Crystallization Process and Microstructural Evolution of Melt Spun Al-RE-Ni-(Cu) Ribbons

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Abstract: The crystallization process, both at the initial and subsequent stages, of amorphous Al88RE4Ni8 alloys (RE = Y, Sm and Ce) has been studied. Additionally, the consequences of adding 1 at.% Cu replacing Ni or Al were studied. The stability of the amorphous structure in melt spun ribbons was thermally studied by differential scanning calorimetry, with Ce alloys being the most stable. The effect of Cu to reduce the nanocrystal size during primary crystallization was analyzed by transmission electron microscopy. This latter technique and x-ray diffraction showed the formation of intermetallic phases at higher temperatures. A clear difference was observed for the Ce alloy, with a simpler sequence involving the presence of Al3Ni and Al11Ce3. However, for the Y and Sm alloys, a more complex evolution involving metastable ternary phases before Al19RE5Ni3 appears, takes place. The shape of the intermetallics changes from equiaxial in the Ce alloys to elongate for Y and Sm, with longer particles for Sm and, in general, when Cu is added to the alloy.

Keywords: amorphous materials; melt spinning; Al-RE-Ni alloys; crystallization

1. Introduction

The disordered or amorphous structure typical of the liquid state can be maintained in solid metals by heating above their melting temperature and very quick cooling to room temperature [1,2]. Only particular compositions are appropriate to form metallic glasses, and depending on the composition, the necessary cooling rate varies. The importance of this process resides in the outstanding attained properties of different character such as mechanical, electrical, magnetic, or chemical, all of them of great engineering importance [3,4].

Regarding compositions, aluminum mixtures with at least other two chemical elements (although it is also possible with only one, i.e., [5,6]) can reach the amorphous state by quick cooling, attaining high strengths with respect to conventional aluminum alloys [7]. Properties such as elastic strain [8], or the possibility of superplastic forming in a supercooled liquid region [9] are remarkable.

Initially, Al amorphous alloys with a composition Al-Fe-Si [10,11] showed brittleness problems that were subsequently solved by changing for instance to Al-Ni-Zr compositions [12]. Later, the development of compositions Al-RE (rare earth: La, Y, Ce)-TM (transition metal: Fe, Co, Ni) [6,13–16] allowed very high strengths (up to 1100 MPa) to be reached. This strength can be increased for some alloys by allowing the formation of up to a 50 vol.% of α-Al nanocrystals [17,18] at the expense of the amorphous phase. For nanocrystallized Al-RE-Ni alloys, ultimate tensile strengths can increase up to 1560 MPa [19,20], three times the value of crystalline alloys, even showing improved ductility and
toughness [21]. The nanocrystal size, number, and spatial distribution are key parameters in the obtained properties. These parameters are better controlled by annealing from an amorphous structure obtained after quenching [17].

Intermetallics appearing after heating are unfavorable for the properties of the alloys [22]. Understanding this process, which should be initially avoided to preserve the best properties, is of great importance. One of the important aims of working with amorphous alloys is to obtain thick parts [2], and the formation of intermetallics is a concern in this process. Many bulk metallic glasses (BMG) have been prepared, with lots of them still to be discovered [23]. The formation of BMG can be theoretically predicted according to the physical properties of the amorphous and crystalline phases, or on known rules such as the confusion principle. However, for Al-based alloys, the glass forming ability (GFA) does not adequately follow the expected criteria, and the production of Al-based BMG is relatively difficult and limited to low Al contents [24]. Under these circumstances, Al-based alloys are bad candidates for BMG processing; and the amorphous ribbons require further processing by powder metallurgy methods. For instance, spark plasma sintering, one of the multiple electric field assisted sintering techniques, has been used to consolidate Al-Y-Ni amorphous powders [25,26].

