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Temperature dependent crystallization of amorphous Y₆₇Fe₃₃ studied by kinetic small angle neutron scattering

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Abstract

Temperature-resolved small angle neutron scattering has been used to study the nucleation, growth kinetics and crystallite morphology in the Y-Fe system. Crystallization from amorphous $Y_{67}Fe_{33}$ to the YFe₂ Laves phase via a novel 'YFe' intermediate phase has been followed to completion as a function of temperature from 180°C to 500°C. The SANS results agree well with published kinetic neutron diffraction data. Below 390°C, diffraction data suggest that SANS arises solely from the contrast between crystalline Y and the Fe-rich amorphous matrix. Between 390°C and 410°C all temperature variables are seen to form a sharp peak. This suggests that critical scattering occurs at $T_c \approx 400$ °C. This critical scattering implies full crystallization of $Y_{67}Fe_{33}$ occurs over a very narrow (~20°C) temperature range.

1. Introduction

Studying the kinetics of processes such as crystallization, phase formation and grain growth in metallic systems leads to a deeper understanding of these fundamental physical phenomena, and allows the possibility of tailoring the microstructures of materials for their particular applications. Time-resolved neutron scattering is a particularly powerful technique with which to measure any kinetic process where the neutron scattering lengths of the starting and final materials are sufficiently different to be distinguished. Rare earth-transition metal (RE-TM) alloys are frequently studied because of the interesting magnetic properties they often exhibit [1-2], and their potential technological applications (high-density permanent magnets, magnetic storage media, and giant magnetoresistive materials). The large number of crystallographic phases, observed in RE-TM systems provide the opportunity to vary magnetic exchange, anisotropy, coercivity, and conduction electron density making them ideal candidates to test theories of magnetic properties of intermetallics, such as the fundamentals of spin fluctuation, moment formation, and magnetic order.

The crystallization processes in rare earth-transition metal binary systems and the associated equilibrium phase diagrams are thought to be well established for most compounds. Phase diagrams are generally constructed using a quenching technique, where samples are heated to the desired temperature and then cooled rapidly. It is assumed that the crystallographic state existing at elevated temperature is 'frozen' into the system. However, this approach has limitations, as was demonstrated by Kilcoyne *et al* in 2001 with the discovery of a new 'YFe' intermediate phase, prepared by carefully annealing amorphous $Y_{67}Fe_{33}$ ribbons whilst monitoring the crystallization processes as a function of time with neutron diffraction [3]. From the results of the study by Kilcoyne *et al*, the crystallization of amorphous $Y_{67}Fe_{33}$ can be fully described. Firstly partial crystallization occurs at 300°C when elemental Y begins to form. Crystalline Y exists within an iron-rich amorphous matrix, until 390°C when the whole sample crystallizes. At 390°C the intermediate 'YFe' phase starts to form, coexisting

with elemental Y. This phase only exists over a narrow temperature range (approximately 60°C). By 450°C, no Bragg peaks arising from the intermediate 'YFe' phase are observed. Lastly, at 450°C the YFe₂ Laves phase forms. This phase coexists with pure Y until completion of the reaction at 490°C.

Observation of the intermediate phase indicated that this was a previously unreported phase which clearly required further investigation. Kilcoyne *et al* managed to stabilize the phase at room temperature by rapidly cooling ribbons from 375°C. Diffraction patterns were collected regularly during cooling, to confirm that the phases present at elevated temperature were unaltered at room temperature [3]. The crystallographic parameters of the 'YFe' intermediate phase were determined using pattern matching analysis with the FullProf program [4] and are given in Table 1.

In this paper we present the results of a small angle neutron scattering (SANS) study of nucleation and grain growth during crystallization in the Y-Fe system as a function of temperature. The results are linked to the diffraction data obtained on the D2B diffractometer at ILL, Grenoble and presented in [3].

