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Rapid prototyping of 3D Organic Electrochemical Transistors by composite photocurable resin

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ABSTRACT

Rapid Prototyping (RP) promises to induce a revolutionary impact on how the objects can be produced and used in industrial manufacturing as well as in everyday life. Over the time a standard technique as the 3D Stereolithography (SL) has become a fundamental technology for RP and Additive Manufacturing (AM), since it enables the fabrication of the 3D objects from a cost-effective photocurable resin. Efforts to obtain devices more complex than just a mere aesthetic simulacre, have been spent with uncertain results. The multidisciplinary nature of such manufacturing technique furtherly hinders the route to the fabrication of complex devices. A good knowledge of the bases of material science and engineering is required to deal with SL technological, characterization and testing aspects. In this framework, our study aims to reveal a new approach to obtain RP of complex devices, namely Organic Electro-Chemical Transistors (OECTs), by SL technique exploiting a resin composite based on the conductive poly(3,4-ethylenedioxythiophene):polystyrene sulfonate (PEDOT:PSS) and the photo curable Poly(ethylene glycol) diacrylate (PEGDA). A comprehensive study is presented, starting from the optimization of composite resin and characterization of its electrochemical properties, up to the 3D OECTs printing and testing. Relevant performances in biosensing for dopamine (DA) detection using the 3D OECTs are reported and discussed too.

Introduction

3D Stereolithography (SL), belonging to the larger 3D Printing or Additive Manufacturing (AM) family, has delivered a new conceptualization of object design and fabrication with beneficial implications also for Organic Electronics ^{1–3}. When a new technology frontier approaches, researchers are stimulated to push their effort to overcome its limitations. The translation of this concept for SL is the development of new architectures and materials able to add functionalities, and hence implementing and validating new smart objects ⁴ and devices ⁵ with enhanced performances ⁶ and unconventional properties ⁷. Three-dimensional expansion of devices brings new solutions and capabilities integration, providing improvements in many fields of application: sensors for wearable electronics ⁸, energy storage in building ⁹ and biomedical devices for medicine ¹⁰. In the latter case, a class of organic materials showing peculiar properties of biocompatibility and conformability, along with a mixed ionic-electronic charge transport, was developed. Emerging applications are focused on neuromorphic devices ¹⁰, biosensors ^{11–13} and cell monitoring ¹⁴. Among these devices, Organic Electro-Chemical Transistors (OECTs) show multifunctional operation in so far as they allow implementing both a transistors-like and a memristive-like response ^{15,16}.

Recently, a new composite with electrical conducting properties has been proposed by Scordo et al. ¹⁷. This is a SL resin based on poly(3,4-ethylenedioxythiophene):polystyrene sulfonate (PEDOT:PSS), which provides the conducting properties, and Poly(ethylene glycol) diacrylate (PEGDA), acting as is the photo curable matrix in the composite. Of course, the material alone represents only the first milestone for a device development, since its physical properties should be accurately studied and the processing protocols need to be engineered with the aim of obtaining a significant throughput. The passage from the material synthesis to a practical exploitation often requires some decades. In this respect, an exhaustive example is given by carbon nanotubes and/or graphene that, although at the basis of a large set of device prototypes, still lack of an actual killer application to go out from laboratories. Indeed, the Rapid Prototyping (RP) approach grants the extraordinary opportunity to speed up the passage from the materials' development to their

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use in well performing devices, ensuring at the same time a highly repeatable fabrication process. This is another aspect that authors had in mind during the design and developing of the presented experimental activity. On this basis, the present work demonstrates, for the first time, that a RP of 3D OECTs, which in this case have been demonstrated to be efficient in biosensing applications, can be effectively obtained by SL. 3D printing methods for the development of electronic devices actually provide 2D structures (e.g. ink-jet¹⁸, aerosol-jet¹⁹) or 3D structures implemented upon multistep processes²⁰ and by Fused Deposition Modeling (FDM) and micro-dispensing ^{21–23}. This work introduces a non-trivial novelty consisting in demonstrating the fabrication of multi-materials 3D objects, which are intrinsically equipped with specific physical properties and can be manufactured in-line by combining 3D SL techniques and engineered resins.

