# Toward Stretchable Self-Powered Sensors Based on the Thermoelectric Response of PEDOT:PSS/Polyurethane Blends

Prospero J. Taroni, Giovanni Santagiuliana, Kening Wan, Philip Calado, Manting Qiu, Han Zhang, Nicola M. Pugno, Matteo Palma, Natalie Stingelin-Stutzman, Martin Heeney, Oliver Fenwick, Mark Baxendale, and Emiliano Bilotti\*

P. J. Taroni, G. Santagiuliana, K. Wan, Dr. H. Zhang, Prof. N. M. Pugno, Dr. O. Fenwick, Dr. E. Bilotti School of Engineering and Material Sciences Queen Mary University of London Mile End Road, E1 4NS London, UK E-mail: e.bilotti@qmul.ac.uk

P. Calado

School of Physics Imperial College London London SW7 2AZ, UK

Dr. M. Baxendale, M. Qiu

School of Physics and Astronomy Queen Mary University of London Mile End Road, London E1 4NS, UK

Prof. N. M. Pugno

Laboratory of Bio-inspired & Graphene Nanomechanics Department of Civil, Environmental and Mechanical Engineering University of Trento Via Mesiano 77, 38123 Trento, Italy

Ket-Lab Edoardo Amaldi Foundation Italian Space Agency Via del Politecnico snc, 00133 Rome, Italy

Dr M. Palma

School of Biological and Chemical Sciences Queen Mary University of London Joseph Priestley building, Mile End Road, London F1 4NS. UK

Prof. N. Stingelin-Stutzman

School of Materials Sciences Georgia Tech 771 Ferst Drive, J. Erskine Love Building, Atlanta GA 30332-0245, USA

Prof. M. Heeney

School of Chemistry and Centre of Plastic Electronics Imperial College London London SW7 2AZ, UK

The development of new flexible and stretchable sensors addresses the demands of upcoming application fields like internet-of-things, soft robotics, and health/structure monitoring. However, finding a reliable and robust power source to operate these devices, particularly in off-the-grid, maintenance-free applications, still poses a great challenge. The exploitation of ubiquitous temperature gradients, as the source of energy, can become a practical solution, since the recent discovery of the outstanding thermoelectric properties of a conductive polymer, poly(3,4-ethylenedioxythiophene)- poly(styrenesulfonate) (PEDOT:PSS). Unfortunately the use of PEDOT:PSS is currently constrained by its brittleness and limited processability. Herein, PEDOT:PSS is blended with a commercial elastomeric polyurethane (Lycra), to obtain tough and processable self-standing films. A remarkable strainat-break of ≈700% is achieved for blends with 90 wt% Lycra, after ethylene glycol treatment, without affecting the Seebeck voltage. For the first time the viability of these novel blends as stretchable self-powered sensors is demonstrated.

#### 1. Introduction

The fast developing field of wearable electronics for autonomous monitoring is combining smart textile technology with state of the art wireless sensing communication, and it could soon experience exponential growth as new applications are being considered for fields such as internet-of-things, soft robotics, medicine, defence, advertising, fashion, and many others.[1–5] On the other hand, numerous challenges are still to be overcome, including the need for improved materials that are not only responsive to specific stimuli but can also conform to the objects monitored, for instance being flexible and stretchable, as well as durable.[6] Another important requirement is to make the sensors autonomous and selfpowered, particularly desirable in off-grid, maintenance-free conditions, where battery replacement is not practical, economical or feasible.[5] There have been different research strategies for finding sustainable, small scale energy harvesting systems to recharge batteries or to directly power sensor devices, with long lasting and reliable supplies. This has been demonstrated so far using triboelectric, [7– 11] photovoltaic,[5,12] piezoelectric,[5,13,14] radiofrequency,[14] thermoelectric (TE) systems, [14-18] and others. [19-21] In the case of TE technology, conventional systems have been considered as a power source for integrated sensors of pressure, [22] corrosion, [23] heat flow,[24] vibration,[25] heart rate,[26] and other stimuli.[27-31] The utilization of conjugated polymers as the active component of a TE device is a relatively recent concept, [32-34] but it is rather attractive where flexibility is needed, enabling a new generation of novel, low-cost, lowpowered wearable, or even remote, sensors. There are many desirable attributes driving the investigation of polymer TE materials, such as low toxicity, ubiquity, and abundance of constitutive elements, as well as ease in processing by various established coating or printing techniques. Poly(3,4-ethylenedioxythiophene)-poly(styrenesulfonate) (PEDOT:PSS) is at the centre of the effort to achieve thermoelectric power conversion efficiencies comparable to those of inorganic thermoelectric materials. In 2003, PEDOT was reported to have a dimensionless figure of merit (zT value) as high as 0.42.[35] This result is within close reach of the best commercial inorganic materials used for room temperature applications, such as bismuth telluride compounds, with zT value around 1.[36–38] Promising results were also obtained by combining PEDOT, or other conductive polymers (e.g., P3HT and PANi), with carbon nanotubes and other nanoparticles.[39-42] PEDOT was further studied as a p-type TE material,[43-53] and shown to be sensitive to stimuli like temperature and pressure.[54] However, a number of challenges still restrict its commercial exploitation, including low toughness and tear resistance, in addition to processing constraints, limited to film casting from water solution, while bulk (greater than tens of microns thick) processing is needed to fulfill current TE device design requirements. Here we propose a novel strategy for overcoming the above limitations, consisting of blending PEDOT:PSS with a commercial polyurethane (Lycra), followed by post-treatment of the films with ethylene glycol (EG), a polar solvent known to improve the thermoelectric behaviour of PEDOT:PSS.[37] After a systematic study of blend composition, film morphology, as well as mechanical and TE characterization, a new set of materials were obtained. They are stretchable and tough, of relative low cost and potentially compatible with bulk industrial

