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3D Printing/Interfacial Polymerization Coupling for the Fabrication of Conductive Hydrogel

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E. Fantino,* I. Roppolo, D. Zhang, J. Xiao, A. Chiappone, M. Castellino, Q. Guo, C. F. Pirri, J. Yang*.....1700356

3D Printing/Interfacial Polymerization Coupling for the Fabrication of Conductive Hydrogel SD PRINTING

3D printing is coupled with interfacial polymerization to obtain electroactive hydrogels with complex and defined ge-ometry. Conductive hydrogels are created through a two-step procedure: first a digital light processing 3D printing system is used to fabricate 3D structures and then pyrrole is oxidized to polypyrrole, exploiting an interfacial polymerization mechanism, thus creating a conductive phase directly in 3D printed structure.

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FULL PAPER



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3D Printing/Interfacial Polymerization Coupling for the Fabrication of Conductive Hydrogel

Erika Fantino,* Ignazio Roppolo, Dongxing Zhang, Junfeng Xiao, Annalisa Chiappone, Micaela Castellino, Qiuquan Guo, Candido Fabrizio Pirri, and Jun Yang*

In this study, 3D printing is coupled with interfacial polymerization to obtain electroactive hydrogels with complex and defined geometry. Conductive hydrogels are created through a two-step procedure: first a digital light processing 3D printing system is used to fabricate poly(ethylene glycol)diacrylate 3D structure and then pyrrole is oxidized to polypyrrole (PPY), exploiting an interfacial polymerization mechanism thanks to which PPY can be formed into the poly(ethylene glycol) matrix, thus creating a conductive phase.

1. Introduction

26 Recently, conductive hydrogels have attracted much interest in 27 the field of biomaterials science being able to combine highly hydrated polymer structures with intriguing electronic function-28 alities.^[1,2] These hybrid materials, merging the hydrogel charac-29 30 teristics with the advantages of the conductive components such as electrical conductivity and electrochemical-redox properties, 31 32 have found applications across a range of biomedical applications such as biosensors, drug delivery, and tissue engineering.^[3,4] 33 In these composite hydrogels, conductivity was achieved either 34 incorporating electrically conductive fillers, such as graphite, 35 36 metallic particles, and carbon nanotubes,^[5-7] or integrating 37 intrinsically conductive polymers within the hydrogel matrix.

Conductive polymers are macromolecules with conjugated backbone; examples are polyacetylene, polythiophene, polypyrrole (PPY), or polyaniline. In recent years they have received great interest owing to their ability to conduct electricity and they have been used in several biomedical applications.^[4,8–11]

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Conductive hydrogels exploiting 12 intrinsically conducting polymers allow 13 obtaining materials that further expand 14 the application area of hydrogels^[12–14] 15 driving researchers to develop new fabrication methods. 17

The fabrication of conductive hydrogels 18 with particular geometry and controlled 19 features is still a big issue. 3D printing 20 is an additive manufacturing (AM) pro- 21 cess that enables to construct 3D objects 22 directly from a digital model. It has 23

received a great deal of attention from a diverse range of fields 24 including electronics,^[14] biomedics and regenerative medi-25 cine,^[15-17] and microfluidics.^[18] Objects are constructed layer-26 by-layer, enabling the creation of complex parts with tailored 27 morphology and functionality.^[19] There exist several different 28 3D printing techniques that differ from each other for the kind 29 of polymer that is used (i.e., thermoplastic or thermosetting) 30 and for the technology beyond the building process.^[16] One of 31 the most widely used 3D printing processes, namely, vat poly-32 merization, is based on photopolymerization. Stereolithography 33 (SLA), a well-established AM technology, is used to create ther-34 moset objects with features <100 µm by selectively scanning an 35 ultraviolet (UV) laser beam across a reservoir of photopolymer 36 resin.^[19] Another 3D printing process based on spatially con-37 trolled solidification of a liquid resin by photopolymerization is 38 digital light processing (DLP). Instead of using a laser to "draw" 39 the object under a point-by-point manner like in SLA, DLP 40 equipment projects an entire slice of an object using a digital 41 projector.^[20] Both techniques provide powerful tools to fabricate 42 complex 3D polymeric structures, with good resolution and fast 43 production times especially in the case of DLP. This opens infi-44 nite possibilities in their design and applications.^[21-25] 45

