Evolution of microstructure and elevated-temperature properties during hot rolling and post-rolling annealing process in Al-Mg-Si 6082 alloys

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Abstract. In the present work, the hot rolling at 400 °C and post-rolling annealing at 500 °C were applied on heat-treated Al-Mg-Si 6082 alloys with different Mn contents to study the evolution of microstructure and elevated-temperature properties. During the pre-heat treatment before rolling (400 °C/2h), a number of fine rod-like dispersoids formed in Mn-containing alloy while only high volume of larger Mg2Si particles was observed in the matrix of base alloy free of Mn. After hot rolling, the morphology of dispersoids was transferred from rod-like to spherical with finer size and increased number density while the dispersoids gradually coarsened during post-rolling annealing in Mn-containing alloy. The full recrystallization was completed after 1-2 h during annealing in the base alloy, while only partial recrystallization was observed in Mn-containing alloy. The micro-hardness at room temperature and the tensile yield strength at 300 °C firstly increased from as-rolled condition to the initial stage of annealing (1 h) for both alloys, which was likely attributed to the dissolution of Mg2Si during the beginning of annealing. With further increasing annealing time (2-8 h), both the microhardness at room temperature and the elevated-temperature strengths of the base alloy remained similar, while they were slightly decreased in Mn-containing alloy owing to the partially recrystallization and coarsening of dispersoids. However, the elevated-temperature strengths were always higher in Mn-containing alloy than the base alloy while their differences between two alloys was reducing with prolonging the post-rolling annealing time. The tensile fracture surface was observed to be ductile for all the conditions of both alloys but the dimples in Mn-containing alloy were finer and much more uniformly distributed.

Keywords: Al-Mg-Si 6082 alloy; hot rolling; post-rolling annealing; dispersoids, elevated-temperature properties

1. Introduction

In recent years, the increasing demands in the legislation about the exhaust emissions place higher requirement on the elevated-temperature (250-350 °C) performance of materials in automobile industrials, in which the Al-Mg-Si 6xxx aluminum alloys are widely used. As well-known, the mechanical properties of precipitation-strengthening 6xxx aluminum alloys will sharply drop at elevated temperature (e.g. 300 °C) under traditional heat treatment (solution treatment followed by artificial aging treatment) due to the fast-coarsening process of precipitates [1]. Therefore, how to improve the elevated-temperature properties of Al-Mg-Si 6xxx aluminum alloys has been highlighted as one of the industrial key concerns. On the other hand, the introduce of thermal-stable dispersoids during the proper heat treatment is proved to greatly improve the elevated-temperature properties in 3xxx and 4xxx alloys [2-5]. Though the evolution of dispersoids are also investigated in Al-Mg-Si 6xxx alloys [6-8], most of them focused on the influence of dispersoids on the recrystallization behaviors at room temperature and the role of dispersoids on the elevated-temperature mechanical properties of 6xxx aluminum alloys is rarely discussed.

Recently, our study has shown that the formation of Mn-containing dispersoids in Al-Mg-Si 6082 alloys is of greatly benefit for the enhancement of Yield Strength (YS) at 300 °C, which is increasing from 30 MPa in alloy free of dispersoids to 70 MPa in alloy containing dispersoids after treated at 400 °C/2h [9]. Meanwhile, it is also proved that the formation the dispersoids can greatly inhibit the dynamic recrystallization during the hot rolling process [9, 10]. However, limited work has been performed on the elevated-temperature properties as well as the evolution of dispersoids during the hot-rolling and the post-rolling annealing process in Al-Mg-Si 6xxx alloys, which is of significance to their industrial applications.

In the present work, the emphasis was put on the evolution of microstructure and YS at 300 °C of Al-Mg-Si 6082 alloy containing various Mn during the hot rolling and post-rolling annealing process with the aid of optical and electron microscopy. Meanwhile, the fracture surface after tensile test at 300 °C was analyzed to
discover the role of dispersoids on the failure mode at elevated temperature.

2. Experimental

In the present work, two Al-Mg-Si 6xxx alloys, which are free of Mn (Alloy A) and containing 0.5 wt.% Mn (Alloy B), were cast (Table 1) and then heat treated at 400°C/2h according to our previous study [9, 11]. The hot-rolling was performed at 400°C with the reduction of 90% (from 30 mm to 3 mm). More details about the hot-rolling process can be found in [9]. After rolling, the sample were further annealed at 500°C for 1, 2, 4 and 8h, which was defined as post-rolling annealing process.

