Self-Heating-Induced Deterioration of Electromechanical Performance in Polymer-Supported Metal Films for Flexible Electronics

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The retention of electrical performance under the combined conditions of mechanical strain and an electrical current is essential for flexible electronics. Here, we report that even below the critical current density required for electromigration, the electrical current can significantly deteriorate the electromechanical performance of metal film/polymer substrate systems. This leads to a loss of stretchability, and this effect becomes more severe with increasing strain as well as increasing current. The local increase of electrical resistance in the metal film caused by damage, such as localized deformations, cracks, etc., locally raises the temperature of the test sample via Joule heating. This weakens the deformation resistance of the polymer substrate, accelerating the necking instability, and consequently leading to a rapid loss of electrical conductivity with strain. To minimize such a current-induced deterioration of the polymer-supported metal films, we develop and demonstrate the feasibility of two methods that enhance the deformation resistance of the polymer substrate at elevated temperatures: increasing the thickness of the polymer substrate, and utilizing a polymer substrate with a high glass transition temperature.

Flexible electronics is an emerging technology that has potential in a wide variety of applications, including flexible displays1–4, artificial skins5, stretchable circuits6, solar cells7, and wearable sensors8. In flexible electronics, most components lie on a compliant polymer substrate and are connected with each other using metallic interconnects. Thus, their functionality and usability rely entirely on the electromechanical performance of the metal film/polymer substrate couple under operational conditions. In this sense, sustaining a large deformation while maintaining a low and constant electrical resistance (i.e., achieving high stretchability) is considered an essential requirement for the implementation of flexible electronics. Considerable effort has been devoted in this regard to date9–15. It is noted, however, that flexible electronic devices are subjected to electrical currents as well as mechanical strain under working conditions. The electrical current can influence the electromechanical performance of the metal film/polymer substrate system in two ways: 1) the loss of conductivity of the metal film by electromigration16–18 and 2) the improvement of the metal film ductility by Joule heating19,20 or electroplasticity21–23. The current density, usually required in state-of-the-art flexible electronic devices, is less than ~5 × 10^4 A cm^-2, somewhat below the critical value required for electromigration24; hence, this effect is expected to be negligible. In addition, considering the high melting temperatures of metal films (e.g., 961 °C for a silver (Ag) film25) the temperature rise caused by Joule heating appears not to have a significant effect on ductility enhancement. For these reasons, the effect of the electrical current on electromechanical behavior has been ignored in the design of metal film/polymer substrate systems. However, the trend toward smaller and more functional and powerful devices demands an increasing current density, together with a much higher stretchability. The high current density will raise the possibility of electromigration under operational conditions. It is known that large deformations in the metal film give rise to inhomogeneities (i.e., localized deformation, formation of cracks, etc.)10,26,27. These will lead
to localized Joule heating, causing a temperature rise in both the metal film and polymer substrate. These points therefore suggest that the electrical current may play an important role in the electromechanical performance of future flexible electronic devices.

In this study, we investigated the electromechanical behavior of Ag film/polyethylene terephthalate (PET) substrate systems under electrical currents. Tensile tests were performed in combination with electrical resistance measurements, inside a scanning electron microscope (SEM) or under an infrared (IR) camera, while applying a constant electrical current. The current-induced deterioration of electromechanical performance was analyzed in a quantitative manner by defining the stretchability of the material system. An underlying mechanism of the deterioration was inferred, focusing on the Joule heating-induced acceleration of the necking instability in the PET substrate. We further suggested a viable solution to prevent (or reduce) the current-induced deterioration of electromechanical performance by using a polymer substrate with a high glass transition temperature, and demonstrated its feasibility with a polyimide (PI) substrate.

