Quantitative TEM study of the precipitation microstructure in aluminium alloy Al(MgSiCu) 6056 T6
François Delmas, Marie-José Casanove, Philippe Lours, Alain Couret, Armand Coujou

To cite this version:

HAL Id: hal-01716116
https://hal.archives-ouvertes.fr/hal-01716116
Submitted on 4 Dec 2019

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Quantitative TEM study of the precipitation microstructure in aluminium alloy Al(MgSiCu) 6056 T6

F. Delmas a, *, M.J. Casanove a, P. Lours b, A. Couret a, A. Coujou a

a CEMES/CNRS, 29 rue Jeanne Marvig, B.P. 4347, 31055 Toulouse Cedex, France
b CROMEP, Ecole des Mines d’Albi-Carmaux, Campus Jarlard, 81013 Albi Cedex 09, France

Abstract

The precipitate microstructure in the last-generation aluminium alloy 6056 T6 [AlMgSiCu] is investigated using three complementary techniques of transmission electron microscopy (TEM) with a special focus on the density and volume fraction of strengthening particles. High-resolution TEM allows the identification of the precipitates and the measurement of the precipitate sizes to be performed. Conventional TEM is used to evaluate the number of precipitates in the investigated area as well as their distribution in the matrix. In situ TEM straining, via the analysis of the dislocation slip traces, permits to determine precisely the thickness and the volume of the foil in the region where the precipitates are analysed. Taking into account the shape and the dimensions of precipitates with respect to the foil thickness, a novel methodology for measuring the volume density and the volume fraction of precipitates is proposed.

Keywords: Aluminium alloys; Precipitation hardening; Density and volume fraction; In situ straining experiments

1. Introduction

The improvement of the performances of aluminium alloys used in aeronautics, in relation to increasingly demanding specifications, is based on a thorough control of the alloys microstructure. In the case of precipitation hardening alloys, the distribution, the morphology and the crystal structure of the strengthening particles strongly depend on the alloy composition and the thermal treatment. Alloy 6056 T6 (AlMgSi) of the last generation, with optimised composition and thermal treatment [1], shows a good compromise in term of mechanical strength, weldability and corrosion resistance. As a consequence, this alloy is candidate to substitute for standard 2024 alloy (AlCu) for the manufacturing of the fuselage skin of the forthcoming A380 Airbus.

In order to better understand the interaction between the precipitates and the dislocations responsible for the mechanical properties of the material, it is important to determine parameters such as the precipitate crystal structure and chemical composition as well as the precipitate morphologies. It is generally admitted that tempering results in the precipitation of intermetallic ternary or quaternary nanometric second phase particles [2]. However, the strengthening microstructure of AlMgSiCu alloys is still not well established crystallographically and remains matter of controversy, see for instance [3–6]. This is partly due to the difficulties inherent to the study of the structure of such small particles. In the case of 6056 T6* alloy corresponding to the first generation with slightly different composition and tempering condition than the alloy under investigation in this paper, SANS and transmission electron microscopy (TEM) studies report the precipitation of lath and needle shaped particles lying along ⟨1 0 0⟩ directions of the aluminium matrix [2,7].

In this paper, focus is placed on the quantitative characterisation of the morphological parameters such as the precipitate mean size and the density and volume fraction of the finely distributed precipitated phase. For this, an original methodology based on various and complementary TEM techniques is developed. High resolution transmission electron microscopy (HRTEM) is used to determine the
2. Experimental details

Alloy 6056 T6 is produced by Pechiney Centre de Recherche de Voreppe in the shape of polycrystalline 6 mm thick sheets. The composition (wt.%) of the material is given in Table 1. First part of the alloy treatment consists in solutionizing at 550 °C, water quenching, and finally a tensile plastic stretching (2%). Strengthening of the alloy is then promoted by T6 temper consisting in maintaining the alloy at 190 °C during 4 h. The resultant average grain size is 30 μm.

