

SOLIDIFICATION MODELING OF Nb BEARING SUPERALLOYS

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ABSTRACT

The solidification behavior of experimental Ni base and Fe base superalloys containing Nb, Si, and C was studied using differential thermal analysis (DTA) and microstructural characterization techniques. The solidification reaction sequences responsible for microstructural development were found to be similar to those expected in the Ni-Nb-C ternary system, where the solute-rich interdendritic liquid exhibited two eutectic-type reactions at the terminal stages of solidification: $L \rightarrow (\gamma + NbC)$ and $L \rightarrow (\gamma + Laves)$. A pseudo ternary γ -Nb-C approach was developed to provide a quantitative description of solidification behavior for these experimental alloys. Solute redistribution calculations in the model are based on a previous approach developed by Mehrabian and Flemings, with modifications made to account for the high diffusion rate of C in the solid. Solidification parameters for Nb and C were determined through DTA and electron probe microanalysis techniques and used as inputs to the model. Reasonable agreement is found between calculated volume fractions of the γ /NbC and γ /Laves constituents and those measured experimentally. The modelling results permit detailed descriptions of the relation between alloy composition and microstructural evolution during solidification.

INTRODUCTION

Ni base and Fe base Nb-strengthened alloys represent a significant portion of superalloys currently in use. Commercial examples include alloys IN625, IN706, IN718, IN903, IN909, and Thermo-Span, to name a few. These alloys are often used for components which are fabricated by solidification processes such as casting and fusion welding. Previous studies conducted on commercial superalloys [1,2,3] have shown that minor additions of Nb, Si, and C strongly affect the type and amount of secondary phases which form during the terminal stages of solidification. In particular, two types of secondary phases, NbC and Laves, are known to form. Although several detailed investigations have been conducted on specific commercial alloy systems, no general model has been developed to quantitatively predict microstructural evolution during solidification. Thus, a solidification model was developed and verified on a wide range of experimental Fe base and Ni base alloys containing systematic variations in Nb, Si, and C. This article will focus on the effects of the two most important solute elements: Nb and C. More details, which describe the effects of Si and results of solidification cracking studies, can be found elsewhere [4].

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EXPERIMENTAL PROCEDURE

The experimental alloys (Table 1) exhibit factorial variations in Fe, Nb, Si, and C at two levels. The alloys were prepared at Sandia National Laboratory by investment casting. Differential thermal analysis (DTA) was conducted on a Netzsch STA 409 differential thermal analyzer using 500 to 550 mg samples. The DTA system was calibrated to melt within 2 °C of the melting point of pure Ni (1455 °C). Samples were melted and solidified under flowing argon in alumina crucibles using pure Ni as the reference material. Scanning Electron Microscopy was conducted on autogenous GTA weld metals using a JEOL 6300 Field Emission Gun Scanning Electron Microscope (FEG-SEM) at an accelerating voltage of 15kV. Quantitative Image Analysis (QIA) was performed on GTA welds and selected DTA samples. Area fractions of total eutectic type contents and, where possible, individual γ /NbC and γ /Laves eutectic type constituents were measured on each sample with at least ten SEM photomicrographs. Area fractions were assumed to be equivalent to volume fractions. Electron Probe Micro-Analysis (EPMA-WDS) was conducted on a JEOL 733 Probe at an accelerating voltage of 15kV and beam current of 20 nA. The K_{α} lines were used for Fe, Ni, Cr, and Si while the L_{α} line was used for Nb. Raw data were reduced to weight percentages using the ZAF algorithm.

Table 1. Alloy compositions. All values in weight percent.

Alloy	Fe	Ni	Cr	Nb	Si	C	P	S
1	10.49	68.53	18.90	1.93	0.08	0.017	0.004	0.003
1.5	10.75	67.95	19.21	2.00	0.03	0.052	0.004	0.003
2	11.12	68.20	19.12	1.95	0.06	0.132	0.004	0.002
3	10.70	68.11	19.02	1.82	0.38	0.010	0.004	0.003
3.5	10.39	66.80	19.29	1.94	0.41	0.075	0.004	0.003
4	10.72	67.60	19.08	1.91	0.40	0.155	0.004	0.001
5	10.84	65.79	18.98	5.17	0.05	0.013	0.005	0.010
6	10.88	65.22	18.89	4.87	0.08	0.161	0.005	0.007
7	10.70	65.53	19.30	4.86	0.52	0.010	0.005	0.009
7.5	10.82	63.93	18.54	4.92	0.46	0.081	0.005	0.004
8	10.80	64.96	18.90	4.72	0.52	0.170	0.005	0.007
9	46.03	33.56	19.31	1.66	0.10	0.003	0.006	0.003
10	46.69	32.80	19.70	1.66	0.01	0.108	0.006	0.002
11	45.38	32.80	19.53	1.77	0.57	0.004	0.006	0.002
11.5	47.38	31.05	19.64	1.84	0.67	0.116	0.006	0.001
12	45.28	32.39	19.89	1.93	0.61	0.079	0.006	0.002
13	44.55	31.24	19.63	4.42	0.02	0.015	0.007	0.003
14	44.05	31.93	19.52	4.51	0.08	0.210	0.006	0.002
15	45.40	30.03	19.54	4.88	0.66	0.010	0.007	0.003
16	44.47	30.89	19.45	4.77	0.64	0.216	0.006	0.002

