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Dendrite spacings in directionally solidified superalloy PWA-1480

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Abstract

Primary dendrite spacings and side-branch coarsening kinetics were examined in specimens of the single-crystal multicomponent commercial superalloy PWA-1480, which were directionally solidified in a positive thermal gradient. The experimentally observed dependence of primary dendrite spacings and side-branch coarsening kinetics on growth rate and thermal gradient were in agreement with the behavior predicted by analytical models developed for binary alloys.

1. Introduction

A complex branched dendritic microstructure is commonly produced during solidification of binary alloys. The breakdown of a planar liquid–solid interface first results in the formation of cells, the fingers of the solid protruding into the solidifying melt. These fingers are called primary dendrites when increasing constitutional supercooling in the intercellular melt results in the formation of side branches (secondary arms). Dendrite spacings have a significant influence on mechanical properties. The solidification parameters which control the formation and distribution of primary and secondary dendrites have, therefore, been extensively studied. Several analytical models have been proposed in the literature for predicting the dependence of the primary dendrite spacing $\lambda_1$ in binary alloys on the solidification rate $R$, the thermal gradient in the melt at the liquid–solid interface $G_l$ and the alloy solute content $C_0$ [1–3]. Hunt [1], using the approach taken by Bower et al. [4] for modeling microsegregation for a primary dendrite array, obtained the following relationship for $\lambda_1$

$$\lambda_1^2 = 4\sqrt{2} \left( \frac{G_1}{G_l} - 1 \right) \left( D_s / R \right) r_i$$

where $D_s$ is the solute diffusivity in the melt and $r_i$ is the radius at the tip of the primary dendrite. Neglecting the solutal diffusion in the solid, the concentration gradient in the melt at the dendrite tip $G_1$ during steady-state directional solidification is equal to $RC_l(k-1)/D_s$, where $C_l$ is the solute content of the melt at the dendrite tip and $k$ is the solute partition coefficient. Because of the assumptions of local equilibrium and constant thermal gradient, the interdendritic composition gradient $G_c$ is taken to be equal to $G_i/m_l$, where $m_l$ is the liquidus slope. Hunt used $C_l$ and $r_i$ predictions from the dendrite growth model of Burden and Hunt [5] in the above relationship, and showed that for a variety of binary alloys, the experimentally observed primary arm spacings are proportional to $G_1^{-0.5}$ and $R^{-0.25}$, as predicted from eqn. (1). However, detailed quantitative comparisons show that this approach underestimates the experimentally observed primary dendrite spacings [3, 6–8]. The dendrite growth model of Burden and Hunt [5] treats an array of primary dendrites and assumes minimum undercooling at the tips to predict $r_i$ and $C_l$. Trivedi [3] incorporated the $C_l$ and $r_i$ predictions from his model (which assumes a marginally stable paraboloidal dendrite tip) in eqn. (1) in order to predict dependence of primary dendrite spacings on the thermal gradient and growth rate. This approach shows a good quantitative agreement with the experimental data for succinonitride-acetone [3] and other binary alloy dendrites [6]. An equally good agreement with
the experimental data is obtained when eqn. (1) is used with the \( r \) and \( C_i \) predictions from the dendrite growth models of Laxmanan [6]. While extensive \( G \), \( R \) and \( \lambda \), data are available in the literature from directional solidification experiments on binary alloys, where actual thermal gradients were carefully measured by thermocouples placed in the melt, such data for commercial alloys are very limited [9, 10]. The feasibility of using the above models for predicting quantitatively the experimentally observed primary dendrite spacings in commercial multicomponent alloys is relatively unexplored.