processing (e.g., extrusion, melt spinning), while maintaining the same Seebeck coefficient values as pure PEDOT:PSS. The sensing efficiency of these blends were then studied with the use of an external power source, where the resistance is determined and correlated with the changes in film geometry when responding to strain. Furthermore, with a temperature difference across the end of the film, the resultant thermovoltage was found to produce a measurable sensing signal which was used to detect changes in strain as well as ambient temperature and/or air flow. The results presented in this paper promise to be a viable solution for applications such as selfpowered sensors.

#### 2. Results and Discussion

In this study, freeze-dried PEDOT:PSS and the elastomeric Lycra yarn were separately dispersed in dimethyl sulfoxide (DMSO), then mixed together at different proportions, aiming at maximizing mechanical behavior and strain-sensing capabilities. Free-standing films were produced by drop casting onto glass substrates. Post-treatment was performed by immersing the films in an EG bath.

#### 2.1. Mechanical Properties

The mechanical properties of the blend films have been studied using quasi-static tensile tests and presented in Figure 1a,b, showing typical stress—strain responses as a function of Lycra content. The strain-at-break value increases with increased concentration of elastomer, from  $4\pm1\%$  at 0 wt% Lycra (pure PEDOT:PSS) to  $300\pm230\%$  at 90 wt%, while the elastic modulus is inversely correlated, going from  $1300\pm250$  MPa for pure PEDOT:PSS to  $23\pm11$  MPa at 90 wt% Lycra. Thus blending clearly improve the ductility of PEDOT:PSS. A rather large deviation in results for films with high elastomer content was observed. This is possibly due to microstructural inhomogeneity of films related to subtle variation in the casting temperature, which was demonstrated to significantly affect phase-segregation of the components (Section S1, Supporting Information). Some of the samples containing 90 wt% Lycra were also immersed in an EG bath after casting. They exhibited outstanding improvements in elastic modulus and strainat-break, with values of  $75\pm65$  MPa and  $700\pm150\%$ , respectively. These mechanical properties are unprecedented for a TE material, and closely approach the ones of neat elastomeric yarns (Section S2, Supporting Information).

## 2.2. Thermoelectric Properties

A simplified way to quantify the ability of a given material to generate useful thermoelectricity, is the measurement of the power factor (PF), expressed in Equation (1) PF S2 =  $\times \sigma$  (1) where S is the Seebeck coefficient, the voltage difference generated by unit of temperature, and  $\sigma$  is the electrical conductivity. The power factor, electrical conductivity, and Seebeck coefficient of all the blends are presented in Figure 1c. The electrical conductivity of the blended films decreases nearly linearly with increasing proportion of elastomer, from 635  $\pm$  8 S cm-1 for unmodified

