HRTEM study of austenite and martensite in splat-cooled and nanoscale thin film Ni-Al

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Abstract. Selected results of electron microscopic investigations on the atomic and microstructure of Ni-Al austenite and martensite in new types of materials are presented. It is shown that splat-cooled material behaves similar as normal homogenised bulk material whereas in evaporated thin films the austenite is strongly stabilised.

1. INTRODUCTION

These days new types of materials of conventional alloys are investigated in view of their martensitic and shape memory characteristics. For Ni-Al the martensitic behaviour in the bulk material has been properly documented by several experimental as well as theoretical studies (for a recent review see [1]). In order to obtain new insight in the influence of the existing defects and the dimensions of the material on the atomic and microstructures occurring during this transformation, Ni-Al splat-cooled material and nanoscale thin films were investigated by conventional and high resolution transmission electron microscopy (CTEM and HRTEM). In the first case the rapid quench from the melt yields a very fast crystallisation process which implies the existence of a higher number of point and other defects when compared with the conventional arc-melting procedure used to obtain bulk material. Due to the very short quenching times, splats are usually very homogenous with respect to compositional variation on a macroscopic scale but they are also crystallised in many small grains. For the thin films, the nanoscale dimension of the thickness presents an opportunity to investigate the influence of an anisotropic environment.
Samples with a Ni content of 62.5 and 65 at.% Ni were splat-cooled at a rate of $5 \times 10^5$ K/s. During the splat-cooling a droplet of the melt is pressed to a thin film by two Cu pistons that are shot against each other. During the contact of the pistons with the melt heat is transferred from the melt into the pistons causing the melt to solidify. The resulting discs have a diameter of approximately 20 mm and a thickness of about 200 µm. From these 3 mm EM samples were cut with their normal parallel with the normal to the splat-cooled discs. These samples were then further thinned by the double-jet electropolishing technique using the conventional solution and parameters for bulk Ni-Al [2].

Figure 1. [001] HRTEM image of the precursor domains in a B2 grain of splat-cooled Ni$_{62.5}$Al$_{37.5}$. The inset shows the typical <110> diffuse intensity streaks around the Bragg reflections.

In splat-cooled material of 62.5 at.% Ni the majority of the grains are a few micron in diameter and consist completely of the B2 phase. Adjacent grains are often observed to have a very similar orientation. This is shown in the overview and SAED patterns in fig. 4 of ref. [3]. All investigated B2 grains are in the precursor state which is clear from the typical <110> diffuse intensity streaks in reciprocal space shown in the inset of fig. 1. The distorted domains in this precursor state do not, however, reveal the micromodulation observed in bulk material of this composition [2]. Indeed, from the HRTEM image of fig. 1 a contiguous configuration of distorted domains, each with an averaged diameter of approximately 2-3 nm, plus strong strain contrast is observed. This observation is similar to the one in bulk material with a higher Ni content [4,5]. In the domains revealing a well resolved dot pattern the shear distortion occurs on the <110> planes edge-on in this orientation and can be seen as a displacement of the dot rows
when viewing the image along a grazing 110 or 110 incidence. Depending on the actual location
the B2 ordering or the basic lattice is visible. The exact origin of this strong difference at such a
local scale is unclear. It could be an imaging effect because of different thicknesses or it could
indicate microscopic inhomogeneities in the ordering. In some regions a line instead of a dot
pattern is observed. When these lines are parallel with the <100> directions in the image plane, as
was the case for the bulk examples [4,5], this feature is explained by shear planes inclined with
respect to the electron beam which yields imperfect atom columns along the viewing direction.
However, in the present splat-cooled material these lines are often parallel with the <110>
directions in the image. In this case they are probably due to severe internal distortions as a result
of the many dislocations observed in CTEM images. Because of the strain contrast and the local
changes in the HRTEM image these dislocations are difficult to recognise in these high
magnification images. The lack of the micromodulation corresponds with the missing of any
intensity enhancements or satellites in the diffuse intensity streaks in reciprocal space.