Because of the limited achievable tilting angles of the sample holder, the analysis of precipitates by HRTEM requires the orientation of the investigated grain to be close to a (1 0 0) crystal axis. To address the statistical distribution of grains and aid for choosing locations for specimen extraction, the crystal texture of specimens is investigated using X-ray diffraction (XRD). XRD experiments reveal that the normal of the sheet shows a strong {2 0 0} preferential orientation in the region close to the outer surface [8]. This texture gradually changes to {2 2 0} within the bulk of the sheet. Consequently, specimens are preferentially taken from regions close to the surface.

All MET and HRTEM photos, presented in this paper, were digitalised with Photoshop 6 software. Obtained images have a mean resolution of 3 pixels/nm.

Table 1

| Chemical composition (in wt.%) of aluminium alloy 6056 |
|-------------|---------|
| Mg          | 0.75    |
| Si          | 0.93    |
| Cu          | 0.60    |
| Mn          | 0.63    |
| Fe          | 0.10    |
| Zn          | 0.15    |
| Zr          | 0.03    |
| Cr          | 0.03    |
| Ti          | 0.04    |

3. Characterisation of the precipitation

3.1. Typology of precipitates

The [0 1 0] zone axis HRTEM image in Fig. 1 shows the precipitation microstructure in alloy 6056 T6. Only one type of precipitates is observed: they are elongated laths with long axis lying along the {1 0 0} directions of the aluminium matrix. The precipitates are viewed “side-on” (respectively “edge-on”) when their long axis is parallel to either [1 0 0] or [0 0 1] (respectively, parallel to [0 1 0]). Note the presence of side-on precipitates (for instance C1, C2, C3) and edge-on precipitates (for instance D1, D2, D3) in Fig. 1.

The transverse cross-sections of edge-on precipitates are shown in Fig. 2. Cross-section of the laths D1 and D2 are closely rectangular with long edge perfectly straight and aligned in the [0 0 1] direction showing clear crystal pattern (Fig. 2(a)). Cross-sections of D4 and D5 are, respectively, 8.3 nm × 2 nm and 7.8 nm × 3 nm. Truncated side-on precipitate C1, with long axis parallel to [0 0 1] and a thickness of 3 nm, is also imaged in Fig. 2(a). In the [0 0 1] crystal direction, the precipitate show high coherency with the aluminium matrix.

In Fig. 2(b), the precipitates cross-sections show much more irregular shape (D3, D4, D5) and do not exhibit clear crystallographic pattern. This second type of precipitates represents only about 5% of the whole population and their contribution for the measurement of volume fraction can be neglected.

3.2. Spatial distribution

Fig. 3 is a [1 0 0] dark-field image done by selecting, in the diffraction pattern, the superstructure diffuse lines due to the precipitated phase [2,7,9]. The low magnification of the image allows to investigate a larger volume of material and to confirm the presence of both edge-on and side-on precipitates. The distribution of precipitates is quite homogeneous except in areas marked with dashed rectangles where there are many strings of precipitates. They have probably nucleated during heat treatment on residual dislocations. Typically in this material, this kind of heterogeneous precipitation results in some misorientation of the precipitate cross-sections with respect to the (1 0 0) directions of the matrix.

3.3. Precipitate cross-sections

Features of HRTEM images such as Figs. 1 and 2 are processed using Optimas software to derive the dimensional characteristics of each precipitate. Cross-sections s, major axis b and minor axis a (a < b) are measured for 300 individual edge-on particles either resulting from homogeneous or heterogeneous precipitation (values corresponding to heterogeneous precipitation are given in brackets). Statistical data, derived from the distribution histograms of those parameters, are given in Table 2. Note that the mean values...
of $a$, $b$ and $s$ for both distributions of precipitates are very close. As a consequence, the two types of precipitation will not be differentiated in the following. The variation of $a$ versus $b$ plotted in Fig. 4(a) shows that there is no significant correlation between the two parameters indicating that the growth mechanisms in the two directions are not interdependent. This is confirmed by the widely spread histogram of the distribution of $a/b$ values in Fig. 4(b): the mean value is 0.5 and the standard variance is of the order of magnitude of the spectral width.