Results and discussion

PEGDA:PEDOT resin

A large number of protocols aimed at enhancing PEDOT:PSS conductivity through the combination of thermal curing and acid treatments are reported in literature ^{24–26}. H₂SO₄ 0.5 M treatment is used to induce a significant structural change on PEDOT:PSS layer, because it removes the excess of PSS counter-ion, forming crystallized nanofibrils ²⁵. A similar approach was applied in this study to obtain the high conductive agglomerates of PEDOT:PSS starting from the commercial PEDOT:PSS solution eluted with sulfuric acid (hereinafter PEDOT:PSS treated by H₂SO₄ will be only referred to as PEDOT). Since PEDOT agglomerates size can affect printing accuracy, a fractionation was carried out by dispersing them in a proper solvent. Two different dispersing solvents, ethanol and H₂SO₄ 0.5 M solution, were compared aiming to enhance the fractionation of the PEDOT agglomerates. It was found that the solvent affected also the conductivity of the resin. The polymerized PEGDA:PEDOT resin fractionated using ethanol shows a mean conductivity value of 5·10⁻² S/cm compared to 1·10⁻³ S/cm of the one fractionated using H₂SO₄. Moreover, an optimal dispersion of PEDOT agglomerates was achieved in ethanol because its low density helps to better suspend them in comparison to acid solution. This optimized treatment allows to encapsulate the PEDOT in a PEGDA 575 matrix following the method previously reported¹⁷.

The electrochemical behavior of the resin was investigated by Cyclic Voltammetry (CV), carried out using a well-known electrochemically reversible redox system, i.e. the potassium ferricyanide, in PBS (Phosphate-buffered saline) solution. Voltammograms in the case of SPEs (Screen Printed Electrodes) with Working Electrode, (WEs) bare and modified (upon composite resin drop casting) are reported in Figure 1a and Figure 1b, respectively. The curve for PBS is also reported, showing a behavior similar to that shown elsewhere ²⁷.

Figure 1b shows that the electroactive properties of the composite resin are clearly revealed by the amplification of the electrochemical signals acquired in the case of modified WEs. Again, the curve profile is similar to that found elsewhere in the case of potassium ferricyanide in PBS solutions ^{28,29}.

The enhanced performance for modified SPEs can be rationalized by recurring to the Randles-Sevcik equation:

$$I_p = (2.69 \times 10^5) A D^{\frac{1}{2}} C v^{\frac{1}{2}}$$
 (1),

where A is the electrode surface area (cm²), D is the diffusion coefficient (cm²/s), C is the concentration of the electroactive species in the bulk solution (mol/cm³) and v is the scan rate.

The Randles-Sevick equation describes the effect of ν on the peak current (I_p) for a reversible process and predicts a linear dependence of the anodic (I_{pa}) and cathodic (I_{pc}) peak currents with respect to the square root of the scan rate. The linear trends of $I_p \, \nu s. \, \nu'^2$ in Figure 1d indicate a diffusive electrochemical process (i.e. a process controlled by the mass transport velocity of the electroactive species moving towards the WE surface due to a concentration gradient), even if a mild separation of the peak potentials upon increasing the scan rate in the case of modified SPEs (black arrow in figure 1b) indicates a slightly lower electron transfer kinetics than that of bare Au WEs. Nevertheless, while a reversible behavior is found for the bare SPE, where the anodic-to-cathodic ratio (I_{pc}/I_{pa}) is almost 1 (Figure 1c), the composite resin shows a quasi-reversible behavior. In fact, as predicted by equation (1) a linear trend is still obtained, but the composite resin shows an I_{pc}/I_{pa} varying from 0.97 (at 25 mV/s) to 0.84 (at 300 mV/s). Values significantly different from 1 are typically associated with chemical reactions coupled with the electron transfer due to the electroactivity of coverages (in our case, this role is ascribable to the PEDOT counterpart).

