Influence of Small Content Elements Additions on the Glass Forming Ability of Zr-based Bulk Metallic Glasses Alloys

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Received: December 12, 2017; Revised: April 24, 2018; Accepted: August 19, 2018

The present work reports investigation on the Glass Forming Ability (GFA) of Zr-based Bulk Metallic Glasses (BMG’s) by microalloying with early transition metals. GFA was measured as the amorphous fraction formed in samples with different diameters using optical microscopy (OM) and image analysis techniques. Samples with the highest GFA had their oxygen content measured in a Leco RO-400. This study shows that additions of Molybdenum or Iron in a Zr48Cu(48-x)Al4Mx (M = Mo or Fe) alloys resulted in a minor GFA improvement, but far from the results reported in the literature with Nb additions in this system. Niobium microalloying to a Zr62Cu15.5Al10Ni (12.5-x)Nb x and (Zr55Cu30Al10)100-xNb x alloys have a deleterious effect on the GFA. Oxygen measurements of (Zr55Cu30Al10)99Nb1 and Zr62Cu15.5Al10Ni12.4Nb0.1 alloys have shown similar oxygen content, indicating that oxygen was not a limiting factor on this study. The amorphous fraction quantification from OM image analysis maintains the same fraction ratio from the heat released at the crystallization event from heating DSC analysis.

Keywords: Bulk Metallic Glasses, Rapid-solidification, Metallography, Glass Forming Ability, Oxygen.

1. Introduction

The application of Bulk Metallic Glasses (BMG) as engineering materials is the goal of many research teams spread around the globe. BMG have outstanding mechanical properties, compared to the conventional alloys, due to high strength1, resilience2, wear resistance3 and corrosion resistance4. In order to use BMGs in structural applications the research groups are investigating ways of increasing the Glass Forming Ability (GFA) and ductility of such alloys.

The literature shows that GFA and ductility can be both improved with the use of minor addition of elements, or microalloying technique5. Microalloying has improved the GFA of many BMG systems by stabilizing the amorphous phase6, removing deleterious elements from the glass matrix6 or altering its crystallization kinetics7. The ductility of several alloys was increased by the compositional change8,9 or by the formation of a second phase10-11 embedded in the amorphous matrix.

The present work reports our investigation of microalloying effects on the GFA of Zr-based BMG, sharing some data to other groups and supporting the development of commercial BMG. Zr-based BMG was one of the first BMG to achieve amorphous diameters larger than 10 mm20 and nowadays it is the only commercially feasible, known as Vitreloy21. There is a vast literature of microalloying Zr-based BMG with early22 and late23 transition metals but very few studies with elements from the groups 5-8. This study has the objective of report the data of the microalloying effect of Molybdenum, Iron and Niobium upon the GFA of Zr-based BMG. The GFA was measured as the amorphous fraction formed in samples with different diameters by image analysis of optical microscopy and correlated with differential scanning calorimetry (DSC) and scanning electron microscopy (SEM). Oxygen was also measured showing that not only the GFA was improved but the oxygen deleterious effect was minimized.

2. Experimental Procedure

The present study was made considering glassy alloys of various compositions based on the Zr-Cu-Al system described by the general formulas (at. %): Zr48Cu(48-x)Al4Mx with M = Mo or Fe; Zr55Cu15.5Al10Ni12.5xNb x and (Zr55Cu30Al10)100-xNb x where x could be equal to 0.1; 0.5; 1.0; or 1.5. This set of values was chosen in order to analyze the influence of microadditions of the presented elements on the GFA of the glassy alloys.

All alloys used in the present study were produced by arc-melting from their respective constituent high purity raw elements (99%+), in a high purity argon atmosphere, on a copper hearth cooled bed with circulating room temperature water. Each initial ingot was arc-melted at least five times and flipped between steps to assure complete homogeneity. Afterwards, cylindrical specimens with diameters varying from 2 to 7 mm were produced by suction casting of the liquid metal into copper molds.