Some of these grains are partially transformed into the microtwinned L10 martensite,
occasionally revealing a stacking close to that of the 14M structure [2,6]. Due to the smallness of
the grains and the fact that the transformation is apparently caught midway the hysteresis different
interfacial configurations can be investigated. In fig. 2 an example of a martensite-austenite
interface is shown. The B2 austenite is observed along its [111] direction with one of the edge-on
{110} families of planes being transformed into the close packed planes of the martensite. In the

![Figure 2. Low magnification HRTEM image of an austenite-martensite interface in splat-
cooled Ni62.5Al37.5 showing a change of habit plane. The star indicates the orientations
of the {110} families of planes in the B2 area observed along a [111] direction. The inset
shows a magnification of the rectangular area revealing the atomic resolution near the
interface.](image)
Fig. 3 shows a macrotwin interface between two martensite plates with a relatively short stacking sequence indicated on the image. This sequence, however, is not very well defined which could be due to the incomplete situation of the transformation. From this image it can be concluded that the interface is parallel with the (020) plane of the largest microtwin variants on both sides and which used to be a (110) plane of the austenite. At most locations the microtwin variants of one given plate stop at a microtwin of the other plate which yields a zig-zag shaped interface. The angle between both microtwin planes is 118°, originating from a 120° between the (110) planes in the austenite [7].
Samples with a 65 at.% Ni content generally completely transform to microtwinned martensite, usually as one microtwinned plate within a given grain but sometimes as self-accommodating groups. Some examples are shown in fig. 5 of ref. [3]. Occasionally indications of an irregular 14M-like structure are found. An example of the latter at a boundary with a normally microtwinned plate is shown in fig. 4 with both plates observed along the [110]_{L1_0} direction. The interface is viewed edge-on and perpendicular to the microtwin planes of the 14M plate. From a lower magnification (not shown) it is also seen to be close to parallel with the microtwin planes of the larger plate which are not visible but parallel with the horizontal close packed planes of this variant. In the present case the tips of the thinnest microtwins of the 14M-like stacking reveal more strain contrast than in fig. 2 but again no dislocations are observed in this projection. The microtwin planes do, however, exhibit a severe curvature of approximately 5° when crossing the boundary and becoming non-twinned close packed planes in the other plate, as indicated on fig. 4. The fact that the interface is not a symmetry plane for the configuration of both plates indicates that it was formed by coalescence and not by simultaneous growth.

In plates with wider microtwin variants irregular arrangements of microtwins exist due to specific interfaces between adjacent B2 grains. An example of such a configuration is shown in fig. 6 of ref. [3].

![HRTEM image of a martensite-martensite interface in splat-cooled Ni65Al35 showing strain contrast at the tips of the smallest twins. Severe curvature of the close packed planes is noted.](image-url)
Until now most studies of martensitic transformations in thin metallic films only considered films with a thickness in the order of microns. In this case the microscopic behaviour of the film still follows the normal bulk conditions. However, when the film thickness is decreased to a few nanometers the anisotropic nature of the film dimensions can play an important role in the martensitic transformation. Indeed, several investigators have already noted difficulties for transforming a thinned bulk sample in-situ. Also the increased number of lattice defects when compared with well homogenised bulk material will influence this transformation.

In the present study the aim is to produce monophase Ni-Al austenite films with a thickness in the nanometer range which can directly be investigated by the different EM modes. Two compositions, 62.5 and 65 at.% Ni (further referred to as 62.5Ni and 65Ni), are currently being used. In the bulk the $M_s$ temperature of 62.5Ni is reported to be between -100 °C and 0 °C while that for 65Ni is around 400 °C [8,9]. The evaporation source can be an alloy with this composition or a combination of separate Ni and Al sources with the proper ratio. In the latter case both elements are always evaporated simultaneously. Typical evaporation rates are a few Å/s and films of a few tens of nanometers are obtained. As substrate single crystals of ex-situ cleaved (001) NaCl and KCl and which can be heated during deposition up to 550 °C are used. NaCl presents the best epitaxial configuration for the preferred B2 structure but KCl allows for the highest substrate temperature. The vacuum of the evaporation instrument is within the $10^{-5}$ mbar range. After deposition the films are floated of by dissolving the substrate in water and then placed on a Cu EM grid.

