Fracture and Delamination of Chromium Thin Films on Polymer Substrates

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New emerging technologies in the field of flexible electronic devices require that metal films adhere well and flex with polymer substrates. Common thin film materials used for these applications include copper (Cu) with an adherence interlayer of chromium (Cr). Copper can be quite ductile and easily move with the polymer substrate. However, Cr is more brittle and fractures at lower strains than Cu. This study aims to examine the fracture and subsequent buckling and delamination of strained Cr films on polyimide (PI). In-situ scanning electron microscope (SEM) straining is used to systematically study the influence of film thickness on fracture and buckling strains. Film fracture and delamination depend on film thickness, and increases in crack and buckle density with decreasing thickness are explored by a shear lag model.

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1. INTRODUCTION

Over the past few years, metal and ceramic films on polymer substrates have led to the emergence of deformable electronics, such as foldable displays, sensor skins, and electronic textiles. The typical route to create deformable devices on polymer substrates is to fabricate stiff islands, commonly made with silicon-nitride, where active cells (thin film transistors) are fabricated and connected by metal lines. These interconnect lines must deform with the substrate and survive strains greater than 10 pct while maintaining electrical conductivity. Several studies have examined the tensile strength of blanket copper (Cu), aluminum, and gold films on polymer substrates as well as gold lines. Further research on ceramic coatings on polymers for barrier gas coatings has also been completed. Compared to bulk metals, these films on polymer substrates can have small rupture strains (less than 2 pct) but high strengths. However, when the film is well bonded to the substrate, it will have high rupture strains due to the lack of localized deformation that is suppressed by the strong adhesion. When the film is brittle like silicon nitride, no plastic elongation takes place and fracture is caused by the breaking of an array of atomic bonds. Failure of the films by cleavage fracture is similar to previous studies of ceramic films on metal substrates pulled in tension.

With the periodic cracking method that is based on a shear lag analysis, the maximum shear traction that is supported by the interface can be calculated. The periodic cracking method was first developed to study the shear properties of metal-ceramic interfaces, with similar periodic cracking of other material systems having also been studied. With the metal-ceramic system, plastic deformation in the metal substrate is accommodated by elastic deformation of the ceramic film until the fracture stress is reached. With metal-polymer interfaces, the plastic deformation of the metal is accommodated by the elastic deformation of the polymer substrates until the fracture stress of the metal film is reached. At fracture, the in-plane traction in the film is transferred as shear traction to the interface adjacent to the free edge of the crack. The shortest distance between cracks is the characteristic relaxation distance of the shear traction. This leads to defining the maximum shear traction supported by the interface, \( \tau_{\text{max}} \), as a function of the fracture stress of the film, \( \sigma_{\text{frac}} \), as

\[
\tau_{\text{max}} = \frac{3}{4} \frac{h}{\lambda} \sigma_{\text{frac}}
\]

where \( h \) is the film thickness and \( \lambda \) is the mean spacing between cracks. The fracture stress can be estimated with Eq. [2], assuming that the film behaves elastically,

\[
\sigma_{\text{frac}} \approx \frac{E_f \epsilon_{\text{frac}}}{h}
\]

where \( E_f \) is the elastic modulus of the film and \( \epsilon_{\text{frac}} \) is the fracture strain. It should be noted that Eq. [2] does not take into account the residual in-plane stress of the film.
after deposition. It is easily included, as shown by Andersons and Leterrier et al.\textsuperscript{[13–15]} Similar shear lag analyses have been performed with SiO\textsubscript{2}-PET film systems as well as SiN\textsubscript{x}-Kapton film systems.\textsuperscript{[13–15]} However, some differences are found between the preceding approach and others\textsuperscript{[17,18]} but both are based on the shear lag analysis of Kelly and Tyson.\textsuperscript{[20]}