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The samples were cut in the cross-section using a diamond saw disc equipment for metallographic preparation. The samples were then embedded in polymeric resin and prepared for optical microscopy (OM), by grinding and polishing according to ASTM E3-11. The polishing step was performed on an automatic machine EcoMet 250 BUEHLER® using 1 µm alumina suspension. Subsequently, the samples had their surfaces etched by immersing them in an etching solution of HF (0.8% wt.), HNO₃ (14.6% wt.) and H₂O (84.6% wt.) for five seconds.

The OM analyses occurred by image acquisition of the samples etched surfaces. For that purpose, it was used the software Axion Vision AxioVs40 V4.8.2.0 in a Scope A.1 Zeiss microscope. The images obtained had their analysis based on ASTM E1245-95. The method consisted in the use of automatic software-controlled segmentation procedure to select and measure areas of interest in the images. In this case, these areas corresponded to the bright amorphous region. Once they were measured, it was possible to calculate its percentage considering the whole sample. Fig. 1 shows an example of measurement by image analysis. The red region corresponds to the amorphous fraction, while the green one is a region excluded from the analysis, i.e. sample mount and pores. The same quantification procedure was performed by 4 different trained operators in order to reduce the bias in the measurement method.

DSC analysis was made in four different alloys to evaluate and compare the results from image analysis with Enthalpy released during crystallization of these alloys. SEM back-scattered (BSE) images were acquired from two different alloys to confirm the presence of the amorphous phase detected by optical microscopy and quantified by image analysis.

The samples with the higher GFA, measured by their amorphous fraction, had the oxygen content measured by hot extraction using the inert gas fusion principle in a LECO RO-400 equipment following ASTM-E-1019. For that purpose, 9 pieces of around 0.1 g each were used in an oxygen free Ni basket. The oxygen levels of the samples were compared in order to evaluate any statistically significant difference among the samples.

3. Results and Discussion

3.1 Optical microscopy analysis

Table 1 shows the samples synthesized relating their composition with the respective diameters produced, in mm. The OM results of the amorphous fraction by the content of microalloying added for the Zr₄₈Cu₄₈₋ₓAlₓMₓ (M = Mo or Fe) samples are shown in Fig. 2. Figure 2(a) correspond to M = Fe and Figure 2 b) to M = Mo. Figure 2(a) shows clearly that the addition of Fe has a negative effect on the GFA, independently of the amount of Fe added in the alloy, where by increasing the amount of Fe it leads to a decrease
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Figure 2. a) $\text{Zr}_{48}\text{Cu}_{48-x}\text{Al}_4\text{Fe}_x$ with $x = 0.1$, 0.5, 1.0 and 1.5 % at.; b) $\text{Zr}_{48}\text{Cu}_{48-x}\text{Al}_4\text{Mo}_x$ with $x = 0.1$, 0.5, 1.0 and 1.5 % at.

Figure 3. Correlation between amorphous fraction and content of elements added for samples of general composition $\text{Zr}_{48}\text{Cu}_{48-x}\text{Al}_4\text{M}_x$ (M = Fe, Mo, Nb). Collected from 10,12-13

Figure 4. Correlation between amorphous fraction measured by OM and content of elements added for samples of general composition $\text{Zr}_{62}\text{Cu}_{15.5}\text{Al}_{10}\text{Ni}_{(12.5-x)}\text{Nb}_x$, with $x = 0$.

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In the amorphous fraction. Figure 2(b), on the other hand, shows that an increase of the GFA happened with additions of Mo 0.5% at. For both cases, additions of Fe or Mo, when the value of microalloying was increased from 1.0 to 1.5 (at%), even for 2 mm samples it was not observed a significant amorphous fraction, therefore, the parent composition was the limiting factor.

Figure 3 shows the comparison of our results with the same based alloy studied with different Nb amounts 10,12-13. All the points were fixed for samples with 3 mm in diameter, as a base for comparison. It is possible to see that our samples with the addition of Fe and Mo resulted in alloys with the least amount of amorphous phase formed. This difference between our results and the literature has many possible explanations. The first, and most obvious, is the alloy composition which indicates that the Nb additions have a positive influence in the alloy’s GFA over Fe and Mo additions. The other explanation can be related to the oxygen content in the samples. It is well-known that the oxygen has a strong influence on GFA of Zr-based BMG 14-17. Once the literature used for the present comparison lacks the information about the oxygen content of the samples it is impossible to make an affirmation based only on the sample composition. Comparing the effect on GFA of our samples with the literature, the addition of 1 at% of Nb on the $\text{Zr}_{62}\text{Cu}_{15.5}\text{Al}_{10}\text{Ni}_{(12.5-x)}\text{Nb}_x$ seems to be the best option to enhance the GFA of this alloy.

