Electromechanical study of polyurethane films with carbon black nanoparticles for MEMS actuators

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Abstract
Pure polyurethane and nanocomposite carbon black (CB) polyurethane solutions were deposited by spin-coating on a silicon substrate using gold as the adhesion layer and electrode. Different test structures were achieved for electrical and mechanical characterizations. The incorporation of CB nanoparticles in the polyurethane matrix has a significant influence on the dielectric permittivity of the material with an increase of about one third of its value. The Young’s modulus of PU and nanocomposite PU films was determined by different characterization methods. Nanoindentation experiments have pointed out a Young’s modulus gradient through the film thickness. By performing mechanical tests (tensile, bulge, point deflection) on freestanding films, an average Young’s modulus value of about 30 MPa was found as well as a residual stress value of about 0.4 MPa. However, no influence of the presence of the nanoparticles was found. Finally, several MEMS actuators were realized and characterized. At their fundamental resonance frequency, the actuation of the nanocomposite membranes is more efficient than that of pure polyurethane. However, the time constant of the material seems to provide a major barrier for the development of high-frequency PU-based micro-actuators.

Keywords: MEMS, actuator, polyurethane, carbon black, electrostriction

1. Introduction
Electroactive polymer (EAP) transducers are an emerging technology that offers great opportunities for the development of actuators [1–3]. Among them, dielectric EAP for actuators (DEAs) have some advantages over some traditional electroactive materials such as electro-ceramics or magnetostRICTive materials due to their high strain, flexibility, low energy consumption and low cost [4]. DEA actuation is driven by electric field or Coulomb force. They can be used to develop thin film-based actuators that consist of a polymer film sandwiched between two compliant electrodes. Two mechanisms contribute to the actuation phenomenon: Maxwell’s effect and the true electrostrictive effect [3, 5]. Maxwell stress is a result of the electrostatic interaction between oppositely charged electrodes. The
where $Q$ is the real part of the dielectric constant and $\varepsilon_0$ the vacuum dielectric permittivity. True electrostriction is due to the reorientation and mutual attraction under an applied electric field of induced or permanent dipoles present in the dielectric. This microscopic movement induces macroscopic deformation in the polymer:

$$S_E = -Q\varepsilon_0^2 (\varepsilon_r - 1)^2 E^2,$$

where $Q$ (negative) represents the coefficient of true electrostriction. The total deformation of the electrostrictive polymer under an electric field is

$$S_{tot} = S_{Max} + S_E = ME^2,$$

where $M$ is the apparent electrostrictive coefficient in the direction of the electric field. All dielectric elastomers are subjected to both effects, however one mechanism may prevail over the other. In amorphous and low-polar elastomers, like polydimethylsiloxane (PDMS), Maxwell stress dominates electrostriction. However, polar elastomers, or semi-crystalline elastomers having numerous interfaces between nanodomains of different dielectric permittivity, true electrostriction can prevail [5]. Some types of polyurethane elastomers possess interesting electrostrictive properties, as they comprise soft segments [6], responsible for the elasticity of the polymer, and hard polar segments that orient themselves in the presence of an electric field [7]. Therefore, the electrostrictive effect is due to true electrostriction as well as the trapping of charges at the interfaces between soft and hard-segment domains. Several studies have compared the electro-mechanical properties of various types of polyurethane (PU) and demonstrated that true electrostriction is widely responsible for the electromechanical response of the material and the Maxwell stress is neglected [7, 8]. In addition, it was experimentally observed that $Q$ is inversely proportional to the product of the Young’s modulus and the dielectric permittivity [7]:

$$Q \propto \frac{1}{\varepsilon_0 \varepsilon_r Y}.$$  

The apparent electrostrictive coefficient is then proportional to dielectric permittivity and inversely proportional to the Young’s modulus (the assumption $\varepsilon_r \gg 1$ leads to simplification) [9]:

$$M \approx -Q\varepsilon_0^2 (\varepsilon_r - 1)^2 \propto -\frac{1}{\varepsilon_0 \varepsilon_r Y} \varepsilon_0^2 (\varepsilon_r - 1)^2 \rightarrow M \propto -\frac{\varepsilon_0 \varepsilon_r}{Y}.$$  

