
Jianbiao Pan, Member, IEEE, Robert M. Paechek, Frank F. Judd, and Jason B. Baxter

Abstract—Gold metallization on wafer substrates for wire/ribbon bond applications requires good bond strength to the substrate without weakening the wire/ribbon. This paper compares the ribbon bondability of Cr–Au and Ti–TiN–Pt–Au metallization systems for an optoelectronic application. Both Chromium and Titanium are used to promote adhesion between semiconductor substrates and sputtered gold films. However, both will be oxidized if they diffuse to the gold surface and result in the degradation of the wire/ribbon bondability. Restoring bondability by ceric ammonium nitrate (CAN) etch was investigated. Experiments were conducted to investigate the effect of Cr–Au and Ti–TiN–Pt–Au, annealing, and CAN etch processes, on 25.4 × 254 μm (1 × 10 mil) ribbon bonding. All bonds were evaluated by noting pull strengths and examining specific failure modes. The results show that there is no significant difference in bondability between Cr–Au and Ti–TiN–Pt–Au before the annealing process. At this point, excellent bond strength can be achieved. However, wire/ribbon bondability of Cr–Au degraded after the wafers were annealed. The experimental results also show that a CAN etch can remove Cr oxide, and that the improvement in wire/ribbon bondability of Cr–Au depends on the CAN etch time. It is further demonstrated that the same annealing process does not have a significant effect on the bondability of Ti–TiN–Pt–Au metallization on the same type substrate materials. Auger electron spectroscopy was used to investigate the causes of the difference in bondability between these two metallizations.

Index Terms—Chromium, gold, metallization, ribbon bonding, titanium, wire bonding.

I. INTRODUCTION

Future developments in silicon waveguide devices such as thermo-optics require multilevel metallization in addition to the deposition of doped silica layers. There are several available metallization schemes under consideration that could meet the requirements for resistance and stability. The metallization systems investigated in this paper were Chromium–Gold (Cr–Au) and Titanium–Titanium Nitride–Platinum/Gold (Ti–TiN–Pt–Au).

In the Cr–Au metallization scheme, Cr is used to promote adhesion between the semiconductor substrate and the sputtered gold film because the gold generally will not form a mechanical bond with the substrate. However, Cr can diffuse through grain boundary up to the gold surface after annealing and then oxidize to form Cr₂O₃ [1]. The Cr₂O₃ on Au surface will degrade wire/ribbon bondability. For example, thermocompression bondability of 3-μm-thick Au films degraded after heating Cr–Au films for 2 h at 250 °C [2] or for 2 h at 300 °C [3]. Higher temperature can increase Cr diffusion speed and reduce the diffusion time to Au surface significantly [1]. It has been shown that these bonding problems can be eliminated either by etching the gold surface using KI + I₂ so that the attached oxide is removed, or by etching with ceric ammonium nitrate (CAN) to remove the oxides without affecting the gold [3].

Titanium (Ti) is often used as a glue layer between the semiconductor substrate and a conductor layer such as Au [4]. In the Ti–Au bilayer thin film, interdiffusion will happen and form TiAu₂, TiAu, and Ti₃Au during annealing at over 250 °C [5]. To prevent the interdiffusion between Ti and Au, a third metal Platinum (Pt) or Palladium (Pd) is commonly used as a barrier layer [4], [6]. In the Ti–Pt–Au metallization scheme, however, Ti can diffuse to the Pt layer through grain boundaries as well. The diffusion process model proposed by Tisone and Drabek [4] in which Ti diffuses along Pt grain boundaries and forms TiPt intermetallic compounds was verified by Garceau et al. [7]. Garceau et al. [7] further showed that the incorporation of a thin (150 Å) layer of Titanium Nitride (TiN) between the Ti (850 to 1000 Å) and Pt (1500 to 2000 Å) layers could inhibit the formation of TiPt intermetallic compounds effectively. Thompson et al. [8] reported the bondability problems associated with Ti–Pt–Au metallization. Based on the authors’ knowledge, no study was yet published on the Ti–TiN–Pt–Au bondability.

