Investigation of High-Strength Electroformed Ni for Microprobes

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We have developed a microprobe that achieves low contact resistance under low contact force only for gold pads. However, in the case of Al pads, an oxide layer formed on the aluminum pad surface obstructs stable contacting, so higher contact force with a strong probe is required. The present study attempts to enhance the strength of the probe material by improving its mechanical properties. It is said that grain downsizing, functionally alloying, or impurity addition can increase material strength. Our study has adopted impurity addition to the electroforming bath because the process can be controlled. Thus, high-strength electroformed Ni has successfully been obtained. Improved Ni has a high Vickers hardness of Hv600 compared with Hv450 for conventional nickel, and a high Young’s modulus of $E = 200\ \text{GPa}$ compared with $E = 150\ \text{GPa}$ for conventional nickel.

**Key Words:** Electroforming, Sodium Allylsulfonate, Grain Size

1. Introduction

The probe card is a tool used in testing each IC chip on a silicon wafer on which ICs have been formed. The probe card electrically connects each IC chip to the test system. To this end, it comprises a printed circuit board (PCB) to be connected with the test system, and probes to be connected with the external contact terminals (bonding pads) on IC chips, as shown in Fig. 1.

The probe card is expected to satisfy various requirements, such as low and stable contact resistance, high-frequency response, and fine-pitch probes. Of these, the need for finer-pitch probes is especially strong, due to the recent trend toward ever-smaller IC chips, and the resultant decrease in the size of the bonding-pad. With probes produced by conventional mechanical fabrication processing, however, the finest pitch achievable is $35\ \mu\text{m}$. Development of a new type of microprobe has been long awaited.

Against this background, we have been studying the applicability of LIGA micro machining technique(1) to probe-card fabrication. In our past study(2), we successfully fabricated three-dimensional nickel microprobes each extending from tip to beam all at once in one cycle of the LIGA X-ray lithography process. Testing with the experimentally fabricated microprobe has verified its low contact resistance for Au pads. For Al-Cu pads on which an oxide film exists, however, the probe is required to break the film to achieve a low and stable contact resistance, which means the need for contact-force improvement. The probe contact force can be improved by various methods, such as increasing the probe thickness, or increasing Young’s modulus. If the probe thickness is excessively large, however, the tip can deviate to the side when the probe card is moved. Increasing the thickness is therefore impractical. We focused on an approach of increasing the Young’s modulus of the probe material. In the LIGA process, microprobes are fabricated by electroforming. Among various methods available for increasing the strength of electroformed material, such as alloying.
and grain downsizing, the present paper studies bath additives that are considered to be effective in suppressing crystalline precipitation.

2. 3-Dimensional Microprobe

The microprobe has been developed to respond to the recent trend toward smaller IC chips. It is fabricated by forming a probe mold using LIGA X-ray lithography and Si anisotropic etching(3), and electroforming a nickel probe in the mold. Figure 2 shows the steps in the microprobe fabrication process. First, a Si substrate is anisotropically etched to create a slope that defines the probe-mold shape in the Z-axis direction. Next, a thick resist layer is formed over the substrate surface. Then, X-ray lithography is performed to form the probe-mold shape in the X and Y directions. Since this process uses X-rays, it is virtually free from the influence of diffraction, thus enabling accurate three-dimensional, which would be difficult with UV lithography. Figure 3 shows SEM photographs of microprobes experimentally fabricated by the abovementioned process. Each microprobe is 20 µm wide and 50 µm thick, with the tip portion being 180 µm thick. Very fine three-dimensional probes, each having a tip and beam, can successfully be produced in one cycle of the LIGA X-ray lithography process. Using X-ray lithography makes it possible to fabricate microprobes whose beam is 50 µm thick.

Despite the very small overall size, each probe allows tip deflection by 80 µm while maintaining a contact force of 3 mN. With the aim of further improving the probe contact force for Al pads, we studied how to enhance the strength of the probe material. We set a Vickers hardness of HV600 and Young’s modulus of $E = 180$ GPa as the material’s target mechanical property values to be achieved.