Many types of hydrogels have been reported in the literature 46 including natural materials or synthetic ones.[16,26,27] Among 47 the synthetic polymers, poly(ethylene glycol) (PEG) based 48 ones have been used extensively for the fabrication of hydro-49 gels thanks to their well-known hydrophilicity and biocompati-50 bility.^[28,29] With acrylated or methacrylated moieties, PEG mon-51 omers can be photocrosslinked in the presence of appropriate 52 initiating agents and the use of SLA or DLP has also been 53 reported as a successful method for fabricating complex func-54 tional 3D structures based on PEG monomer materials.^[23,30-33] 55 The choice of relatively low molecular weight ($M_{\rm w}$ 500–700) 56 PEG-based acrylate monomers allows preparing ractive formu-57 lations with sufficietly low viscosity to be processed with a fast 58 and low-cost DLP apparatus giving self-standing and robust 59

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structures thanks to the crosslinkind density reached in this 1 kind of network.^[34] Herein, the strategy we developed within 2 this study consists in coupling interfacial polymerization^[8,35–37] 3 4 with 3D printing in order to obtain electroactive hydrogel 5 with complex and defined geometry. After 3D printing of the 6 hydrogel structures, a chemical oxidative polymerization of pyr-7 role (PY) was performed for preparing conductive hydrogels. By 8 tuning the reaction conditions, the conductive components can 9 permeate spontaneously and exclusively into the hydrogels to achieve required conductivity.[5,38-40] 10

11 Moreover the dye used for obtaining the 3D printed structures was employed as dopant for PPY, enhancing the elec-12 trical conducitvity.^[6,41,42] If compared with other methods 13 proposed in the literature,^[12,13,43] this two-step approach can 14 provide a higher control over the structure of the conductive 15 16 hydrogel thanks to the formation of the conductive phase in a 17 dedicated step that allows maintaining the defined 3D microarchitecture of the printed hydrogel, bringing to very precise 18 19 and complex structures with electrical features. This approach has the potential to be extended to a wide variety of photo-20 sensitive hydrogels. The proposed strategy, coupling hydro-21 gels, 3D printing, and conductive polymers, could open new 22 23 paths for the development of new bioelectrical interfaces for several biomedical applications, like complex systems for drug 24 25 delivery or scaffolds for regenerative medicine but also in bio-26 energy fields or sensors.

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2. Experimental Section 30

2.1. Materials 31

32 33 Poly(ethylene glycol)diacrylate (PEGDA) with a molecular weight of 700 g mol^{-1} and PY reagent grade 98% 34 were purchased from Sigma-Aldrich and used as received. 35 Phenylbis(2,4,6-trimethylbenzoyl)phosphine oxide, selected as 36 photoinitiator (PI) for its fair absorbing characteristics in the 37 38 deep blue to near UV, and the dye, methyl orange (MO), were purchased from Sigma-Aldrich and used as received. Iron(III) 39 40 chloride hexahydrate (FeCl₃ 6H₂O) was purchased from Alfa Aesar.

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44 2.2. Fabrication of 3D PPy/PEGDA Structure

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3D printable mixtures containing PEGDA, MO 0.2 phr (per 46 47 hundred resin), and distilled water (dH2O) in different concen-48 tration (40, 50, 60, and 70% w/w) were prepared. 2 phr PI with 49 respect to the PEGDA amount was added in the formulation. 50 The 3D printing was performed on a PICOplus39, Asiga, with 51 an X-Y resolution of 39 μ m and light intensity of 30 mW cm⁻². 52 The digital models of structures were designed and converted 53 to sterolithography (STL) file format for 3D printing. The layer 54 thickness was set to 25 μ m and the exposure time varied from 2 55 to 15 s for increasing amounts of water.

56 Pyrrole monomers were dissolved in cyclohexane (0.5 M PY) 57 while ferric chloride was dissolved in deionized water (0.3 M 58 FeCl₃). First the 3D parts were soaked for 1 h in the aqueous 59 solution containing ferric chloride. The hydrogel was then carefully blotted and subsequently soaked in the organic solu-1 tion containing the pyrrole monomers. The monomer pyrrole 2 and the oxidant FeCl₃ diffuse at the aqueous/organic interface, 3 4 the polymerization occurs, and the PPY/PEGDA composite hydrogel is formed. The same procedure was conducted on the 5 thin film for electrical measurement.

