Modeling of tip kinetics of undercooled Ti dendrites with consideration of forced flow and oxygen impurity effects

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Abstract

Tip velocities of undercooled dendrites in electromagnetically levitated melt droplets of pure Ti were modeled using a theory on three-dimensional dendritic growth with fluid flow and a dilute solute. The modeling shows that a forced flow due to electromagnetic stirring depresses tip velocities of the dendrites at low undercoolings and that its effect becomes negligible at high undercoolings. In contrast, an oxygen impurity dissolved in liquid Ti depresses tip velocities of the dendrites over a broad range of undercooling while it coarsens tip radii. Such modeling results allowed for reconciliation of discrepancies in literature data by considering an oxygen impurity effect. The modeling also predicts that the effect of the oxygen impurity becomes insignificant when its concentration is reduced below 50 ppm in atomic fraction.

Keywords: Dendrites; tip kinetics; modeling; liquid flow; oxygen impurity

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1. Introduction

Metal additive manufacturing technologies have received extensive attention because of advantages in producing geometrically complex or chemically graded parts over traditional technologies (see a recent review [1,2]). There is growing interest in an understanding of non-equilibrium solidification kinetics of a small melt pool for the purpose of online control of microstructure and defect formation and improvement of mechanical properties of as-built products [3,4]. In traditional non-equilibrium solidification processing, the cooling rate of a liquid is treated as the key process parameter. In additive manufacturing technologies, the scanning speed of a power laser or an electron beam is treated as a key process parameter instead. From a fundamental point of view, it is liquid undercooling that plays a critical role in determining metastable microstructure formation [5–8]. In situ observations using ultrafast X-ray imaging [9] demonstrated that a maximum solidification velocity of 0.75 m/s was attained under a simulated condition of laser additive manufacturing technologies of Ti-6Al-4V alloys. Such a high solidification velocity should be driven only by a high liquid undercooling. However, it is challenging to measure this quantity directly because of a high temperature gradient in the melt pool [10]. To bypass this difficulty, it is plausible to evaluate liquid undercooling by comparing an actual solidification velocity to an established crystal growth velocity-undercooling relationship. However, crystal growth velocities in an undercooled liquid are apt to fluid flow and impurities dissolved. The effects of fluid flow and impurities are not well determined for most of metals and alloys. Pure Ti is an example of this kind.

Ti is chemically reactive in its liquid state and a high undercooling of its bulk liquid can be accessed only by containerless melting and solidification [11–15]. Ti also has a high affinity to oxygen in solid and liquid states [16]. Thus, pure Ti commercially available typically has an oxygen impurity concentration of about 2500 ppm in atomic fraction (1000 ppm in mass fraction) [17]. After special purification, high-purity Ti could have an oxygen concentration of 89 ppm in atomic fraction (30 ppm in mass fraction) [18]. In solidification, the oxygen impurity dissolved in liquid Ti may slow down dendritic growth kinetics due to a solute drag effect. For this reason, measured tip velocities of undercooled Ti dendrites could be dependent on bulk concentration of an oxygen impurity. Walder and Ryder [11] found that tip velocities of Ti dendrites in electromagnetically levitated melt droplets were comparable to those of undercooled Ni dendrites under similar experimental conditions. Compared to pure Ti, pure Ni can easily reach a low oxygen impurity concentration [8]. Thus, observations of similar tip velocities of the two metals suggested a low concentration of the oxygen impurity in their Ti samples. Bassler [12] measured tip velocities of Ti dendrites using a similar experimental set-up. Their data showed a depression of tip velocities as compared to those measured by Walder and Ryder [11]. It is unclear if the discrepancy is due to a difference in the oxygen impurity because impurity concentrations were not determined in either study. Very recently, Kavousi et al. modeled tip kinetics of undercooled Ti dendrites [19]. Their molecular dynamics simulation suggested an interfacial kinetic coefficient similar to that of pure Ni, but their phase field modeling suggested tip velocities even lower than those measured by Bassler [12]. Thus, there remain discrepancies about the tip velocities of Ti dendrites in literature. Although a revisit to tip velocities in electromagnetically levitated melts may provide a straightforward approach to reconciling the discrepancies, it is easier and instructive to carry out modeling of observed tip velocities with consideration of effects of fluid flow and impurities [8,20].

