Influence of Mg/Si Ratio on Nanocluster Formation in Al-Mg-Si Alloys with Constant Mg + Si Concentration

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Two types of nanoclusters are formed during low temperature aging and play significantly important roles in age-hardening of Al-Mg-Si alloys. The formation behavior of nanoclusters markedly depends on the alloy composition. In this paper, alloy specimens with different Mg/Si ratios in the constant Mg + Si concentration of 1.1, 1.3 and 1.5 mol% were used in order to investigate the influence of the Mg/Si ratio on the nanocluster formation using differential scanning calorimetry (DSC), hardness and electrical resistivity measurements. Two overlapped peaks obtained by the DSC analysis are separated using the Gaussian function method and it is clarified that the volume fraction and formation kinetics of nanoclusters change with the Mg/Si ratio even though in the constant Mg + Si concentration. It is found that the most favorable Mg/Si ratio for the nanocluster formation is almost 1.0 even for the different values of the Mg + Si concentration. The influence of the Mg/Si ratio on the nanocluster formation is discussed in terms of interactions among the solute Mg and Si atoms and vacancies in the matrix.

(Received April 16, 2014; Accepted August 6, 2014; Published October 18, 2014)

Keywords: aluminum-magnesium-silicon alloys, nanoclusters, differential scanning calorimetry, Gaussian function method, Mg/Si ratio

1. Introduction

Al-Mg-Si alloys (6xxx series aluminum alloys) have been extensively used for body panels of automobiles due to their attractive properties, such as good formability which is required for forming processes and high corrosion resistance. In addition, these Al-Mg-Si alloys are well known to be extensively used for body panels of automobiles due to their attractive properties, such as good formability which is required for forming processes and high corrosion resistance. In addition, these Al-Mg-Si alloys are well known to be extensively used for body panels of automobiles due to their attractive properties, such as good formability which is required for forming processes and high corrosion resistance.

The chemical compositions (mol%) of the used Al-Mg-Si composition.⁹⁻¹³ Serizawa et al.¹² proposed the characteristics of the two types of nanoclusters, i.e., Cluster (1) and Cluster (2), which have strong roles in the age-hardening. The chemical composition of Cluster (1) formed at around RT is Si-rich at the early stage of NA. Mg atoms are gradually incorporated at the longer NA time to change composition. On the contrary, Cluster (2) approaches a certain Mg/Si ratio with increasing its size, which is very close to that of the β⁺-(Mg,Si) phase.¹⁴ Serizawa et al.¹⁴ also confirmed that the “negative effect of two-step aging” is caused by the formation of Cluster (1), which cannot transform into the β⁺ phase due to the high thermal stability. In addition, Kim et al.¹⁵ explained that the formation of Cluster (2) prior to the formation of Cluster (1) effectively suppresses the negative effect on the BH response. That is, the control of nanoclusters becomes extremely essential in terms of the formation kinetics. It is reported that the Si and Mg concentration of Al-Mg-Si alloys has a marked effect on the formation of nanoclusters.¹⁶⁻¹⁷ Kim et al.¹⁷ especially, revealed that the formation of Cluster (1) is more correlated with the Si concentration than Mg concentration, whereas both the Mg and Si concentrations are important for the formation of Cluster (2). In addition, they clarified that the Mg/Si ratio of the alloy composition is also analyzed.

The aim of this study is, therefore, to investigate the dependence of nanoclusters on the alloy composition and to clarify the most favorable composition for the two types of nanoclusters, Cluster (1) and Cluster (2). The activation energy for the nanocluster formation is also analyzed.

