Fabrication of High-strength Cu40ZnSnTi Brass via Powder Metallurgy and Hot Extrusion†

LI Shufeng*, IMAI Hisashi*, ATSUMI Haruhiko**, KONDOH Katsuyoshi***

Abstract

The super-saturated solid solution of Ti in water-atomized BS40-1.0Ti (Cu-40wt%Zn1.0wt%Ti) brass powder creates a high chemical potential for the precipitation reaction. However, Ti readily segregates in the primary particle boundaries at elevated temperatures, which detrimentally affects the mechanical properties of the extruded BS40-1.0Ti brass via powder metallurgy (P/M). Accordingly, Sn was proposed as an additive in BS40-0.6Sn1.0Ti (Cu-40wt%Zn0.6wt%Sn1.0wt%Ti) to inhibit Ti segregation. Consequently, the Ti precipitates were retained as CuSn3Ti5 compound in both interior grains and grain boundaries rather than in the primary particle boundaries. This result demonstrates that Sn addition can effectively obstruct Ti segregation in the primary particle boundaries, and showed a significant grain refinement and mechanical strengthening effect on BS40-0.6Sn1.0Ti brass.

KEY WORDS: (Brass) (Powder metallurgy) (Extrusion) (Microstructure) (Mechanical property)

1. Introduction

Solid-solution strengthening and precipitation hardening are the primary methods used for the development of high-strength brasses 1-3). Our previous study 4, 5) involved the addition of Ti to Cu-40wt%Zn brass, which favored grain refinement by increasing the formation rate of nucleation and recrystallization centers and by retarding the subsequent grain growth by fine precipitates. Unfortunately, precipitation strengthening effects continue to weaken with increased processing temperature because of segregation and the coarsening of Ti precipitates in the primary particle boundaries by the powder metallurgy route. In this study, Sn was proposed as an additive for the Cu40Zn-1.0wt%Ti brass from the viewpoint of alloy designation and processing method. The addition of Sn to brass is expected to impede the segregation of Ti and to stabilize the mechanical properties of Ti brass at high processing temperatures via powder metallurgy.

2. Experimental

The as-received water-atomized BS40-1.0Ti (Cu-40wt%Zn 1.0wt%Ti), BS40-0.6Sn1.0Ti (Cu-40wt% Zn 0.6 wt%Sn1.0wt% Ti) and BS40 (Cu-40wt%Zn) pre-alloyed powders (Nippon Atomized Metal Powders.) were used as the raw materials. The chemical compositions of the raw powders are shown in Table 1. For sintering, 300g of raw powder was loaded into a cylindrical graphite die and sintered by SPS (DR.Sinter/SPS-1030; Sumitomo Coal Mining) at 400 °C – 600 °C with an interval of 100 °C. The obtained sintering billets with 41 mm diameter were subsequently extruded, using a hydraulic press with a load of 2000 kN (SHP-200-450; Shibayama Machine). Before hot extrusion, the billets were preheated to 650 °C for 1.8 ks under a nitrogen gas atmosphere. The final diameter after extrusion was 7 mm. The extruded round bar was machined into test samples with diameter of 3 mm in accordance with ICS 59.100.01. Tensile tests were conducted on a universal testing machine (Autograph AG-X 50kN; Shimadzu) with a strain rate of 5×10⁻⁴ s⁻¹. The phase compositions of the samples were identified using X-ray diffraction (Labx, XRD-6100; Shimadzu). The microstructure evolutions were conducted using a field-emission scanning electronic microscope (FE–SEM, JEM-6500F; JEOL) and a transmission electron microscope (TEM; JEM-3010, JEOL). The phases were examined using an energy-dispersive X-ray spectrometer (EDS) equipped with the SEM. Crystal orientation information, which contained grain size and phase information, was analyzed by the electron backscatter diffraction (EBSD) technique. EBSD was performed using a TSL (TSL DigiView IV; EDAX) instrument.

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attached to the FE-SEM, which was operated at 20 kV.

