**In-situ** measurements of dendrite tip shape selection in a metallic alloy

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The size and shape of the primary dendrite tips determine the principal length scale of the microstructure evolving during solidification of alloys. **In-situ** x-ray measurements of the tip shape in metals have been unsuccessful so far due to insufficient spatial resolution or high image noise. To overcome these limitations, high-resolution synchrotron radiography and advanced image processing techniques are applied to a thin sample of a solidifying Ga-35wt.%In alloy. Quantitative **in-situ** measurements are performed of the growth of dendrite tips during the fast initial transient and the subsequent steady growth period, with tip velocities ranging over almost two orders of magnitude. The value of the dendrite tip shape selection parameter is found to be $\sigma^* = 0.0768$, which suggests an interface energy anisotropy of $\varepsilon_4 = 0.015$ for the present Ga-In alloy. The nonaxisymmetric dendrite tip shape amplitude coefficient is measured to be $A_4 \approx 0.004$, which is in excellent agreement with the universal value previously established for dendrites.

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**I. INTRODUCTION**

Dendritic growth is of vital importance in the formation of microstructures during solidification of metal alloys. The key to understanding dendrite patterns is to study the shape and stability of the dendrite tips, since they stand at the beginning of the morphological evolution [1]. The radius of curvature of a dendrite tip, $R$, represents the initial length scale of a dendrite during growth. The selection criterion [1,2]

$$\sigma^* = \frac{2Dd_0}{RV}$$

provides a fundamental link between the radius $R$ and growth velocity $V$ of a dendrite tip via the dimensionless selection constant $\sigma^*$ that is independent of the growth conditions, where $D$ and $d_0$ are the diffusion coefficient of the melt and chemical capillary length, respectively. According to microscopic solvability theory (MST) [1,2], in a low undercooling regime $\sigma^*$ depends solely on the interface energy anisotropy $\varepsilon_4$ for a $n$-fold crystalline symmetry.

Very near the tip, the shape of a dendrite can be closely approximated as an axisymmetric paraboloid. Further away from the tip, the anisotropic interface energy increasingly affects the shape. Therefore, fins start to develop perpendicular to the growth axis that are located at azimuthal angles where the interface energy has a maximum. Soon after, the growing fins develop transversal instabilities that ultimately lead to the characteristic side branches of dendrites. Ben Amar and Brener [3] provided an analytical solution for the universal shape of dendrite tips before side branches form. For a four-fold crystalline symmetry, the shape in a plane containing the fins is given by

$$\frac{z}{R} = \frac{1}{2} \left( \frac{x}{R} \right)^2 - A_4 \left( \frac{x}{R} \right)^4,$$

where the coordinates $z$ and $x$ are centered at the tip [cf. Fig. 1(c)] and have orientations opposite and perpendicular to the growth direction, respectively. The last term in Eq. (2) represents the deviation from the isotropic paraboloidal shape by the fins. The fourth-order amplitude coefficient $A_4$ does not depend on any material properties or growth conditions. Several experimental and numerical studies [4–6] have confirmed that Eq. (2) provides a good approximation to the dendrite tip shape. However, the value of $A_4 \approx 0.004$ found in these studies deviates significantly from the theoretical solution $A_4 = 1/96 \approx 0.01$, the reasons for which are not yet entirely clear [4].