In this work, tip kinetics of undercooled Ti dendrites was modeled using a theory on three-dimensional dendritic growth [21] with consideration of a fluid flow effect and an oxygen impurity effect. The modeling suggested that a forced flow due to electromagnetic stirring has a significant effect on tip kinetics at low undercoolings but that this effect becomes negligible at high undercoolings. The modeling showed that the effect of a typical oxygen impurity of pure Ti on tip kinetics is significant at low and high undercoolings. Such modeling results reconciled the discrepancies about the tip velocities of Ti dendrites in electromagnetically levitated melts [11,12]. They also shed light onto effects of an oxygen impurity on microstructure formation in other materials. For convenience, we give a brief introduction to the theory first.

2. Theory

The interface undercooling at a primary tip of an undercooled dendrite can be divided into three parts [7]: a curvature undercooling ΔT_r , a constitutional undercooling ΔT_c , and a kinetic undercooling ΔT_k . There is an additional undercooling, thermal undercooling ΔT_t , in the liquid close to the tip. A sum of the interface and the thermal undercooling defines a bulk undercooling, ΔT that drives tip growth [7], namely

$$\Delta T = \Delta T_t + \Delta T_t + \Delta T_c + \Delta T_k. \tag{1}$$

Alexandrov and Galenko analyzed effects of fluid flow and a dilute solute on tip kinetics of an undercooled dendrite at an arbitrary Péclet number. Their theory, termed AG theory in short, considered three-dimensional dendritic growth with a laminar flow incident to a primary tip [21]. In the AG theory, the four partial undercoolings are expressed as

$$\Delta T_t = (Q/C_p) I v (P_t, P_f), \qquad (2)$$

$$\Delta T_r = 2\Gamma/R,\tag{3}$$

$$\Delta T_{c} = \frac{m_{e}C_{0}(1-k_{e})P_{c}\exp(P_{c}+P_{cf})I_{c}(\infty)}{1-(1-k_{e})P_{c}\exp(P_{c}+P_{cf})I_{c}(\infty)},$$
(4)

$$\Delta T_k = V/\mu,\tag{5}$$

where Q is the fusion of heat of the solid, C_p is the specific heat of the liquid, Γ is the Gibbs-Thompson coefficient of a solid-liquid interface, R is the dendritic tip radius, m_e is the liquidus slope, C_0 is the solute concentration (impurity concentration), k_e is the equilibrium partition coefficient, V is the dendritic tip velocity, and μ is the interfacial kinetic coefficient. $Iv(P_t, P_f)$ and $Iv(P_c, P_{cf})$ are flow-dependent Ivantsov functions for heat transfer and mass transfer in the liquid, respectively and can be expressed as

$$Iv(P_t, P_f) = P_t \exp(P_t + P_f) I_t(\infty), \tag{6}$$

$$Iv(P_c, P_{cf}) = P_c \exp(P_c + P_{cf}) I_c(\infty),$$
(7)

where P_t is a thermal Pélect number, P_f is a flow-dependent thermal Pélect number, P_c is a chemical Péclet number, and P_{cf} is a flow-dependent chemical Péclet number. These Pélect numbers are defined as follows:

$$P_t = VR/(2D_t),\tag{8}$$

$$P_f = UR/(2D_t),\tag{9}$$

$$P_c = VR/(2D_c),\tag{10}$$

$$P_{cf} = UR/(2D_c),\tag{11}$$

where D_t is thermal diffusivity in the liquid, D_c is chemical diffusivity in the liquid, and U is flow velocity. The two integrals of Equations (6) and (7) can be expressed as

$$I_{t}(\eta) = \int_{1}^{\eta} \exp\left[2P_{f} \int_{1}^{\eta'} \frac{g(\eta'')}{\sqrt{\eta''}} d\eta'' - (P_{f} + P_{t})\eta'\right] \frac{d\eta'}{\eta'},$$
(12)

$$I_{c}(\eta) = \int_{1}^{\eta} \exp\left[2P_{f} \frac{D_{t}}{D_{c}} \int_{1}^{\eta'} \frac{g(\eta'')}{\sqrt{\eta''}} d\eta'' - (P_{f} + P_{t}) \frac{D_{t}}{D_{c}} \eta'\right] \frac{d\eta'}{\eta'},$$
(13)

where the hydrodynamic function $g(\eta'')$ is given by

$$g(\eta'') = \frac{\sqrt{\eta''} E_1(\operatorname{Re} \eta''/2)}{2E_1(\operatorname{Re}/2)} + \frac{\exp(-\operatorname{Re}/2) - \exp(-\operatorname{Re} \eta''/2)}{\sqrt{\eta''} \operatorname{Re} E_1(\operatorname{Re}/2)}.$$
 (14)

Re is the Reynolds number and given by

$$Re = RU\rho_l/v \tag{15}$$

where ρ_l and v are mass density and dynamical viscosity, respectively.

