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Published in:
IEEE Transactions on Components, Packaging, and Manufacturing Technology, Part C

Link to article, DOI:
10.1109/3476.681391

Publication date:
1998

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):

DTU Library
Technical Information Center of Denmark

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Oxidation and Reduction Kinetics of Eutectic SnPb, InSn, and AuSn: A Knowledge Base for Fluxless Solder Bonding Applications

Jochen F. Kuhmann, Andrea Preuss, Barbara Adolphi, Karsten Maly, Thomas Wirth, Werner Oesterle, Wolfgang Pittroff, Gerd Weyer, and Marco Fanciulli

Abstract—For microelectronics and especially for upcoming new packaging technologies in micromechanics and photonics fluxless, reliable and economic soldering technologies are needed. In this article, we consequently focus on the oxidation and reduction kinetics of three commonly used eutectic solder alloys:
1) SnPb;
2) InSn;
3) AuSn.

The studies of the oxidation kinetics show that the growth of the native oxide, which covers the solder surfaces from the start of all soldering operations is self-limiting. The rate of oxidation on the molten, metallic solder surfaces is significantly reduced with decreasing $O_2$ partial-pressure. Using in situ Auger electron spectroscopy (AES) it could be shown for the first time, that $H_2$ can reduce Sn-oxide as well as In-oxide at moderate heating duration and temperatures.

In the second part of this study, the results, obtained by the investigation of oxidation and reduction kinetics, are applied to flip-chip (FC) bonding experiments in vacuum with and without the injection of $H_2$. Wetting in vacuum is excellent but the self-alignment during flip-chip soldering is restricted. The desired, perfectly self-aligned FC-bonds have been only achieved, using evaporated and reflowed AuSn(80/20) and SnPb(60/40) after the introduction of $H_2$.

Index Terms—Auger electron spectroscopy, AuSn, flip-chip bonding, fluxless soldering, InSn, oxidation kinetics, SnPb.

I. INTRODUCTION

Fluxless soldering has been a topic for many researchers in the last decade. Environmental concerns and reliability issues have been the main drive [1]–[3]. In the last few years, other topics appeared which again increased the impetus to develop fluxless soldering techniques: The assembly of micromachined [4], [5] and photonic devices [6], [7] and low cost flip-chip (FC) solder bonding on organic substrate materials [8], [9].

In the case of micromachined components, fluxes can clog nozzles and air gaps and destroy very thin and sensitive membranes. Cleaning of fluxes in order to remove corrosive flux residues will be aggravated by capillary forces and might in the case of sensors with membranes not be tolerable at all. For the photonic assembly the major concern in using fluxes is attenuation or deflection of the optical signal and degradation of unpassivated devices. The need to use underfiller in order to compensate for the higher thermal mismatch between organic substrates and silicon dies are the main driving forces in the area of low cost, high volume application. Here the residues of fluxes can lead to poor adhesion of the underfiller and subsequently to failure of the joint when subjected to thermal cycling.

Despite all efforts to circumvent the use of fluxes for sensitive applications too little is known about solder oxides, their growth during soldering in ambient and under reduced $O_2$ partial-pressures. In addition, the potential of $H_2$ to reduce oxides on solder surfaces at typical bonding temperatures is misconceived [10]–[12]. Therefore, in this article we present both new and previously published data about fluxless, self-aligned FC-solder bonding investigations [13]–[16] together with the study of oxidation and reduction kinetics on eutectic InSn [17], SnPb [18], and AuSn.

The studies to determine the scale growth of the oxide were carried out using Auger electron spectroscopy (AES). The thickness of the films, formed on eutectic InSn and SnPb were measured as a function of temperature and $O_2$ partial-pressure. In the case of SnPb(60/40) the native oxide layer was additionally characterized using Moessbauer spectroscopy (MS) and transmission electron microscopy (TEM).

Furthermore, for all solders the results of in-situ AES measurements will be given, which prove the potential of $H_2$ to reduce the solder oxides. The applicability of $H_2$ as an effective flux substitute during bonding at moderate temperatures and heating cycles will be demonstrated by photographs, recorded during FC-bonding experiments.