The purpose of this study was to compare the ribbon bondability of Cr–Au and Ti–TiN–Pt–Au metallization systems for an optoelectronic application. The specific metallization systems were 2000/6000 Cr–Au (2000 Å of Chromium and 6000 Å of Gold on the wafer surface), and 1000/250/2000/6000 Ti–TiN–Pt–Au (1000 Å of Titanium, 250 Å of Titanium Nitride, 2000 Å of Platinum, and 6000 Å of Gold on the wafer).

II. EXPERIMENTAL DESIGN AND ANALYSIS

Two experiments were designed to examine the effects of these two metallization schemes on ribbon bondability. The first experiment focuses on metallization without annealing.
The second experiment includes effects of the annealing and the CAN etch. Samples for both experiments were prepared by dicing wafers and mounting them on a suitable metal package with epoxy so that the wafers were clamped well during bonding.

The bondability was based on evaluations using gold ribbon 25.4 μm (1 mil) thick and 254 μm (10 mil) wide. The bondability of wide gold ribbon was believed to be more sensitive to the surface contamination than a round 25.4-μm (1-mil) diameter or 33-μm (1.3-mil) diameter wire due to larger bonding area, based on the authors’ experience. Ribbon samples were tested destructively on a Dage pull tester. Pull strengths and failure modes were recorded. The ribbon bonds were made using an automated Palomar 2470V bonder using row bond mode.

### A. Experiment 1

The first experiment was constructed from wafers without annealing. A two factor two level factorial design was selected in this experiment. The samples were split into four cells: Cr–Au, one with 10 min of UV ozone cleaning and one without; and Ti–TiN–Pt–Au, one with 10 min of UV ozone cleaning and one without. The purpose of UV ozone cleaning is to remove the surface contamination from wafer surfaces.

To achieve the highest pull strength, bonding process parameters were optimized and are shown in Table I. Each package was bonded with 100 ribbons. Thermal cycling was used to evaluate the reliability of the ribbon bonds. After thermal cycling (0 °C–85 °C, four cycles), all ribbons were pulled. The thermal cycle parameters of 0 to 85 °C, and four cycles, are representative of typical manufacturing stresses required in the assembly of optoelectronic devices. Pull strengths and failure modes are documented in Table II.

The failure modes indicated in Table II are typical of those seen in developing a good quality ribbon bonding process for optoelectronic devices. The “nonstick” failure occurs when the bonding process is insufficient to achieve adhesion of the ribbon to the substrate surface. When this happens, the ribbon bonder itself, during the act of ribbon tear, lifts the foot of the ribbon off the substrate surface. The “foot lift” failure occurs when the foot of the ribbon is pulled away from the substrate surface during the pull testing, but the ribbon stuck to the substrate surface before the pull test. The “heel break” occurs at the interface between the ribbon foot and the ribbon, and the foot of the wire still remains at the bond pad after destructive pull test. The “ribbon break” is a failure that occurs at any point in the ribbon above the heel during the pull testing. In terms of ribbon bondability performance, heel break and ribbon break are desirable failure modes, while foot lift and nonstick are undesirable failure modes.

Bond pull strength testing was accomplished in accordance with MIL-STD-883E, method 2011.7. Per MIL-STD-883E, the minimum acceptable bond pull strength for a 25.4 μm × 254 μm (1 mil × 10 mil) gold ribbon bonds is 20 g. Since pull strength is known to be loop dependent [1], all loop heights in this study were made at 203 μm (8 mil), and loop lengths were made at 508 μm (20 mil). The two bonds were at the same level on wafer surface and the pull hook was at the center of the loop as shown in Fig. 1. The tensile strength in the first bond and the second bond can be calculated according the following equation [1]:

$$F_1 = F_2 = \frac{F}{2} \sqrt{1 + \left(\frac{d}{2h}\right)^2}$$

where $F_1$ and $F_2$ is the tensile strength at the first bond and second bond, respectively, $F$ is the pull strength at the hook, $h$ is the loop height, and $d$ is the distance between the first bond and the second bond. The loop profile used in this study results in a break strength at the first bond being the same as the break strength at the second bond. These break strengths are about 0.8 times of the pull strength at the hook.