Al-RE-Ni alloys were studied in this work. The study of the crystallization kinetics and competition of the crystalline phases [27-30], the expected GFA [31-33], and the theoretical modeling of the adequate solute range for amorphous forming [34-36] shows that ternary alloys with a solute content around 10-15 at.% are good candidates to attain amorphous structures. Thus, the compositions Al<sub>x</sub>-RE<sub>4</sub>-Ni<sub>8</sub> were selected (with RE = Y, Sm, and Ce, three well-known candidates to reach the amorphous state). The influence of small amounts of alloying elements in the primary crystallization of Al-RE-Ni systems has been previously studied [37-44]. In this work, it will be extended to include the subsequent appearance of intermetallic phases, which are of great importance during the consolidation process of amorphous ribbons.

2. Materials and Methods

The alloys Al<sub>x</sub>-RE<sub>4</sub>-Ni<sub>8</sub> (RE = Y, Sm, and Ce) were selected in this work. Moreover, Al or Ni was replaced by just 1 at.% Cu, resulting in the alloys Al<sub>x</sub>-RE<sub>4</sub>-Ni<sub>8</sub>-Cu<sub>1</sub> or Al<sub>x</sub>-RE<sub>4</sub>-Ni<sub>8</sub>-Cu<sub>4</sub>. Pure metals (Al 99.9%, Ni 99.95%, Y 95%, Sm 99.9%, Ce 99.9%, and Cu 99.9%) were arc melted in vacuum to prepare the master alloys Al<sub>x</sub>-Cu<sub>10</sub> and Al<sub>x</sub>-RE<sub>4</sub>. Pure metals and master alloys in the adequate proportion were repeatedly remelted to obtain the final alloys with an adequate homogeneity. The final materials were obtained in the form of ribbons after heating the alloy at 1000 °C in a melt spinning chamber filled at a 200 mbar He atmosphere. The liquid was ejected through a round nozzle 0.85 mm in diameter on a Cu wheel rotating to reach 40 m/s on its surface. A distance nozzle-wheel of 1.5 mm resulted in ribbons with a thickness of 25-30 µm and width of 1-2 mm.

The amorphous character of the melt-spun ribbons was examined by x-ray diffraction (XRD, model PW 1729, Philips, Amsterdam, Netherlands, with Cu-Kα radiation for 2θ = 20-80°). Transmission electron microscopy (TEM, model CM20, Philips, Amsterdam, Netherlands, working at 200 KV) was used to characterize heat treated ribbons, after electropolishing at ~30 °C in 10% perchloric acid, 90% ethanol solution. In general, the Al nanocrystals were especially attacked with this technique, being necessary to ion mill the heat treated ribbons after cooling at ~30 °C.

The thermal stability of the ribbons was studied by differential scanning calorimetry (DSC, model 2010, TA Instruments, New Castle, DE, USA) heating at 20 °C/min, on times combined with final isothermal treatments, under an Ar atmosphere. Cooling was always carried out by switching off the equipment, without particular control of the cooling rate.

3. Results and Discussion

3.1. Melt-Spun Ribbons

XRD patterns of as-spun materials are shown in Figure 1. A broad peak at 2θ ~38° can be observed, being representative of having reached an amorphous structure in Al alloys. Regarding the
Y alloy, a small shoulder appeared for $2\theta \approx 44^\circ$, because of the presence of some local order [41,45,46], corresponding to the initial stages of disappearance of the amorphous structure, while still being maintained. According to the confusion principle [47], the low atomic radius difference between Al and Y when compared to Al-Sm or Al-Ce [48] reduces the GFA. Al-RE-Ni alloys showed a strong linear dependence for the GFA on the size of RE elements, with bigger atoms facilitating the glass formation [49]. Nevertheless, the obtained structures can be considered amorphous from the XRD analysis.

![X-ray diffraction (XRD) patterns of Al-RE-Ni-(Cu) melt-spun ribbons.](image)

**Figure 1.** X-ray diffraction (XRD) patterns of Al-RE-Ni-(Cu) melt-spun ribbons.

XRD also showed that, in general terms, patterns corresponding to Al$_{87}$-RE$_4$-Ni$_9$-Cu$_1$ seemed to be flatter, and those of Al$_{88}$-RE$_4$-Ni$_7$-Cu$_1$ tended to contain sharper peaks. On the other hand, the diffuse peak position shifted to lower angles for the Ce alloys. According to the Ehrenfest relation [50], this position was lower for a higher value of near-neighbor distance, occurring on the Ce-containing alloy, with a higher atomic radius than Y and Sm.