PEDOT:PSS to 62 ± 4 S cm-1 for 90 wt% Lycra. These relatively high values can be explained by the effect of the chosen solvent (DMSO), which is known to improve the electrical conductivity of PEDOT:PSS.[55] Further improvements can be achieved if films are treated with an EG bath, giving up to 26% increase in electrical conductivity for composites containing 90 wt% Lycra, reaching an averaged value of 79 ± 5 S cm-1 . Indeed, EG treatment consistently resulted in improved conductivity, as summarized in Section S3 in the Supporting Information. EG has been previously shown to help controlling the film morphology through removal of excess insulating PSS.[48] The X-ray diffractograms presented in Section S4 in the Supporting Information confirm extraction of excess PSS from the film through a reduction of signal intensity for the peak at  $2\theta \approx$ 18°. The Seebeck coefficient stayed relatively constant, invariant with the proportion of Lycra (Figure 1c). This is of great importance because blends with high proportions of elastomer will be able to produce a similar thermovoltage as neat PEDOT:PSS (useful for self-power capability), while gaining significant elasticity and toughness (as demonstrated in Section 2.1), Furthermore, the Seebeck coefficient of PEDOT:PSS is apparently not reduced by EG treatment (Section S5, Supporting Information). As for the electrical conductivity, also the power factor follows a similar declining trend with increasing proportion of Lycra. Therefore, in this case, the electrical conductivity has a stronger influence than the Seebeck coefficient on PF.

## 2.3. Morphology

For a better understanding of the mechanical and thermoelectric properties presented in this paper, a systematic study of blend composition and film morphology was undertaken. Optical transmission studies of the films with varying composition revealed no evidence of any chemical reaction upon blending (Section S6, Supporting Information). The study of atomic force microscopy (AFM) imaging, in quantitative nanomechanical analysis mode, was used to investigate the morphology of the films, after EG treatment, shown in Figure 2. These images depict the relative elastic moduli of the film surfaces and clearly show the dispersion of Lycrarich regions (low elastic modulus) into PEDOT:PSS-rich regions (high elastic modulus), with a continuous change in morphology between the two pure materials. Nanomechanical imaging offers in this case an improved contrast as compared to topographymode, owing to the significant difference in elastic modulus between the two polymers. Further studies of AFM imaging revealed the Lycra-rich regions to exhibit greater deformation, adhesion, and dissipation than PEDOT:PSS-rich regions (Section S7a, Supporting Information). A low degree of connectivity between domains, for films with less than 60 wt% Lycra, is also shown in Figure 2, which we would expect to have detrimental effects on elasticity and toughness, as the elastomer may not be forming a continuous network, leading to more brittle and rigid materials. Yet this is not what we observed in macroscopic tests. Since these are surface images, so we cannot rule out an extended Lycra component embedded within the film structure. The study of topography-mode AFM imaging (Section S7b, Supporting Information) revealed a root mean square (rms) roughness of 540 nm for blended films of thickness ≈10 µm, which is more than an order of magnitude greater than that of commercial PEDOT:PSS cast from aqueous suspension. For the pure Lycra films, a value of 350 nm was found. The top surface and cross-sectional area of the blended films,

before and after EG treatment, were also studied by scanning electron microscopy (SEM), presented in Section S8 in the Supporting Information. A trend toward compact structure with increasing Lycra content is evident, which could explain the decrease in roughness observed by AFM. No distinct microstructural differentiation is apparent in the images of films before and after EG treatment. Some isolated PEDOT:PSS domains are visible in the cross-section of films with 90 wt% Lycra, although the conductive pathway may not be a sole result of these apparently discontinuous sites. As previously discussed in Section 2.1, it is possible that the components partly phasesegregate during casting of the films.

# 2.4. Externally Powered Strain Sensing

Sensitivity to strain was determined by a bespoke test set-up, in which the electrical resistance (R) is measured by passing a current from an external power supply through the sample while simultaneously subjecting it to a given strain profile. These tests were performed on films containing 90 wt% Lycra, previously subjected to EG treatment, owing to the high values of elasticity and electrical conductivity. As shown in Figure 3a,b, the modulation of resistance is up to 20% and 80% for strains of 10% and 50%, respectively, and this behavior can be explained with a simple geometrical effect. By assuming incompressibility (Poisson ratio of 0.5),[56] and a constant resistivity, one can calculate the reduction of sample cross-sectional area for a given extension, and hence a theoretical resistance (RT), (Section S9, Supporting Information). The variation of R with strain is very well predicted by this simple model. Nonetheless, for strains higher than 50%, the values of R are higher than RT, indicating change in intrinsic resistivity may also be involved (Section S10, Supporting Information). This can be explained by a possible modification of the conductive network. At relatively high strains (>80%), the sample exhibited plastic deformation, with a permanent increased resistance. A strategy to overcome this limitation, while increasing the strain sensitivity further still, can come from engineering the conductive network morphology. It has been noticed that some samples with lower elastomer content (50 wt% Lycra) and prepared from poorly dissolved Lycra presented some intact short filaments from the original yarn in the final composite film (Figure 3c). In these cases, the filaments indirectly control the morphology of the conductive network, in terms of excluding a volume in which PEDOT:PSS can be dispersed, but also in guiding its dispersion, which seems to preferentially coat the filaments (Figure 3d). Once the composite film cracks, after exceeding a threshold strain of ≈10%, filaments coated with PEDOT:PSS were still observed connecting the two extremities of the cracked sample (fibre bridging), bearing the load. The strain sensing tests for these samples show an interesting bimodal behavior. For strains lower than the threshold, the material is not sensitive, while for higher strain (after crack formation), the strain, sensitivity suddenly increases (with an outstanding Gauge Factor of ≈17 000-20 000 when 10% strain is applied). The strain, localised on the bridging filaments, determines the pathway for charge carrier transport (Figure 3e). Interestingly, this change in resistance is reversible, with the normalised resistance going back to initial values before cracking, when the strain decreases to zero. In future, we envisage the development of new materials inspired by these intriguing results, where, for instance, Lycra fibres are coated with PEDOT:PSS, which can then be used for