Major progress in understanding the tip shape selection mechanism was achieved during the 1980s and 1990s [1]. An important factor in this success was the use of transparent substances in high-precision solidification experiments, e.g., [5–8]. These materials permit **in-situ** observation of dendritic growth features near room temperature using an optical microscope. Nonetheless, the dendrite tip shape selection measurements available for transparent substances do not confirm MST conclusively [1,9,10]. At the same time, the experimental analysis of dendritic microstructure formation in (opaque) metals remained limited to post-mortem observations [11]. In a classical analysis of micrographs, Liu et al. [12] determined the selection parameter $\sigma^*$ for a quenched Al-Cu alloy sample. Only with the development of improved synchrotron x-ray facilities and sensors starting in the late 1990s has **in-situ** imaging of dendritic growth in metallic alloys become possible [13]. Although there is a vital interest in extending the experimental validation of dendrite growth...
In transmission radiography there exists a strong trade-off between time resolution and image noise [14]. Even thin layers of metal cause strong absorption of the illuminating x-rays, which inherently limits the number of photons that can be captured by the image sensor. Since the tip region exhibits the strongest dynamics and smallest length scales within a dendritic structure, its imaging is accompanied by significant noise, which prevents a straightforward use of standard analysis methods. A critical role in the design of x-ray imaging experiments is played by the initial solute concentration of the solidifying alloy. While higher concentrations usually lead to a better image contrast due to larger density differences between the solid and liquid phases, the dendritic structures become finer and more difficult to resolve. When using lower concentrations, the dendrites become larger, but the contrast is reduced. Larger structures are also more prone to confinement effects caused by the limited thickness of the samples.

Despite these difficulties, a limited number of studies have performed in-situ radiography measurements of dendrite tips in metals. Several experiments focused on the measurement of tip velocities in Al-Cu alloys, e.g., [15,16], but provided no data on the tip shape. In-situ measurements of both the velocity and the radius of dendrite tips have only recently been reported [17,18]. Mirihanage et al. [17] measured time varying tip velocities and radii for directional solidification of an Al-15wt.%Cu-9wt.%Si alloy and successfully compared measured and predicted solute concentration fields ahead of the dendrite tips. However, dendrite tip shape selection was not examined and no values of \( \sigma^* \) and \( A_4 \) were determined. Clarke et al. [18] measured tip velocities and radii for an Al-Cu alloy and compared their results with phase-field simulations. They found that the measured tip radii essentially follow the \( R \sim V^{-1/2} \) relationship from MST [Eq. (1)], but their radii distribution range at a given tip velocity was between a factor of 2 and 3. This large uncertainty was attributed to a lack of contrast in the radiographic images. Consequently, no values of \( \sigma^* \) and \( A_4 \) were obtained. Their phase-field results indicate that in their experiment the effect of melt convection on tip radius selection was negligibly small.

In the present study, in-situ measurements of dendrite tip shape selection are made for a hypereutectic Ga-35wt.%In metallic alloy, which solidifies near room temperature. It should be noted that the primary phase in this alloy (In) has a body-centered tetragonal (bct) crystal structure, which corresponds to a slightly distorted face-centered cubic (fcc) system with fourfold crystalline symmetry. The evolution of the dendrite tip shape is observed over a large range of growth velocities by high-resolution synchrotron radiography imaging. The region around the tip is extracted by a relatively small window that follows the tip position over time. The image quality is substantially improved using a temporal averaging filter. Another aspect of the present study is that the parameter group \( Dd_0 \) in the definition of the selection parameter, Eq. (1), is obtained directly from previous measurements of the universal pinching dynamics during dendrite side-branch detachment [19]. This enables the tip selection parameter \( \sigma^* \) to be predicted much more accurately, as the measurement of the individual material properties needed to evaluate the group \( Dd_0 \) usually requires elaborate experimental tests that are available only for a small number of the most frequently analyzed alloys [20–22].

**II. EXPERIMENTAL SETUP AND PROCEDURE**

Figure 1(a) shows the test cell used in the present solidification experiment. The cell was previously employed in a different study carried out by means of a microfocus x-ray tube [23]. The Ga-35wt.%In alloy is prepared from 99.99% Ga and 99.99% In. The alloy is melted and filled into the cell, which has a liquid metal volume of 22 \( \times \) 22 \( \times \) 0.2 mm\(^3\).
As shown in Fig. 1(a), the heating/cooling system consists of two sets of Peltier elements in thermal contact with the bottom and top ends of the solidification cell. The Peltier elements are connected to a control unit that allows for the independent adjustment of the cooling rate and temperature gradient across the cell. The distance between the heater and the cooler is approximately 23 mm. Temperatures are measured using two miniature type-K thermocouples that are in thermal contact with the outer surface of the cell near the edge of the Peltier elements. The accuracy of the temperature control is ±0.2 K. The vertical temperature gradient is calculated from the temperature difference measured between these two thermocouples. The alloy is directionally solidified from top to bottom with a constant temperature gradient of 2 ± 0.4 K/mm and cooling rate of 0.002 K/s.

