Change in Crystal Orientations of a {100} (001) Pure Aluminum Single Crystal during Accumulative Roll Bonding*

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A {100} (001) pure aluminum single crystal specimen was deformed by accumulative roll bonding (ARB) up to 9 cycles ($\varepsilon_{eq} = 7.18$) with lubrication. The 5- and 9-cycle processes developed weakened textures composed of {123} (634) and {112} (111). The {100} (001) areas existed even after the 9th cycle. The area fraction of {100} (001) was almost the same as that of the 2-cycle processed specimen. The average area fraction of {100} (001) was 4.8%. The cross slips by primary slip pairs may play an important role in maintaining the {100} (001) area. In the 5-cycle processed specimen, {123} (634) bands were observed in not only the center layer but also the one-eighth thickness layers close to the specimen surfaces. After the 7th and 9th cycles, {112} (111) was observed to form by further crystal rotation from {123} (634). [doi:10.2320/matertrans.L-MZ201109]

(Received September 27, 2010; Accepted February 3, 2011; Published April 13, 2011)

Keywords: sever plastic deformation, aluminum, single crystal, rolling, local texture

1. Introduction

Ultra grain refinement is achieved by severe plastic deformation techniques such as HTP (High Pressure Torsion),^{1–3)} ECAP (Equal-Channel Angular Pressing),^{1,4)} CEC (Cyclic Extrusion and Compression),^{5,6)} and ARB.^{7,8)} Among these processes, ARB has the advantage that it can be used to produce large bulky materials. The development of microstructures^{9–11)} and the characteristic of mechanical properties using the ARB process have been studied.^{12,13)} Ultra fine grain structures are generally considered to form by accumulation of abundant strain imposed into the material.^{14,15)}

In FCC metals such as aluminum, the crystal orientation {100} (001) is of great interest in texture control.¹⁶ When a heavy cold-rolled aluminum specimen is annealed under certain heat treatment conditions, the recrystallization texture contains a main {100} (001) component. One of the origins of {100} (001) recrystallization nuclei is considered to be the {100} (001) band existing in the deformed structure, called "the cube band", which is arrayed between deformation regions with major rolling texture components such as {123} (634) and {112} (111).¹⁷ It was clarified that the stability of {100} (001) area during rolling was enhanced by the constraint of grain boundaries,¹⁸ and the preferred nucleation of {100} (001) recrystallized grain was dependent on both orientation gradient and stored energy.¹⁹

One of the present authors investigated the evolution of texture and microstructure in cold-rolled $\{100\}$ $\langle 001 \rangle$ aluminum single crystals.^{20,21)} When the thickness was reduced by more than 50%, a banded structure consisting of two matrices was observed on a mid-thickness plane parallel to the rolling plane. The transition bands between the matrices corresponded to the cube bands. It was empirically demonstrated that the banded structure easily develops when the initial orientation of the single crystal deviates by 5° from

the ideal $\{100\}$ $\langle 001 \rangle$ orientation. We have further interest in the stability of cube bands during severe plastic deformation more than 99% thickness reduction.

In the former paper, {100} (001) aluminum single crystals were heavily deformed using ARB to clarify whether {100} (001) deformed areas can survive upon severe plastic deformation. The change in rolling texture components was also clarified.²²⁾ In this paper, further results were added, and deeper discussions were carried out, especially in order to discuss the distribution of deformed areas with β -fiber texture components and {100} (001) through the specimen thickness.

2. Experimental Procedure

A {100} $\langle 001 \rangle$ single crystal plate was made from a 99.99% aluminum ingot using the modified Bridgeman method. Specimens for rolling had dimensions of 60 mm × 16 mm × 4 mm; they were cut from the plate by an electric discharge machine. The surfaces of the specimens were mechanically and electrolytically polished.

The single crystal specimens had an initial orientation that deviated by 1.5° from the ideal {100} (001). A schematic illustration depicting primary slip systems on eight {111} slip planes is shown in Fig. 1. The designation for slip systems follows the notation proposed by Bishop and Hill,²³ where alphabets a-d and numbers 1-6 indicates {111} slip planes and (110) slip directions, respectively. Assuming that a single crystal during rolling is deformed in compression along the normal direction (*ND*) and in tension along the rolling direction (*RD*), slip system pairs of a2–d2 and b2–c2 become the primary slip systems. Here, these slip system pairs have a cross-slip relationship to each other.