RESULTS AND DISCUSSION

Solidification Sequences

The general solidification sequence of these alloys can be described by a three step process: 1) Primary $L \rightarrow \gamma$ solidification in which the interdendritic liquid becomes enriched in Nb and C, followed by 2) a eutectic-type $L \rightarrow (\gamma + NbC)$ reaction which depletes the interdendritic liquid of C, and 3) termination of solidification by a second eutectic-type reaction $L \rightarrow (\gamma + Laves)$. A typical DTA trace (for Alloy 16) showing this solidification sequence is presented in Figure 1. The microstructure of a GTA weld in Alloy 16 is shown in Figure 2, where the γ/NbC and $\gamma/Laves$ eutectic-type constituents are readily apparent. Identification of the NbC and Laves phases was confirmed through EPMA measurements on coarse phases in DTA samples and backscattered electron kikuchi patterns collected from finer phases in weld metal samples [4]. The Ni base alloys with high C/low Nb (Alloys 2, 3.5, and 4) represent the only exception to this general solidification sequence. These alloys did not exhibit any of the Laves phase, indicating solidification terminated with the $L \rightarrow (\gamma + NbC)$ reaction. The individual amounts of the γ/NbC and $\gamma/Laves$ constituents were high enough to quantify by QIA in a number of alloys, and these results are shown in Figure 3. The results from Ni base and Fe base alloys are aligned to permit comparisons among alloys with similar solute levels. Carbon additions promote formation of the γ/NbC constituent while Fe and Si increase the amount of the $\gamma/Laves$ constituent. The effect of Nb depends on the C content. At high C levels, Nb additions will promote formation of γ/NbC . When the C level is low, Nb generally promotes formation of the $\gamma/Laves$ constituent. These effects have been documented in commercial alloy systems as well [1,2].

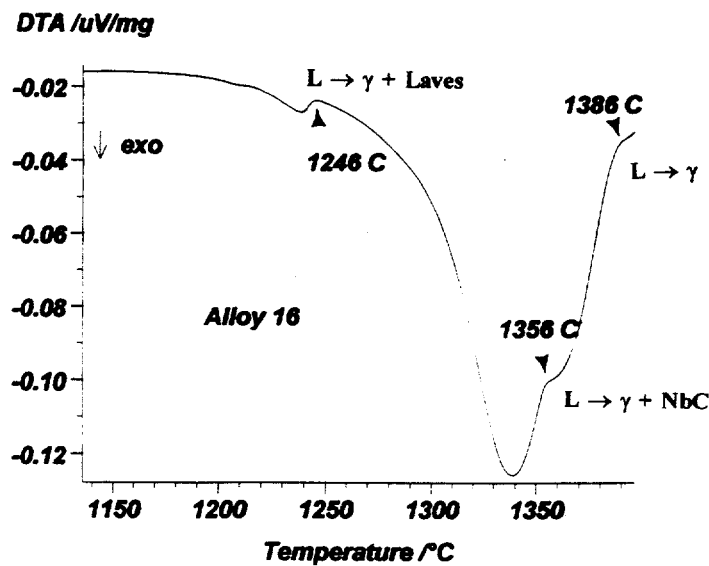


Figure 1. DTA solidification scan for Alloy 16.

The solidification reactions in these experimental multi-component alloys are similar to those expected in the pure ternary Ni-Nb-C system [5,6]. The liquidus projection for this system is shown in Figure 4. The liquidus projection exhibits three primary phase fields which are of interest here: γ , NbC, and Ni_3Nb . A primary C (graphite) phase field exists at high C contents which is not of importance. Additions of Fe, Cr, and Si to the Ni-Nb system are well known to promote Laves at the expense of Ni_3Nb in commercial superalloys as well as the experimental alloys utilized in this work [7,8]. Thus, by replacing Ni_3Nb with Laves, the Ni-Nb-C liquidus projection can be utilized as a guide in developing a description of the solidification reactions in these alloys.

Solidification begins with formation of primary γ dendrites which, upon forming, reject Nb and C to the liquid. As solidification proceeds, the liquid composition moves away from the Ni corner, becoming progressively richer in Nb and C until the line of two fold saturation between γ and NbC is reached. At this point, the maximum solubility of Nb and C in the austenite matrix is exceeded and the γ and NbC phases form simultaneously from the liquid by a eutectic-type reaction as the liquid

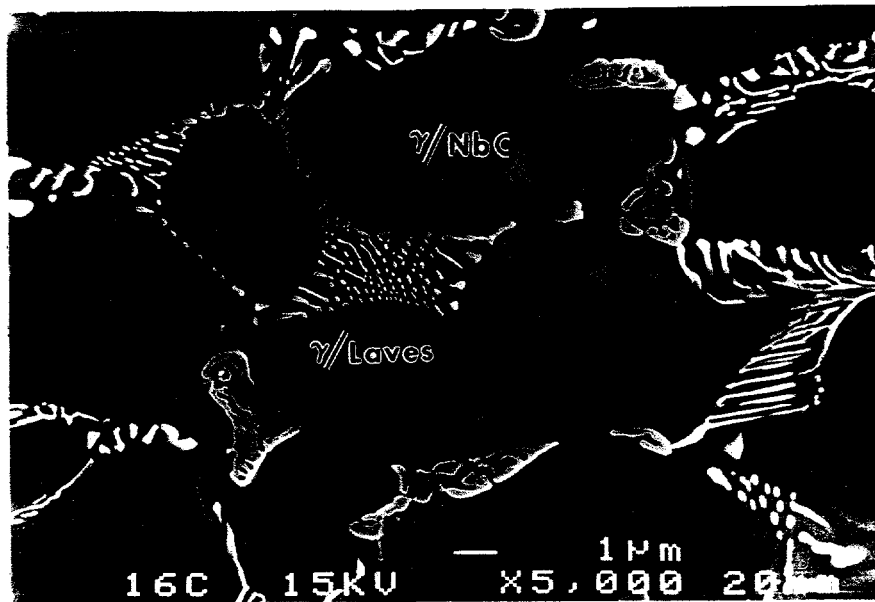


Figure 2. SEM photomicrograph of γ/NbC and γ/Laves constituents in GTA weld of Alloy 16.