The experiment is performed at the ID19 beamline of the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. The solidifying sample is exposed to a monochromatic, parallel x-ray beam with a photon energy of 40 keV. Conventional transmission radiographs are obtained by means of a scintillator that is coupled to a high speed scCMOS camera (PCO.edge) with 2048 × 2048 pixels, yielding an effective pixel size of 0.72 µm. This imaging equipment leads to a field of view of about 1.5 × 1.5 mm² [24]. Radiographs are recorded continuously with a frame rate of 2 s⁻¹. To change the location of the observation window, the position of the solidification cell is manipulated with respect to the x-ray beam by means of a motorized positioning system.

Before solidification is initiated, the Ga-In alloy is heated to a temperature of 70 °C and held at this temperature for a few minutes. The liquidus temperature of the alloy is about 45 °C. During this stage, images are taken by real-time radiography to ensure that the alloy is homogenously mixed before the cooling process is started. Dark field and flat field images of the completely molten alloy are also recorded for subsequent data processing. The cooling of the cell is initiated after recording these reference images. Soon after the first appearance (t = 0) of solid alloy at the upper cold end of the sample, the solidification front is searched for dendrites that are suitable for analysis. Dendrites are selected by requiring that the underlying crystallographic orientations are well aligned with the sample plane and the viewing direction, as indicated by the angles of the dendrite side branches with respect to the main stem.

The selected dendrite tips are then followed by shifting the cell position in regular intervals along the growth direction to ensure that the tips remain within the field of view. Figure 1(b) shows an example stack of images that consists of the last frames captured at seven successive cell positions. Each image is outlined by a different color dashed line. The dashed black rectangle in Fig. 1(a) illustrates the approximate position of the image stack of Fig. 1(b) within the test cell. In the present study, four tips are selected for detailed analysis, as indicated by the labels Tip 1–4 in Fig. 1(b). The measured tip trajectories are displayed as solid lines, where the colors correspond to the cell positions. To observe changes in the global dendritic structure at some intermediate times, the tip-following scans are interrupted to perform mesh scans of the entire solidification cell. This is the reason for the measurement gaps that are apparent in the data presented below.

III. IMAGE PROCESSING AND ANALYSIS

Noisy images pose a formidable challenge to reliably measuring the fine morphological details that are associated with a dendrite tip. In the present image processing approach, the results are achieved. In the present image processing approach, a comoving frame of reference is used to detect stationary tip features. Then, for convenience, the background intensity of all frames is normalized to some predefined value vbg. This is achieved by scaling of the image contrast in each of the individual frames.

The following processing steps are performed individually for each dendrite tip. A narrow rectangular image region SX,Z,t) is defined that contains the entire growth path [shown as black outline in Fig. 1(b) for Tip 2]. Figure 1(c) shows a closeup of the rectangular region near the tip, where its coordinate system [X,Z] is fixed to the sample plane. To improve robustness, the image data are slightly smoothed using a spatial Gauss filter with a standard deviation of one pixel.

Next, the dendrite tip position Z*(t) is determined as a function of time. The denser solid has a lower intensity than the (liquid) background. Thus, the progression of the tip parallel to the Z coordinate can be tracked by performing a minimum projection P(Z,t) = minZ[S(X,Z,t)] along the lateral direction X. In the resulting image P(Z,t), the tip position Z*(t) is then identified as the border separating the high and low intensity regions. This is illustrated in Fig. 1(e), where the profile of P is shown at t = 748 s.

The tip velocity is calculated as the time derivative of Z*(t) using a finite difference approximation followed by locally estimated scatterplot smoothing (LOESS), which is based on second degree polynomials. Here, particular care is taken to properly handle the time gaps in the measurement data that are mentioned in the previous section. Furthermore, Z*(t) is used to define a small subimage T(x,z,t) of S that only contains a narrow region around the tip. In this comoving window, the tip location remains fixed at the origin of the local coordinate system [x,z].