2.3. Characterization Methods

Real-time rheological measurements were performed using an 11 Anton Paar rheometer (Physica MCR 302) in parallel plate mode 12 with a Hamamatsu LC8 lamp with a visible bulb and a cutoff 13 filter below 400 nm (light intensity was set to 30 mW cm⁻²). 14 The UV curing set up is equipped with a lower plate in quartz 15 that allows the irradiation of the sample during the measure-16 ment. The gap between the two plates was set to 0.1 mm and 17 the sample was kept at a constant temperature (25 °C) and 18 under constant shear frequency of 10Hz, light was turned 19 on after 120 s in order to stabilize the system. Concomitant 20 changes in viscoelastic material moduli during polymerization 21 were measured as a function of exposure time. The experiment 22 was performed in the linear viscoelastic region with a strain 23 amplitude of 0.5%. The morphological characterization of the 24 3D printed materials after the interfacial polymerization of PY 25 was carried out by scanning electron microscopy (SEM, Hitachi 26 27 TM3030Plus). ATR spectra were collected on a Tensor 27 FTIR Q3 spectrometer (Bruker). The averaged signal was collected with 28 a resolution of 2 cm⁻¹ from 4000 to 400 cm⁻¹. X-ray photoelec-29 tron spectroscopy (XPS) was carried out by using a PHI 5000 30 VersaProbe (Physical Electronics) system. The X-ray source was 31 a monochromatic Al K α radiation. Depth profile, by means of 32 an Ar+ flux at 2 kV accelerating voltage, was performed on the 33 sample in an alternate mode with sputtering cycles of 1 min 34 each. Spectra were analyzed using Multipak 9.7 software. 35 All core-level peak energies were referenced to C1s peak at 36 284.5 eV (C-C/C-H) and the background contribution in HR 37 38 scans was subtracted by means of a Shirley function. Differential scanning calorimetry (DSC) measurements were performed 39 with a Netzsch DSC 204 F1 Phoenix instrument, equipped with 40 41 a low-temperature probe, between -80 and 60 °C with a heating rate of 10 °C min⁻¹ in nitrogen atmosphere. For each sample, 42 the same heating module was applied two times and the final 43 heat flow value recorded during the second heating cycle. The 44 T_{g} was defined as the midpoint of the heat capacity change 45 observed in the DSC thermogram. Thermogravimetric analysis 46 (TGA) was performed using a Netzsch TG 209 F1 Libra instru-47 ment in the range between 25 and 700 °C, with a heating rate 48 of 10 °C min⁻¹ in nitrogen. Prior to the measurement all the 49 50 samples were dried overnight in a vacuum oven in order to avoid the drop of weight relative to the water/moisture eventu-51 ally present in the sample. Compression tests were carried out 52 53 using a dynamometer (Deben Microtest) equipped with a load 54 cell of 200N. Alveolar structures were tested, and at least three specimens for each sample were tested. Samples resistivity 55 was measured by using a Keithley-238 High Current Source 56 Measure Unit (voltage range ± 10 V, step 0.1 V), realizing a two-57 point contact setup placing copper electrodes on the two oppo-58 site basal sides. The data showed were obtained by multiple 59



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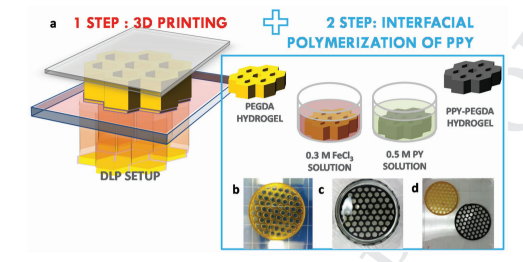


Figure 1. a) Sketch of the process. b) 3D printed PEGDA structure. c) PEGDA structure in PY/CYH solution. d) 3D PEGDA and 3D PEGDA/PPY structures.

measurements on different samples (three for each formulation). The electrical measurements on 3D printed structures were carried out with an apparatus composed of a Keithley 2750 multimeter and realizing a two-point contact setup, placing one copper electrode on each side of the structure.

3. Results and Discussion

In this work hybrid PPY/PEGDA conductive hydrogels were created through a two-step procedure. First a DLP 3D printing system was used to fabricate PEGDA microstructures and then PY was oxidized to PPY, exploiting an interfacial polymerization mechanism. Following this route, PPY could be formed into the PEGDA matrix, thus creating a conductive 3D printed hydrogel (Figure 1a).

The developed process is sketched in Figure 1a. 3D PEGDA hydrogel was printed by a DLP machine and then the 3D printed parts were soaked for 1 h in an aqueous solution containing ferric chloride: thanks to the high swell ability of PEGDA networks, the oxidant agent was incorporated in the 3D printed structure. The hydrogels were then carefully blotted and subsequently soaked again in an organic solution containing pyrrole monomers. The polymerization to PPY involves the oxidation of the pyrrole monomer with ferric ions, the pyrrole monomers and the initiator diffuse at the aqueous/ organic interface, and thus the polymerization is trigged. The reaction of PY with aqueous ferric chloride (FeCl₃) is rapid and 26 the PPY product is in the form of a black powder. After 1 h, the 3D parts are removed from the solution and immersed in dH_2O for at least 1 d to leach out low molecular weight components.