Table 1 Chemical composition of alloys in present work

| Alloy # | Element (wt. %) |
|--------|-----------------|
|        | Mg  | Si  | Mn | Fe  | Al   |
| A      | 0.99 | 0.66 | 0  | 0.26 | Bal. |
| B      | 1.07 | 0.67 | 0.52 | 0.27 | Bal. |

The YS at 300°C under various conditions were measured from the tensile tests using Gleeble thermomechanical testing unit and the average value of 3 samples was calculated. The heating rate was controlled at 2°C/S with a holding time of 3 min at target temperature and the strain rate was set as 0.001 S⁻¹. The Vickers microhardness was measured at room temperature with a load of 10g and a dwelling time of 20s. The indentations were assured on the interdendrite cell to evaluate the effect of precipitation/dispersoids on the properties and the average value was calculated from 15 measurements for each condition.

The evolution of microstructure during the rolling and annealing processes, such as the precipitation of Mg₂Si and dispersoids, were characterized by Optical microscopy (OM), Scanning electron microscopy (SEM) and Transmission electron microscopy (TEM) while the fracture surface after tensile test was observed with SEM. Samples were etched with 0.5% HF for 30s to clearer display the large Mg₂Si precipitates and dispersoids. The grain structure during rolling and annealing process was mapped and analyzed by Electron Back-Scatter Diffraction (EBSD) with a step size of 2 μm.

3. Results and Discussion

3.1 Microstructure before and after rolling

According to our previous study [9, 11], it is proved that the dispersoids have been fully precipitated to the stable condition after 400°C/2h. Therefore, Fig.1 shows the microstructure of experimental alloys after 400°C/2h, which was defined as the pre-treatment before rolling. As shown in Fig. 1a, high volume of large Mg₂Si particles have precipitated in the matrix besides the Al-Fe-Si intermetallics in Alloy A free of Mn (base alloy), which can be due to the relative low heat treatment temperature (400°C).

However, the volume of Mg₂Si precipitates in the matrix was greatly reduced in Alloy B (Fig.1 b) and they were dominantly located at the dispersoids free zone (DFZ, more details about DFZ can be found in [9]) while some smaller Mg₂Si were occasionally observed in the interdendrite cell. On the other hand, high volume of fine dispersoids in dispersoids zone (DZ) was observed in Alloy B. As shown in Fig. 1c, high quantity dispersoids were precipitated with the morphology of cubic and rods, which were reported to be α-Al(MnFe)Si dispersoids [9]. The dispersoids was reported to nucleate on the pre-formed Mg₂Si and then decrease the volume of Mg₂Si during the heat treatment [6, 12], explaining the less Mg₂Si in the interdendrite cell in Alloy B. However, little dispersoids precipitated at the DFZ due to the lower super saturated solution level of Mn [2], therefore, some larger Mg₂Si can still be observed in DFZ (Fig. 1b).

Fig. 1. Microstructure of experimental alloys after 400°C/2h (after etched): (a) Alloy A and (b, c) Alloy B

After pre-heat treatment (400°C/2h), the experimental alloys were hot-rolled at 400°C and the as-rolled microstructure was illustrated in Fig. 2. For Alloy A (Fig. 2a-2b), the Al-Fe-Si intermetallics were fragmented to smaller particles while high volume of Mg₂Si can still be observed (Fig. 2b). As shown in Fig. 1a, the size of Mg₂Si before rolling was so bigger that they were hardly to be dissolved into matrix during the rolling. However, it seems they were also fragmented to smaller size due to the high deformation during the rolling.

Similar to Alloy A, the Al-Fe-Si intermetallics were also fragmented in Alloy B (Fig. 2c). However, much less Mg₂Si can be observed while the dispersoids were still present in the matrix (Fig. 2d). As shown in Fig. 1b, though lower volume of Mg₂Si were remaining in Alloy B after 400°C/2h, their size was much smaller than that in Alloy A (Fig. 1a), leading to it easier for them to dissolve into matrix during the hot-rolling process. Meanwhile, it is interesting to be noted that the dispersoids after hot-rolling (Fig. 2e) is greatly different with that before rolling (Fig. 1c) that the morphology of dispersoids were much more spherical and their size was finer with higher number density after rolling (Fig. 2e). The reason of this evolution of dispersoids is still under
investigation but the favourable conditions promoting precipitation induced by the dislocations or sub-grain boundaries formed during the hot rolling was a plausible cause [13].