### 3.2. Crystallization Process

DSC results for the as-spun ribbons are shown in Figure 2. The general behavior is a crystallization process occurring in three stages, although a small fourth stage can also appear at temperatures higher than 400 °C, as shown for the Cu-free Sm-containing alloy. Crystallization can therefore be quite dependent on composition. Even the composition [51] and DSC heating rate [5,42,45,52–54] can determine the transformation temperatures, and limit the capacity to detect them when taking place very close. Thus, Y-containing alloys with a similar range of compositions as those studied in this work are usually found to crystallize from the amorphous structure in three stages [34,41,45,55–59], but four stages have also been reported [27,48,58,60]. Likewise, synchrotron XRD revealed the presence of two stages in Al$_{87}$-Y$_4$-Ni$_9$ and Al$_{86}$-Y$_4$-Ni$_{10}$, but three stages for Al$_{83}$-Y$_4$-Ni$_{13}$ [29]. For Sm-containing alloys, the first three DSC reactions have been reported [51,54,61–64], but a fourth reaction has also been found for similar alloys [53,61,65–68], and up to five reactions have been identified [69]. Regarding Ce-containing alloys, the general behavior depends on the composition,
but three reactions have generally been reported for similar compositions to the one now studied [49].

![Differential scanning calorimetry (DSC) traces at a heating rate of 20 °C/min of Al-RE-Ni-(Cu) melt-spun ribbons.](image)

**Figure 2.** Differential scanning calorimetry (DSC) traces at a heating rate of 20 °C/min of Al-RE-Ni-(Cu) melt-spun ribbons.

Comparing the three RE elements, the first reaction was quite similar for the three RE-containing alloys, only at some higher temperatures for Ce, and some starting earlier for the Y-containing alloys. Regarding the second and third reactions, the Y and Sm-containing alloys showed a very similar behavior, although the third reaction seemed to be of greater importance in Y alloys. Ce-containing alloys, however, showed a second reaction much smaller with regard to the third one, and even nearer to each other when compared with the two other cases. These different crystallization stages will be analyzed next.

### 3.3. Primary Crystallization

The crystallization process starts for all of the alloys with a broad exothermic reaction. The onset temperatures were 150 °C, 165 °C, and 180 °C for the Cu-free Y, Sm, and Ce-containing alloys, respectively. According to Figure 1, the Y alloys seemed to reach a less amorphous state, and Ce patterns appeared flatter and therefore more amorphous, which is now reflected in the ease in which to start crystallization. DSC showed that increasing the total amount of solute when 1 at.% Cu replaces Al resulted in slightly more stable alloys. In contrast, when Ni was replaced, therefore maintaining the total amount of solute, and with the atomic radius of Cu being more similar to the Al than to the Ni one, a reduction of the amorphous structure stability was found.

XRD and TEM results from ribbons heated to the end of the first reaction at 250 °C showed the formation of α-Al nanocrystals. Figure 3 shows the results for the Sm-containing alloys, with similar results for the Y and Ce alloys.
Figure 3. Study of the first DSC reaction of Sm alloys after heating at 20 °C/min to 250 °C and immediate cooling: (a) XRD patterns of ribbons, with peaks corresponding to α-Al nanocrystals, and (b) transmission electron microscopy (TEM) dark field images and diffraction patterns (DP).