3. Experimental Method

3.1 Preparation of electroforming bath

Improving the microprobe contact force without changing its shape and size requires an increase in material strength. Since the microprobe is Ni-electroformed, research was conducted to increase the strength of the electroformed material. The strength of the electroformed material can be increased by alloying, grain downsizing, etc. Our study focused on allylsulfonate sodium, an additive considered to be effective in downsizing grains and controlling crystal orientation(4). For comparison, research was also carried out with sodium saccharin, a commonly used additive. Tables 1 and 2 specify the bath compositions and electroforming conditions for sodium allylsulfonate and saccharin additives, respectively.

3.2 Method for evaluating mechanical properties

For the electroformed Ni layer produced using the abovementioned bath, we measured the hardness and Young’s modulus, major mechanical properties important for microprobes. Although the hardness of the electro-
Table 2  Bath composition and electroforming condition
(saccharin)

| Bath composition                        | Value   |
|-----------------------------------------|---------|
| Ni(II) sulfate hexahydrate (g/L)        | 240     |
| Ni(II) chloride hexahydrate (g/L)       | 45      |
| Boric acid (g/L)                        | 30      |
| Sodium lauryl sulfate (g/L)             | 0.04    |
| Saccharin (g/L)                         | 1       |

| Electroforming condition               | Value   |
|----------------------------------------|---------|
| PH                                     | 4.0     |
| Bath temperature (K)                   | 313     |
| Current density (A/dm³)                | 2       |
| Anode plate                            | Nickel  |

Table 3  Mechanical properties of experimental Ni

| Ni                  | Vickers hardness | Young’s modulus   |
|---------------------|------------------|-------------------|
| (Sodium allylsulfonate) | 610              | 180–200 GPa       |
| (Saccharin)          | 435              | 150–175 GPa       |

3.3 Method for evaluating crystal grain size and orientation

For experimentally produced electroformed Ni material, we evaluated the influence of crystal grain size and orientation on the mechanical properties. In general, the grain size of electroformed Ni material enhanced in strength by adding an impurity to the bath is so small (several tens of nanometers) that it is difficult to measure under an ordinary microscope. We therefore used X-ray diffraction analysis, since it can measure the grain size and preferred orientation simultaneously. Electroformed Ni samples were prepared specially for the purposes of mechanical property evaluation. The Rigaku RINT-1500 X-ray diffractometer was used as the analytical equipment. From the analysis results, the grain size was calculated using the Scherer equation.

4. Experimental Results and Discussion

4.1 Vickers hardness and Young’s modulus

The hardness of each evaluation sample was measured using a micro-Vickers hardness testing machine (HM-114, Akashi). Young’s modulus was measured by the method described below, using the test system shown in Fig. 6. The evaluation sample was fixed at its wider end to a carbide jig and secured to the test system so that the sample was cantilevered. A displacement load was applied to this sample while the reaction force was measured. Young’s modulus was calculated from the results obtained. To ensure an accurate calculation, the sample thickness and distance from the cantilever fixing point to the point of action, measured in advance, were taken into account in the calculation. Table 3 gives the Vickers hardness and Young’s modulus of each sample. Those of the electroformed Ni sample produced with sodium allylsulfonate added to the bath are considerably higher than those of the sample produced with saccharin added to the bath.
clearing the initially set target values.

4.2 Heat resistance of electroformed Ni

In assembling microprobes into a probe card in a later stage, it will be necessary to solder the probes to an intermediate substrate and resin-bond the substrate to a PCB. These processes should involve heating the probes. We therefore evaluated the heat resistance of electroformed Ni material. Each sample prepared for mechanical properties evaluation was heat-treated (annealed) on a hot plate at various temperatures from 373 K to 573 K (ordinary joining temperature) in increments of 50 K, and the Vickers hardness of each sample was measured at each temperature. The annealing duration was set at 1 h, taking into account the bonding-resin cure time. Figure 7 shows the relationship between Vickers hardness and annealing temperature. The hardness of electroformed Ni produced with saccharin added to the bath continues to increase until the annealing temperature reaches 423 K, but thereafter decreases sharply. In contrast, the hardness of electroformed Ni produced with sodium allylsulfonate added to the bath remains mostly constant until the temperature reaches 523 K and thereafter decreases rapidly. This result supports the superiority of sodium-allylsulfonate-using Ni to saccharin-using Ni, in terms of heat resistance, promising freedom in subsequent assembly processing.