Table 1. Particle size and chemical compositions of BS40, BS40-1.0Ti and BS40-0.6Sn1.0Ti alloy powders.

<table>
<thead>
<tr>
<th>Powders</th>
<th>Median size (μm)</th>
<th>Mean size (μm)</th>
<th>Composition (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Zn</td>
</tr>
<tr>
<td>BS40</td>
<td>248</td>
<td>279</td>
<td>40.00</td>
</tr>
<tr>
<td>BS40-1.0Ti</td>
<td>206</td>
<td>250</td>
<td>41.19</td>
</tr>
<tr>
<td>BS40-0.6Sn1.0Ti2256</td>
<td>285</td>
<td>40.09</td>
<td>1.03</td>
</tr>
</tbody>
</table>

3. Results

3.1 X-ray diffraction

The X-ray diffraction data of the raw powders and the SPS compacts of BS40-1.0Ti and BS40-0.6Sn1.0Ti are presented in Fig. 1. For BS40-1.0Ti SPS compacts prepared at 400 °C, α peaks appear at 32.324°, 49.52°, and 72.54°, which indicate that the α phases precipitate and the parts of β phases are transformed to the α phases after being subjected to thermal impact at 400 °C during sintering. The phase volume fractions of the α and β phases in the samples prepared at different temperatures are calculated using the XRD diffraction peak intensities. The α-phase ratio in the sintered BS40-1.0Ti compact increases from 50.7% to 55.9% when the temperature is increased from 400 °C to 500 °C, then decreases to 54.2% as the temperature is increased to 600 °C. For the sintered BS40-0.6Sn1.0Ti compact, the α-phase ratio shows a trend similar to that of the BS40-1.0Ti SPS compact, increased from 50.5% to 52.3%, then decreased to 50.5% as the temperature is increased from 400 °C to 600 °C. The α-phase ratio reaches a maximum value of 55.9% for BS40-1.0Ti and 52.3% for BS40-0.6Sn1.0Ti at 500 °C. The shift of the maximum value of the α-phase ratio from 55.9% for BS40-1.0Ti to 52.3% for BS40-0.6Sn1.0Ti verifies that the addition of 0.6 wt% Sn is favorable to the formation of β-phase. Because the intermetallic β-phase exhibits higher hardness and better hot workability compared to the α-phase, the appropriate increment of β-phase volume fraction created by the addition of Ti and Sn alloying elements is favorable to the mechanical properties of the brass alloys.

In addition, minor peaks at 37.3°, 38.1° and 39.2° are observed in the XRD patterns of BS40-0.6Sn1.0Ti SPS compacts sintered at different temperatures as shown in Fig. 1. These peaks are indexed by spacings characteristic of the ternary compound CuSn3Ti5. Two ternary compounds, CuSn3Ti5 and CuSnTi, and their crystal-structure data have been reported in the Cu-Sn-Ti system 6-8). The ternary compound CuSn3Ti5 is in equilibrium with a liquid at 900 °C and evolves during cooling 6). The peak intensity of CuSn3Ti5 increases with increased sintering temperatures, which indicates that the volume of CuSn3Ti5 increases with temperature from 400 °C to 600 °C.

Microstructure

Fig. 2 SEM micrographs of BS40-1.0Ti and BS40-0.6Sn1.0Ti SPS compacts sintered at different temperatures. (a) BS40-0.6Sn1.0Ti, 400 °C; (b) BS40-0.6Sn1.0Ti, 600 °C; (a') BS40-1.0Ti, 400 °C; (b') BS40-1.0Ti, 600 °C.

