

Modeling and numerical simulations of dendritic crystal growth

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A simple phase field model for one component melt growth is presented, which includes anisotropy in a certain form. The formation of various dendritic patterns can be shown by a series of numerical simulations of this model. Qualitative relations between the shapes of crystals and some physical parameters are discussed. Also it is shown that noises give a crucial influence on the side branch structure of dendrites in some situations.

1. Introduction

Crystal growth in one of the most fascinating phenomena of spontaneous pattern formations in nature. For instance, snowflakes have various types of very beautiful and complicated shapes, although they are formed in almost uniform circumstances. Also dendritic structure is commonly seen in solidification of metals [14] or crystalization in supersaturated solutions [11,12]. It's an exciting problem how such complex patterns can be formed by the systems which seem to be too simple to yield them. The aim of the present paper is to give a simplest model which can describe the dendritic pattern formations. There might be some fluid dynamical effects like the convection induced by gravity or Marangoni effect which give influences on the growth rate. Also the expansion or shrinking caused by the phase transition may affect the crystallization process. But these are not considered here, because our approach is not to present a model describing the phenomena minutely but to show the minimal set of factors which can yield variety

of typical dendritic patterns essentially. For this reason, all the computations and all the discussions in this paper are devoted to the morphology of the crystals.

There are two types of the crystal forms; such as equilibrium and growth. The equilibrium forms are determined so as to minimize the surface energy and therefore the shape is a sphere if the surface energy is isotropic and is some polyhedron if anisotropy exists. For this meaning, dendrites are not equilibrium forms but growth ones, which is formed under the balance of the surface tension and the thermodynamical driving force, say supercooling or supersaturation. However various dendritic patterns can not be obtained by only these two factors. Anisotropy, as is shown later, affects the shape of the crystal drastically. In our model, we consider the growth process that the only one diffusion field limits the growth rate of crystals, and I concentrate on the melt growth of pure material, in which there are molecules sufficiently dense in the surrounding phase (liquid phase) and they attach to the solid/liquid interface collectively. Therefore the process limiting the growth rate of the crystal is a removing process of latent heat from the interface, which keeps the interfacial temperature below the equilibrium temperature.

From the computational view point, we have two ways to handle such moving boundary problems. The one is to keep the data of the interfacial position explicitly and move it due to the interfacial equation of motion. (See [1-4] for example.) But we don't take this kind of method here because the complicated manipulation is needed when the topology of the interface changes - merging or coming off, and also it must be hard to extend such methods to the three dimensional space. Thus we'll adopt the other way, in which the interface is expressed implicitly using some function defined on the whole region. One of such methods was proposed by Osher and Sethian [5], in which the interface is expressed as a contour line (or surface) of some function and it is driven by Hamilton Jacobi equation. This is a numerical technique to solve the interface equation in which the interface thickness is considered to be zero.

The model that we introduce here is a kind of phase field model in which the interface has a finite thickness although it is very thin and is expressed as a steep internal layer of a phase indicating function. The typical isotropic version of such model was proposed by Fix [6] and Collins and Levin [7], also it was justified mathematically by Caginalp [8] and Caginalp and Fife [9]. Also Caginalp presented relations between the phase field model and its singular limit equations [10]. A similar model including anisotropy was introduced and proved to be able to describe realistic dendritic patterns numerically Kobayashi [15] and Mimura, Kobayashi and Okazaki [16]. In the following section, it will be shown that the model equations introduced in [15,16] can exhibit various realistic dendritic patterns by the numerical simulations. And the relations between the growth forms and some physical parameters such as dimensionless latent heat or strength of anisotropy are discussed. This article is essentially an explanation of the simulations presented in [17].

