Temperature Effects on the Whiskers in Rare-Earth Doped Sn-3Ag-0.5Cu-0.5Ce Solder Joints

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The whisker growth in Sn-3Ag-0.5Cu-0.5Ce solder joints after storage at various temperatures is investigated. After reflow, cluster-shaped CeSn$_3$ precipitates form in the solder matrix. It is much easier for the surface of the CeSn$_3$ phases to oxidize than for the solder matrix. The oxidation of the outer surface of these CeSn$_3$ precipitates is accompanied by rapid growth of tin whiskers. Storage temperature affects the morphology and growth kinetics of the whiskers, which are also correlated with the oxidation rates of the CeSn$_3$ surface at various temperatures. The results showed that the driving force for the growth of tin whiskers in this Sn-3Ag-0.5Cu-0.5Ce solder is the compressive stress induced by the diffusion of oxygen into the CeSn$_3$ precipitate clusters, which results in the lattice expansion in the oxidized CeSn$_3$ phase.

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1. INTRODUCTION

Due to the chemical activity of rare-earth (RE) elements, they have been effectively applied for the refining of nonwettable materials [3,4,5] Recently, RE elements have been added into solder alloys to improve their physical and mechanical properties. Ma et al. added 0.05 wt pct mixed metal (Ce, La) to a Sn-40Pb solder and observed that the intermetallic compounds at the interfaces of solder joints were inhibited. [8] In addition, their thermal fatigue life increased by a factor of 3. Chen et al. found that the addition of 0.2 wt pct mixed metal (Ce, La) to a Sn-3.8Ag-0.7Cu solder caused the refining of its microstructure and the formation of needlelike Ag$_3$Sn precipitates, through which the creep rupture life rose seven folds. [7] Dudek et al. added 0.5 wt pct La into a Sn-3.9Ag-0.7Cu solder and showed that the strain to failure of samples increased nearly 150 pct over that of the undoped alloy. [8] Wu et al. also reported that a Sn-3.5Ag solder containing 0.25 wt pct mixed metal possessed better wettability, tensile strength, and creep resistance than undoped alloys. [9] For the Sn-0.7Cu solder, Wu et al. further indicated that alloying with 0.5 wt pct mixed metal could refine its grain size and provide a fine and uniform distribution of Cu$_5$Sn$_3$ intermetallic compounds in the solidified matrix, which resulted in significant improvements of its tensile strength, microhardness, and creep resistance. Upon aging at 150 °C for 20 hours, the microstructure of such a Sn-0.7Cu-RE solder is more stable than that of the undoped Sn-0.7Cu alloy. [10] The beneficial effects of RE addition have also been shown to be effective for Sn-9Zn solder. Wu et al. reported that the Sn-9Zn-RE solder exhibited finer β-Sn grains and higher tensile strength than the undoped Sn-9Zn alloy. [11] In addition, the wetting properties of Sn-9Zn-RE solder were greatly improved due to the decrease of surface tension. Summarizing these experimental results, Wu et al. concluded in a review article that RE elements would play an important role in providing better electronic interconnections. [12]

However, an amazingly rapid growth of tin whiskers has been observed in a Sn-3Ag-0.5Cu-0.5Ce solder joint after room temperature storage in air for just a few days. [13] Because the appearance of tin whiskers in solder joints will cause short circuits, the addition of RE elements to solder alloys could be risky for electronic devices in actual application. Furthermore, during the operation of electronic devices, the temperature of solder joints is increased. The morphology and growth kinetics of tin whiskers might be affected by the storage temperature. The focus of this study has thus been the temperature effects on the whisker growth in this RE element containing Sn-3Ag-0.5Cu-0.5Ce solder.