2. Experimental Procedure

The chemical compositions (mol%) of the used Al-Mg-Si
alloys are described in Table 1. The alloys were DC-cast, and then the alloy billets were extruded to 2 mm thickness at 500°C at Sumitomo Light Metal company (now, UACJ). The specimens contain different Mg (0.31 to 0.91 mass %) and Si (0.20 to 1.14 mass %) concentration in order to investigate the influence of the Mg/Si ratio. The letters of M and S in the first column in Table 1 are abbreviation of Mg and Si. The numbers in front of M and S represent mass % of Mg and Si, respectively. Three kinds of specimens with the different Mg/Si ratios as well as constant total solute concentration (Mg + Si) of about 1.1, 1.3 and 1.5 mol % are shown in Fig. 1. The cold-rolled to the thickness of 1.3 mm specimens were solution heat treated at 560°C for 1.8 ks in a salt bath, followed by quenching into ice-water at 0°C for 60 s. These specimens are called as the as-quenched (this is abbreviated to A.Q.) alloy in this paper. Differential scanning calorimetry (DSC), an effective technique to detect nanostructures, was carried out using a Rigaku DSC8230 instrument under a purified argon gas atmosphere after direct quenching. The specimens for the DSC measurement were made of disc shapes with 30 mg. Pure Al with 99.99% purity was used as the reference sample. After putting a specimen into the furnace, the furnace was firstly cooled down to −50°C using a liquid nitrogen controller in order to keep the stable state at the low temperature, and then heated up to 500°C with the different heating rate of 2, 5, 10, 15 and 20°C/min, respectively. After the solid solution heat treatment, alloy specimens were naturally aged at room temperature and aged at 100°C using an oil bath up to 2419.2 ks (1 month) in order to investigate the specific formation behavior of Cluster (1) and Cluster (2), respectively. Micro-Vickers hardness was measured with a load of 200 or 300 g for 15 s using Mitsutoyo HM-102. The specimens were polished within 0.3 ks after each fixed aging time during natural aging and aging at 100°C. Seven indentations were measured from each specimen and cut off the maximum and minimum measured values and plotted on the averaged values. Furthermore, electrical resistivity measurements were also carried out by a four-probe method at −196°C using liquid nitrogen with 120 mA direct current. The specimens were prepared as wires with the diameter of 1.0 mm and the gage length of 300 mm.

3. Results

3.1 Differential scanning calorimetry (DSC)

3.1.1 DSC curves for as-quenched alloys

Figure 2 shows DSC results of the as-quenched alloys with constant Mg + Si concentration about 1.5. Peak I: formation of nanoclusters of Mg and Si, Peak II: dissolution of pre-formed nanoclusters into the matrix, Peak III: precipitation of $\alpha$Al, Peak IV: precipitation of $\beta'$, Peak V: precipitation of $\beta$, Peak VI: U1, U2 and $\alpha$ phase (only in excess Si alloys).
the Peak (1) and Peak (2) indicated in Fig. 3 are caused by
like III, IV and V are attributed to the precipitation of the needle-
by means of DSC and atom probe
Serizawa two types of nanoclusters, i.e., Cluster (1) and Cluster (2), by
After that, these Peak (1) and Peak (2) are designated by
They also proposed that the U1, U2 and B
formations of the nanocluster formation is mainly focused and discussed
0 to 150°C), III (240 to 280°C), IV (280 to 300°C), V (400 to
Another exothermic peak designated by VI (nearby 340°C) in Fig. 2
is observed only for the 3M12S alloy with excess Si. Peak I
detected at a relatively low temperature range represents that
the nanoclusters consisted of Mg, Si atoms and vacancies are
The DSC curves for the nanoclusters formed at low temperature in the as-quenched alloys of (a) Mg + Si ≈ 1.5,
(b) Mg + Si ≈ 1.3 and (c) Mg + Si ≈ 1.1. Exothermic peaks of Peak (1) and Peak (2) are detected in all used alloys.

In this paper, the exothermic peak I corresponding to the nanocluster formation is mainly focused and discussed
because nanoclusters have a significant effect on the later strengthening precipitates of β′.
The DSC curves for the nanoclusters formed between 0 to
150°C are enlarged and shown in Fig. 3 for the alloys with
the constant Mg + Si concentration of (a) 1.5, (b) 1.3 and (c) 1.1. Two overlapped exothermic peaks indicated by Peak (1) and
Peak (2) are observed in all the examined alloys. Peak (1)
and Peak (2) obtained in the present DSC curves were also
detected in the low-temperature adiabatic calorimetry.22)
After that, these Peak (1) and Peak (2) are designated by
two types of nanoclusters, i.e., Cluster (1) and Cluster (2), by
Serizawa et al.11,12) From these viewpoints, it is found that
the Peak (1) and Peak (2) indicated in Fig. 3 are caused by
the formation of Cluster (1) and Cluster (2), having different
characteristics each other.
Comparing with the DSC curves in Fig. 3(a), (b) and (c), it is
found that the variation of heat evolution is different
depending upon the Mg/Si ratio as well as the Mg + Si
concentration. In order to analyze these variations in more
detail, two overlapped peaks are fitted to separate using the
Gaussian function method. The fitting of the DSC curves to
separate into two peaks using the Gaussian function method
is well done as is already reported in our previous paper.17)
The formation behavior of nanoclusters is analyzed using the parameter of the peak area and peak temperature of the
separated Cluster (1) and Cluster (2) peaks. In the present
paper, it is regarded that the peak area directly corresponds
to the volume fraction (Vf) of the formed nanoclusters
because this peak area is caused by the heat evolution of
an exothermic reaction. It is also assumed that the peak
temperature is that at which the formation of nanoclusters is
most accelerated. In other words, the peak temperature shifts
to lower or higher temperature depending on the formation
rate of nanoclusters.