	Phase 1 Y	Phase 2 'YFe'
Space Group	P6 ₃ /mmc	P6 ₃ /mmc
а	3.628(3)Å	12.90(1)Å
с	5.739(5)Å	1.171(1)Å
c/a	1.58	0.91
γ	120°	120°
R _(Bragg)	2.76	1.60

Table 1 Refinement parameters from D2B refinement and pattern matching (taken from [3]).

2. Experimental Procedure

Polycrystalline $Y_{67}Fe_{33}$ ingots were produced using an argon arc furnace. High purity elements (99.9% purity Y, and 99.99% purity Fe) were melted together in an argon atmosphere from which any gaseous impurities were removed by first melting a titanium "getter". To ensure homogeneity, ingots were melted and turned several times and their masses were limited to 10g. The ingots were then cut into small pieces (0.7 to 1.0g) in preparation for melt spinning. Once melted in a quartz tube, a burst of Ar gas forced the molten sample onto the rapidly rotating copper wheel, cooling it at a rate of ~10⁶ Ks⁻¹. Several grams of amorphous ribbon, ~20 µm thick and several cm long, were produced using this method.

Approximately 2g of melt spun $Y_{67}Fe_{33}$ ribbons were mounted in a flat, circular vanadium cell. The sample cell had a radius of 10mm and a thickness of 2mm. The cell was mounted in a standard vanadium-element vacuum furnace on the SANS instrument D22 at the Institute Laue Langevin, France. The set up used allowed a Q range of 0.0018\AA^{-1} to 0.015\AA^{-1} to be explored. The ribbons were first heated to 150° C for one hour to establish thermal equilibrium, and then heated from 180° C to 500° C (i.e. the temperature range in which all three stages of crystallization can be observed) at 1°C min⁻¹ whilst SANS patterns were collected every minute. In order to compensate the measured intensity to account for the efficiency of the detector and for the detector surface being planar rather than spherical, a standard sample must be measured. In our experiment where relative cross-sections, rather than absolute values are required Perspex, which has a uniform cross-section in the Q range of interest, is a suitable standard.

3. Results and Discussion

Figure 1 shows the variation of I(Q) with Q for several temperatures between 180°C and 500°C plotted on an x-y logarithmic scale. It can be seen from this figure that the Q and temperature dependence of the intensity is not straightforward. In the high Q range, (Q>0.01Å⁻¹) I(Q) begins to increase at around 330°C. Between 330°C and

380°C, I(*Q*) increases in the high *Q* region, whereas at very low *Q*, (Q<0.004Å⁻¹) it remains unchanged. Between 380°C and 400°C I(*Q*) increases in both the low *Q* (Q<0.006Å⁻¹) and high *Q* regions. Above 400°C, I(*Q*) continues to increase at low *Q* but starts decreasing at high *Q*.

Figure 2 shows the variation of I(Q) with temperature at several Q values. This plot also illustrates the marked difference in the behaviour of I(Q) at low Q and at high Q as a function of temperature. At high Q, I(Q) starts to increase at around 300°C and continues to increase until ~410°C where it peaks. It then decreases between 410°C and 430°C. From 430°C to 500°C I(Q) stays approximately constant in the high Q region. At low Q, I(Q) stays constant until ~390°C. From 390°C to 420°C it increases sharply, and from 420°C to 500°C it continues to increase but at a lower rate. At intermediate Q, $0.006Å^{-1} < Q < 0.01Å^{-1}$, the temperature dependence of I(Q) shows behaviour from both the high Q and low Q regimes.



Figure 1 log I(Q) vs log Q at various temperatures during heating at 1°C min⁻¹ of amorphous Y₆₇Fe₃₃. Errors lie within the data points. The y-axis has been logarithmically scaled to fit all the curves on the same axes.



Figure 2 Temperature dependence of the SANS intensity at various values of Q, heating amorphous Y_{67} Fe₃₃ at 1°C min⁻¹. Errors lie within the data points.