conductive film reinforcement or as a sensing device on its own. Initial tests have already started with encouraging initial results (Section S11, Supporting Information).

## 2.5. Self-Powered Strain Sensing

To demonstrate the concept of a self-powered strain sensor based on the thermoelectric effect, a film containing 90 wt% Lycra was exposed to a constant temperature difference of 30 °C, while uniaxially deformed (Figure 4a). When the sample was series-connected to a 10 k $\Omega$  load resistor, a thermocurrent was induced by the Seebeck voltage. It is noted that the load resistance of 10 kΩ was selected to match the internal specimen resistance at high strains, hence maximizing the power output. However, the internal resistance does change with the applied strain (Section S10, Supporting Information). So, for a more efficient self-powered strain sensor, the load resistance should ideally be modulated with the strain. Nevertheless, Figure 4c demonstrates that large strains, above 80%, can clearly be sensed by a simultaneous change in voltage and current. In order to explain this, it is observed that the thermovoltage generated is assumed constant with strain (Section S12, Supporting Information), while the resistance of the sample varies. Hence the thermocurrent generated, passing through the external circuit, also varies. Considering the equivalent circuit in Figure 4b, the voltage measured across the load resistor is proportional to the load resistance only, and therefore it changes proportionally to the thermocurrent variation, in agreement with measurements presented in Figure 4c. But the selfpowered sensor does not respond only to strain. In fact it shows sensitivity to other stimuli, for instance, ambient temperature and/or air currents (Section S13, Supporting Information). It is noted that the sample in Figure 4, can harvest a power output of up to few nW (Section S14, Supporting Information) for a ΔT of 30 °C. Increasing the power output to the level of 10–100 μW, necessary for an integrated self-powered system (including data trasmission for instance) can be achieved by: (i) increasing the temperature gradient, (ii) connecting our p-type TE material in series with a n-type one, to form a p-n junction, (iii) connecting many p-n junctions in series.

#### 3. Conclusion

This work demonstrates a strategy to develop stretchable self-powered sensors, via blending an elastomeric polyurethane (Lycra) with the best current organic thermoelectric material, PEDOT:PSS. In doing so, the main technological constrains of PEDOT:PSS, namely brittleness, processability, and costs, have, at the same time, been overcome. An unprecedented strain at break of 700  $\pm$  150% has been reached, for a blend containing 90 wt% Lycra, after EG bath, while maintaining high electrical conductivity (79  $\pm$  5 S cm-1 ) and Seebeck coefficient (16  $\pm$  1  $\mu$ V K-1 ). This novel material has shown sensitivity to different stimuli including strain, as well as ambient temperature and/or air flow.

