving force information is sorted out and shown underneath the graph. Next this diagram will be analyzed and elaborated from the kinetic view.

For this purpose, a physical kinetic parameter named the interface permeability for redistribution flux or the inverse interface resistivity, $P$, originally defined in the phase-field model with finite interface dissipation, is directly used here. According to refs 8,9, $P$ has the units of an inverse action density ($\text{cm}^3/(\text{Js})$), which is used to characterize the ability of atoms to overcome resistance during migration inside interface. When interface permeability $P$ equals to zero, it indicates that the rearrangement resistance is huge enough to nail the solute atoms. While on the contrary, if the interface permeability $P$ approaches to infinity (or larger enough), the solute rearrangement will be almost free from resistance, and the diffusion potential difference will approach zero instantaneously. Then the impact of atomic mobility on dissipation of the solute rearrangement and driving force for interface migration can be explained expediently. The discussion will be divided into three cases on the basis of the solidification conditions:

**Case 1 (point d).** The interface migration from liquid to solid is torpid, which implies that a liquid atom has enough time and enough trials to move to find its place in the new crystal structure of solid phase. While the permeability approaches to be infinite, the driving force consumed for the motion of the solute atoms is inexistente but dissipated in the phase transition completely. That is to say, $\Delta G_t = 0$ while $\Delta G_m$ is maximum here. With the permeability decreasing, the atomic mobility resistance will increase, the driving force consumed in solute rearrangement will increase and the available driving force for phase transition will be reduced. Furthermore, when the resistance is large enough, the driving force will be exhausted entirely by the rearrangement and motion of solute atoms over the interface, i.e., that $\Delta G_t$ maximum and $\Delta G_m = 0$.

**Case 2 (point a).** The interface will be pushed quickly toward liquid phase. There is a certain chance that some solute atoms will be pinned before migration hence the atom mobility is highly sensitive to the resistance in this situation. Therefore, $\Delta G_t$ shows a drastically declination compared to the slow solidification since it is positively associated with the product of resistance and the number of moving atoms. If $P$ is extremely large, the composition of the precipitate phase will be located at somewhere the diffusion potentials of the two phases are consistent. Furthermore, with the decrease of $P$, the resistance will increase and the quantity of moving solute atoms will decrease, thus the dissipation $\Delta G_t$ will increase from zero (i.e., that the resistance is non-existent) to a certain peak and then decreases to zero (i.e., that the solute atoms are all frozen). Besides, the loss of $\Delta G_m$ will increase with the deepening of the atomic pinning until a minimum $\Delta G_m$ reaches at the point $\Delta G_t = 0$, as illustrated in Fig. 3. It is possible for the clusters to inherit the parental component here.

**Case 3 (points b and c).** The situation can be treated analogously as Case 2, but the solidification process is slower, $\Delta G_m$ is smaller while $\Delta G_t$ is larger. Furthermore, it is worth noting that while the melt concentration locates on $T_0$, the permeability reduces to zero, both $\Delta G_m$ and $\Delta G_t$ are equal to zero, which suggests that the system maintains a delicate balance in this situation, where a slight disturbance will cause its rupture.

Next a quantitative determination of the energy dissipation and the driving force for nucleation will be provided.

It is universally acknowledged that the difference of diffusion potential between solid and liquid phases will drive the rearrangement and motion of solute atoms over the interface. The diffusion flux for solute $B$ that crosses the atomistic physical interface can be expressed as:

$$ J_B = \frac{M_{\text{inter}}}{V_m} \nabla \mu_B $$

(5)

$M_{\text{inter}}$ is the atomic mobility over the interface. Based on ref 9, one can have $M_{\text{inter}} = a \eta P/8$, where $a$ is the size of the physical solid-liquid interface, and can simply be assumed to be in the same scale of the unit cell of the solidified solid phase, $\eta$ is the thickness of the interface. $V_m$ is the molar volume and $\nabla \mu_B$ is the gradient in diffusion potential over the interface, which can be estimated as $\Delta \mu_B/\eta$ for the interface with a thickness of $\eta$. Alternatively, the diffusion flux for solute $B$ along the moving interface can also be defined as:

$$ J_B = \nu \cdot \Delta x $$

(6)

$$ \nu = M \cdot \Delta G_m $$

(7)

where $\nu$ is the average velocity of moving interface, $\Delta x$ is the composition difference of solute in solid and liquid phases over the interface, and $M$ is the interface mobility for migrating interface.

Integrating Eqs (4–7), $\Delta G_t$ is related to $\Delta G_m$ via,

$$ \Delta G_t = \frac{aP(\Delta \mu_B)^2}{8V_m M \Delta G_m} $$

(8)

Now with the third equation (i.e., Eq. (8)) besides Eqs (3) and (4), one can solve all the unknown quantities in Eqs (3) and (4).