Results and Discussion

Electromechanical behavior and its variation under electrical currents. Figure 1 shows the normalized electrical resistance—strain curves of 400 nm Ag/188 μm PET with and without an electrical current (I). The reproducibility was confirmed by employing each test condition more than three times; representative curves are presented. The electrical resistance generally increased with increasing strain, irrespective of the application of an electrical current. Notably, the rates of the resistance increase with strain were distinctly different. When no electrical current was applied, the electrical resistance gradually increased with strain up to ε ~ -40%, indicating that no severe damage (e.g., cracks) developed in the Ag film (refer to Figure S1 in the Supplementary Information for the SEM micrographs showing the surface morphologies of this sample at six strain levels). However, the application of an electrical current caused deviating behavior. Once the divergence between the curves with and without an electrical current started to occur, the extent of deviation became more pronounced with further strain, accelerating the loss of electrical conductivity and consequently leading to an early failure of the sample. The onset strain at which the curves began to deviate decreased with increasing electrical current (ε = 19% for I = 0.1 A; ε = 17% for I = 0.2 A; ε = 10% for I = 0.3 A; ε = 6% for I = 0.4 A; and ε = 3% for I = 0.5 A), and the extent of the deviation became more severe with increasing electrical current. The current has a negligible effect on electromechanical behavior at the early stage of deformation, as shown by the matching curves in Fig. 1 in this region. It is clear, though, that its effect becomes important with increasing strain as well as increasing electrical current, deteriorating the electromechanical performance of the Ag film/PET substrate. The data for the 100 nm Ag/188 μm PET and 200 nm Ag/188 μm PET systems are provided in Figure S2, and consistent results with 400 nm Ag/188 μm PET were also confirmed.

To quantitatively analyze the current-induced deterioration of electromechanical performance, we defined the stretchability of the Ag film/PET substrate as the strain value at which the normalized electrical resistance, R/R₀, reached 1.5, i.e., a 50% increase relative to the initial resistance value. As shown in Fig. 1 and Figure S2, samples subjected to a high electrical current failed mechanically before reaching a value of R/R₀ = 1.5. In these cases, the stretchability was defined as the mechanical failure strain of the sample. Figure 2a shows the variation of stretchability with electrical current. For all three systems with different Ag film thicknesses, the stretchability slowly decreased with increasing strain at low electrical currents, but a steep reduction was observed beyond a certain strain. The higher stretchability of the thicker Ag film without an applied electrical current is consistent with previous studies. The electrical current at which a sharp decline in stretchability occurred was proportional to the Ag film thickness, and the rate of stretchability degradation was more severe in the 100 nm Ag/188 μm PET system than in the 200 nm Ag/188 μm PET and 400 nm Ag/188 μm PET systems. To systematically analyze
the effect of the Ag film thickness on the current-induced deterioration of stretchability, the “stretchability loss” parameter was defined as follows:

$$\text{stretchability loss} = \frac{\text{stretchability}_{[J=0A]} - \text{stretchability}_{[J]} \to \text{stretchability}_{[J=0A]}},$$