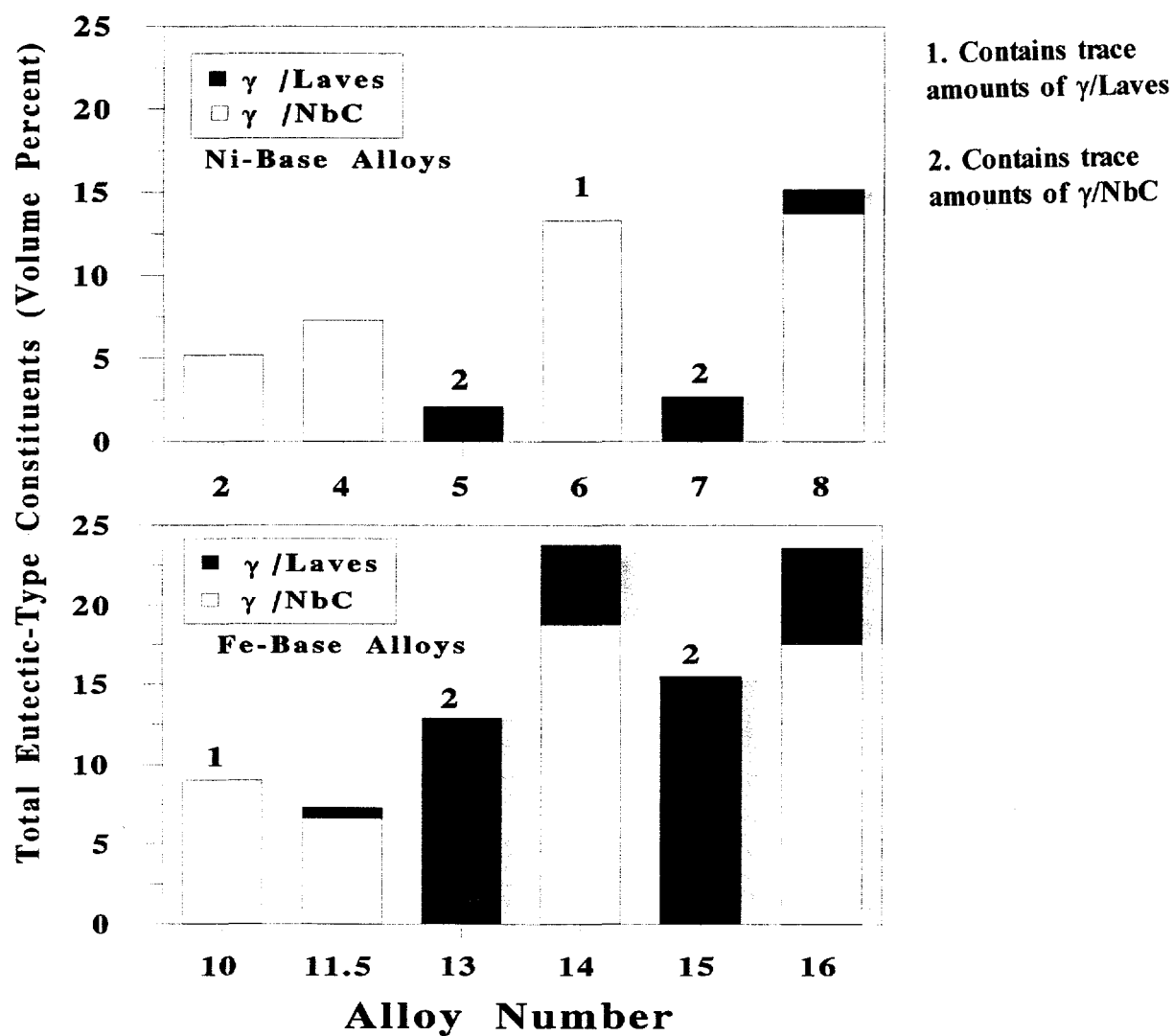


Figure 3. Quantitative image analysis results.

composition travels down the line of two fold saturation. Due to its' high C content (~ 9.5 wt%, [9]), the formation of NbC depletes the liquid of C while the Nb content of the liquid continues to increase. For Alloys 2, 3.5, and 4, solidification is completed along the line of two fold saturation between γ and NbC, as no Laves phase was observed in these alloys. The initial reaction sequence is similar for the remaining alloys, except that solidification does not terminate with the $L \rightarrow (\gamma + NbC)$ reaction. Instead, the liquid composition continues to travel down the line of two fold saturation between γ and NbC, becoming progressively enriched in Nb (and depleted in C). According to the proposed Ni-Nb-C liquidus projection, solidification should terminate with the ternary $L \rightarrow (\gamma + NbC + Laves)$ reaction where the liquidus surface is at a minimum. Under this condition, the Laves and ternary γ and NbC phases should be intermixed. However, this type of structure is not observed in any of the alloys which form both the NbC and Laves phases (e.g., Figure 2). Instead, the γ /NbC and γ /Laves eutectic-type constituents are always distinctly separated. This suggests that the pseudo-ternary liquidus projection for the alloys in this work is more properly represented by a class II reaction [10]. In this case, the local minimum on the liquidus surface occurs where the line of two fold saturation separating the γ and Laves phases intersects the Ni-Nb "binary" side of the diagram. The liquidus surface in Figure 4 has been drawn to reflect this proposed change. The solidification sequence for alloys with this type of surface would occur as follows. After primary γ solidification, the liquid composition travels along the line of two fold saturation separating the γ and NbC phases during the $L \rightarrow (\gamma + NbC)$ reaction. This process continues until the class II reaction is reached, at which point the NbC stops forming as the $L \rightarrow (\gamma + NbC)$ reaction is replaced by the $L \rightarrow (\gamma + Laves)$ reaction. Solidification is completed at the Ni-Nb "binary" side of the diagram by the $L \rightarrow (\gamma + Laves)$ reaction. This solidification sequence accounts for the two spatially separate γ /NbC and γ /Laves eutectic-type constituents which are observed experimentally.

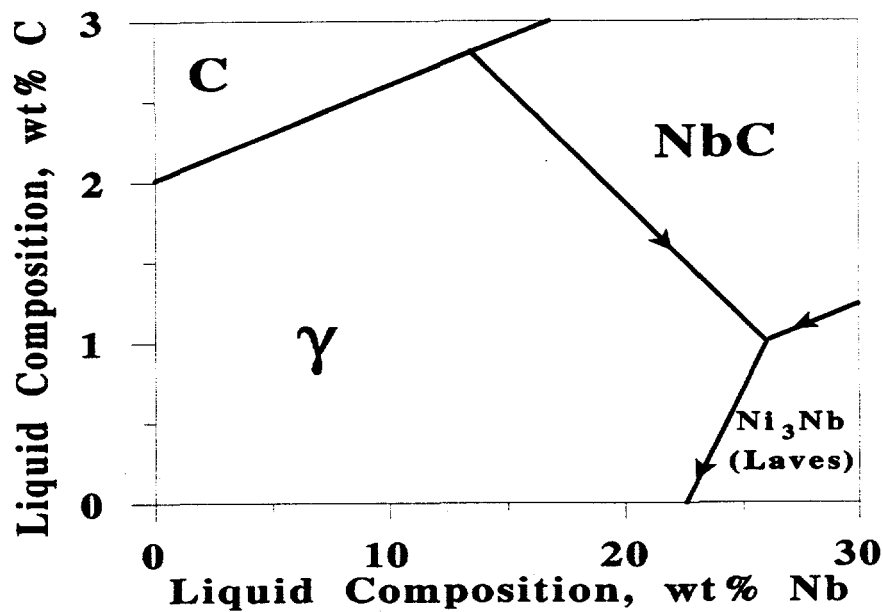


Figure 4. Ni-rich corner of the Ni-Nb-C liquidus projection.