Insert Eqs (3) and (4) into Eq. (8), and the expression for the nucleus composition $x^*$ is emerged:
where \( \Delta \) shows a downward trend. The more complex case lies in that, as the composition moves from the liquidus to the Ti line, the nucleation potential increases from positive to negative, leading to the appearance of the second nucleus. Moreover, the position of the nucleus will not change as the calculation results also show that when \( P \) decreases from infinity to zero, the composition of the second nucleus will move from \( x_{\text{quasi}} \) to \( x^2 \).

Based on Eq. (9), one can arrive at the following conclusions:

### Benchmark test

Based on thermodynamic and kinetic analysis demonstrated above, the nucleation composition can be unambiguously determined for a given initial melt composition and interfacial permeability in a specific alloy. In this section, different Al-Si binary alloys were chosen as benchmark test for the above thermodynamic and kinetic analysis. For the sake of simplicity, the linear phase diagram of the binary Al-Si system was utilized, from which the Gibbs energy, chemical/diffusion potentials of both liquid and solid phases can be evaluated. All the relevant thermophysical parameters used in the calculations are listed in Table 1, except for the values of the interface permeability \( P \) which was calculated to keep the interface movement in the diffusion-control regime. The calculated compositions of nucleus are labeled in Fig. 4. Based on the rigorous analysis of the results in Fig. 4, the following conclusions that the influence of permeability on the new phase composition can be divided into three cases can be drawn.

#### Case 1: Initial concentration is on the left side of \( T_0 \) line. When \( P \) varies from infinity to zero, the composition of the first nucleus will move from \( x_{\text{quasi}} \) to \( x^1 \). According to the analysis in the previous section, one know \( \Delta G_m \geq 0 \) here, and hence it is possible for nucleus to form inside the entire composition range from \( x_{\text{quasi}} \) to \( x^1 \).

#### Case 2: Initial concentration is between \( T_0 \) line and liquidus. With the decrease of \( P \), the nucleation driving force shows a downward trend. The more complex case lies in that, as the composition moves from \( x_{\text{quasi}} \) to \( x^1 \), the nucleation driving force is reduced from positive to negative, and the nucleus are generated only when the nucleation driving force is greater than zero, leading to the narrowed concentration range of nucleus. Moreover, the calculation results also show that when \( P \) goes from infinity to a certain threshold greater than zero, a critical component point \( x_{\text{quasi}} \) which is in good agreement with the Hillert-Rettenmayr thermodynamic construction will be exposed, and the composition of the nucleus will travel from \( x_{\text{quasi}} \) to \( x^1 \) while \( P \) is less than the minimum limit, the nucleus will not appear.

#### Case 3: Initial concentration is on the right side of liquidus. The nucleation driving force is less with all the \( P \)s. Thus the formation of nucleus is impossible. The shaded part in Fig. 4 is the theoretical range given by Hillert et al. Obviously, the calculated results are consistent with the theoretical construction, and the composition of first nucleus can be determined uniquely by \( P \) within the theoretical range.

| Parameters                        | Symbols | Values  |
|-----------------------------------|---------|---------|
| interface mobility                | \( M \) | \( 3.84 \times 10^{-2} \text{ cm}^2/\text{s} \) |
| molar volume                      | \( V_m \) | \( 10.06 \text{ cm}^2/\text{mol} \) |
| melting temperature of solvent Al | \( T_m \) | \( 933.6 \text{ K} \) |
| isothermal temperature            | \( T \) | \( 900 \text{ K} \) |
| liquidus slope                    | \( m_t \) | \( -618.42 \text{ K/at.fraction} \) |
| equilibrium partitioning coefficient | \( k_p \) | \( 0.105 \) |

Table 1. List of all the relevant parameters used in the present calculation of nucleus compositions.
Nucleation model and its application. Based on the previous sections on the evaluation of nucleus composition during isothermal solidification, a pragmatic nucleation model is thus proposed for isothermal solidification in alloys, as demonstrated in the following: During the isothermal solidification, the liquid concentration is not absolutely uniform. The local slight fluctuation in consistency with the Gaussian distribution exists throughout the solidification process. Moreover, once the initial local liquid composition and the interfacial permeability are given, the corresponding nucleus composition and its driving force can be evaluated according to Eqs (3), (4) and (9). Thus, the uneven driving forces for different local liquid compositions will lead to different probabilities of the nucleus in the melt. According to the classical nucleation theory \(^{12}\), if the driving force somewhere is large enough to satisfy the condition

\[
\sigma{\Delta}\geq \frac{G_{d\alpha x}^2}{m}
\]

the nucleus with the radius of \(dx\) and the concentration of \(x\) will appear in the melt. After that, the microstructure during isothermal solidification will evolve. Since the concentration fluctuations in the melt always exist during solidification, the nucleus can form continuously as well.