Taking into account that the surface of the WE is $0.126~\text{cm}^2$ and that the $K_3[Fe(CN)_6]$ concentration is 2.5×10^{-5} mol/cm³, a linear fit to the I_{pa} vs. $v\frac{1}{2}$ curve in Figure 1c allows extracting the diffusion coefficient for potassium ferricyanide in PBS from equation (1). In this case $D=1.7\times10^{-6}~\text{cm}^2/\text{s}$ (for instance, the diffusion coefficient of potassium ferricyanide calculated by more sophisticated electrochemical time of flight measurements³0 is $7.3\pm0.7\times10^{-6}~\text{cm}^2/\text{s}$). Using the extracted diffusion coefficient value, an effective surface area of $0.73~\text{cm}^2$ is assessed for SPEs covered by PEGDA:PEDOT.

This outcome (i.e. active WE surface upon modification enhanced by a nearly six-fold ratio with respect to that of the bare SPE) demonstrates that the PEGDA:PEDOT resin induces a strongly enhanced electrochemical performance. The hypothesis is that the composite coverage, on one hand enhances the density of redox sites due to the presence of the electroactive component (PEDOT) while, on the other hand, to some extent PEGDA takes the role of PSS in

PEDOT:PSS, i.e. it favors the diffusion of ionic species through the PEDOT bulk. This outcome points out the composite resin as ideally suitable for OECTs applications.

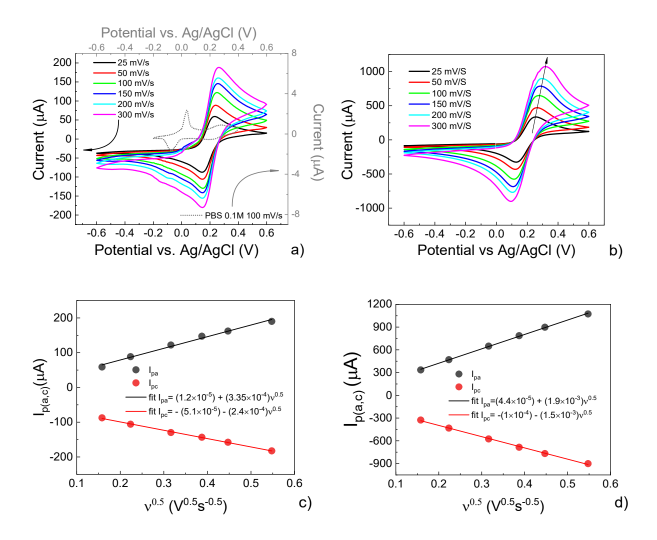


Figure 1: Cyclic voltammetry of Potassium Ferricyanide (K₃[Fe(CN)₆]), 25 mM, in PBS (pH 7.4), 100 mM (scan rates 25, 50, 100, 150, 200 and 300 mV/s) carried out using (a) a SPE equipped with Au working and counter electrodes and Ag/AgCl reference electrode; (b) the same measurement as in (a) using a homologue SPE where the working electrode is covered by a PEGDA:PEDOT layer; (c) anodic and cathodic peak positions extracted from (a) as a function of the square root of the scan rates (related fitting curves, solid lines); (d) anodic and cathodic peak positions extracted from (b) as a function of the square root of the scan rates (related fitting curves, solid lines).

Rapid prototyping of 3D OECTs

The PEGDA:PEDOT resin was optimized to work with a customized SL printer (Microla Optoelectronics s.r.l). The printer works in a top-down configuration and its accuracy is about $80 \mu m$ on the XY plane (depending on the resin optical properties) and $50 \mu m$ along the vertical axis.

