Focused ion beam preparation of microbeams for in situ mechanical analysis of electroplated nanotwinned copper with probe type indenters

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Key words. Chunk lift-out, focused ion beam, in situ, micromechanical testing, nanotwinned copper, P-FIB.

Summary

A site-specific xenon plasma focused ion beam preparation technique for microcantilever samples (1–20 µm width and 1:10 aspect ratio) is presented. The novelty of the methodology is the use of a chunk lift-out onto a clean silicon wafer to facilitate easy access of a low-cost probe type indenter which provides bending force measurement. The lift-out method allows sufficient room for the indenter and a line of sight for the electron beam to enable displacement measurement. An electroplated nanotwinned copper (NTC) was cut to a 3 × 3 × 25 µm microbeam and in situ mechanically tested using the developed technique. It demonstrated measured values of Young’s modulus of 78.7 ± 11 GPa and flow stress of 0.80 ± 0.05 GPa, which is within the ranges reported in the literature.

Introduction

The introduction of twins is a promising method for the strengthening of copper without reducing its ductility or electrical conductivity (Lu et al., 2004; Hasegawa et al., 2015; Cheng et al., 2017). Orientated and nanotwinned copper (NTC) may be more capable of resisting electromigration and is promising for future high-density and/or three-dimensional integrated circuitry packaging (3D-IC) (Chang et al., 2016; Sun et al., 2018a). Key to NTC’s application to 3D-IC is understanding its mechanical properties, thus far NTC has been found to be dependent on twin frequency, strain rate and crystal orientation (Lu et al., 2009; Ye et al., 2012; Kobler et al., 2015).

Micromechanical testing offers the ability to test small structures and features while providing well-defined stress fields (Haque & Saif, 2003; Hemker & Sharpe, 2007; Pantano et al., 2012). From microbend testing flexural modulus, yield strength, flow stress, and fracture toughness can be determined (Di Maio & Roberts, 2005; Motz et al., 2005; Pantano et al., 2012).

Probe type indenters present a specific set of requirements for micromechanical sample preparation. The challenges are to provide space for indenter access and line of sight from the probe tip to the electron beam for displacement measurement. In the case of FIB prepared samples for probe type indenters, corner or edge sample preparation techniques are commonly used (Liu & Flewitt, 2013; Liu et al., 2014; Robertson et al., 2017). Due to the deficiencies of the edge/corner cutting techniques, a xenon plasma FIB (P-FIB) chunk lift-out based methodology is presented which can be used to test areas of interest from any location or multiple locations on a sample and allow batch mechanical testing.

Experimental details

A FEI G4 CXe P-FIB-SEM (FEI, Oregon, USA) instrument at the University of Loughborough characterisation centre was used for the preparation of the microbeam chunk lift-out. The P-FB was equipped with an Easy Lift system and multichem gas injection system. A gallium FIB FEI Nova 600 Nanolab (FEI, Oregon, USA), equipped with a FMT 120, Kleindiek Nanotechnik probe (Kleindiek Nanotechnik, Reutlingen, German) was used to make the force measurements. The experiments were performed through the following procedures. A summary of the steps required is shown in Table 1.
Table 1. Summary table of steps required for microcantilever preparation including accelerating voltage, current and sample tilt (currents may vary dependent on the sputter rate of the material used).

| Step   | Process and patterning function                  | Accelerating voltage | Current  | Sample tilt |
|--------|--------------------------------------------------|----------------------|----------|-------------|
| Step 1 | Pt deposition                                     | 16 keV               | 20 nA    | 52°         |
| Step 2 | Sample side cutting (regular cross-section)       | 30 keV               | 2.5 µA   | 52°         |
| Step 3 | Under cutting (box milling)                       | 30 keV               | 60 nA    | 0°          |
| Step 4.1| Probe attachment (Pt deposition)                 | 30 keV               | 0.1 nA   | 0°          |
| Step 4.2| Chunk retaining cut (box milling)                | 30 keV               | 4 nA     | 0°          |
| Step 5 | Silicon cut to shape                              | 30 keV               | 2.5 µA   | 52°         |
| Step 6.1| Cut cantilever to shape (cleaning cross-section)  | 30 keV               | 2.5 µA–15 nA | 52° ± 1°   |
| Step 6.2| Under cutting the cantilever (cleaning cross-section) | 30 keV               | 2.5 µA–15 nA | −38°       |
| Step 7 | Cut holding position (box milling)                | 30 keV               | 2.5 µA   | 0°          |
| Step 8 | Repeat steps 4.1–4.2 to move the sample to the holding position | 16 keV               | 3 nA     | 52°         |

Step 1: At 52° a pad platinum/carbon composite was deposited at 16 keV with a current density of 30 pA µm⁻² to a thickness of 3 µm. The platinum carbon deposit is used to reduce damage and improve cutting performance. The area of the pad could be scaled depending on the microbeam required, the pad width should be 5–7 times the required width of the beam and pad length is deposited at \( \approx \)1.5 times the required length of the microbeam (see Fig. 1A).

Step 2: A ‘G’ shaped cut was made around the Pt pad consisting of cross sections leaving a tab of material linking the sample to the substrate. Cuts were made at 30 keV and 2.5 µA. Each cut was 20 µm width and 100 µm depth. A schematic representation of this cutting procedure can be seen in Figure 1(B).