Next, the proposed nucleation model is applied in a real alloy together with the phase-field simulation. Here, one hypoeutectic alloy, Al-2.89 at. % Si, was chosen as the target, and the phase-field model with finite interface dissipation \(^{8}\) was utilized in the present work. The evolution equation of the phase field and individual phase concentrations are given \(^{8}\):

\[
\frac{\partial \phi_\alpha}{\partial t} = \nabla \cdot \left( \phi_\alpha D_\alpha \nabla c_\alpha \right) + P \phi_\alpha \phi_\beta \frac{\partial f_\alpha}{\partial c_\beta} - \frac{\partial f_\alpha}{\partial c_\alpha} + \phi_\alpha \phi_\beta (c_\beta - c_\alpha)
\]

\[
\phi_\beta \frac{\partial \phi_\beta}{\partial t} = \nabla \cdot \left( \phi_\beta D_\beta \nabla c_\beta \right) + P \phi_\alpha \phi_\beta \frac{\partial f_\beta}{\partial c_\beta} - \frac{\partial f_\beta}{\partial c_\alpha} + \phi_\alpha \phi_\beta (c_\alpha - c_\beta)
\]

where \(c_\alpha\) and \(c_\beta\) are the compositions of \(\alpha\) and \(\beta\) phases, while \(D_\alpha\) and \(D_\beta\) are the chemical diffusivities of \(\alpha\) and \(\beta\) phases. The solid-liquid interface energy and interface mobility during solidification are strongly anisotropic and thus the anisotropy in the phase-field simulation of the dendritic growth needs to be taken into account. The anisotropy equation used in the phase-field simulation is as follows \(^{13}\):

\[
\sigma^+ = \sigma(1 + 3e_\sigma - 4e_\sigma (n_4^2 + n_5^2 + n_6^2))
\]

\[
\mu^+ = \mu(1 - 3e_\sigma + 4e_\sigma (n_4^2 + n_5^2 + n_6^2))
\]

where \(\sigma\) and \(\mu\) are the average interface energy and interface mobility, while \(e_\sigma\) and \(e_\sigma\) are the anisotropic coefficients of interface energy and interface mobility. The anisotropic coefficients of both interface energy and interface mobility were set to be 0.4. Here, \(n_4, n_5, n_6\) are the normal vectors in the direction of the axis. All the material and
numerical parameters used in the present three-dimensional (3-D) phase-field simulation are listed in Table 2, except for the values of the interface permeability $P$. The interfacial energy $\sigma$ was directly adopted from14, while the interface mobility $M$ was calculated according to11 for keeping in the diffusion-controlled regime. $D_s$ and $D_l$ were simply set to be constants15. The simulation domain was chosen to be of $128 \times 128 \times 128$ grid points. The boundary conditions for all the phase fields and concentrations were set to be periodic.

First, the impact of permeability on the nucleation process is explored, as already explained above, and variation of permeability will lead to changes in the nucleus concentration and the nucleation driving force, even if the melt concentration is the same. Furthermore, the concentration distribution in the melt is represented in Fig. 5 (Note that here it is only intended to exhibit the practicability of the model, not to provide a result with high precision), and the dashed lines corresponding to different permeabilities are the calculated nucleation driving force curves. It is apparent that the nucleation driving force will be inhibited with the reduction of permeability, hence the change of permeability has noticeable impact on the nucleation events.

Then, the formation of the nucleus is followed by its dendritic growth. The 3-D phase-field simulation for grain growth evolution in Al-2.89 at. % Si alloy corresponding to different permeabilities are enumerated in Fig. 6. Figure 6(a) shows the morphology evolution of microstructure while Fig. 6(b) displays the concentration evolution of the nucleus. Obviously, the permeability has a great influence on the initial concentration of nucleus whereas it is almost impervious for both the grain growth and the subsequent concentration evolution of nucleus.

Table 2. List of the numerical and materials parameters used in the present phase-field simulations.

| Parameters          | Symbols | Values                |
|---------------------|---------|-----------------------|
| grid spacing        | $\Delta x$ | 50 nm                 |
| interface width     | $\eta$  | $4\Delta x$           |
| interface energy    | $\sigma$ | $1.69 \times 10^{-5}$ J/cm$^2$ |
| interface mobility  | $M$     | $1.23 \times 10^{-1}$ cm$^4$/Js |
| diffusivity of liquid phase | $D_l$ | $1 \times 10^{-7}$ cm$^2$/s |
| diffusivity of solid phase | $D_s$ | $1 \times 10^{-9}$ cm$^2$/s |

Figure 5. Concentration distribution in the melt and nucleation driving force with respect to different permeabilities.