## 4. Experimental Section

PEDOT:PSS Clevios PH1000 (from Heraeus GmbH) was freeze dried (using a Benchtop Pro with Omnitronics from SP Scientific), then dispersed in anhydrous DMSO > 99.9% (from Sigma Aldrich) by homogenisation (Heidolph SilentCrusher M set at 500 rpm for 5 min) followed by probe sonication (Sonics VibraCell, amplitude 35%, 1 s on and 1 s off, for 20 min), making up a dispersion with 1 wt% solid content. This process could not be reproduced as efficiently if the PEDOT:PSS was dried by applying heat to the solution, which leads to a compacted dry material. On redispersing with DMSO, a nonhomogeneous casting solution was formed, as compared to the ones using high surface area foam (Section S15A, Supporting Information). Spandex yarn (Lycra Invista) (Section S15B, Supporting Information) was also dispersed in anhydrous DMSO by stirring at 80 °C for 12 h, to afford a dispersion with 1 wt% solid content. Both Lycra and PEDOT:PSS dispersions were mixed together at different proportions to make up blends of 1 wt% load, with Lycra content varying from 0% to 100% (Section 15C, Supporting Information). All mixtures were magnetically stirred for a further 24 h at room temperature, drop cast and dried at least ten consecutive times on precleaned glass slides, using an oven set at 80 °C. Films were then mechanically removed from the glass substrate. The drying temperature was chosen on the basis of thermal gravimetric analysis results using a TA Instruments Q500 Thermogravimetric Analyzer, flushed with air (60 mL min-1), shown in Section S16 in the Supporting Information, with further consideration to the effect of different casting temperatures on the morphology and electrical conductivity (Section S1, Supporting Information). Mechanical tests were performed using a quasi-static tensile tester (Instron 5566, with a rate of 0.5% per minute, then 10% per minute for 90 wt% Lycra blends after the first 5% strain, in this way to control the timing of the test). Films were prepared as described above, then cut at ≈50 mm × 5 mm surface area. The results for % strain at break and elastic modulus were averaged out of 6 to 10 individual samples, whereas the presented curves were selected from representative results for each composition. Fouriertransform infrared spectroscopy (FTIR) spectrum of solution blends were recorded using a Bruker Tensor 27, with the DMSO signal eliminated from background. UV-vis spectra of blends were recorded using a Perkin Elmer Lambda35. In this instance, the DMSO is not eliminated from background spectrum. AFM images were obtained with a Bruker Dimension Icon operating in quantitative nanomechanical analysis mode on samples drop casted a single time, having films with thickness of about 10 μm. SEM images were taken with a FEI Inspect F equipped with energy dispersive x-ray spectrometry (EDX). Samples were prepared by consecutively drop casted films (about 10%), with thickness of about 100 μm. X-ray diffraction (XRD) signals recorded with a diffractometer system XPERT-PRO using K-alpha of 1.54 Å. A home built four point probe (0.25 mm probe space) is used for electrical conductivity measurements on the substrate, at room temperature and ambient atmosphere using an Agilent 6614 System DC power supply integrated with a Keithley 6485 picoammeter, connected to the external probes. The voltage generated when current passes through the sample is then measured (high impedance Keithley 2000 Multimeter) between the two internal probes. For this purpose, films are prepared by single drop cast, with thickens measured using a profilometer (Bruker Dektak Vision 64), and calculated electrical conductivity averaged out of four films from each blend composition, and measured five times at different points within the centre of the film. A Seebeck System was used for

Seebeck measurements at 300 K under nitrogen atmosphere (MMR, SB100 digital Seebeck controller and K20 digital temperature controller), with self-standing films cut to the dimensions required by the instrument (≈1 mm × 5 mm surface area). The averaged result is calculated out of three films made from each blend composition and measured at least ten times each. The electrical conductivity and Seebeck coefficient were used for calculation of the power factor, based on Equation (1). For cyclic test, as well as the self-powered strain sensing experiments, films were prepared as described above for mechanical test. These prototype materials were then fitted with copper electrodes on the extremities, at a distance of about 40 mm apart, then padded using carbon conductive tabs (12 mm diameter from Agar Scientific), to avoid the material being pulled from the electrode during stretching. A constant voltage of about 0.5 V was applied during the cyclic tests using an Agilent 6614 System DC power supply, while current variations with strain recorded using a coupled Agilent 34401A 6 ½ digital multimeter. Details of theoretical calculation model for resistance changes during the cycles were explained in Section S9 in the Supporting Information. For self-powered tests, heat is applied to one side of the device to generate a thermovoltage, by using a silicon heat pad (RS 12V, 7.5W, item 245–556) connected to a power supply (Farnell D30-2 set at 12V). This thermovoltage is recorded using a high impedance voltameter (Keithley 2000 Multimeter) and the current recorded using a Keithley 6485 picoammeter. Variable strain rates were used, depending on the % applied, aimed for controlling the time of test.

# **Acknowledgements**

The authors are grateful to the EPSRC Doctoral Training Centre in Plastic Electronics (grant number EP/G037515) for funding this project, with additional financial support from European Thermodynamics Ltd. O.F. acknowledges the Royal Society for support in the frame of his University Research Fellowship. N.M.P. acknowledges support from the European Commission H2020 under the Graphene FET Flagship (WP14 "Polymer Composites" No. 696656) and under the FET proactive ("Neurofibres" No. 732344).