Nanocrystal size measurements of the ribbons heated beyond the first DSC reaction (i.e., at 250 °C) were carried out on TEM micrographs. The similar area under DSC curves ensured a similar amount of nanocrystals formed. Results obtained by line intercept showed that the presence of Cu acts to diminish the nanocrystal size. The crystals’ size mean value was thus reduced to about 9 nm when Cu was present in Sm-containing alloys, with 10.7 nm for the Cu-free alloy, and in a similar way for the Y and Ce alloys. One of the possible reasons for this is the reduction of the nucleation barrier, as reported for Al-Y-Ni-Cu [38] and Al-Ce-Ni-Cu alloys [70]. Nevertheless, for these alloys, Cu has also been found to be uniformly distributed [61,71], besides causing composition variations [70,72]. In Al-Sm-Ni alloys, heterogeneities that can act as nucleation sites have been reported after the addition of Cu [73,74]. In summary, there is not a clear consensus for the actual reasons for this reduction.

3.4. High Temperature Crystallization Stages

The second DSC crystallization reaction for the Y and Sm-containing alloys appeared with an onset at approximately 315 °C and at 335 °C for the Ce alloy (see Figure 2). This reaction ends only about 35 °C later for the Y and Sm alloys, and 20 °C for the Ce-containing alloy, and before it has totally finished, the third reaction starts (this being more pronounced for the Ce alloy).

A higher separation between the second and third reactions, allowing a better study of the processes taking place, can be attained by heating the ribbons up to 5 °C before the onset of the second DSC reaction, then maintaining the temperature in an isothermal heat treatment. Figure 4 shows the DSC traces thus obtained.
DSC traces at 20 °C/min in Figure 2 showed an asymmetry at the beginning of the second reaction for the Y and Sm alloys, not being so clear for the Ce-containing alloy. This peculiarity of the second reaction is separated in different reactions (we will call them subreactions 1 and 2) when holding the temperature in Figure 4, always more clearly for the Cu-free alloys. This different behavior for the Ce-containing alloys could be caused by a different crystallization process, which is discussed next. The third reaction remains during isothermal heating as only one, starting up to 10 min after the end of the second one. The start of this third reaction is quicker for a lower separation with respect to the second one in continuous heating, something taking place very clearly for the Ce alloys, and in general for the Cu-free alloys.

The second reaction is usually a source of discrepancies in previous studies, mainly for the Y and Sm alloys. The reason is that even after heating the ribbons to make this reaction take place, identifying the phases present in the alloys is challenging. XRD patterns of ribbons isothermally treated to the end of the two subreactions of the second crystallization stage (Figure 5 shows the

**Figure 4.** DSC traces (heating rate of 20 °C/min up to 5 °C before the onset of the second reaction and then maintaining the temperature) of the Al-RE-Ni-(Cu) alloys (vertical lines mark the heating regime change.)
results for the Y alloys, being similar for Sm and Ce alloys) seem to show that in both cases, only α-Al nanocrystals were found. Likewise, TEM diffraction showed no spots from new phases once the second DSC reaction finished.

Undoubtedly, something is happening in the second crystallization stage. The only difference with previous XRD patterns (i.e., after the primary crystallization) was an asymmetry at the left of the main Al peak, which could be related to the formation of new phases probably just nucleating, but their identification is not possible. A rough estimation can be done after (i) taking into account that the intermetallic formation requires an activation energy 25% higher than α-Al nanocrystals [48], (ii) that the amount of intermetallics and primary nanocrystals are similar, and (iii) considering the areas of the reactions under the DSC curves. Thinking of an α-Al nuclei about 10 nm in diameter, the equivalent nucleation and growth for the intermetallics would lead during the first and second subreactions of this second stage to particles of about 6 nm and 5 nm respectively, in the case of an homogeneous process. However, this same volume of new particles led to layers of 0.4 nm and 0.25 nm forming around the existing α-Al nanocrystals (values in the same order of magnitude result by computing the areas in the DSC curves of the Sm alloys). These very thin and not easily detectable layers could be the explanation for the almost similar results, therefore a heterogeneous nucleation surrounding previous α-Al nanocrystals can be proposed.