The AG theory [21] includes a linear analysis of tip stability and determines the tip stability parameter as

$$\sigma^{*} = \frac{\sigma_{0}\beta^{\frac{7}{4}}}{1 + b(\alpha\beta^{-3/4})^{\frac{1}{1}}} \left[\frac{1}{(1 + a_{1}\sqrt{\beta}P_{t})^{2}} + \frac{1}{(1 + a_{2}\sqrt{\beta}P_{t}D_{t}/D_{c})^{2}} \frac{2m_{e}C_{i}(1 - k_{e})D_{t}}{(Q/C_{p})D_{c}} \right]$$
(16)

where σ_0 and b are material-dependent stability parameters, $\beta = 15\varepsilon_4$ is the anisotropy of the liquid/solid interfacial energy with a four-fold symmetry ε_4 , $a_1 \approx 0.381\sigma_0$ and $a_2 \approx 0.505\sigma_0$. The parameter α is given by

$$\alpha = \frac{d_0 Ua(\text{Re}) \left[1 + D_c / (2D_t) \right]}{4RV \left[D_c / (2D_t) + m_e C_i (1 - k_e) C_p / Q \right]},$$
(17)

where d_0 is the thermocapillary length and C_i is the solute concentration of the liquid at the dendritic tip. a(Re) is a flow dependent function and given by

$$a(\text{Re}) = \frac{\exp(-\text{Re}/2)}{E_1(\text{Re}/2)}$$
(18)

where $E_1(q) = \int_q^{\infty} u^{-1} \exp(-u) du$ is the first exponential integral function. The solute concentration C_i is given by

$$\Delta T_c = \frac{m_e C_0 (1 - k_e) P_c \exp(P_c + P_{cf}) I_c(\infty)}{1 - (1 - k_e) P_c \exp(P_c + P_{cf}) I_c(\infty)}.$$
(19)

Gao et al. [22] showed that the AG theory can provide a reasonable description of tip kinetics of undercooled succinonitrile dendrites under the influence of natural convection on the ground and in space. They further showed that the AG theory can also provide a reasonable description of tip kinetics of undercooled dendrites of pure Fe and pure Ni under the influence of strong magnetohydrodynamic flows [20,23]. Following such studies, it was supposed that a modeling of tip kinetics of undercooled Ti dendrites using the AG theory may help understand the discrepancies in literature [11,12]. Modeling results are shown below.

3. Modeling of tip velocities of undercooled Ti dendrites

Parameter	Symbol	Value	References
Heat of fusion (J mol ⁻¹)	Q	14300	[13]
Liquid specific heat (J mol ⁻¹)	C_p	45.5	[13]
Melting temperature (K)	T_m	1943	[13]
Thermal diffusivity (m ² s ⁻¹)	D_t	9.5×10 ⁻⁶	[15]
Density of liquid (kg m ⁻³)	$ ho_l$	4130	[15]
Kinetic viscosity in liquid (Pa s)	ν	3.3×10 ⁻³	[15]
Gibbs-Thompson coefficient (K m)	Г	2.77×10 ⁻⁷	*
Interfacial kinetic coefficient (m s ⁻¹ K ⁻¹)) μ	0.37	present work
Anisotropy of surface tension	E 4	0.04	[19]
Tip stability parameter	σ_0	0.04	present work
Tip stability parameter	b	0.1	[20–23]
Flow velocity (m s ⁻¹)	U	-0.3	[20]

Table 1 Parameters used in modeling of tip kinetics of thermal dendrites of Ti.

* Calculated in terms of a solid/liquid interfacial energy of $\sigma_{s/l} = 0.176 \text{ J m}^{-2}$ [19].