An approach to improve the electromechanical response of an electrostrictive elastomer is to disperse conducting or semi-conducting nano-objects in the polymer matrix [9]. The goal is to increase the dielectric permittivity and to create nanodomains. Wongtimnoi et al [10] have shown that incorporation of carbon black in a PU matrix at a concentration of 1.25 vol% (in the vicinity of the percolation threshold) yields a threefold increase in the absolute value of the apparent electrostriction coefficient $M$ at low frequency, compared to pure PU. Using nanoscale particles instead of larger ones, at low concentration, should not have a significant impact on the Young’s modulus of the polymer; one objective of this study was to check this assumption.

Works dealing with the integration of electrostrictive elastomers in MEMS are not numerous in literature. Most of them are based on PDMS [11]. Indeed, PDMS possesses interesting processing properties such as soft-lithography and O$_2$ plasma surface activation, for optical [12] and fluidic applications [13]. However, electrostriction is dominated in PDMS by Maxwell stress and important electric fields are necessary to achieve significant deformation. As explained previously, PU elastomers and related composites present interesting electrostrictive properties but to our knowledge, there is no published work on its integration in MEMS. This study focuses on the electrical and mechanical characterization of pure and carbon black composite polyurethane films. The realization and the characterization of a first MEMS demonstrator are shown.

2. Pure and composite solution preparation

The polymer is a polyether-based thermoplastic PU, ref. Estane 58888 NATO21 Lubrizol. Hard segments comprise the 4,4’-methylene bis(phenyl isocyanate) (MDI) and 1,4-butanediol (BDO) and soft segments are poly(tetramethylene oxide) (PTMO). The PU88 has a density of 1.13 g cm$^{-3}$, a hardness of 88 Shore A and the hard-segment content is about 46%. Carbon black (CB), Vulcan XC72R (Cabot Corp.), has a specific surface area of 254 m$^2$ g$^{-1}$. The primary CB nanoparticles have an average diameter of 30 nm and a specific gravity of 1.80. Pure and nanocomposite films were prepared from solutions. For pure solution, the granules were put in N,N-dimethylformamide (DMF, Sigma-Aldrich D158550, 99%); 20 wt% PU into DMF and mechanically stirred for 24 h to complete dissolution.

In the case of the nanocomposite solution, the PU granules were heated at 80°C for 3 h before use. They were then put in N,N-dimethylformamide (DMF, Sigma-Aldrich D158550, 99%); 40 wt% PU into DMF. The solution was maintained at 80°C for 2 h. At the same time, optimal CB volume fraction (1.25 vol% [10]) was dispersed in DMF using an ultrasonic processor (Hielsher UP400S, 400 W, 24 kHz, 7 mm diameter sonotrode) under the following experimental conditions: 20 min duration, 80% amplitude, 80% duty cycle. Subsequently, the two solutions were mixed together, heated and stirred until a homogeneous and viscous solution was obtained. This solution was degassed for 24 h at room temperature before use. Solutions were then spin coated on 4'' silicon wafer. The films were then placed on a hotplate at 115°C for 15 min to remove the solvent.

3. Electrical characterizations

Capacitance measurements were done on test samples with a HP® 4284A impedance meter coupled to a probe station.
PU solutions (20 wt%) were spin-coated (film thickness ranging from 19.2 to 25.1 μm) and baked (115 °C—15 min) on a 2" silicon wafer previously metalized with Cr (10 nm) and Au (100 nm) layers. Top gold electrodes were evaporated through a shadow mask, to avoid any PU film damage which can occur with a photoresist-based process. Square shaped capacitors were achieved with the following side dimensions: 500 μm, 1 mm, 2 mm and 3 mm (figure 1). For each sample, the dielectric constant was determined from the slope of the capacitance (electrode surface) curve. A 2 V amplitude sinusoidal test signal was applied with a frequency ranging from 100 Hz to 10 kHz. Four pure PU and four composite PU samples were analyzed. The error bars in figure 2 take into account both the scatter between the data and the uncertainties in the determination of the dielectric constant. At 100 Hz, the composite dielectric constant is 1.41 times that of the pure one, reaching 10.4. At 10 kHz, the multiplication factor becomes 1.36 and the dielectric constant reaches 8.9 for composite PU. Using a parallel RC model, electrical conductivity was also determined. Conductivity of the composite PU (1.25 vol%) is very close to that of pure PU, ranging from 4.5 × 10⁻⁹ S m⁻¹ at 100 Hz to 4 × 10⁻⁷ S m⁻¹ at 10 kHz. The dissipation factor was between 0.04 and 0.07. All these data are in very good agreement with those already published in [10] and they confirm that the composite PU at 1.25 vol% concentration is just below the percolation threshold.

4. Mechanical characterizations

Several methods were employed to evaluate the Young’s modulus and stress state of pure PU and nanocomposite PU films: nanoindentation measurements on films deposited on a thick substrate (surface mechanical characterization) and mechanical testing on free-standing polyurethane membranes (average in-plane elastic modulus and residual stress).

4.1. Nanoindentation measurements

Nanoindentation experiments were performed on an Agilent® XP nanoindenter (dynamic contact module head). A CSM (continuous stiffness measurement) device was used for a continuous evaluation of the reduced modulus between 150 and 450 nm of indentation depth, in order to avoid substrate influence on the measurements, by following the Oliver and Pharr model [14]. A Poisson’s ratio value of 0.49 was assumed. The characteristics of the four PU88 polyurethane coatings that were analyzed are summarized in table 1. For three samples, the short 115 °C—15 min annealing (after film deposition by spin-coating) was followed by a long annealing over several hours in order to simulate the thermal budget of the realization of a MEMS device (KOH etching of the silicon substrate to achieve membranes). For all samples, figure 3 shows a large variation of the Young’s modulus as a function of the indentation depth, as for a layered solid. This phenomenon, previously reported by Wongtimnoi [15], is related to the film thickness and depends on the annealing conditions after film deposition. The hard-segment domain size may change through the film thickness as a function of the solvent evaporation kinetic. At the film surface, where rapid solvent evaporation favors hard and soft segment mixing, the hard-segment domain size is smaller than at the film/substrate interface leading to a larger density of cross-linking nodes, then a higher material stiffness. When considering measurement errors, carbon black nanoparticles do not seem to have any...
significant influence on the modulus value. Presumably the concentration of CB nanoparticles on the first few hundred nanometers is not large enough to change the elasticity of the material.

4.2. Tensile tests

Micro-tensile experiments were performed on an experimental device already described elsewhere in [16, 17]. The main steps for the realization of polyurethane specimens are shown in figure 4 and pictures of a processed wafer and a finished specimen are shown in figure 5. As polyurethane film cannot be patterned, the beam geometry was defined by means of a dicing saw (Disco®). A thin film (about 50 nm) of Au/Cr was used as an adhesion layer before PU deposition (21 μm). This adhesion layer was patterned by lift-off in order to define anchor pads under the PU film. The pads were also used to guide the dicing blade through the transparent polymer film. Unfortunately, that was not possible for composite films that were non-transparent and black in color. That is the reason why tensile tests were only performed on pure polyurethane films. After substrate etching, the excess PU film was peeled away from the substrate. Finally, the fixing holes of the specimens were released by hand using a cutting tool. The overall dimension (length × width × thickness) of each specimen was 20 mm × 14 mm × 450 μm whereas the PU beam (gage) dimension was 3 mm × 1 mm × 21 μm.

The obtained stress–strain experimental curves are given in figure 6. The displacement speed was set to 1 μm s⁻¹ (first test) or 50 μm s⁻¹ (second and third tests), and the applied stress remains uniaxial, allowing direct evaluation of the mechanical properties. During each measurement, a loop was performed at half cycle in order to observe any damage phenomenon. The change in the mechanical properties resulting from the first extension is clearly visible and is called Mullins’ effect. From the beginning of the first curve, a Young’s modulus value of (28.0 ± 0.7) MPa was calculated.

4.3. Measurements on free-standing membranes

4.3.1. Fabrication process of polyurethane membranes.

Devices were achieved from thermally oxidized (1.5 μm) double-side polished (100) silicon wafers. On the back side, the oxide layer was used as masking layer for KOH etching of silicon. On the front side, SiO₂ played the role of an etching stop layer to protect PU or metallic layers. First, the back side SiO₂ was structured by photolithography and BOE etching in order to obtain square openings of 2 mm (figure 7(a)). On the front side, a Cr/Au (20 nm/40 nm) layer was deposited using an electron-gun evaporator and patterned by lift-off to achieve square openings aligned with those on the back side (figure 7(b)). The gold surface morphology was modified by aqua regia chemical etching (HCl (37%): HNO₃ (68%), 3:1, v:v) to improve PU adhesion. Indeed, we observed qualitatively a better adhesion of PU on Au with such a treatment than without any treatment. PU film was then...
4.3.2. Point membrane deflection. The principle consists of applying a concentrated transversal load at the center of a freestanding membrane and measuring the resulting deflection. The stress value and possibly the Young’s modulus of the material can be extracted from the load–deflection relationship. The Young’s modulus, Poisson ratio and residual stress of the material, respectively. $C_1$ and $C_2(\nu)$ are dimensionless coefficients that depend on the membrane shape, and $C_2(\nu)$ also depends on Poisson’s ratio. For a square membrane, $C_1 = 3.393$ [23]. A Poisson ratio of 0.5 was used, the corresponding $C_2(\nu)$ value was calculated as $C_2(\nu) = (0.8 + 0.062\nu)^{-3} = 1.74$ [24]. The displacement response was measured using a Wyko® NT1100 white-light interferometric microscope. The experimental setup was previously detailed in [25]:

$$P = C_1 \frac{\sigma_0}{a^2} h + C_2(\nu) \frac{f}{a^2 \frac{1}{1-\nu}} h^3$$  \hspace{1cm} (6)
Table 2. Bulge test results.

| Membrane Type | Type   | \(a\) (\(\mu\)m) | \(t\) (\(\mu\)m) | \(\sigma_0\) (MPa) | \(Y\) (MPa) |
|---------------|--------|-------------------|-----------------|----------------|-------------|
| PU1           | PU88   | 840 ± 50          | 47.4 ± 0.3      | 0.49 ± 0.01    | 28.1 ± 11.4 |
| PU2           | PU88   | 840 ± 50          | 47.7 ± 0.3      | 0.48 ± 0.01    | 20.1 ± 2.2  |
| PU3           | PU88   | 840 ± 50          | 46.8 ± 0.3      | 0.55 ± 0.01    | 22.4 ± 1.8  |
| CB-PU         | PU88 + CB | 1000 ± 50       | 61.5 ± 0.3      | 0.40 ± 0.00    | 17.7 ± 0.2  |

Table 3. MEMS characteristics.

| Device Type   | Type   | \(a\) (\(\mu\)m) | \(t\) (\(\mu\)m) | \(f_0\) (kHz) |
|---------------|--------|-----------------|-----------------|--------------|
| CB-PU88-MEMS 1 | PU88 + CB | 840 ± 50       | 48.2            | 7.4          |
| CB-PU88-MEMS 2 | PU88 + CB | 840 ± 50       | 48.2            | 7.6          |
| PU88-MEMS 3   | PU88   | 840 ± 50        | 55.0            | 11.2         |
| PU88-MEMS 4   | PU88   | 840 ± 50        | 55.0            | 8.2          |

Figure 9. Pressure–deflection curves for pure and composite (carbon black) PU88 membranes.

The pressure–deflection curves for three similar pure polyurethane membranes and one composite CB-polyurethane membrane are shown in figure 9. The composite membrane was larger and thicker than pure polyurethane ones. The membrane dimensions and the best-fit residual stress and Young’s modulus values are reported in table 2. The errors given for \(Y\) and \(\sigma_0\) are only related to the curve-fit algorithm. These results are very similar to those obtained with the other characterization method shown previously: the Young’s modulus values lie between about 10 and 30 MPa, whereas the residual stress value is about 0.5 MPa. As noted above, the determination of the Young’s modulus was imprecise except for the composite membrane due to its lower stiffness. Finally, the presence of carbon black nanoparticles seems to have no significant influence on the Young’s modulus value. But further experiments should be carried out in order to minimize measurement errors on \(Y\).

5. Microsystem fabrication and characterization

The fabrication process for microsystems starts as membrane fabrication for the bulge test and point deflection method, but electrodes must be integrated on both faces of the PU membrane. After lift-off of the Cr/Au layer, the bottom Au electrode (40 nm) was deposited. The PU film was spin-coated. The top electrode (side length about 1.0 mm) was deposited through a mask to avoid a patterning step. Finally, the Au/PU/Au membrane was obtained after standard silicon deep etching in an aqueous KOH solution (34%, 80 °C) (figure 10(a)). After dicing, each sample was mounted on a TO8 header and connections were made by wire-bonding. A top view of a pure PU microsystem is reported in figure 10(b).

The membrane was excited at its fundamental resonance frequency by applying a sinusoidal signal (electrostrictive mode) from a waveform generator (Agilent® 33220A connected to a Trek® 609 D-6 voltage amplifier) to the upper electrode, with the bottom electrode held at ground potential. Another driving mode, called the pseudo-piezoelectric mode, consisted of applying a dc bias in addition to the ac excitation in order to induce permanent polarization and pre-stress the material (the dc bias voltage was chosen equal to the ac voltage amplitude). Displacements were measured at the center of the upper electrode by means of a laser vibrometer (Polytec® OFV-505 with OFV-5000 controller). The main characteristics of the microsystems are reported in table 3, where \(a\), \(t\) and \(f_0\) are the membrane half side length, thickness and fundamental resonance frequency, respectively. Figures 11 and 12 show the measured center deflections for pure and composite PU membranes, normalized to their thickness, as a function of the squared applied electric field. Curves are quite linear; in a first approximation, as the membrane deflection is very small compared to its thickness, the membrane has a behavior similar to that of a free film and depends on the square of the electric field, as shown in equation (3). Deflections in the pseudo-piezoelectric mode are about four times as large as those in the electrostrictive mode. It can be explained by the fact that the electric field has a constant direction in pseudo-piezoelectric mode. The induced rotation of dipoles in the material remains limited and the actuation is more efficient. Finally, deflections of composite membranes are about twice as large as those for pure PU membranes. The presence of
CB nanoparticles has a positive influence on the microsystem actuation. However, deflections are only limited to a few hundred nanometers because of the relatively large thickness and stiffness of the membranes. In addition, the clamping effect of the stiff metal electrodes might be important, despite their low thickness (40 nm) [26]. However, the deflections measured in this study are about one order of magnitude larger than those obtained in equivalent microsystems integrating irradiated P(VDF-TrFE) film as the membrane [27]. Another barrier to the design of PU-based actuators is related to the response time of the polymer. Beside harmonic characterizations, we also measured the voltage step-response of the microsystems and a time constant of about 800 ms was found. A value of the same order of magnitude (700 ms) was previously measured by Diaconu et al [6]. At the fundamental resonance frequency of the membranes, it can be assumed that the true electrostrictive effect in the material is negligible and that the membrane actuation is only due to the electrostatic interaction between the two electrodes.

6. Conclusion

The integration of carbon black nanoparticles in the polyurethane matrix has a significant influence on the dielectric permittivity of the material with an increase of about one third of its value. The Young’s modulus of PU and nanocomposite PU films was determined by different characterization methods. Nanoindentation experiments have pointed out a Young’s modulus gradient through the film thickness that might be related to post-deposition annealing conditions (temperature and time) and inhomogeneous solvent evaporation. By analyzing the behavior of freestanding films, an average in-plane Young’s modulus value of about 30 MPa was found as well as a residual stress value of about 0.4 MPa. However, no influence of the presence of the nanoparticles was found. Finally, several MEMS actuators were realized and characterized. At their fundamental resonance frequency, the actuation of the nanocomposite membranes is more efficient than that of pure polyurethane. However, the time constant of the material seems to provide a major barrier for the development of high-frequency PU-based micro-actuators.

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