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As-quenched Microstructures of Cu₃₋ₓMnₓAl (x=0.1, 0.2, 0.3, 0.4) Alloys.

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The as-quenched microstructure of the alloy A (Cu₂.₉Mn₀.₁Al) was D₀₃ phase containing plate-like γ’ martensite; whereas the as-quenched microstructures of both the B (Cu₂.₈Mn₀.₂Al) and C (Cu₂.₇Mn₀.₃Al) alloys were a mixture of (D₀₃+L₂₁+L-J) phases. However, when the manganese content was added up to 9.73 at%, the as-quenched microstructure of the alloy D (Cu₂.₆Mn₀.₄Al) was a mixture of (D₀₃+L₂₁+L-J) phases. The fine D₀₃ phase existing in the as-quenched alloy A, B and C were formed through the A₂→B₂→D₀₃ continuous ordering transition during quenching. The A₂→B₂ and B₂→D₀₃ transitions produced a/₄<111> APBs, respectively. It is noted that a/₄<111> APBs have never been found by other workers in the Cu-Mn-Al alloy system before. Similarly, no evidence of the a/₄<111> APBs could be observed in the as-quenched alloy D. This shows that the energy of the a/₄<111> APBs was increased with increasing the manganese content.

Key words: Cu-Mn-Al alloy, Anti-phase boundary, Phase transformation, Continuous ordering transition

Introduction

By using thermal analysis method, M. Bouchard and G. Thomas had established the Cu₃₋ₓMnₓAl (0 ≤ X ≤ 1) metastable phase diagram [1]. In this phase diagram, it is seen that when the Cu₃₋ₓMnₓAl alloys with 0 ≤ X ≤ 0.8 were solution heat-treated in the single β phase (disordered body-centered cubic (bcc)) region and then quenched into iced brine rapidly, a β (A₂)→B₂→D₀₃+L₂₁ phase transition would occur during quenching. The crystal structure of the L₂₁ (Cu₂MnAl) phase is similar to the D₀₃ (Cu₃Al) phase, and the only difference between them is that manganese atom replaces the copper atom at a specific lattice sites with eight nearest copper atoms in the D₀₃ structure [1]. When the manganese content in the Cu₃₋ₓMnₓAl alloy was increased to 25 at. % (X=1), the as-quenched microstructure became a single L₂₁ phase. In addition, the as-quenched microstructures of the Cu₃₋ₓMnₓAl (0.5 ≤ X ≤ 1.0) alloys were also examined by using transmission electron microscopy [2-5]. These investigations have confirmed the results proposed by M. Bouchard and G. Thomas.

Recently, we made transmission electron microscopy observations on the phase transformation of a Cu₂.₉Mn₀.₃Al alloy. [6] Consequently, our experimental result indicated that the as-quenched microstructure of the Cu₂.₅Mn₀.₃Al alloy was a mixture of (D₀₃+L₂₁+L-J) phases. It is worthwhile to note here that the L-J phase had never been found previously by other workers in the Cu-Al, Cu-Mn and Cu-Mn-Al alloy systems. However, to date, all of the transmission electron microscopy examinations were focused on the Cu₃₋ₓMnₓAl alloys with 0.5 ≤ X ≤ 1. Little information concerning the Cu₃₋ₓMnₓAl alloys with lower manganese content has been provided. Therefore, the purpose of the present study is to investigate the as-quenched microstructure of the Cu₃₋ₓMnₓAl alloys with X < 0.5.

Experimental Procedure

Four alloys, Cu₂.₉Mn₀.₁Al (alloy A), Cu₂.₈Mn₀.₂Al (alloy B), Cu₂.₇Mn₀.₃Al (alloy C) and Cu₂.₆Mn₀.₄Al (alloy D), were prepared in a vacuum induction furnace under a controlled protective argon atmosphere by using 99.99% copper, 99.9% manganese and 99.9% aluminum. The melts were chill cast into a 30x50x200-㎜ copper molds. After being homogenized at 900℃ for 72 hours, the ingots were sectioned into 2-㎜ thick slices. These slices were subsequently solution heat-treated at 900℃ for 1 hour (in the single β-phase state) and then quenched into iced brine rapidly.

Electron microscopy specimens were prepared by means of a double-jet electropolisher with an electrolyte of 70 % methanol and 30 % nitric acid. The polishing temperature was kept in the range from -30℃ to -15℃, and the current density was kept in the range from 3.0x10⁴ to 4.0x10⁴ A/m². Electron microscopy was performed on a JEOL JEM-2000FX scanning transmission electron microscope operating at 200 KV.

Results and Discussion

Figure 1(a) is a bright-field (BF) electron micrograph of the as-quenched alloy A, clearly exhibiting that a second phase with a plate-like morphology was formed within the matrix. Figures 1(b) and (c) show two selected-area diffraction patterns (SADPs) taken from a plate-like phase and its surrounding matrix. In these SADPs, it is seen that in addition to the reflection spots corresponding to the D₀₃ phase [1,7], the diffraction patterns also consist of extra spots caused by the presence of the second phase. Compared with the previous studies in Cu-Al and Cu-Al-Ni alloys [7-10], it is clear that the positions and streak behaviors of the extra spots are the same as those of the γ’₁ (2H) martensite with internal twins [7, 9-10]. The γ’₁ martensite has an orthorhombic structure with lattice parameters a=0.440 nm, b=0.534 nm and c=0.422 nm [9, 11]. Figure 1(c) is a (1Σ₁) γ’₁ dark-field (DF)
electron micrograph, clearly revealing the presence of the plate-like $\gamma'_1$ martensite. Accordingly, it is concluded that the as-quenched microstructure of the alloy A was D0$_3$ phase containing plate-like $\gamma'_1$ martensite.

When the manganese content was increased to X=0.2, no evidence of the $\gamma'_1$ martensite could be detected and a high density of extremely fine precipitates with a mottled structure could be observed within the D0$_3$ matrix. A typical example is shown in Figure 2. Figure 2(a) is a BF electron micrograph of the alloy B in the as-quenched condition. Figures 2(b) and (c) show SADPs of the as-quenched alloy B. When compared with our previous studies in the Cu$_2$Mn$_{0.8}$Al and Cu-14.6Al-4.3Ni alloys [6,12], it is found in the SADPs that the extra spots with streaks showed derive from the L-J phase with two variants. Figure 2(d) is a (002) D0$_3$ DF electron micrograph of the same area as Figure 2(a), revealing the presence of the small B2 domains with a/4<111> anti-phase boundaries (APBs). Figure 2(e), a (111) D0$_3$ DF electron micrograph, shows the presence of the fine D0$_3$ domains with a/2<100> APBs. In Figures 2(d) and (e), it is seen that the sizes of both B2 and D0$_3$ domains are very small. Therefore, it is deduced that the D0$_3$ phase existing in the as-quenched alloy was formed by an A2→B2→D0$_3$ continuous ordering transition during quenching [13-16]. Figure 2(f) is a (0$\overline{2}$0$\overline{2}$) L-J DF electron micrograph, exhibiting the presence of the extremely fine L-J precipitates. Based on the above observations, it is concluded that the as-quenched microstructure of the present alloy B was D0$_3$ phase containing extremely fine L-J precipitates, where the D0$_3$ phase was formed by the A2→B2→D0$_3$ continuous ordering transition during quenching.
Fig. 3 Electron micrographs of the as-quenched alloy C. (a) BF, (b) and (c) two SADPs. The zone axes of the D0\textsubscript{3} phase are (b) [001] and (c) [110], respectively, (d) and (e) (002) and (1\overline{1}1 \ ) D0\textsubscript{3} DF, respectively, (f) (0 \overline{2}0 \ ) L-J DF.

Figure 4(a) is a BF electron micrograph of the as-quenched alloy D, exhibiting a modulated structure. Shown in Figure 4(b) is an SADP of the as-quenched alloy. In this Figure, it is seen that in addition to the reflection spots with streaks of the L-J phase, the superlattice reflection spots with satellites lying along <001> reciprocal lattice directions could be clearly observed. In the previous studies [1,6], it is confirmed that these superlattice reflection spots with satellites were attributed to the coexistence of the (D0\textsubscript{3}+L\textsubscript{2\textsubscript{1}}) phases. Figure 4(c), a (002) D0\textsubscript{3} DF electron micrograph, indicates that no evidence of the a/4<111> APBs could be examined. Figures 4(d) and (e) are (1\overline{1}1\ ) D0\textsubscript{3} and (0 \overline{2}0 \ ) L-J DF electron micrographs of the alloy D in the as-quenched condition, revealing the presence of the D0\textsubscript{3} domains with a/2<100> APBs and L-J precipitates, respectively. As a result, the as-quenched microstructure of the alloy D was a mixture of (D0\textsubscript{3}+L\textsubscript{2\textsubscript{1}} +L-J) phases.

Fig. 4 Electron micrographs of the as-quenched alloy D. (a) BF, (b) an SADP. The zone axis of the D0\textsubscript{3} phase is [001]. (hkl= D0\textsubscript{3} + L2\textsubscript{1}, phase, hkl= L-J phase,). (c) and (d) (002) and (1\overline{1}1\ ) D0\textsubscript{3} DF, respectively, (e) (0 \overline{2}0 \ ) L-J DF.

On the basis of the preceding results, some discussion is appropriate. In the Cu-Al, Cu-Al-Mn, Fe-Al and Fe-Al-Mn alloys[13-18], it is well-known that if the D0\textsubscript{3} phase was formed by continuous ordering transition during quenching, it was always occurred through an A2 (disordered body-centered cubic)→B2→D0\textsubscript{3} transition. The A2→B2 transition produced the a/4<111> APBs and the B2→D0\textsubscript{3} transition produced the a/2<100> APBs [13-16]. However, to date, no a/4<111> APBs could be investigated by other workers in the Cu-Al-Mn alloys [1-7,17-18]. In the present study, it is indeed found that no evidence of the a/4<111> APBs could be observed in the alloy D, which the manganese content is X=0.4. However, when the manganese content was decreased to X=0.3 or below, the a/4<111> APBs could clearly be observed. This result seems to imply that in the Cu-Al-Mn alloys with higher manganese (e.g. X=0.4), the size of the B2 domains could be equivalent to whole grain size. However, the decrease of the manganese content could decrease the B2 domain size significantly. Therefore, the a/4<111> APBs became visible, as shown in Figures 2(d) and 3(d).

In the Cu\textsubscript{3-x}Mn\textsubscript{x}Al metastable phase diagram established by M. Bouchard and G. Thomas, it is seen that when the Cu\textsubscript{3-x}Mn\textsubscript{x}Al alloys with 0.5\leq X\leq1 were solution heat-treated in the β phase region and then quenched into iced-brine rapidly, a β→B2→D0\textsubscript{3}+L\textsubscript{2\textsubscript{1}} phase transition would occur during quenching. By comparing Figures 2 and 3, two important experimental results are given below. (1) The amount of the extremely fine L-J precipitates was increased with increasing the manganese content. Correspondingly, the intensity of the reflection spots and streaks of the L-J precipitates was also increased with increasing the manganese content. (2) The sizes of both the B2 and D0\textsubscript{3} domains were increased with increasing the manganese content.

Conclusions

(1) In as-quenched condition, the microstructure of the alloy A was D0\textsubscript{3} phase containing plate-like γ\textsubscript{1} martensite with internal twin.

(2) With increasing manganese content, the D0\textsubscript{3} matrix would be changed to (D0\textsubscript{3} + L\textsubscript{2\textsubscript{1}} ) with a modulation structure.

(3) The Ms temperature was decreased with increasing the manganese content. On the contrary, the amount of the L-J particles was increased with increasing the manganese content.

(4) No evidence of the a/4<111> APBs could be determined in the alloy D. However, the a/4<111> APBs were clearly observed in the both alloy B and alloy C. This result seems to suggest that the increase of the manganese content in the Cu-Mn-Al alloys could increase the B2 domain size.
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References