The 3D printing of the OECT with the PEDOT-filled resin started from a simple geometry to assess the properties of the resin and its suitability for printing a sensing element. To this end, a 500 μ m thick, 2 mm long straight channel was designed. At the channel extremities, two squared contact pads were printed using the same material as the channel ³¹. With a view to miniaturized 3D printed sensors, the present OECT was firstly designed to have the smallest channel width achievable with the reported setup and material. Therefore, taking into account the size of PEDOT agglomerates inside PEGDA:PEDOT resin and the printer lateral resolution (around 80 μ m), the minimum channel width was set to be 300 μ m in order to ensure both accuracy and mechanical stability. A wider channel was also printed (700 μ m wide, 2 mm long) to investigate the transistor response at different channel widths. An insulating frame was then added to ensure always the same channel length (2 mm) to be involved in the doping/de-doping process, also ensuring the OECTs response repeatability. Two wells were built, on an additional insulating support, to contain a maximum electrolyte volume of 175 μ l. As a result, the 3D printed OECT includes four main parts: an insulating base, the electrically conductive parts, an insulant support for the frame and wells (as reported in Figure 2). The insulating parts

were printed with the commercial SpotHT resin while the conductive parts with the PEGDA:PEDOT resin. The insulating base, the transistor part and the insulating support were $500 \, \mu m$ thick, while the wells walls were $8 \, mm$ high.

Top view A Wells B Insulating frame C Insulating support D PEGDA:PEDOT E Insulating base

Figure 2 CAD drawing of the OECT to be printed.

Once the device was printed, it was exposed to a thermal curing to post-process the 3D printed part, as previously reported ³². In addition, this thermal treatment turned out to enhance the 3D OECTs performance with a similar effect observed for the annealing of PEDOT:PSS thin films ^{33,34}. The thermal treatment was performed by means of a slow heating process followed by a likewise cooling process to gradually release internal stresses and thus avoid undesired fractures.

The printability of the PEGDA:PEDOT was clearly demonstrated by RP of 3D OECTs with the two different channel widths of 300 µm and 700 µm, respectively (Figure 3a). The electrical characterization of the final device indicated that the major constrain related to prototyping was not due to the SL intrinsic limitation, but rather to the reproducibility of the OECT response. In fact, the characterization of the 3D printed devices highlighted that the 300 µm wide channel had not a reproducible response like the one by the 700 µm wide channel (Figure 3b). Therefore, the latter geometry was selected as the best performing one in terms of reproducibility and, hence, considered for the following characterizations. This could be ascribed to the need of a sufficient concentration of active material in the channel in order to properly work as OECT. Nevertheless, the increasing of the PEDOT content in the resin slightly reduces the printability. Hence, to get reproducible and well performing devices the best compromise was achieved using the PEGDA:PEDOT (55:45 wt%) formulation and the 700 µm wide channel.

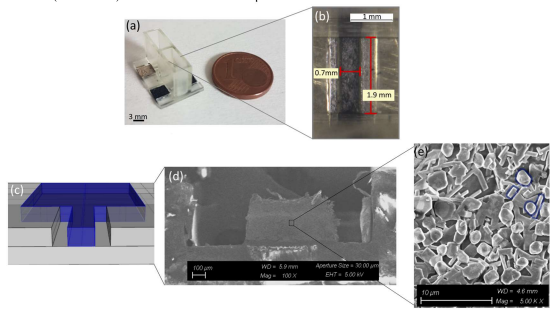


Figure 3: 3D printed OECT. (a) Device after printing; (b) microscope image of the 700 μm wide channel; (c) section view of the 700 μm wide channel geometry; (d) FESEM image of the same section view reported in (c); (e) FESEM image of the PEGDA:PEDOT polymerized resin in which the PEDOT particles (some are highlighted in blue) are clearly visible.