3.4. Side-on precipitates

The observation of side-on precipitates by HRTEM images such as Fig. 1 allows the measurement of the precipitate lengths $L$ and widths $l$. The distribution histograms of the lengths, ranging from 10 to 193 nm with a mean value of 63 nm, and the width of the side-on precipitates are, respectively, given in Fig. 5(a) and (b).

<table>
<thead>
<tr>
<th>Table 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cross-sections, lengths of the major and minor axis of the precipitates seen edge on (homogeneous precipitation)</td>
</tr>
<tr>
<td>Mean value</td>
</tr>
<tr>
<td>-----------------</td>
</tr>
<tr>
<td>Cross-section $s$ (nm$^2$)</td>
</tr>
<tr>
<td>Length of the major axis $b$ (nm)</td>
</tr>
<tr>
<td>Length of the minor axis $a$ (nm)</td>
</tr>
</tbody>
</table>

Values between brackets correspond to the precipitate strings (heterogeneous precipitation).

4. Volume density and volume fraction of precipitates

4.1. Measurement of the foil thickness

The absence of through-thickness defects such as twin boundaries or stacking fault ribbons makes the foil thickness measurement by conventional TEM very difficult. In addition, usual techniques such as convergent beam microscopy or Electron Energy Loss Spectroscopy failed to give reliable results. The first technique is not useful in this type of alloy because of the too high density of precipitates which prevents measurement of foil thickness in the matrix and the second gives only a mean value of the thickness for a too large region of the foil [8].

So, an original method based on the TEM in situ straining technique, has been developed. Samples are strained in the MET to cause dislocation motions. The occurrence of slip traces at the surfaces of the specimen induced by the dislocation motions allows the accurate calculation of the
foil thickness. Indeed, by measuring the apparent distance between two slip traces on the images and taking into account imaging conditions, thickness of the foil can be deduced using simple trigonometric relationships as illustrated in Fig. 6. The relation between the apparent distance between slip traces and the thickness is given by the following relation:

\[ e = d_{\text{app}} \frac{\sin(\alpha)}{\cos(\alpha - \beta)} \]

where \( \alpha \) represents the angle between the slip plane and the sample plane and \( \beta \) is the tilting angle of the sample in the MET. A TEM image of an 6056 T6 sample is presented in Fig. 7, showing an example of slip traces where a thickness measurement has been done.

The major interest of this method is that the [100] zone axis TEM images are taken for precipitates precisely located in the region where the thickness is accurately calculated. In the present paper, all measurements, precipitate counting and calculations are performed in a 50 nm thick region.

4.2. Visibility of side-on precipitates

The estimation of the volume density of precipitates requires to count the number of precipitates accurately. However, in all [010] zone axis images analysed, the number of side-on precipitates is well lower than the number of edge-on precipitates. Indeed, for a given orientation of side-on precipitates, either lying along [100] or [001], the ratio between the number of side-on and edge-on precipitates \( n_s/n_e \) is always equal to 0.08. As the probability of nucleation and early growth of precipitates is the same in the three \( \langle 100 \rangle \) directions, the ratio should be 1. This result indicates that not all side-on precipitates are visible in the TEM images. The discrepancy between the experimental value of \( n_s/n_e \) and 1 may be understood considering visibility criteria of precipitates. The visibility is directly related to the dimensions of the precipitates with respect to the foil thickness through two distinct parameters:

- The precipitate thickness \( \Delta e \) crossed by the electrons which must be high enough relatively to the foil thickness.
to result in significant beam diffraction and consequently to an image formation (relative thickness).

- The volume $V'$ that must be considered to count the edge-on precipitates (volume of counting) which is greater than the foil volume truly observed in TEM.