where stretchability_{[J=0A]} and stretchability_{[J]} are the stretchability values without and with an electrical current, respectively. The stretchability loss measures the current-induced reduction of stretchability with respect to the reference value without an electrical current. As shown in Fig. 2a, the effect of the Ag film thickness on stretchability loss was negligible at low electrical currents up to ~0.15 A, but it was clearly evident beyond this value. The thinner Ag film showed a pronounced stretchability loss and an increased degradation rate with increasing electrical current. This behavior suggests that the current-induced deterioration of electromechanical performance was accelerated in the thinner Ag film. However, when the stretchability data are analyzed in terms of the current density (J), the opposite results are obtained (Fig. 2b). The current density at which the stretchability started to rapidly decrease was inversely proportional to the Ag film thickness. Additionally, the rate of the stretchability decrease (or the stretchability loss increase) with electrical current was larger in the thicker Ag film. These trends indicate that the current-induced deterioration of electromechanical performance was accelerated in the thicker Ag film. These contrary results appear to arise from the fact that neither the electrical current nor the current density is a suitable parameter for explaining the current-induced deterioration of electromechanical performance. The electrical current does not account for the effect of sample size, particularly the cross-sectional area of the Ag film. The current density is considered a key parameter influencing the electrical performance deterioration of metal film/polymer substrate systems via electromigration. Nevertheless, the SEM images showing the surface morphologies of the tested samples revealed that electromigration does not play a significant role in the present study; a detailed discussion in this regard will be presented later. Another factor that can be considered is the current-induced Joule heating effect. As the temperature rise is proportional to the electrical power applied, we introduced a parameter associated with the electrical power, \(P/t_o\), which is equal to the initial power divided by the factor \(\rho L_o/w_o\). \(\rho\) is the electrical resistivity, and \(L_o\) and \(w_o\) are the length, width, and thickness of the Ag film, respectively, at the unstrained state. Note that \(\rho\) is a constant value (~2 \(\mu\Omega\)-cm in the film thickness range of ~80 to 1,000 nm)\(^\text{34,35}\). \(L_o\) and \(w_o\) are fixed for all samples, and only \(t_o\) varies, indicating that the initial power applied to the sample is directly proportional to \(P/t_o\). The analyzed results are presented in Fig. 2c. Similar to the results obtained from the electrical current and current density measurements, stretchability loss occurred with increasing initial power (i.e., \(P/t_o\)), but it occurred gradually and there was no drastic transition from the slow to steep increase. More importantly, all the data for the three different Ag film thicknesses resulted in a single stretchability loss – \(P/t_o\) curve, indicating that the Ag film thickness does not affect the current-induced deterioration of electromechanical performance. These findings indirectly support that the deterioration of the Ag film/PET substrate system observed in the present study arises from the current-induced Joule heating that leads to a temperature rise in both the film and substrate. The stretchability loss averaged over the three Ag film thicknesses was ~3% at \(P/t_o = 25 A^2\) mm\(^{-1}\) (corresponding to \(J = 2.5 \times 10^4 A\) cm\(^{-2}\) for 400 nm Ag/188 μm PET) and increased to ~6% at \(P/t_o = 100 A^2\) mm\(^{-1}\) (\(J = 5 \times 10^4 A\) cm\(^{-2}\)). It was ~15% at \(P/t_o = 225 A^2\) mm\(^{-1}\) (\(J = 7.5 \times 10^4 A\) cm\(^{-2}\)) and ~40% at \(P/t_o = 400 A^2\) mm\(^{-1}\) (\(J = 10^5 A\) cm\(^{-2}\)). The relationships between \(J, I, P\), and \(P/t_o\) for the samples are given in Table 1. The current density used in state-of-the-art flexible electronic devices is in the range of ~5 \(\times 10^4 A\) cm\(^{-2}\) and electromigration requires the sustained application of a high current density of >10^5 A cm\(^{-2}\) (note that the duration of each test in the present study was shorter than 12 min)\(^\text{34}\). Considering this, our results clearly show that, even below the current density required for electromigration, electrical current can induce the deterioration of electromechanical performance. This effect will eventually impair (or stop) device performance and should thus be carefully considered in the design of metal film/polymer substrate systems used in flexible electronic devices.