During the ARB process, the single crystal specimens were deformed by a two-high rolling mill with rolls of 310 mm diameter at room temperature with lubrication. The principle of the ARB process has previously been described in detail.⁷ The thickness reduction per cycle was 50% except for the

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Fig. 1 Schematic illustration depicting primary slip systems on eight {111} slip planes.

Table 1 The thickness reduction, equivalent strain and total thickness of specimen in each cycle. At the 1st and 2nd cycles, conventional rolling was conducted. Roll-bonding was carried out after the 3rd cycle.

Cycle	1	2	3	4	5	7	9
Thickness reduction	0.5	0.75	0.875	0.9375	0.9688	0.9922	0.998
Equivalent strain	0.8	1.6	2.4	3.2	4	5.6	7.18
Total thickness (mm)	2	1	1	1	1	1	1

1st cycle, where the 4 mm thick specimen was rolled down to 2 mm in two passes. The 1st and 2nd cycles did not use roll-bonding but conventional rolling. Cutting, stacking and roll-bonding were conducted after the 3rd cycle. Just after roll-bonding, the rolled specimen was immediately cooled into water. By repeating the ARB process, the 9-cycle processed specimen was obtained. Table 1 shows the thickness reduction and equivalent strain during the ARB process. The thickness reduction and equivalent strain imposed on a specimen after the 9th cycle were r = 99.8% and $\varepsilon_{eq} = 7.18$, respectively.

The crystal orientations through the thickness of the processed specimens were measured by a field emission type scanning electron microscope (FEI XL30S) equipped with an EBSD system. The observation plane was normal to the transverse direction (TD plane). The scan step for crystal orientation measurement was carried out at 1 µm. In the 9-cycle processed specimen, the crystal orientations were measured at 0.5 µm steps for more precise orientation analysis. The crystal orientations were analyzed by commercial software (TSL-OIM analysis). The high angle boundary was defined as that having misorientation larger than 15°. Crystal orientations of $\{100\}$ (001), $\{102\}$ (201), and $\{100\}$ (011) that had a tolerance within 15° from the ideal orientation were represented as the crystal orientation components. In this study, the tolerance value for classifying the crystal orientations was 10° for $\{110\}$ $\langle 112 \rangle$, $\{123\}$ (634), and $\{112\}$ (111) to avoid overlapping representation of crystal orientations.

3. Experimental Results

Figure 2 shows the {111} pole figures constructed from the EBSD data of the 1-, 5-, and 9-cycle processed speci-



Fig. 2 {111} pole figures of the 1-, 5-, and 9-cycle processed specimens.



Fig. 3 Change in area fractions of $\{100\}$ (001), $\{102\}$ (201), $\{100\}$ (011) and $\{110\}$ (011) with increasing the number of cycles. The tolerance value for classifying the crystal orientation was 15° .

mens. These textures reveal the crystal orientations viewed from the *ND* plane. In 1-cycle processed specimens, the main orientation rotated about the *TD* with respect to the initial orientation. The 5- and 9-cycle processes developed weakened textures composed of {123} $\langle 634 \rangle$ and {112} $\langle 111 \rangle$. In the 9th cycle pole figure, crystal rotation about *TD* axis was observed relative to the texture components of the 5-cycle processed specimen.

Figure 3 shows the change in area fractions of {100} $\langle 001 \rangle$, {102} $\langle 201 \rangle$, {100} $\langle 011 \rangle$ and {110} $\langle 001 \rangle$ with increasing the number of ARB cycles. The tolerance value used for classifying the crystal orientation was 15°. The area fraction of {100} $\langle 001 \rangle$ was 82.3% after the 1st cycle. It abruptly decreased to 6.9% after the 2nd cycle. Furthermore, the ARB cycles produced little significant change in the area fraction of {100} $\langle 001 \rangle$. After the 9th cycle the area fraction of {100} $\langle 001 \rangle$ was 5.8%. The area fraction of {102} $\langle 201 \rangle$ reached its maximum in the specimen after the 2nd cycle and decreased with further ARB processing. The {100} $\langle 001 \rangle$ and {110} $\langle 001 \rangle$ orientations did not develop in the {100} $\langle 001 \rangle$ aluminum single crystal deformed by the ARB process under lubricated conditions.



