TEM microstructural analysis of as-bonded copper ball bonds on aluminum metallization

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TEM Microstructural Analysis of As-bonded Copper Ball Bonds on Aluminum Metallization

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Abstract
In this study, the nano-scale interfacial details of ultrasonic copper ball bonding to an aluminum metallization in the as-bonded states were investigated using high resolution scanning/transmission electron microscopy with energy dispersive spectroscopy. Our results showed that ultrasonic vibration swept aluminum oxide and copper oxide in some regions of contacting surface, where an approximate 20 nm Cu-Al intermetallics (i.e., CuAl2) formed. In the regions where oxide remained, aluminum oxide layer connected with copper oxides layer. No nano-level voids or gaps were observed at the central area of the interface, including the regions with oxide. Calculation of interfacial temperature showed that the ultrasonic vibration increased the flash temperature up to 465 °C which was believed to improve the interdiffusion for the formation of Cu-Al intermetallics.

1. Introduction
Thermosonic copper wire-ball bonding is an absorbing interconnection technology that serves viable and cost saving alternative to thermosonic gold wire-ball bonding [1]. Its excellent mechanical and electrical characteristics make copper ball bonding attractive for high-speed, high-power devices and fine-pitch applications. There have been many studies on the effects of bonding process parameters on bondability, which was also evaluated through shear/pull tests, [2-4] and Cu-Al interfacial evolution via thermal cycling/aging [5-7]. Also, several models were established to explain the bonding mechanism, including fretting [8] and micro-slip model [9]. Fretting mechanism was proposed by Hull et al. [8] who illustrated that interfacial sliding between wire and pad cleaning and heating surface was the key for bonding. Micro-slip model [9] predicted slip only existed at the periphery of the contacting interface, while two metals stick in the central regions, thus bonding was preferential at the periphery. However, all these models were lack of experimental supports as to the fundamental mechanisms. The details of atomic features at the bonding interface have not yet been observed to provide any supporting evidence.

IMCs formation and potential voids and cracks have a significant effect on the strength and reliability of the bonds. A proper amount of IMCs formation in a wire bond increases the bonding strength but its excessive growth can result in the performance degradation of the bond. It has been reported that a layer of Au-Al intermetallics was formed during bonding process for gold ball bonded on aluminum metallization, typically 200-500 nm thick [10, 11]. Compared with the rapid growth of Au-Al IMCs, the growth of Cu-Al IMCs has been suggested to be much slower [12]. SEM observation of interfaces by Wulf et al. [6] found no Cu-Al IMCs was formed in the as-bonded Cu-Al bonds. However, the conclusion was not firm due to limitations of analytical tool they used to carry out the study. Therefore, an advanced nano-level analysis tool becomes a key to ascertain interfacial metallurgical characteristic of as-bonded Cu-Al bonds. The importance of interfacial analysis at the atomic level is necessary in order to obtain the direct evidence of a fundamental mechanism of copper thermosonic bonding.

In this paper, a 0.4 µm 99.99% in purity copper wire was bonded on a 3 µm thick Al metallization using an ASM angle 60 automatic thermosonic ball/wedge bonder with a frequency of 138 kHz. High resolution scanning/transmission electron microscopy (HR SEM/TEM) with energy dispersive spectroscopy (EDS) were conducted to ascertain the interfacial metallurgical characteristics of as-bonded Cu-Al bonds. Site-specific TEM samples were prepared using a dual-beam focused ion beam (FIB) system. The thermosonic copper ball bonding mechanism was then proposed.

2. Experimental Procedures
Bonding process
In the experiment, a 0.4 µm copper wire (99.99% purity) was bonded onto a 3 µm thick Al metallization pad on a silicon chip using an ASM Eagle 60 ball/wedge automatic bonder with an ultrasonic frequency of 138 kHz.