Table II shows that there is no statistically significant difference in ribbon bondability between the Cr–Au and the Ti–TiN–Pt–Au metallization systems without annealing. Moreover, both metallization systems without annealing provide excellent bondability. The average pull strength is more than 18 sigma above the minimum acceptable bond pull strength defined by MIL-STD-883E. Although Table II shows that there were 2% and 1% foot lifts on Cr–Au metallization without UV Ozone cleaning, and UV Ozone cleaning for 10 min, respectively, the bonding process could be adjusted to eliminate this undesired failure mode. Generally speaking, there is a correlation between bond strength and wire/ribbon deformation [9]. Increasing bonding ultrasonic power can deform wire/ribbon more, which tends to make stronger bond between bond foot and the substrate [9]. However, excessively deformed bonds will weaken the heel, a critical region in the bond, and result in low bond strength. The UV ozone clean did not improve

### Table I

<table>
<thead>
<tr>
<th>PROCESS PARAMETERS</th>
<th>1st bond</th>
<th>2nd bond</th>
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<tbody>
<tr>
<td>Force (grams)</td>
<td>120</td>
<td>130</td>
</tr>
<tr>
<td>Time (msec.)</td>
<td>500</td>
<td>500</td>
</tr>
<tr>
<td>Ultrasonic Power</td>
<td>60</td>
<td>60</td>
</tr>
<tr>
<td>Ultrasonic Mode</td>
<td>voltage</td>
<td>voltage</td>
</tr>
<tr>
<td>Ultrasonic Type</td>
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<td>ramp</td>
</tr>
<tr>
<td>Delay time (msec.)</td>
<td>250</td>
<td>250</td>
</tr>
<tr>
<td>Workholder temp.</td>
<td>150°C</td>
<td></td>
</tr>
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### Table II

<table>
<thead>
<tr>
<th>FAILURE MODES AND PULL STRENGTHS FROM EXPERIMENT 1</th>
<th>Pull strength for 100 ribbons</th>
<th>Pull strength for 100 ribbons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Failure mode for 100 ribbons</td>
<td>Heel break</td>
<td>Foot lift</td>
</tr>
<tr>
<td>Without UV cleaning</td>
<td>Cr/Au</td>
<td>93%</td>
</tr>
<tr>
<td>Ti/TiN/Pt/Au</td>
<td>97%</td>
<td>0</td>
</tr>
<tr>
<td>UV cleaning</td>
<td>Cr/Au</td>
<td>98%</td>
</tr>
<tr>
<td>10 min</td>
<td>Ti/TiN/Pt/Au</td>
<td>96%</td>
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</table>

![Fig. 1. Ribbon bonds pull test geometry.](image-url)
bonding performance indicating that there was not much contamination on the wafer surface in this experiment.

B. Experiment 2

It is clear from the results of the previous experiment, that both Cr–Au and Ti–TiN–Pt–Au metallization systems are acceptable for a 25.4 μm × 254 μm (1 mil × 10 mil) ribbon bonding process. The purpose of the second experiment was to determine if an additional wafer annealing process has a detrimental effect on ribbon bonding performance. The purpose of the annealing process is to stabilize the metallization system. In this experiment, the annealing process added during wafer fabrication was a 400 °C air bake for 30 min to pronounce the effect.

The second experiment was split into 16 cells as illustrated in Fig. 2. The two metallizations each had one annealed and one nonannealed sample with each of these being with or without CAN etch. After ribbon bonding, half of the bonds were pulled...
immediately and half were pulled after the temperature cycling of 0°C–85°C for four cycles.

The CAN etch time used in this experiment was 1 min. The solution for CAN etch consists of 65%–75% water, 20%–30% ceric ammonium nitride, and 1%–5% acetic acid. In addition, all of the samples received the 10-min UV ozone cleaning after the CAN etch.