2. Model

The model includes two variables; one is a phase field p(r, t) and the other is a temperature field T(r, t). The variable p(r, t) is an ordering parameter at the position r and the time t, p = 0 means being liquid and p = 1 solid. And the solid/liquid interface is expressed by the steep layer of p connecting the values 0 and 1. Fig. 1 shows how the shape of crystal is described by the phase field p. In order to keep the profile of p such form and to move it reasonably, we consider the following Ginzburg-Landau type free energy including m as a parameter:

$$\Phi[p;m] = \int \frac{1}{2} \varepsilon^2 |\nabla p|^2 + F(p;m) dr,$$

where ε is a small parameter which determines the thickness of the layer. It is a microscopic interaction length and it also controls the mobility of the interface. F is a double-well potential which has local minimums at p=0 and 1 for each m. Here we take the specific form of F as follows:

$$F(p;m) = \frac{1}{4}p^4 - (\frac{1}{2} - \frac{1}{3}m)p^3 + (\frac{1}{4} - \frac{1}{2}m)p^2,$$

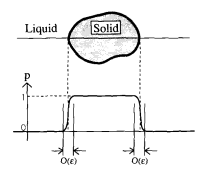


Fig. 1. Expression of the solid/liquid interface by the phase indicating function p.

where $|m| < \frac{1}{2}$. The profile of F and its dependency on m are shown in fig. 2. The difference of the two minimum values is proportional to m, which gives a difference of chemical potentials of both the phases.

Anisotropy can be introduced by assuming that ε depends on the direction of the outer normal vector at the interface. So ε is represented as a function of the vector $\mathbf{v} = (\mathbf{v}_i)$ satisfying $\varepsilon(\lambda \mathbf{v}) = \varepsilon(\mathbf{v})$ for $\lambda > 0$. The outer normal vector is represented by $-\nabla p$ at the interface. Thus we consider

$$\Phi[p;m] = \int \frac{1}{2} \varepsilon (-\nabla p)^2 |\nabla p|^2 + F(p;m) \, \mathrm{d}r \,. \quad (1)$$

From the formula $\tau \partial p/\partial t = -\delta \Phi/\delta p$, we have the following evolution equation:

$$\tau \frac{\partial p}{\partial t} = -\nabla \cdot \left(|\nabla p|^2 \varepsilon \frac{\partial \varepsilon}{\partial v} \right) + \nabla \cdot (\varepsilon^2 \nabla p) + p(1-p)(p - \frac{1}{2} + m),$$
(2)

where τ is a small positive constant and $\partial \varepsilon / \partial v = (\partial \varepsilon / \partial v_i)_i$. The parameter m gives a thermodynamical driving force. Especially in two dimensional space, we can take $\varepsilon = \varepsilon(\theta)$ where θ is an angle between v and a certain direction (for example the positive direction of the x-axis). Thus we have

$$\tau \frac{\partial p}{\partial t} = -\frac{\partial}{\partial x} \left(\varepsilon \varepsilon' \frac{\partial p}{\partial y} \right) + \frac{\partial}{\partial y} \left(\varepsilon \varepsilon' \frac{\partial p}{\partial x} \right) + \nabla \cdot (\varepsilon^2 \nabla p) + p(1-p)(p - \frac{1}{2} + m) ,$$
(3)

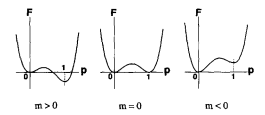


Fig. 2. Double-well potential F controlled by the parameter m. $\Delta F = F(0; m) - F(1; m) = m/6$.

where the prime means $d/d\theta$. Although (2) and (3) have a field equation form, they can be called interface equations, since they have a practical meaning only in the neighborhood of the interface. In fact, every term of them almost equals to zero in bulk.

In order to understand intuitively how eq. (3) moves the interface, let us derive the singular limit equation in two dimensional case. Put $\varepsilon = \bar{\varepsilon}\sigma(\theta)$ where $\bar{\varepsilon}$ is a mean value of ε and $\sigma(\theta)$ represents anisotropy. Taking $\tau = b\bar{\varepsilon}$ and $m = \bar{\varepsilon}f/\sqrt{2}$, we obtain an interface equation in the limit of $\bar{\varepsilon}$ tending to zero, as follows (See appendix):

$$bV = \sigma(\theta)f - [\sigma(\theta)^2 + (\frac{1}{2}\sigma(\theta)^2)'']\kappa , \qquad (4)$$

where V is the normal velocity and κ is the mean curvature of the interface. Note that the limiting equation (4) is defined only on the interface while (3) is defined on the whole region. By taking τ and m as stated above, we can regard eq. (3) as a family of equations parametrized by $\bar{\varepsilon}$. Thus the parameter $\bar{\varepsilon}$ is not a physical parameter but a computational parameter corresponding to the thickness of the layer which expresses an interface, or an indicator how well (3) approximates (4). Eq. (4) includes thermodynamical driving force, surface tension effect and anisotropy. Therefore eq. (3) also contains these factors. Anisotropy expressed by (4) will be discussed later.