Step 3: At 0° tilt, the ion column is incident at 38° to the sample, depending on the size of the chunk required, and propensity for the material to redeposit cutting from both or one side can be conducted to free the chunk. Undercuts were made along the long edges of the chunk with box type cuts of width 5 µm and depth of 100 µm. The sample was rotated, and undercuts repeated. This process resulted in a free-standing chunk with an ‘inverted house’ cross section (see Fig. 2A).

Step 4: The sample was then Pt welded to the EasyLift probe and the tab connecting the chunk to the bulk cut (see Fig. 2B for detail).

Step 5: A freshly fractured piece of silicon is then used as a mounting point for the sample. At the corner of the silicon, a post is machined to receive the sample to allow for undercutting of the microbeam. The chunk can then be docked on the post and Pt welded in a manner similar to that which is used for TEM lift outs.

Step 6: Sample machining to final dimension was completed using the cleaning cross-section at 52° at 30 keV and 2.5 µA (as shown in Fig. 2D). Once the beam is thinned to 1.5 times the required thickness the sample can be undercut. Tilting the stage to -38° allows the ion beam to be parallel to the microbeam, the cantilever can then be machined to the final dimension with concurrently lower ion beam currents, in this case from 2.5 µA to 15 nA. To ensure parallelism of the beam faces it is advised that final thinning be conducted at ±1° to the milling angle.

Step 7: To ensure a sufficient bond between the chunk and Si substrate a ‘holding’ point was milled using a cleaning cross section cut at 0° tilt 30 keV 2.5 µA. The cantilever was placed in the holding station in a similar manner to step 5. Thick (3 µm) Platinum welds were used to ensure the sample
was fixed to the silicon substrate. The easy lift probe was cut from the sample. The final machined beam is shown in Figure 2(E).

Bend testing was completed inside the Ga+ FIB due to the additional pole piece working distance. The cantilever and probe were aligned. The Y stage of the microscope was scripted to index in 0.1 μm steps towards the indenter bending the cantilever. A video and time stamped images were recorded to link force and displacement.

Following microbend testing an in-plane transmission electron microscope lift-out was prepared from the deformed region including base of the cantilever.

Results and discussion

The microcantilever fabricated had a width of 3.45 ± 0.05 μm, height of 3.72 ± 0.15 μm and length of 25.0 μm. The microbeam was found to significantly plastically deform (see Figs. 3A–D).

No deflection of the base material relative to the silicon substrate was measured indicating the Pt welds held the sample sufficiently. Based on the images and force-time data recorded a force-displacement plot was produced (see Fig. 3E). Standard square cross-section bending equations were used to determine modulus based on the elastic portion of the force deflection graph 17. The Young’s modulus of the copper nanotwins was found to be 78.7 ± 11 GPa (error calculated based on cross section variation). The Young’s modulus of copper at the macro scale is widely accepted as 117 GPa; however, depending on the crystallographic orientation of individual grains, the copper’s modulus can vary from 59–77 GPa in the [001] direction up to 160–202 GPa in the [110] direction (Armstrong et al., 2009; Ye et al., 2012).

The twins in the sample were on the (111) plane with a frequency ranging from several nm to 100 nm with a mean frequency of 22 nm (see Fig. 4B) (Sun et al., 2018a, Sun et al., 2018b). The bend test was conducted parallel to the (111) plane, and the cantilever was polycrystalline as such no single value for the crystal orientation tested can be determined. Following testing the tensile face exhibited cupping deformation (see Fig. 4A). This cupping deformation demonstrates the high ductility of the copper nanotwin material. The flow stress of the copper microbeam was calculated to be 0.80 ± 0.06 GPa based on the methodology presented by Motz et al. (2005). The estimated flow stress of the nanotwinned copper is higher than those reported for equiaxed copper microbeams of a comparable size (3.5 × 3.5 μm) in the literature, being 0.334 GPa 13. This increase in flow stress can be attributed to the strengthening effects of the twins in the sample blocking dislocation movement. STEM images from the ion beam milled tensile face of the cantilever do not show any significant modification, or amorphisation of the surface (Fig. 4D). Thus, it is expected that any modification in mechanical properties through FIB machining small when compared to the errors from SEM deflection measurements and force measurements. Motz et al. reported 0.5–1 GPa hardness increases to the first 300 nm of gallium ion implanted copper, such changes were deemed negligible when measuring >1 μm² cantilevers (Motz et al., 2005). Furthermore, xenon ion milling has been shown to produce significantly less
Fig. 3. (A)–(C) The progress of deformation during microbend testing. (D) The plastically deformed copper nanotwinned cantilever. (E) The force deflection plot from the bend test.

Fig. 4. (A) An electron beam image recorded at 45° tilt showing the cupping deformation of the tensile face of the microbeam. (B) Cross section bright field scanning transmission electron micrograph of the deformed copper cantilever. (C) A magnified section from image (B) of the tensile ion beam machined face, no clear modification of the surface of the cantilever is observable.

amorphisation when compared to gallium ion milling (Kelley et al., 2013).

Conclusion
An efficient site-specific chunk lift-out methodology has been presented and applied to copper nanotwins, and a 3 × 3 × 25 µm beam cut parallel to the growth direction of the electroplated nanotwinned copper was produced. The microbeam was bend tested, the flexural modulus was found to be 78.7 ± 11 GPa, and the flow stress was 0.80 ± 0.06 GPa. The chunk lift-out methodology was ideal for site specific micromechanical testing with a probe type indenter.

Acknowledgements
The authors thank EPSRC EP/R032203/1 for funding and acknowledge the Loughborough Materials Characterisation Centre for access to the Helios PFIB, funded by EPSRC grant EP/P030599/1.

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