Fig. 2. As-rolled microstructure of experimental alloys (after etching): (a-b) Alloy A and (c-d) Alloy B

Fig. 3 displays the as-rolled grain structure of experimental alloys from the EBSD mapping analysis. It can be found that the as-rolled grain in Alloy A is almost equiaxed while some elongated grains in the rolling direction can still be observed (Fig. 3a), indicating that the partial or close-fully recrystallization during the rolling process in Alloy A occurred. This was also confirmed by the scattered and weaker texture from the 

\{111\} pole figure in Fig. 3a. As shown in Fig. 1a and 2b, only coarse Mg2Si particles and fragmented Al-Fe-Si particles present in Alloy A, which have limited retarding effect on recrystallization during the hot-rolling process. Therefore, the recrystallized microstructure (equiaxed grains) was mostly observed in as-rolled condition of Alloy A.

However, the as-rolled microstructure was different in Alloy B (Fig. 3b) that the lamellar structure was dominate with deformed bands and highly elongated grains. Meanwhile, the \{111\} pole figure also showed the strong rolling texture, which was similar to the Brass and S texture [14]. This can be attributed to the precipitation of high volume of dispersoids in Alloy B, which have the strong pinning effect on the recrystallization during the hot-rolling [2, 9].

Fig. 3. As-rolled grain structure and the \{111\} pole figure from EBSD mapping of experimental alloys

3.2 Microstructure during the post-rolling annealing process

As shown in Figs. 2 and 3, different microstructure and grain structure was observed in Alloy A and Alloy B, which will also affect the microstructures during the post-rolling annealing process. Fig. 4 shows the annealing microstructure of experimental alloys after 1h and 8h at 500°C. For Alloy A, it can be seen that only the Al-Fe-Si intermetallics remains in the alloy while almost all the Mg2Si particles present in Fig. 2b has dissolved into matrix after 1 h annealed at 500°C, as shown in Fig. 4a. The microstructure keeps similar even after 500°C/8h (Fig. 4b) that the matrix is clean with only fragmented Al-Fe-Si intermetallics.

Fig. 4. Annealed microstructure of experimental alloys after 1h and 8h at 500°C
Compared with the as-rolled microstructure of Alloy B in Fig. 2d, all the Mg$_2$Si in Alloy B after 500°C/1h has also been fully dissolved into matrix (Fig. 4c), which is similar to Alloy A. This can be attributed to the higher annealing temperature (500°C). Meanwhile, it is observed that the dispersoids coarsened during the annealing process. As shown in Fig. 4d of annealed after 500°C/8h, the color of dispersoids zone was much darker, which was further confirmed by the TEM observation in Fig. 4e and 4f. Compared with the dispersoids in as-rolled condition (Fig. 2e), the dispersoids was gradually coarsened after 500°C/1h (Fig. 4e) and they are even much bigger after 500°C/8h (Fig. 4f). The coarsening process of dispersoids during the annealing process can be related to the formed and stored dislocations or subgrains during the hot-rolling, which was favorite for the nucleation and coarsening of dispersoids [12, 13].

Fig. 5 shows the evolution of grain structure of experimental alloys during the post-rolling annealing process, in which the differences can be observed between two alloys. As shown in Fig. 5a-5d, the equiaxed grains were always present in Alloy A free of Mn and the fully recrystallization was assumed to complete only after 500°C/2h (Fig. 5b), representing as the uniformed distributed equiaxed grains and they were gradually growing with the increase of annealing time, as shown in Fig. 5e-5d.

However, different grain structures from Alloy A were observed in Alloy B containing dispersoids due to the addition of Mn. As shown in Fig. 5e-5g, the similar lamellar structure to the as-rolled condition (Fig. 3b) with elongated grains and deformation bands was dominate until to 500°C/4h (Fig. 5g) thought some fine recrystallized grains can be observed after 500°C/2h (Fig. 5f) and 500°C/4h (Fig. 5g). With prolonged annealing time (500°C/8h in Fig. 5h), more recrystallized grains were observed between the elongated grains, indicating the partial discontinuous recrystallization occurred [15]. However, the volume fraction of recrystallization was still lower, even after 500°C/8h (< 20 vol.%), and the elongated recovery grains were still dominant structure in Alloy B during the post-rolling annealing process, which is greatly different with the fully recrystallized structure in Alloy A.