All bonding parameters and loop parameters were the same as in Experiment 1 except the ultrasonic power increased from 60 to 65. The purpose of the change was to eliminate the footlift failure mode. Some gold peel-off was observed on annealed wafers with Cr–Au metallization during dicing.

The pull strengths and failure modes are summarized in Table III. The data show that the failure mode for bonds on annealed Cr–Au metallization before the CAN etch was 100% nonstick, which means that no bond was able to be made. Although different bonding parameter combinations were tried to improve bonding performance, nonstick still remained the dominant failure modes on annealed Cr–Au metallization before the CAN etch. Even maximum ultrasonic power that could be offered by the bonder was unable to make bonds stick. CAN etch improved bondability slightly and switched the dominant failure mode from nonstick to foot lift.

C. Data Analysis

The data were analyzed using analysis of variance (ANOVA). The ANOVA Table for the average pull strength is shown in Table IV. It shows that metallization and annealing have statistically significant effects on the mean pull strength and strong interaction exists between metallization and annealing. The interaction plot of metallization and annealing in Fig. 3 shows that annealing has significant effect on Cr–Au metallization, but not for Ti–TiN–Pt–Au metallization. The bondability of Cr–Au degraded dramatically after the wafers were annealed. The ANOVA analysis in Table IV also shows that CAN etch and temperature cycles do not have statistically significant effect on mean pull strength. There is interaction between annealing and CAN etch at 90% confidence level. The interaction plot of annealed and CAN etch is shown in Fig. 4. Table III shows that CAN etch improved the ribbon bond pull strength on annealed Cr–Au metallization. However, the failure mode is 80% foot-lift and 20% nonstick, which is not acceptable for good
To further investigate what caused the bondability difference between these two metallizations, auger electron spectroscopy (AES) was used to analyze all wafer surfaces. The AES surface survey for annealed no CAN-etched Cr–Au wafer is illustrated in Fig. 5 and that for annealed CAN-etched Cr–Au wafer is illustrated in Fig. 6. It is clearly shown that significant Cr was present on annealed no CAN-etched wafer and the Cr contamination level was reduced after CAN etch. The AES surface analysis results are summarized in Table V. Chromium was detected on annealed Cr–Au metallization, and the chromium–gold ratio decreased from 3.7 to 0.3 after CAN etch for 1 min. Panousis and Bonham [2] reported that no chromium was detected after CAN etch for 10 min. This implies that the CAN etch time in this

<table>
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<th>Metallization</th>
<th>Annealed</th>
<th>CAN etched</th>
<th>Elements detected except C, O, N, Au</th>
<th>Cr/Au ratio</th>
<th>C/Au ratio</th>
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<tr>
<td>Cr/Au</td>
<td>No</td>
<td>No</td>
<td>None</td>
<td>--</td>
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</tr>
<tr>
<td>Cr/Au</td>
<td>No</td>
<td>Yes</td>
<td>None</td>
<td>--</td>
<td>2.5</td>
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<tr>
<td>Cr/Au</td>
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<td>Cr</td>
<td>3.7</td>
<td>7.8</td>
</tr>
<tr>
<td>Cr/Au</td>
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<td>Yes</td>
<td>Cr</td>
<td>0.3</td>
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<tr>
<td>Ti/TiN/Pt/Au</td>
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<td>No</td>
<td>None</td>
<td>--</td>
<td>3.2</td>
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<tr>
<td>Ti/TiN/Pt/Au</td>
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<td>Yes</td>
<td>None</td>
<td>--</td>
<td>4.0</td>
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<tr>
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<td>No</td>
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<td>--</td>
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Fig. 7. AES profile for annealed no CAN-etched Cr–Au.

Fig. 8. AES profile for no annealed no CAN-etched Cr–Au.
experiment was not long enough to eliminate the surface Cr contamination. Increasing etch time should remove all chromium and chromium oxide on the gold surface. It was expected that the amount of chromium diffusing on the gold surface would increase with higher annealing temperature, longer annealing time, and thinner gold thickness. Therefore, CAN etch time, for manufacturing purposes, should be determined according to the annealing process (annealing temperature and annealing time) and the gold thickness.