4.3 Crystal grain size

The abovementioned results verify the superiority of electroformed Ni produced with sodium allylsulfonate added to the bath, for all of the Vickers hardness, Young’s modulus and heat resistance. Predicting that these superior properties might be due to the crystal grain size and orientation, we measured the crystal grain size and orientation using X-ray diffraction. For comparison purposes, this measurement was also carried out for electroformed Ni produced using additive-free baths. Figure 8 shows the X-ray diffraction patterns of respective electroformed Ni samples. While the electroformed Ni produced using the additive-free bath has (220) preferred orientation, those produced with saccharin and sodium allylsulfonate added to the bath have (200) and (111) preferred orientations, respectively.

From these X-ray diffraction patterns, grain size was calculated using the following Scherer equation:

\[ t = \frac{0.9 \lambda}{B \cos \theta} \]  

where \( t \) is the grain size (nm), \( \lambda \) the wavelength (nm), \( B \) the half bandwidth (rad), and \( \theta \) the diffraction angle.

Table 4 shows the calculated results. The grain sizes of electroformed Ni samples produced using additive-containing baths are smaller than that produced using the additive-free bath, indicating the higher hardness of the former. However, no significant difference was observed in grain size between saccharin-using Ni and sodium-
Table 4  Grain size of various kinds of electroforming Ni

| Additive            | Grain Size (nm) |
|---------------------|-----------------|
| Additive free       | 40              |
| Saccharin           | 15              |
| Sodium allylsulfonate | 18             |

![SEM photographs of fractured surface](image)

allylsulfonate-using Ni, despite a far higher Vickers hardness of the latter, as shown in Table 3. This cannot be explained by the Hall-Petch rule, which states that material hardness increases with a decrease in crystal grain size, since dislocations caused by plastic deformation more readily come in contact with boundaries, thus making it harder for grains to move\(^{6}\). Presumably, some unknown additive factor is working so that sodium-allylsulfonate-using Ni provides less dislocating motion than saccharin-using Ni, resulting in higher hardness despite the same grain size. In addition, the Vickers hardness of sodium-allylsulfonate-using Ni begins to decrease at an annealing temperature of 573 K, while that of saccharin-using Ni begins to decrease at 473 K, as shown in Fig. 7. This implies the higher effectiveness of sodium allylsulfonate than saccharin in suppressing grain growth.

To observe the grain growth status due to annealing temperature, each Ni sample annealed at 1 h was split by bending, and analyzed the fractured surface using scanning electron microscopy (SEM). Figure 9 shows the results.

The SEM analysis revealed the following: (1) when the annealing temperature is 573 K, crystal grains are obviously coarser in the saccharin-using Ni than in sodium-allylsulfonate-using Ni (c, f). These results confirm that sodium allylsulfonate added to the electroforming bath is effective in suppressing crystal grain growth during the annealing process.

From the X-ray diffraction patterns shown in Fig. 8, it is possible to identify the preferred orientation, but not the definite advantage of sodium-allylsulfonate-using Ni in terms of mechanical properties.

5. Conclusions

The microprobes of a probe card used in testing semiconductor IC chips are fabricated by lithography and Ni-electroforming. To improve the microprobe contact force, we attempted to increase the strength of electroformed Ni by adding an impurity to the bath. Evaluation of experimentally produced electroformed Ni revealed the following:

(1) The electroformed Ni produced with sodium allylsulfonate added to the bath has a Vickers hardness of 610 and Young’s modulus of 180 to 200 GPa, providing superior mechanical properties compared with those of the electroformed Ni produced with sodium saccharin added to the bath.

(2) A heat resistance test conducted on electroformed Ni samples revealed that whereas the Vickers hardness of saccharin-using Ni begins to decrease at an annealing temperature of 473 K, that of sodium-allylsulfonate-using Ni remains virtually constant until the annealing temperature reaches 523 K.

(3) X-ray diffraction analysis for crystal orientation and grain size revealed that sodium-allylsulfonate-using Ni has the (111) preferred orientation. The grain size is mostly the same in the saccharin-using Ni material and sodium-allylsulfonate-using Ni material, ranging between 15 and 18 nm.

(4) The fractured surface of each of these electroformed Ni samples, observed after annealing under the same conditions, shows that sodium allylsulfonate is effective in suppressing crystal grain growth.

In the future, we will apply this high-strength electroformed Ni to microprobes to verify its practical usability.

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