Here we assume that m is a function of the temperature T, for example, $m(T) = \gamma(T_e - T)$ where T_e is an equilibrium temperature, which means that the driving force of interfacial motion is proportional to the supercooling there. But in the following simulations, we used the form $m(T) = (\alpha/\pi) \tan^{-1} [\gamma(T_e - T)]$ where α is a positive constant satisfying $\alpha < 1$, since this assures $|m(T)| < \frac{1}{2}$ for all values of T. Also m(T) is almost linear for T near T_e . To take anisotropy into account, let us specify σ to be

$$\sigma(\theta) = 1 + \delta \cos[j(\theta - \theta_0)].$$

The parameter δ means the strength of anisotropy and j is a mode number of anisotropy.

The equation for T is derived from the conservation law of enthalpy as

$$\frac{\partial T}{\partial t} = \nabla^2 T + K \frac{\partial p}{\partial t} . \tag{5}$$

cT is non-dimensionalized so that the characteristic cooling temperature is 0 and the equilibrium temperature is 1^{*1} . K is a dimensionless latent heat which is proportional to the latent heat and inversely proportional to the strength of the cooling. For simplicity, the diffusion constant is set to be identical in both of solid and liquid regions. (5) is a heat conduction equation having a heat source along the moving interface, since $K \partial p/\partial t$ has non-zero value only when the interface passes through the point.

Here we refer to the difference between the dynamical term of our model equation (2) or (3) and the one of common phase field model (see [6–10]). In the common models, the dynamical term has a summation form essentially like $p(1-p)(p-\frac{1}{2}) + \text{const.} \times (T_e - T)$. Then the equilibrium value of p depends on the value of T, while they are fixed to be 0 or 1 in our model. Thus if the temperature of a point in bulk changes (in fact, it may change from the cooling temperature to one near T_e in supercooling solidification), the value of p at that point varies corresponding to the temporal change of T. Consequently the term $K \partial p/\partial t$ has non-zero value in bulk, which means that latent heat is released there. Therefore we should adopt the form of dynamical term such that the equilibrium values of p are fixed in bulk, if we use $K \partial p/\partial t$ as a latent heat term. Otherwise we should change the latent heat term $K \partial p/\partial t$ to $K \partial g(p)/\partial t$, where g(p) is a non-decreasing smooth function satisfying $g(p) \equiv 0$ near p = 0 and $g(p) \equiv 1$ near p = 1.

3. Simulations

In this section, two dimensional simulations are presented which reveal various types of formations of dendritic pattern. All the simulations were computed by using the model equations (3) and (5). Dimensionless latent heat K, strength of anisotropy δ and mode number of anisotropy j are changed, and other parameters are fixed #2. The characteristic cooling temperature was taken as the temperature of the cooling wall in the simulation without supercooling, and the initial temperature of the liquid in the supercooling solidification. Note that small noises were added on the interface all through the computations. This might be somewhat realistic and we can examine the stability of the shape of the interfaces against the noise. Noises were given to the system by using a random number sequence χ in the form $ap(1-p)\chi$, where a is an amplitude of the noise and χ is uniformly distributed on the interval $\left[-\frac{1}{2}, \frac{1}{2}\right]$. This term was added to the dynamical term of (3) on the whole region, which is equivalent to adding the noises $a\chi$ to the term m(T) at the interface. Denote that the boundary condition of p is always the zero flux condition. In all the simulations, the parameters are taken as follows: the domain size is 9.0×9.0 for the square region and 12.0×3.0 for the rectangular region, and the mesh numbers are 300×300 and 400×100 , respectively. The fixed parameters are evaluated as follows, $\bar{\varepsilon} = 0.01$, $\tau = 0.0003$, $\alpha = 0.9$, $\gamma = 10.0$ and a = 0.01. Time mesh size is taken to be 0.0002. In the following simulations, no special technique was used for solving eqs. (3) and (5). A simple explicit scheme is used for (3) and a simple implicit scheme for (5). Note that the last picture of each figure does not mean the last state of the process.