A section of the OECT channel (as sketched in Figure 3c) was observed by Field Emission Scanning Electron Microscopy (FESEM) to assess that the insulating SpotHT resin didn't accidentally cover the active PEGDA:PEDOT

resin, thus passivating it. Indeed, a clear separation between the two materials was observed, as reported in Figure 3d. Finally, the FESEM imaging also revealed small particles (in the order of a few microns) inside the OECT channel, ascribable to the PEDOT:PSS particles (Figure 3e).

3D OECTs characterization

In order to perform the electrical characterization, the fluidic chamber of the OECT was filled with 150 µL of NaCl 100 mM, while an Ag/AgCl pellet electrode was placed in the electrolyte and exploited as gate electrode. Transfer characteristics (I_{ds} vs V_{gs}) were measured at different V_{ds} by sweeping V_{gs} in forward (from -750 mV to +1 V) and backward (from +1V to -750 mV) mode (Figure 4 a,b). The drain current decreases upon increasing the gate voltage, as expected for a p-type OECT working in depletion mode³⁵. A strong hysteresis is present in the transfer curves due to the diffusion of electrolyte ions inside the PEGDA:PEDOT resin. The I_{on}/I_{off} ratio was calculated as the ratio between the current values recorded at V_{gs} = -750 mV (I_{on}) and V_{gs} = +1 V (I_{off}), respectively, for four different devices. As a result, I_{on}/I_{off} was found to increase from (1.21 \pm 0.40) \cdot 10³ at V_{ds} = -100 mV to (2.79 \pm 1.6) \cdot 10³ at V_{ds} = -600 mV. The gate current has been continuously monitored throughout the measurements (dashed lines in Figure 1b) and it was typically two orders of magnitude lower than the drain current in the ON state, with an average value around 10 µA. Figure 4 c shows the differential transconductance $g_m = \frac{\partial I_{ds}}{\partial V_{gs}}$ as a function of V_{gs} . The maximum of g_m linearly increases from 550 μ S at $V_{ds} = -100$ mV to 2.5 mS at $V_{ds} = -600$ mV where it saturates, while the peak position moves towards negative V_{gs} , as expected from the high volume capacitance due to the thick active layer of the device ^{34,36}. As reported in Table 1, it is expected that maximum transconductance can vary a lot depending on the 3D printing fabrication process, therefore the found values are comparable or even higher with respect to others reported in literature 19,20, while higher values of g_{max} are achieved for lower thickness 21,37. In our case, the thickness of the channel is higher than planar one but, the assessed low transconductance value is clearly due to the properties of the resin composite that works on the basis of the PEDOT particles interconnections inside the insulating PEGDA matrix. In addition, high transconductances for OECTs based on PEDOT:PSS have been attributed to the high volumetric capacitance showed by this diphasic organic conductor³⁸. Indeed, the volumetric capacitance in the case of PEDOT:PSS has been attributed to the presence of sulfonate anion-hole pairs, acting as capacitive elements, at the interface between anionic PSS shells and hole-rich PEDOT clusters³⁹. In the case of the resin composite, the PSS phase has been replaced by the PEGDA one and as a consequence capacitive sites with a lower intrinsic capacitance can be attributed to sites constituted by holes (PEDOT) and non-polar groups (PEGDA) at the PEDOT:PEGDA interface."

Table 1. Reported main parameters for 3D printed OECT in literatures: channel dimensions, g_{max} , I_{ON}/I_{OFF} . ND indicates not declared parameter.