Fig. 8 is the histogram of the minor axis of the cross-sections of edge-on precipitates derived from the parameters measured and presented in Section 3.4. This distribution compared to that of the widths of side-on precipitates (Fig. 5(b)) allow the determination of the minimum relative thickness of a precipitate to be visible.

Fig. 9 shows the two possible orientations of the cross-section $(a \times b)$ of side-on precipitates with either major (configuration 1) or minor (configuration 2) axis parallel to the electron beam. These two configurations are equi-probable and consequently, the width distribution of side-on precipitates should correspond, in term of mean values, extreme values and variance to the distribution of cross-section axis plotted without distinguishing $a$ and $b$. This former distribution, which corresponds to the sum of lines 2 and 3 of Table 2, ranges from 1 to 17 nm with a mean value of 5 nm very different from the statistical parameters of the width distribution (Fig. 5b). On the contrary, the distribution of width $l$ of side-on precipitates (Fig. 5b) is statistically very close to the single distribution of the minor axis $a$ of the edge-on precipitates (Fig. 8). This suggests that almost all side-on precipitates currently imaged in the TEM micrographs are in configuration 1, configuration 2 being invisible.

As a consequence, the histogram of side-on precipitates width allows the determination of a visibility criterion. Indeed, only precipitates in configuration 1 with their cross-section major axis $b$ greater than $\Delta e = 7.1$ nm, corresponding to the highest value of $a$ measured, are visible (Table 2). For the $e = 50$ nm thick investigated region, this visibility criterion corresponds to a minimum relative precipitate thickness $\Delta e/e$ of 15%.

Over the whole measured edge-on precipitates, only 40% have a cross-section major axis greater than 7.5 nm. In addition, configuration 1 being equi-probable to configuration 2, only 20% of precipitates are visible side-on for a given (1 0 0) orientation of precipitates.

Fig. 10 presents various situations for edge-on precipitates within a foil with surface $S$ and thickness $e$, and corresponding visibility criterion $\Delta e$. Precipitates, assumed to have all the same length $L$, are located in the foil using their centre of gravity. Precipitates A and B with length included in the foil greater than $\Delta e$ are both visible in TEM though B has its centre of gravity out of the foil. Precipitates C, intercepting the foil over a length smaller than $\Delta e$ is invisible as well as D corresponding to the limit case where the particle loses contact with the foil. According to these considerations, visible edge-on precipitates are included in a volume $V' = Se'$ where $e' = e + 2(L/2) - 2\Delta e$ greater than the volume truly observed $V = Se$. With the experi-
mental values $e = 50$ nm, $\Delta e = 0.15e$ and $L = 63$ nm, one obtains $e' = 98$ nm. The volume of counting for edge-on precipitates $V' = Se'$ is about two times larger than the volume of counting $V = Se$ for side-on precipitates resulting in a number of side-on precipitates two times lower.

The combination of the effect of $\Delta e$ and $V'$ shows that the visible side-on precipitates of a given $\langle 100 \rangle$ orientation represent only 10% of the total number of edge-on precipitates in fairly good agreement with the experimental value of 8%.

4.3. Volume density

The volume density is estimated from TEM micrographs such as Fig. 3. The volume of counting $V'$ is derived from the measurement of the foil thickness and the dimensions of the precipitates as indicated in the previous section. In Fig. 3, both edge-on and side-on precipitates lying along the $\langle 100 \rangle$ directions of the matrix are present. According to the definition of the volume of counting, only the edge-on precipitates must be considered to calculate the volume density. Assuming an isotropic distribution of particles along the three $\langle 011 \rangle$ directions, the total number of precipitates $N$ is consequently three times the number of edge-on precipitates. The volume density of precipitates $N/V'$, calculated from several TEM images, is about $2.8 \times 10^{16}$ cm$^{-3}$.