^{*1} The parameter γ in m(T) is non-dimensionalized corresponding to the non-dimensionalization of T. Hereafter we use the same notation γ for the dimensionless value.

^{*2} To control the values of K fixing the one of γ means to consider various materials with different values of latent heat, not to control the strength of supercooling for one material.

3.1. No supercooling – isotropic

Initially the vessel is filled with a liquid of equilibrium temperature, and the vessel is cooled from its walls simultaneously. Solidification occurs from the walls and the crystal grows inward. Latent heat is taken away through the solid phase. The result of the simulation is shown in fig. 3. The shape of the interface is rounded as it moves and disappears, in the end. It corresponds to making an ice cube in an ice tray. Next we examine the stability of the advancing flat interface. The vessel is filled with liquid of equilibrium temperature, and along the left wall the solid exists and its interface is deformed as fig. 4 (time = 0.0) although it is rather artificial. The vessel is cooled from the left wall, and no heat flux is allowed at other walls. The crystal grows to the right, and latent heat is taken away through the solid phase to the left. The shape of the interface becomes flat very quickly as shown in fig. 4. In both cases, the interface never becomes complicated, and it acquires a shape with higher symmetry. This is because not only the surface tension but also the temperature field make the shape smooth and simple. In the anisotropic case, the situation is essentially the same,

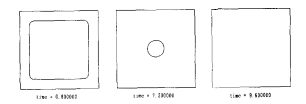


Fig. 3. Crystal grows inward from the walls. The system is cooled by the surrounding walls. Isotropic. K = 1.0.

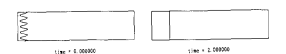


Fig. 4. Crystal grows from the left wall to the right. The system is cooled only by the left wall. Isotropic. K = 1.0.

although the results of simulations are not shown here. These are not interesting from the point of view of pattern formation (not of practical industry). We are interested in the complicated pattern formation like dendrites. In the following simulations, we consider the supercooling solidification.

In the subsequent simulations, the whole region is initially filled with uniformly supercooled melt (the initial temperature is taken to be zero as stated before). Then the nucleation occurs and the solidification process follows under adiabatic conditions. In such situation, the whole liquid will change to crystal for K < 1. If K > 1, 1/K of the whole region will be solidified, and then lose the supercooling in the end. After that the interface moves only by the surface tension, thus very slowly.

3.2. Directional solidification – isotropic

Solidification occurs simultaneously from the left wall to the right. Fig. 5 shows the results of the simulations for the various values of K. The flat interface is stable for K = 0.8 and a little bit destabilized for K = 0.9. A weak cellular structure is observed for K = 1.0 and 1.1. For the latter parameter the slits between cells sometimes remain behind the front and they become holes by surface tension. For $K \ge 1.2$, tip splitting occurs and the splitting branches tend to spread widely. But there is a limited space in the channel region and therefore they have to compete with each other. Consequently the branches differentiate to the winner and the loser. The loser cannot grow any more because it is screened by the winner. The winners can continue to grow and the tip splitting occurs at the top of them, then the new branches compete with each other again. This process succeeds until the front reaches the right wall. As K becomes larger, the branches come to have a stronger tendency to spread widely. The growth rate is drastically decreasing for $K \ge 1.6$.

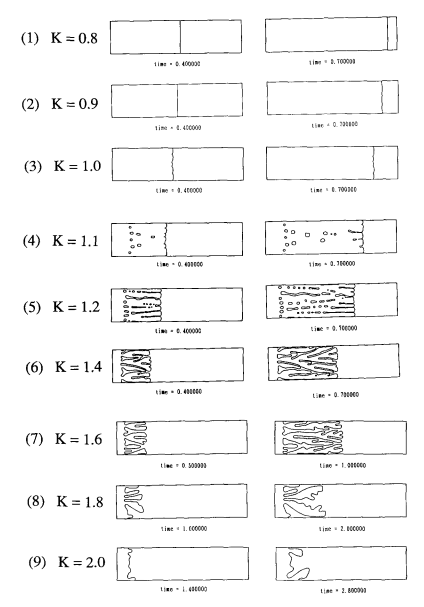


Fig. 5. Crystal grows from the left wall toward supercooled melt under adiabatic condition for various values of K. Isotropic.