As mentioned before, no clear results have been reported regarding the second and subsequent crystallization stages of these alloys. For Y alloys, Al₃Ni, Al₁₉Y₃Ni₅, and unidentified phases have been indicated to appear [43,45,55–57,75,76]. The Al₁₆YN₁₃ phase is, in more recent years, not referred, and Al₁₃Y₃Ni₅ and Al₁₅Y₅N₁₃ have been proposed [27,29,77–79], the former on times considered a metastable phase, and both of them with a very similar structure to the well-known Al₃YN phase [80,81]. This metastable character is not discarded according to the Al-Y-Ni intermetallic enthalpies of formation [31,78,82], lower for Al₁₃Y₃Ni₅ than for Al₁₅Y₅N₁₃. However, Al₁₅Y₅N₁₃ [29] and also the Al₁₃YN₁₃ [27,29] have been proposed as stable phases of this system.
Regarding Sm alloys, the second stage is related to the crystallization of Al₃Sm (cubic + hexagonal) [51,53], Al₁₁Sm₃ [65], or Al₃Ni and Al₃Sm [69]. Unknown metastable intermetallics have also been suggested [51,53,54,64,68]. After the third reaction, unknown and Al₁₁Sm phases [51,53], only binary phases such as Al₃Sm and Al₃Ni [67] (together to Al₁₁Sm and Al₁₁Sm₃ [69]), or binary and ternary phases together such as Al₁₁Sm₃ and Al₃SmNi₃ [54,62,64] are reported. Additionally, the continuity of orthorhombic Al₁₁Sm₃ and the appearance of tetragonal Al₁₁Sm₃ and Al₃SmNi₃, which then transform from orthorhombic Al₁₁Sm₃ to orthorhombic Al₁₁Sm, has been proposed [65].

Ce alloys have been reported to lead to unknown metastable phases and (AlNi)₁₁La₃-like phases [48], or Al₃Ni, Al₃Ce, and unidentified phases [83] as well as to Al₃Ni, Al₃Ce, Al₃Ce, AlCeNi, and unidentified phases [84].

According to these studies, it can be concluded that both Al₃Ni and a certain ternary phase are causing the two subreactions in the second DSC reaction of the Y and Sm-containing alloys. On the other hand, for Ce-containing alloys, Al₃Ni and probably a binary Al-Ce phase should appear.

Back to the present study, further heating the second DSC reaction, intermetallics clearly appeared for all the alloys. Figure 6 shows the XRD patterns after heating at 20 °C/min to the peak of the third DSC reaction (at about 355–365 °C), to 375 °C (end of the reaction) held for 30 min, to 450 °C and to 600 °C.
Although some differences were previously observed in the DSC traces for the Ce alloys, the first big difference between the different RE elements studied was now found. The final microstructure, according to the XRD results, was quite similar for the Y and Sm-containing alloys, but different for the Ce alloys.

Focusing first on the Y and Sm alloys, the Cu-free Y alloy showed the same pattern for the different heat treatments, with peaks identified as Al, Al$_3$Ni, and Al$_{19}$Y$_3$Ni$_5$. Nevertheless, the quite small amount of Al$_3$Ni, and the coincidence of the XRD peaks with those of the other phases, makes its identification difficult. When Cu is added to the Y alloy, the high-temperature microstructure is the same, but it seems to be attained through the evolution of a metastable phase, which tends to transform to the Al$_{19}$Y$_3$Ni$_5$ phase. It is only possible to detect the metastable phase when the third DSC reaction is not completed (i.e., for the treatment to the peak temperature of the third reaction). According to this, the metastable phase tends to grow, and then transforms to the stable phase during the third DSC reaction, probably after appearing during the second DSC reaction. Whether the metastable phase also appears in the Cu-free alloy cannot confirmed from the XRD results because the proximity of the second and third DSC reactions does not make it possible to detect it before the Al$_{19}$Y$_3$Ni$_5$ phase has been totally formed, even when heating to the peak temperature of the third reaction.

