BEHAVIOR OF ELECTRICAL STRESSED FLEXIBLE RESISTIVE LAYER

PART I: CARBON FILLED POLYMER

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There is a necessity to include sensors (resistors) in the design of organic electronic devices in order to extend the range of possible applications. It is essential to identify potential resistive materials, the processes and methods to structure them and to analyze their resistive properties on flexible substrates. Two materials widely used in organic electronics are carbon filled polymer (CFP) and the intrinsically conductive polymer (ICP) poly (3, 4-ethylendioxythiophene) doped with polystyrene sulfonate acid (PEDOT:PSS). In this paper we focus on the DC and pulsed stress behavior of these conductive polymers on flexible substrates and the resulting changes of their resistive properties. The paper is presented in two parts. Part one deals with carbon filled polymers (CFPs) and part two analyzes the behavior of intrinsically conductive polymers (ICPs).

Keywords: Thick Film Flexible Resistors, Carbon filled polymer, PEDOT:PSS, DC- and Pulsed Stress

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1. Introduction

There is a necessity to include sensors, based on resistive effects, in the design of organic electronic devices in order to extend the range of possible applications, mentioned in the last iNEMI roadmap. It is essential to identify potential organic resistive materials, the processes and methods to structure them and to characterize their resistive properties on flexible substrates including their reliability.

One of the most important issues is their stability under different electrical and non electrical influences such as mechanical, thermal stress together with humidity. Highly isolating flexible substrates make the devices even more susceptible to transient electrical stress events, like Electro Static Discharge (ESD).

Passive components are becoming susceptible to ESD events as their geometries are reducing. Integrating and embedding makes them even more vulnerable. Thick film is one main technology in producing resistive passive components on flexible substrates used in polytronics.

Two materials widely used in organic electronics are carbon filled polymer (CFP) and the intrinsically conductive polymer (ICP) poly (3, 4- ethylendioxythiophene) doped with polystyrene sulfonate acid (PEDOT:PSS).

Dispersing electrically conductive fillers into an isolating plastic matrix above a threshold concentration (percolation threshold) will result after curing in a conducting layer [1], [2], [3] and [4]. Carbon black (CB) is a widely used conductive filler which also improves dimensional and ultraviolet light stability, covering a wide square resistance range from ohms to Megohms.

The scientific importance, the properties and the growing applications of PEDOT:PSS, a polythiophene derivative, are presented in [5], [6] and [7]. This material has unique electrical (high conductivity) [5], [6], optical (high visible-light transmission)[6], [8], thermal [6], [8], and mechanical (high stability) [10], [11] properties. Due to this fact there is a wide application range for this polymer, like antistatic coatings, transparent conductor, conducting layer in capacitors, conducting layers for printed wiring boards, hole injection layer in organic light emitting diodes (OLED), conductor and semiconductor layer in organic thin film transistors (OFET), thermal and mechanical sensors.

The ESD behavior of thick film resistors has been reported in several studies [12]-[15], using RC- pulses, which lead to HBM-like discharges. They all report a resistance change after pulse application. A thermal failure model for thick film printed resistors is developed by D.C. Wunsch [15], similar to that for semiconductor devices. Overheating is considered to be the principle failure mechanism.
An efficient technique for electrostatic discharge (ESD) measurements is to use square pulses. Rectangular pulses from Transmission Line Pulser (TLP) are of particular interest for ESD measurements. They have been well established for the analysis of ESD-protection structures [16], [17] and [18]. This analytical test technique on the basis of square pulses allows the in-situ monitoring of the voltages and currents at the DUT, during pulsing and helps to gain fundamental insights into the electrical behavior.

In [19] and [20] the authors made an introduction to the TLP-measurement technique on thick film resistors. In [21] the TLP characterization of off-chip polymer ESD protection devices is presented. In this work the analysis is made on the flexible carbon based polymer thick film resistors and on intrinsically conductive polymer PEDOT:PSS layer, during DC-stress, single and multiple pulse stress, showing the behavior before, during and after the stress.

The mentioned conductive polymer layers are realized by screen printing in a roll-to-roll process on foil.

2. Test setup

The test setup for pulsing the polymer resistive layers is shown in Fig. 2.1. It consists of the pulse generator, the DC measurement and the test fixture with the DUT.

For the characterization of flexible thin film polymer layer the Time Domain Reflection (TDR) configuration of the TLP was used, with 100 ns wide pulses. Fig. 2.2 presents a simplified schematic of the TLP system for this configuration.

![Fig. 2.1. Principle of flexible resistive layer stressing and measuring](image)
An incident voltage pulse, defined by the length of charging transmission line (TL1), travels from the pulse generator to the device under test (DUT) and may be reflected at the DUT. The voltage of the incident and reflected pulse is measured with a passive voltage probe close to the DUT, so that both pulses can be recorded. The current of the incident and reflected pulse is measured by a current probe.