Assuming a monodispersed system of particle sizes, Guinier plots of log I(Q) versus Q^2 for spherical particles were used to estimate the mean particle size [5]. The Guinier approximation is only valid in a limited low Qrange up to $QR_G \sim 1.2$ where R_G is the radius of gyration of the spherical particles. In this work, Guinier plots do not give the expected straight line suggesting that a range of particle sizes exists in the Y-Fe system. This is not unexpected since from kinetic neutron diffraction experiments it is known that the crystallization of amorphous $Y_{67}Fe_{33}$ follows a complex multi-stage, multi-phase process. One or more crystalline phases or an amorphous phase coexist at any given temperature and these different phases will have different correlation lengths. Linearity is seen however, in plots of 1/I(Q) versus Q^2 and $1/I(Q)^{1/2}$ versus Q^2 which suggests that a Lorentzian or Lorentzian squared distribution would be a more appropriate model for the data. Although strictly only valid in the Guinier region, the generalized Lorentzian function given in Equation 1 appears to provide a good description of the data for the entire Q range.

$$I(Q) = \frac{A(\kappa^2)^z}{\left(\kappa^2 + Q^2\right)^z} \tag{1}$$

where A is the scattering amplitude, which is related to the number of particles and the contrast in scattering length density in the material. The range parameter $1/\kappa$; is the average length-scale of fluctuations in scattering length density, and gives the particle size or the distance between particles. *z* is the power of the Lorentzian term. A power of Lorentzian expression, is observed when scattering arises from a Maxwellian distribution of geometrically similar particles [6]. Equation 1 was used by Boardman *et al* to fit similar structural SANS data collected on D11 (ILL, Grenoble) during the crystallization of amorphous Bi₂Sr₂CaCu₂O_x [7].

Equation 1 fits the SANS data collected during the crystallization of amorphous $Y_{67}Fe_{33}$ extremely well across the whole *Q* range and temperature range. The fits to the data collected at several temperatures between 300°C (the start of Y-crystallization) and 500°C, are shown in Figure 3, while Figure 4 shows the temperature dependence of the fit parameters *A*, $1/\kappa$, and *z*.



Figure 3 SANS data fitted to the general Lorentzian function given in (1).

In can be seen in Figure 4 that for T<300°C the scattering is effectively squared Lorentzian in form (z~1.9), indicating that a broad distribution of structural inhomogeneities exist in the melt-spun ribbons [8]. The range parameter suggests that these inhomogeneities are ~170Å in size. At 300°C, the power of the Lorentzian term, z, starts decreasing non-linearly reaching ~1.3 at 390°C, and then drops rapidly between 390°C and 410°C almost reaching unity at 410°C. A and 1/ κ do not alter between 300°C and 390°C. Between 390°C and 410°C however, while z decreases rapidly, the scattering amplitude A, and the range parameter 1/ κ both increase dramatically.

 $1/\kappa$ reaches a peak of 700Å at 410°C, which is still within the resolution of D22 for our experimental setup. Between 410°C and 450°C *z* begins to increase and returns to its initial value of ≈1.9 by 450°C. Over the same temperature range, *A* decreases to a local minimum and $1/\kappa$ drops rapidly reaching a minimum of 300Å at 450°C. Above 450°C, *z* remains approximately constant at 1.9 while *A* and $1/\kappa$ both start to increase again, with $1/\kappa$ reaching a value of 410Å when the experiment was terminated at 500°C.



Figure 4 Generalized Lorentzian parameters for fits to SANS data. The errors lie within the data points.

A comparison of temperature dependent SANS data with temperature dependent neutron diffraction data (from Kilcoyne *et al* [3]) during crystallization of amorphous $Y_{67}Fe_{33}$ is given below. Features are seen in both the diffraction data and SANS data at 300°C, between 390°C and 450°C, and above 450°C.