II. OXIDATION AND REDUCTION IN VACUUM

A. Thermodynamics of Oxidation

The change in the Gibbs free energy $\Delta G$, associated with a chemical reaction gives the answer, whether a reaction as, e.g., the oxidation of tin (1) will ($\Delta G < 0$) or will not occur ($\Delta G > 0$).
The following equations exemplify the thermodynamic calculations for the oxidation of tin to stannic oxide

\[ \text{Sn} + \text{O}_2 = \text{SnO}_2 \]  

(1)

Whether oxidation or decomposition of the oxide occurs is determined by the Gibbs free energy \( \Delta G^0 \) at the relevant temperature. For equilibrium condition \( (\Delta G = 0) \) \( \Delta G^0 \) is given by

\[ \Delta G^0 = R T \ln(k_{p2}) \]  

(2)

in which \( R \) is the molar gas constant and \( k_{p2} \) the equilibrium constant. Equation (2) can also be expressed in terms of \( \text{O}_2 \) partial-pressure \( (p_{\text{O}_2}) \), normalized by the \( \text{O}_2 \) partial-pressure under standard conditions \( (p_{\text{O}_2}^{\text{st}}) \) as

\[ \Delta G^0 = R T \ln \left( \frac{p_{\text{O}_2}}{p_{\text{O}_2}^{\text{st}}} \right). \]  

(3)

\( p_{\text{O}_2} \) at 250 °C for the oxidation of Sn to stannic oxide as described in (1) is given as

\[ p_{\text{O}_2} = 10^{-43} \text{ Pa}. \]  

(4)

Partial-pressures in this range can clearly not be realized by vacuum equipment. Therefore a chemical reaction is used to reach \( \text{O}_2 \) partial-pressures in that order of magnitude

\[ 2\text{H}_2 + \text{O}_2 \rightarrow 2\text{H}_2\text{O}(g). \]  

(5)

According to (2) the equilibrium constant \( k_{p2} \) of water formation (5) at a specific temperature defines the \( \text{O}_2 \) partial-pressure as

\[ p_{\text{O}_2} = \frac{p(\text{H}_2\text{O})^2}{k_{p2} \cdot p(\text{H}_2)^2}. \]  

(6)

From a thermodynamic point of view Sn should be predominantly oxidized in AuSn and SnPb solders whereas in InSn alloys In would be oxidized preferentially. Table I gives an overview of the available \( \Delta G^0 \) values for the possible oxides. Table II lists the \( p_{\text{H}_2}/p_{\text{H}_2\text{O}} \) which mark the threshold between oxide formation and decomposition \( (\Delta G = 0) \).

It is of great importance to realize that meeting the boundary conditions for reduction is mandatory in order to reduce the oxide layers. This means, that the ambient needs to be very well controlled. The desorption of water during heating or contamination of the solder stemming from photoresist or co-deposits during electroplating have a detrimental effect on the reduction. These are the reasons why we believe the potential of \( \text{H}_2 \) to reduce Sn-oxide has been misjudged as stated before.
Fig. 2. Scale growth on liquid nonreduced samples during heating in air at 200 and 250 °C.

Fig. 3. Scale growth on liquid SnPb(60/40) during 10\(^{-4}\) Pa at 200 and 250 °C.

Fig. 4. Scale growth on liquid SnPb(60/40) during different pressures at 200 °C.

Fig. 5. Scale growth on liquid SnPb(60/40) during different pressures at 250 °C.

Films carrying the native oxide did not yield any diffraction patterns but could neither be characterized by EDX. Moessbauer Spectroscopy was therefore carried out to determine the binding state within the native oxide: with this method the oxide could be identified unambiguously as SnO\(_2\). No stannous oxide was detected.

2) InSn(51.2/48.8): Eutectic InSn was oxidized at 22, 100, 150, and 250 °C. For the oxidation treatment at 4 × 10\(^{-8}\), 7 × 10\(^{-5}\), 7 × 10\(^{-4}\) Pa partial-pressure of O\(_2\), which was carried out inside the AES chamber, the samples were sputtered prior to oxidation to remove the native oxide. This was not possible for the samples treated in ambient (O\(_2\) partial-pressure = 2 × 10\(^{4}\) Pa). Here the native oxide was left on top of the samples. Figs. 6–9 give the measured scale growth of the samples as a function of O\(_2\) partial-pressure and temperature. The scale growth of the samples treated below the melting point of the solder (120 °C) could be fitted to a logarithmic law whereas above the melting point of the solder a parabolic growth law was observed.

The characterization of the oxide films were carried out by X-ray diffraction (XRD) and X-ray photon spectrosocpy (XPS). Despite the intermetallic phases InSn\(_4\) and In\(_2\)Sn no oxide was found on the samples which have been stored at room temperature. On samples treated at temperatures above the melting point In\(_2\)O\(_3\), In(OH)\(_3\), SnO, and SnO\(_2\) were detected by XRD.