Examples of a tip-centered window for Tip 3 at three different times are provided in the top row of Fig. 2(a). As shown in Fig. 3(a) below, these times correspond to a large range of tip growth velocities. As the most important step in reducing image noise, a uniform temporal filter (moving average) is now applied to T over a range of t ± 70 frames (±35 s). The resulting images in the center row of Fig. 2(a) show a very well-defined dendrite tip, indicating that the tip shape is nearly stationary within the chosen time interval for averaging. Since no noise-free reference images are available, the peak-signal-to-noise ratio (PSNR) [26] is estimated for the noisy and time-averaged images. An improvement from about 34.5 ± 0.4 dB (top row) to 55.3 ± 0.1 dB (center row) is achieved.
in Eq. (2) is neglected by setting $A_4 = 0$, which corresponds to a purely parabolic shape. Fit 2 is then used to measure the anisotropic deviation from the parabolic shape as quantified by the fourth-order amplitude coefficient $A_4$. For this fit, a larger fit range of $8\mu m$ is applied (blue arrow). This range is still close enough to the tip that the asymmetries in the dendrite shape and side branches visible in the lower portion of the images in Fig. 2(a) do not affect the fit. To suppress unreliable measurements, fitting data that have a coefficient of determination of $r^2 < 0.9$ are neglected. Based on the confidence intervals of the two fits, the mean uncertainties in $R$ and $A_4$ are 5.5% and 33.6%, respectively.

IV. RESULTS AND DISCUSSION

The four neighboring dendrites selected for analysis [Fig. 1(b)] are part of a larger grain that had previously nucleated near the cold upper end of the sample. The selected dendrites are tilted at a uniform angle of $\sim55^\circ$ with respect to the vertical sample axis. As can be seen in Fig. 1(b), the side branches are longer on the downward facing side of the dendrites, where the undercooling is larger. The temperature of the tips decreases from about 37 °C to 27 °C during the time span considered in the analysis.

A. Tip velocity, radius, and shape

The measured growth velocities and radii for the four selected dendrite tips are shown in Fig. 3. The tip velocities vary by almost two orders of magnitude over the roughly 6000 s long measurement period. Initially, the growth velocity is high because of the relatively large initial undercooling of the melt. The rapid growth results in a relatively small tip radius of about 1 $\mu m$. The tips then relax towards a slower, steady growth regime with a larger tip radius. Towards the end of this transition, at about 1500 s, the tip velocity reaches a local minimum and the tip radius a maximum close to 5 $\mu m$. Subsequently, the tip velocity experiences a slight increase and the tip radius a decrease. After about 2500 s, the tip growth is almost completely steady, except for Tip 1 which continues to slow down.

At the beginning of the measurements, the growth rate is highest for Tip 1 and decreases towards Tip 4. This can be explained by the fact that Tip 1 is closest to the cold top boundary and experiences the largest undercooling. After a short time, however, the velocity ranking among the tips becomes inverted. The trailing tips (lower number) experience a stronger deceleration, and their velocity eventually falls below that of the leading tips (higher number). This may be explained by diffusive interactions between the dendrites. A trailing tip has a reduced undercooling because of solute diffusion from the neighboring dendrite that is ahead. The velocity decrease of Tip 1 after 1500 s can also be attributed to diffusive interactions.

The measured fourth-order amplitude coefficients $A_4$ are shown in Fig. 4. The results for the four dendrites exhibit no systematic trend with tip number, and differences among them consist of statistical noise only. Therefore, instead of providing the $A_4$ values individually for each of the four dendrites, only a moving average of their median value is shown (solid...
FIG. 4. Measured fourth-order amplitude coefficient as a function of time. The black line is a moving average of the median values, while the gray areas represent the range of the smallest and largest values among the four tips. Density distributions of the $A_4$ values measured for each of the four tips are shown in the plot on the right side (colored lines). The black dashed line represents the mean of the peaks in the density distributions.