3.3. Directional solidification-anisotropic

Next the 4-mode anisotropy is assumed; δ is fixed to be 0.050 and the horizontal and the vertical directions, are the ones at which ε has maximums (i.e. j=4 and $\theta_0=0$). The initial state is just the same as in the isotropic case. Fig. 6 shows results. For K=0.8, the flat interface is

already destabilized slightly, whereas it was stable in the isotropic situation. For K = 0.9 and 1.0, weak cellular structure is seen. For K = 1.1, 1.2 and 1.4, slits are observed and they remain a long time because the branches have almost no tendency to spread. And their fronts advance faster than the ones of the istropic case. For $K \ge 1.6$, we can see that the branches compete

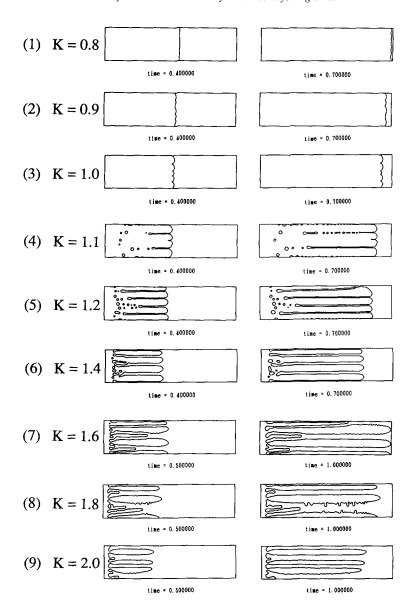


Fig. 6. Crystal grows from the left wall toward supercooled melt under adiabatic condition for various values of K. There is 4-mode anisotropy. $\delta = 0.050$.

with each other in a way different from the isotropic one. The branches going faster than the adjacent ones suppress the growth of the slower ones by becoming thick or having side branches. The growth rates are completely larger than the isotropic growth. The branches sometimes come off at their root by surface tension.

3.4. Dendrite growth

In the following simulations, we will show the relations between the strength of anisotropy and the shapes of crystals. Nucleation occurs at the center of the bottom edge, and it triggers the growth process. All the parameters except δ are

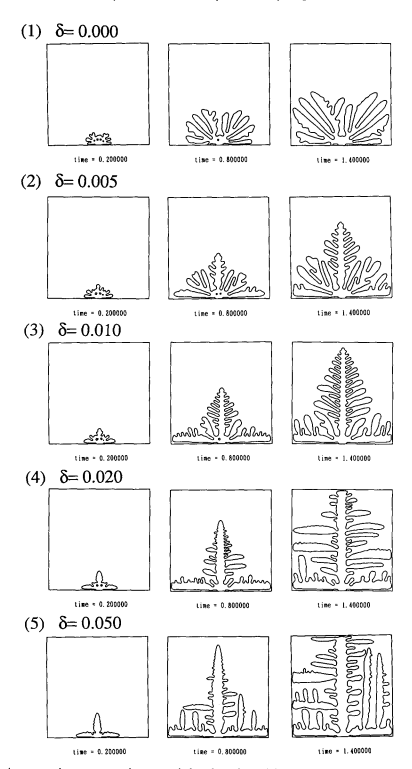


Fig. 7. After nucleation, crystal grows toward supercooled melt under adiabatic condition for various values of δ . There is 4-mode anisotropy. K = 2.0.

fixed and δ is increased gradually from zero. The horizontal and vertical direction is the one at which ε has maximum value (i.e. j=4 and $\theta_0=0$). Dimensionless latent heat K is fixed to be 2.0. The results are shown in fig. 7.