The different recrystallization behavior between two experimental alloys can be explained by the formation of dispersoids in Alloy B since Mg$_2$Si were already dissolved into matrix in both alloys (Fig. 4). In Alloy A, no precipitation or dispersoids were present (Fig. 4a-b) in the matrix and then the recrystallization was easier to occur and even it has fully completed after 500°C/2h. However, there were high density of fine dispersoids formed after pre-heat treatment (Fig. 1c) and they were even much finer with increased number density after rolling (Fig. 2e), which were reported to strongly inhibit the recrystallization through increasing the Zener drag force due to its finer size and higher number density [4, 16]. Therefore, the recrystallization in Alloy B was greatly retarded (Fig. 4e) while it was gradually happened after 500°C/8h due to the reduced Zener drag force from coarsened of dispersoids (Fig. 4f).

### 3.3 Evolution of elevated-temperature properties during the post-rolling annealing process

As shown in Figs. 4 and 5, it can be observed that the grain structure as well as the formation of precipitates and dispersoids were greatly different between Alloy A and Alloy B after rolling and post-rolling annulling process, which would also lead to the various mechanical properties. In the present work, the tensile tests at 300°C were performed. According to the evolution of microstructure shown in Figs. 4 and 5, the tensile train-stress curves under conditions of as-rolled, 500°C/1h and 500°C/8h were shown in Fig. 6. It can be found the total strain at failure under all these conditions of experimental alloys was in the range 0.3-0.4 (except Alloy A after 500°C/1h), indicating the excellent ductility of experimental 6xxx aluminum alloys at 300°C. Meanwhile, it was also observed that the tensile stress in Alloy B was always higher than that in Alloy A during the tests in all the conditions. As shown in Fig. 6a, the ultimate tensile strength (UTS) at 300°C of Alloy B was measured to be 78 MPa, which was the almost doubled of that in Alloy A (42 MPa). During the post-rolling annealing process, it seemed that the extent of differences on the UTS was decreasing with increasing annealing time. For instance, the differences of UTS between two alloys decreased to 13 MPa after 500°C/1h (Fig. 6b) and further to 7 MPa after 500°C/8h (Fig. 6c).
Furthermore, it was more interesting to notice that the total tensile strain of Alloy B was still higher than Alloy A, especially at as-rolled condition (Fig. 6a) and the beginning of annealing process (Fig. 6b) while the strength of Alloy B was also higher than Alloy A, which was similar to the results in Al-Si 356 alloys that Mo-containing dispersoids improved both the strength and ductility [17].

Fig. 6. Tensile strain-stress curves at 300°C of experimental alloys at: (a) as-rolled; (b) 500°C/1h and (c) 500°C/8h

In order to fully evaluate the evolution of mechanical properties of experimental alloys, Fig. 7 displayed the evolution of microhardness at room temperature (RT) as well as the tensile YS at 300°C experimental alloys during the rolling as well as the post-rolling annealing process. As shown in Fig. 7, the evolution of microhardness at room temperature and the tensile YS at 300°C was generally similar for both experimental alloys that it was firstly increased from as-rolled condition to 500°C/1h and then stabilized (Alloy A) or gradually decreased (Alloy B) with prolonging of the post-rolling annealing time, which was agreed with the tendency shown in Fig. 6. For instance, the microhardness of Alloy A increased from 56 HV at as-rolled condition to 77 HV after 500°C/1h and then reduced to 74-75 after 500°C/8h while the YS at 300°C was initially increased from 35 MPa at as-rolled condition to 56 MPa after 500°C/1h and then stabilized at 56-57 MPa until to 500°C/8h. However, it was observed that both the microhardness at RT and the YS at 300°C in Alloy B were always higher than that in Alloy A, especially at as-rolled condition. As shown in Fig. 7, the as-rolled microhardness at RT and YS at 300°C of Alloy B were 80 HV and 54 MPa, respectively, which were much higher than that of Alloy A (56 HV and 35 MPa). However, the differences of mechanical properties between two experimental alloys were reducing with prolonging the annealing time during the post-rolling annealing process. For instance, the differences of YS at 300°C between Alloy A and Alloy B was decreasing from 19 MPa at as-rolled condition to 11 MPa after 500°C/1h and 7 MPa after 500°C/2-4h and then 5 MPa after 500°C/8h.

