Effect of Deformation on the Transformation Temperatures of Martensite and Bainite Structures in Cu–9.97%Al–4.62%Mn Alloy

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At present study of Cu–9.97%Al–4.62%Mn alloy system is used. Some physical properties of martensitic and bainite phase transformations in Cu–9.97%Al–4.62%Mn (wt%) alloy were investigated by means of scanning electron microscopy and differential scanning calorimetry. In scanning electron microscope observations of Cu–9.97%Al–4.62%Mn alloy were noted two kinds of martensitic phases. These phases were defined as $\beta'$ (M18R) and $\gamma'$ (2H) martensites structures. According to differential scanning calorimetry measurement results, the deformation induced temperature is expected to be higher than the thermally induced temperature in Cu–9.97%Al–4.62%Mn alloy for martensite and bainite structures.

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1. Introduction

Cu-based shape memory alloys are economical materials, therefore they can be used in many automotive, aerospace, robotic and biomedical fields [1]. Particularly, high temperature shape memory alloys have been studied for actuators in various aero-engine applications [2].

When the Cu-based shape memory alloys go through ageing on $\beta$ parent phase, a series of transformations can take place, such as martensite, bainitic and eutectoid transformation. The Cu-based shape memory alloys have found that during the quenching procedure the chemical ordering sequence $A2 \rightarrow B2 \rightarrow DO_3-L_2$ [3]. In copper-based shape memory alloys, composition and heat treatment plays an important role in determining their transformation temperatures and other characteristics. The previous study is showed that the transformation temperature decreases with an increase in the aluminum and manganese contents. Also, precipitates and dislocations effects the transformation temperatures [4]. The purpose of this work is to investigate the effects of heat treatments and deformation.

2. Experimental

The Cu–9.97%Al–4.62%Mn (wt%) alloy was chosen for the present study, which was prepared by vacuum induction melting under an argon atmosphere from pure (99.9%) alloying elements. First group of samples were homogenized at 800°C for 1 h and then quenched into ice water, the second group of samples were homogenized at 800°C for 1 h and furnace cooled to 25°C at a rate of approximately 2°C/min and then air cooled. These samples are denoted as A1 and A2, respectively. For scanning electron microscope observations, the samples were mechanically polished and etched into a solution composed of 2.5 g FeCl$_3$·6H$_2$O and 48 ml methanol in 10 ml HCl for 10 min. Scanning electron microscope observations were made using a JEOL 5600 scanning electron microscope, operated at 20 kV. Subsequently, homogenized samples were plastically deformed by 7% with an INSTRON 8510-type machine, operated at a constant strain rate of 0.2 mm/min at room temperature. These samples are designated as A3 and A4, respectively. The samples of alloy were prepared, and DSC curves were obtained. A Perkin-Elmer Sapphire model DSC equipment was used for all measurements. The transformation temperatures were determined using a differential scanning calorimetry (DSC) measurements from room temperature to 600°C by heating/cooling the samples at the rate of 10°C/min.

3. Results and discussion

In our previous studies, the crystal structures and microstructures determined for samples A1, A2, A3 and A4 [5, 6]. As can be seen from Fig. 1, the samples consists of microstructures at room temperature. In samples A1 and A3 formed $\beta'$ and $\gamma'$ of martensite structures from DO$_3$ parent phase. The bainite and $\gamma'$ martensite variants formed by effects of thermally and deformation in sample A4 [6]. The transformation temperatures for phase transformations of Cu–9.97%Al–4.62%Mn alloy have also been determined by differential scanning calorimetry. To compare the transformation characteristics of this alloy, samples were prepared as the undeformed and deformed.

Figure 2a,b shows the differential scanning calorimetry heating and cooling curves of samples A1 and A3. Three
calorimetry measurement results, sample A3 temperature is expected to be higher than the sample A1 temperature for Cu–9.97%Al–4.62%Mn alloy. These results are shown in Fig. 2b, which indicates that the transformation temperatures are raised by an increase in stress. In our study, the driving force occurred in sample A3 during the deformation is larger than that of sample A1. On the other hand, both kind of the produced martensite structures show different transformation characteristics. Their hysteresis temperatures differ by approximately 20°C [9, 10]. The changes on the transformation temperatures are attributed to the relative stability of the product phase and closely related to the corresponding thermodynamic properties such as enthalpy. These results are in agreement with the observation made by previous works [13, 14].

Differential scanning calorimetry curves of the alloy samples for both forward and reverse transformations are presented in Fig. 3a,b differential scanning calorimetry curves exhibit a prominent multi-peak behavior for the samples. During the cooling from 600°C to room temperature, the bainitic transformation starts ($B_s$) at 481°C for sample A2 and at 486°C for sample A4. The bainitic transformation finishes ($B_f$) at a temperature of 466°C and 470°C for samples A2 and A4, respectively. During heating of the samples, the reverse transformation starts at a temperature of 497°C ($A_s$) and finishes at 513°C ($A_f$) for sample A2. It was determined that reverse transformations starts at 500°C and finishes at 518°C for sample A4. In addition, the sample A4 observed a sharp peak after the deformation. This sharp peak corresponds to 536°C ($\beta \rightarrow \alpha + \gamma_2$) eutectoid decomposition [8, 9].
It is believed that both the nucleation and growth mechanisms of bainite are shear, which are governed by solute atom diffusion. It was found that the activation energy for bainite nucleation is proportional to the driving force available for transformation [15, 16]. According to differential scanning calorimetry measurement results, sample A2 temperature is expected to be higher than the sample A4 temperature for Cu–9.97%Al–4.62%Mn alloy. This behavior can be explained by taking account of crystal defects brought about by plastic deformation such as dislocations [17]. These defects are increased in transformation temperatures [18].

4. Conclusion

The copper-based alloy are important for technological applications and the transformation temperatures vary with the variation applied by various thermal treatments and deformation. The transformation temperatures values of samples are increased with applied deformation. In this study, it has been observed that the transformation temperatures are high for the alloy studied. Therefore, Cu–9.97%Al–4.62%Mn alloy can be suitably designed for use in different fields where high temperature applications such as in actuators and automotive components. Our study give rise to new application opportunities and this is particularly true in the development of new materials.

References