as an experimental setup similar to the present one. A value of $Dd_0 = 0.122 \pm 0.0026 \mu m^3 s^{-1}$ was found. To evaluate if the same value can be used for the present experiments involving a Ga-35wt.%In alloy and a temperature range of $27–37^\circ C$, it is necessary to consider the temperature dependence of each property in the product $Dd_0$. Using the definition of the chemical capillary length, $Dd_0$ is given by

$$Dd_0 = \frac{DI}{m |l1 - k| C_{eq}^2},$$

where $\Gamma$ is the Gibbs-Thomson coefficient, $m$ is the liquidus slope, $k$ is the partition coefficient, and $C_{eq}$ is the equilibrium solute concentration of the melt at a given temperature. Based on the data summarized in Ref. [19], a careful examination of the temperature dependence of each material property in Eq. (3) revealed that only the diffusion coefficient $D$ and the liquidus slope $m$ show strong variations in the near-eutectic region. However, the combined effect on the product $Dd_0$ is quite small because the individual variations tend to compensate each other in Eq. (3). It is estimated that the maximum deviation from the previously measured value for $Dd_0$ for the current temperature range is less than $\pm4\%$. Therefore, the value of $Dd_0 = 0.122 \mu m^3 s^{-1}$ is adopted in the following. The overall uncertainty in this value is estimated to be 6.1%.

Figure 5 shows the time variation of the tip selection parameter $\sigma^*$ calculated from Eq. (1) based on the measured $V$ and $R$ and the above value for $Dd_0$. The figure is designed the same way as Fig. 4 above. It can be seen that the tip selection parameter is largely constant over time. This result is expected, since theory predicts that $\sigma^*$ is independent of the growth conditions. It is remarkable nonetheless, because the tip growth velocities vary by almost two orders of magnitude over the course of the measurements. The density distributions for $\sigma^*$ are largely consistent among the different tips. Taking the mean location of the peaks of the density distributions for the four tips gives $\sigma^* = 0.0768 \pm 0.0034$, where the uncertainty is again the standard error for a 95% confidence level. Based on the uncertainties in the tip radius and the product $Dd_0$ provided above, and a negligible uncertainty in the measured tip velocity, the mean uncertainty in a single measurement is estimated to be 12.6%.

The value for $\sigma^*$ determined here is compared to other experimental results and MST in Fig. 6. The line showing the dependence of the tip selection parameter on the fourfold

black line) together with their minimum-maximum range at each measurement time (gray area). As expected from theory, the median $A_4$ values are, despite some fluctuations, essentially constant over time and, therefore, independent of the growth conditions. Density distributions of the $A_4$ values for each tip are provided on the right side of Fig. 4. The four density curves are largely consistent in their shape and the location of their maximum. The density maxima (peaks) are relatively sharp, further confirming that a single value of $A_4$ is preferred throughout the measurements. It can be seen that the density distributions are somewhat skewed towards lower $A_4$ values. These deviations from a symmetrical normal distribution imply that using the mean value and standard deviation of all combined data would likely result in a poor estimate of $A_4$ and its uncertainty. Instead, the fourth-order amplitude coefficient is estimated here based on the location of the peaks in the density distributions. The mean location of the four peaks (dashed line in Fig. 4) is given by $A_4 \approx 0.00406 \pm 0.00039$, where the uncertainty is the standard error for a 95% confidence level. Note that this uncertainty is much smaller than the mean uncertainty of a single measurement, which is 33.6% (see above). The present value of $A_4 \approx 0.004$ is in excellent agreement with other measurements and numerical results reported in the literature [4–6]. These previous results are for different alloy systems, indicating that the value of the fourth-order amplitude coefficient is indeed universal. The ability to determine such a small coefficient to reasonable accuracy provides considerable confidence in the present tip shape measurements.