Fig. 6. Schematic showing the methodology of samples thickness measurement.
4.4. Volume fraction

The calculation of the volume fraction of precipitates requires the knowledge of the volume of precipitates. Unfortunately, TEM does not permit to image simultaneously the section and the length of a precipitate. However, those techniques allow the determination of the histograms of sections of edge-on precipitates and lengths and width of side-on precipitates independently and with a good accuracy. From these data, statistical approaches can serve to the calculation of the volume fraction.

A first method (method 1) consists in estimating a mean volume using the mean values of the section $\bar{S}$ and the length $\bar{L}$

$$\hat{V}_1 = \bar{S}\bar{L} = \frac{1}{n_{SL}} \sum_{n=1}^{n_S} S_n \sum_{n=1}^{n_L} L_n$$

with $n_S$ and $n_L$ the number of precipitates for which the section or the length has been measured, respectively.

A second way to calculate the volume fraction is the statistical “boustrap” method [10] (method 2). Amongst the experimental values reported in the histograms, a section
$S_i$ and a length $L_i$ are selected at random with a probability dependent on the shape of the histogram. However, this method does not take into account the relationship between the length $a$ of the minor axis of the section and the length $L$ of the precipitates. As discussed in Section 4.2, the width of side-on precipitates $l$ corresponds to the minor axis of edge-on precipitates $a$. Consequently, the histogram of $a/L$ may be plotted from the measurement of the widths and lengths of side-on precipitates (Fig. 11). The histogram shows that $a$ is between 1 and 11% of the length $L$. As a consequence, the method has been refined to force the computer program used for the calculation to select only couples $(S_i, L_i)$ that do occur experimentally. For a given $S_i$, associated to a certain $a_i$, a value $L_{id}$ is derived at random from the histogram of the aspect ratio $a/L$. The mean volume obtained by this method is defined by:

$$\bar{V}_2 = \frac{1}{N} \sum_{j=1}^{N} V_j \quad \text{with} \quad \bar{V}_j = \frac{1}{n} \sum_{i=1}^{n} S_i L_{id}$$

$V_j$ is the mean volume for $n$ couples $(S_i, L_i)$ of a given draw and $V_2$ is the mean values for all $N$ draws. The calculation was performed for $n = 200$. For the bootstrap method, a satisfactory convergence is obtained for $N > 5000$. Results of mean volumes ($\bar{V}_1$ and $\bar{V}_2$) and volume fractions $F_v$ are given in Table 3.

The two methods lead to very similar results. However, method 2 is beneficial as it allows the calculation of the volume of each precipitate, data that cannot be straightforwardly measured experimentally.

Concerning the quantitative analysis of microstructural precipitation, similar methods have been developed for the case of Nickel-based superalloys containing spherical precipitates [11–13]. In this former case, particular attention was paid to take into account both the precipitate diameters and the thickness of the thin foil. In the present work, improved measurement of the foil thickness using data derived from in situ straining experiments, more accurate at least for the case of aluminium alloys, has been carried out. Additional difficulties have raised from the lath shape of precipitates, whose length is of the order of magnitude of the thin foil thickness. This lath shape indeed yields to the impossibility of measuring all the precipitate dimensions by one technique alone applied to a single precipitate. Consequently, precipitates cross-sections and lengths are, respectively, measured on HRTEM micrographs and dark field images. Introducing statistical methods has cleared up this first difficulty with the aim to have an estimation of the precipitate volumes. New corrective methods have been subsequently introduced to take into account that the precipitate length is of the order of magnitude of the foil thickness.