3.1.2 Peak area and peak temperature for nanocluster formation
Figure 4 shows peak areas of Cluster (1) and Cluster (2) in
the alloys with the constant Mg + Si concentration of 1.1, 1.3
and 1.5 mol% as a function of the Mg/Si ratio. These results
are obtained based on the peaks observed in Fig. 3 in a
constant heating rate of 0.167 K/s (10°C/min). It is found
that the peak areas of Cluster (1) and Cluster (2) are gradually
enhanced with the increasing Mg + Si concentration.
Furthermore, this result shows that the maximum value of the
peak area is most enhanced when the Mg/Si ratio is close to
1.0 for Cluster (1) and Cluster (2). That is, the volume
fractions of Cluster (1) and Cluster (2) are increased with the
Mg/Si ratio closer to approximately 1.0. It is clarified that
these characteristics are markedly observed with the higher
Mg + Si concentration.

Fig. 3 DSC results showing the nanoclusters formed at low temperature in the as-quenched alloys of (a) Mg + Si ≈ 1.5,
(b) Mg + Si ≈ 1.3 and (c) Mg + Si ≈ 1.1. Exothermic peaks of Peak (1) and Peak (2) are detected in all used alloys.
In order to investigate the influence of the Mg/Si ratio at different heating rates on the nanocluster formation, the DSC measurement was performed at various heating rates of 2, 5, 10, 15, and 20°C/min. Figure 5 shows the results of the 7M8S alloy for the nanocluster formation with the different heating rates from 2 to 20°C/min. Other alloys with the constant Mg + Si concentration of 1.5 mol% were investigated in the same way as shown in Fig. 5. The peak area and peak temperature of Cluster (1) and Cluster (2) obtained from the DSC curves are represented as shown in Fig. 6 and Fig. 7. From the results of the peak area in Fig. 6, it is noted that the volume fraction of (a) Cluster (1) and (b) Cluster (2) is most increased in the case of the Mg/Si ratio close to 1.0 for the different heating rate. As can be seen in the results in Fig. 7, the peak temperature has a tendency to shift to the lower temperature when the Mg/Si ratio increases. The lowest peak temperature is obtained when the Mg/Si is close to 1.0 for Cluster (1). However, the lowest peak temperature for Cluster (2) is 1.7 of Mg/Si ratio as seen in Fig. 7 (b). It is noted that the tendency is not changed even for the different heating rate. This shift of the peak temperature for the formation of Cluster (1) and Cluster (2) with the different heating rates is confirmed in all the examined alloys, indicating that the formation of Cluster (1) and Cluster (2) is kinetically controlled by the heating rate.

3.1.3 Formation kinetics of Cluster (1) and Cluster (2)

The formation kinetics with the different Mg/Si ratio can be estimated by comparing the activation energy, which is the...
energy barrier required for the formation of nanoclusters. There are several methods to determine the activation energy ($E_a$) for the precipitation process using peak temperature of DSC results.23-25 In this paper the generally used Kissinger method for the experimentally obtained DSC curves is applied to evaluate the activation energy for the nanocluster formation. The Kissinger equation can be written as:

$$\ln\left(\frac{\gamma}{T_p^2}\right) = \frac{-E_a}{R} \left(\frac{1}{T_p}\right) + C$$

(2)

where $\gamma$, $T_p$, $E_a$, $R$ and $C$ represent the heating rate, peak temperature, activation energy, gas constant and an constant, respectively. Figure 8(a) and (b) show the plot of $\ln(\gamma/T_p^2)$ as a function of $1000/T_p$ ($K^{-1}$) obtained by eq. (2) for Cluster (1) and Cluster (2). Based on the slope of the plots (given by the least square fitting), the activation energies ($E_1$ and $E_2$) for Cluster (1) and Cluster (2) with alloy composition were calculated and summarized in Table 2. The determined activation energies for Cluster (1) and Cluster (2) have a different value with the Mg/Si ratio of the alloy composition. Furthermore, the much lower value for Cluster (1) formation is shown in the 7M8S alloy (72.83 kJ/mol), whereas in the 9M6S alloy (77.02 kJ/mol) for the Cluster (2) formation. In other words, the reaction rates of Cluster (1) and Cluster (2) are accelerated when the Mg/Si ratio is close to 1.0 and 1.7, respectively. These results correspond to the results of peak temperature as shown in Fig. 7. The activation energy is most decreased when the peak temperature has minimum value. The 7M8S alloy with the Mg/Si ratio of approximately 1.0 has a largest volume fraction of nanoclusters and simultaneously low activation energy and is found to be most favorable for the formation of Cluster (1) and Cluster (2).

3.2 Hardness and electrical resistivity changes during natural aging

In this section, natural aging was performed to investigate the isothermal aging behavior of Cluster (1) in the alloys with the constant Mg + Si concentration of 1.5 mol%. Figure 9(a) and (b) show the results of electrical resistivity and hardness changes during natural aging for the different Mg/Si ratio...
alloys. In Fig. 9(a), the hardness change ($\Delta HV$) is determined by $HV_{\text{measured}} - HV_{\text{A.Q.}}$, indicating that how much Cluster (1) is formed from the supersaturated solid solution during the natural aging process. The hardness is almost unchanged at the initial stage within 0.6 ks (10 min) and fast hardening up to about 14.4 ks (4 hour), then slow down at longer time. In addition, the influence of the Mg/Si ratio on Cluster (1) is confirmed from the increment of hardness changes. It is found that the 7M8S alloy with the Mg/Si ratio close to 1.0 exhibits the most increased increment of hardness. This result for the Mg/Si ratio is also confirmed by the electrical resistivity changes during natural aging as shown in Fig. 9(b). The electrical resistivity is strongly affected by the amount of solute atoms in the Al matrix and the nanocluster formation. As is seen in Fig. 9(b), the electrical resistivity changes are increased with the natural aging time due to the formation of Cluster (1), the solute atoms of Mg and Si in the Al matrix decrease simultaneously. These results of hardness and electrical resistivity changes are in good agreement with the DSC results shown in Fig. 5 and Fig. 6.

### 3.3 Hardness and electrical resistivity changes during aging at 100°C

In order to investigate the isothermal aging behavior of Cluster (2), aging was performed at 100°C for the alloys with the constant Mg + Si concentration of 1.5 mol%. Figure 10(a) and (b) show the results of hardness and electrical resistivity changes during aging at 100°C. The hardness and electrical resistivity changes are increased with the aging time due to the formation of the large amount of Cluster (2). The alloys with more than 0.66 of the Mg/Si ratio represent almost similar behavior until about 86.4 ks (1 day) and after that different behavior in both the hardness and electrical resistivity changes. Furthermore, the electrical resistivity and hardness increase more in the alloys with over 0.66 of the Mg/Si ratio compared to the 3M12S alloy (Mg/Si ratio is 0.31). In other words, the formation of Cluster (2) is greatly accelerated in the alloys with more than 0.66 of the Mg/Si ratio with increasing aging time.