B. Tip selection parameter $\sigma^*$

As indicated by Eq. (1), the determination of the dendrite tip selection parameter $\sigma^*$ requires not only the measurement of the tip velocity and radius, but also the knowledge of the product of the diffusion coefficient and the capillary length, $Dd_0$. The individual measurements necessary to determine this property product are far from trivial and only available for a small number of materials. Recently, some of the present authors proposed a method to obtain $Dd_0$ as a single parameter based on measurements of the pinching dynamics during the detachment of dendrite side branches [19]. In this previous study, the product $Dd_0$ was determined for a Ga-25wt.%In alloy at a temperature of approximately 15 °C using
interface energy anisotropy $\varepsilon_4$ is the linearized MST result in the limit of small tip growth Péclet numbers [27]. The MST curve suggests an interface energy anisotropy of $\varepsilon_4 \approx 0.015$ for the present Ga-In alloy. Unfortunately, no independent measurement of $\varepsilon_4$ for a Ga-In alloy was found in the literature. The $\sigma^*$ measured postmortem for an Al-Cu alloy [12], which is the only other data point available for metals, is in excellent agreement with MST. For illustrative purposes, two representative experimental results are included in Fig. 6 for transparent alloys; a more complete comparison for all available $\sigma^*$ measurements in transparent substances can be found in Refs. [9,10]. The measured $\sigma^*$ for NH$_4$Br–H$_2$O [28] falls almost exactly on the MST line. The experimental $\sigma^*$ values reported in Ref. [5] for succinonitrile (SCN)–acetone (ACE) alloys are multiplied by a factor of 2 to conform to the $\sigma^*$ definition given by Eq. (1). The original $\sigma^*$ values of Ref. [5] correspond to an alternative definition that is uniformly applicable to alloys and pure substances [29]. The $\varepsilon_4$ for SCN was taken from Ref. [22]. The data in Ref. [5] show a strong dependence of $\sigma^*$ on the undercooling, and the vertical bar in Fig. 6 indicates the range of values measured. The relatively large discrepancy between SCN measurements and MST was already noted in Ref. [27], and no explanation has emerged since.

During the present experiment, the diffusion length at the dendrite tips $l_D = D/V$ increases approximately from 50 $\mu$m to 1500 $\mu$m. Therefore, the solute diffusion field around the tips will become affected by the sample walls, which are 200 $\mu$m apart. Nonetheless, no significant effect on the selection of the tip shape can be noted. This can be explained by the fact that the tip radius, which is the essential length scale in the tip selection problem, remains small compared to the dimensions of the test cell.

Moreover, it is likely that melt convection is present in the test cell during solidification. The nature and intensity of such convection would be highly variable since the amount and shape of the solid in the test cell is constantly evolving. Although such convection will cause changes in the dendrite tip growth velocity and radius compared to a purely diffusive environment, it does not affect the dendrite tip shape selection. Otherwise, the tip selection parameter would not be as constant as shown in Fig. 5. This finding agrees with previous theories [30] and experiments in transparent alloys [5] that low to moderate intensity convection has no effect on the selection parameter $\sigma^*$.

V. CONCLUSIONS

In-situ measurements of the shape selection of dendrite tips are performed during solidification of the metallic alloy Ga-35wt.%In using a combination of high-resolution synchrotron radiography and advanced image processing techniques. Tip radii down to 1 $\mu$m are accurately evaluated. Tip growth velocities range over almost two orders of magnitude. The parameter group $Dd_0$ is obtained from previous measurements of the universal pinching dynamics during dendrite side-branch detachment. By means of this approach reliable in-situ measurements of dendrite tip shape selection in metals could be conducted. A key result of this work is the value of the dendrite tip shape selection parameter, which is found to be $\sigma^* = 0.0768 \pm 0.0034$. Based on microscopic solvability theory, this $\sigma^*$ value suggests an interface energy anisotropy of $\varepsilon_4 = 0.015$ for the present Ga-In alloy. The fourth-order amplitude coefficient describing the nonaxisymmetric shape of a dendrite tip is determined to
be $A_4 = 0.00406 \pm 0.00039$, which is in excellent agreement with the universal value previously established. The interface energy anisotropy for the present alloy should be measured independently to fully verify MST. Considering the importance of dendritic growth in establishing the microstructure in almost all commercially important metals and alloys [1], and the relatively uncertain nature or limitations of present analytical theories [2,3] and computer simulations [4], the present experiments should provide new motivation for making progress in this field.

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