\textit{In-situ} tensile tests performed inside a scanning electron microscope (SEM) allow one to determine the system’s fracture strain, where cracks first initiate, as well as any subsequent buckling and delamination that may occur.\textsuperscript{[19]} This type of experiment can provide a wealth of information that can be used to improve existing adhesion and deformation models currently being implemented\textsuperscript{[11–25]} to predict material failure and lifetimes of the material systems. Current thin film systems being examined for the connecting lines of flexible devices include Cu or Au with an adhesion layer of chromium (Cr). It has been well known that Cr can increase the adhesion energy of metal films on polymer substrates, for example, Cu on PI\textsuperscript{[27]} and Au on PDMS.\textsuperscript{[13]} It has also been observed that Cr adhesion layers (10 to 20 nm in thickness) will fracture before cracks are viewed in the overlying Cu films, and the failure occurs via channel cracking perpendicular to the tensile loading direction.\textsuperscript{[24]} This study aims to examine the fracture behavior of bare Cr films on PI substrates to better understand the mechanisms that lead to failure of this adhesion layer. Of importance is whether fracture strain is influenced by the film thickness.

\section*{II. EXPERIMENTAL PROCEDURE}

Chromium films were deposited onto cleaned 50-\textmu m-thick UPLIEX brand PI films ($E = 8.5$ GPa). The PI film was cleaned by soaking it in a 10 ppt aqueous solution of RBS 50 (a laboratory cleaning concentrate with high pH) for 24 hours at room temperature, followed by a deionized water rinse and a 1 MHz ultrasonic cleaning process. The Cr layers were deposited by e-beam evaporation in a Balzers BAK 550 evaporation machine with the vacuum at $2.10 \times 10^{-7}$ mbar and using a deposition rate of 0.5 nm/s. Four film thicknesses were made and studied: 50, 100, 200, and 500 nm. It is known that adhesion layers are commonly 50 nm or less; however, in order to fully study the failure mechanism, an order of magnitude of film thickness was used. The 100-, 200-, and 500-nm films have residual tensile stresses on the order of 1 GPa (1.2, 1, and 0.8 GPa, for the 100, 200, and 500 nm, respectively), as determined by X-ray diffraction using the sin$^2\Psi$ method. The same method was also used on the 50-nm film but was unsuccessful in measuring a stress because the film was too thin to get a reliable peak shift. In the analysis, a 1-GPa residual stress is assumed for the 50-nm film. Tensile cracks formed before straining experiments of the 200- and 500-nm films.

The structure of the films was determined with electron microscopy techniques. By careful sample preparation, cross-sectional transmission electron microscopy (TEM) images and diffraction patterns of the 50- and 500-nm films were taken. Figure 1 shows a columnar grain structure with peculiar peaks for both film thicknesses. The peaks are also visible in scanning electron microscopy images of the film surfaces (not shown). Diffraction patterns indicate that the films are polycrystalline. The average grain size of the films is on the order of 50 nm for all film thicknesses.

The films were pulled in tension using a small scale tensile device (Kammrath and Weiss, Dortmund, Germany) that has a maximum displacement of 6 mm. The sample geometry used was approximately 1-cm wide with a 3-cm gage length. Each test was carried out using a 10-\textmu m/s displacement rate. The tensile device was used to perform both \textit{ex-situ} and \textit{in-situ} tests inside the SEM. \textit{In-situ} tests were initially stopped every 100 \textmu m of displacement until the first cracks perpendicular to the tensile direction were observed (approximately 600-\textmu m total displacement) in order for SEM images to be taken of the samples at various magnifications. After the first 600 \textmu m of displacement, the experiment was stopped every 200 \textmu m of displacement. The images were then correlated to the strain. Comparisons of \textit{ex-situ} and \textit{in-situ} tests indicate that the electron beam and the vacuum do not alter the results notably.