## **Keywords**

organics thermoelectricity, polymer blends, self-powered, sensors

#### References

- [1] R. F. Service, Science 2003, 301, 909.
- [2] M. J. Cima, Nat. Biotechnol. 2014, 32, 642.
- [3] W. Zeng, L. Shu, Q. Li, S. Chen, F. Wang, X. M. Tao, Adv. Mater. 2014, 26, 5310.

- [4] A. Tricoli, N. Nasiri, S. De, Adv. Funct. Mater. 2017, 27, 1605271.
- [5] T. Dias, Electronic Textiles: Smart Fabrics and Wearable Technology. Woodhead Publishing, Cambridge, UK 2015.
- [6] Stanford Wearable Electronics Initiative, https://wearable.stanford.edu (accessed: July 2017).
- [7] S. Y. Kuang, J. Chen, X. B. Cheng, G. Zhu, Z. L. Wang, Nano Energy 2015, 17, 10.
- [8] M. Ha, J. Park, Y. Lee, H. Ko, ACS Nano 2015, 9, 3421.
- [9] H. Zhang, Y. Yang, Y. Su, J. Chen, C. Hu, Z. Wu, Y. Liu, C. Ping Wong, Y. Bando, Z. L. Wang, Nano Energy 2013, 2, 693.
- [10] Q. Shi, H. Wang, T. Wang, C. Lee, Nano Energy 2016, 30, 450.
- [11] J. Wang, S. Li, F. Yi, Y. Zi, J. Lin, X. Wang, Y. Xu, Z. L. Wang, Nat. Commun. 2016, 7, 12744.
- [12] R. Vyas, V. Lakafosis, H. Lee, G. Shaker, L. Yang, G. Orecchini, A. Traille, M. M. Tentzeris, L. Roselli, IEEE Sens. J. 2011, 11, 3139.
- [13] Y.-K. Fuh, H.-C. Ho, Nanotechnology 2016, 27, 95401.
- [14] N. S. Hudak, G. G. Amatucci, J. Appl. Phys. 2008, 103, 101301.
- [15] A. R. M. Siddique, S. Mahmud, B. Van Heyst, Renewable Sustainable Energy Rev. 2017, 73, 730.
- [16] E. A. Mondarte, V. Copa, A. Tuico, C. J. Vergara, E. Estacio, A. Salvador, A. Somintac, Mater. Sci. Semicond. Process. 2016, 45, 27.
- [17] Y. Yang, Z. H. Lin, T. Hou, F. Zhang, Z. L. Wang, Nano Res. 2012, 5, 888.
- [18] J. J. Kuchle, N. D. Love, Measurement: J. Int. Meas. Confed. 2014, 47, 26.
- [19] L. Xie, M. Cai, IEEE Pervasive Comput. 2014, 13, 42.
- [20] J. Zhong, Q. Zhong, Q. Hu, N. Wu, W. Li, B. Wang, B. Hu, J. Zhou, Adv. Funct. Mater. 2015, 25, 1798.
- [21] A. Dewan, S. U. Ay, M. N. Karim, H. Beyenal, J. Power Sources 2014, 245, 129.
- [22] B. Sümer, E. K. San, K. Sancakdar, Procedia Eng. 2016, 168, 63.
- [23] N. N. Aung, E. Crowe, X. Liu, ISA Trans. 2015, 55, 188.
- [24] I. Korhonen, R. Lankinen, Measurement: J. Int. Meas. Confed. 2014, 58, 241.
- [25] C. Wang, K. Zhao, Q. Guo, Z. Li, IET Circuits, Devices Syst. 2016, 10, 147.
- [26] V. Leonov, P. Fiorini, T. Torfs, R. Vullers, C. Van Hoof, Microelectronics Journal 2011, 42, 579.