Experiments on directionally solidified alloys have shown that the initial side-branch spacing near the tip of a primary dendrite is about 2-3 times the dendrite tip radius [11, 12]. This ratio has been observed to be 3 for succinonitrile [13] and 4.7 for water-ammonium-bromide solution [14], during growth of dendrites in supercooled melts. The spacing between neighboring side branches is observed to increase with increasing distance from the dendrite tip. Several factors have been proposed in the literature which may be responsible for the secondary arm coarsening. The coarsening may be due to simultaneous dissolution of smaller arms with sharper tip radii and growth of larger arms with less curvature [15, 16]. It may be due to the process of dendritic separation, i.e. the secondary arm dissolves and detaches from the primary dendrite (owing to the sharp curvature which reduces the local liquidus temperature) [16]. The coalescence of the neighboring side-branches can also lead to coarsening [17]. Kirkwood [18] has recently proposed a simple analytical model to predict the side-branch coarsening kinetics in binary alloys. This model is especially suited to partial directional solidification and quenching experiments where measurements of the side-branch spacing from the tip of the primary dendrites to their base in the quenched mushy region of the microstructure can be carried out [12, 18]. The analytical model assumes a constant temperature and composition of the interdendritic melt. It treats the coarsening process as the simultaneous dissolution of smaller arms (spheres) with sharper tip radii and growth of larger arms with less curvature.

This paper is part of a detailed investigation into the possible effects of reduced convection on microsegregation and dendritic morphologies in directionally solidified single crystals of a nickel base superalloy, PWA-1480. This alloy, with a nominal composition of Ni–12Ta–10.4Cr–5Co–5Al–4W–1.5Ti (in weight per cent) is used as single-crystal blades in gas turbine aeroengines. Understanding the dependence of the primary and secondary dendrite spacings and the microsegregation behavior on the processing parameters is of great importance in modeling the formation of solidification-related defects, such as microporosity and macrosegregation, especially freckling. The purpose of this study was to measure the dependence of primary dendrite spacings on thermal gradients and growth speeds, and to explore the feasibility of predicting it quantitatively using the analytical models available for binary alloys. Similar examination of the side-branch coarsening kinetics was the second purpose. This alloy has been selected because the solidification behavior of PWA-1480, despite its being a multicomponent system, is similar to that of a binary alloy. The final solidification at the base of primary dendrites in this alloy occurs in a narrow temperature range (about 5–7 K) [19]; it can be considered nearly isothermal. Unlike other commercial superalloys, it does not form any interdendritic precipitates in the mushy zone. The side branch coarsening analysis is therefore not complicated by the formation of carbide precipitates between the liquidus and eutectic temperatures. In addition, the solutal partition coefficients in PWA-1480 are independent of temperature in the range 1610–1585 K [19].

2. Experimental details

Single-crystal cylindrical specimens (length 7 cm, diameter 0.5 cm) of the superalloy PWA-1480, were directionally solidified in alumina crucibles under flowing argon atmosphere in a Bridgman-type furnace. The growth direction was within \( \pm 8^\circ \) of the [100] orientation of the f.c.c. nickel based gamma matrix in these specimens. Details of the directional solidification furnace are presented in ref. 20. Directional solidification was carried out by translating the furnace (gradient zone) at constant speeds varying from 0.001 to 0.023 cm s\(^{-1}\). Two Pt–Pt–13%Rh thermocouples kept inside the specimen (located along the specimen length with a distance of about 0.5 cm between thermocouple beads) were used to record the \textit{in situ} thermal profile. When the thermocouple extending farther into the melt was at a predetermined temperature, usually near the liquidus temperature of
the alloy, 1610 K, the furnace was quickly withdrawn, positioning a quench nozzle over the solid–liquid zone and quenching by spraying water on the crucible surface.

The specimens were metallographically polished and etched (33 acetic acid, 33 nitric acid, 33 water and 1 hydrofluoric acid by volume) to correlate microstructural features, such as the tip and base of primary dendrites, with the temperatures measured by the two thermocouples. The dendrite tip radius was measured for one sample by serial sectioning of the primary dendrites on longitudinal sections (parallel to the alloy growth direction), and fitting a parabola to the midsection of the dendrite near its tip. Primary dendrite spacings were measured on transverse sections (perpendicular to the growth direction). The spacings reported in this paper are equal to 

\[ \frac{A}{N} \] 

where \( A \) is the total specimen cross-section area (transverse section) and \( N \) is the number of primary dendrites on that cross-section. Depending on the growth conditions, 200–600 primary dendrites were observed and counted on the corresponding specimen cross-sections (18.58–20.14 mm²).

Secondary arm spacings were measured by averaging the distance between five adjacent side branches on the longitudinal section (parallel to the alloy growth direction) of a primary dendrite as a function of distance from the dendrite tip. Each of the side branch spacing data reported here is the average of secondary arm spacings from four or five primary dendrites. The main source of scatter in this method of obtaining the secondary arm spacing data is the misalignment of primary dendrites, in the mushy region, with respect to the growth direction (growth direction is in the plane of metallographic polishing). This causes larger scatter in samples with thinner primary dendrites, i.e. those solidified at higher growth speeds.