The results for the composition of $\text{Zr}_{62}\text{Cu}_{15.5}\text{Al}_{10}\text{Ni}_{(12.5-x)}\text{Nb}_x$ can be seen in Fig. 4. For diameters above 4 mm, the amorphous fraction measured was not significant, below 30%. When considering 3 and 4 mm diameter samples however, it was possible to notice that the glass content decreased as the value of the microalloying element increased. In other words, more additions of Nb tend to affect negatively the GFA. It is possible to assume that the critical cooling rate (Re) value for this composition increases drastically. The base alloy $\text{Zr}_{62}\text{Cu}_{15.5}\text{Al}_{10}\text{Ni}_{(12.5-x)}\text{Nb}_x$, with $x=0$, is reported
to be feasible to produce a fully amorphous rod up to 5 mm in diameter\textsuperscript{18}. In this study, we have found around 30% of amorphous fraction for the case of Nb addition (0.1 at%) at 5 mm. Therefore, it is possible to extrapolate that the addition of Nb on the base alloy has not a positive effect.

Fig. 5 displays the results of OM quantification of (Zr\textsubscript{55}Cu\textsubscript{30}Al\textsubscript{10})\textsubscript{100-x}Nb\textsubscript{x} samples. As can be seen, 5, 6 and 7 mm diameter samples presented amorphous content smaller than 30% for practically any content of Nb. A good result in terms of amorphous fraction was only observed for Nb = 1.0% and 2 mm diameter sample, meaning that our equipment was able to achieve cooling rates closer to the Rc of the alloy. When analyzing 3 and 4 mm samples, the amorphous fraction was measured between 47.5 and 67.5% for Nb = 1.0%. One drawback related to OM quantification in samples with larger diameters is that the resolution decreases with the increase of sample size.

3.2 Oxygen content measurement

The compositions with the higher GFA had their oxygen content measured and the results are shown in Fig. 6. For each sample, there were 9 measurements of oxygen content. The difference in average oxygen measured in these 2 different alloys cannot be based on the difference of the Zirconium masses between the two alloys as proposed previously\textsuperscript{19}. The two sample t-test failed to reject the null hypothesis. The alloy (Zr\textsubscript{55}Cu\textsubscript{30}Al\textsubscript{10})\textsubscript{99}Nb\textsubscript{1} has an average of 922 ppm of oxygen and a large dispersion compared to the alloy Zr\textsubscript{62}Cu\textsubscript{15.5}Al\textsubscript{10}Ni\textsubscript{12.4}Nb\textsubscript{0.1}, with an average of 981 ppm of oxygen. These results shows that both alloys have the same oxygen content within a 95% confidence interval.

3.3 Scanning electron microscopy and differential scanning calorimetry

Figure 7 shows two BSE images from 3 mm samples of the alloys a) Zr\textsubscript{48}Cu\textsubscript{47.9}Al\textsubscript{4}Fe\textsubscript{0.1} and b) Zr\textsubscript{48}Cu\textsubscript{47.5}Al\textsubscript{4}Mo\textsubscript{0.5}. From the images it is clear the presence of a brighter amorphous matrix and darker crystalline phases. The same samples were analyzed by DSC and the results are showed at Table 2 and Figure 8, where the ratio of heat released in the crystallization event is the same as from the ratio of amorphous fraction quantification by OM image analysis. The samples from 3 mm (Zr\textsubscript{55}Cu\textsubscript{30}Al\textsubscript{10})\textsubscript{98.5}Nb\textsubscript{1.5} alloys are also being showed at Table 2 and Figure 8, and the same effect of ratio in the heat released in the crystallization event was the same as from the ratio of amorphous fraction quantification by OM image analysis in these samples. This supports that the amorphous fraction quantification by OM image analysis is a cheaper and valid process for measuring the GFA from the alloys that maintains the same fraction ratio from the heat released at the crystallization event from heating DSC analysis.