Fig. 7. Evolution of mechanical properties of experimentally alloys during the post-rolling annealing process: (a) Microhardness at RT and (b) tensile YS at 300°C
The evolution of mechanical properties was greatly related to the change of microstructures during the post-rolling annealing process. As shown in Fig. 2 and Fig. 4, high volume of Mg$_2$Si was observed to fully dissolve into the matrix after 500°C/1h, which can be the likely reason for the initial increase of microhardness and YS from as-rolled condition to 500°C/1h. However, it can be found that much higher volume of Mg$_2$Si in Alloy A were dissolved than Alloy B after 500°C/1h, explaining the higher increasing degree of mechanical properties of Alloy A at this stage. After 500°C/1-2h, the fully recrystallization was observed in Alloy A followed by a minor growth of grain, leading to the stabilization of microhardness at RT and YS at 300°C while the partially recrystallization and coarsening of dispersoids was occurred in Alloy B with increasing post-rolling annealing time, resulting in its micro reduced mechanical properties during the annealing process. However, the contribution of dispersoids on the mechanical properties can be clearly demonstrated by the difference between both the microhardness at RT and YS at 300°C. For instance, the difference of YS at 300°C at as-rolled condition between Alloy B and Alloy A was 19 MPa due to the high quantity of finer dispersoids as well as the higher Mg and Si solid solution strengthening induced by the formation of dispersoids (Figs. 2e and 4e) and it was still 5 MPa after 500°C/8h though the dispersoids has greatly coarsened (Fig. 4f). It should not be underestimated this “5 MPa” different on the YS at 300°C since every “1 MPa” counts for the mechanical properties at elevated temperature. For instance, it was reported that the creep resistance of materials could be improved one order with increasing of 3 MPa on the threshold stress at 300°C [2, 18].

The fracture surface after tensile test at 300°C of experimental alloys was analyzed using SEM and shown in Fig. 8.

It can be found that the failure mode of both experimental alloys at all conditions was ductile due to the high tensile temperature (300°C), representing by the formation of dimples at the fracture surface. However, the dimples in Alloy B were smaller and more uniform than that in Alloy A, which can be attributed to the homogenous deformation due to the presence of dispersoids [17]. During the tensile test, the effect of dispersoids-dislocation interactions became more remarkable due to the increase of dislocation density [19]. It was reported that dispersoids can change the slip system by means of cross-slip at high temperature when the dislocations were trapped by dispersoids, making the homogenization of slip [20], explaining of much more uniformly distributed dimples in Alloy B. Besides, it was also proposed that the homogenization deformation can be highly promoted by converting the dislocation movement from the planar slip to a wavy slip mode through cross slip [21], which would also benefit for the higher ductility. Therefore, the formation of dispersoids was the likely reason for both higher strength and better ductility in Alloy B.

4. Conclusions

In the present work, the evolution of microstructure and elevated-temperature strength in 6xxx alloys containing various Mn was investigated during hot-rolling and post-rolling annealing processes and the following conclusions can be drawn:

(1). After the pre-heat treatment before rolling (400°C/2h), high volume of larger Mg$_2$Si was precipitated in the base alloy free of Mn while a number of fine rod-like dispersoids were dominate precipitates with lower volume and smaller Mg$_2$Si in Mn-containing alloy.

(2). During the hot rolling, the dispersoids was transfer from rod-like to the spherical and most of smaller Mg$_2$Si has dissolved into matrix in Mn-containing alloy. However, high volume of fragmented larger Mg$_2$Si was still remaining in base alloy free of Mn.

(3). During the post-rolling annealing process at 500°C, fully recrystallization was observed after 1-2h in base alloy free of Mn while only partial recrystallization was observed in Mn-containing alloys, even after 8h.

(4). The elevated-temperature strength and microhardness at room temperature was firstly increase from as-rolled to 500°C/1h followed by a plateau or a gradually reduction in both alloys. However, they were always higher in Mn-containing alloy than the base alloy and the biggest difference was observed at as-rolled condition while the difference was becoming smaller with increasing post-rolling annealing time.

(5). Tensile fracture surface at 300°C was observed to be ductile with the presence of dimples under all the conditions of both experimental alloys, which is
confirmed by the high total strain at failure (~ 0.35). However, the dimples were smaller and more uniformly distributed in Mn-containing alloy, leading to its higher ductility.

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