Fig. 4 Change in area fractions of $\{110\} \langle 112 \rangle$, $\{123\} \langle 634 \rangle$, and $\{112\} \langle 111 \rangle$ with increasing the number of cycles. The tolerance value for classifying the crystal orientation was 10° .

Figure 4 shows the area fractions of $\{110\}$ $\langle 112 \rangle$, $\{123\}$ $\langle 634 \rangle$, and $\{112\}$ $\langle 111 \rangle$ within 10° from the ideal orientations. These crystal orientations are typical texture components in rolled FCC metals, which could be distinguished after the 3rd cycle. In the 5-cycle processed specimen, $\{123\}$ $\langle 634 \rangle$ had the maximum area fraction of 14.2%. The area fraction of $\{112\}$ $\langle 111 \rangle$ increased up to 6.9% after the 7th cycle, while the area fraction of $\{123\}$ $\langle 634 \rangle$ was reduced to 8.3%. The area fraction of $\{112\}$ $\langle 111 \rangle$ in the 9-cycle processed specimen was 7.2%, which was second next to that of $\{123\}$ $\langle 634 \rangle$.

Figure 5 shows a crystal orientation map for each processed specimen. This map depicts $\{100\}$ $\langle 001 \rangle$, $\{102\}$ (201), and $\{100\}$ (011) within 15° from the ideal orientations and $\{110\}$ $\langle 112 \rangle$, $\{123\}$ $\langle 634 \rangle$, and $\{112\}$ $\langle 111 \rangle$ within 10° . In the 2-cycle processed specimen, the matrix bands of M1 and M4 had $\{102\}$ $\langle 201 \rangle$, which were obtained by crystal rotation about the TD axis with respect to the initial orientation. The macroscopic subdivision of the single crystal and the crystal orientations in four matrix bands were described in detail in the former paper.²²⁾ The transition bands between the four matrix bands were colored by red, indicating $\{100\}$ (001). The 3-cycle process developed thick and thin bands consisting of $\{100\}$ $\langle 001 \rangle$ and $\{102\}$ $\langle 201 \rangle$. The $\{123\}$ (634) areas started to form after the 3rd cycle. The $\{123\}$ (634) bands became thinner with increasing the number of ARB cycles, and the number of the bands increased. In the 5-cycle processed specimen, the {123} (634) bands were observed in one-eighth thickness layers close to the specimen surfaces.

Figure 6 shows a magnified deformation structure in the 9-cycle processed specimen. The lamellar band structure parallel to the *RD* was composed of $\{100\}$ (001), $\{102\}$ (201), $\{123\}$ (634), and $\{112\}$ (111).

The mean grain size along the *ND* in the 5-, 7- and 9-cycle processed specimens had sizes of 1.60, 1.42, and 0.88 µm, respectively. The grain sizes after 5 and 7 cycles were smaller than a mean grain size of 1.80 µm after the 6th cycle with lubrication in {110} (112) pure aluminum single crystals.²⁴⁾ In ARB experiments using pure aluminum polycrystal, it was reported that mean grain sizes after the 8th ARB cycle ($\varepsilon_{eq} = 6.4$) were about 0.25 µm.^{10,14,15)} The mean grain size in the single crystal specimens was larger than that in the

polycrystal specimens. The difference in grain size could be ascribed to the effect of grain boundaries which exist in polycrystalline material before rolling.

4. Discussion

Ishida *et al.* conducted an ARB experiment using a $\{112\}$ $\langle 110 \rangle$ aluminum single crystal²⁴⁾ and clarified the macroscopic grain subdivision. The present study especially focuses on the change in rolling texture component developed by the ARB process.

It has been previously reported that a weak rolling texture was developed by the ARB process, nevertheless the material was highly strained.⁷⁾ As shown in Fig. 2, the rolling texture obtained by using the single crystals also showed weak rolling textures. The 5- and 9-cycle processed specimens contained $\{123\}$ $\langle 634 \rangle$ and $\{112\}$ $\langle 111 \rangle$, which are typical rolling texture components in rolled FCC metals.