In fig. 5 a series of SAED patterns comparing the different films prepared under slightly different conditions are presented. From the ring nature of these patterns it is immediately clear that most films consist of very small grains while the actual appearing structures can easily be concluded from the measured ring diameters and ratios between those. In practise the structures of the B2 and L1$_2$ phases are recognised depending on the actual evaporation and substrate conditions. For convenience a schematic of the observed rings is included in fig. 5a.

When comparing the experimental ring patterns with the schematic in fig. 5a, it is clear that in fig. 5b both ring systems exist while all other patterns only show the B2 phase. Comparing this with other results it is concluded that the main parameter here is the substrate temperature. Thus, if one wants to start with a monophase film the substrate temperature should not exceed 400 °C. It is also clear that the best epitaxy is obtained in fig. 5d which corresponds with the fact that the NaCl cubic lattice parameter only has a discrepancy of approximately 2 % with that of the B2, whereas this discrepancy becomes about 8 % for KCl. As a result the substrate-film correlation usually becomes (001)//(001), [100]//[100] as seen from the most intense reflections in the 110 (second) and 200 (fourth) rings, although some weaker reflections in the 100 ring also reveal the existence of a (001)//(001), [100]//[110] epitaxial relationship.
Figure 5. SAED patterns of Ni-Al films evaporated under different conditions: (a) schematic of B2 and L1₂ rings, (b) 62.5Ni on KCl at 450 °C, (c) 62.5Ni on KCl at 300 °C, (d) 62.5Ni on NaCl at 300 °C, (e) 65Ni on KCl at 400 °C, (f) 65Ni on NaCl at 400 °C.

In fig. 6 a HRTEM image of the same film as in fig. 5d is shown. The grains are a few nanometers in diameter and the central one clearly shows the [001] B2 type of lattice image. The edges of the grain are primarily parallel with (100) or (110) planes. Inside the grain the lattice image changes from site to site yielding a domain structure which can be compared with that of the precursor distortions as shown in fig. 1. Again no extra micromodulations as in bulk material of this composition are recognised. As expected from the SAED patterns, 65Ni films do not show grains of this size.

Figure 6. HRTEM image of 62.5Ni film evaporated onto NaCl at 300 °C. The [001] zone orientation of the central grain is clearly visible.
When both the 62.5Ni and 65Ni films were cooled to 173 °C no martensitic transformation was observed, which is clearly different from the bulk transformation. Different reasons for this changed behaviour could be suggested. First it is well known that a deposited film can have a composition different from that of the source and knowing the strong dependency of $M_S$ on composition, this parameter is very important. Different EDX measurement, however, indicate a slight enhancement of Ni with respect to the source which would in fact increase and not decrease $M_S$. An alternative explanation for the increased stability of the austenite could lie in the increased number of point defects [10] and the increased yield strength of the austenite by grain refinement [11]. Indeed, in view of the low diffusion kinetics at the deposition temperatures, a high density of point defects such as vacancies and antisite atoms can be expected. Also the two-dimensional nature of the film could allow for some extra strain relief without the need for a martensitic transformation.

![SAED and microstructure of a 62.5Ni film heated in-situ up to 1000 °C.](image)

In an attempt to overcome these austenite stabilising features the 62.5Ni film was heated in the electron microscope in the hope to enlarge the domains and to induce some atom diffusion to decrease the number of point defects. Unfortunately, in this respect the nanoscale film behaves in a similar way as the bulk material, i.e. it follows the bulk phase diagram. As a result, the film starts to decompose into B2 and L1₂ grains at 750 °C, the latter maintaining a similar epitaxial relationship over large distances. As a result the remaining B2 is further depleted of Ni and its $M_S$ drops even further. In fig. 7 the result of such a heating experiment is shown, here after reaching 1000 °C. At higher temperatures the film starts to decompose. From the SAED pattern in fig. 7 it
is evident that both structures exist simultaneously and that the L12 grains indeed exhibit the cubic orientation as well yielding a (001)//(001),[100]//[110] relation with the B2.

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References