For 100 ns and wider pulses the measured incident and reflected voltages and currents overlap, if the measurement points are close enough to the DUT (TDR-O configuration if TL3 is of short length). Voltages and currents of the DUT can be directly measured. An example of the pulsed voltage across and the pulsed current into the DUT for a TLP of 100ns pulse width, in the TDR-O configuration, is presented in Fig. 2.3.

Fig. 2.3. Pulsed voltage across and pulsed current into the DUT for 100 ns TLP (TDR-O).
The used Transmission Line Pulser (TLP), presented in Fig. 2.4, is a Celestron-I type from Thermo Fisher Scientific Inc. It is a flexible, two terminal bench top system for fast, accurate and reliable characterization of advanced semiconductor structures [17]. The system is capable of delivering up to 10 A into a 50 Ω load, representing a dissipated power per pulse of 5 kW.

Prior to pulsing, the initial DC value of the resistor was recorded. During this measurement heating of the resistor was avoided, measuring well below its maximum DC power ratings.

During the TLP measurement the pulse amplitude was stepwise increased, as shown in Fig. 2.5. The resistors voltage and current transients were recorded for each pulse, leading to the pulsed current-voltage (I-V) characteristic of the DUT.

Additionally, after each pulse, a DC-spot measurement was performed in order to detect any resistance change. Fig. 2.6 shows the choosing of the DC spot measurement point to avoid self-heating of the resistive layer between pulses. The DC-spot measurement leads to the DC resistance vs. voltage (DC R-V) characteristic, as well as to the relative resistance change compared to the starting value (DC $\Delta R/R_0$).

From the pulsed I-V characteristic the R-V behavior during the pulses is calculated (pulsed R) as well as the pulsed relative resistance change with respect to the starting value (pulsed $\Delta R/R_0$).

To see whether the shift of the resistance after the first pulsed measurement remains constant, successive pulsed measurements were performed, unless the devices failed after the first measurement.
Applying high voltage pulses with constant amplitude gives information about the robustness of the DUT to TLP stress.

![Fig. 2.5. Principle of stressing and measurement technique with successive pulsing.](image)

In order to connect the flexible thin film devices to the TLP, a special DUT test fixture has been developed, using a 50 Ω environment.

### 3. Carbon based polymer resistive layer

The resistive test structures were realized in thick film technology in a screen printing roll-to-roll process. The printed polymer resistor paste (ESL RS-12112) is designed for the use on polyimide foils and ceramic substrates. Fig. 3.1 gives an optical view and a 3D scan of the tested flexible thick film carbon-based polymer resistor on polyimide (PI) foil, (Upilex S). The nominal sheet resistance...
The processed devices had an electrical stress resistance of $R_S = 100 \, \Omega/\square$, the resistor had a width $W$ of 1.4 mm, and a length $L$ of 3.55 mm, leading to an aspect ratio of $L/W = 2.5$. The resistor thickness was approx. 15 $\mu$m.

For contacting the resistive layer, a silver-based polymer paste (ESL 1109-S) was used.

Drying the deposited film was done at 125 °C for 15 minutes, followed by the curing process at 150°C for 2 hours.

Fig. 3.2 shows the cured carbon-based polymer resistive layer a) below the percolation threshold, where no conducting chains were formed and b) upon percolation threshold with CB grains embedded in the polymer matrix. As can be seen, to lower the resistivity, silver particles were added to the polymer paste.

In order to investigate the behavior at high current densities, a DC-four point measurement technique was applied to the resistive layer on flexible substrates using a professional semiconductor parameter analyzer, Agilent 4156C. During the DC-stress a thermal image of the layer was taken in order to characterize the thermal behavior. For DC characterization two kinds of measurements were performed, current sweep and current sampling.
3.1. DC electrical stress

To determine the DC behavior of the resistive layer, a double DC sweep was performed. Fig. 3.3 shows the area where the resistance changes due to excessive current rating. Due to Joule-heating the resistor is heated up and the resistance increases because of the positive TCR of the thick film polymer resistor. The double sweep of the measurement shows finally the permanent change of the initial value. Because of the excessive heating the resistivity of the layer was permanently reduced. Structural changes in the polymer matrix occurred.

Fig. 3.3. DC resistance during current sweep, flexible thick film polymer resistor on foil, $R_0 \, \Omega/□$ (permanent changes).

For reference measurements, test structures were made on a rigid substrate of Al$_2$O$_3$. Fig. 3.4 shows that the resistive layer has a similar behavior as on the flexible substrate, but the resistor on the rigid inorganic substrate can dissipate
much more power. The changes that occur during the double sweep are similar. The backward sweeping reveals the permanent changes that occur in the resistive layer, underlined by additional low current measurements.