Pseudo-Ternary Solidification Model

The Ni-Nb-C ternary system forms the basis for developing a model for quantitative prediction of microstructural evolution during solidification of these multi-component alloys. The main objective of such a model is to predict the composition and relative fractions of each solid phase forming during the solidification process. Several key elements are required to construct the solidification model. These include the equilibrium distribution coefficients, k , for the solutes Nb and C, a liquidus surface to identify the lines of two fold saturation separating primary phase fields, and an appropriate solute redistribution model which describes solutal transport in the liquid and solid during solidification. The measurement of k values and liquidus surfaces for these alloys are described briefly below (more detail can be found in [4]) followed by a description of the solute redistribution model.

Equilibrium Distribution Coefficients. The equilibrium distribution coefficients are summarized in Table 2. These values were determined by a combination of EPMA and DTA techniques [4]. The k values for Nb and C are less than unity, indicating they segregate preferentially to the liquid during solidification. In contrast, the k values for the "solvent" elements (Fe,Ni,Cr) are all close to unity, indicating they show little preference for segregation to either the solid or liquid. It is interesting to note that the value of k_{Nb} is significantly lower in the Fe base alloys ($k_{Nb} = 0.25$) compared to the Ni base alloys ($k_{Nb} = 0.46$). Reference to the Ni-Nb phase diagram [11] indicates that the maximum solid solubility of Nb in γ -Ni is 18.2 wt% Nb (at 1286°C). In contrast, γ -Fe can only dissolve a maximum of 1.5 wt% Nb at a comparable temperature of 1210°C [11]. Based on these differences, it might be expected that iron additions to these alloys would decrease the solubility of Nb in the γ -(Fe,Ni,Cr) matrix. Thus, as the Fe-rich γ dendrites begin to form, less Nb is taken into the solid and, as a result, more is rejected to the liquid (in comparison to the Ni base alloys). This effect would induce a concomitant decrease in the value of k_{Nb} . In fact, a general correlation between k_{Nb} and nominal iron content has recently been observed [4] over a wide range of matrix compositions.

Pseudo-Ternary Solidification Surfaces. Since the alloys in this research contain six elements (Fe,Ni,Cr,Nb,Si,C), the term "ternary liquidus projection" can not be properly applied. In addition, the experimental data utilized to construct such projections were derived from welds prepared under non-equilibrium conditions [4]. Thus, the term "pseudo-ternary solidification surface" more appropriately describes these diagrams. The distribution coefficients for the matrix elements (Fe,Ni,Cr) in commercial systems [1,3] and these experimental alloys are similar and take on a value close to unity, indicating they all behave in a comparable manner and exhibit little tendency for segregation. Thus, the (Fe,Ni,Cr) elements in the γ matrix are all grouped together to form one "component" of the diagram. Based on the similarity between reaction sequences observed in these multi-component alloys and those expected in the Ni-Nb-C system, Nb and C are chosen as the solute elements in this pseudo-ternary solidification model. (Effects from Si are ignored here, but are described in more detail in reference [4].) Lastly, the Ni_3Nb phase is replaced by the Laves phase. The system can now be represented by a γ -Nb-C solidification diagram which exhibits primary phase fields of γ , NbC, and Laves. The experimentally determined boundaries separating the γ /NbC and γ /Laves phase fields are shown in Figure 5. The position of these boundaries was determined with a combination of QIA and EPMA techniques. (The position of the NbC/Laves boundary was not determined as it is not needed for the present analysis. The line drawn is provided only for clarity.)

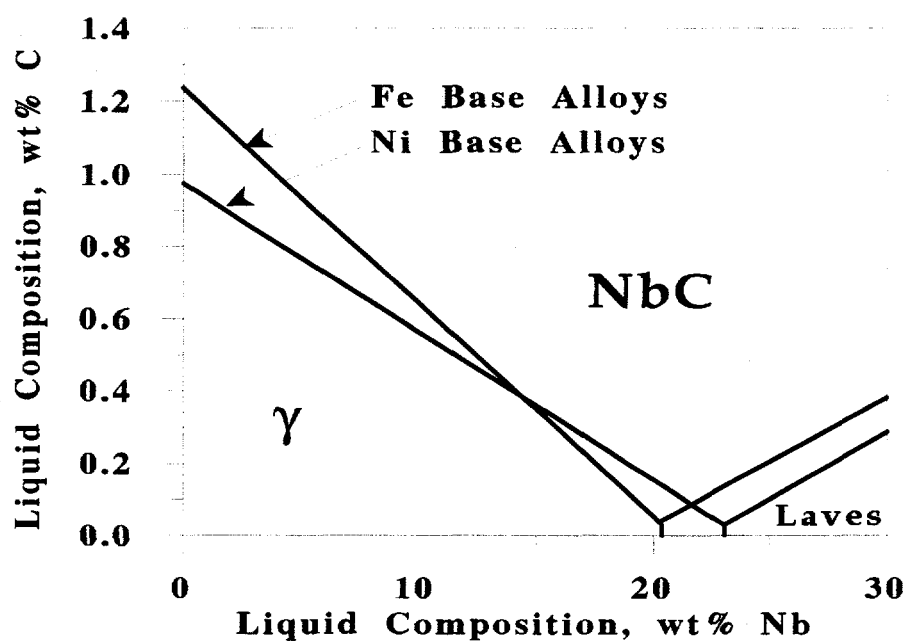


Figure 5. Pseudo-ternary γ -Nb-C solidification surfaces for Ni base and Fe base alloys.

Table 2. Distribution coefficients.

Alloys	Fe	Ni	Cr	Nb	C
Ni-Base	1.00	1.02	1.06	0.46	0.27
Fe-Base	1.06	1.00	1.02	0.25	0.27*

*Estimated from the Ni base alloy data

Solute Redistribution Model Quantitative treatment of solute redistribution during solidification of these alloys requires two sets of expressions: one to describe the solidification path (i.e., variation in liquid composition) during primary $L \rightarrow \gamma$ solidification and one to describe the variation in fraction liquid, f_l , with liquid composition, C_l , during the $L \rightarrow (\gamma + NbC)$ eutectic-type reaction. The model developed below is based on a previous treatment proposed by Mehrabian and Flemings [12], with modifications being made to account for the high diffusion rate of carbon in the solid. More details of the model are provided in [4].

The solidification path during primary $L \rightarrow \gamma$ solidification is obtained by first writing the expressions describing the relation between f_l and C_l for each solute (Nb and C). For Nb, diffusion in the solid is negligible [3,4] and the relation between f_l and $C_{l,Nb}$ is given by the well-known Scheil equation [13].

$$f_l = \left(\frac{C_{l,Nb}}{C_{o,Nb}} \right)^{\frac{1}{k_{Nb}-1}} \quad (1)$$

Eq. (1) also assumes that dendrite tip undercooling is negligible, thermodynamic equilibrium is maintained at the solid/liquid interface, and diffusion is infinitely fast in the liquid. For transport of C, diffusion in the solid is typically fast enough to maintain equilibrium [4,14], and the equilibrium lever law provides the relation between f_l and $C_{l,C}$

$$f_l = \frac{C_{o,C} - k_C C_{l,C}}{(1 - k_C) C_{l,C}} \quad (2)$$

The f_l values given by eqs. (1) and (2) must be equivalent. Thus, by equating eqs. (1) and (2) and solving for $C_{l,Nb}$, a solidification path relation is obtained

$$C_{l,Nb} = C_{o,Nb} \left[\frac{C_{o,C} - k_C C_{l,C}}{(1 - k_C) C_{l,C}} \right]^{k_{Nb}-1} \quad (3)$$

Eq. (3) describes the solidification path of the liquid during primary solidification of γ assuming that Nb diffusion in the solid is negligible while C diffuses fast enough to maintain equilibrium. This relation is physically consistent at the limits of solidification. At the beginning of solidification, the liquid composition should equal the nominal composition for both solutes. Insertion of $C_{l,C} = C_{o,C}$ into eq. (3) yields $C_{l,Nb} = C_{o,Nb}$. Under equilibrium conditions for carbon, the liquid will reach a maximum concentration of $C_{o,C}/k_C$ when solidification is complete while the Nb concentration in the liquid should tend toward infinity as predicted by the Scheil equation. Insertion of $C_l = C_{o,C}/k_C$ into equation (3) yields $C_{l,Nb} = \infty$.

When the liquid composition intersects a line of two fold saturation on the liquidus surface, the remaining liquid (f_l) will transform to the eutectic-type constituents (i.e., γ/NbC and $\gamma/Laves$). Thus, the value of f_l at that point represents the total amount of eutectic-type constituent which exists in the final solidification microstructure. In order to calculate the total volume fraction of eutectic-type

constituent, the line of two fold saturation on the liquidus surface is first represented by a linear equation. For the two solutes of interest here:

$$C_{1,C} = a + bC_{1,Nb} \quad (4)$$

Here, a is the C content at the line of two fold saturation between γ and NbC at 0 wt% Nb and b is the slope of the line. In order to find the intersection of the primary solidification path and the line of two fold saturation, eq. (4) is first solved in terms of $C_{1,Nb}$

$$C_{1,Nb} = \frac{C_{1,C} - a}{b} \quad (5)$$

and eqs. (3) and (5) are equated

$$\frac{C_{1,C} - a}{b} = C_{o,Nb} \left(\frac{C_{1,C}}{C_{o,C}} \right)^{\frac{k_{Nb} - 1}{k_C - 1}} \quad (6)$$

Eq. (6) provides a unique value of carbon content in the liquid, $C_{1,C}$, in terms of the nominal alloy composition ($C_{o,Nb}$ and $C_{o,C}$) and distribution coefficients (k_{Nb} and k_C) for the two solutes. This value of $C_{1,C}$ defines the C content in the liquid where the solidification path intersects the line of two fold saturation. The corresponding Nb content in the liquid, $C_{1,Nb}$, is given by substituting this value of $C_{1,C}$ into either eq. (3) or (5). Once this particular value of $C_{1,Nb}$ is known, it is used in eq. (1) to calculate the remaining fraction of liquid which transforms to eutectic-type constituents.

Once the primary solidification path intersects the line of two fold saturation between γ and NbC, additional terms are needed in the mass balance equations to account for formation of two solid phases ($\gamma + \text{NbC}$) from the liquid. This has been considered in detail [4] for the present set of conditions (i.e., negligible solid state diffusion of Nb and infinitely fast diffusion of carbon) and the final relation between fraction liquid (f_1) and liquid composition ($C_{1,C}$, $C_{1,Nb}$) is given by

$$\frac{df_1}{dC_{1,Nb}} = -\frac{1}{(1-k_{\gamma,Nb})} \frac{f_1}{C_{1,Nb}} - \left[\frac{(k_{NbC,Nb} - k_{\gamma,Nb})}{(1-k_{\gamma,Nb})} \right] \left[\frac{bk_{\gamma,C}C_{o,C} - bC_{NbC,C}[f_1 + k_{\gamma,C}(1-f_1)]}{(C_{NbC,C} - k_{\gamma,C}C_{1,C})^2} \right] \quad (7)$$

Where k_{ij} is the distribution coefficient (given by the ratio of solid to liquid composition) for element j in the i phase. The distribution coefficients for Nb and C in γ are assumed to be constant at the values shown in Table 2. The NbC phase is a stoichiometric compound so the composition of NbC, $C_{NbC,Nb}$, and $C_{NbC,C}$ are constant at 90.5 wt% Nb and 9.5 wt% C, respectively [4,9]. Thus, the distribution coefficients for Nb and C in the NbC phase can be calculated from knowledge of the liquid composition at any point during solidification.

Eq. (7) can be used to calculate solute redistribution during the eutectic-type $L \rightarrow (\gamma + \text{NbC})$ reaction using a finite difference technique in the following manner: The first values of f_1 , $C_{1,Nb}$, and $C_{1,C}$ are calculated from the intersection of the solidification path and line of two fold saturation, as given by eqs. (3), (5), and (6). These values are used to calculate the first value of $(df_1/dC_{1,Nb})_1$. The value of $(C_{1,Nb})_1$ is incremented by a very small amount, $\Delta C_{1,Nb}$, along the line of two fold saturation to a new value of $(C_{1,Nb})_2 = (C_{1,Nb})_1 + \Delta C_{1,Nb}$. The corresponding value of $(C_{1,C})_2$ is determined through eq. (4). The corresponding value of $(f_1)_2$ is calculated by a small extrapolation of the value of $(df_1/dC_{1,Nb})_1$ by

$$(f_1)_2 = (f_1)_1 + \left(\frac{df_1}{dC_{1,Nb}} \right)_1 \Delta C_{1,Nb} \quad (8)$$

The values of $(C_{1,Nb})_2$, $(C_{1,C})_2$, and $(f_1)_2$ are then put back into eq. (7) and the calculations are repeated. The values of $k_{NbC,Nb}$ and $k_{NbC,C}$ are calculated at each step by knowledge of the NbC and liquid compositions. When the values of $C_{1,Nb}$ and $C_{1,C}$ reach the class II reaction point where the $L \rightarrow (\gamma + NbC)$ transformation is replaced by the $L \rightarrow (\gamma + Laves)$ reaction, the remaining fraction liquid at that point transforms to the γ /Laves eutectic-type constituent and solidification is complete. This procedure provides the relation between the fraction liquid and liquid composition at any point along the line of two fold saturation which, in turn, permits calculation of the volume fractions of each eutectic-type constituent present in the as solidified microstructure. More details of the calculation scheme are presented in [4].

Modeling Results

The parameters used to calculate the compositions and fractions of each phase forming during solidification are summarized in Table 3. The values of $C_{Nb,L \rightarrow (\gamma+Laves)}$ and $C_{C,L \rightarrow (\gamma+Laves)}$ listed in Table 3 represent the point of the class II reaction on the solidification. This point was determined through EPMA data [4]. When the Nb and C contents in the liquid reach this point, the remaining liquid (f_1) transforms to the γ /Laves constituent ($f_{\gamma Laves}$) and the calculations are terminated. The constants a and b were determined by linear regression analysis from the point of $(C_{Nb,L \rightarrow (\gamma+Laves)}, C_{C,L \rightarrow (\gamma+Laves)})$ to the data points on the line of two fold saturation.

Table 3. Summary of solidification parameters used in finite difference calculations.

QUANTITY	Ni Base Alloys	Fe Base Alloys	UNITS
$k_{\gamma Nb}$	0.46	0.25	unitless
$k_{\gamma C}$	0.27	0.27	unitless
a	0.98	1.24	wt% C
b	-0.04	-0.06	wt% C/wt% Nb
$C_{Nb,L \rightarrow (\gamma + Laves)}$	23.1	20.4	wt% Nb
$C_{C,L \rightarrow (\gamma + Laves)}$	0.03	0.03	wt% C

Primary $L \rightarrow \gamma$ Solidification The solidification path of each alloy was calculated via eq. (3) and plotted on the respective solidification surface. The results of the calculations are presented in Figure 6. Figures 6a and 6b plot typical results for the Ni base and Fe base alloys that have similar Nb levels, but variations in C content. (The nominal alloy composition is easily read by the start of the solidification path.) Figure 6c compares the solidification paths of Ni base alloys with similar C contents and variations in Nb, and Figure 6d compares the solidification paths of a Ni base and Fe base alloy with similar levels of solute elements (Nb, Si, and C). The alloy number of each path is denoted at the intersection point between the primary solidification path and line of two fold saturation. The solidification path for each alloy initiates at the nominal composition and generally progresses toward the line of two fold saturation between γ and NbC as the liquid becomes enriched in Nb and C. When the primary $L \rightarrow \gamma$ path strikes the line of two fold saturation, solidification then proceeds along the line of two fold saturation in the direction of decreasing temperature (i.e., toward the γ - NbC "binary"). The amount of solid formed during primary solidification is proportional to the total distance of the primary solidification path. This relation is useful, as it allows for fast, qualitative interpretations of alloying effects by simple visual inspection of the results.

These diagrams provide several key points concerning the solidification behavior of these alloys. First, it is readily apparent that the point of intersection between the primary solidification path and line of two fold saturation is a strong function of C content. Although C additions are intuitively expected