<u>At 300°C</u>

From the diffraction data, it is known that the sample remains amorphous until 300°C when Y grains start to precipitate out. Using the Scherrer equation on the FWHM of Bragg peaks arising from the Y phase, Kilcoyne *et al* found that Y grains first appear at 300°C and are 170Å in size. The same value of 170Å is found for the range parameter, $1/\kappa$; at the start of heating on D22, which suggests that the Y₆₇Fe₃₃ sample used on D22 was not completely amorphous and that Y grains were already present in the ribbons. Unfortunately it was not possible to check the amorphicity of ribbons by diffraction before starting the experiment, however previous studies of Y₆₇Fe₃₃ have shown that small quantities of crystalline Y can be present in melt spun ribbons [9]. The value of the power of a Lorentzian parameter, *z*, was found to be ~1.9 at 300°C. Dierker *et al* [8] stated that Lorentzian squared scattering (*z*=2) can arise from a crystalline phase existing in an amorphous matrix with sharp interfaces between them, agreeing with the argument that nanocrystalline Y grains are present in the amorphous matrix of the melt spun Y₆₇Fe₃₃ ribbons.

Between 390°C and 450°C

All three variables in the SANS fits (A, $1/\kappa$ and z) peak between 390°C and 410°C. This suggests that critical scattering occurs in this temperature range and suggests that there is a critical point that defines the transition between one state and another [10]. Ornstein and Zernike demonstrated that during critical scattering, the scattering distribution is Lorentzian in shape [11, 12]. In addition, Birgeneau *et al* [13] showed that the Fourier transform of the correlation function C(Q) could be written as:

$$C(Q) \propto \frac{\kappa^p}{\left(\kappa^2 + Q^2\right)} \tag{2}$$

where p is the critical exponent and $1/\kappa$ is the correlation length (or range parameter). For the case of z=1, Equations 1 and 2 are equivalent with a critical exponent of p=2.

Generally, there are two possible causes for the critical scattering seen in structural SANS data. The first is that a percolation threshold has been reached at a critical temperature, T_c . The second is that critical scattering is occurring near T_c in a continuous, higher order phase transition. The neutron diffraction data collected by Kilcoyne *et al* [3] show that at ~390°C there is a large drop in the background signal as the whole sample crystallizes. It is clear from this that the Y₆₇Fe₃₃ sample fully crystallizes extremely rapidly at ~390°C whilst at the same temperature the new 'YFe' phase appears. This points to the critical scattering seen in the SANS data being caused by the sudden crystallization of the whole sample – i.e. a percolation threshold. The rapid increase in $1/\kappa$ as the temperature approaches T_c therefore suggests that there is an increase in the average grain size as the crystallites grow and impinge on one another prior to the effectively infinite crystalline cluster being formed at the percolation threshold. A value of $T_c \approx 400$ °C can be extracted for the critical temperature of the full crystallization of amorphous $Y_{67}Fe_{33}$.

Above 450°C

From diffraction data, 450°C is the temperature at which the new 'YFe' phase transforms into YFe₂. Also above 450°C is when the Y grains grow most rapidly. SANS above T_c at T>450°C can therefore be explained by the phase transformation from the new 'YFe' phase to YFe₂ as the atoms reorder and the Y and YFe₂ grains grow.

4. Conclusions

Small angle neutron scattering has been used to study the temperature dependent crystallization of amorphous $Y_{67}Fe_{33}$. The results can be described extremely well by a general Lorentzian function, and the data are in good

agreement with the neutron diffraction study by Kilcoyne *et al.* At low temperatures, diffraction data suggest that SANS arises solely from the contrast between crystalline Y and the Fe-rich amorphous matrix. At T~400°C critical scattering is observed corresponding to the full crystallization of the sample. Above 410°C, trends in *A*, $1/\kappa$ and *z* can be followed as a function of temperature, but having a complex multi-phase system makes it is difficult to isolate the individual range parameters, $1/\kappa$, for each crystallographic phase (Y, 'YFe' or YFe₂).

Until now the lowest reported Fe concentration in a Y-Fe compound was $\sim 67\%$, (YFe₂) but, through the formation of this phase, this has now been reduced to 50%. It is hoped that it will be possible to isolate this phase, and through understanding how the magnetic properties relate to the crystallographic properties and growth morphologies we will be able to tailor micro- and nano-structured materials with interesting technological applications.

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