Different hydrogel formulations were prepared; MO was 30 used both as dopant for the PPY^[44,45] and as dye to improve 31 the 3D printing resolution.^[22-24] dH₂O was added in different 32 amounts (40, 60, and 70% w/w) in the formulations in order to adjust the concentration (Figure 2a); the higher is the amount 34 of initial water present in the formulation and, thus in the 35 printed part, the higher will be the hydrogel ability of swelling in the subsequent steps (oxidant solution) and therefore the interfacial polymerization will be probably more efficient. 38 Before printing the formulations, their reactivity was inves- 39 tigated by means of photorheology aiming to check the influ- 40 ence of the high amount of water added to the monomers on 41 the light-induced crosslinking reaction. As expected, the higher 42 the amount of water, the slowest the curing process as shown 43 by the time-dependent elastic modulus (*G*) values reported in

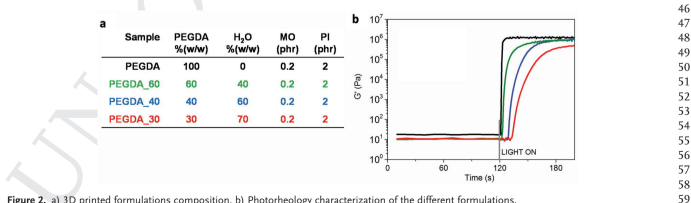
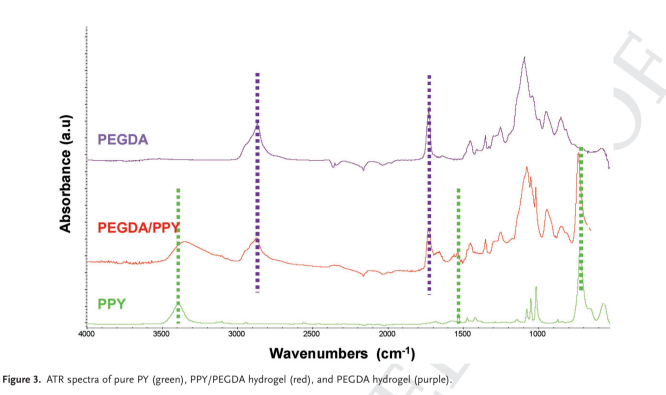


Figure 2. a) 3D printed formulations composition. b) Photorheology characterization of the different formulations.



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Figure 2b. While the G curve relative to neat PEGDA starts increasing almost instantaneously after turning on the light and reaches its postcuring plateau in few seconds, the other formulations show a delay in the beginning of the reaction and a slower kinetic revealed by the lower slope of the curves. This effect results in the need of a longer exposition time during the printing process when the formulations containing higher amounts of water are printed. The use of higher amounts of water also induces a slight reduction of the G modulus after curing; in all the cases the obtained values are in line with those reported for other crosslinked hydrogels measured in similar conditions (amplitude and frequency).^[46]

ATR spectroscopy was used to characterize the interfacial polymerization; the spectra are shown in Figure 3. The spectrum of neat PPY illustrates absorption peaks at 3402, 1528, 1418, and 1046 cm⁻¹ that could be assigned to the N-H, C-C, C-N stretching vibrations, and C-H in-plane vibrational bands of the pyrrole ring, respectively, and the peak at 1309 cm⁻¹ is assigned to C-N bonds.^[7] In the PEGDA spec-trum, the strong peak at 1724 cm⁻¹ could be assigned to the -C=O stretching from ester bonds, while the broad band cen-tered at 2862 cm⁻¹ can be assigned to the C-H (-CH, -CH₂, and CH₃) stretching vibration. In the FTIR of the PEGDA/ PPY hydrogel, it is possible to identify characteristic peaks of both polypyrrole, at 3396, 1528, 1465 cm⁻¹, and PEGDA, peak at 1724 and 2862 cm⁻¹, indicating the presence of both poly-mers blended. This confirms the successful formation of PPY on PEGDA matrix (Figure 3).

55 Detailed images of 3D printed structures with optical micro-56 scopy and SEM are shown in **Figure 4**. Well-defined architecture 57 and good building accuracy of complex objects were achieved; 58 details down to 200 μ m were achieved. This value represents 59 the experimental limit observed for this kind of water-based

formulations. Moreover, the PPY postpolymerization pro-cess on the printed structures does not worsen the resolution achieved during the printing step: the structures in fact remain intact without cracks. The surface after the polymerization of PPY was evaluated by SEM. While the surface of PEGDA struc-tures is smooth, a certain roughness was evident after PPY poly-merization, with a flowerlike pattern and big porosity (Figure 4c). Probably this could be related to the high concentration of PY monomers in the solution, which leads to a faster polym-erization rate and thus a more uneven distribution of PPY.^[47] Thicker samples (4 mm) were also produced and their cross section was observed by SEM analyses in order to evaluate the PPY distribution into the sample. As visible in Figure 4d,e, PPY forms spherical island in all the thickness of the sample showing a complete interpenetration of the two polymers.

