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Effect of Deformation and Thermal Treatment of NiTi Alloy on Transition Sequence

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Abstract: Transformation sequences in a NiTi shape memory alloy after deformation and subsequent annealing were studied using various methods. Low temperature annealing of the deformed alloy caused that the R-phase preceded the martensitic transformation. The R-phase formation during both cooling and heating depends strongly on the deformation degree and temperature of annealing. On DTA curves an additional peak in the martensitic transformation range was observed for specimens annealed at 500 and 550°C. It is suggested that this peak was caused by presence of dislocation cell structure and subgrains whose boundaries interact with growing martensite plate.

1. INTRODUCTION

Deformation and subsequent annealing of NiTi alloys offers great possibilities of controlling temperatures and sequences of transformations. Todoroki and Tamura [1] showed that, depending on annealing temperature, changes in transformation sequences in deformed alloys occur during both heating and cooling. Abujudom et. al. [2] found that the deformation degree significantly influences both course and characteristic temperatures of the transformations. The authors of [3-8] studied the effect of both deformation degree and annealing temperatures on the course of transformations. In [4,5] and [8] the influence of deformation and annealing on shape recovery phenomenon was investigated. It was shown [8] that large deformations have a positive influence on shape recovery. However, the results mentioned above show certain discrepancies, especially in variations of characteristic temperatures versus annealing temperature and in sequences of transformations. The aim of this paper was to establish the influence of dislocation density and configuration on the characteristic temperatures and sequences of transformations.

2. EXPERIMENTAL PROCEDURE

A commercial NiTi shape memory alloy containing 50.6 at % Ni was used in this study. Specimens in shape of strips were quenched from 800°C after annealing for 1h and then cold-rolled by 5%, 10%, 20% or 25% reduction. The deformed samples were then annealed at various temperatures between 300-550°C. The transformation behaviour and the characteristic temperatures were determined using the DTA method on the TA1 Mettler instrument and the internal friction method on an acoustic relaxator with the logarithmic decrement of damping.

The phase composition of the specimens was established using the X-ray phase analysis on a Philips diffractometer equipped with a temperature attachment. CuKα radiation was used. The alloy structure was studied on thin foils using JEM 200B transmission electron microscope.
3. EXPERIMENTAL RESULTS

3.1. Effect of deformation

The studied alloy, after quenching from 800°C, consisted of the parent B2 phase only as was shown by X-ray studies. Deformation by cold-rolling caused strengthening of the alloy as evidenced by increase in its hardness. The 110 B2 diffraction line widened due to increase in lattice strain and density of dislocations (Fig. 1).

Fig. 1 Hardness, $M_s$, temperature and half-width of the 110 B2 versus deformation degree

Fig. 2 Changes of enthalpy of reverse transformation and amount of martensite versus deformation degree

Fig. 3 The DTA curves obtained during cooling of the alloy deformed by 10%

Fig. 4 The DTA curves obtained during heating of the alloy deformed by 10%
This plot shows that the $M_s$ temperature decreases with the increase in deformation degree. Plastic deformation of the parent phase causes the creation of martensite in a quantity increasing with increase in deformation (Fig. 2). After deformation by 20-25% the DTA cooling curve spreads to such a degree that the martensitic transformation is hardly visible. This indicates that the quantity of material undergoing transformation was small. More distinctive effects were observed on the heating curve. From the DSC curves obtained for these specimens the enthalpy of the reverse transformation $\Delta H_{M\rightarrow B_2}$ was calculated. Enthalpy changes versus deformation degree are shown in Fig. 2.

3.2 Effect of annealing

The effect of annealing of the deformed alloy on the course of transformation was studied using the DTA method. Fig. 3 shows the DTA cooling curves for the alloy deformed by 10%, and then annealed at various temperatures. The curve of the deformed alloy shows a strongly spread peak of martensite transformation on which overlapping of two effects may be distinguished, i.e. that of R-phase formation and, at lower temperatures, of martensite formation. Annealing of the alloy at 300°C causes distinct separation of these two effects. The separation becomes more visible when the annealing temperature increases up to 550°C. Figure 3 also illustrates changes of the characteristic temperatures of both transformations versus the annealing temperature. Increase of annealing temperature up to 600°C causes disappearance of the R-phase transition peak.

The DTA curves obtained during cooling of specimens annealed at 400-550°C (except for the specimen annealed at 450°C) show three distinct peaks, one from R-phase formation and two in the range of the martensitic transformation. Fig. 4 presents the DTA curves obtained during heating of the same specimens. The reverse transformation for specimens annealed at 350°C and 400°C also consists of two distinct effects from the $M\rightarrow R$ and $R\rightarrow B_2$ transitions.

Fig. 5 shows the influence of annealing temperatures on half-width of the 110 $B_2$ line for the alloy deformed by 10%. The decrease in this parameter with increase in annealing temperature has a form similar to that for decrease in alloy hardness and is characteristic for the recovery process. Annealing at a temperature higher than 550°C does not cause any changes in this parameter. Disappearance of defects after annealing at these temperatures is also indicated by $M_s$ increase, whereas the $M_s$ decrease for specimens annealed at 400-500°C shows that besides lowering of defects density, there are other factors influencing the temperature. One of these could be the formation of dislocation cell structure. Thus, although total dislocation density decreases this structure inhibits the interface mobility and lower the $M_s$ temperature. Also the occurrence of the R-phase transition in the specimen can influence the martensitic transformation in such a way that the $M_s$ temperature decreases.

