Research Article

Fluxless Direct Soldering of Transparent Conductive Oxides (TCOs) to Copper

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Due to the combined advantages of low cost, good soldering properties, and appropriate melting temperature range, novel Sn8Zn3Bi1Mg active solder was developed for direct soldering of transparent conductive oxide (TCO) ceramic targets with oxygen-free copper at 200 °C in air. The TCO specimens have aluminum-doped zinc oxide (AZO) and zinc oxide (ZnO) ceramics. The direct soldering process was performed without the need for flux or pre-metallization of the two transparent conductive oxides. The microstructure, phase constitution, melting characteristics, and soldering properties of the Sn8Zn3Bi1Mg active solder were investigated. The liquidus temperature of the Sn8Zn3Bi1Mg active solder was 198.6 °C, which was very close to the binary Sn-Zn eutectic temperature of 198.5 °C. The effect of temperature on the bonding strength of the solder joints was evaluated. The shear strengths of AZO/Cu and ZnO/Cu joints soldered with Sn8Zn3Bi1Mg active solder were 10.3 and 7.5 MPa at room temperature, respectively. Increasing the temperature from room temperature to 180 °C reduced the bonding shear strengths of AZO/Cu and ZnO/Cu joints to 3.3 and 3.7 MPa, respectively.

1. Introduction

Transparent conductive oxide (TCO) layers such as indium-tin-oxide (ITO), aluminum-doped zinc oxide (AZO), and zinc oxide (ZnO) are widely applied in optoelectronic devices such as huge touch panels, flat-panel displays, photovoltaics, electrically heatable glass, opto-electrical interfaces, and circuitry [1]. Due to their lower cost than that of ITO film, wide band gap, and large exciton binding energy, AZO and ZnO films have attracted much attention. Among the multiple methods of depositing TCO thin film layers onto desired substrates, such as evaporation, spray pyrolysis, chemical vapor deposition, sol-gel, and sputtering, magnetron sputtering is the most attractive technique because of its good reproducibility, high deposition rate, and best deposition results in terms of the conductivity and transparency of thin film layers [2, 3]. The backing plate provides electrical conductivity and heat transfer, and it also supports the target material during sputtering to keep the target from falling off and to avoid cracking of the brittle ceramic target. Thus, the ceramic sputter target must be bonded to copper backing plates of high conductivity. The integrity of the bond between the sputtering target and the backing plate is critical to the performance of the sputtering. Metallic bonds inherently exhibit high thermal and electrical conductivity and are also used as a thermal interface material for reducing the thermal resistance between the TCO sputtering target and copper backing plate. As a result, solder bonded targets are able to operate at higher power densities and deposition rates can be enhanced without overheating or cracking of the targets. However, ceramics are usually ionic, covalent, or ionic-covalent bonded materials. The high energy of ion and covalent bonds makes ceramics chemically inert. Thus, most fillers used for
Active soldering, which is not toxic, low cost, and highly durable, is a promising alternative to the commonly used Sn-Ag-Cu alloy and can improve the wettability, reduce the liquidus temperature to one close to that of Sn-Ag-Cu. 8Vhe active soldering process, which is usually carried out in air, requires mechanical activation such as edge abrasion, brushing, and ultrasonic vibration to destroy the oxide layer that forms on the molten filler metal and thereby to permit the interfacial interaction of the active elements, rare earth elements, titanium, or magnesium with the ceramic substrate to permit the interfacial interaction of the active elements, rare earth elements, titanium, or magnesium with the ceramic substrate. 8Vhe active soldering of SiC/SiC ceramic by using Sn-Zn-Al active solder. Kolenak et al. [12, 13] studied the direct bonding of Al2O3/Cu and silicon using Sn3La and Sn-Ag-Ti (Mg) solders, respectively. Their results demonstrated that adding active elements into the active solder can effectively enhance the soldering ability and that satisfactory joining with ceramics can be achieved. Prior study [14] and the studies by Zhao et al. [15] and Gorjan et al. [16] have demonstrated that adding an active element such as Ti and trace rare earth elements to lead-free Sn-Ag, Sn-Ag-Cu, Sn-Zn, and Sn-Bi solders significantly enhances their wettability, and such fillers have been used to join ceramic and metal at low temperature. The active soldering process, which is usually carried out in air, requires mechanical activation such as edge abrasion, brushing, and ultrasonic vibration to destroy the oxide layer that forms on the molten filler metal and thereby to permit the interfacial interaction of the active elements, rare earth elements, titanium, or magnesium with the ceramics [14].