XPS of samples treated at room temperature and at 190 °C showed the surfaces to have a similar composition of the oxide: In\(_2\)O\(_3\), In\(_2\)O, SnO, and with increasing depth also metallic In and Sn. A distinct surface segregation of In, which increased with increasing temperature, could be found on all the investigated samples.
3) AuSn(80/20): AuSn samples were only characterized as to their native oxide. Similar to eutectic SnPb and InSn also AuSn exhibits a difference between surface and bulk concentration of the solder: Sn segregates to the surface and is, of course, the only element to be oxidized in this solder. By AES the oxide covering eutectic AuSn was found to have approximately the same thickness as the oxide which is covering eutectic SnPb (~3 nm).

C. Reduction

Reduction experiments were carried out for all three solders by in situ AES. Sn and In, which are the predominantly oxidized species in the investigated alloys, have a distinct energy difference of several eV between the oxidized and the metallic state spectra. The reduction was therefore monitored by this energy shift. The thermodynamical conditions for the decomposition of the oxide at temperatures relevant for soldering operations are given in Table II. The samples, covered with their native oxide were introduced into the AES chamber which was then pumped down to 10^-9 Pa. Then the samples were heated up and H₂ was introduced, allowing the pressure to augment the initial base pressure by the factors given in Table II. The typical AES spectra of tin peaks in SnPb(60/40) and AuSn(80/20) before and after treatment with H₂ can be seen in Fig. 10. At approximately 250 °C a distinct energy shift was observed between the oxidized and metallic state of tin. The energy shift was recorded within 2 min. It is interesting to note that the reduction of Sn-oxide on AuSn(80/20) occurred, while the solder was still solid.

Fig. 11 describes the reduction of In-oxide by H₂. In₂O₃ is much more stable than SnO or SnO₂ and therefore also higher H₂/H₂O ratios are required. This is the reason, why in the case of this solder alloy the temperature needed to be elevated to 250 °C in order to meet the required thermodynamic boundary conditions (see Table II).

III. REFLOW AND BONDING EXPERIMENTS

Reflow experiments with solder bumps of all three investigated materials were done in vacuum. Very smooth bumps with the desired shape of a truncated sphere were obtained when using SnPb and also AuSn (see Fig. 12). In both cases the solder material was deposited by evaporation technologies. In the case of eutectic InSn the bumps were deposited by electroplating. The surface as well as the homogeneity of the electroplated InSn solder bumps was of lower quality compared to the material, deposited by evaporation (see Fig. 13).

Bonding experiments with evaporated SnPb(60/40) and AuSn(80/20) in vacuum always showed excellent wetting.
Fig. 11. Energy shift during in situ AES between In-oxide and metallic In, observed during reduction of In-oxide on eutectic InSn by H₂ at 250 °C.

Nevertheless, the passive alignment of the samples was not achieved in the narrow tolerances which are needed, e.g., for the assembly of photonic devices. In that area, using pre-reflowed SnPb and AuSn solder bumps, deposited by evaporation, it was shown that the desired bonding results only were achieved with the use of H₂ as a safe flux substitute [15].

Figs. 14 and 15 show two cross sections of samples bonded in vacuum (p_{total} = 10⁻⁴ Pa): from Fig. 14 it can be seen, that the bump-shape of the reflown solder bumps partly remained intact. Fig. 15 shows the solder bumps still remain in their tilted position after bonding. Using evaporated and reflown SnPb(60/40) bumps the alignment was not in the tolerable range of 2 μm.

In contrast to that, bonding experiments at 250 and 320°C (t ≤ 2 min), carried out with H₂ as a reducing agent (p_{H₂}/p_{H₂O} values see Table II), using eutectic SnPb and AuSn bumps led to solder joints exhibiting cross-sections of a barrel shaped, symmetrical contour. With this new FC-bonding process an accuracy of 1.4 ± 0.8 μm (n = 35) was achieved.