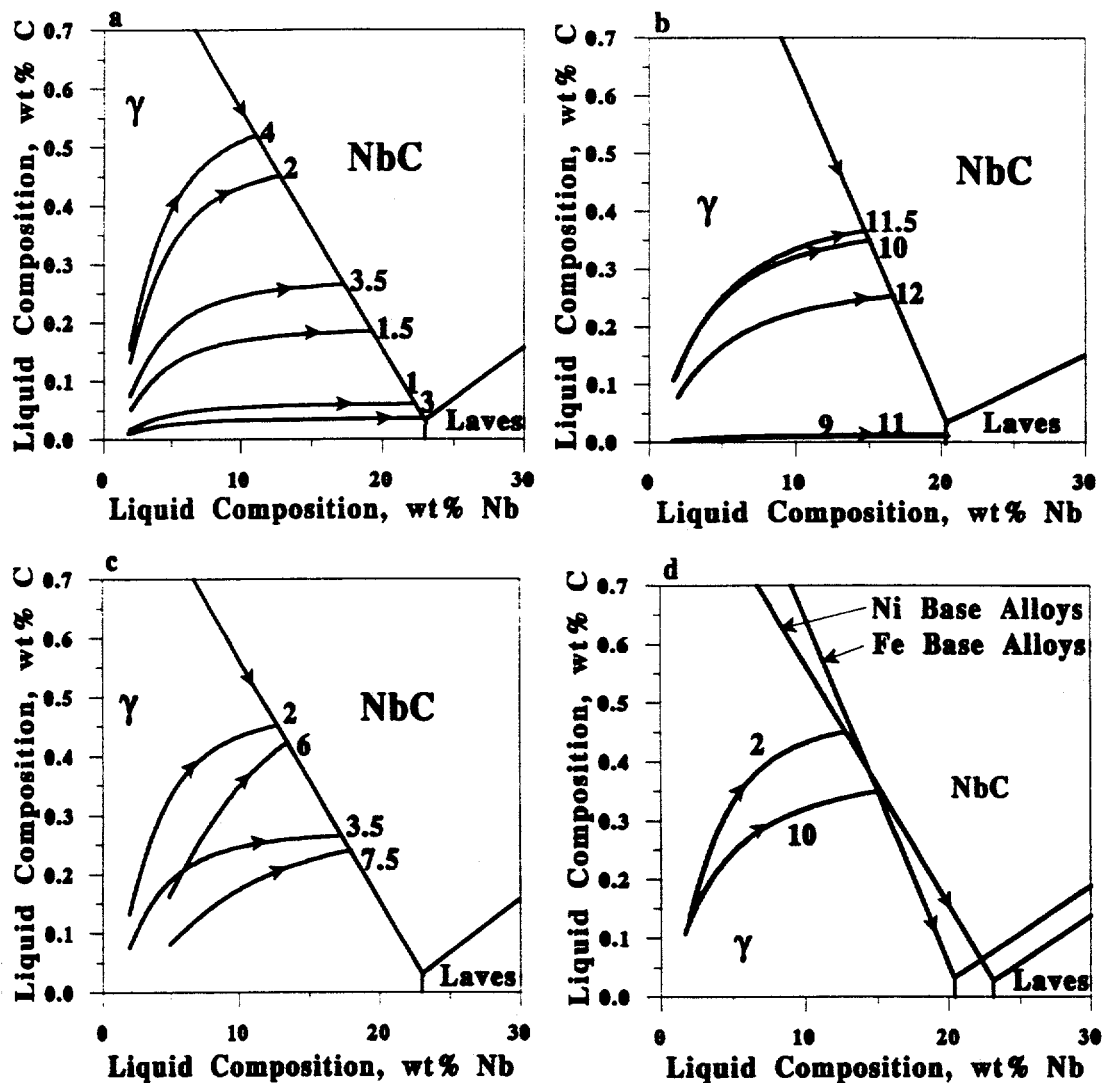


Figure 6. Calculated solidification paths.

to promote the γ /NbC eutectic type constituent, this analysis provides a quantitative rationale for the observed behavior. As the C content is increased, the intersection point occurs at higher C contents. As a result, the liquid composition must "travel" a long distance down the $L \rightarrow (\gamma + \text{NbC})$ line of two fold saturation, forming γ /NbC as it travels, before the γ /Laves constituent can possibly form. At a given C content, the Nb concentration also has a significant effect on the primary solidification path. This is revealed by comparing Alloy 2 to 6 and Alloy 3.5 to 7.5 in Figure 6c. When Nb is low, the C content in the liquid increases relatively quickly during the early stage of solidification. The rate of C build-up then decreases as the solute becomes depleted in the liquid. Since the length of the primary solidification paths for the high Nb alloys are shorter, these alloys will exhibit more liquid at the intersection point (i.e., more total eutectic-type constituent). The effect of matrix composition is presented in Figure 6d. Alloys 2 and 10 have very similar levels of solute content, but different matrix compositions. Thus, the main difference here is in the value of k_{Nb} (k_{Nb} in the Ni base Alloys = 0.46 while k_{Nb} in the Fe base Alloys = 0.25). As k is reduced, Nb segregates more aggressively to the liquid. Thus, at any given level of C in the liquid, the Fe base alloys will always possess more Nb. As a result, the Fe base alloys will have more liquid remaining after primary solidification, and the Nb content in that remaining liquid will be higher. These two effects from the reduced k_{Nb} value are the primary reasons for the higher amount of total eutectic-type constituents observed in the Fe base alloys. The accuracy of this pseudo-ternary approach for the primary $L \rightarrow \gamma$ reaction portion of solidification can

be evaluated by comparing the values of calculated and measured amounts of total eutectic-type constituent. The results are plotted in Figure 7, and there is good agreement between calculated and measured total eutectic-type constituent.

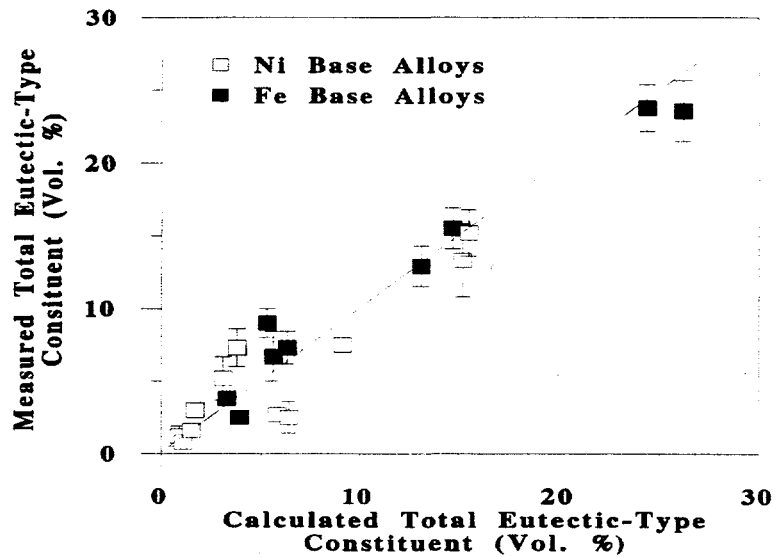


Figure 7. Comparison of calculated and measured amounts of total eutectic-type constituent.