Among the many Pb-free solders developed to date, Sn-Zn alloys are regarded as promising candidates due to their merits of good mechanical performance and low cost. The addition of 3 wt.% Bi into eutectic Sn-9Zn solder alloy reduces the liquidus temperature to one close to that of eutectic Sn-Pb alloy and can improve the wettability, mechanical properties, and corrosion resistance [17]. Moreover, ZnO-based TCO is a promising alternative to the commonly utilized ITO, which is nontoxic, low cost, and highly durable against hydrogen plasma [18]. In the present work, a novel Sn8Zn3Bi1Mg active solder was developed and used to join AZO and ZnO targets with oxygen-free copper backing plates. The fluxless direct active soldering process was carried out in air without flux, thus further reducing the cost of the bonding. During the sputtering process, large ion currents were generated and caused very intense and localized heating of the target. Generally, a higher power input causes the temperature of the target to increase sharply [19], which in turn limits sputtering rates and final film properties. Therefore, the effect of temperature on the joint strengths was evaluated by shear testing at 120, 150, and 180°C in this study.

2. Experimental

The materials joined in this study were oxygen-free copper and aluminum-doped zinc oxide (AZO) composed of a mixture of 98 wt.% ZnO and 2 wt.% Al2O3 and ceramic targets of zinc oxide (ZnO). The active solder Sn8Zn3Bi1Mg was prepared from Sn, Zn, and Bi of 99.99% purity in the shape of slugs and flaky AZ31 magnesium alloy in a vacuum arc furnace under a high-purity argon atmosphere. The chemical composition of AZ31 magnesium alloy is provided in Table 1. To ensure a homogenous composition within the alloy, the Sn8Zn3Bi1Mg active solder was remelted at least 3 times. After the solder alloy was solidified in a water-cooled copper mold, the cast ingot was rolled to a thickness of 50 μm. The chemical composition of the active solder used in this study was determined by an X-ray fluorescence spectrometer (XRF, Olympus Innov-X Delta) and is given in Table 2.

Differential scanning calorimetry (DSC 204 F1 Phoenix, Netzsch) was used to determine the melting range (liquidus and solidus) of the active solder at a heating rate of 10°C/min from room temperature to 250°C in an Ar atmosphere. An electron probe microanalyzer (EPMA, JEOL TXA-8600SX) equipped with wavelength-dispersive X-ray spectrometer (WDS) was used to analyze the chemical composition and chemical distributions of the elements of the solder alloy. The microstructure of the solder alloy was observed with a scanning electron microscope (FE-SEM, Philips XL40).

The TCO specimens supplied for soldering were processed to a size of 7 mm × 7 mm × 5 mm, as shown in Figure 1(a). Prior to soldering, all bonding surfaces of AZO, ZnO, and copper and the surfaces of the solder alloy were wet-ground with SiC paper down to grade 1200. In active soldering, the AZO and ZnO specimens were pre-heated to 200°C on a heating plate for about 5 minutes in air. Subsequently, the 50 μm thick foil of the active solder was then placed on the AZO and ZnO surfaces. The molten solder alloy was agitated for about 30 seconds for wetting on the surfaces, and then the joints were tightly held with about 1 kg force on the aluminum plate and naturally cooled in air, leading to the solidification of the Sn8Zn3Bi1Mg active solder. For the metallographic observations of the joint interfaces, the soldered specimens were cross-sectioned and characterized with a scanning electron microscope and energy dispersive X-ray (EDS). The joining strengths were measured at room temperature, 120, 150, and 180°C by shear testing, which was performed on a tensile testing machine (Hung Ta HT-2102Bi). At least 4 shear tests were performed, and the highest bonding strengths values were obtained from the valid test results. The dimensions and geometry of the joints subjected to shear testing are shown in Figures 1(a) and 1(b).

3. Results and Discussion

The microstructure of the cast Sn8Zn3Bi1Mg active solder consisted of stripe-like Sn-Zn eutectic phase, needle-like Zn-rich phase (black), Bi-rich phase particles (bright), and granular Mg-Sn intermetallic compound particles, as shown...
in Figure 2. The EPMA analyses indicated that the chemical composition of the Mg-Sn intermetallic compound was Mg: Sn = 65.6:34.4 (at.%). According to Sn-Mg-Zn ternary phase diagram [20] and the compositional analysis, the intermetallic compound corresponds to the Mg$_2$Sn phase.