In the first computation, where $\delta = 0.000$ i.e. perfect isotropic growth is considered, patterns similar to the viscous fingering are obtained. Tip splitting is seen as the crystal grows, and roughly speaking, the shapes of the curve which connects the tops of branches are half circles until the interaction between branches and walls occurs. The shape of the crystal for $\delta = 0.005$ has both features of the isotropic and of the dendritic structure. We can see the dendritic structure in the vertical branch and the viscous finger-like structure in the other branches. The results for $\delta = 0.010$ show one typical dendritic structure whose side branches are shifted from the direction of anisotropy. Roughly speaking, their direction is perpendicular to the isothermal lines,

since it is the easiest direction to grow and the anisotropy is weaker than this effect. Side branches are formed at the top of the principal branch having the constant spacing. For δ less than 0.010, we sometimes observe the mutual side branching. This non-symmetric side branching occurs more often for smaller values of δ . For $\delta = 0.020$, we can see other typical dendritic structures whose side branches almost coincide with the direction of anisotropy. They occur from the destabilization of the side surface of the principal branch near the top. For $\delta = 0.050$, the process of forming side branches is the same as the previous one, but the side branch structure is a little different from it. Competitions and screening clearly occur between the branches which have perpendicular directions. And also when the side branch grows rather longer than other side branches, new side branches grow from it.

Comparing these simulations, it is obvious that

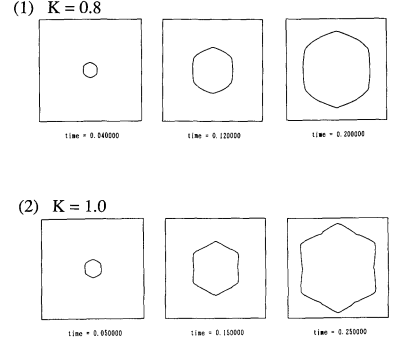


Fig. 8. After nucleation, crystal grows toward supercooled melt under adiabatic condition for various values of K. There is 6-mode anisotropy. $\delta = 0.040$.

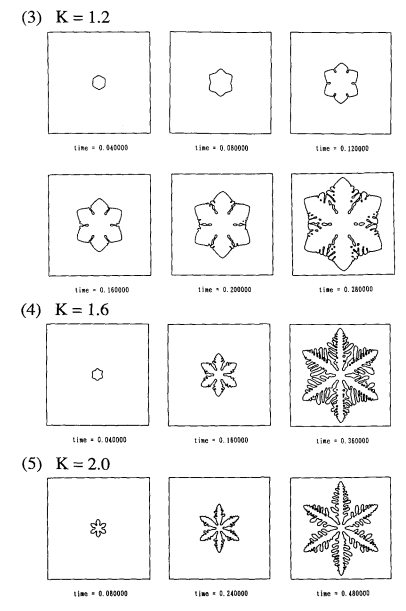


Fig. 8 (cont.).

the velocity of the principal branch increases as δ increases. Also it is clear that the branch structure is very sensitively dependent on δ .

3.5. Ice dendrites

At last, snowflake-like patterns are presented in fig. 8. Although patterns observed here are

similar to snowflakes, the model equations (3) and (5) do not describe the formation of snow-flakes, since snowflakes are formed by vapor growth which has a different process from melt growth. In order to simulate such a process, the formation of a facet and its destabilization should be considered [19]. So the following simulations correspond to the supercooling solidifica-

tion of water. Nucleation occurs at the center of the vessel and the solidification follows. In these simulations, only the parameter K is changed and the other parameters are fixed as $\delta = 0.040$, j = 6 and $\theta_0 = \frac{1}{2}\pi$.

First, the crystal is hexagon which is strictly convex for K = 0.8. But as K increases, the convexity is broken. In fact, for K = 1.0, we can find the dents at the center of the crystal edges. For K = 1.2, more pictures are shown than for the other parameters, since it presents various patterns during its growth process. At first we can see a hexagonal shape and then the dents at the centers of the edges. After that the dents become slits and next the tuck appears at the top of the slit. At last it shows a thick branching pattern, which is an intermediate pattern between the essentially hexagonal patterns and the branching patterns. For K = 1.6, the crystal shows a branching pattern. As K increases, the branches become thinner and we can obtain a typical snowflake-like pattern for K = 2.0. Thus we can observe the change from the hexagonal shape to the branching patterns as K increases.

4. Discussions

One may feel curious that the shape of the crystal goes simpler as K decreases, as shown in figs. 5, 6, and 8. It is because these simulations are in the parameter area in which K is close to one. (See [13].)