Then, solidification processes with different interface permeabilities (i.e., 1000 cm$^3$/Js, 5000 cm$^3$/Js and 10000 cm$^3$/Js) in Al-2.89 at. % Si alloy were simulated by the 3-D phase-field model and are exhibited in Fig. 7. From the three evolutionary graphs it can be seen clearly that the nucleus are precipitated continuously with the liquid concentration undulating, nucleation is almost impossible to occur in Fig. 7(a) while the occurrence of nucleation is very easy in Fig. 7(c), thus it can be said without exaggeration that a high interface permeability is likely to increase the total number of nucleation events, fully affirmed our speculation.

Next, the nucleation rate curves corresponding to different permeabilities are fitted by a recently modified function similar to the LSW size distribution16:

$$f(n) = a \cdot \left(\frac{1}{1 - \tau}\right)^b \cdot \left(\frac{1}{1 + \tau}\right)^c \cdot \exp\left(-\frac{1}{1 - \tau}\right)$$  \hspace{1cm} (15)

Here, $a$, $b$, $c$ are the adjustable parameters, the nucleation rate ($f(n)$) and the time ($\tau$) are normalized values respectively (Here, the total number of nucleus are $N_{P=1000}^{total} = 5$, $N_{P=5000}^{total} = 67$ and $N_{P=10000}^{total} = 1175$ respectively). As predicted in Fig. 8, the nucleation rate shows an upward trend in the beginning because a small number of nucleus will precipitate faster in the place where the driving force is larger. Then the nucleation rate shows a
downward trend. The reason for this phenomenon may be that, the saturation of melt will increase with the growth of crystals, therefore the subsequent nucleation potency will be restrained.

Besides, the time to reach the peak and the quantities of nucleus are affected seriously by the permeability as well. The average volume of the grains is sketched schematically in Fig. 9, calculated by:

**Figure 6.** Grain growth evolution corresponding to different permeabilities during isothermal solidification (a) morphology evolution; (b) concentration evolution of the nucleus.

**Figure 7.** Three-dimensional phase-field simulated nucleation process of Al-2.89 at. % Si alloy during isothermal solidification with different interfacial permeabilities: (a) \( P = 1000 \text{ cm}^3/\text{Js} \); (b) \( P = 5000 \text{ cm}^3/\text{Js} \); (c) \( P = 10000 \text{ cm}^3/\text{Js} \).

**Figure 8.** Normalized nucleation rates corresponding to different interfacial permeabilities.
\[ V = \frac{V_{\text{total}}}{N} \]  
(16)

where \(V_{\text{total}}\) is the total volume of the solid phase. It can be seen clearly that at the beginning a larger permeability will result in a larger average grain volume, however, as the evolution proceeds, the average grain volume for the largest permeability is smaller compared to the other two smaller permeabilities. A feasible explanation may be that when the nucleus density is high, the growth of grains is inhibited by other grains, and with the growth of enormous amounts of nucleus, solute atoms will be released into the melt, leading to a saturation rapidly which will restrain the grain growth as well. Thus, it can be concluded that the increase in permeability can accelerate the process of solidification mainly through the promoted occurrence of nucleation.

**Conclusions**

A kinetic view into the Hillert-Rettenmayr thermodynamic analysis was performed, and demonstrated to determine nucleus composition during isothermal solidification, which still represents a challenge nowadays. With the introduction of the interface permeability, the energy dissipation of the solute rearrangement at the liquid-solid interface can be evaluated, and the driving force for nucleation can be unambiguously determined.

A pragmatic nucleation model was proposed, and then validated using a 3-D phase-field simulation of nucleation and subsequent dendritic growth in one hypothetic Al-Si alloy. The simulation results indicate that the permeability affects the nucleation driving force of metastable clusters by influencing their composition, which has a great effect on the nucleation rate and finally affects the entire solidification process.

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Acknowledgements
The financial support from the National Natural Science of Foundation of China (Grant No. 51474239) and Project of International Cooperation and Exchanges from the National Natural Science Foundation of China (Grant No. 51611130058) is acknowledged. Lijun Zhang acknowledges the Huxiang Youth Talent Plan released by Hunan Province, China, and the project supported by State Key Laboratory of Powder Metallurgy Foundation, Central South University, Changsha, P.R. China.

Author Contributions
L.Z. conceived the idea for this paper. L.Z. and X.K. conducted the theoretical analysis. X.K. performed the three-dimensional phase-field simulation. X.K., L.Z. and S.S. wrote the manuscript. All authors reviewed and commented on the manuscript at all stages.

Additional Information
Competing Interests: The authors declare no competing interests.

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