The SEM micrographs of the BS40-1.0Ti and BS40-0.6Sn1.0Ti SPS compacts sintered at different temperature are shown in Figs. 2. The BS40-1.0Ti samples sintered at 400 °C contains ultrafine precipitates distributed evenly throughout the structure, as shown in Fig. 2(a). The laminar-structured α phase coalesced and formed large island structures in the β-phase matrix. The Ti precipitates concentrated and formed a layered structure in the primary particle boundaries. Few precipitates were observed in the interior of the α and β phases in the BS40-1.0Ti compact sintered at 600 °C (Fig. 2(b)). In the BS40-0.6Sn1.0Ti compact sintered at 400 °C (Fig. 2(a')), ultrafine, nano-scale-sized Ti precipitates dispersed at grain boundaries and formed net-like structures, and a partial α phase grew that passed through the grain boundaries. After the compact was sintered at 600 °C (Fig. 2(b')), Ti precipitates were observed to grow coarser and were retained inside the particles instead of completely segregating completely in the primary particle boundaries. The precipitates were confirmed to be CuSn3Ti5 IMC according to their XRD pattern, as shown.
in Fig. 1, Figures 3 depict the TEM micrographs of the BS40-0.6Sn1.0Ti SPS compact sintered at 400 °C. Titanium precipitates with a size of 50 nm were observed in the interiors of grains and at grain boundaries, as shown in the bright field (BF) image of Fig. 3(a). These precipitates were also discernible in the dark-field (DF) image of Fig. 3(b).

Fig. 3 Transmission electron micrographs of a BS40-0.6Sn1.0Ti SPS compact consolidated at 600 °C; (a) bright-field image; (b) dark-field image.

2.2 Mechanical properties

Figure 4 presents the true stress versus true strain curves of the extruded BS40, BS40-1.0Ti and BS40-0.6Sn1.0Ti brasses, which are consolidated at different temperatures in the range of 400 °C – 600 °C. All depictions involving mechanical properties represent the extruded materials. The average values of the ultimate tensile strength (UTS), the yield strength (YS), and the elongation are summarized in Table 2. For the samples sintered at 400 °C, the UTS of the BS40-1.0Ti is 617 MPa, which is 23.4% higher than that of the BS40 prepared under identical conditions. When the sintering temperature is increased from 400 °C to 600 °C, the UTS shows a dramatic decreasing trend, that is, a decrease of 19.2% from 617 MPa to 498 MPa, as shown in Fig. 4; this latter value is approximately the same as that of BS40 at 600 °C. The UTS of the BS40-0.6Sn1.0Ti exhibits a distinctly different tendency than does the BS40-1.0Ti; it presents almost the same UTS value as BS40-1.0Ti at 400 °C, shows a slight decreasing trend when the sintering temperature is increased from 400 °C to 600 °C, decreasing by 2.8% from 615 MPa to 598 MPa showing that the addition of elemental Sn has a significantly positive effect on the proportion of the the degradation of the UTS of BS40-0.6Sn1.0Ti at elevated temperatures. The values of σYS for all brasses decrease gradually when the sintering temperature is increased. The σYS value of 390 MPa and 392 MPa are obtained for BS40-1.0Ti and BS40-0.6Sn1.0Ti sintered at 400 °C, respectively, which are 28.7% and 31.3% higher than the 303 MPa σYS value of BS40. These results verify that the addition of Ti makes a valuable contribution to the yield strength of brass, whereas the addition of Sn does not. The behavior of BS40-1.0Ti is, interestingly, different from that of BS40-0.6Sn1.0Ti prepared at an elevated sintering temperature. The σYS of BS40-1.0Ti exhibits a rapid drop from 390 MPa to 248 MPa, in which 36.4% of the σYS is lost when the sintering temperature is increased from 400 °C to 600 °C. This result indicates that BS40-1.0Ti shows approximately the same σYS value as that of BS40. In contrast, the σYS value of BS40-0.6Sn1.0Ti exhibits a moderate reduction of 20.8%. The decreased rate of the σYS degradation at elevated temperatures is clearly attributable to the addition of Sn.

Fig. 4 True stress–strain curves of extruded BS40, BS40-1.0Ti and BS40-0.6Sn1.0Ti brasses; the billets are prepared by SPS at different temperatures of 400 °C – 600 °C. The inset shows a photograph of tensile test sample.