An electronic flame-off (EFO) process was applied to produce a spherical ball as the first step of the bonding process circle. The initial copper ball was achieved with a current of 150 mA and a gap voltage 5500 V with a discharge time of 1.3 ms. In order to prevent the copper ball from oxidation in the EFO process, a protective shielding gas of mixture - 95%N2+ 5%H2 - was supplied and maintained at a flow rate of 0.8 L/min. The copper ball was then bonded to an aluminum metallization using a combination of heat, transverse ultrasonic vibration, and a normal force. The detailed bonding parameters for the copper ball bonds are shown in Table 1 after the process window was optimized by means of pull/shear tests and dimension tests.

Table 1 Optimized bonding parameters for copper ball bonds

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ultrasonic power (DAC)</td>
<td>90</td>
</tr>
<tr>
<td>Bonding force (gf)</td>
<td>110</td>
</tr>
<tr>
<td>Bonding time (ms)</td>
<td>28</td>
</tr>
<tr>
<td>Substrate temperature (°C)</td>
<td>220</td>
</tr>
</tbody>
</table>

Characterization
The shear strength of the wire-bonds was measured using a Dage-4000 shear tester, using lateral movement of a shear tool positioned 3µm above the aluminum metallization. The maximum shear value was measured by a strain gauge transducer.

A dual-beam FIB workstation was used to prepare site-specific TEM specimens across the Cu-Al interfacial zone. First, the milled ball was thinned by bombardment of ion
beam, till the height from the top of the thinned ball to Cu-Al interface was below 10 μm. Two holes were etched by focused ion beam, while an approximate 2 μm width foil remained between them, as shown in Fig. 1. This foil was the rough TEM sample. Compared with traditional preparation methods, such as ion beam etching and electropolishing, the FIB microscope has been used as a high-resolution imaging tool. Its secondary electron images provide enhanced crystallographic contrast similar to the electron channeling contrast in SEM. Therefore, interesting zones can be located and selected accurately, making it especially suitable for the multi-interface observation in high resolution analysis. The in-situ FIB lift-out technique was used to obtain these site-specific TEM specimens. An on-stage in-situ tungsten (W) lift-out probe was used. The W lift-out probe was attached to the specimen by ion beam assisted Pt chemical vapor deposition (CVD). The TEM specimen was then lifted out from the bulk using the W probe, as shown in Fig. 2. Before the specimen can be adhered to the TEM grid, a semicircular Mo TEM grid was positioned on the sample stud. A Mo grid instead of Cu grid which was usually used was applied to avoid x-ray energy overlaps with Cu from Cu ball.

After the sample was lift-out, the grid was moved into the field of view. The probe was lowered and loaded onto the TEM grid, as seen in Fig. 3. Then the foil was attached to the grid using the ion beam Pt CVD. Once the specimen has been attached to the TEM grid, the in-situ W probe was milled free by focused ion beam. The focus ion beam with lower current was then used to thin the specimen to electron transparency (100 - 200 nm) as shown in Fig. 4. Through all the sample preparation process, the site-specific TEM samples were stress-free and contamination-free.

Fig. 3. TEM foil touched a Mo grid, followed by ion beam Pt CVD to make adhesion.

Fig. 4. TEM foil was finally thinned using focused ion beam for electron transparency, i.e. 100 - 200 nm.

HRSEM was conducted to view the morphology of the interface in as-bonded Cu-Al bonds, using the secondary electron of the double beam FIB workstation.

HRTEM analysis was done at 200 kV using a JEOL 2100 field emission gun TEM. The resulting HR TEM images were subjected to a fast Fourier transform (FFT) analysis, so as to get the diffraction patterns. Several d-spacing values were measured from the HR TEM images using commercial software ImageJ. Phase structure can be identified by the obtained diffraction patterns and D-spacing. EDX quantification was using the mode of scanning TEM (STEM)-EDX, using a 0.6 nm diameter electron beam.