Fig. 9 shows a typical dendrite whose side branches' direction almost coincides with the direction of anisotropy ($\delta = 0.020$). As a rough sketch, we can distinguish two regions by connecting the tops of the side branches, region I and region II. In region I, all of the young side branches are growing and restrain each other, thus the growth rate is small. In region II, the competition is already over and the winners grow faster than the branches in region I, because they can have more cooling than the branches in region I. It is just like the situation where the thinned out plants grow faster by obtaining more light or nourishment. While the growth rates of the side branches are different in regions I and II, the velocity of the top of the principal branch is almost constant. Thus we have the two lines of different slopes.

Here let us consider the influence of noises. If the noises are larger, the competition between the young side branches may end earlier because the noises are partial to some branches, then region I becomes smaller. If the noises are smaller, it takes a longer time to select winners and therefore region I will be larger. This is proved by fig. 9, in which the noises are added in the different amplitude a. The parameter a is becoming smaller from left to right. Region I clearly goes larger as a decreases. Comparing the pictures in fig. 9 we have the essential two properties. The one is that the velocity of the top of the main branch is exactly the same in each case. And the other is that the side branch structure is

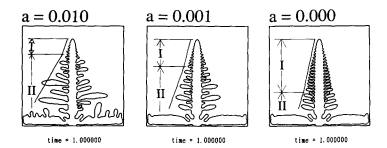


Fig. 9. The relation between the side branch structure and the amplitude of noise in the non-oscillating growth. K = 2.0 and $\delta = 0.020$.

strongly affected by the existence or amplitude of the noises.

Fig. 10 shows crystal forms for $\delta = 0.010$, in which the parameter a is controlled in the same way as in fig. 9. Their side branch structures are less affected by the change of the amplitude of noise than the ones in fig. 9. It is because the growth rate of the top of the principal branch oscillates in fig. 10. while it does not in fig. 9 [18]. As the oscillation is a strong mechanism for making side branches, they are less affected by noises. In fig. 9, the side branches are strongly affected by noises, since they come from the destabilization of the side surface of the principal branch which is a more delicate process than the oscillation.

Here let us consider the singular limit equation (4), which can be deformed to

$$bV = \sigma(\theta) \left[f - d_0(\theta) \kappa \right], \tag{6}$$

where $d_0(\theta) = \sigma(\theta) + \sigma''(\theta) + [\sigma'(\theta)]^2/\sigma(\theta)$. When δ is small, $(\sigma')^2/\sigma$ is negligible since it is of order δ^2 , while $\sigma + \sigma''$ is $1 + \mathcal{O}(\delta)$. For all the values of δ that I took in the simulations here $(\delta \le 5 \times 10^{-2})$, $(\sigma')^2/\sigma$ is completely smaller than $\sigma + \sigma''$. Fig. 11 shows graphs of $\sigma(\theta)$ and $d_0(\theta)$ for the parameters used in the simulations corresponding to fig. 7 (5) and fig. 8. If $d_0(\theta)$ is always positive (of course, δ values for fig. 7(1)–(4) are smaller than the one of (5), thus $d_0(\theta)$ is always positive for all the simulations in fig. 7), no corner appears along the interface as shown

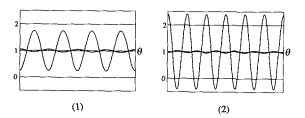


Fig. 11. The curve with smaller amplitude is a graph of $\sigma(\theta) = 1 + \delta \cos \left[j(\theta - \theta_0) \right]$ and the one with larger amplitude $d_0(\theta)$. (1) $\delta = 0.050$, j = 4 and $\theta_0 = 0$, corresponding to fig. 7(5). (2) $\delta = 0.040$, j = 6 and $\theta_0 = \pi/2$, corresponding to fig. 8.

in fig. 7. If $d_0(\theta)$ is negative for some interval of θ , we can observe the formation of corners as shown in fig. 8. The parameter δ gives crucial influence on the growth form of crystal even if it is rather small, because $d_0(\theta)$ includes a term σ'' which magnifies anisotropy by m^2 times in our setting of anisotropy.

As shown in the previous part, the model equations (3) and (5) can exhibit various solutions which represent the crystal growth realistically, although they are obtained for only a small part of the total range of the parameter values. Thus we would like to conclude that eqs. (3) and (5) are a very good model for simulating dendritic crystal growth.