A mechanism for the current-induced deterioration of electromechanical performance. As described earlier, the temperature rise in the samples caused by the current-induced Joule heating appears to be responsible for the deterioration of electromechanical performance. To test this hypothesis and understand the
underlying mechanism, we measured the temperature distribution over the sample surface with an IR camera and observed the variation in surface morphology of the tested sample with strain using a SEM. The results for 400 nm Ag/188 μm PET under an electrical current of 0.3 A are presented in Figs 3 and 4. A uniform temperature existed at the center of the sample at low strains of <7%, which spanned ~7 mm or almost 50% of the sample gauge length (Fig. 3b); note that heat transfer from the sample reduced the temperature near the grips. However, a local temperature rise started to occur at ε = 7%, with the temperatures at locations 1 and 3 reaching 48 °C and 46 °C, respectively. This nonuniform temperature distribution became much more pronounced with further strain: the temperatures were 54 °C and 49 °C at locations 1 and 3 for ε = 11%, and increased to 77 °C and 55 °C at ε = 16%. According to the SEM observations (Fig. 4), there were no marked changes in surface morphology up to ε = 5%, but small cracks started to form near location 1 for ε = 7.5%. The formation of these small cracks will have induced a loss of electrical conductivity at location 1, thereby increasing the electrical resistance and leading to localized Joule heating. This interpretation provides a rationale for the local temperature rise at ε = 7%, shown in Fig. 3, and further suggests that the localized cracking in the Ag film is closely associated with the non-uniform temperature distribution. The localized cracking process proceeded further with strain, which caused severe damage around location 1 and enhanced the local rise of temperature at this site, eventually accelerating the mechanical failure of the sample there (refer to the SEM micrographs at ε = 15% in Fig. 4). It is noted that the surface morphologies at locations 2 and 3 remained almost unchanged even at ε = 15%, which supports the observation that the cracking was localized near location 1. Such a current-induced acceleration of the damage process in the sample was also reflected in the electromechanical behavior. As shown in Fig. 1, the electrical resistance–strain curve of I = 0.3 A started to deviate from that collected without an electrical current at ε = ~10%. According to previous studies, oxidation or electromigration in the Ag film could be responsible for the loss of electrical conductivity (i.e., the increase of resistance). However, the SEM micrographs and scanning X-ray photoelectron spectroscopy analysis of the tested sample surface, shown in Figs 4 and S3, revealed no evidence of oxides or electromigration, the latter of which would be indicated by the accumulation of material (hillocks) at the anode and depletion (voids) at the cathode. We confirmed this by measuring the variation in electrical resistance of 400 nm Ag/38 μm PET and 400 nm Ag/188 μm PET with time under an electrical current of 0.3 A in the unstrained condition. As shown in Figure S4, the electrical resistance and temperature in the samples remained almost constant for ~30 min. This indicates that the electrical current itself does not cause any damage. Considering that the time taken for each test was less than ~12 min, we can conclude that the effects of oxidation and electromigration are negligible in the present study.

| $I^2/t_o$ (A²/mm) | 100 nm Ag | 200 nm Ag | 400 nm Ag |
|-------------------|-----------|-----------|-----------|
| I (A)             | J (A/cm²) | I (A)     | J (A/cm²) |
| 25                | 0.05      | 5 × 10⁴   | 0.07      | 3.55 × 10⁴ |
| 100               | 0.1       | 10⁵       | 0.14      | 7.1 × 10⁵ |
| 225               | 0.15      | 1.5 × 10⁵ | 0.21      | 1.06 × 10⁵ |
| 400               | 0.2       | 2 × 10⁵   | 0.28      | 1.42 × 10⁵ |
| 625               | 0.25      | 2.5 × 10⁵ | 0.35      | 1.77 × 10⁵ |
|                   | 10⁶       | 5 × 10⁵   | 0.5       | 1.25 × 10⁵ |

Table 1. The relationships between $I$, $I^2/t_o$ (=initial power/[ρL_o/w_o]) in the samples.

Figure 3. Temperature distribution of 400 nm Ag/188 μm PET during the tensile test with an electrical current of 0.3 A. (a) IR image and (b) the temperature distribution evolution with strain.
As for the crack generation mechanism, we have reported this in our previous study where the electromechanical characteristics of the identical material systems (i.e., Ag/PET and Ag/PI) were studied under no electrical current. According to the results, the tractions exerted by the Ag film on the interface arise in the vicinity of imperfections (e.g., any defects or discontinuities) and this causes local delamination of the Ag film from the substrate. Once delaminated, the Ag film becomes locally freestanding, allowing strain localization (i.e., local necking) and eventually leading to crack formation. The two processes, delamination and strain localization, promote each other and coevolve, ultimately leading to Ag film failure. It is noted that as described earlier, the electrical current does not have a meaningful effect at the early stage of tensile deformation, indicating that the crack generation mechanism, suggested for no electrical current condition, is still valid under an electrical current condition; the effect of electrical current becomes significant after the Ag film is damaged. 