Two experimental observations were used to ascertain that the steady-state growth conditions were achieved before the specimen was quenched: the variation in primary dendrite spacings along the length of the directionally solidified sample and the thermal profiles from the two thermocouples. Primary and secondary arm spacings reported in this paper correspond to the specimen portions where primary arm spacings were constant along the specimen length. In this portion of the sample, the thermal gradients measured at the tips of primary dendrites \( G_t \), assumed to be at the alloy liquidus temperature (1610 K), from the two thermocouples were within ±6%. The growth speed is therefore assumed to be identical with the furnace translation speed in the present analysis. The side branch coarsening time \( t \) was obtained as the ratio of the distance from the quenched primary dendrite tips (the liquidus temperature) along the specimen length in the mushy region and the growth speed. Table 1 lists the growth parameters of the specimens examined in this study. A cooling rate typically about 50 K s⁻¹ was recorded during quenching by the thermocouple located near the dendrite tips.

### 3. Results and discussion

Figure 1(a) shows a typical longitudinal section (parallel to the growth direction) through a directionally solidified specimen. This specimen was solidified at a growth speed of 0.013 cm s⁻¹ (\( G_s = 170 \text{ K cm}^{-1} \)). As mentioned earlier, the temperatures shown in this figure were obtained by correlating the location of the thermocouple tips in the longitudinal microstructures and the corresponding thermal profiles. The use of single-crystal specimens has resulted in primary dendrites having excellent alignment with the growth direction. This reduces the contribution to the experimental scatter in the dendrite spacings from their poor alignment in the microstructure. A distinct change in the microstructure of the quenched interdendritic region, observed at a

<table>
<thead>
<tr>
<th>Sample</th>
<th>( R ) (cm s⁻¹)</th>
<th>( G_s ) (K cm⁻¹)</th>
<th>( \lambda_t ) (cm)</th>
<th>Cell length (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4G</td>
<td>0.0033</td>
<td>120</td>
<td>0.0273</td>
<td></td>
</tr>
<tr>
<td>9G</td>
<td>0.0100</td>
<td>85</td>
<td>0.0215</td>
<td></td>
</tr>
<tr>
<td>11G</td>
<td>0.0133</td>
<td>90</td>
<td>0.0219</td>
<td></td>
</tr>
<tr>
<td>11K</td>
<td>0.0233</td>
<td>—</td>
<td>0.0152</td>
<td></td>
</tr>
<tr>
<td>12G</td>
<td>0.0017</td>
<td>100</td>
<td>0.0335</td>
<td></td>
</tr>
<tr>
<td>13G</td>
<td>0.0010</td>
<td>120</td>
<td>0.0368</td>
<td>0.605</td>
</tr>
<tr>
<td>10G</td>
<td>0.0067</td>
<td>120</td>
<td>0.0245</td>
<td>0.581</td>
</tr>
<tr>
<td>1K</td>
<td>0.0233</td>
<td>140</td>
<td>0.0138</td>
<td>0.482</td>
</tr>
<tr>
<td>8K</td>
<td>0.0233</td>
<td>150</td>
<td>0.0132</td>
<td>0.447</td>
</tr>
<tr>
<td>25G</td>
<td>0.0233</td>
<td>180</td>
<td>0.0142</td>
<td></td>
</tr>
<tr>
<td>13K</td>
<td>0.0133</td>
<td>150</td>
<td>0.0161</td>
<td>0.424</td>
</tr>
<tr>
<td>17K</td>
<td>0.0133</td>
<td>170</td>
<td>0.0146</td>
<td>0.425</td>
</tr>
<tr>
<td>27G</td>
<td>0.0133</td>
<td>110</td>
<td>0.0185</td>
<td></td>
</tr>
</tbody>
</table>
temperature corresponding to about 1550 K [19], was used to locate the base of the primary dendrite array, where the last interdendritic liquid solidified just before quench. The experimentally observed lengths (cell length) of the primary dendrite arrays protruding into the melt, varying from 0.45 to 0.6 cm, are also listed in Table 1. Figure 1(b) shows a typical transverse microstructure in the directionally solidified specimen. As seen in this figure, the distribution of primary dendrites was observed to be very uniform across the entire specimen cross-section.