Diffusion-controlled growth of thermal dendrites of pure Ti in stagnant flow conditions was modeled using the parameters listed in Table 1 and compared to the experimental data by Walder and Ryder [11]. For thermal dendrites, the constitutional undercooling $\Delta T_c = 0$. Because forced flows have a negligible effect on tip velocities at high ΔT [20], least-square fitting of the AG theory to the experimental data in this undercooling region allowed for unambiguous determination of interfacial kinetic coefficient and tip stability parameter. In fitting, the relative error of an observed tip velocity was considered, which was defined as its percentage deviation from a modeled tip velocity at a given ΔT . A sum of the square of the relative error of the experimental data over high ΔT was achieved by choosing an interfacial kinetic coefficient of $\mu = 0.37$ m/s/K and a tip stability parameter of $\sigma_0 = 0.04$. Provided that these two values are accepted, the AG theory gives a reasonably good prediction of the tip velocities observed at $\Delta T > 50$ K. That is, the experimental data had a r.m.s. deviation of up to 20% only. Although μ is only a half of the value predicted by molecular dynamics simulation [19], it is comparable to a value of 0.30 m/s/K predicted and verified for pure Fe [23, 24]. This similar magnitude is reasonable because Fe and Ti have similar melting temperatures and both crystallize into a body-centered cubic structure. The molecular dynamics simulation [19] may have overestimated μ . On the other hand, σ_0 determined is by a factor of four larger than that of Fe. This difference is understood in terms of a larger anisotropy of liquid/crystal interfacial energy [19] and higher tip velocities of Ti dendrites. There is a discrepancy between the modeling and the experiment in a low undercooling region. The r.m.s. deviation of the experimental data amounted to 110 % at $\Delta T < 50$ K. Although solidification paths of the sample were not controlled by triggering of crystal nucleation, this discrepancy cannot be solely attributed to experimental errors. The thermal conductivity used in the modeling has high accuracy because it was measured by containerless processing of liquid and solid Ti in an ultrahigh vacuum [15]. Thus, it is neither the reason for the discrepancy. A forced flow effect due to electromagnetic stirring was determined to be significant on tip velocities of Ni dendrites at low ΔT [20]. Thus, it may be the reason for the discrepancy. To check this hypothesis, we considered the effect of a similar flow velocity of U = -0.3 m/s [20]. To check this hypothesis, we considered the effect of a similar flow velocity of U = -0.3 m/s [20]. The flow was supposed to point to the tip growth direction and has the negative sign. According to Hyers [25], forced flows in an electromagnetically levitated Ni melt have a Re larger than 600 in the ground-based gravity condition and have a turbulent nature. Such a large Re is defined on a length scale equal to the size of the levitated melt and provides a reason for a maximum flow velocity of up to 0.3 m/s. In the AG theory, Re is defined on a length scale equal to primary tip radius of an equiaxed dendrite (see Eq. 15) and has a much smaller value, e.g. Re = 0.68 at $\Delta T \sim 26$ K. Thus, the forced flow was treated as a laminar flow in the modeling. Including this convective transport effect did not further reduce the deviations of the experimental data at higher ΔT . Rather, it reduced the r.m.s. deviation of the experimental data to 46% at $\Delta T < 50$ K. In other words, it brought about an improved agreement between the experiment and the modeling. A remaining difference can be attributed to reasonable experimental errors estimated from the scatter of the observed tip velocities at low ΔT .



Fig. 1. Modeling of tip velocities of undercooled dendrites in electromagnetically levitated pure Ti. Solid and open circles are literature data [11,12].

However, even when the flow effect is included, there is a discrepancy between the modeling of thermal dendrites and the tip velocities observed by Bassler [12]. There is no physical reason for consideration of a flow velocity higher than 0.3 m/s to reconcile this discrepancy. Rather, it is more reasonable to consider an impurity effect. Although the species of impurities of the Ti material investigated by Bassler was not known, the majority impurity was most likely to be oxygen following other studies of pure Ti [17,18]. When oxygen is dissolved in levitated melts as the majority impurity, undercooled Ti dendrites crystallized therein should be treated as chemical dendrites. In other words, the ΔT_c of the dendritic tip is not zero and needs to be accounted for. Using parameters listed in Table 2, we modeled the tip velocities of chemical dendrites of Ti. Because a smaller atom usually diffuses faster than a large one, we adopted a value of $D_c = 1 \times 10^{-8}$ m²/s for chemical diffusivity of oxygen in liquid Ti. This value is higher than that of typical solute in a liquid metal [26]. As a first approximation, thermophysical properties of dilute Ti–O alloys

including μ were supposed to be equal to those of oxygen-free Ti. The only unknown parameters then are C_0 and σ_0 . Treating these parameters as free parameters, least-square fitting of the AG theory to the tip velocities observed by Bassler [12] yielded $\sigma_0 = 0.0078$ and $C_0 = 269$ ppm (atomic fraction). As these values are accepted, a r.m.s. deviation of the experimental data relative to the modeling is reduced down to 8%. Such a small deviation indicates a fairly good agreement between the AG theory and the experiment by Bassler [12]. Note that the determined σ_0 value is by a factor of 5 smaller than that used in the modeling of the data measured by Walder and Ryder [11]. Such a large difference is due to the depression of tip velocities by the oxygen impurity (see Fig. 1). On the other hand, the determined C_0 is close to a reported value of 95 ppm (mass fraction) for high-purity Ti [27]. For such reasons, the values of the two fitting parameters are reasonable.