Sampling measurements with currents covering the same current range as for sweeping measurements will underline the findings of the sweeping measurement. Fig. 3.5 shows the resistance behavior during successive current sampling measurements for the carbon based polymer resistive layers. Due to self heating, the steady state is reached after approx. 4 - 6 sec, representing the thermal time constant of the resistor. The permanent changes after this DC stress can also observed from this Fig.. After the last sampling with 20 mA and after resistor down- cooling, the permanent resistance decrease that occurred is in the range of approx. 2%.

Sampling measurements on the reference structure on Al₂O₃ substrate, represented in Fig. 3.6 result in even higher permanent changes in the resistive layer (-25%), because of higher currents used, which generate more heating of the resistive layer. But the behavior is similar to that on resistors on polyimide foils.
Infrared (IR) measurements during current sampling give the temperature distribution during self-heating of the polymer resistor, as represented in Fig. 3.7. It also shows the temperature profile across the length and width of the resistive layer. Because of the low thermal conductivity of the PI foil (0.29 W/mK) in the center of the resistor occurs a hot-spot. For a sampling current of 12 mA the hot-spot reaches a temperature of 91°C.

To prove this non-uniform surface temperature distribution an electro-thermal direct coupled simulation of the studied resistive structure was performed in ANSYS. Fig. 3.8 represents the simulation results, which are in good agreement with the measurements indicating the hot-spot in the center of the resistive layer.
Fig. 3.8. Simulation of temperature distribution during self-heating of flexible thick film polymer resistor on PI foil, showing hot spot in the center of the resistive layer.

Fig. 3.9. a) DC resistance during current sweep, flexible thick film polymer resistor on PI foil, b) power dissipation in resistive layer, $R_{\text{S}}$ 100 $\Omega/\square$ (resistor damage).

Fig. 3.10. a) Resistor burnout during DC stressing, caused by hot spot in the center of the flexible thick film polymer resistor on PI foil, b) zoomed damaged area $R_{\text{S}}$ 100 $\Omega/\square$. 
Applying multiple current sweeps up to 20 mA, as represented in Fig. 3.9 a), will finally lead to the burn-out of the resistive layer in the area of the hot spot. The dissipated power during burn-out, shown in Fig. 3.9 b) reaches 1.6 W, corresponding to a power density of 32 W/cm² of resistive area, which is roughly four times the rated value. Fig. 3.10 shows the burn-out region in the center of the resistive layer. Although the resistive layer is damaged in the center, there is still a current flow. This is illustrated in Fig. 3.11 by the temperature distribution of the damaged resistor, which now has two hot spots.

Fig. 3.11. Temperature distribution during self-heating of partially damaged flexible thick film polymer resistor on PI foil, two hot spots in the center of the resistive layer.

For thermal measurements the current through the resistive layer on PI foil was increased from 2 mA up to 20 mA, in 2 mA steps. For each current value a thermal images with an IR-camera was made and analyzed.

![Fig. 3.12. Resistance versus temperature during self-heating of flexible thick film polymer resistor on PI foil.](image)

Fig. 3.12. shows the resistance change of the carbon based polymer layer versus the corresponding measured layer temperature.
3.2 Pulsed electrical stress

The behavior of the carbon based resistive layer during pulsed stress was studied applying 100 ns wide pulses from transmission line pulser with increasing and constant amplitudes.

3.2.1 Influence of pulse amplitude on resistance

To determine the influence of pulse amplitude on the resistance, TLP measurements were performed with stepwise increasing the open source TLP pulse amplitudes. Recording the measured voltage and current transients at the resistor during each voltage step gives a 3-D representation of the measurement, as shown in Fig. 3.13 a) and, respectively b). During the 100 ns pulses there is little change in voltage and current during transients, while sweeping the pulse amplitude.

![Fig. 3.13. 3D representation of a) pulsed voltage at and b) pulsed current through the flexible thick film polymer resistor on PI-foil for TLP pulse width of 100 ns, TLP-voltage 0-1000 V, 2 V step.](image)

During the pulses, the pulsed resistance is calculated from each corresponding point of the current- and voltage- transients. Fig. 3.14 shows this behavior for two TLP voltages, 100 V and 900 V, indicating a resistance decrease with higher applied TLP voltages.
Fig. 3.14. Pulsed resistance of a flexible thick film polymer resistor on PI-foil ($R_s$ 200 $\Omega$/□) for TLP pulse width of 100 ns, TLP-voltage 100 V and 900 V.

Fig. 3.15 depicts the pulsed resistance behavior of three successive pulses of TLP voltages of 900 V, 902 V and 904 V. The same slope of the pulses indicates resistors down-cooling between pulses, suggesting the adiabatic behavior during this pulsed stress.