II. EXPERIMENTAL

The Sn-3Ag-0.5Cu-0.5Ce solder used for this study was prepared by melting a Sn-6.6 wt pct Ce master alloy at 1000 °C under 10$^{-4}$ Pa vacuum and then remelting it with the addition of the elements Sn, Ag, and Cu under the same conditions. The ingot was rolled into a 180-μm-thick plate and punched into a disk of 400 μm in diameter. The solder disks, after being dipped in rosin mildly activated flux, were placed on the Ag surface-finished Cu pads of a ball grid array package. The specimens were then reflowed in a hot-air furnace using a temperature profile with the soaking temperature and peak temperature at 150 °C and 240 °C, respectively. The surface tension of the molten solder used for this study was prepared by melting a Sn-6.6 wt pct Ce master alloy at 1000 °C under 10$^{-4}$ Pa vacuum and then remelting it with the addition of the elements Sn, Ag, and Cu under the same conditions. The ingot was rolled into a 180-μm-thick plate and punched into a disk of 400 μm in diameter. The solder disks, after being dipped in rosin mildly activated flux, were placed on the Ag surface-finished Cu pads of a ball grid array package. The specimens were then reflowed in a hot-air furnace using a temperature profile with the soaking temperature and peak temperature at 150 °C and 240 °C, respectively. The surface tension of the molten solder caused the solder disks to become solder balls after solidification. The reflowed specimens were cut from cross sections of a.
row of solder balls, ground with 2000 grit SiC paper, and polished with 0.3-μm Al₂O₃ powder. Some of the cross-sectioned solder joints were further aged for various storage durations at room temperature and at 50 °C, 100 °C, and 150 °C in an air furnace. Both the reflowed and aged specimens were observed via scanning electron microscopy, and their chemical compositions were analyzed using an electron probe microanalyzer (EPMA).

### III. RESULTS AND DISCUSSION

After reflow, large-sized intermetallic clusters with a composition of CeSn₃ form in the solder matrix, as shown in Figure 1(a). Figure 1(b) reveals that after 3 days of further storage at room temperature in air, many thread-shaped whiskers appear in the region of the CeSn₃ precipitates. The lengths of some whiskers are over 30 μm, and their diameters are about 0.1 to 0.3 μm. The whiskers grow very quickly after room-temperature storage for longer time durations (Figure 1(c)). However, the diameters remain unchanged during the growth of the whiskers. The EPMA analyses show that the chemical compositions of these threadlike whiskers are near pure tin, with small amounts of oxygen. Figure 2 shows that in the case of air storage at 50 °C, the whiskers in Sn-3Ag-0.5Cu-0.5Ce solder joints have a morphology similar to that observed for the specimens aged at room temperature. However, some hillock-shape whiskers can be found at the edges of the CeSn₃ precipitates. The size of these hillocks is about 5 μm, with the length much shorter than that of the thread-shaped whiskers.

When the aging temperature is increased to 100 °C, the hillock-type whiskers become dominant, as shown in Figure 3. However, a small number of thread-type whiskers can still be observed. These fine whiskers can

![Fig. 1](Image)(a) Whisker growth in a Sn-3Ag-0.5Cu-0.5Ce solder joint after air storage at room temperature for various times: (a) as-reflowed, (b) 3 days, and (c) 10 days.

![Fig. 2](Image)(a) Whisker growth in a Sn-3Ag-0.5Cu-0.5Ce solder joint after air storage at 50 °C for various times: (a) as-reflowed, (b) 2 days, and (c) 21 days.
nevertheless grow quickly to a length of over 40 μm in just 2 days, as indicated in Figure 3(b). When the storage temperature is raised further to 150 °C, thread-type whiskers can scarcely be found in the specimens. In contrast, the edges of the CeSn₃ phase are full of hillock-type whiskers, as revealed by Figure 4. Figure 4 also shows that these hillocks appear in as short a time as 0.25 hours. However, both the diameter and length of these hillock-type whiskers cease to increase when the specimens are further stored in air for longer periods. The occurrence percentages of hillock-type whiskers and the average whisker lengths in Sn-3Ag-0.5Cu-0.5Ce solder joints are plotted in dependence of the storage temperatures (Figure 5). It can be seen that, along with the rise of storage temperature, the occurrence percentage of hillock-type whiskers increases from 20 to 86 pct, and the average length of the whiskers decreases from 52.6 to 18.6 μm.

In comparison to the morphology of CeSn₃ phases observed in the Sn-3Ag-0.5Cu-0.5Ce solder joints after air storage, tin whiskers are absent when specimens have been aged at room temperature under 10⁻² Pa vacuum for various time periods up to 21 days, as illustrated in Figure 6. A similar result has also been obtained in the case of 150 °C storage in a vacuum furnace of 10⁻² Pa for 3 hours.