An ARB experiment was carried out by Kamikawa et al. using 99.99% pure aluminum polycrystal deformed by ARB for up to 6 cycles without lubrication.^{25,26)} The grain size along the ND, misorientation of grain boundary, and change in crystal orientation were analyzed on the TD plane by the EBSD method. The local textures between the surface and inner layers were quite different. The inner layer had local textures consisting of $\{110\}$ $\langle 112 \rangle$, $\{123\}$ $\langle 634 \rangle$, and $\{112\}$ (111), which were termed "rolling texture components". In this study, it is called " β -fiber texture components". In contrast, the surface layers which had the guarter-thickness of specimen showed the local textures of $\{100\}$ $\langle 011 \rangle$, $\{111\}$ $\langle 112 \rangle$, and $\langle 111 \rangle \langle 011 \rangle$, called "shear texture components". The crystal orientation distribution through the specimen thickness did not correspond with the accumulation of strain imposed by ARB processes over the 1st to 6th cycles. Kamikawa et al. produced a 7-cycle ARB processed specimen and compared the distribution of the crystal orientation through the specimen thickness with that of the 6-cycle processed specimen. The shear texture components observed in the quarter-thickness layers close to the surfaces of the 6cycle ARB processed specimen were destroyed and rotated into the β -fiber texture components in the inner layer after the 7th ARB cycle. Conversely, the β -fiber texture components observed in the inner layer of the 6-cycle ARB processed specimen were rotated into the shear texture components in the quarter-thickness layers of the 7-cycle ARB processed specimen. This result indicates that the local textures formed through the specimen thickness were easily destroyed by the next ARB process. According to their conclusion, under the unlubricated condition the changes in local textures after every ARB cycle may reduce the intensity of the rolling texture and create the characteristic local texture through the specimen thickness.

As shown in Fig. 4, $\{123\}$ $\langle 634 \rangle$ began to form in the 3-cycle processed specimen. The maximum area fraction was observed in the 5-cycle processed specimen. Figure 5 shows that $\{123\}$ $\langle 634 \rangle$ bands were distributed through the specimen thickness except for regions close to the specimen surfaces. The distribution of $\{123\}$ $\langle 634 \rangle$ observed in this study substantiates the destruction and rotation mechanism of local textures in every ARB cycle proposed by Kamikawa



Fig. 5 Crystal orientation maps in processed specimens. These maps show {100} (001), {102} (201), and {100} (011) within 15° from the ideal orientations and {110} (112), {123} (634), and {112} (111) within 10° from the ideal orientations. The stars indicate one-eighth thickness positions from the surfaces of the 5-cycle processed specimen.



Fig. 6 Magnified deformation structure in the 9-cycle processed specimen.

et al. However, the effect of friction between the specimen and the rolls under lubricated condition in the present study is much smaller than that in the previous study using unlubricated condition.^{25,26)} In this study, the use of lubricant may have allowed the formation of $\{123\}$ (634) bands in the one-eighth thickness layers of the 5-cycle processed specimen.

The 5-cycle processed specimen showed the maximum $\{123\}$ (634) area fraction. However, the area fraction decreased after the 7th cycle. Such a local maximum area fraction of $\{123\}$ (634) at intermediate ARB cycles around $\varepsilon_{eq} = 4$ was also observed in an ARB experiment using commercial-purity aluminum polycrystal.²⁷⁾ In this experi-

ment, the volume fraction of {123} (634) measured by an X-ray diffraction method reached its maximum value at a true strain of 3.5 ($\varepsilon_{eq} = 4$). It subsequently decreased with increasing true strain. Alternatively, the volume fraction of {112} (111) gradually increased. After a true strain of 6 has been reached, the volume fractions of {112} (111) and {123} (634) remained relatively constant. The fraction of {112} (111) was higher than that of {123} (634), but this relation is different from the result of the present our study in terms of a β -fiber texture component with the highest area fraction (see Fig. 4). Except for this difference, the previous study²⁷) and our study indicate that in ARB process {123} (634) is not stable at intermediate cycles around $\varepsilon_{eq} = 4$.

In our study, since the area fraction of $\{123\}$ $\langle 634 \rangle$ decreased after the 7th cycle and conversely the area fraction of $\{112\}$ $\langle 111 \rangle$ increased, $\{112\}$ $\langle 111 \rangle$ was nominated for the crystal orientation resulting from further rotation from {123} (634). In the ARB experiment performed by Heason and Prangnell,²⁷⁾ the polycrystalline specimen was deformed by ARB without lubrication up to a true strain of 10. {100} $\langle 011 \rangle$ orientation produced in the surface layers rotated toward $\{112\}$ $\langle 111 \rangle$, when the $\{100\}$ $\langle 011 \rangle$ surface layer was transferred into the center layer by the next ARB process. The crystal rotation from $\{100\}$ $\langle 011 \rangle$ to $\{112\}$ $\langle 111 \rangle$ was explained by the plane strain compression in the center layer. However, in this study using lubricated condition, {100} (011) could barely be detected during ARB (see Fig. 3). Therefore, it is unlikely that the route in crystal rotation from $\{100\}\ \langle 011 \rangle$ to $\{112\}\ \langle 111 \rangle$ is dominant. We concluded that the {123} (634) orientation observed in the 5-cycle processed specimen rotated toward $\{112\}$ $\langle 111 \rangle$ in the 7- and 9-cycle processed specimens under the lubricated condition.

After the 2nd cycle, the area fraction of $\{100\} \langle 001 \rangle$ showed a constant amount of about 6.9%. $\{100\} \langle 001 \rangle$ areas were observed even in the 9-cycle processed specimen

(r = 99.8% and $\varepsilon_{eq} = 7.18)$, and the area fraction was 5.8%. The average area fraction of $\{100\}$ (001) from the 2nd to 9th cycles was 4.8%. We think that $\{100\}$ (001) areas existing in the deformation structure were subjected to not only grain subdivision but also dynamic recovery; this was because the area fraction of $\{100\}$ (001) was constant over a wide range of the 2nd to 9th cycles. The stability in the area fraction of $\{100\}$ (001) may be ascribed to the special slip geometry. In general, dynamic recovery behavior for FCC metals with high stacking fault energy such as aluminum can be explained by cross slip.²⁸⁾ The cross slip is considered to be a stress relief mechanism in plastic deformation.²⁹⁾ Cross slip easily occurs when a crystal is deformed along $\langle 100 \rangle$, while it rarely occurs when deformed along $(111)^{.30}$ Assuming that a $\{100\}$ (001) single crystal is deformed in tension along the RD and compression along the ND, cross slips occur between two slip system pairs: one pair is a2 and d2, and the other b2 and c2. In highly strained specimens such as the 7- and 9-cycle processed specimens, cross slips occur in abundance in $\{100\}$ (001) areas, keeping the balance of operations between the two slip system pairs. Therefore, $\{100\}$ (001) areas can be maintained even in the 9-cycle processed specimen, which had almost the same area fraction as the 2-cycle processed specimen.

The {100} $\langle 001 \rangle$ areas were partially adjacent to the {123} $\langle 634 \rangle$ or {112} $\langle 111 \rangle$ grains after the 9th cycle, as shown in Fig. 6. The {100} $\langle 001 \rangle$ single crystal deformed by ARB up to 9 cycles with lubrication was shown to produce both the {100} $\langle 001 \rangle$ areas resulting from the cross slips and the {123} $\langle 634 \rangle$ and {112} $\langle 111 \rangle$ grains formed by the crystal rotations.

5. Conclusions

- (1) The weakened texture composed of {123} $\langle 634 \rangle$ and {112} $\langle 111 \rangle$ developed in the 9-cycle processed specimens (r = 99.8% and $\varepsilon_{eq} = 7.18$).
- (2) After 2 cycles, the area fraction of {100} (001) reaches a steady state. The average area fraction over the 2nd to 9th cycles was 4.8% with a tolerance value of 15° in crystal orientation. The {100} (001) areas existed even in the 9-cycle processed specimen because of the dynamic recovery caused by cross slip.
- (3) Under lubricated conditions, the development of {112} (111) after the 7th and 9th cycles was ascribed to further crystal rotation of {123} (634), which achieved the maximum area fraction for the 5-cycle processed specimen. {123} (634) is not stable at intermediate cycles around ε_{eq} = 4 in ARB process.

Acknowledgement

One of the authors (K.K.) acknowledges the financial

support provided by the Light Metal Educational Foundation Inc., and N.T. acknowledges the financial support by the Grant-in-Aid for Scientific Research on Innovative Area, "Bulk Nanostructured Metals", through MEXT, Japan.

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