Fig. 1—Bright-field TEM images of the (a) 50- and (b) 500-nm-thick films. The arrows in (a) indicate regions of possible columnar porosity.
III. RESULTS AND DISCUSSION

The load vs strain behavior of the Cr films on PI is shown in Figure 2(a) for ex-situ experiments. Both ex-situ and in-situ experiments illustrate that there is no pronounced thickness dependence on the load and strain response. This may be explained by the small grain size of 50 nm, which is for all thicknesses the smallest dimension and thus expected to control the mechanical response of the films. The ex-situ samples were all strained to 11 pct, while in-situ tests were performed to 12 to 18 pct strain. With the in-situ experiments, cracks perpendicular to the loading direction were observed to initiate at less than 1 pct strain for all thicknesses (Table I). Measured crack densities increase with decreasing film thickness (Figure 2(b)). The crack density of the thickest film (500 nm) reaches a steady state at about 4 pct strain, which is at a lower strain value than the thinner films. The 100- and 200-nm films reach a steady state at approximately 5 pct strain. Figure 3 illustrates the cracking found in all four thicknesses at 11 pct strain. Crack density as a function of strain of the films 100 nm and greater reaches a clear plateau in the data indicating the steady state. The thinnest film, 50 nm, does not reach such a plateau within the prescribed straining limit of the apparatus. Rather, a change in slope is reached at about 7 pct strain and then continues to gradually increase, up to 18 pct global strain. This indicates that cracks are still forming in the very thin brittle films where cracks stop forming in the thicker films. The continued crack growth could be of significance and may aid in increasing the strain a system can withstand. The average crack spacing at saturation can also be found in Table I.

Due to the fact that the fracture strains of the films can be observed with the in-situ experiments, it is easy to calculate the maximum shear stress of the interface, \( \tau_{\text{max}} \), assuming that the fracture stress can be estimated based on Eq. [2] with an elastic modulus of 280 GPa. Since the grain size is homogenous for the various film thicknesses and too small to allow considerable plasticity, this approximation is justified. There is a wide distribution of observed fracture strains but a trend of decreasing fracture strain with increasing film thickness is found, as shown in Table I. The fracture strain is defined when the first new cracks perpendicular to the tensile direction are observed with the SEM during the in-situ experiment. Using Eq. [1] and the fracture stress values from Eq. [2], the maximum shear stress of the interface also decreases with increasing film thickness, as shown in Table I. The calculated values for the Cr-PI system are lower (119 to 31 MPa) than that of a metal-ceramic interface (1.7 GPa for Cu-SiO\(_2\)) and 0.4 to 0.8 GPa for Cu-Al\(_2\)O\(_3\)), but the values are reasonable. It should be noted that the residual stresses in the thicker films (>100 nm) have not been considered with this model. Rather, the formation of new cracks due to the tensile loading is considered to be the over-riding mechanism. When the residual stress of about 1 GPa of the films is included in the analysis, the fracture stresses range between 3.1 and 1.2 GPa and the maximum interface shear strength values increase to 175 to 93 MPa, respectively, for the 50- to 500-nm film. These values are shown in brackets (i.e., [175]) in Table I. In either case, the data show a tendency for the thinner films to have higher interfacial shear strengths, which is most likely caused by the decrease in fracture stress due to a scaling with the film thickness because the grain size is constant for all thicknesses. A possible explanation could be the presence of columnar porosity, commonly observed with columnar grained films. This type of defect may be present in these films, as shown in Figure 1(a) (arrows) at the column boundaries. Further TEM examination is required in order to say with
certainty how columnar porosity should be taken into account. Larger defects in the thicker films, e.g., cracks caused by residual stresses from deposition, will cause a lower fracture stress and as a consequence the simple model provides only an upper limit of the interfacial shear strength. However, the large crack density and lack of a steady-state regime may be a cause for overlayers of Cu or gold to fail.