It is interesting that EPMA analyses indicate that the hillocks at the edge of the CeSn₃ phase contain a certain amount of Ag, which is quite different from the chemical composition of the thread-type whiskers. In a few cases, Ag₃Sn particles are even found to be embedded in the surface of the hillocks, which results in the Ag content in the hillocks reaching an amount of 24 at. pct. This result implies that the surrounding hillocks have sprouted from the Sn-3Ag-0.5Cu-0.5Ce solder matrix, rather than from the CeSn₃ precipitates.
Liu et al. studied whisker growth along with the electromigration in Sn thin films, observing that both needle-shaped and hillock-type whiskers grow at the anode when the films are stressed at room temperature and at 50 °C. However, only hillock-type whiskers have been observed when the films are stressed at 75 °C and 100 °C. They suggested that the result may be attributed to the softer surface oxide formed at higher temperatures.\cite{14} Tu also reported on the dominant formation of hillock-type whiskers at higher temperatures.\cite{15} To model the hillock growth, he assumed that whiskers grow at weaker spots on the Sn surface where the surface oxide has been broken, and these spots become localized stress relief centers surrounded by a long-range stress gradient needed for whisker growth.

In order to clarify the temperature effects on the morphology and growth kinetics of tin whiskers in this study, the chemical compositions of the oxide layers on CeSn$_3$ phases after the storage of specimens at various temperatures have been analyzed. Figure 7 reveals that the surface of CeSn$_3$ intermetallic clusters in the as-reflowed Sn-3Ag-0.5Cu-0.5Ce solder joints contains only a slight amount of oxygen. However, the oxygen content on the surface of the CeSn$_3$ phase rises to about 50 at. pct after air storage for 15 days and then remains almost constant with further increases in aging time at room temperature. Along with the rise of the oxygen content, the Ce content also increases slightly, accompanied by the plummeting of the Sn content. The composition (at. pct) of the surface of the CeSn$_3$ intermetallic phase after long-term storage at room temperature is Ce:Sn:O = 40.0:7.5:52.5. The oxygen content on the surface of the CeSn$_3$ phase rises to its maximum value (49 at. pct) after air storage at 50 °C for 3 days, as shown in Figure 8. The final composition after long-term storage at 50 °C is Ce:Sn:O = 45.0:7.0:48.0. When the storage temperature is raised further to 100 °C and 150 °C, the time durations needed for the oxygen content on CeSn$_3$ surfaces to attain its...
maximum value are 1 and 0.5 hours, respectively, as indicated by Figures 9 and 10. The final compositions of oxide on CeSn$_3$ surfaces after long-term storage at 100 °C and 150 °C are Ce$_{46.0}$Sn$_{4.0}$O$_{50.0}$ and Ce$_{45.0}$Sn$_{4.5}$O$_{50.5}$, respectively. Collating the observations of whisker growth with the surface composition analyses of the CeSn$_3$ phase, it is observed that the tin whiskers have stopped sprouting before the attainment of the maximum oxygen content.

Because EPMA analysis gives a maximum depth resolution of about 1 μm, the time duration needed for attaining the maximum oxygen content (Figures 7 to 10) represents the time that the oxide layer takes to grow to a thickness of about 1 μm. The reciprocal square root of time duration ($t^{-1/2}$) for attaining the maximum oxygen content is plotted in an Arrhenius diagram (log $t^{-1/2}$ vs $1/T$), as shown in Figure 11, which indicates an activation energy of 12.9 kJ/mol. This value is quite consistent with the activation energy (15.1 kJ/mol) for the lattice diffusivity of trace radioactive O$^{18}$ in cerium oxide, as measured by Steele and Floyd. It indicates that the...
growth kinetics of the oxide layer on the CeSn₃ phase are controlled by the lattice diffusion of oxygen through the cerium oxide. The results confirm that the growth of tin whiskers in Sn-3Ag-0.5Cu-0.5Ce solder joints has a close relationship with the diffusion of oxygen into CeSn₃ precipitates during the storage of the packages. In contrast to the composition profiles of the surfaces of CeSn₃ intermetallic clusters in air-stored Sn-3Ag-0.5Cu-0.5Ce solder joints, the contents of cerium, tin, and oxygen on the CeSn₃ of specimens aged at room temperature under 10⁻² Pa vacuum remain almost constant, as evidenced in Figure 12.