3.1. Primary dendrite spacings

At high constitutional supercooling gradients, i.e. for growth conditions where $G_1$ is much larger than $G_c$ or $G_1/m_1$, the dendrite tip radius predictions from dendrite growth models for binary alloys are nearly the same [6, 21]. Incorporation in eqn. (1) of the dependence of the tip radii on the growth rate and thermal gradient, as predicted from these models, yields the following relationship for the primary dendrite spacings

$$\lambda_i = (B64 \sigma D_i k\Delta T_0/\Delta S)^{0.25} G_1^{-0.5} R^{-0.25}$$

where $\sigma$ is the liquid–solid surface energy, $\Delta T_0 = m_1 C_0/(k-1)/k$ is the alloy freezing range, $\Delta S$ is the entropy of fusion per unit volume and $B$ is a constant. The value of $B$ is respectively 1, 16 or 28 for the dendrite growth models of Burden and Hunt [5], Laxmanan [21] and Trivedi [3]. As mentioned earlier, despite its multicomponent nature, PWA-1480 shows a solidification behavior which is similar to a binary alloy. We will therefore compare the experimentally observed primary arm spacing data with those predicted from analytical models, even though the models are strictly only valid for simple binary alloys directionally solidified in a positive thermal gradient.

Figure 2 shows the dependence of the primary dendrite spacings $\lambda_i$ on the thermal gradient $G_1$ and growth rate $R$ in directionally solidified single-crystal specimens of PWA-1480. The abscissa in this figure was selected to be $R^{-0.25} G_1^{-0.5}$, with $R$ measured in centimeters per second, and $G_1$ in degrees Kelvin per centimeter, as suggested by eqn. (2). The solid line in this figure, described by the following relationship, is the linear least-squared fit to the data.
solidified single-crystal linear least-squared fit to the experimental data. The broken line is the prediction from Trivedi’s model [3].

\[
\lambda_1 = (-0.00128 \pm 0.0013) + (0.0728 \pm 0.0040)R^{-0.25}G_i^{-0.5}
\]

(3)

where \( \lambda_1 \) is in centimeters, \( R \) in centimeters per second, and \( G_i \) in degrees Kelvin per centimeter. A high value of correlation coefficient, 0.94, and very low value of relative standard deviation for this fit, 3%, denote an excellent fit of the experimental data to the above linear relationship. The linear dependence of primary dendrite spacings on \( G_i^{-0.5}R^{-0.25} \), predicted by Hunt’s model, is also valid for our complex multicomponent superalloy, PWA-1480.

We will now estimate the solutal diffusivity in the melt by equating \( \sqrt{B64\sigma D_k \Delta T_0 / \Delta S}^{0.25} \) (eqn. (2)) to the experimental value of the slope, 0.0728 (eqn. (3)). The alloy freezing range, \( \Delta T_0 \), is 35 K (for PWA-1480 liquidus 1610 K, and solidus 1575 K). The value of \( \sigma \) and the heat of fusion per unit volume of PWA-1480 is assumed to be the same as that of nickel, 255 erg cm\(^{-2}\) [22] and \( 2.77 \times 10^{10} \) erg cm\(^{-3}\) [23] respectively. The entropy \( \Delta S \) value used here is that calculated at the alloy liquidus temperature, 1610 K. For \( B=28 \), as used in Trivedi’s model [3], the \( D_k \) value thus obtained is \( 3.0 \times 10^{-5} \) cm\(^2\) s\(^{-1}\). The experimentally observed partition coefficients for the various solutes in superalloy PWA-1480 are in the range 0.4–1.7 [19]. Thus the solute diffusivity in the melt \( D_k \) is predicted to be in the range 1.8–7.0 \( \times 10^{-5} \) cm\(^2\) s\(^{-1}\). This is in reasonable agreement with the reported solutal diffusivity, about 2–8 \( \times 10^{-5} \) cm\(^2\) s\(^{-1}\) [9], obtained from estimations based on the use of the constitutional supercooling criterion for plane front solidification of several superalloys. Based on these physical parameters, the primary dendrite spacings predicted from eqn. (2) are plotted in Fig. 2. The broken line in Fig. 2 represents the values predicted from the approach used by Trivedi [3] for \( D_k = 8 \times 10^{-5} \) cm\(^2\) s\(^{-1}\) and \( k = 0.6 \) corresponding to tantalum in PWA-1480 [19]. It should also be mentioned that the predictions obtained by combining the models of Laxmanan [21] and Hunt [1] are also approximately similar (within 15% of the values indicated by the broken line in Fig. 2). Considering the uncertainties in the various physical properties and the assumption of treating PWA-1480 as a binary alloy, the experimental data are in good quantitative agreement with the predictions from the analytical models.