A similar evolution was found for the Sm-containing alloys. The existence of the Al$_{19}$Sm$_3$Ni$_5$ phase has not been established in previous studies, but according to the peaks’ positions, and through a comparison with the Y alloys, this can be proposed as the stable phase appearing in the Sm alloys. Based on the similarity of the Al$_{19}$Sm$_3$Ni$_5$ and Al$_{19}$Y$_3$Ni$_5$ lattice parameters (1.5939 nm, 0.40967 nm, 1.8320 nm, and 113.09° for the former [85], and 1.5836 nm, 0.4068 nm, 1.8311 nm, and 112.97° for the latter [77]), this could be an acceptable assumption. Conversely, the metastable phase prevails in the Sm alloys for tougher treatments (with a similar evolution in the Cu-free alloy to that of the Cu-

\[ \text{Figure 6. XRD patterns of (a) Y, (b) Sm, and (c) Ce-containing alloys after heat treating to the peak of the third DSC reaction (355–365 °C) to 375 °C (end of the reaction) held for 30 min, to 450 °C and to 600 °C.} \]
containing Y alloy), with the presence of the metastable phase being detected, even when heat treating beyond the third DSC reaction. The metastable phase partially transforms to Al$_{19}$Sm$_3$Ni$_5$ after heating to 450 °C, and totally for 600 °C. Therefore, in Sm-containing alloys, the transformation from the metastable to the Al$_{19}$Sm$_3$Ni$_5$ phase must take place after the third peak, which now appears as a separate reaction. This reaction was found in Figure 2 at about 400 °C for the Cu-free alloy, and must be taking place at higher temperatures for the Cu-containing alloys, although it is not clearly detected in DSC traces.

Figure 7 shows the XRD patterns simulations (with PowderCell software [86]) and the experimental traces for Y and Sm-containing Al$_{87}$-RE$_4$-Ni$_8$-Cu$_1$ alloys after heating to 600 °C. For the Y alloy, Al, Al$_{19}$Y$_3$Ni$_5$, and Al$_3$Ni in proportions 63/35/2 were simulated (these are just rough qualitative values to replicate the experimental pattern). The crystalline data for Al$_{19}$Sm$_3$Ni$_5$ have been taken in accordance with the Al$_{19}$Y$_3$Ni$_5$ phase (space group Cmcm, Pearson symbol oC108 and lattice parameters 0.403 nm, 1.598 nm, and 2.689 nm [81]), although small variations of the lattice parameters to 0.410 nm, 1.598 nm, and 2.689 nm have been considered for a better reproduction of the experimental pattern (with Al, Al$_{19}$Sm$_3$Ni$_5$, and Al$_3$Ni in proportions 58/40/2).

![Figure 7](image_url)

**Figure 7.** XRD patterns of the Al$_{87}$-Y$_4$-Ni$_8$-Cu$_1$ and Al$_{87}$-Sm$_4$-Ni$_8$-Cu$_1$ alloys, heat treated to 600 °C. The considered phases summed to give the simulated patterns were Al, Al$_{19}$RE$_4$Ni$_8$, and Al$_3$Ni in proportions 63/35/2 and 58/40/2 for the Y and Sm-containing alloys, respectively.

According to the results in Figure 7, the similarities between the Al$_{19}$Y$_3$Ni$_5$ and Al$_{19}$Sm$_3$Ni$_5$ phases are possible. On the other hand, it must be noted that the simulated pattern could reproduce the experimental one, even without considering the presence of Al$_3$Ni. It can be concluded that the presence of Al$_3$Ni, which seems quite certain according to the literature, decreases with heating, therefore probably contributing to the formation of the Al$_{19}$RE$_4$Ni$_8$ phases.

Taking into account the results of the Y and Sm alloys, a plausible description of the process starts with the nucleation of both Al$_3$Ni and the metastable phase during the second DSC reaction
(the direct appearance of the stable phase for the Y-containing alloys cannot be discarded, at least partially, as indicated in [27,29]). The third reaction should correspond to the growth of the nucleated phases, with the ternary phase then transforming to the stable Al$_{19}$Y$_3$Ni$_5$ or Al$_{19}$Sm$_3$Ni$_5$ phases. This later transformation must take place during this same third reaction in the Y alloys (which according to Figure 2 showed a bigger reaction), or during the small fourth DSC reaction in Sm-containing alloys, which however was not clearly found in the DSC traces when Cu is present.