- [27] V. Leonov, T. Torfs, P. Fiorini, C. Van Hoof, IEEE Sens. J. 2007, 7, 650.
- [28] S. Dalola, M. Ferrari, V. Ferrari, M. Guizzetti, D. Marioli, A. Taroni, IEEE Trans. Instrum. Meas. 2009, 58, 99.
- [29] D. Davila, A. Tarancon, C. Calaza, M. Salleras, M. Fernandez-Reglez, A. San Paulo, L. Fonseca, Nano Energy 2012, 1, 812.
- [30] M. Guan, K. Wang, D. Xu, W.-H. Liao, Energy Convers. Manage. 2017, 138, 30.
- [31] Y. Shi, Y. Wang, Y. Deng, H. Gao, Z. Lin, W. Zhu, H. Ye, Energy Convers. Manage. 2014, 80, 110.
- [32] P. J. Taroni, I. Hoces, N. Stingelin, M. Heeney, E. Bilotti, Isr. J. Chem. 2014, 54, 534.
- [33] O. Bubnova, X. Crispin, Energy Environ. Sci. 2012, 5, 9345.
- [34] N. Dubey, M. Leclerc, J. Polym. Sci., Part B: Polym. Phys. 2011, 49, 467.
- [35] G. Kim, L. Shao, K. Zhang, K. Pipe, Nat. Mater. 2013, 12, 719.
- [36] H. Julian Goldsmid, J. Electron. Mater. 2017, 46, 2599.
- [37] T. J. Seebeck, Abh. Akad. Wiss. 1822, 289.
- [38] G. J. Snyder, E. S. Toberer, Nat. Mater. 2008, 7, 105.
- [39] C. Cho, K. L. Wallace, P. Tzeng, J. H. Hsu, C. Yu, J. C. Grunlan, Adv. Energy Mater. 2016, 6, 1502168.
- [40] C. Cho, B. Stevens, J. H. Hsu, R. Bureau, D. A. Hagen, O. Regev, C. Yh, J. C. Grunlan, Adv. Mater. 2015, 27, 2996.
- [41] C. Bounioux, P. Diaz-Chao, M. Campoy-Quiles, M. S. MartinGonzález, A. R. Goñi, R. Yerushalmi-Rozene, C. Müller, Energy Environ. Sci. 2013, 6, 918.
- [42] B. Dörling, J. D. Ryan, J. D. Craddock, A. Sorrentino, A. El Basaty, A. Gomez, M. Garriga, E. Pereiro, J. E. Anthony, M. C. Weisenberger, A. R. Goñi, C. Müller, M. Campoy-Quiles, Adv. Mater. 2016, 28, 2782.
- [43] A. M. Nardes, M. Kemerink, R. A. J. Janssen, J. A. M. Bastiaansen, N. M. M. Kiggen, B. M. W. Langeveld, A. J. J. M. van Breemen, M. M. deKok, Adv. Mater. 2007, 19, 1196.
- [44] K.-C. Chang, M.-S. Jeng, C.-C. Yang, Y.-W. Chou, S.-K. Wu, M. A. Thomas, Y.-C. Peng, J. Electron. Mater. 2009, 38, 1182.
- [45] C. Liu, B. Lu, J. Yan, J. Xu, R. Yue, Z. Zhu, S. Zhou, X. Hu, Z. Zhang, P. Chen, Synth. Met. 2010, 160, 2481.
- [46] T.-C. Tsai, H.-C. Chang, C.-H. Chen, W.-T. Whang, Org. Electron. 2011, 12, 2159.

- [47] A. M. Nardes, R. a. J. Janssen, M. Kemerink, Adv. Funct. Mater. 2008, 18, 865.
- [48] X. Crispin, S. Marciniak, W. Osikowicz, G. Zotti, A. W. D. van Der Gon, F. Louwt, M. Fahlman, L. Groenendaal, F. De Schryver, W. R. Salaneck, J. Polym. Sci., Part B: Polym. Phys. 2003, 41, 2561.
- [49] O. Bubnova, M. Berggren, X. Crispin, J. Am. Chem. Soc. 2012, 134, 16456.
- [50] O. Bubnova, Z. U. Khan, A. Malti, S. Braun, M. Fahlman, M. Berggren, X. Crispin, Nat. Mater. 2011, 10, 429.
- [51] T. Park, C. Park, B. Kim, H. Shin, E. Kim, Energy Environ. Sci. 2013, 6, 788.
- [52] T. Murakami, Y. Mori, H. Okuzaki, Trans. Mater. Res. Soc. Jpn. 2011, 168, 165.
- [53] J. Kim, J. Jung, D. Lee, J. Joo, Synth. Met. 2002, 126, 311.
- [54] F. Zhang, Y. Zang, D. Huang, C. Di, D. Zhu, Nat. Commun. 2015, 6, 8356.
- [55] Q. Wei, M. Mukaida, Y. Naitoh, T. Ishida, Adv. Mater. 2013, 25, 2831.
- [56] H. J. Qi, M. C. Boyce, J. Mech. Phys. Solids 2004, 52, 2187.