Figure 3 presents the differential scanning calorimetry (DSC) curves of Sn$_8$Zn$_3$Bi$_1$Mg active solder, indicating the endothermic reactions associated with melting. An endothermic process up to approximately 181.1°C was found in the DSC curve of the alloy and attributed to the existence of a nearly eutectic composition of Sn-Zn-Mg alloy [20]. Sn$_8$Zn$_3$Bi$_1$Mg active solder showed a large endothermic peak at higher temperature, approximately 198.6°C, corresponding to the eutectic temperature of the Zn-Sn binary alloy.

Cross-sectional SEM micrographs of the bonding interfaces of AZO/Cu and ZnO/Cu joints joined with the Sn$_8$Zn$_3$Bi$_1$Mg active solder at 200°C are shown in Figures 4 and 5, respectively. Those figures demonstrate the good wettability of the active filler metal and satisfactory bonding interfaces both between AZO and copper and between ZnO and copper. The copper dissolved irregularly to form a thin intermetallic compound layer with a thickness of about 0.5 μm at the solder/copper interface. The EDS analysis indicated that the composition of the intermetallic layer was Cu:Sn = 41.1:58.9 (at.%), which corresponds to the γ-Cu$_5$Zn$_8$ IMCs. At the AZO/solder and ZnO/solder interfaces, no reaction products were observed after the bonding.

The AZO and ZnO are too brittle and easily break during shear test. After 4 shear tests in each condition, the highest bonding shear strengths values were obtained from the valid test results. The shear strengths of AZO/Cu and ZnO/Cu joints bonded with the Sn$_8$Zn$_3$Bi$_1$Mg active solder were 10.3 and 7.5 MPa, respectively. Figure 6 shows the joining strengths of AZO/Cu and ZnO/Cu joints measured at room temperature, 120, 150, and 180°C. These results indicated that the joint strengths decreased with increases in testing temperature. The lower shear strengths of the AZO/Cu and ZnO/Cu joints at high temperatures could be attributed to the lower strength of
the solder alloy at temperatures above 120°C. The joint strength remained at about 4 MPa at 180°C, very near the liquefaction temperature of the alloy (180.4°C). Figures 7 and 8 present the fractography of AZO/Cu joints with Sn₈Zn₃Bi₁Mg active solder after shear tests at room temperature and at 180°C. The fractography of ZnO/Cu joints with Sn₈Zn₃Bi₁Mg active solder after shear tests at room temperature and at 180°C is shown in Figures 9 and 10, respectively. All fractured surfaces of the copper were covered with a Sn₈Zn₃Bi₁Mg active solder layer, which indicated that both joints fractured along the AZO/Cu interface.
Figure 7: Fractography of an AZO/Cu joint with Sn$_6$Zn$_3$Bi$_1$Mg active solder after shear tests at room temperature: (a) the Cu side; (b) the AZO side.

Figure 8: Fractography of an AZO/Cu joint with Sn$_6$Zn$_3$Bi$_1$Mg active solder after shear tests at 180°C: (a) the Cu side; (b) the AZO side.

Figure 9: Fractography of a ZnO/Cu joint with Sn$_6$Zn$_3$Bi$_1$Mg active solder after shear tests at room temperature: (a) the Cu side; (b) the ZnO side.
solder and ZnO/solder interfaces. Few of active solder was on the fractured surfaces of the AZO or ZnO side after the shear test at 180°C.

4. Conclusion

In this study, AZO/Cu and ZnO/Cu were joined at 200°C in air by using a novel Sn8Zn3Bi1Mg active solder. The liquidus temperature of the Sn8Zn3Bi1Mg active solder was very close to the binary Sn-Zn eutectic temperature (198.5°C). Satisfactory joints of AZO/Cu and ZnO/Cu were realized at 200°C in air without pre-metallization for AZO and ZnO. It was found that during the soldering process, copper dissolved and diffused into the solder, forming Cu5Zn8 intermetallic compounds. The shear strengths of the AZO/Cu and ZnO/Cu joints with Sn8Zn3Bi1Mg active solder at room temperature were 10.3 MPa and 7.5 MPa, respectively. The joint strength decreased to 3.3 MPa of AZO/Cu and 3.7 MPa of ZnO/Cu, respectively, as the testing temperature increased to 180°C. This active soldering process provides a low-cost, high-quality, and easy direct ceramic soldering technology with environmental benefits.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

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