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3. Results
3.1. As-bonded Copper Bonds on Aluminum Metallization
Bonds were subject to the thermal and electrical impact during operation service, especially at the interface. Most of the device fails as a result from the reliability eliminated by the bonding joints [1,2]. Fig. 5 shows typical copper ball bonds on aluminum metallization on an IC chip after thermosonic bonding interconnection with optimized parameters in this study. From the bond appearance, the bonds were well formed and there is no evident of cracking. The failure in the shear test typically occurs in the mashed copper ball rather than at the interface between the ball and the aluminum metallization pad (see Fig. 6); it typically occurs as a neck break in the pull test (see Fig. 7), indicating that the copper ball and aluminum metallization are successfully bonded together with a high bonding strength for above optimized process parameters shown in Table 2.

Fig. 5. SEM image of typical copper wire ball bonds on aluminum metallization on an IC by thermosonic bonding

Fig. 6. Failure position in mashed copper balls after shear test showing the copper ball strongly bonded on the aluminum pad (average shear force 145 gf)

At present, it remains unclear as to how a strong bond can be formed within such a short time (e.g. 25 ms), and whether melting or sufficiently high temperature may cause IMCs or interdiffusion region at the interface leading to a strong interconnection. In order to ascertain the interfacial metallurgical characteristics of as-bonded Cu-Al bonds, high-resolution HRSEM and HRTEM with EDS are employed to provide the details at the interface.

3.2. Nano-scale interfacial morphology of Cu-Al bonds
Copper ball was successfully bonded on aluminum metallization with the optimized bonding parameters shown in Table 1. The typical interfacial morphology by SE SEM was shown in Fig. 8. The details of the highlighted area A and B in Fig. 8 are given in Fig. 9 and Fig. 10 by HRSEM, respectively.

Fig. 7. Failure position in necks after pull test confirmed good bondability (average pull force 41 gf)

Fig. 8. Secondary electron micrograph of the Cu-Al interface formed with optimized parameters shown in Table 1

Fig. 9. Details of the highlighted area A in Fig. 8 showing some reactants were formed at some regions - not entire regions - of the Cu-Al interface.

Fig. 9 shows that some reactants formed at some regions of the interface (e.g. area A in Fig. 9), although they did not exist in some regions (e.g. area B in Fig. 9). These reactants suggested metallurgical bonding. However, it was difficult for HRSEM to determine weather it is Cu-Al intermetallics, solid-solution or contaminants. Identification of these reactants is needed by HRTEM-FFT as discussed in the following.

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paragraphs. It is unclear if the metallurgical bonding is present at the interface where no such reactants were formed. It is expected that observing the nano features of such interface using HRTEM may provide an insight.

Fig. 10 illustrates that some micro-cracks were formed at the edge of the bonding interface. It was interesting that the crack was in some extent inside of aluminum and copper, as shown in area A and B in Fig 10, respectively. This suggested that metallurgical bonding also exists at the edge of the interface, and crack was formed after the Cu-Al metallurgical joining due to the long-drawn ultrasonic vibration.

Fig. 10. Details of the highlighted area B in Fig. 8 showing micro-crack present at the edge, which was in some extent inside of aluminum and copper.

Fig. 11 is a bright field (BF) diffraction contrast TEM micrograph of the interface region of a bond between a copper ball and aluminum metallization. The difference between the atomic number of Cu and Al results in large contrast changes between the Al and the Cu, and in the black contrast from the Cu. Two typical features were found at the interface, as shown in region A and region B in Fig. 11, respectively. Both of these features will be discussed in details subsequently using HRTEM.

Fig. 11. BF TEM micrograph presenting the morphology of the interface region of a Cu-Al bond. The Al has a white contrast, due to it's relatively low mass-thickness compared to Cu.

The details of region A in Fig. 11 were shown in Fig. 12. It is clearly shown that several layers were presented between the copper and aluminum. Layer A and layer B were amorphous phase and layer 3 was crystalline phase. In order to obtain the composition of these layers, STEM-EDX was conducted, and results were presented in Table 2, which indicated that layer A was aluminum oxide and layer B consists of aluminum-rich amorphous structure. The amorphous aluminum oxide layer (approximately 10-15 nm) could be the initial layer on the surface of the aluminum.

Fig. 12. Details of the highlighted region A in Fig. 11 showing multi-layers at the Cu-Al interface.

<table>
<thead>
<tr>
<th>Spots</th>
<th>OK Aᵀ%</th>
<th>AlK Aᵀ%</th>
<th>CuK Aᵀ%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4.8</td>
<td>93.1</td>
<td>2.1</td>
</tr>
<tr>
<td>2</td>
<td>11.2</td>
<td>87.4</td>
<td>1.4</td>
</tr>
<tr>
<td>3</td>
<td>55.3</td>
<td>43.0</td>
<td>1.8</td>
</tr>
<tr>
<td>4</td>
<td>42.5</td>
<td>50.1</td>
<td>7.4</td>
</tr>
<tr>
<td>5</td>
<td>46.5</td>
<td>48.0</td>
<td>5.4</td>
</tr>
<tr>
<td>6</td>
<td>44.8</td>
<td>11.7</td>
<td>43.5</td>
</tr>
<tr>
<td>7</td>
<td>6.3</td>
<td>3.3</td>
<td>90.4</td>
</tr>
</tbody>
</table>

Table 2 EDS results of spots 1-7 in Fig. 12.

Fig. 13 shows the details of region A in Fig. 12. The diffraction pattern and lattice orientation of region A and region B were reconstructed from HR TEM image using FFT, as shown on the top right corner and top left corner, respectively, in Fig. 13. The values of spacing-d were measured using image analysis software ImageJ, and measured d-spacing = 0.209 nm, 0.210 nm and 0.181 nm for region A and d-spacing = 0.269 nm, 0.260 nm and 0.231 nm for region B. Thus, the crystalline phase in region A and region B were identified to be copper (lattice constant $a = 0.362$ nm) and copper oxide (i.e. CuO) (lattice constants $a = 0.465$ nm, $b = 0.341$ nm and $c = 0.511$ nm), respectively. Those results were consistent with that obtained by STEM-EDX as shown in Table 2. The copper oxide was thought to be the initial layer formed during copper ball formation when copper was melted by electronic sparking. This copper oxide grew preferentially along the [0 0 1] and was approximate 3 nm in thickness. It was surprising that the amorphous

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aluminum oxide layer and copper oxide layer connected very well at atomic scale.

Fig. 13. Bright field HRTEM and lattice images of interface between amorphous aluminum oxide layer and crystalline phase. It was confirmed region A was copper and region B was CuO.

The details of another feature at the interface in region B of Fig. 11, was analyzed by HRTEM, the results are shown in Fig. 14. The thick amorphous aluminum oxide layer did not exist; in such a case, one approximate 20 nm crystalline layer, labeled as A, was found between the copper and aluminum.

Fig. 14. Details of region B in Fig 11 showing an approx. 20 nm crystalline layer present between the aluminum and copper, and the thick amorphous aluminum oxide disappeared.

Further study on region D of Fig 14 by HRTEM, the results are shown in Fig. 15. Analysis of this HRTEM image and FFT reconstruction on region A confirmed the formation of CuAl$_2$, with lattice constants $a$ = 0.6067 nm and $c$ = 0.4877 nm, that grew preferentially along the [2 1 0]. The diffraction pattern and lattice orientation was shown on the top right corner in Fig. 15, and measure d-spacing = 0.239 nm, 0.230 nm and 0.229 nm. Region B has the same diffraction pattern as region A, and measured d-spacing were 0.242 nm, 0.233 nm and 0.231 nm, thus this represents the CuAl$_2$ in region B with a possible slightly aluminum solute.

Fig. 15. Zoom-in image of region D in Fig. 14. HRTEM-FFT confirmed the crystalline phase between aluminum and copper was CuAl$_2$.

4. Discussion

Strong Cu-Al bond was made through thermosonic bonding process which was implemented within 25 ms with the application of ultrasonic vibration for each bond. HRSEM provided overview of interfacial morphology of as-bonded Cu-Al bonds. Further study by HRTEM elaborated the nano-features of Cu-Al interface. Two typical locations were found at the interface, as shown in Fig. 12 and Fig. 14, respectively. One consists of aluminum oxide layer and copper oxide layer, another one was free of oxide layer. As aluminum oxide and copper oxide were the initial layers at the surface of aluminum and copper, respectively, it suggested that the ultrasonic vibration swept some regions of the interface, while in some other regions, the oxides remained there.