TGA analyses were performed in order to determine the influence of PPY created in the interfacial polymerization step on thermal stability of the object and characterize the amount of PPY deposited (Figure 5). The acidic dopants of the PPY are volatilized upon heating,[48] which explains why the PEGDA/ PPY hydrogels show lower thermal stability than that of PEGDA ones below 350 °C. As expected, PEGDA/PPY hydrogel left more residue at 700 °C in comparison to the sample without PPY (Figure 5). Moreover this residue increases as the increasing of water in the initial formulation. This could be related to the fact that the more the structure is swellable, the more free space is available for PPY infiltration. This relationship between cross-linking density and the amount of water in the initial formulation could also be observed in the decrease of T_{σ} measured by DSC. In fact, the more the hydrogel is swelled during its formation (which means the higher its water con-tent), the lower is the cross-linking density.[49] DSC measure-ments were also performed on treated PEGDA/PPY samples.



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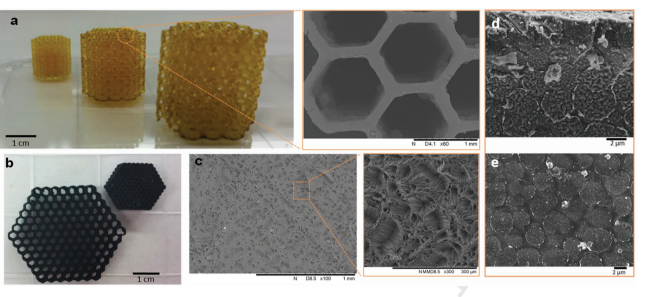


 Figure 4. a) Example of a honeycomb structure printed in different dimension (PEGDA_40) and SEM detail of the features obtained. b) 3D printed
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 PEGDA-PPY honeycomb structures. c) SEM images of the pyrrole surface. d,e) Cross section of a thick sample (4 mm).
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The presence of a second polymeric phase hinders PEGDA chain mobility resulting in the increase of T_{g} .

XPS measurements have been performed on sample PEGDA 30_PPY surface in order to obtain information regarding the chemical composition of its first layers (≤ 10 nm). In **Figure 6**a we report the survey spectrum of the sample: we detected C (59.4 at%), O (26.0 at%), N (5.7 at%), and some impurities due to Cl (0.5 at%) and Si (8.5 at%). We then performed high resolution analysis on the C1s peak (as reported in Figure 6b).

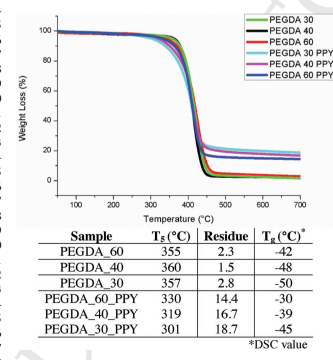


Figure 5. TGA plots and thermal properties of the compositions. $*T_5$ represents the temperature at which the 5% in weight is lost.

We attributed four components to the raw signal in order to fit properly the experimental curve. The first component at 284.5 eV has been attributed to C–C/H or α -carbons bonds,^[50] the second at 286.1 eV to α -carbons, the third one at 287.4 eV 27 to α -carbons adjacent to positively charged N atoms (C α *), and 28 the last one at 288.3 eV at carbon atoms bonded with oxygen species (-C=0 or -COO).^[51] The same procedure has also been applied to the N1s peak (Figure 6c), obtaining four com-ponents: imine -N = (398.2 eV), amine -NH- (399.4 eV), protonated amine -NH2+- (400.1 eV), and protonated imine = NH- (401.5 eV).^[52]