### 3.2.2 Influence of successive pulsing on resistance

Applying successive pulses with stepwise increasing voltages, up to the same final, non destructive voltage, gives information about the irreversible changes that take place in the resistor. Fig. 3.16 depicts the pulsed current-voltage characteristic for six successive measurements, M1 – M6. During each measurement the amplitude of the 100 ns wide pulses was stepwise increased, in
2 V steps, up to 1000 V of the TLP charging voltage. The different shapes of these characteristics indicate permanent and reversible changes that take place in the resistor during pulsing. This is confirmed by the absolute DC- resistance changes shown in Fig. 3.17. The final value of a measurement is the starting value for the following one, indicating irreversible changes in the resistor material. The decrease behaves like a reverse exponential decay, shown by the fit in Fig. 3.17.

Fig. 3.16. Pulsed current-voltage characteristic of flexible thick film polymer resistor on PI foil for successive measurements (pulse width 100 ns, Rs 200 Ω/□).

Fig. 3.17. DC-resistance change versus applied pulse voltage for successive measurements of flexible thick film polymer resistor on PI foil (pulse width 100 ns, Rs 200 Ω/□).
Fig. 3.18. DC-resistance change versus applied pulse voltage for successive measurements of thick film polymer resistor on Al₂O₃ substrate (pulse width 100 ns, Rₛ 100 Ω/□).

Reference measurements of test structures on a rigid substrate of Al₂O₃. Fig. 3.18 shows that the resistive layer has a similar behavior as on the flexible substrate, the inorganic substrate does not influence the pulsed behavior because of the adiabatic regime of the stressing.

\[ R_{\text{pulsed}}, \text{TLP 100 ns, 2-1000V, step 2V, Polymer Resistor on PI Foil 12, ESL 12112, R78, 200}\Omega/\square, \text{temp. M1-M6} \]

\[ y = -1.2019 \times 10^{-9}x^4 + 1.9426 \times 10^{-6}x^3 - 1.1533 \times 10^{-3}x^2 + 1.4476 \times 10^{-1}x + 9.6905 \times 10^2 \]

\[ R^2 = 0.9919 \]

Fig. 3.19. Pulsed resistance change versus applied pulse voltage for successive measurements of flexible thick film polymer resistor on PI foil (pulse width 100 ns, Rₛ 200 Ω/□).
Fig. 3.20. Pulsed resistance change versus applied pulse voltage for successive measurements of thick film polymer resistor on Al$_2$O$_3$ (pulse width 100 ns, $R_S$ 100 $\Omega$/$\square$).

From the corresponding pulsed I-V characteristic the R-V data during the pulses were calculated and represented in Fig. 3.19., for polyimide substrate, and in Fig. 3.20 for the reference on Al$_2$O$_3$ substrate. The resistance decrease during pulsing indicates, besides the irreversible resistance change, dynamic effects, which are reversible after each measurement. The starting points for the successive measurements have the same variation as for the DC- spot measurements and confirm the irreversible changes in the resistive layer.

### 3.2.3 Influence of pulse number on resistance

Applying multiple high voltage pulses, below the breakdown value, and with constant amplitude gives information about the robustness of the resistor to the TLP stress. Up to 4000 pulses of constant amplitude were applied with 100 ns wide pulses. For a TLP- voltage of 850 V the DC- and pulsed resistance decreases and stabilizes after 2500 applied pulses, withstanding the stress of further 1500 pulses, as depicted in Fig. 3.21, and Fig. 3.22 respectively. The resistance behavior is a double exponential decay, shown by the fit in the diagram.
Fig. 3.21. DC-resistance change versus number of applied pulses with constant voltage (TLP-voltage 850V, pulse width 100 ns, R_s 200 Ω/□).

\[
R(n) = R_0 + R_1 \exp(-n/n_1) + R_2 \exp(-n/n_2)
\]

- \( R_0 = 908.0 \) Ω
- \( R_1 = 8.0 \) Ω
- \( n_1 = 55 \)
- \( R_2 = 8.0 \) Ω
- \( n_2 = 800 \)

Fig. 3.22. Pulsed resistance change versus number of applied pulses with constant voltage (TLP-voltage 850V, pulse width 100 ns, R_s 200 Ω/□).

\[
R_{pulsed}(n) = R_0 + R_1 \exp(-n/n_1) + R_2 \exp(-n/n_2)
\]

- \( R_0 = 820.0 \) Ω
- \( R_1 = 8.0 \) Ω
- \( n_1 = 55 \)
- \( R_2 = 8.0 \) Ω
- \( n_2 = 800 \)

NOTE
The second part of the paper analyzes the behavior of electrical stressed intrinsically conductive polymers (ICPs): - Part II: Intrinsically Conductive Polymer. This second part is going to be published in the next issue of the Scientific Bulletin of UPB.

REFERENCES