Increasing the strain of the film system leads to delamination and buckling at the interface. Buckling is caused by the lateral compressive deformation of the film on the substrate due to the different Poisson’s ratios of Cr and PI (0.21 and 0.3, respectively). This occurs above ~10 pct strain along the tensile direction, with the exact amount of strain depending on the film thickness. The nature of the buckling is also dependent on film thickness. With the 500-nm film, buckles form with a crack at the apex and at the edges of the buckle, forming a triangle buckle, which can lead to immediate spalling of the film. On the other hand, the 50-nm film

### Table 1. Measured and Calculated Values of Fracture Strain and Stress, Average Crack and Buckle Spacing, and Maximum Interface Shear Strength for All Performed In-Situ Experiments; Values that Include the Residual Tensile Stress of 1 GPa are Shown in Brackets

| Film Thickness | Pct Fracture Strain | Fracture Stress (GPa) | Average Crack Spacing (μm) | Maximum Interface Shear Strength (MPa) | Pct Buckle Strain | Average Buckle Spacing (μm) |
|----------------|---------------------|-----------------------|----------------------------|---------------------------------------|------------------|-----------------------------|
| 50 nm          | 0.58                | 1.6 [2.6]*            | 1.9 ± 0.1                  | 100 [163]*                            | 8                | 4.4 ± 1.5                   |
|                | 0.76                | 2.1 [3.1]*            | 2.1 ± 0.2                  | 119 [175]*                            | 13               |                             |
| 100 nm         | 0.32                | 0.9 [2.1]             | 4.1 ± 0.5                  | 52 [120]                              | 13               | 11 ± 6                      |
| 200 nm         | 0.23                | 0.64 [1.6]            | 4.8 ± 0.4                  | 64 [161]                              | 17               | 12 ± 5.9                    |
| 500 nm         | 0.15                | 0.42 [1.2]            | 15.5 ± 0.5                 | 31 [93]                               | 10               | 62 ± 35                     |
|                | 0.19                | 0.54 [1.3]            | 16.7 ± 1                   | 38 [94]                               |                  |                             |

*The residual stress of the 50-nm film could not be measured and was assumed to be 1 GPa and was based on the stress results of the other films.

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![Fig. 3](image-url)

**Fig. 3**—The SEM micrographs at the same magnification of all four films strained to 12 pct. The effect that thickness has on crack spacing is evident as well as the first stages of buckling in the 500-nm film. (a) 50-nm, (b) 100-nm, (c) 200-nm, and (d) 500-nm.
buckles are much smaller and form straight buckles between the initial cracks with a cracked apex as well. Figure 4 shows examples of the buckling with the two film thicknesses. Inspection of the buckling behavior may be a way to determine the work of separation of the film system. Further experiments combined with modeling are currently under way to examine this behavior.

IV. CONCLUSIONS

The failure of blanket chromium films commonly used as adhesion layers to increase the interfacial strength of metal-polymer film systems has been investigated. Under tension, Cr films fracture and delaminate from the PI substrate with a brittle cleavage fracture behavior. Initial crack formation perpendicular to the loading direction occurs at strains less than 1 pct, much lower than the fracture strain of the Cu films whose adhesion is being improved in applications. The cracking behavior can be compared to ceramic films on metal substrates pulled in tension, and a periodic cracking model for calculating the maximum interface shear strength was used. The average interface shear strength is $67 \pm 35 \text{ MPa}$. The spacing of the cracks reaches a steady state at approximately 4 to 5 pct strain for thick films ($\geq 100 \text{ nm}$). For 50-nm-thick films, the crack density does not reach a steady-state region even after 18 pct global strain has been induced. Crack and buckle spacings also increase with increasing film thickness, with buckles occurring at 8 to 17 pct strain depending on film thickness. These buckles can be used to measure adhesion energies by taking the shape and possible cracks of the buckles into account. This study has shown that thin adhesion interlayers continue to form cracks during tensile deformation up to high strains. The continued fracture process and large crack density may cause metal bilayer systems on polymers (e.g., Cu film on Cr adhesion layer) to be less reliable than a thicker adhesion layer with a lower crack density.

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