Regarding Ce alloys, the behavior was significantly different, with Al$_8$Ni and Al$_8$Ce$_3$ being the only phases identified for any of the heat treatments. These phases must probably appear again during the second DSC reaction, and grow during the third reaction.

Regarding the Y and Sm alloys, it is worth further investigating the peculiarity previously indicated. The fourth DSC reaction is a very small reaction, at times undetectable (coincident with the third one), causing the metastable to the stable phase transformation. According to this, a very easy process must be taking place, and the metastable phase should be very similar in composition and structure to the stable one, not even making a new nucleation and growth process necessary, but an easier transformation that just adapts to the new lattice parameters, probably with a very similar composition. In order to check this hypothesis, the Al$_{87}$Sm$_4$Ni$_8$Cu$_1$ diffraction pattern was simulated for 375 °C + 30 min (considering a metastable phase with very similar crystalline structure to the Al$_{19}$Sm$_3$Ni$_5$) and compared with the experimental one. The initial lattice parameters to consider were obtained from a metastable particle, studied by TEM microdiffraction in the Al$_{87}$Sm$_4$Ni$_8$Cu$_1$ ribbon treated to 375 °C + 30 min (Figure 8), where the metastable phase is the only one present.

**Figure 8.** TEM analysis of a metastable intermetallic particle in an Al$_{87}$Sm$_4$Ni$_8$Cu$_1$ ribbon treated to 375 °C + 30 min. Angles between patterns and diffraction spot distances served to index DP.

The real lattice parameters obtained from spot distances of the reciprocal lattice in the DP of Figure 8 resulted in values for the metastable phase of about 0.41 nm, 1.86 nm, and 2.81 nm, certainly similar to those values considered for the Al$_{19}$Sm$_3$Ni$_5$ phase (0.410 nm, 1.598 nm, and 2.689 nm). Although measurements were directly made on TEM DP images, the simulation after heat treating
the ribbons to 375 °C + 30 min, as shown in Figure 9, was very similar to the experimental one, therefore corroborating the previous ideas.

Figure 9. XRD patterns of the Al$_{87}$-Sm$_4$-Ni$_8$-Cu$_1$ ribbon treated to 375 °C + 30 min. The simulated pattern considered proportions of 40/40/20 for Al, Al$_3$Ni, and the metastable phase.

3.5. Microstructural Identification

Another aspect that was studied in this work was the shape of the intermetallics, depending on the RE used in the alloy. Al and Al$_3$Ni particles always appeared with a relatively equiaxed morphology, however, some other differences can be found in the ternary phases.

Figure 10 shows the TEM micrographs of the different alloys after heating to 375 °C and held for 30 min. The Cu-free Y alloy showed relatively equiaxial shapes, with the Sm alloy with more elongated shapes. Adding Cu to these alloys makes them appear as elongated shapes or increases their length, reaching up to about 300 nm. On the other hand, Ce alloys always showed equiaxial shapes, independently of adding Cu or not.
Figure 10. TEM bright field images of heat treated ribbons to the 375 °C held for 30 min of (a) Y, (b) Sm, and (c) Ce-containing alloys.

According to Figure 10 and the XRD evolution shown in Figure 6, it could be established that for the Y and Sm-containing alloys, the longer the time the metastable phase is present in the alloy, the higher the trend to develop elongated shapes. As previously explained, the stable Al_{19}RE_{3}Ni_{5} phase of these alloys appeared without new nucleation and shape change from the metastable alloy, and therefore, the one growing with elongated shapes must be the metastable phase. The two extreme conditions were the Y-containing Cu-free alloy, where the metastable phase was not even detected by XRD and indeed, only equiaxial shapes were found, and the two Cu-containing Sm alloys, in which elongated shapes had more time to grow. For the Ce-containing alloys, in which no metastable phase was detected, the shape found was always equiaxial.