To find the CAN etch time, it is necessary to know the diffusion rate. An understanding of the diffusion mechanism is important to determine the diffusion rate. The possible mechanisms of Cr diffusion to Au surface include grain boundary, dislocation pipes, bulk diffusion, and surface diffusion [10]. Thomas and Hass [10] reported that the diffusion rate for Cr into Au films in thin-film system, where dislocation pipes is the dominant diffusion mechanism, depends on the Au film deposition rates. Huang et al. [11] observed Cr diffused markedly into Au layer when annealed at 200 °C for 30 min or 250 °C for 5 min. The specific metallization they used was 110-nm-thick Au on 20-nm-thick Cr layer. The mechanism of Cr diffusion was not documented in their paper. Weinman et al. [12] derived the grain boundary diffusion coefficient for Cr in Au at 300 °C is $4 \times 10^{-15}$ cm$^2$s$^{-1}$. However, the grain boundary diffusion coefficient for Cr in Au at 400 °C could not be found from published literature.

To determine why Au peeled off from annealed Cr–Au wafers during dicing, AES was used to do ion sputtered depth profiles, which is an analytical technique capable of identifying elemental concentration profiles. Ion sputtered depth profile of annealed no CAN-etched Cr–Au wafer is shown in Fig. 7 and that of nonannealed no CAN-etched Cr–Au wafer is shown in Fig. 8. Note that Au was found in the Cr layer (1.5–2 units of Au and 10 units of Cr in Fig. 7) of the annealed Cr–Au metallization and little Au was found in Cr layer (0.5 units of Au and 10 units of Cr in Fig. 8) of nonannealed samples. For comparison, ion sputtered depth profiles of annealed CAN-etched Cr–Au is shown in Fig. 9. The function of chromium in the metallization is to promote adhesion between the substrate and the sputtered gold film. The fact that Au diffused to the Cr layer may be the reason that Au peeled off during dicing.

Another possibility that caused Au peeled off could be the chromium oxides under the gold results in a weakened adhesion of the Au film. Since the annealing process was performed
in the air, oxygen can diffuse into the Au layer and form \( \text{Cr}_2\text{O}_3 \) under the Au layer. Weinman et al. [12] detected the presence of oxygen throughout the gold layer using a depth profile and confirmed the Cr remaining in the grain boundaries was oxidized using X-ray photoelectron spectroscopy (XPS). The metallization they used was 200 Å of chromium on 1000 Å of gold.

The C/Au ratio is one of the ways to determine the organic contamination level on Au surface. The higher the C/Au ratio means more organic contamination on gold surface. Table V shows that the no-annealed Ti–TiN–Pt–Au wafers have higher C/Au ratio than that with annealed. This could explain why the no-annealed Ti–TiN–Pt–Au wafers have more foot lift bonds than the annealed ones.

Ti was not detected either on nonannealed or on annealed samples. That means that Ti diffusion was blocked by the TiN layer or the Pt layer. Fig. 10 shows the Ion sputtered depth profile of annealed no CAN-etched Ti–TiN–Pt–Au. That is why annealing at 400 °C for 30 min does not degrade ribbon bondability of Ti–TiN–Pt–Au metallization.

IV. CONCLUSION

The following conclusions can be drawn based on this study.

1) Without annealing, both Cr–Au and Ti–TiN–Pt–Au metallization systems have good wire/ribbon bondability.

2) The ribbon bonding performance degraded dramatically on Cr–Au wafers after they were annealed at 400 °C for 30 min. CAN etch for 1 min partially removed Cr oxide on Au surfaces. A longer time could restore the bondability for Cr–Au metallization.

3) Annealing at 400 °C for 30 min does not degrade ribbon bondability of Ti–TiN–Pt–Au metallization.

4) Temperature cycling from 0 °C to 85 °C with four cycles does not degrade Au wire/ribbon to Au metallization bonds.

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REFERENCES


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