Effects of Ca Addition on Solidification Structure of Cu-Sn-Zn Bronze Castings

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The substitution of Ca for a part or all of Pb in Cu-Sn-Zn alloys of 88-8-4 type was carried out in order to decrease solidification defects. The relation between cooling rate and segregation of alloying elements or solidification defects was investigated using steps shaped ingots. Finer dendrite structure was obtained due to decrease in the diffusion coefficient and mass transfer of Sn or Zn in the Cu-liquid for Ca addition alloys. The concentration of Ca and the amount of the micro shrinkage cavity in each step was almost constant in Ca added alloys. The same segregation course in the solidification path showed in both 88-8-4-Pb (CAC402) and 88-8-4-Pb-Ca alloys. In Ca added alloys, the enrichment of Sn in the residual liquid and decrease in final solidification temperature were suppressed by expenditure of Sn due to formation of the compounds consisting of Sn and Ca at higher temperatures compared with the melting point of Pb, which resulted in the decrease of the micro shrinkage cavity. Hardness of Ca containing alloys was higher than that of Ca free alloys. The tensile properties were improved by addition of 0.25 mass% Ca, because of the decrease in micro shrinkage. [doi:10.2320/matertrans.48.799]

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1. Introduction

It is well known from the phase equilibrium of Cu-Sn system that shrinkage cavities and blow holes are easily formed in bronze castings due to the wide melting range.1) Zn and Pb are added in the Cu-Sn bronze because of improvement in castability and machinability, respectively, which results in promotion of formation of defects by solidification.2) An appropriate counterplan has not been made against the formation of defects mentioned above. To investigate the segregation of alloying elements is important, because the formation of defects relates to characteristics in solidification.

Bronze and brass castings including a little amount of Pb have been used as main materials for pipe arrangements of water-supply. Pb in castings is easily eluted in water. The third edition of guidelines for drinking-water quality was published by World Health Organization, WHO, in 2004.3) Effort for decrease or removal of Pb in these alloys has been carried out in Japan.

Bronze melts were often poured into a furan resin mold in this study are listed in Table 1. 88 mass%Cu-8 mass%Sn-4 mass%Zn, 88-8-4 alloy, defined as CAC402 in JIS4) was selected as basic alloy. Limit of Pb as impurities is 1 mass% in CAC402. Ca of 0.02 to 1.0 mass% was added in the 88-8-4 alloy. Molten alloys of Cu-Sn-Zn system were firstly prepared using mother alloy of Cu-12 mass%Sn, pure Cu and pure Zn, and were kept at 1533 K in air. Ca or Pb was added in these molten alloys. A Ca-44 mass%Cu alloy was used for addition of Ca, because its affinity for oxygen was higher. In order to achieve different cooling rate, four and five steps shaped molds which were made of the cast iron and furan resin sand, respectively, were used in solidification of alloys, and the shapes of cast ingots were shown in Figs. 1 and 2.

2. Experimental Procedure

2.1 Preparation of bronze castings

Chemical or nominal compositions of bronze ingots cast in this study are listed in Table 1. 88 mass%Cu-8 mass%Sn-4 mass%Zn, 88-8-4 alloy, defined as CAC402 in JIS4) was selected as basic alloy. Limit of Pb as impurities is 1 mass% in CAC402. Ca of 0.02 to 1.0 mass% was added in the 88-8-4 alloy. Molten alloys of Cu-Sn-Zn system were firstly prepared using mother alloy of Cu-12 mass%Sn, pure Cu and pure Zn, and were kept at 1533 K in air. Ca or Pb was added in these molten alloys. A Ca-44 mass%Cu alloy was used for addition of Ca, because its affinity for oxygen was higher. In order to achieve different cooling rate, four and five steps shaped molds which were made of the cast iron and furan resin sand, respectively, were used in solidification of alloys, and the shapes of cast ingots were shown in Figs. 1 and 2.