Parameter	Symbol	Value	References
Heat of fusion (J mol ⁻¹)	Q	14300	[13]
Liquid specific heat (J mol ⁻¹)	C_p	45.5	[13]
Melting temperature (K)	T_m	1943	[13]
Thermal diffusivity (m ² s ⁻¹)	D_t	9.5×10 ⁻⁶	[15]
Density of liquid (kg m ⁻³)	$ ho_l$	4130	[15]
Kinetic viscosity in liquid (Pa s)	ν	3.3×10 ⁻³	[15]
Gibbs-Thompson coefficient (K m)	Г	2.77×10 ⁻⁷	*
Interfacial kinetic coefficient (m s ⁻¹ K ⁻	¹) μ	0.37	present work
Anisotropy of surface tension	84	0.04	[19]
Tip stability parameter	σ_0	0.0078	present work
Tip stability parameter	b	0.1	[20-23]
Chemical diffusivity (m ² s ⁻¹)	D_c	1×10 ⁻⁸	present work
Oxygen concentration (at.%)	C_{0}	0.0269	present work
Equilibrium partition coefficient	k_e	1.6	[16]
Liquidus slope (K/at.%)	me	10	[16]

Table 2 Parameters used in modeling of tip kinetics of chemical dendrites of Ti.

* Calculated in terms of a solid/liquid interfacial energy of $\sigma_{s/l} = 0.176 \text{ Jm}^{-2}$ [19].



Fig. 2. Comparison of partial undercoolings of thermal dendrites and chemical dendrites of Ti.

Given such a good agreement between the Bassler's data [12] and the AG theory, the modeling provided insights into the effect of the oxygen impurity on the tip velocities of undercooled Ti dendrites. Like that modeled for thermal dendrites, the flow effect is significant for chemical dendrites at low ΔT (not shown in Fig. 1) and becomes insignificant at high ΔT . Thus, attention was focused onto the impurity effect. Fig. 2 shows a comparison of partial undercoolings of the thermal and chemical dendrites of Ti. The free growth of the thermal dendrites of Ti is controlled largely by heat diffusion in the undercooled liquid. While this is true up to a maximum undercooling of 250 K accessed in the experiment [11], interface kinetics becomes important at $\Delta T > 150$ K. The curvature undercooling ΔT_r plays a sub-critical role at $\Delta T < 20$ K but it becomes smaller than ΔT_k at higher ΔT . The dissolution of the oxygen impurity in the liquid brings about significant changes of tip kinetics and partial undercoolings. As suggested above, the oxygen impurity introduces a non-zero ΔT_c to the ΔT . The ΔT_c is saturated at $\Delta T \sim 10$ K and its contribution is significant only at $\Delta T < 50$ K. Because of this non-zero ΔT_c , tip velocities are depressed at low ΔT . However, the low undercooling region is also where the flow effect is significant. The depression of tip velocities induced by the dissolved oxygen impurity is comparable to that by the forced flow in the tip growth direction. Because of this similarity, it becomes

technically difficult to distinguish the two effects in the low undercooling region. The two effects have to be distinguished by extending a modeling of tip velocities observed in the high undercooling region to the low undercooling region. The dissolved oxygen impurity has opposite effects on other partial undercoolings. Unlike a reduction of ΔT_r and ΔT_k , ΔT_t gains an increase. Such opposing effects on partial undercoolings remain significant as ΔT increases. This is the very reason why the data measured by Bassler [12] became lower than those measured by Walder and Ryder [11].