The density and volume fraction of precipitates is finally calculated, taking into account their morphology and visibility, which leads to more accurate results than those previously reported in the literature [2,7]. The experimental evaluation of the above parameters still suffers some uncertainties. The uncertainty on the measurement of the foil thickness, about 1%, is majored to 5% because of the thickness fluctuation in a given investigated zone. In any case, it remains lower than 10% estimated for EELS measurements by Myhr et al. [14]. The absolute uncertainty on the dimension of a precipitate imaged in HRTEM is of the order of magnitude of the inter-atomic distance. Added to the effect of the binarization of the TEM images (3 pixels/nm), this leads to a relative uncertainty of 6 and 2%, respectively, for the widths and the lengths. Finally, the exhaustive counting of precipitates is also subject to some approximation, estimated to 8%, due to the difficulty to image and thus to

**Table 3**

<table>
<thead>
<tr>
<th>Method</th>
<th>Mean volume</th>
<th>$F_v$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$\bar{V}_1 = 1323 \text{ nm}^3$</td>
<td>3.7</td>
</tr>
<tr>
<td>2</td>
<td>$\bar{V}_2 = 1423 \text{ nm}^3$</td>
<td>4.0</td>
</tr>
</tbody>
</table>

5. Discussion

In the present paper, the precipitation microstructure in aluminium alloy 6056 T6 of the last generation has been studied. In particular, a new method for the quantitative analysis of the precipitation microstructure based on the combined use of three electron microscopy techniques providing complementary information, has been developed. In the first section of this part, the method of precipitates counting will be discussed with a special attention to its originalities and to the associated experimental uncertainties. In the two following sections, the question of the influence of thermal treatment on the precipitation and that of the correlation between microstructural precipitation and dislocation behaviour will be successively addressed.

5.1. Method of precipitates counting

The determination of the correlation between microstructural precipitation and dislocation behaviour will be successively addressed.

**Fig. 11. Histogram of $a/L$.**
number the most little edge-on particles. All together, the relative uncertainty on the volume density is about 15%.

5.2. Influence of thermal treatments on the microstructural precipitation

According to literature, the morphology of precipitates obviously depends on their structure [4]: needle or rod shaped particles correspond to a metastable ternary phase $\beta$' or $\beta''$ while laths correspond to the quaternary phase Q' (AlMgSiCu) precursor of the stable Q phase. Caron [15–17] reports the presence of laths with either unclear or clear crystal patterns corresponding, respectively, to the hexagonal phases QP and QC, precursor of Q' and stable Q phases. By analogy, the experimental observations in 6056 T6 presented here suggest that the microstructure of precipitation is essentially quaternary and the population of precipitates is composed of QP, QC and Q' particles.

Table 4 summarizes a comparison between the present results and those obtained previously in a 6056 T6 of the first generation [2,7,9]. The two alloys differ just by the temperature and the duration of the ageing, which is longer but performed at a lower temperature in the first generation of alloy. This difference is quantified by the equivalent time $t_{eq}$ introduced by Shercliff and Ashby [18]

$$t_{eq} = t \exp \left( -\frac{Q}{RT} \right)$$

where $t$ is the duration of the ageing, $Q = 130$ mJ/m$^2$ is the self-diffusion energy and $R$ is the ideal gas constant.

In first generation 6056 T6* alloy, it has been observed as well these two types of precipitate with average lengths 60 nm and average cross-sections 7 nm$^2$ [2,7,9]. In the 6056 T6 alloy investigated here, only lath shaped precipitates are observed with similar length but higher cross-sections (20 nm$^2$). This change can be explained by either the non-formation of needle precipitates at 190 °C, or by their dissolution during ageing. According to Cayron’s results in 6061 T6 alloy (175 °C per 8 h) [15–17], showing that no change in precipitate size occurs following 600 h long-term exposure at 175 °C, it is assumed that temperature plays the most important role in precipitate growth.

Concerning precipitate dimensions, the major difference is a strong increase of precipitate cross-section with the increase of the equivalent time whereas the length is not affected. This suggests that the growth processes are not identical along the three (0 0 1) directions of the aluminium matrix, probably because of different interface structures. Consistently with this growth mechanisms, the precipitate density decreases by a factor of 2 as the result of an Ostwald-like ripening process. Note also the increase of the volume fraction from 2.5 to 3.7%, indicating that additional atoms previously in solid solution and/or in Guinier-Preston zones are involved.