The real contacting surfaces are rough [14], thus, the wear is preferential at the tips of asperities. In the valley of the surface, aluminum oxide and copper oxide can remain. However, we found aluminum oxide and copper oxide still connected at atomic scale.

At the regions where oxides were cleaned by ultrasonic vibration, a layer of Cu-Al intermetallics was formed and it was confirmed that this first nucleation of intermetallics was CuAl$_2$. The thickness of this intermetallics layer was approximate 20 nm. Previous studies on the Cu-Al bonds showed that no Cu-Al intermetallics formed at the as-bonded state [6], because of the small size of the intermetallics, which have been observed in our study by HR TEM and at the relative high resolution.

It has been reported that the Cu-Al intermetallics growth at the Cu-Al bond followed the parabolic law during thermal aging [5, 12]. Assuming that Cu-Al intermetallics growth during bonding process was diffusion controlled, obeying by

$$x = (Dt)^{1/2}$$

$$D = D_0 \exp(-\frac{Q}{RT})$$

where $x$ is a IMC thickness at time $t$, $D$ is the growth rate constant, $D_0$ is a pre-factor, $Q$ is the activation energy, $R =$ 2008 10th Electronics Packaging Technology Conference
8.314 J/(K×mol) is the Boltzmann constant, and \( T \) is the absolute temperature.

Literature data [5] for Cu-Al intermetallics growth lead to \( Q = 97.1 \) kJ/mol; with \( x = 20 \) nm and \( t = 0.025 \) s for bonding process, the interfacial temperature can be calculated to be 465 °C. The previous study of in-situ measurement of the interfacial temperature using K-type thin film thermocouples showed the measured temperature was 320 °C [15], and the authors supposed that the real temperature on some spots or regions should be higher due to the big sensor used (20 μm in width) and the measured temperature was an average temperature. Thus, our calculated temperature of 465 °C for the regions with intermetallics where the tips of asperities is reasonable. Therefore, the ultrasonic vibration not only cleaned the contact surface, but also increased the interfacial temperature, which improved atomic interdiffusion between Cu and Al.

To summarize, thermosonic copper ball bonding mechanism can be described as follows: ultrasonic vibration sweeps aluminum oxide and copper oxide locally to enable Cu/Al contacts with the application of bonding pressure. At such regions where oxide is removed, pure metals expose and contact, and atomic interdiffusion proceeds. The flash temperature at these regions increases up to 465 °C due to the cyclic relative motion induced by ultrasonic vibration. As a result, a layer of approximate 20 nm Cu-Al intermetallics (i.e., CuAl2) are formed at the Cu/Al contact areas. The formation of intermetallics greatly improves bonding strength. At the regions where oxide remains, no solid-solution or intermetallics present, however, the aluminum oxide and copper oxide still remains contact and free of voids or gaps between them.

5. Conclusions

Nano-scale interfacial characteristics of as-bonded Cu-Al bonds were observed by HR TEM which provided the direct evidence of a fundamental mechanism of thermosonic copper ball bonding.

(1) Ultrasonic vibration swept aluminum oxide and copper oxide in some regions of contacting surface; while in other regions, oxides remained.

(2) No voids or gaps were found at the interface, including the regions with and without oxide, except at the edge. At the regions with oxide, aluminum oxide and copper oxide remains connected.

(3) In the cleaned regions, an approximate 20 nm Cu-Al intermetallics was formed. The formation of the intermetallics improved the bonding strength. The first nucleated intermetallics was identified to be CuAl2.

(4) Calculation of interfacial temperature showed that the ultrasonic vibration increased the flash temperature at the regions without oxide up to 465 °C which improved the diffusivity for the formation of intermetallics.

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