4. Discussion

The above modeling revealed a significant effect of the oxygen impurity on tip kinetics of Ti dendrites at low and high ΔT . There is a question at which concentration level the effects of oxygen impurity on the tip kinetics of Ti dendrites become insignificant. To answer this question, we extended the modeling of chemical dendrites to lower oxygen concentrations. For simplicity, we neglected the oxygen concentration of Walder and Ryder's samples and considered a power law dependence of σ_0 on C_0 :

 $\sigma_0 = 0.04 \exp(-60.86C_0), \tag{20}$

where $C_0 = 0-269$ ppm. Using Eq. 20, tip velocities and tip radii of chemical dendrites of Ti at varied C_0 were calculated in a diffusive growth condition. As shown in Fig. 3, tip velocities of chemical dendrites are reduced generally as C_0 is increased. Meanwhile the tip radii of chemical dendrites are increased. When C_0 is increased from 0 to 50 ppm (atomic fraction), the tip velocity at $\Delta T = 26$ K is depressed by about 25%. This small depression is not significant compared to uncertainties of the data measured by Walder and Ryder [11]. Thus, an oxygen concentration of $C_0 = 50$ ppm can be accepted as a technical threshold for distinguishing the oxygen impurity effect. Strictly speaking, the oxygen impurity concentration of the samples investigated by Walder and Ryder was unlikely to be zero. Our modeling with a reasonably low oxygen impurity concentration, e.g. $C_0 = 100$ ppm in atomic fraction, still supports the conclusion of the forced flow effect and allows for the reconciliation of the discrepancies in literature [11,12].

The above modeling also provided insights into oxygen impurity effects observed in other metals. Battersby et al. [28] showed that an oxygen concentration of 2400 ppm (600 ppm in mass fraction) brought about a more than 50% reduction of tip velocities of Cu



Figure 3. Modeled tip velocities (a) and tip radii (b) of Ti dendrites for different O concentrations. No flow effects are considered in the modeling for simplicity.

dendrites in a glass-fluxed melt, where convective flow is weaker than in a levitated melt [20]. Such a reduction of the tip velocities by the oxygen impurity can be understood in terms of the modeling results shown in Fig. 3a. In addition to tip velocities, the oxygen impurity may have a significant effect on tip radii of chemical dendrites and therefore on microstructure formation in non-equilibrium solidification. As shown in Fig. 3b, the change of the tip radii at medium ΔT due to an oxygen impurity is most significant. The introduction of an oxygen concentration of $C_0 = 269$ ppm as modeled above is shown to

increase the tip radii by one order of magnitude at medium ΔT . According to a model for grain-refinement in undercooled solidification [29], this huge change of the dendritic tip radii may increase a critical time required for break-up of a primary dendrite in a mushy zone and therefore shifts the formation of a grain-refined microstructure to a higher ΔT or a higher tip velocity. Consideration of this oxygen impurity effect can explain why the literature data of a critical ΔT for formation of a grain-refined microstructure in pure Cu and pure Ni showed a scatter [30, 31].

As far as dendritic growth in an additive manufactured material of pure Ti is concerned, an oxygen impurity may have multiple effects on microstructure formation. Firstly, a depression of the tip velocity by the oxygen impurity may increase the time available for another dendrite to be nucleated from the liquid such that microstructure formation becomes more complex due to competition between neighboring dendrites [32]. Secondly, the depression of the tip velocity may also increase the sensitivity of dendritic growth to a liquid flow affecting local microstructure and defect formation. Lastly, coarsening of dendritic tip radii due to the impurity effect may increase secondary arm spacings because of a change of local thermal and chemical gradients. All of these effects on local or general microstructure formation may have an impact on mechanical properties of as-built material of pure Ti and thus, need to be investigated in the future.

5. Conclusions

Tip velocities of undercooled dendrites in electromagnetically levitated Ti melts have been modeled using a three-dimensional theory of dendritic growth with consideration of effects of a forced flow in the tip growth direction and an oxygen impurity. The modeling has shown that a typical oxygen impurity concentration in nominally high-purity Ti material causes a depression of tip velocities and coarsening of tip radii. Consideration of this oxygen impurity effect has reconciled the discrepancies in literature [11,12]. The modeling has also shown that the oxygen impurity effect on tip velocities of Ti dendrites is similar to that of a forced flow due to electromagnetic stirring at low undercoolings and that distinguishing of them requires an extension of the modeling of tip velocities at high undercoolings down to low undercoolings. The modeling has predicted that the effect on tip velocities becomes indiscernible when the oxygen impurity concentration of pure Ti is reduced below 50 ppm (atomic fraction). Therefore, the oxygen impurity could have a significant influence in additive manufacturing technologies, the consequences of which need to be addressed in the future.

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