But it still has some problems. The phase field p cannot describe a structure which is finer than or the same size as the layer thickness. So we must take ε sufficiently small if we want to simulate the fine structure. The computation does not go well if the physical parameters re-

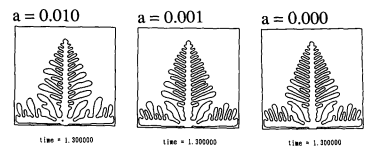


Fig. 10. The relation between the side branch structure and the amplitude of noise in the oscillating growth. K = 2.0 and $\delta = 0.010$.

quire a finer structure than the size which the phase field can express with given ε . For example, as seen in fig. 6(4), the narrow slits are buried easier than the real physical system, since the interfacial thickness is not so small compared with the width of slits. One more problem is a computational one, which comes from the fact that (3) includes small parameters. The values of $\varepsilon/\delta x$ and $\tau/\delta t$ (δx is a space mesh size and δt is a time mesh size) should be taken large in order to obtain good approximations quantitatively. But in the limited CPU power, we cannot take space mesh size and time mesh size so small. In fact, in the simulations presented here, we selected the space mesh size small such that anisotropy induced by the computational lattice does not appear, but not so small such that the interfacial velocity of (3) is precisely approximated. This is because we intended to see the qualitative properties of the whole crystal structure here. If we want quantitatively precise results, we should restrict the size of the computational region and take the space mesh size and the time mesh size small enough.

Appendix

Eq. (3) is rewritten as

$$b\bar{\varepsilon}^2 \frac{\partial p}{\partial t} = \bar{\varepsilon}^2 \left(-\frac{\partial}{\partial x} \left(\frac{1}{2} \sigma^2 \right)' \frac{\partial p}{\partial y} + \frac{\partial}{\partial y} \left(\frac{1}{2} \sigma^2 \right)' \frac{\partial p}{\partial x} \right.$$
$$+ \nabla \cdot (\sigma^2 \nabla p) + p(1 - p(p - \frac{1}{2} + \bar{\varepsilon}f/\sqrt{2}),$$

where the prime means $d/d\theta$. Here let us assume the axi-symmetric solution p, and deform the terms in the bracket $\{\ \}$ using polar coordinates as follows:

$$-\frac{\partial}{\partial x} \left(\frac{1}{2}\sigma^2\right)' \frac{\partial p}{\partial y} + \frac{\partial}{\partial y} \left(\frac{1}{2}\sigma^2\right)' \frac{\partial p}{\partial x} = \left(\frac{1}{2}\sigma^2\right)'' \frac{1}{r} \frac{\partial p}{\partial r} ,$$

$$\nabla \cdot (\sigma^2 \nabla p) = \sigma^2 \left(\frac{\partial^2 p}{\partial r^2} + \frac{1}{r} \frac{\partial p}{\partial r}\right) .$$

Thus (3) is deformed to

$$b\bar{\varepsilon}^2 \frac{\partial p}{\partial t} = \bar{\varepsilon}^2 \left(\sigma^2 \frac{\partial^2 p}{\partial r^2} + \frac{1}{r} \left[\sigma^2 + (\frac{1}{2}\sigma^2)'' \right] \frac{\partial p}{\partial r} \right) + p(1-p)(p-\frac{1}{2} + \bar{\varepsilon}f/\sqrt{2}).$$

Stretching the coordinate r near the interface by the factor $\bar{\epsilon}\sigma$ and assuming the traveling front solution in the stretched interval, we have

$$\frac{\mathrm{d}^2 p}{\mathrm{d}\eta^2} + \frac{\bar{\varepsilon}}{\sigma} \left\{ bV + \left[\sigma^2 + \left(\frac{1}{2} \sigma^2 \right)'' \right] k \right\} \frac{\mathrm{d}p}{\mathrm{d}\eta} + p(1-p)(p-\frac{1}{2} + \tilde{\varepsilon}f/\sqrt{2}) = 0,$$

where η is a traveling coordinate in the stretched interval. Solving this non-linear eigenvalue problem (see [20]), we have (4) as a singular limit equation. For the general case, we can obtain the same result by taking the polar coordinate system for each point on the interface such that the origin coincides with the center of curvature.

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