The TEM study after heating to higher temperatures corroborates this idea. Figure 11 shows the images after heating to 450 °C.
Figure 11. TEM bright field images of heat treated ribbons up to 450 °C of (a) Y, (b) Sm, and (c) Ce-containing alloys.

As expected, after reviewing the XRD changes in Figure 6, the micrographs in Figure 11 of Y-containing alloys did not show evident changes, and neither did the Cu-free Sm alloy. This is because no changes were detected by XRD between both heat treatments. However, Cu-containing Sm alloys showed even longer shapes because at 450 °C, the stable phase was still present, and therefore the metastable phase had more time to grow. It has to be noted that the microstructures shown in Figure 11 correspond, in general, to the stable phases, and, as expected, the shapes were maintained during the change from the metastable to the stable phase. The similarities in structure make nucleation of the stable phase unnecessary, but just an easy adaptation to the new lattice parameters and composition.

Finally, previous ideas concerning the unidentified metastable phase to Al6RE5Ni5 transformation were confirmed by TEM microdiffraction on particular grains of heat treated ribbons to 450 °C (Figure 12). Indeed, the measurements on DP of the studied particles confirmed the expected lattice parameters.
Figure 12. TEM bright field images of Al<sub>88</sub>RE<sub>4</sub>Ni<sub>8</sub> particles in ribbons treated to 450 °C, and microdiffraction patterns of (a) Al<sub>88</sub>Y<sub>4</sub>-Ni<sub>8</sub>-Cu, and (b) Al<sub>88</sub>Sm<sub>4</sub>-Ni<sub>8</sub> alloys. Reciprocal lattice parameters are used in DP.

4. Conclusions

Amorphous structure formation in Al<sub>88</sub>-RE<sub>4</sub>-Ni<sub>8</sub> alloys was quite similar for the Y, Sm, and Ce-containing alloys, although the Sm and Ce alloys seemed to show a higher GFA. This was confirmed by the stability of the amorphous phase when heat treated. A 1 at.% Cu addition to the Al<sub>88</sub>-RE<sub>4</sub>-Ni<sub>8</sub> amorphous alloys affected their GFA and very clearly their stability. Al substitution by Cu increased the amorphous structure stability, whereas Ni substitution by Cu decreased it.

Destruction of the amorphous structure started with a primary crystallization process leading to the formation of α-Al nanocrystals that were smaller when Cu was present in the alloy.

At higher temperatures the crystallization process continued in different ways for the Y and Sm-containing alloys on one side, and the Ce-containing alloys in the other. For the Y and Sm-containing alloys, Al<sub>3</sub>Ni and unidentified metastable Al-RE-Ni intermetallic particles nucleated in a second crystallization process, subsequently growing and transforming to the stable ternary phase Al<sub>19</sub>RE<sub>3</sub>Ni<sub>5</sub> in the final stages of crystallization. The transformation to the stable phase was easier in the Y-containing alloys and also easier for the Cu-free alloys in both the Y and Sm alloys.

On the other hand, Ce-containing alloys seemed to evolve with the formation of Al<sub>3</sub>Ni and Al<sub>11</sub>Ce<sub>3</sub> during the second reaction, and growth during the third reaction.

Regarding the shape of these intermetallic phases, for the Y and Sm-containing alloys, the metastable ternary phase appeared to have an elongated shape, with a higher stability in the Sm alloys allowing them to reach even longer shapes. Once transformed to the stable Al<sub>19</sub>RE<sub>3</sub>Ni<sub>5</sub> phases, the shape reached in the metastable phase was maintained. This means that the transformation takes place without a nucleation process, but is only a structural transformation because of a similar structure in both the metastable and stable phases. According to the prevalence of the metastable phase, Cu-free Y alloys presented a clear equiaxial aspect, whereas phases in Cu-containing Sm alloys appeared with an elongated acicular morphology. On the other hand, Ce-containing alloys only showed phases with equiaxial shapes.

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