4. Conclusions

The present study performed on Zr\textsubscript{48}Cu\textsubscript{48-x}Al\textsubscript{M}x, where M = Fe or Mo, showed that the addition of these elements decrease the alloy’s GFA. The GFA of the Zr\textsubscript{48}Cu\textsubscript{48-x}Al\textsubscript{M}x system have been successfully improved with M = Nb as reported in the literature\textsuperscript{10,12-13}. Therefore, our results show that the use of Fe or Mo is not a viable alternative. In the second part of our study, we tried to investigate whether the GFA improvement by the Nb microalloying could be extended to other Zr-based systems. Our results show that the addition of Nb from 0.1 at.% to 1.5 at.% in the Zr\textsubscript{48}Cu\textsubscript{48-x}Al\textsubscript{10}Ni\textsubscript{12.5-x}Nb\textsubscript{x} and the (Zr\textsubscript{55}Cu\textsubscript{30}Al\textsubscript{10})\textsubscript{100-x}Nb\textsubscript{x} alloys have a deleterious effect on the alloys’ GFA. Thus, the GFA improvement by the Nb microalloying is not an inherent feature on Zr-based BMG.

The oxygen measurements of the (Zr\textsubscript{55}Cu\textsubscript{30}Al\textsubscript{10})\textsubscript{99}Nb\textsubscript{1} and Zr\textsubscript{62}Cu\textsubscript{15.5}Al\textsubscript{10}Ni\textsubscript{12.4}Nb\textsubscript{0.1} alloys have shown that the higher content of Zr in the later alloy do not increase the tendency to be contaminated with oxygen. This hypothesis has been raised in a previous study\textsuperscript{19} where the higher oxygen content...
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Figure 7. SEM BSE images from a) Zr_{48}Cu_{47.9}Al_{4}Fe_{0.1} and b) Zr_{48}Cu_{47.5}Al_{4}Mo_{0.5}.

Table 2. DSC analysis showing the characteristic temperatures from the alloys, where the temperatures $T_g$ = glass transition; $T_x$ = crystallization; $\Delta T_x$ = supercooled liquid region; $\Delta H_x$ = enthalpy at crystallization.

| Alloys                     | D (mm) | $T_g$ (K) | $T_x$ (K) | $\Delta T_x$ (K) | $\Delta H_x$ (J/g) |
|----------------------------|--------|-----------|-----------|------------------|-------------------|
| Zr_{48}Cu_{47.9}Al_{4}Fe_{0.1} | 3      | 656       | 744       | 88               | 30.2              |
| Zr_{48}Cu_{47.5}Al_{4}Mo_{0.5}   | 3      | 650       | 744       | 94               | 41.7              |
| (Zr_{55}Cu_{30}Al_{10})_{99}Nb_{1.0} | 3      | 665       | 733       | 68               | 54.4              |
| (Zr_{55}Cu_{30}Al_{10})_{98.5}Nb_{1.5} | 6      | 662       | 735       | 97               | 27.8              |

Figure 8. DSC heating curves for 3 mm Zr_{48}Cu_{47.9}Al_{4}Fe_{0.1}, Zr_{48}Cu_{47.5}Al_{4}Mo_{0.5}, (Zr_{55}Cu_{30}Al_{10})_{99}Nb_{1.0}, and 6 mm (Zr_{55}Cu_{30}Al_{10})_{98.5}Nb_{1.5}.

of the alloys with higher Zr content would be explained by the Zr gettering nature.

The amorphous fraction quantification by OM image analysis is a cheaper and valid process for measuring the GFA from the alloys that maintains the same fraction ratio from the heat released at the crystallization event from heating DSC analysis.

5. Acknowledgements

The authors would like to thank Boeing Research & Technology Brazil (BR&T-Brazil) for the support for this research and for support of the University of Sao Paulo - Sao Carlos School of Engineering (EESC).

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