Based on these findings, we can speculate on the mechanism responsible for the current-induced deterioration of electromechanical performance in the Ag film/PET substrate system. At low strains, before any notable localized damage (localized deformations, cracks, etc.) develops in the sample, the deformation and hence the Joule heating occurs uniformly over the sample. With increasing strain, however, the deformation or crack formation is local to certain regions of the sample. This gives rise to a local increase of electrical resistance, thereby inducing Joule heating that locally raises the temperature both in the Ag film and PET substrate. Considering the high melting temperature of the Ag film (961 °C), the temperature rise of <100 °C observed in the present study does not appear to have a significant effect on the ductility enhancement of the Ag film. It will, however, influence the deformation behavior of the PET substrate because of its low glass transition temperature ($T_g = \sim 75$ °C). PET is therefore capable of facilitating deformation in the temperature range of the present study. The temperature dependence of the PET deformation behavior, obtained by conducting tensile tests inside a thermal chamber, is presented in Figure S5. The stress required to accommodate the strain was reduced by ~17% at 50 °C and ~25% at 70 °C in the strain range of 3–7%, indicating that PET deformed readily with increasing temperature. In the Ag film/PET substrate systems used here, the PET substrate was at least ~500 times thicker than the Ag film (470 times thicker in 400 nm Ag/188 μm PET and 1,880 times thicker in 100 nm Ag/188 μm PET). Therefore, the overall deformation behavior of each system was dominated by that of the PET substrate itself. The local rise of temperature will facilitate the deformation of the PET substrate at this location, inducing a localized deformation (i.e., necking instability), and this will accelerate cracking in the Ag film, consequently leading to a rapid loss of
electrical conductivity with strain. A schematic illustrating the fracture mechanism of the Ag film/PET substrate during tensile tests performed with or without an electrical current is presented in Fig. 5.

**Strategy to improve the current-induced deterioration of electromechanical performance.** It is known that a metal film on a polymer substrate can sustain a much larger deformation compared to its freestanding counterpart because of the suppression of the necking instability. The superior deformation capability of the polymer substrate suppresses the occurrence of localized deformation in the metal film, retarding the onset of necking, when the adhesion between the metal film and polymer substrate is strong enough to prevent delamination. However, our results showed that such a beneficial effect is impaired under an electrical current due to the local Joule heating in the polymer substrate, which accelerates its necking instability and eventually leads to the reduction of stretchability. Considering the operational conditions of flexible electronic devices and the demand for increasing current densities together with a much higher stretchability in future devices, this current-induced deterioration of electromechanical performance is a critical issue that should be resolved for their commercialization.

According to our results, as the local heating and weakening of the polymer substrate is a primary cause of the current-induced deterioration of electromechanical performance, we can expect to improve the electromechanical performance of the polymer-supported metal film system under an electrical current by enhancing the deformation resistance of the polymer substrate at elevated temperatures. Two methods can be followed for achieving this purpose. One is to increase the thickness of the polymer substrate and the other is to utilize a polymer substrate with a high glass transition temperature. To demonstrate these methods, we varied the substrate thickness and material: 400 nm Ag/38 μm PET, 400 nm Ag/25 μm PI, and 400 nm Ag/125 μm PI samples were additionally fabricated and tested. Figure 6 compares the test results. Regarding substrate thickness, as shown in Fig. 6a, the electromechanical behavior of the 400 nm Ag/PET or 400 nm Ag/PI samples was almost identical irrespective of the substrate thickness when the electrical current was not applied. With the application of an electrical current, though, the deterioration of electromechanical performance (i.e., the increase of electrical resistance) with strain was accelerated for both the thin PET and PI substrates. The onset strain at which the curve of \( I = 0.3 \) A started to deviate from that of \( I = 0 \) A was \( \varepsilon = 2.9\% \) for 400 nm Ag/38 μm PET, \( \varepsilon = 6.5\% \) for 400 nm Ag/188 μm PET, \( \varepsilon = 8.0\% \) for 400 nm Ag/25 μm PI, and \( \varepsilon = 9.5\% \) for 400 nm Ag/125 μm PI. The stretchability (or stretchability loss) data, shown in Fig. 6b, also confirmed that the current-induced deterioration became more severe with a thinner substrate; the thick PET or PI substrates always possessed a higher stretchability (or a lower stretchability loss) than the thin versions in the electrical power range of \( P/I_e = 25 \) A² mm⁻¹ (\( J = 2.5 \times 10^3 \) A cm⁻²) to \( P/I_e = 625 \) A² mm⁻¹ (\( J = 1.25 \times 10^4 \) A cm⁻²). The underlying mechanism of the improved current-induced deterioration with increasing substrate thickness can be understood by considering that the higher heat capacity of the thick substrate mitigates the temperature rise of sample during the tensile test. This leads to a high deformation resistance, which consequently suppresses the necking instability in the thick substrate. It is noted that the effect of substrate thickness on the current-induced deterioration was noticeable with the Ag film/PET substrate samples, while it was minor in the Ag film/PI substrate samples. As for the effect of substrate material, the Ag films on the PI substrates with a high \( T_g \) of ~400 °C showed excellent electromechanical characteristics under electrical currents, compared to the Ag films on the PET substrates with a low \( T_g \) of ~70 °C (Fig. 6). Note that there was no marked difference in the electromechanical characteristics between the Ag films on PET and PI substrates without the application of an electrical current: the stretchability was ~25% for both 400 nm Ag/38 μm PET and 400 nm Ag/188 μm PET, 26% for 400 nm Ag/25 μm PI, and 27% for 400 nm Ag/125 μm PI. As explained earlier, the onset strain for the 400 nm Ag/25 μm PI system at which the \( I = 0.3 \) A and \( I = 0 \) A curves started to deviate was 8.0%, even higher than that of 400 nm Ag/188 μm PET (6.5%). The curves of 400 nm Ag/25 μm PI and 400 nm Ag/125 μm PI with \( I = 0.3 \) A did not significantly deviate from the curves of \( I = 0 \) A (Fig. 6a). Such an
Conclusions

The electromechanical characteristics of polymer-supported metal films for flexible electronics were investigated under their operational conditions, i.e., a combination of mechanical deformation and electrical currents, by conducting electromechanical tensile tests inside a SEM or under an IR camera, while applying a constant electrical current. Our results showed that electrical current alone did not cause any noticeable loss of electrical conductivity with increasing strain. To lessen such a current-induced deterioration of electromechanical performance became more severe with increasing electrical current as well as increasing strain.

The in situ measurements of the surface morphology and temperature distribution over the samples during testing unveiled an underlying mechanism of the current-induced deterioration of electromechanical performance. The damage-induced local increase in the electrical resistance of the metal film, caused by localized deformations, cracks, etc., locally raised the temperature of the sample via Joule heating. This weakened the deformation resistance of the polymer substrate, accelerating the necking instability, and consequently leading to a rapid loss of electrical conductivity with increasing strain. To lessen such a current-induced deterioration of electromechanical performance in the polymer-supported metal films, we demonstrated the feasibility of two methods to enhance the deformation resistance of the polymer substrate at elevated temperatures: increasing the thickness of polymer substrate and utilizing a polymer substrate with a high glass transition temperature. Our results further suggest that the current-induced deterioration of electromechanical performance would be a critical issue in the development of polymer-supported metal films for flexible electronics.
methods previously developed to improve the stretchability of various polymer-supported conductive materials for flexible electronics (e.g., creating serpentine, non-planar buckling or metal mesh structures, using nanowire networks, etc.) because they are mostly based on increasing the heterogeneity in the conductive material. The methods developed in the present study can be an effective means to resolve this issue.

Methods
Preparation of Ag film/PET substrate and Ag film/PI substrate samples. The polymer substrates used were PET films with thicknesses of 38 and 188 μm (Toray XG532, Toray Industries, Japan) and PI films with thicknesses of 25 and 125 μm (Dupont Kapton HN, DuPont, USA). The PET films were primer-coated on one side to improve their adhesion to the Ag film, while a primer coating was not needed on the PI films because of their good adhesion. These PET and PI substrates were patterned into 28 × 1 mm rectangles using a cutting plotter (Graphtec FC8000, Graphtec America, USA). Ag films with different thicknesses were deposited on the patterned PET and PI substrates using an electron-beam evaporator (KVET-C500200, Korea Vacuum Tech., South Korea) at a rate of 0.15 nm s⁻¹. Ag films with thicknesses of 100, 200, and 400 nm Ag were deposited on 188-μm-thick PET substrates, and 400-nm-thick Ag films were deposited on the 38-μm-thick PET and 25- and 125-μm-thick PI substrates. The electrical resistance (R) in Figs 1 and 6a), measured over a length of 14 mm and a width of 1 mm (i.e., the gauge section of the test sample), was 1.02 ± 0.13 Ω for the 100 nm Ag film, 2.56 ± 0.16 Ω for the 200 nm Ag film, and 6.10 ± 0.25 Ω for the 400 nm Ag film.

Electromechanical tensile tests inside a SEM or under an IR camera. The gauge length of the above rectangular (28 × 1 mm) test samples excluding the two gripping parts was 14 mm. The electromechanical tensile tests were conducted using a commercial micro-tensile tester (Microtest, Deben UK Ltd., UK) in combination with simultaneous electrical resistance measurements. The strain developed in the sample was calculated by measuring the grip-to-grip distance whilst the strain rate was fixed at 6 × 10⁻⁴ s⁻¹. A DC power supply (Keithley 2000, Keithley Instruments, USA) was used to apply a constant electrical current to the sample in the range of 0–0.5 A. To measure the variation of electrical resistance in the sample during the test, a voltage signal was collected with a digital multimeter (Keithley 2400 Multimeter, Keithley Instruments, USA) and was divided by the applied electrical current; two electrodes were located at the gripping parts, giving the 14 mm gauge length.

To observe the evolution of the surface morphology (i.e., damage) of the sample during the electromechanical tensile test, the test was conducted inside a SEM (Quanta FEG 650, FEI, USA). The instrument was operated at an accelerating voltage of 10 kV, at room temperature under high vacuum conditions of <5 × 10⁻⁵ mbar; the micro-tensile tester was installed in the sample holder of the SEM. In addition, the evolution of the temperature distribution over the sample surface during the test was measured using an IR camera (FLIR X6500sc, FLIR Systems, Inc., USA). The surface of each sample was coated with a black dye to enhance the temperature measurements and a 5 × magnification lens was used for obtaining the IR images. The temperature calibration of the IR camera system was performed by measuring the temperature of the sample with a K-type thermocouple and data acquisition unit (Keysight 34970 A, Keysight Technologies, USA) attached to the bottom of the polymer substrate; as the electrical current flowed in the Ag film during the test, it was impossible to attach the thermocouple directly to the Ag film. According to finite element analysis, the temperature difference between the top (Ag film) and bottom (polymer substrate) surfaces was negligible because of the thinness of the polymer substrate; for the 400 nm Ag/188 μm PET sample, the difference was less than 1 °C at a Ag film temperature of ~95 °C (refer to Figure S6). The schematic of the electromechanical tensile testing system operating inside the microscope or under the IR camera is presented in Figure S7.

Data Availability. All data generated or analyzed during this study are included in this published article (and its Supplementary Information files).

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**Author Contributions**
S.G.H. and S.B.L. suggested the research idea and designed the experiments. The experiments were performed by D.W.J., J.W.L., and A.K. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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