A comparison of the experimentally observed tip radius of 12 \( \mu m \) for one specimen, \( G_i = 120 \) K cm\(^{-1}\), \( R = 0.001 \) cm s\(^{-1}\), with predictions from dendrite growth models also shows a good agreement. The tip radius value predicted from Trivedi’s model [3] is 7.2 \( \mu m \) for \( D_k = 2 \times 10^{-5} \) cm\(^2\) s\(^{-1}\) and 14.4 \( \mu m \) for \( D_k = 8 \times 10^{-5} \) cm\(^2\) s\(^{-1}\).

### 3.2. Side-branch coarsening kinetics

In the following, the experimentally observed side-branch coarsening behavior for the superalloy PWA-1480 will be compared with predictions from Kirkwood’s analytical model [18], treating the multicomponent superalloy as if it were a binary alloy. As mentioned earlier, this model is especially suited to partial directional solidification and quenching experiments. The model predicts the following behavior:

\[
d^3 = [128\sigma(T/H)/(D_k/\Delta T_0)]t
g,(H/T)
\]

(4)

where \( d \) is the side-branch spacing corresponding to the coarsening time \( t \) and \( H/T \) is the entropy of fusion per unit volume.

Figure 3(a) plots \( \log_{10}(d) \) vs. \( \log_{10}(t) \) data obtained from several experiments. The corresponding growth speeds are indicated in the figure. The straight line drawn through the data is the linear least-squared fit. The fit is given as

\[
\log_{10}(d) = (1.021 + 0.014) + (0.33 + 0.001)\log_{10}(t)
\]

(5)

where \( d \) is measured in micrometers and \( t \) in seconds.

There is considerable scatter in the data, but the experimentally observed slope of the curve, 0.33, is in agreement with that expected from the above coarsening relationship, eqn. (4). The same
Fig. 3. Side-branch coarsening kinetics in superalloy PWA-1480, the line is the linear least squared fit through the data: (a) log-log plot of side branch spacing $d$ vs. coarsening time $t$; (b) plot of the cube of the side branch spacing $d$ vs. the coarsening time $t$.

The following conclusions can be drawn from this study on single-crystal nickel base superalloy PWA-1480 specimens directionally solidified in a positive thermal gradient. In this study the primary dendrite spacings and side-branch coarsening kinetics were investigated. The alloy PWA-1480, nominal composition Ni-12Ta-10.4Cr-5Co-5Al-4W-1.5Ti (weight per cent), has been treated as a simple binary alloy for comparison of the experimental data with predictions from analytical models.

1. The dependence of the primary dendrite spacing $\lambda_1$ on the growth rate $R$ and thermal gradient $G_l$ is given as:

$$\lambda_1 = (-0.00128 \pm 0.0013) + (0.0728 \pm 0.0040) R^{-0.25} G_l^{-0.5}$$

with $\lambda_1$, measured in centimeters, $R$ in centimeters per second, and $G_l$ in degrees Kelvin per centimeter. This relationship is the same as that predicted from the analytical model developed by Hunt [1] for binary alloys. A reasonable quantitative agreement is obtained between the experimental data and the predictions from the analytical model of Trivedi [3].

2. The secondary arm coarsening kinetics in PWA-1480 obeys the following relationship

$$\log_{10}(d) = (1.021 \pm 0.014) + (0.33 \pm 0.001) \log_{10}(t)$$

where $d$ (in micrometers) is the side-branch spacing corresponding to the coarsening time $t$ (in seconds). This behavior is in agreement with predictions from the model of Kirkwood [18] developed for binary alloys.
Acknowledgments

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References