### 4. Discussion

The various Al-Mg-Si alloys with the different Mg/Si
ratios at the constant total solute concentrations of Mg + Si (mol%) were used in order to investigate the influence of the alloy composition on the nanocluster formation. In the previous research, Kim et al. reported that the Si and Mg concentration of Al-Mg-Si alloys have a marked effect on the formation of nanoclusters. The formation of Cluster (1) is more correlated with the Si concentration, whereas both of the Mg and Si concentrations affect the formation of Cluster (2). These results are closely related with the results reported by Yamada et al. and Serizawa et al., indicating that the Cluster (1) is Si-rich clusters and Cluster (2) is Mg-Si clusters. In this paper, the alloys with the fixed constant Mg + Si concentrations were examined in order to investigate the effect of the Mg/Si ratio more precisely. The distributions of volume fraction of Cluster (1) and Cluster (2) with the Mg and Si concentration obtained by the DSC curves are summarized in Fig. 11, including the previous results with the 9M2S-9M12S-3M12S alloys. The shaded circles are represented as a normalized value by the maximum value. The locations of the binary Al-Mg and Al-Si alloys are also represented by arrows for understanding. From these results, it is found that the volume fraction of Cluster (1) and Cluster (2) is most enhanced when the Mg/Si ratio is close to 1.0. In other words, a certain amount of the solute Si and Mg atoms are required for the nanocluster formation. There are a large number of structures and compositional changes in Al-Mg-Si alloys from the initial supersaturated solid solution to the equilibrium β phase. However, the reason why the nanocluster formation is most accelerated when the value of the Mg/Si ratio is 1.0 is not clear at the moment. The previous research has suggested that the positron lifetime related with the quenched-in excess vacancies in the binary Al-Mg and Al-Si alloys is rapidly decreased, comparing to the slight decrease in the Al-Mg-Si alloys. These results are correlated with the fact that no exothermic clustering reaction occurs in the as-quenched Al-Mg27,28 and Al-Si21 alloys. One of the possible mechanisms is that the thermodynamically metastable state for the nanocluster formation is achieved with the matrix concentration closer to the Mg/Si ratio of 1.0. It is most likely that the nanocluster formation is closely correlated with the interaction energies among solute Mg and Si atoms. As the possible mechanism in the kinetics viewpoint, the activation energy, which is the combination of vacancy formation energy, migration energy and bonding energy between solute Mg and Si atoms and quenched-in excess vacancies, can be considered. It is assumed that the vacancy formation energy is less important due to the existence of quenched-in excess vacancies in the matrix. Therefore, the bonding and migration energies of the Si/Mg/vacancy are significant factors for the nanocluster formation. The diffusion kinetics is favorable to produce nanoclusters when the Mg/Si ratio of the matrix is close to 1.0 due to the interactions among solute atoms and vacancies. The further research is required to clarify these points in more detail, especially the relationship between the alloy composition and nanocluster composition. Furthermore, the vacancy behavior for the nanocluster formation should be also investigated. Pashley et al. firstly proposed a model taking into account the solute atom clusters, and then, Dutta and Allen reported that Si-clusters are formed during low temperature aging based on the DSC analysis. In order to examine the early stage of decomposition from the supersaturated solid solution, there have been many attempts to use a more direct technique using one-dimensional atom probe field ion microscopy (APFIM) and three-dimensional atom probe (3DAP). They revealed the direct evidence of the presence of Si-clusters, Mg-clusters and/or Mg-Si co-clusters. However, the relationship between the alloy composition and cluster composition is still not clear. The alloy compositions were different in the previous researches, the difficult situation to compare. In addition, it is also complicated because of the different heat treatment temperature and time as well as the parameters of measurements. The obtained previous results by the APFIM and 3DAP are briefly summarized in Table 3, including the composition of Cluster (1) and Cluster (2), respectively. According to these summarized results, Cluster (1) has initially various Mg/Si ratios in the alloy close to Mg/Si = 1.0, whereas much more Si-rich clusters exist than the Mg-rich clusters in the excess Si alloys. Mg atoms are incorporated into the Si-rich clusters.
with the NA time, and then Mg-Si co-clusters with the Mg/Si ratio close to 1.0 are formed. In the case of Cluster (2), the Mg/Si ratio becomes 1.0 in a short time at approximately 100°C, different from Cluster (1) formation behavior. These results possibly explain the reason why the most favorable alloy composition is when the Mg/Si ratio is close to 1.0.

In order to overcome the “negative effect of two-step aging” which occurs in the manufacturing of automobile body panels, it is well understood that the transition behavior from the pre-formed nanoclusters into the strengthening phase is the most favorable for the formation of Cluster (1) and Cluster (2). It is closely related with the interaction energies among the solute Mg and Si atoms and vacancies in the matrix.

5. Conclusions

The influence of the Mg/Si ratio on the nanocluster formation with the constant total solute concentrations Mg + Si of about 1.1, 1.3 and 1.5 mol% was investigated in the present paper by means of DSC, hardness and electrical resistivity measurements. From these results conclusions are drawn as follows:

(1) Two exothermic peaks are detected using DSC measurements in the all examined alloys having the different Mg/Si ratio with the constant Mg + Si concentration. It is clarified that the heat evolution, the shape and position of two overlapped exothermic peaks due to the formation of Cluster (1) and Cluster (2) change with the Mg/Si ratio change even though in the constant Mg + Si concentration. These overlapped peaks are well separated using the Gaussian function method.

(2) The formation behavior of nanoclusters strongly depends on the Mg/Si ratio more than the Mg + Si concentration. The alloys with the Mg/Si ratio of approximately 1.0 produce the largest volume fraction of nanoclusters and the lowest activation energy for the nanocluster formation, indicating that the alloy composition is the most favorable for the formation of Cluster (1) and Cluster (2). It is closely related with the interaction energies among the solute Mg and Si atoms and vacancies in the matrix.

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