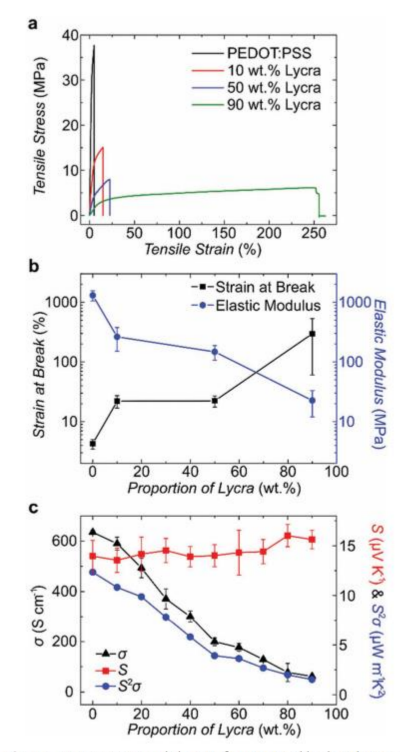


Figure 1. a) Stress vs strain behavior of PEDOT:PSS blends with various proportions of Lycra, b) elastic modulus and strain-at-break for the same blends, and c) electrical conductivity, Seebeck coefficient, and power factor of films of PEDOT:PSS blends with various proportions of Lycra (all graphical results are before EG treatment).

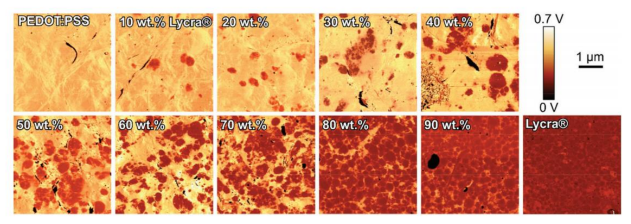


Figure 2. AFM images  $(4 \times 4 \mu m)$  of the PEDOT:PSS/Lycra film surfaces obtained in nanomechanical analysis mode. The colourscale is an indicator of the voltage signal from the instrument, which is proportional to the elastic modulus. Lycra-rich regions appear dark (low elastic modulus) and PEDOT:PSS-rich regions appear bright (high elastic modulus).

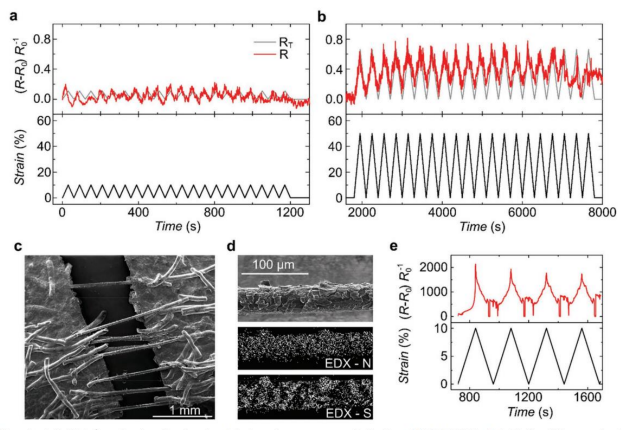


Figure 3. a)  $(R-R_0) R_0^{-1}$  as a function of strain, where  $R_0$  is the resistance at zero strain, for films of PEDOT:PSS blended with 90 wt% Latex previously subject to EG treatment, when 10% strain is applied, followed by 50% (Gauge factor  $\approx$ 1 and 1.3, respectively). The theoretical resistance  $R_T$  is shown in red. b) SEM imaging of a cracked film surface of 50 wt% Lycra used for strain sensing, showing stretched bridging filaments, c,d) SEM and EDX of one stretched filament for detection of sulphur (from PEDOT:PSS) and nitrogen (from Lycra), e) plot of  $(R-R_0) R_0^{-1}$  as a function of strain for this film, after cracking.

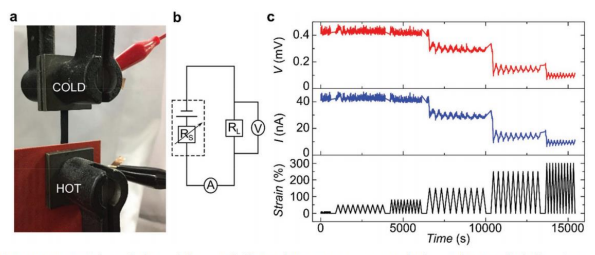


Figure 4. a) Experimental setup for the proof-of-concept of self-powered strain sensor measurements. b) The equivalent circuit for the film under test, the voltage source is the thermovoltage generated by the temperature difference across the ends of the sample,  $R_s$  is the internal resistance of the film and  $R_L$  is the external load resistance. c) The variation of voltage across and current through the load resistor with strain and time.