The potential of H₂ to reduce oxides on solder bumps can be very effectively demonstrated using in situ observation during the self-alignment. The sequence of a FC-bonding process, using evaporated AuSn(80/20) solder bumps (see Fig. 16) clearly proves the applicability of H₂ for a fluxless and highly accurate FC-soldering process. After pre-alignment within an accuracy of ~10 μm the samples were heated up to 320 °C. After the melting point of the solder was reached a spontaneous self-alignment occurred within the first few seconds. This self-alignment did nevertheless not exceed the accuracy of ~5 μm. Then, after 120 s H₂ was introduced into the recipient in order to meet the required H₂/H₂O ratios for reduction of the Sn-oxide. The remaining oxide on the solder surfaces was reduced and the residual offset taken away within 60 s.
**Fig. 16.** Pictures, taken through a glass chip during FC-bonding. During bonding in vacuum self-alignment occurred, but only after injection of \( \text{H}_2 \) a complete overlap of chip metallization and bump footprint can be seen: (a) pre-alignment, \( t = 0 \) s, (b) alignment in vacuum \( (10^{-4} \text{ Pa}) \), \( t = 120 \) s, and (c) alignment in vacuum \(+\text{H}_2\), \( t = 180 \) s.

**IV. CONCLUSION**

This study of oxidation and reduction kinetics in conjunction with reflow and bonding experiments leads to important conclusions regarding fluxless soldering technologies:

1) On both solders a crystalline oxide is formed. For eutectic InSn and SnPb the oxide growth on the molten, metallic solder surface can be effectively reduced in vacuum.

2) The native oxide, grown on the solder surface during storage in ambient is an effective protection. For eutectic InSn and SnPb the oxide is most likely amorphous. In the case of SnPb(60/40) even during heat treatment above the eutectic temperature \( (200 \text{ and } 250 \, \text{C}) \) the oxide did not grow significantly. During storage at room temperature for three days and three months the oxide thickness was in both cases \( \sim 3.5 \, \text{nm} \). For InSn(51.2/48.8) a logarithmic growth of the oxide layer was observed which leads to \( 7-8 \, \text{nm} \) oxide after three days and three month storage at \( 22 \, \text{C} \), respectively. The eutectic AuSn was not characterized as thoroughly as the other solder materials but also here no deviation from the well known logarithmic scale growth is expected.

3) Reflow in vacuum showed good bump formation for all solder alloys, the surface of the InSn alloy being the least smooth. This might be due to the deposition technology, which was in the case of SnPb and AuSn evaporation whereas the InSn solder was electroplated. For AuSn(80/20) and also SnPb(60/40) even at elevated \( \text{O}_2 \) partial-pressures \( (10^{-1} \, \text{Pa}) \) a very good reflow of the evaporated solder to truncated spheres was observed.

4) Bonding experiments in vacuum using evaporated reflowed SnPb(60/40) and AuSn(80/20) bumps showed excellent wetting. Also self-alignment of the samples after wetting occurred. The self-alignment depended on pre-alignment and the rate of reoxidation but never reached the quality which was observed using fluxes. The governing mechanism can be proven by the contour of the cross-sectioned solder joints and by \textit{in situ} observation of the bonding process. Where the misalignment between the chip metallization and the pre-reflowed solder bumps has been moderate, the bump contour indicates, that large surface areas are still covered by the native oxide: the shape of the reflowed bumps remained partly intact. Increasing misalignment between the parts led to tilted solder joints. Here the wetting of the opposite metallization has supported a larger change of the bump contour. In these cases self-alignment clearly could be observed. Nevertheless, the remaining oxide or reoxidation impeded self-alignment within the very close tolerances \( (\leq 2 \, \mu\text{m}) \), needed, e.g., for the assembly of photonic devices.

5) In general the starting point of all soldering operations is a solder covered by its native oxide. This oxide, which for all investigated solder materials is very thin and stable can be taken away by \( \text{H}_2 \) during soldering in vacuum. The FC-bonding results, using evaporated and reflowed AuSn(80/20) and SnPb(60/40) clearly prove the applicability of \( \text{H}_2 \) for a fluxless soldering process. Even high demands on alignment accuracy could be met with this process employing moderate temperatures and short heating cycles. In addition to the bonding results the potential of \( \text{H}_2 \) to reduce the oxides on eutectic InSn, SnPb, and AuSn solder could be proven by means of \textit{in situ} AES measurements. Here the transition of the predominately oxidized species from oxidized to
the metallic state was monitored. Provided the necessary thermodynamic boundary conditions were met the reduction of the oxides occurred within 2 min.

ACKNOWLEDGMENT

The authors gratefully acknowledge the support of numerous colleagues who helped build up the FC-bonder used for the experiments and special thanks are due to G. Urmann, HHI, for performing SEM pictures.

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