Table 2 Mechanical properties of extruded BS40, BS40-1.0Ti and BS40-0.6Sn1.0Ti brasses prepared using SPS in temperature range of 400 °C – 600 °C.

<table>
<thead>
<tr>
<th>Material</th>
<th>Temp/°C</th>
<th>YS/MPa</th>
<th>Devi. UTS/MPa</th>
<th>Devi. Elong.%</th>
<th>Devi.</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS40</td>
<td>400</td>
<td>303.3</td>
<td>0.58</td>
<td>500.1</td>
<td>0.99</td>
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<tr>
<td></td>
<td>500</td>
<td>288.0</td>
<td>1.66</td>
<td>493.7</td>
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<tr>
<td></td>
<td>600</td>
<td>251.3</td>
<td>1.09</td>
<td>483.4</td>
<td>1.37</td>
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<tr>
<td>BS40-1.0Ti</td>
<td>400</td>
<td>390.2</td>
<td>12.94</td>
<td>617.0</td>
<td>12.55</td>
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<tr>
<td></td>
<td>500</td>
<td>292.5</td>
<td>12.24</td>
<td>530.7</td>
<td>32.71</td>
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<tr>
<td></td>
<td>600</td>
<td>247.8</td>
<td>11.27</td>
<td>498.6</td>
<td>34.42</td>
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<tr>
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<td>398.4</td>
<td>5.43</td>
<td>615.1</td>
<td>5.15</td>
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<td>24.84</td>
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<tr>
<td></td>
<td>600</td>
<td>315.2</td>
<td>15.57</td>
<td>597.7</td>
<td>10.96</td>
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</table>

4. Discussion

The Sn element is completely soluble in copper and forms solid solutions with copper. A super-saturated solid solution of 1.03 wt% Ti in BS40-0.6Sn1.0Ti raw powder is formed by the water-atomization method. The difference in atomic radii between Ti, Sn, Cu, and Zn provides a strain field, which is favourable for the solid solution strengthening. Furthermore, rapid cooling from high temperatures is one method for applying stress to cause dislocation motion 9) and thereby creating a high dislocation density and a high precipitation potential in the raw powder. The higher dislocation density of the microstructure provides more sites for precipitation, which provides a thermo-kinetic advantage for the precipitation reaction 10). A super-saturated solution of Ti in the brass matrix creates a high chemical potential for
the precipitation reaction of Ti. The dislocation density increases gradually with increasing deformation strain during hot extrusion, which supplies more potential nucleation sites for Ti precipitation. In fact, Ti precipitates in the form of an ultrafine dispersoid of CuSn3Ti5 with the addition of Sn during sintering. The precipitates distribute in grain boundaries and act as a pinning effect, which increases resistance to grain boundary slip and produces marked work-hardening effects. However, the tiny CuSn3Ti5 precipitates induce significant refinement of the microstructure, which hinders the movement of dislocations so that stronger mechanical properties and higher hardness are achieved at lower sintering temperatures. The BS40-0.6Sn1.0Ti sintered at 600 °C exhibits a yield strength of 315 MPa, a tensile strength of 598 MPa and a micro-hardness of 216 Hv. These values are much higher than those of BS40-1.0Ti. The results show that the addition of Sn has remarkable positive effects. Sn increases hard strengthening by increasing the volume fraction of the harder β phase. Furthermore, it also can sustain the Ti precipitate strengthening by retaining the Ti precipitates in the interior of grains and at grain boundaries in the form of CuSn3Ti5 compound, as opposed to allowing them to segregate at the primary particle boundaries.

5. Conclusion
The Ti precipitates in the form of CuSn3Ti5 at grain boundaries in BS40-0.6Sn1.0Ti, rather than segregating at primary particle boundaries; this result demonstrates significant grain refinement and mechanical strengthening effects on brass BS40-0.6Sn1.0Ti. The results confirm that the introduction of Sn to BS40-1.0Ti brass can effectively impede the segregation of Ti and stabilize the mechanical properties at high processing temperature via powder metallurgy.

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References: