Spray-coated carbon nanotube carpets for creeping reduction of conducting polymer based artificial muscles

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Abstract
During cyclic actuation, conducting polymer based artificial muscles are often creeping from the initial movement range. One of the likely reasons of such behaviour is unbalanced charging during conducting polymer oxidation and reduction. To improve the actuation reversibility and subsequently the long time performance of ionic actuators, we suggest using spray-coated carbon nanotube (CNT) carpets on the surface of the conducting polymer electrodes. We show that carbon nanotubes facilitate a conducting polymer redox reaction and improve its reversibility. Consequently, in the long term, charge accumulation in the polymer film is avoided leading to a significantly improved lifetime performance during cycling actuation. To our knowledge, it is the first time a simple solution to an actuator creeping problem has been suggested.

Keywords: carbon nanotubes, spray coating, conducting polymers, soft actuators, creep

1. Introduction
Ionic conducting polymer based actuators (CPAs) are one of many alternatives to traditional actuators. Due to a low actuation voltage, lightness and compliance, conducting polymer based actuators are biocompatible and attractive for applications in biomedical engineering and biotechnology [1, 2]. Also, CPAs can induce a relatively large force, and only small currents are needed to hold constant strain at DC voltage [2, 3].

The use of conducting polymers as actuators is based on the linear dimensional changes during their electrochemical doping and dedoping process. There are several simultaneous and subsequent processes involved in changing the electrochemical state of the conjugated polymer, such as the charge and mass transport and the conformational changes of the polymer. These processes depend on the material properties, actuation and cycling conditions and affect the performance of the actuators regarding strain, speed, stress and efficiency in a complex and still not fully understood way [4]. For example, macroscopic charge transport of intrinsically conductive polymer films requires ‘hopping’ between polymer chains. Because of this, the composition and the morphology of the polymer film might have a significant influence on the charge transfer [5–9]. Furthermore, the volumetric expansion mechanism needs to take into account the doping of the conducting polymer, and the size and mobility of the anion and cation in the electrolyte and the solvent [10–12]. Finally, even though the flux of ions to and from the material is considered to be the primary mechanism
for producing deformation, conformational changes in the polymer could also play a role [5, 7, 13].

Despite unique advantages that CPAs offer, redox reactions and polymer swelling are the main drawbacks of the conducting polymer based devices [14]. In most cases, actuators fail gradually due to fatigue during the actuation or due to spontaneous self-degradation [15]. Gradual deterioration is observed and reported most often and could be caused by fatigue of the polymer, i.e. change in the mechanical properties, chemical degradation of the material, receding electrolyte or irreversible redox reactions and reduction in the charge transfer [4, 15–19]. The latter is also one of the reasons of creeping [16, 20].

Creeping in CPAs is observed as a continuous displacement of the movement range [21]. It originates from the incomplete recovery of the initial position on consecutive actuation cycles. There are two identified reasons causing such behaviour: (1) cycling induced solvent swelling suggested by Melling et al. [20] and (2) unbalanced charging characterized by Kaneto et al. [22–24]. In the first case, cycling opens the polymer matrix so that with each cycle ions diffuse faster and therefore deeper into the film. Consequently, if present, more solvent can irreversibly accommodate in between polymer chains [7, 20, 23, 25]. In the case of unbalanced charging, both a faster actuation rate during reduction [20] and oxidation [26] were previously observed. This is attributed to the higher electronic conductivity in the oxidised state and the higher ionic conductivity because of the more expanded, open and swollen polymer. In any case, unbalanced charges at the end of an actuation cycle induce accumulation of the ions in the films [23].

Carbon nanotubes (CNTs) are widely used for applications in conducting polymer based actuators, primarily to increase the conductivity, robustness and stiffness of the polymer matrix [27–29]. Besides, incorporation of carbon nanomaterials in actuators was shown to increase the Faradaic capacitance of the conducting polymer, the electrical conductivity and to consequently increase the actuators’ stroke and work per cycle [30–34]. Just as in batteries, incorporation of the CNTs also increases the redox reversibility and significantly reduces creeping [27, 32, 35]. Nevertheless, one of the biggest challenges for using these composites is the low solubility of carbon materials [36]. Only small amounts of the more conductive, non-oxidised CNTs or graphene can be homogeneously dispersed in the polymer solutions [37, 38]. Higher concentrations lead to the formation of agglomerates. When incorporated during the electropolymerization [33], the concentration of carbon nanotubes is difficult to control. Other fabrication methods, i.e. layer-by-layer deposition [34], drop casting [32], soaking [31], that would allow to increase the CNTs amount in the polymer actuators were suggested, but they also require soluble and easily processable CNTs.

In this article, we suggest using spray coated carbon nanotube carpets (CNTCs) to reduce creeping and improve the long-time performance of the actuators. We show that spray-coating allows deposition of large quantities of CNTs leading to well connected and conductive networks. Furthermore, CNTCs are deposited only on the surface and do not create a layered structure that is prone to delamination. Finally, we show that CNT carpets increase the Faradaic capacitance of conducting polymer electrodes, improve charge–discharge reversibility and subsequently lead to a significant decrease in creeping. The effects of CNTCs are particularly advantageous if actuators are needed for cyclic positioning.

2. Materials and methods

2.1. Materials

Hydrophobic polyvinylidene fluoride (PVDF) membranes with a mean pore diameter of 0.1 μm were bought from Merck Millipore Corp (Millipore VVHP04700 Durapore Membrane Filter, thickness - 120 μm and 47 mm in diameter). Poly(3,4-ethylenedioxythiophene) polystyrene sulfonate (PEDOT:PSS) 1.3 wt% conductivity grade dispersion in water (483095 Aldrich), with a PEDOT:PSS ratio of 1:2.5 was bought from Sigma-Aldrich. The ionic liquid emimTFSI was bought from Solvionics (Im0208a).

Double-walled CNTs were synthesized in Cirtimat-CNRS as reported in [39] and provided in water–sodium dodecyl sulfate (SDS) suspension. CNT samples contained 77% of double-walled CNTs, 18% of single-walled CNTs and 5% of triple-walled CNTs that have a length ranging between 1 and 10 μm and a diameter of 1.2 – 3.2 nm. They were shown to exhibit a metallic electrical behaviour [40], 0.1 wt% of CNTs and 1 wt% of SDS were dispersed in deionized water and separated using 150 W ultrasond (VibraCell 75042, Bio-Block Scientific) in an ice bath for 30 min. The suspension was then centrifuged (16000 rpm, 10 min) and the supernatant was collected. Before spray coating, all the suspensions were dispersed using pulsed ultrasound (5 sec on and 5 sec off) for 10 min at 150 W. For spraying SDS, 1 wt% solution was prepared in deionized water.

2.2. Fabrication of PEDOT:PSS/PVDF/PEDOT:PSS actuators

Figure 1 depicts a schematic illustration of materials, as well as the fabrication process of actuators used in this work. We were using PEDOT:PSS/PVDF/ionic liquid actuators made by solvent casting as previously reported and characterized by our group [41, 42]. Nevertheless, CNTs could equally enhance actuators produced in other ways or based on other conducting polymers.

Hybrid PVDF/PVDF-graft-PEGMA membranes were prepared to ensure excellent adhesion between the solvent cast conducting polymer film and the membrane [41]. Grafting of about 0.6 mg of PEGMA on each side of the PVDF membrane leads to 30 μm PVDF-graft-PEGMA layer (hydrophilic) on each surface and 60 μm of hydrophobic PVDF membrane in the center. For the sake of simplicity, in the article, PVDF/PVDF-graft-PEGMA membranes are referred as PVDF. The electrical conductivity of PEDOT:PSS was enhanced by secondary doping with polyethylene glycol (PEG) with average Mn = 400 (PEG400) [43]. 1 vol% of
PEG400 in PEDOT:PSS solution were mixed on a mechanical shaker for 3 hours at 3000 rpm and if needed kept at 4 °C before use. Herein, PEDOT:PSS secondary doped with PEG is called PEDOT:PSS.

For the fabrication of actuators, functionalized PVDF membranes were fixed in a ø45 mm aluminium clamp and 3 ml of PEDOT:PSS were solvent-cast. Membranes were dried in air in a UV-reduced environment for at least 20 hours. The other side of the membrane was processed in the same way. Membranes were thermally annealed in an oven for 1 hour at 70 °C. In order to create a trilayer, the procedure was repeated on the other side of the membrane. Dried actuators were composed of 15 μm thick PEDOT:PSS electrodes, 20 – 25 μm interfacial layer and 60 μm insulating PVDF membrane as shown in the scanning electron microscope (SEM) image on the top right of figure 1. SEM images were obtained using a Hitachi S – 4800 field emission scanning electron microscope (acceleration voltage 800 V, working distance of about 5 mm). Table 1 summarizes the geometry and other properties of actuators used in this work.

Table 1. Geometry and electrical properties of actuators.

<table>
<thead>
<tr>
<th>Without CNT</th>
<th>With CNT</th>
<th>With SDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>h_{dry} (\mu\text{m})</td>
<td>133</td>
<td>142</td>
</tr>
<tr>
<td>h_{el} (\mu\text{m})</td>
<td>15/13</td>
<td>20/10</td>
</tr>
<tr>
<td>w_{dry} (\text{mg})</td>
<td>3.6 ± 0.1</td>
<td>3.5 ± 0.2</td>
</tr>
<tr>
<td>w_{ILs} (\text{mg})</td>
<td>5.8 ± 0.2</td>
<td>6.2 ± 0.3</td>
</tr>
<tr>
<td>R_{el} (\text{k}\Omega)</td>
<td>5.1 ± 3.7</td>
<td>1.6 ± 0.5</td>
</tr>
</tbody>
</table>

\(a\) Thickness of dry trilayer.
\(b\) PEDOT:PSS electrode thickness.
\(c\) Weight of dry actuator.
\(d\) Weight of the actuator with ionic liquid.
\(e\) Resistance of the electrodes.
2.3. Spray coating of CNTs

For post-treatment of actuators with CNTs or SDS, the membrane cast with conducting polymer on both sides was divided into three parts. One part was not modified and was used as a reference. The second part was used for deposition of CNTs and the third part was spray coated with only surfactant (SDS) solution. Spray coating was done under a fume hood and in a glovebox (Captair Pyramid Glovebox). PEDOT:PSS/PVDF/PEDOT:PSS trilayers were stuck to the glass slide, next to two silver electrodes and a piece of paper used as a color indicator as shown in online supplementary figure S1. The following spraying parameters were used: nozzle size of 0.2 mm with associated needle, pressure 1.5 kgf m⁻², valve opening of 1.5 turns, distance of about 30 cm. To ensure homogeneous spraying, the spray was targeted by a laser pointer. CNTs were sprayed until the resistance between silver electrodes decreased to about 10 kΩ corresponding to approximately 6 ml of CNT suspension. A dense and well interconnected CNT carpet was formed as shown in the SEM picture of the surface of PEDOT:PSS in figure 1. After spraying on one side, the membrane was dried under the fume hood at room temperature for 1–2 hours to make sure the surface of the trilayer is dry. The other side of the trilayer was treated in the same way. For post-treatment with a surfactant, 6 ml of SDS solution in water were sprayed. After the spray of CNTs or SDS solutions, the trilayer was again dried in an oven for 1 hour at 70 °C.

2.4. Geometry of PEDOT:PSS/PVDF/PEDOT:PSS actuators

15 x 2 mm² rectangles were cut by a CO₂ laser (45 W power, 100 cm s⁻¹ speed, 1000 Hz, Trotec FineMarker Hybrid) and incubated in ionic liquid for 24 hours before use. Each actuator was then weighted and its thickness measured by digimatic indicator (Mitutoyo Absolute). They were kept in ionic liquid (emimTFSI) for 24 hours before characterisation and then in a closed tube if not used immediately. After soaking, their weight and thickness were measured again and are summarized in table 1. As reported previously by our group [41], incubation in ionic liquid slightly decreases the thickness of PEDOT:PSS to about 12 µm leading to an overall actuator thickness of 137 µm.

Actuators with CNTs might also influence the amount of ionic liquid adsorbed during the same time of incubation (2.7 mg and 2.2 mg with and without CNTs respectively). With ionic liquids, CNTs form gel-like composites (bucky gels) meaning that in addition to storing the ionic liquid in PVDF membrane, some of it could also be adsorbed by CNT carpet on the surface of PEDOT:PSS. If that is the case, it could have interesting implications on redox performance of PEDOT:PSS itself, as counter ions would not need to diffuse through the PEDOT:PSS film to balance the charge. Instead, ions adsorbed by CNTs could participate in a redox reaction. This possibility is not discussed further in the article and requires further investigation.

2.5. Electro-chemo-mechanical characterization of PEDOT:PSS/PVDF/PEDOT:PSS actuators

Resistance and conductivity of PEDOT:PSS were measured between two extremities of each electrode and between two electrodes, by applying 0.1 to −0.5 V voltage (with 0.01 V increments) and measuring current, using a Suss PA200 probe station and an Agilent 4142B tester. Conductivity was then calculated as: \( \sigma = \frac{L}{R \cdot W \cdot h} \), where \( R \) is the measured resistance, \( L, W \) and \( h \) are length (15 mm), width (2 mm) and thickness (20 µm) of the rectangle. We used thickness as measured by SEM and did not include PEDOT:PSS infiltrated in the membrane.

Cyclic voltammetry (CV) measurements were performed using an Autolab potentiostat PGSTAT30 (Metrohm Autolab) coupled with NOVA 1.8 software at room temperature (23 ± 1°C). A two-electrode configuration was adopted (equivalent to the one used for bending characterization), in which the two surfaces of PEDOT:PSS were working and reference electrodes. CV measurements were performed at various voltages and scan rates. The specific capacitance was calculated as \( C = \frac{i \cdot \Delta V}{2m \cdot \Delta t} \), where \( i, \Delta V \) are measured current and applied potential, \( m \) and \( \Delta V \) are the scan rate, the mass of active materials and the potential window respectively. Electrochemical impedance spectroscopy (EIS) measurements were performed with the same equipment and in the actuator working configuration. The magnitude of the alternative signal was 10 mV and the measurements were done in a frequency range between 100 kHz and 10 mHz.

Actuator bending was tracked by laser displacement sensor (optoNCDT 1302, MicroEpsilon) at a position of about 2 mm from the end of the actuator. Scheme of the set-up used for actuation and tracking is shown in figure 2(A). All measurements were performed at room temperature and humidity that were slightly varying. A laser displacement sensor (optoNCDT 1700, MicroEpsilon) was connected to NI-DAQmx for data logging and was also used for signal generation and current measurements. For creeping measurements, displacement measured by laser was used as a feedback to switch actuators position to 1 mm above or below the initial position by changing the polarity of the applied voltage (±1.5 V). The time needed to reach the set position was calculated for each half cycle.

3. Results and discussion

3.1. Harmonic response

A continuous displacement of the movement range, i.e. creeping, can be most easily observed during harmonic actuation as shown in online supplementary figure S2. Actuation starts with cycling in between about +1.2 mm and −1 mm and after 2 minutes or 25 cycles, the position shifts to +1 mm and −1.2 mm. Because solvent was not used for fabrication of our actuators, the most likely reason of creeping is unbalanced charging [22–24]. Transferred charge during harmonic actuation is shown in figure S(B). It was calculated
as integrated current transferred during actuation. In the case of pristine PEDOT:PSS without CNTC significant and fast creeping is observed. In less than 150 s one side of the PEDOT:PSS film continues being charged but is not completely discharged. That shows that charges are being accumulated and stored on one side of the trilayer.

On the other hand, actuators sprayed with CNTs work with lower displacement amplitude, but its position remains relatively stable (online supplementary figure S2). As shown in figure 2(B) harmonic response of the transferred charge for actuators with CNTCs is significantly more reversible and remains in the initial range (±1 mC). A less significant improvement in charging reversibility is also noticeable for actuators with sprayed SDS. After 150 s of actuation, charging and discharging remains in between about 0.8 mC and −2 mC.

The morphology of PEDOT:PSS and its compact structure might be one of the reasons behind irreversible expansion and charge accumulation, as previously suggested by Melling et al [20]. Harmonic cycling opens the polymer matrix so that with each cycle ions can diffuse faster and therefore deeper into the film. When polarity is reversed, some of the ions might remain there. In this case, spraying with a surfactant such as SDS could improve the reversibility of cycling by opening the compact PEDOT:PSS structure and subsequently facilitating ion transport through it.

Nevertheless, CNTCs show much more significant improvement in charge transfer reversibility. To better understand this actuation behaviour, various electro-chemical experiments were performed.

### 3.2. Bending characterization

Actuator performance was evaluated by tracking the displacement at its tip with the laser displacement sensor. The peak-to-peak displacement versus actuation frequency is shown in figure 3(A). Measurements were done by applying square voltage wave with the amplitude of 1.5 V and decreasing frequency of the wave after several cycles from 5 Hz to 10 mHz. At 1.5 V the performance of CNT actuators was poorer in comparison to simple actuators. For instance, at low frequencies, 10 mHz actuation without CNTCs produce 1.5 mm larger displacement amplitude. The results are surprising because most of the previously reported composites with carbon nanomaterials were improving the performance of actuators at all frequencies [34, 44, 45]. Furthermore, CNTCs also reduce the electrical resistance of PEDOT:PSS electrodes from 5.1 ± 3.7 kΩ to 1.6 ± 0.5 kΩ. We think that the increased stiffness of the actuators that was not measured in this study is the most likely explanation for a decrease in their response time [34].

Figures 3(B) and 3(C) show the comparison of the performance of the actuators without and with CNTC (respectively) at voltages in the range from 0.5 V to 2.0 V. In the case of actuators without CNTC, at low voltages below 1 V actuators were reaching a nearly stable position in less than 5 s. This is a consistent response of actuators working due to the charging of a double layer capacitance at the interface between the membrane and PEDOT:PSS as suggested by Okuzaki et al [46]. Only above 1.5 V PEDOT:PSS is starting to be reduced causing the expansion and bending due to the ion incorporation in the polymer film. This can be seen in figure 3(B), where slow continuous bending is shown even after 40 s of actuation. Furthermore, displacement of the actuator tip is less sensitive to the voltage change above 1 V. Results suggest two possibilities: (i) ionic conductivity of PEDOT:PSS is very low (ions do not move through the film), or (ii) charge transfer in PEDOT:PSS is limited.

Different responses to voltage are observed for actuators with CNTCs as shown in figure 3(C). For voltages lower than 1.5 V the actuator responds slower and reaches a slightly lower plateau value. That is, most likely, a consequence of an increased stiffness of the actuator [34]. Nevertheless, CNTCs seem to significantly improve the performance of the actuators when voltages are higher than 1.5 V. At 2 V a displacement of the actuator was significantly larger, 2.6 mm versus 3.2 mm in 50 s without and with CNTCs respectively, and seemed to be further increasing. This is an indication that CNTs positively affect the redox behaviour of PEDOT:PSS. Furthermore, the displacement of the actuators remains sensitive to the higher applied voltages (3.3 mm and 2.3 mm...
after 50 s for 2 V and 1 V respectively). Results suggest that CNTCs also facilitate the ion transport to the film.

### 3.3. Electrochemical characterization

In order to further explore the electrochemical properties of actuators, cyclic voltammetry (CV) measurements were performed in an actuator working configuration and are shown in figure 4(A). As expected from previous results, redox behaviour was noticeable for voltages higher than 1 V. Actuators with CNTC show a significantly larger increase in current at higher voltages confirming their influence in improving redox reaction. The specific capacitance of the actuators is at least 25% higher when CNTCs are used (shown in online supplementary figure S3). Furthermore, the shape of oxidation and reduction peaks is also more similar when CNTCs are used. This suggests improvement in reversibility of the redox reaction. As a reference, CV measurements were also performed for actuators with only surfactant (SDS) sprayed on the surface. SDS does not alter the specific capacitance of the actuator nor show any improvement to redox behaviour. These CV responses imply that CNTs have an influence on improving the redox activity of the conducting polymer based actuators and their reaction reversibility.

As previously mentioned, in addition to redox behaviour, cycling reversibility and creeping could also be influenced by the morphology of PEDOT:PSS films. In order to evaluate ionic conductivity of PEDOT:PSS films with and without a CNTC, EIS was performed at the actuator working configuration with the potential amplitude of AC kept at 10 mV and in the frequency range from 100 kHz to 10 mHz. Results presented as a Nyquist plot are shown in figures 4(B)–(C). Quantitative data fitting and extraction of conductivities were unsuccessful due to the complex quasi-trilayer structure of the actuator [41]. Nevertheless, actuators with CNTCs had lower resistance at high frequency (in a range between 110 Ω and 140 Ω). Our results suggest that a PEDOT:PSS electrode with CNTCs has a higher combined ionic and electronic conductivity.

We have already mentioned that CNTCs reduce the resistance of PEDOT:PSS electrodes from $5.1 \pm 3.7 \text{kΩ}$ to...
1.6 ± 0.5 kΩ. On the other hand spraying PEDOT:PSS with just SDS solution also decreases the resistivity of the film to 1.5 ± 0.4 kΩ. It was previously reported that SDS improves the electrical conductivity of PEDOT:PSS by altering its morphology towards a more swollen and less compact core/shell structure [47]. We think that a less dense morphology also improve the ionic conductivity as it allows easier ion penetration and movement through the PEDOT:PSS film [20]. On the other hand, the difference in EIS response between pristine PEDOT:PSS and PEDOT:PSS with SDS is not significant. That says that even though SDS might play a role in increasing the overall conductivity of actuators, CNTCs (i.e., an improvement in the redox reaction of PEDOT:PSS) is significantly more important.

### 3.4 Creeping reduction

To sum up, we have demonstrated that CNTCs significantly improve the redox behaviour of PEDOT:PSS and its overall electrical and ionic conductivity. Both effects are known to contribute to cycling reversibility and could explain reduced charge accumulation shown in figure 2(B). To demonstrate the importance of reversible cycling on the long term performance of conducting polymer actuators we actuated CPAs with and without CNTCs using feedback from a laser displacement sensor (creeping during harmonic cycling eventually leads to actuation out of the laser measurement range). Actuation asymmetry and creeping were estimated by comparing the time needed for the actuator to do a half cycle, i.e. to bend from one extremum to another. The voltage polarity (±1.5 V) was changed every time the actuator got to the set ±1 mm value. The time needed for the actuator to reach these positions was then extracted and is plotted versus actuation time in figure 5(A).

In the case of the actuator with no CNTC (full red line in figure 5(A), the time needed for the tip of the actuator to move one direction is nearly five times longer than the other. This time difference is further increased and after 1400 s it takes 10 s for the actuator to reach the lower extreme. The difference in time needed to move the actuator upwards or downwards can also be clearly seen in figure 5(B) as a filled area. After less than 2000 s the position of actuator without a CNTC was reset as it was not reaching the lower limit of −1 mm. On the other hand, actuators with a CNTC demonstrated nearly the same time for each half-cycle, and it remained rather constant for more than 2200 cycles or 1 hour of actuation (dashed blue line in figures 5(A)–(B)). This remarkable improvement in actuator performance is in agreement with our previous discussion—CNT carpets increased the redox reaction reversibility that consequently reduced charge accumulation in the PEDOT:PSS film leading to reduced creeping of the actuator.

### 4. Conclusions

We have demonstrated that CNTCs deposited by spray coating could be used to improve conducting polymer based artificial muscles. We suggest that a CNTC increases the redox activity of PEDOT:PSS leading to better performance at voltages above 1.5 V. Furthermore, together with an ionic surfactant, i.e., SDS, CNTs increase the overall electrical and ionic conductivity of PEDOT:PSS, consequently improving the redox cycling reversibility and reducing the charge accumulation in the polymer film. These effects lead to a
significantly reduced electrochemical creeping of conducting polymer based actuators. It is the first time a method for limiting electrochemical creeping of conducting polymer based actuators has been suggested. As creeping is one of the main limitations of conducting polymer based devices, this could be a significant step forward towards their commercial applications. Finally, spray coating is a simple and a versatile technique that can be used as an additional step after any other conducting polymer deposition.

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