Transmission electron microscopy studies of alloy dislocation structure were carried out to investigated the influence of annealing on the course of transformations and on characteristic temperatures. Changes of the alloy structure versus temperature of annealing are shown on Fig. 6. A dislocation cell structure is visible. Dislocation density decreases with increase in annealing temperature. Annealing at 500°C causes partial recrystallization. Characteristic for this stage are areas where the polygonization process

![Fig. 5 Hardness, $M_s$ temperature and the half-width of the 110 $B_2$ line for a specimen deformed by 10%](image-url)
formed regular network of dislocations creating subgrains of low-angle boundaries. Annealing at 600°C causes full recrystallization of the deformed alloy. In order to explain the presence of the two peaks on the DTA curves in the martensitic transformation range, internal friction measurements were carried out. Results of these measurements obtained for alloy deformed by 10% and annealed at 400°C are shown on Fig. 7.

On the internal friction ($Q'$) curve obtained during cooling there are three peaks. The first one, $P_1$, appears at the same temperature as the B2→R DTA peak. The fact that this peak is due to the B2→R transition is confirmed by the minimum on the curve of square of vibration frequency ($f^2$), which is proportional to the elastic modulus. The second internal friction peak $P_2$ does not have any equivalent either on the DTA or on the $f^2$ curves. Thus it is not connected with the martensitic transformation. Its presence may be explained by relaxation of dislocations.

The third internal friction peak corresponds to the first martensite peak on the DTA curve. There is also a small but distinct minimum observed at this temperature on the $f^2$ curve. The internal friction curve obtained during heating consists of two peaks which correspond to the temperatures at which peaks on the DTA curve arise as a result of M → R and R → B2 transformations. The second peak appearing on the DTA cooling curve close to that from the martensite transformation could be caused by interaction between the growing martensite plates and dislocation cell boundaries. For a plate to pass these boundaries additional energy is required and this may result in an additional peak on a DTA curve. This explains the fact that the peak is very clear for specimens annealed at 500°C and 550°C which contain a large number of subgrains formed by small-angle boundaries.

4. DISCUSSION
Plastic deformation of the parent phase produces dislocations, the density of which can be estimated from the increase in half-width of an X-ray diffraction line for the phase. The higher the deformation degree, the wider is the 110 B2 line. The form of the curve representing the half-width versus deformation degree is similar to that for alloy hardness. The $M_s$ temperature decreases with increase in dislocation density, which is due to the fact that the dislocations present obstacles for the moving interfaces. On the other hand, dislocations provide the places where the R-phase nucleates [9]. For small deformations the DTA curve obtained during cooling exhibits a peak corresponding to the R-phase formation. The higher the deformation, the more strain-induced martensite appears in the specimens, and the deformed B2 phase becomes stable. Thus the quantity of the parent phase undergoing thermal martensitic transformation decreases, as is proved by decrease in transformation heat obtained from the DSC measurements. Similar conclusions based on internal friction studies were presented in [10]. The annealing temperature has a vital influence on the course of transformations. Results presented on Figures 3 and 4 are in good agreement with data given in [1,2]. In the annealing range of 300-550°C the transformation occurs in two stages B2→R→M and after annealing at a temperature higher than 550°C only one stage transformation B2→M was observed. The same is true for the reverse transformation. Specimens annealed at low temperatures exhibited two stages of transformation course M→R→B2 while with increase in annealing temperature overlapping of the two peaks is observed. The DTA curves presented on Fig.3 exhibit two peaks in the range of the martensitic transformation for specimens annealed at 400, 500 and 550°C. Similar effects were observed for larger deformations. Todoroki and

Fig. 6 Dislocation structure of the alloy deformed by 10% and annealed at 300°C (a), 400°C (b), 500°C (c), 550°C (d)
Tamura [1] obtained identical results on their DSC curves. Referring to the paper by Monasevich [11], they explained this effect as the formation of two types of martensite. Monasevich and Pascal observed continuous widening of the 111 martensite line during cooling. They concluded that monoclinic martensite formed from the parent phase transforms to a triclinic and established deviations from orthogonality as $\alpha = 0.3^\circ$ and $\gamma = 0.5^\circ$. It is noteworthy that this effect was observed in alloys where no R-phase transformation occurred. Airolldi et al. [12], in specimens after cycling, observed two peaks from the martensite as well as the R-phase peak. They explained this effect in the same way as the authors of [1]. We obtained similar results for specimens after cycling [13], but we also observed that dislocations created during cycling form a cellular structure. Analysis of the DTA and internal friction curves (Fig.7) excluded the possibility of the formation of two types of martensite. Additionally, the results of X-ray studies, not presented in this paper, showed that the transformation sequence is $B2 \rightarrow R \rightarrow B19'$. Thus we may conclude that the specific arrangements of dislocations are responsible for two peaks in the range of the martensitic transformation. Obstacles in the form of low-angle subgrain boundaries inhibit formation of the martensite in these places and cause the occurrence of a second peak at a lower temperature on the DTA and DSC curves.

Throughout the whole range of annealing temperatures, decrease in alloy hardness and in half-width of 110 B2 line is observed. This indicates that strengthening and lattice distortion of the alloy gradually disappears. Annealing at 500°C and higher does not change the density of defects any further, although alloy hardness still decreases. The $M_s$ temperature for this annealing range increases slightly. After annealing in the temperature range 400 - 500°C the defects' density decreases, as does the $M_s$ temperature. Liu and McCormick [14] explained this fact basing on free energy changes analysis. If there occurs the $B2 \rightarrow R$ transition, it shifts the next, $R \rightarrow M$ transformation to a lower temperature.

5. REFERENCES

Fig. 7 The DTA curves (upper part) and the internal friction curves (lower part) for the alloy deformed by 10% and annealed at 400°C.
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