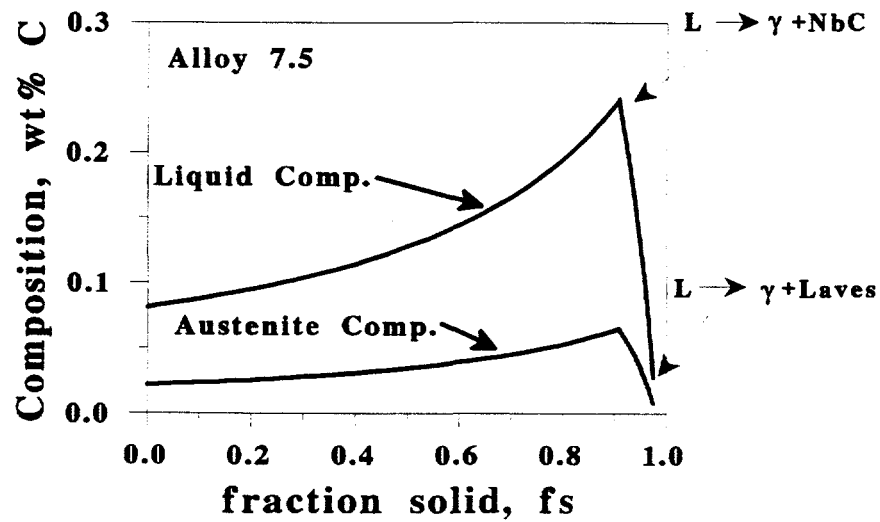


Figure 8. Calculated austenite and liquid carbon contents as a function of fraction solidified.

Eutectic-Type Solidification Figure 8 shows typical results (using Alloy 7.5 as an example) for the entire solidification process, where solute redistribution during primary solidification is calculated with eq. (3) and the eutectic-type $L \rightarrow (\gamma + \text{NbC})$ reaction is calculated with eq (7). The figure shows the variation in carbon content in the γ and liquid phases during solidification. Primary $L \rightarrow \gamma$ solidification occurs until there is 0.10 volume fraction liquid remaining with 0.24 wt% C, at which point the $L \rightarrow (\gamma + \text{NbC})$ reaction is initiated. The $L \rightarrow (\gamma + \text{NbC})$ reaction depletes the liquid of carbon until there is 0.02 volume fraction remaining liquid at 0.03 wt% C, at which point the $L \rightarrow (\gamma + \text{Laves})$ reaction is initiated. The calculations are terminated at this point, and the alloy is predicted to have 0.10 volume fraction of total eutectic (0.08 volume fraction γ/NbC plus 0.02 volume fraction γ/Laves). The measured and calculated values of the individual eutectic-type constituents for all the alloys are compared in the Figure 9. The reasonable agreement between measured and calculated values confirms that the model developed here correctly captures the major factors controlling the solidification behavior of this important class of alloys. In addition, the model is general enough to apply to other alloy

systems in which the diffusion behavior of one solute is high enough to maintain equilibrium during solidification.

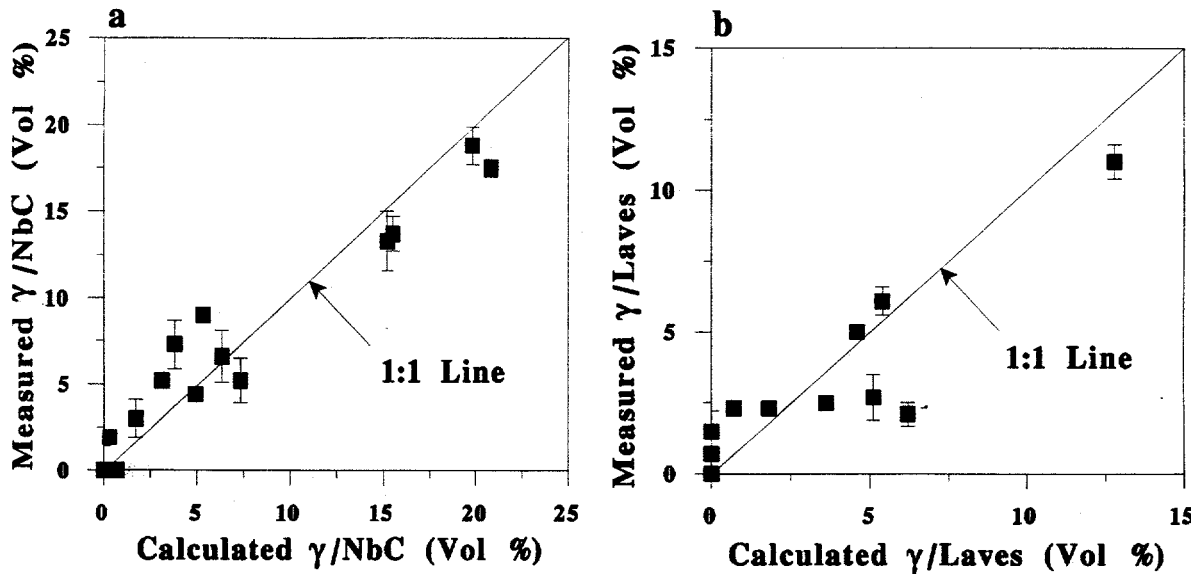


Figure 9. Comparison of measured and calculated values of the individual eutectic-type constituents .

CONCLUSIONS

Solute redistribution and microstructural evolution have been modelled for experimental alloys which simulate compositions of Ni base and Fe base superalloys containing Nb, Si, and C. The multi-component alloys were modelled as a ternary system by grouping the matrix (Fe,Ni,Cr) elements together as the "solvent" to form the γ component of the γ -Nb-C "ternary system". The variation in fraction liquid and liquid composition during the primary $L \rightarrow \gamma$ and eutectic type $L \rightarrow (\gamma + \text{NbC})$ stages of solidification were calculated for conditions of negligible Nb diffusion and infinitely fast C diffusion in the solid phase. The model results permit quantitative interpretations of composition-microstructure relations in these Nb bearing experimental alloys and should provide useful insight into comparable commercial alloy systems as well.

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Acknowledgment

Sandia is a multiprogram laboratory operated by Sandia Corporation, a Lockheed Martin Company, for the United States Department of Energy under Contract DE-AC04-94AL85000.