Also remarkable in the alloy is the occurrence of heterogeneous precipitation on residual dislocations. The misorientation of those precipitates with respect to {100} matrix directions is probably due to the effect of the strain field of the dislocation during precipitate nucleation. Nevertheless, the presence of dislocations does not infer on the precipitates growth kinetics as particles of the heterogeneous and homogeneous distributions have similar dimensions. The formation of dislocations susceptible to initiate such precipitation is generally achieved by room temperature final rolling of sheets (skin-pass).

5.3. A few consequences of the microstructural precipitation on the dislocation behaviour

In situ straining experiments performed on the same alloys have demonstrated that precipitates are by-passed owing to the activation of cross-slip [15]. For cross-slip, the formation of elongated precipitates will offer most efficient obstacles. By the same experimental approach, it has been shown that dislocations are strongly anchored at pinning lines containing series of precipitates close to each others. This effect is probably correlated to the observation of heterogeneous strings of precipitates [20].

From the observation of moving dislocations under stress, the mean distance $\langle d \rangle$ between pinning points anchoring effectively the dislocations was measured [13–21] in the two T6 alloys described in the previous section. In Table 5, $\langle d \rangle$ is compared with the mean distance $d_{111}$ between intercepts

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Ageing Duration (h)</th>
<th>$t_{eq}$ ($\times 10^{-15}$ h)</th>
<th>Density ($\times 10^{16}$ cm$^{-3}$)</th>
<th>Length (nm)</th>
<th>Shape</th>
<th>Section (nm$^2$)</th>
<th>$F_\alpha$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6056 T6*</td>
<td>8</td>
<td>175</td>
<td>5.5</td>
<td>66</td>
<td>Needles</td>
<td>7</td>
<td>2.5</td>
</tr>
<tr>
<td>6056 T6</td>
<td>4</td>
<td>190</td>
<td>8.5</td>
<td>63</td>
<td>Laths</td>
<td>6.5</td>
<td>3.7</td>
</tr>
</tbody>
</table>

$\langle d \rangle$ is the mean distance between obstacles anchoring really the dislocations, $d_{111}$ is the distance between intercepts of precipitates with a {1 1 1} plane, which has been deduced from the present investigations.

Table 5

Comparison of the distance between precipitates for the 6056 T6* and 6056 T6 alloys

<table>
<thead>
<tr>
<th></th>
<th>$\langle d \rangle$</th>
<th>$d_{111}$</th>
<th>$d_{111}/\langle d \rangle$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T6*</td>
<td>30</td>
<td>15</td>
<td>0.5</td>
</tr>
<tr>
<td>T6</td>
<td>40</td>
<td>30</td>
<td>0.75</td>
</tr>
</tbody>
</table>

of precipitates within \{111\} plane. As these distances are not similar, the dislocations are assumed to be not anchored on all precipitates. Table 5 indicates that the ratio $d_{111}/d$ increases with the equivalent time, namely from T6* alloy to T6 alloy. This ratio can be seen as the fraction of precipitates efficient for anchoring of dislocations. As a conclusion, the increase of equivalent time increases the fraction of efficient precipitates but lowers the number of precipitates anchoring the dislocations, as indicated by their mean distance.

6. Conclusion

A method using three complementary TEM techniques has been developed to address statistically the spatial distribution and dimension of precipitates in aluminium alloy 6056 T6. This approach permits to derive relevant experimental parameters of the global microstructure of precipitation such as the volume density and the volume fraction of precipitated particles. Such characteristics, coupled to observations of straining micro-mechanisms, are useful to describe and understand the mechanical properties of the hardening precipitation alloy. Due to the accuracy of the results obtained, this methodology seems to be a useful tool to study hardening in alloys and is currently used to investigate the influence of the tempering treatment on the microstructure of precipitation as well as deformation mechanisms of the 6056 alloy.

Acknowledgements

The authors want to thank C. Roucau for convergent beam microscopy and Y. Khin for EELS experiments. C. Cayron is acknowledged for fruitful discussions.

References