In order to further clarify the whisker growth mechanism in the Sn-3Ag-0.5Cu-0.5Ce solder joints, specimens were cut across the CeSn₃ phase, as shown in Figures 13 and 14. The cross section of a specimen stored at room temperature for 21 days reveals an oxide layer of about 1 µm in thickness on the CeSn₃ phase (Figure 13(b)). On the other hand, Figure 14(b) shows a very thick oxide layer on the outer surface of the CeSn₃ phase after storage at 150 °C for 3 hours. Surrounding the remaining unoxidized CeSn₃ phase, a thinner oxide layer can also be observed. Through the morphological changes of CeSn₃ phase and oxide layer in both cases, the mechanism for the formation of fiber-shaped whiskers and coarse hillocks in Sn-3Ag-0.5Cu-0.5Ce solder joints during the storage at room temperature and at elevated temperatures, respectively, can be schematically interpreted, as shown in Figures 15 and 16. Because the RE element Ce possesses high chemical activity, the CeSn₃ precipitates react predominantly with oxygen and release the tin atoms: 2CeSn₃ + O₂ → 2CeO + 6Sn. The diffusion of oxygen into the CeSn₃ phase leads to the expansion of its crystal lattice. This lattice expansion is constrained by the CeSn₃ phase beneath the outer oxide layer, and the surrounding solder matrix, which introduces a compressive stress, causes the released tin atoms in CeSn₃ precipitates to sprout out of the cerium oxide. At lower temperatures, the oxide on the surface of CeSn₃ phases grows more slowly. In this case, the resulting compressive stress increases slightly, and the released tin atoms are moderately transported through a number of weak spots within the oxide layer (Figure 15(b)). The whiskers tend to be thin in diameter and grow continuously, even to a length of over 300 µm (Figure 15(c)). In contrast, the high diffusion rate of oxygen through the cerium oxide during high-temperature storage causes the oxide layer on the CeSn₃ phase to
grow quickly to a sufficient thickness, without weak spots. The lattice expansion results in a compressive stress outward from the oxide layer, as shown in Figure 16(b). In addition, the oxygen can easily penetrate through the interface between the CeSn$_3$ and solder matrix and react with the CeSn$_3$ precipitates to form an oxide layer around the CeSn$_3$ phase (Figure 16(c)). The lattice expansion caused by the oxygen invasion also induces a compressive stress on the surrounding solder matrix. Both compressive stresses cause the solder alloy around the CeSn$_3$ phase to sprout out of the boundary between the CeSn$_3$ cluster and solder matrix (Figure 16(c)). The resulting whiskers grow in coarse hillock forms and have a short length. The Ag content, as analyzed by EPMA in the surrounding hillocks, supports this interpretation.

**IV. CONCLUSIONS**

In spite of the many advantages reported for solders doped with RE elements, abnormally rapid growth of tin whiskers can occur during storage in air. This growth is attributed to the predominate oxidation of CeSn$_3$ precipitates, which releases the tin atoms. In this study, it is much easier for the surface of the CeSn$_3$ precipitated clusters in a Sn-3Ag-0.5Cu-0.5Ce solder BGA package to oxidize than for the solder matrix. During the diffusion of oxygen into CeSn$_3$ phases, the resulting compressive stress causes the released tin atoms to sprout from the weak spots of oxide layer on the CeSn$_3$ surface. Diverse oxidation rates of the CeSn$_3$ phase at various storage temperatures lead to a diversity of tin whisker morphologies. At lower temperatures, the surface of the CeSn$_3$ precipitates oxidizes more slowly, which causes only a mild increase of the compressive stress. In this case, tin atoms are transported through
many weak spots in the oxide layer, resulting in the formation of long and thin thread-type whiskers. However, the rapid diffusion of oxygen during high-temperature storage induces a compressive stress outward from the oxide layer of the CeSn₃ phase. In addition, the diffusion of oxygen through the interface between the CeSn₃ phase and the solder matrix leads to the formation of an oxide layer around the CeSn₃ phase. The resulting lattice-expansion through both mechanisms of high-temperature storage compresses the solder alloy adjacent to the surrounding oxide layer, causing the formation of hillocks at the edge of the CeSn₃ phase in a short period of time.

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