2.2 Microstructural investigation

Microstructural observations were carried out by an optical microscope and a scanning electron microscope, SEM, using polished surfaces of each step, A to E, in ingots shown in Figs. 1 and 2. Figure 3 shows schematic presentation for measurement paths of each element by electron probe micro analyzer (EPMA). Quantitative analyses for Cu, Sn, Zn, Pb and Ca in more than 200 points on measurement paths (A-A in Fig. 3) between primary dendrite arms through interdendritic region (C in Fig. 3) and grain boundary area (B in Fig. 3) including the solidification defects, were carried out by EPMA for preparation of the solidification path. The quantitative analyses were carried out in center area (300 × 300μm) for part A with the thickness of 5 mm of the furan resin mold. Amounts of Cu, Sn and Zn were plotted in metal and furan resin molds which had a flight of four and five steps, respectively. The relation between cooling rate and segregation of alloying elements or solidification defects was investigated using the proposed alloys.
triangularly compositional diagram, which was utilized as the solidification path showing the segregation course of alloying elements during solidification.

2.3 Evaluation of mechanical properties

Rockwell hardness number on B scale was measured on the center of each step, A–D, shown in Fig. 1. Button-head type tensile specimens with gauge diameters of 5 mm and gauge lengths of 16 mm were produced from the center in the step, B part, with the thickness of 25 mm. Tensile tests were conducted at 293 K under an initial strain rate of $5 \times 10^{-4}$ s$^{-1}$ in air.

3. Results and Discussion

3.1 Microstructures

The macro structure of each step of ingots was shown in Figs. 4 and 5 for 88-8-4-0.5Ca and 88-8-4-1.0Pb, CAC402, castings poured into the metal mold. The finer dendrite structure was observed as cooling rate increased or the thickness of steps in the castings decreased, regardless of the kinds of alloys. In contrast, the finer dendrite structure was obtained by Ca addition in the 88-8-4 alloy. Behavior in inter-dendritic diffusion of Sn showing the inverse segregation was affected by kinds of ternary alloying elements in Cu-8 mass% Sn bronze alloys. It is considered that finer dendrite structure in Ca added alloys was achieved due to decrease in both diffusion coefficient and mass transfer of Sn or Zn in the Cu-liquid. For the part B with the thickness of 25 mm of both ingots, SEM views showing the micro-segregation of alloying elements and concentration profiles of Cu, Sn, Ca or Pb are shown in Figs. 7 and 8. Segregation of Sn was shown in CAC402 such as the red or yellow marked area in its characteristic x-ray image in Fig. 7(b), and enrichment of Pb was also observed near Sn-enriched area. Copper depleted areas which was yellow or blue marked in its image, corresponded to the segregation of Sn. Micro shrinkage cavity was observed near Pb-enriched area, because of the wide melting range and lower melting point. In contrast, segregation of both Sn and Ca was observed in the 88-8-4-0.5Ca alloy as shown in Fig. 8, and both elements

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Table 1 Compositions (mass%) of some 88-8-4 system alloys used in this study.

<table>
<thead>
<tr>
<th>Alloys</th>
<th>Cu</th>
<th>Sn</th>
<th>Zn</th>
<th>Pb</th>
<th>Ca</th>
<th>P</th>
<th>Used mold</th>
</tr>
</thead>
<tbody>
<tr>
<td>88-8-4</td>
<td>Bal.</td>
<td>8.48</td>
<td>4.58</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Metal</td>
</tr>
<tr>
<td>88-8-4-1.0Pb (CAC402)</td>
<td>Bal.</td>
<td>8.40</td>
<td>4.58</td>
<td>0.98</td>
<td>—</td>
<td>—</td>
<td>Metal</td>
</tr>
<tr>
<td>88-8-4-0.25Ca</td>
<td>Bal.</td>
<td>8.41</td>
<td>4.55</td>
<td>—</td>
<td>0.25</td>
<td>—</td>
<td>Metal</td>
</tr>
<tr>
<td>88-8-4-0.5Ca</td>
<td>Bal.</td>
<td>8.54</td>
<td>4.53</td>
<td>—</td>
<td>0.5</td>
<td>—</td>
<td>Metal</td>
</tr>
<tr>
<td>88-8-4-1.0Ca</td>
<td>Bal.</td>
<td>8.37</td>
<td>4.64</td>
<td>—</td>
<td>1.0</td>
<td>—</td>
<td>Metal</td>
</tr>
<tr>
<td>88-8-4-0.5Pb-0.02P (CAC402)</td>
<td>Bal.</td>
<td>8.11</td>
<td>4.65</td>
<td>0.5</td>
<td>—</td>
<td>0.0225</td>
<td>Furan resin</td>
</tr>
<tr>
<td>88-8-4-0.5Pb-0.02Ca</td>
<td>Bal.</td>
<td>8.11</td>
<td>4.65</td>
<td>0.5</td>
<td>0.02</td>
<td>—</td>
<td>Furan resin</td>
</tr>
<tr>
<td>88-8-4-0.5Pb-0.05Ca</td>
<td>Bal.</td>
<td>8.11</td>
<td>4.65</td>
<td>0.5</td>
<td>0.05</td>
<td>—</td>
<td>Furan resin</td>
</tr>
<tr>
<td>88-8-4-0.5Pb-0.1Ca</td>
<td>Bal.</td>
<td>8.11</td>
<td>4.65</td>
<td>0.5</td>
<td>0.1</td>
<td>—</td>
<td>Furan resin</td>
</tr>
</tbody>
</table>
segregated in the same area. The segregation area consisted of Cu, Sn and Ca. It is considered from Fig. 8 that some compounds of Cu-Sn, Cu-Ca and Ca-Sn systems such as Cu₃Sn, Ca₄Cu, CaCu₅, CaSn₃, etc, were formed in segregation area. The Sn-segregated area in the Ca added alloy was smaller than that in CAC402. Micro shrinkage cavity was also observed near the segregation.

For the part A with the thickness of 5 mm of the 88-8-4-0.5Pb-0.05Ca casting poured into the furan resin mold, concentration profiles of Cu, Sn, Ca and Pb are shown in Fig. 9. Segregation of Sn and Ca was observed in the same area and enrichment of Pb was also observed near Sn-enriched area, which corresponded to the area showing the lower Cu level. Any solid solutions are not formed in the Cu-Pb system and neither are any compound formed in the Sn-Pb and Zn-Pb systems. It is considered that Pb was discharged into the residual liquid and crystallized near the Sn-enriched area which was formed in the final stage of solidification. In
contrast, any solid solutions are not formed in the Cu-Ca, Sn-Ca and Zn-Ca systems. It is considered that Ca was also discharged into the residual liquid and segregated with Sn as the compounds in the final stage of solidification.

It is considered that formation of micro shrinkage cavity was promoted due to concentration of solute elements in the residual liquid in the latest stage of solidification, solidification-shrinkage, long solidification-temperature range, slower cooling rate, etc, as seen in Fig. 7. The relation among cooling rate, macro-composition, and area fraction of micro shrinkage cavity was investigated for 88-8-4-1.0Pb (CAC402) and 88-8-4-0.5Ca alloys cast into the metal mold. Chemical analyses for Cu, Sn, Zn, Pb and Ca were carried out by the wet method using parts which were located at center (10 \times 10 \text{ mm}) of each step (A–D) in castings. It is considered that chemical composition of each element was exactly measured by methods mentioned above. For CAC402, the concentration of Pb and Sn increased and decreased, respectively, as the thickness of each step in the casting increased or the cooling rate decreased, as shown in Fig. 10. Sn and Pb showed the inverse and normal segregation, because segregation of Sn and Pb were mainly caused at higher and lower temperatures, respectively. Even at lower temperature the liquid was supplied in the un-solidified part on each step, from a hot top in part “D” showing macro shrinkage porosity, see Fig. 10(a). The concentration of Zn was almost constant because Cu and Zn formed a complete solid solution in the compositional range of less than 39 mass\% Zn, regardless of the difference in the cooling rate. The area fraction of the micro shrinkage cavity increased as the amount of macro segregation of Pb was increased according with the decrease of the cooling rate. It is found that the micro shrinkage cavity was caused by the macro segregation of Pb in CAC402. In contrast, for 88-8-4-0.5Ca alloy, the concentration of Ca and Zn was almost constant, regardless of the difference in the cooling rate, as shown in Fig. 11. Also, inverse segregation of Sn was observed in this figure and its concentration was changed depending on the position of the metal mold. The segregation behavior of each element was affected by its segregation temperature and supply of liquid from a hot top as mentioned above. The concentration of Sn decreased as the cooling rate decreased. The macro segregation of Ca was not found in this alloy, although that of Pb was caused in CAC402, as shown in Fig. 10. The area fraction of micro shrinkage cavity was almost constant, regardless of the difference in the cooling rate. The amount of micro shrinkage cavity in the Ca added alloy was smaller than that in CAC402. Feeding capability of the 88-8-4-0.5Ca melt was improved, because the melt flowed easily in columnar structures which were formed by Ca addition, see Fig. 5. Figure 12 shows micro shrinkage cavity in part D (t = 45) showing lowest cooling rate of both CAC402 and 88-8-4-0.5Ca castings. The Ca containing alloy showed smaller micro shrinkage cavity, although there was almost same number of shrinkage cavity in both castings. It is found from the result mentioned above that the Ca added alloy showed the finer microstructure. Figure 13 shows the secondary dendrite arm spacings measured on some 88-8-4 system bronze castings poured into metal and furan resin molds. The secondary dendrite arm spacings increased as the cooling rate decreased, regardless of the kinds of castings and molds. For the castings poured into the
metal mold, the shorter secondary dendrite arm spacings was shown, compared with those poured into the furan resin mold. Secondary dendrite arm spacings of the Ca added castings were shorter than that of the Ca free 88-8-4 castings. In contrast, the castings poured into the furan resin mold showed the similar tendency as those poured into the metal mold, for the relation among the secondary dendrite arm spacings, Ca addition and cooling rate.

Fig. 7 (a) A scanning electron microscopic image, (b), (c), (d) concentration profiles of Sn, Pb and Cu, respectively, in the vicinity of the solidification defect in the step with the thickness of 25 mm, for the 88-8-4-1.0Pb (CAC402) casting poured into the metal mold.

Fig. 8 (a) A scanning electron microscopic image, (b), (c), (d) concentration profiles of Sn, Pb and Cu, respectively, in the vicinity of the solidification defect in the step with the thickness of 25 mm, for the 88-8-4-0.5Ca casting poured into the metal mold.

Fig. 9 (a), (b), (c), (d) Concentration profiles of Sn, Ca, Cu and Pb, respectively, in the step with the thickness of 5 mm, for the 88-8-4-0.5Pb-0.05Ca casting poured into the furan resin mold.

Fig. 10 Relation among (a) cross sections of steps with 15–45 mm thickness, (b) area fraction of micro shrinkage cavity and (c) chemical composition of alloying elements in each step, for the 88-8-4-1.0Pb (CAC402) casting poured into the metal mold.
3.2 Solidification path

The solidification path was prepared to investigate a segregation course of alloying elements in the step with the thickness of 5 mm for 88-8-4-0.5Pb-0.02P (CAC402) and 88-8-4-0.5Pb-0.05Ca castings poured into the furan resin mold. It of both ingots was shown in Figs. 14 and 15, respectively, using triangularly compositional representation of Cu, Sn and Zn. The average chemical compositions were shown using an open circle symbol in the figures. The same segregation course in the solidification path was shown in both alloys. The concentration of 40 mass% Sn was observed in final stage of solidification for Pb added CAC402, as shown in Fig. 13. The higher concentration of 40–70 mass% in Pb was shown near the average composition of the alloy, because of two melts consisting of Cu-36 mass% Pb and Cu-87 mass% Pb. Pb-concentrated liquid was formed from the Cu-87 mass% Pb melt in the inter-dendritic region. In contrast, the concentration of 30 mass% Sn was observed in final stage of solidification as shown in Fig. 15, and the compounds (Ca$_2$Sn, CaSn and CaSn$_3$) consisting of Ca and Sn were formed depending on their melting points, in the Pb and Ca containing alloy.

In the Pb-containing and Ca-free alloy (CAC402), the solidification was completed at melting point (600 K) of Pb, by crystallization of Pb from the residual liquid showing the higher Sn concentration, compared with that in the Ca added alloy, as shown in Figs. 14 and 15. It is considered that the solidification defect was caused by the enrichment of Pb in the residual liquid and the shrinkage due to the solidification, and when its enrichment in the residual liquid was remarkable, the micro shrinkage cavity became large. In contrast, by
Ca addition, the enrichment of Sn in the residual liquid and decrease in temperature of last stage in solidification were suppressed by expenditure of Sn due to formation of the compounds consisting of Sn and Ca at higher temperatures compared with the melting point of Pb, which resulted in the decrease of the micro shrinkage cavity. Ca enriched area did not exactly correspond to all micro shrinkage cavities, see Figs. 8, 12, which meant the complete completion of solidification by formation of Cu-Sn compounds such as Cu$_3$Sn, in castings at lower temperature, compared with one for formation of compounds consisting of Ca and Sn or Cu. The amount of micro shrinkage cavity was decreased, because the amount of Cu-Sn compounds became lower due to the expenditure of Sn in Ca added castings as shown in Fig. 15.

Inverse segregation of Sn was reported in this alloying system and affected by some factors such as dendrite morphology, solidification-temperature range, evolution of gas bubbles, distribution and diffusion coefficient of the solute, etc. Inverse segregation of Sn was shown in both Ca-free and -added castings as shown in Fig. 10 and 11. The degree of the segregation is affected by the dendrite arm spacings of the primary α phase which is one of factors for the inverse segregation. The dendrite arm spacings are strongly affected by local cooling time in solidification. The same local cooling time was shown among experimental alloys, regardless of Ca-free and -addition alloys. It is considered that the difference in solidification behavior between Ca-free and -addition alloys, was caused by decrease of dendrite arm spacings due to decrease in the diffusion coefficient and mass transfer of Sn and/or Zn in the liquid for Ca addition alloys. The size of segregation decreased by expenditure of Sn and suppression of its diffusion due to formation the compounds consisting of Sn and Ca, which corresponded to the shorter secondary dendrite arm spacings in the Ca-addition alloys, as shown in Fig. 13.

Ca-addition in castings poured into the furan resin mold is hopeful in the view point of the decrease of solidification defects due to three reasons as follows. (1) The lower level in Sn-concentration was observed in latest stage of solidification in the Ca containing castings, compared with the higher one of Ca-free castings, as shown in Figs. 14, 15. (2) Shorter secondary dendrite arm spacings were obtained in Ca containing castings. (3) The formation of defects which were caused by sulfur in the furan resin mold, was suppressed because of desulfurizing power of Ca.

### 3.3 Mechanical properties

For four alloys, 88-8-4-1.0Pb (CAC402), 88-8-4-1.0Ca, 88-8-4-0.5Ca and 88-8-4, poured into the metal mold, Rockwell hardness number on B scale (HRB) was measured and the result was shown in Fig. 16. Hardness decreased as the microstructure coarsened or thickness of steps in the metal mold increased, regardless of the kinds of alloys. The hardness of cast metals correlates with their solidified structure, which agrees with this result. Hardness of Ca-containing alloys was higher than that of Ca-free alloys. Also, the relation between Ca contents in the 88-8-4 system alloys and (HRB) is shown in Fig. 17. Hardness increased as the Ca content increased in alloys, because of the increase of the compounds consisting of Sn and Ca crystallized in interdendritic region.

For three alloys, 88-8-4-1.0Pb (CAC402), 88-8-4-0.25Ca
and 88-8-4-0.5Ca, poured into the metal mold, the 0.2% proof strength, ultimate tensile strength and fracture elongation obtained from tensile test at 293 K were shown in Fig. 18. The 0.2% proof strength, ultimate tensile strength and fracture elongation were increased by the addition of 0.25 mass% Ca, compared with CAC402, because of decrease in the amount of micro shrinkage cavity in Ca-added alloy. In contrast, for the 0.5 mass% Ca added alloy, the lower elongation and same strength were shown, compared with the 0.25 Ca added alloy.

4. Conclusions

(1) Finer dendrite structure was obtained due to decrease in the diffusion coefficient and mass transfer of Sn and/or Zn in the Cu-liquid in Ca-addition alloy.

(2) The concentration of Pb and the amount of the micro shrinkage cavity increased as the cooling rate decreased in CAC402. In contrast, the concentration of Ca and the amount of the micro shrinkage cavity was almost constant in Ca added alloys, regardless of the difference in the cooling rate.

(3) The same segregation course in the solidification path showed in both CAC402 and Ca added alloys.

(4) For CAC402, Sn was segregated in the area solidified in last stage and enrichment of Pb was also observed near Sn-segregated area. Micro shrinkage cavity was affected by crystallization of Pb at its melting point, the solidification-shrinkage, long solidification temperature range, behavior of solute elements and cooling rate. In contrast, by Ca addition, the enrichment of Sn in the residual liquid and decrease in latest solidified temperature were suppressed by expenditure of Sn due to formation of the compounds consisting of Sn and Ca at higher temperatures compared with the melting point of Pb, which resulted in the decrease of the micro shrinkage cavity.

(5) Hardness of Ca containing alloys was higher than that of Ca free alloys. Hardness increased as the Ca content increased in alloys, because of the increase of the compounds consisting of Sn and Ca crystallized in inter-dendrite grains. In contrast, the 0.2% proof strength, ultimate tensile strength and fracture elongation were increased by addition of 0.25% Ca, because of the decrease in the amount of micro shrinkage cavity in the Ca-containing alloy.

REFERENCES