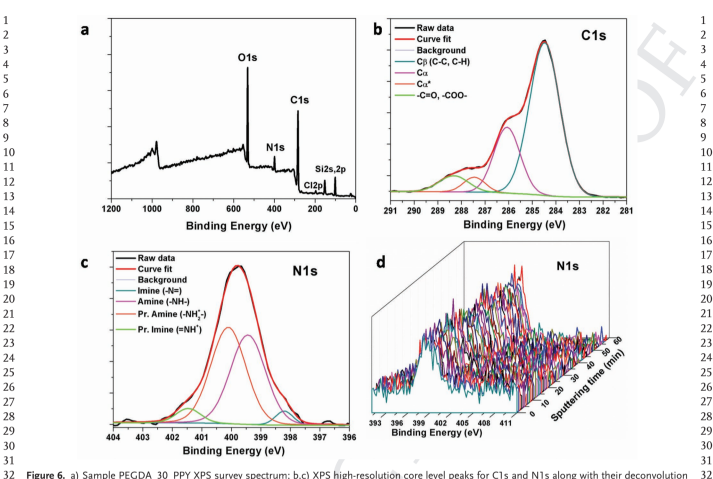
A further analysis has been accomplished to observe the homogeneity of PPy inclusion in the PEGDA matrix, at least in the first layers. To obtain this information we performed a depth profile measurements, using Ar+ as sputtering source. We alternated sputtering cycle (1 min) with HR measurements 39 in the N1s region. We repeated the procedure for 60 cycles 40 (60 min) obtaining the profile reported in Figure 6d. As can be 41 clearly seen, the intensity of the nitrogen signal is almost con- 42 stant during all the measurement. A rough estimation of the 43 polymeric composite depth sputtered can be done according 44 to the literature,^[53] by considering a sputter rate of the order 45 of 10 nm min⁻¹ and then obtaining a 600 nm depth. Along this sputtering depth PPY distribution as expected from SEM observations.

Electrical measurements were performed on films in order to evaluate the influence of PPY on the resistivity. As reported in Figure 7 the presence of PPY decreases the films' resistivity of one order of magnitude or more. This indicates that a con-tinuous PPY network was successfully formed with interfacial polymerization. The resistivity values obtained are in good agreement with values reported in the literature of similar sys-tems.^[54] It is also important to underline that all the new for-mulations showed a resistivity considerably lower than PEGDA 57 photocured without water.^[15] Moreover, the more the water in 58 the initial formulation, the lower the resistivity measured. In

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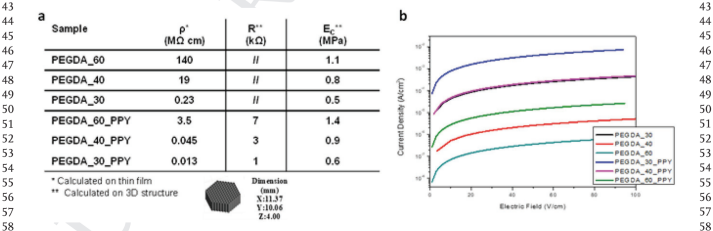


32 Figure 6. a) Sample PEGDA_30_PPY XPS survey spectrum; b,c) XPS high-resolution core level peaks for C1s and N1s along with their deconvolution procedure results; d) XPS depth profile curves for N1s peak (Ar+ source at 2 kV, alternate sputtering with 1 min cycle each). 33 34

35 order to clarify this point, electrical measurements were per-36 formed on the same neat PEGDA samples after drying in a 37 vacuum oven overnight. All the samples showed higher resis-38 tivity (2 M Ω cm), which is consistent with the literature. This let us believe that some water/humidity could remain trapped in 39 the polymeric network during material synthesis, thus reducing 40 resistivity. However, this assumption seems to us not sufficient 41

35 to explain the low resistivity values measured for PEGDA_30 sample and more investigations are ongoing to deepen this 36 point. The same vacuum treatment was performed on the sam-38 ples with PPY, which did not show significant change in the electrical behavior. 39

In order to evaluate the mechanical properties of the samples, 40 honeycomb structures were printed and tested in compression 41



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Figure 7. a) Table reporting electrical resistivity and electrical resistance and Young's modulus; b) semilogarithmic plot of I/V measurements and 59 59 resistance evaluation for the different samples.

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tests. The table in Figure 7a reports the Young's modulus for each formulation under compression. The measurements per-3 formed on the structures showed that the Young's modulus is higher in the presence of PPY compared to PEGDA sample without PPY, which can explain that the presence of a second phase hinders the PEGDA chain mobility as already seen in the DSC. Finally, the resistance of 3D printed conductive hydrogel structures was also measured, confirming the trend observed on films.

12 4. Conclusions 13

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14 In this paper, we reported a novel approach for the fabrica-15 tion of electrically conductive, mechanically tough 3D printed 16 hydrogels by coupling interfacial polymerization of PPY with 17 3D printing technology. It was demonstrated that precise 3D structures of PEG-based hydrogels can be realized even with a 18 19 considerable high amount of water in the initial formulation. The honeycomb structures were successfully infiltrated with 20 21 PPY by interfacial polymerization; ATR, XPS, and SEM results 22 indicate the existence of PPY within the matrix. Moreover the 23 dye used for enhancing 3D printing precision was exploited for PPY doping. The developed structures were electrically conduc-24 25 tive and mechanically tough. This strategy could be applied for 26 developing a new bioelectronics interface for several biomedical 27 applications. 28

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37 **Conflict of Interest** 38

39 The authors declare no conflict of interest. 40

42 **Keywords** 43

3D printing, interfacial polymerization, PEGDA, photopolymerization

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- [1] G. Justin, A. Guiseppi-Elie, Biomacromolecules 2009, 10, 2539.
- 52 [2] C. J. Small, C. O. Too, G. G. Wallace, Polym. Gels Networks 1997, 5, 53 251. 54
- [3] Y.-x. Zhao, K.-f. Ren, Y.-x. Sun, Z.-j. Li, J. Ji, RSC Adv. 2014, 4, 24511. 55
- [4] A. Guiseppi-Elie, Biomaterials 2010, 31, 2701. 56
- [5] Y. Xiao, L. He, J. Che, J. Mater. Chem. 2012, 22, 8076. 57
- [6] J. Ji, et al., J. Macromol. Sci., Part B: Phys. 2015, 54, 1122. 58
- [7] H. P. de Oliveira, S. A. Sydlik, T. M. Swager, J. Phys. Chem. C 2013, 59 117, 10270.

- [8] Y. Wu, et al., J. Mater. Chem. B 2015, 3, 5352.
- [9] S. Naficy, J. M. Razal, G. M. Spinks, G. G. Wallace, P. G. Whitten, 2 Chem. Mater. 2012, 24, 3425 3
- [10] H. Huang, J. Wu, X. Lin, L. Li, S. Shang, M. C. Yuen, G. Yan, Carbo-4 hydr. Polym. 2013, 95, 72. 5
- [11] C. Vallejo-Giraldo, A. Kelly, M. J. P. Biggs, Drug Discovery Today 2014, 19, 88.
- [12] Y. Zhao, B. Liu, L. Pan, G. Yu, Energy Environ. Sci. 2013, 6, 2856.
- [13] J. Stejskal, Chem. Papers, 1.
- 9 [14] Y. L. Kong, I. A. Tamargo, H. Kim, B. N. Johnson, M. K. Gupta, 10 T.-W. Koh, H.-A. Chin, D. A. Steingart, B. P. Rand, M. C. McAlpine, Nano Lett. 2014, 14, 7017. 11
- [15] J. N. Hanson Shepherd, S. T. Parker, R. F. Shepherd, M. U. Gillette, 12 J. A. Lewis, R. G. Nuzzo, Adv. Funct. Mater. 2011, 21, 47. 13
- [16] R. Suntornnond, J. An, C. K. Chua, Macromol. Mater. Eng. 2017, 14 302. 1600266. 15
- [17] M. Zhou, B. H. Lee, L. P. Tan, Int. J. Bioprint. 2017, 3, 2017.
- 16 [18] J. C. McDonald, M. L. Chabinyc, S. J. Metallo, J. R. Anderson, 17 A. D. Stroock, G. M. Whitesides, Anal. Chem. 2002, 74, 1537.
- 18 [19] B. C. Gross, J. L. Erkal, S. Y. Lockwood, C. Chen, D. M. Spence, 19 Anal. Chem. 2014, 86, 3240.
- 20 [20] J. Z. Manapat, Q. Chen, P. Ye, R. C. Advincula, Macromol. Mater. Eng. 2017, 302, 1600553. 21
- [21] X. Wang, Q. Guo, X. Cai, S. Zhou, B. Kobe, J. Yang, ACS Appl. Mater. 22 Interfaces 2014, 6, 2583. 23
- [22] E. Fantino, A. Chiappone, F. Calignano, M. Fontana, F. Pirri, 24 I. Roppolo, Materials 2016, 9, 589. 25
- [23] A. Chiappone, E. Fantino, I. Roppolo, M. Lorusso, D. Manfredi, 26 P. Fino, C. F. Pirri, F. Calignano, ACS Appl. Mater. Interfaces 2016, 27 8.5627.
- 28 [24] E. Fantino, A. Chiappone, I. Roppolo, D. Manfredi, R. Bongiovanni, 29 C. F. Pirri, F. Calignano, Adv. Mater. 2016, 28, 3712.
- 30 [25] S. Stassi, E. Fantino, R. Calmo, A. Chiappone, M. Gillono, D. Scaiola, C. F. Pirri, C. Ricciardi, A. Chiadò, I. Roppolo, ACS Appl. 31 Mater. Interfaces 2017, 9, 19193. 32
- [26] J. Malda, J. Visser, F. P. Melchels, T. Jüngst, W. E. Hennink, 33 W. J. Dhert, J. Groll, D. W. Hutmacher, Adv. Mater. 2013, 34 25, 5011. 35
- [27] J.-Y. Sun, X. Zhao, W. R. Illeperuma, O. Chaudhuri, K. H. Oh, 36 D. J. Mooney, J. J. Vlassak, Z. Suo, Nature 2012, 489, 133. 37
- [28] N. B. Graham, in Poly(Ethylene Glycol) Chemistry: Biotechnical and 38 Biomedical Applications (Ed: J. M. Harris), Springer, Boston, MA 39 1992, pp. 263-281.
- [29] J. M. Harris, Poly (Ethylene Glycol) Chemistry: Biotechnical and Bio-40 41 medical Applications, Springer Science & Business Media, Berlin, Germany 2013. 42
- [30] K. Arcaute, L. Ochoa, F. Medina, C. Elkins, B. Mann, R. Wicker, 43 MRS Online Proceedings Library 2005, 874. 44
- [31] V. Chan, J. H. Jeong, P. Bajaj, M. Collens, T. Saif, H. Kong, R. Bashir, 45 Lab Chip 2012, 12, 88. 46
- [32] V. Chan, P. Zorlutuna, J. H. Jeong, H. Kong, R. Bashir, Lab Chip 47 2010, 10, 2062.
- 48 [33] A. Chiappone, I. Roppolo, E. Naretto, E. Fantino, F. Calignano, 49 M. Sangermano, F. Pirri, Composites, Part B: Eng. 2017, 124, 9.
- 50 [34] X. Wang, M. Jiang, Z. Zhou, J. Gou, D. Hui, Composites, Part B: Eng. 2017, 110, 442. 51 52
- [35] N. Nuraje, K. Su, ACS Nano 2008, 2, 502.
- [36] G. Qi, Z. Wu, H. Wang, J. Mater. Chem. C 2013, 1, 7102.
- [37] J. Bhadra, D. Sarkar, Indian J. Phys. 2010, 84, 1321.
- [38] D. Zhang, F. Di, Y. Zhu, Y. Xiao, J. Che, J. Bioact. Compat. Polym.: 55 Biomed. Appl. 2015, 30, 600. 56
- [39] N. V. Blinova, M. Trchová, J. Stejskal, Eur. Polym. J. 2009, 57 45, 668.
- 58 [40] M. Karbarz, M. Gniadek, M. Donten, Z. Stojek, Electrochem. 59 Commun. 2011, 13, 714.

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- [41] X. Tuo, B. R. Li, C. L. Chen, Z. L. Huang, H. B. Huang, L. Li, Synth. Met. 2016, 213, 73.
- [42] S. Ying, W. Zheng, B. Li, X. She, H. Huang, L. Li, Z. Huang, Y. Huang, Z. Liu, X. Yu, Synth. Met. 2016, 218, 50.
- [43] N. Saengchairat, T. Tran, C.-K. Chua, Virtual Phys. Prototyping 2017, 12, 31.
- [44] J. Feng, W. Yan, L. Zhang, Microchim. Acta 2009, 166, 261.
- [45] X. Yang, Z. Zhu, T. Dai, Y. Lu, Macromol. Rapid Commun. 2005, 26, 1736.
- [46] H. Tai, W. Wang, T. Vermonden, F. Heath, W. E. Hennink, C. Alexander, K. M. Shakesheff, S. M. Howdle, Biomacromolecules 2009, 10, 822.
- [47] A. A. Jatratkar, J. B. Yadav, R. Deshmukh, H. C. Barshilia, V. Puri, J. Phys. Chem. Solids 2015, 80, 78.

- [48] Y. Liao, T. P. Farrell, G. R. Guillen, M. Li, J. A. T. Temple, X.-G. Li, E. M. V. Hoek, R. B. Kaner, Mater. Horiz. 2014, 1, 58.
- [49] L. M. Weber, C. G. Lopez, K. S. Anseth, J. Biomed. Mater. Res., Part A 2009, 90, 720.
- [50] P. M. Carrasco, M. Cortazar, E. Ochoteco, E. Calahorra, J. A. Pomposo, Surf. Interface Anal. 2007, 39, 26.
- [51] N. Garino, A. Sacco, M. Castellino, J. A. Muñoz-Tabares, A. Chiodoni, V. Agostino, V. Margaria, M. Gerosa, G. Massaglia, M. Quaglio, ACS Appl. Mater. Interfaces 2016, 8, 4633.
- [52] S. Golczak, A. Kanciurzewska, M. Fahlman, K. Langer, J. J. Langer, Solid State Ionics 2008, 179, 2234.
- [53] T. Nobuta, T. Ogawa, J. Mater. Sci. 2009, 44, 1800.
- [54] G. Kaur, R. Adhikari, P. Cass, M. Bown, P. Gunatillake, RSC Adv. 2015, 5, 37553.

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