Fabrication Techniques	Channel dimension W/L/t (µm)	Materials	g _{max} (mS)	I _{ON} /I _{OFF}	Ref.
Stereolithopgraphy	700 / 1900 / 500	PEGDA:PEDOT	2.5	2.79±1.6 ·10 ³	This work
Aerosol-jet printing	200 / 200 / 0.2	PEDOT:PSS	0.52	ND	19
Laser sintering	1000 / 15000 / 1000	PEDOT:PSS	2	2	20
FDM / direct writing 3D	1000 / 1600 / 7.1	PEDOT:PSS	31.8	1.33·10³	21
Syringe deposition	3000 / 3000 / 5	PEDOT:PSS	ND	2	23

Fabrication Techniques	Channel dimension W/L/t (µm)	Materials	g _{max} (mS)	I _{ON} /I _{OFF}	Ref.
Melt extrusion / Electrospining	200/1500/200	PEDOT:Nafion	30-40	100	37

The output curves (I_{ds} vs V_{ds}) of the device, measured by varying the drain voltage between 0 V and -600 mV, at a scan rate of 200 mV/s and at fixed gate voltage between -1 V and +1 V (Figure 4d), showed the typical response of a transistor working in linear regime.

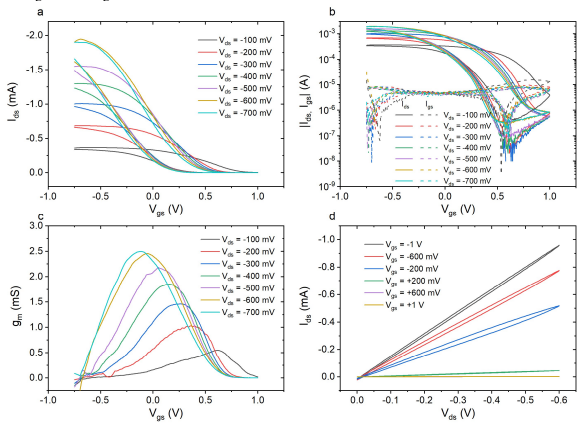


Figure 4 a) Transfer curve measured at different V_{ds} ; b) logarithmic plot of transfer curve (solid lines) and gate leakage current (dashed lines); c) transconductance curve obtained for V_{gs} scanned from -0.75 V to +1 V; d) output characteristics of the device.

The transient behavior of the device has been investigated by holes time of flight (TOF) measurements and by constant gate voltage pulses.

Figure 5a shows the I_{ds} response while driving the device with constant gate current pulses of 15 s at fixed $V_{ds} = -100$ mV. The linear fit of $\frac{dI_{ds}}{dt}$ versus I_{gs} ($r^2 = 0.99$) shown in the inset yielded a hole time of flight $\tau_h = (6.03 \pm 0.19)$ s

and an effective hole mobility $\mu=(6.6\pm0.2)\cdot10^{-2}\frac{cm^2}{V\cdot s}$. The slow recovery of I_{ds} after removal of I_{gs} is associated to a slow back-diffusion of ions from the PEGDA:PEDOT resin in the electrolyte.

In order to extract the volumetric capacitance of the active channel material, the gate current has been measured while applying gate voltage pulses at fixed $V_{ds} = 0$ V (Figure 5b). The inset of Figure 5b shows a linear fit ($r^2 = 0.98$) of the charge accumulated in the channel versus the applied gate voltage, yielding a volumetric capacitance $C^* = (1.28 \pm 0.09) \frac{F}{cm^3}$. This value is one order of magnitude lower than the one reported in literature for PEDOT:PSS ⁴⁰, as expected from the substitution of the anionic PSS sites with non-polar PEGDA sites.

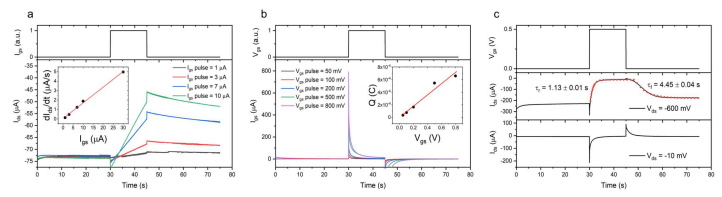


Figure 5: a) Time of flight measurement: device driven by a constant gate current with application of a 15 s pulse with $V_{ds} = -100$ mV. The inset shows the linear fit of $\frac{dI_{ds}}{dt}$ vs I_{gs} ($r^2 = 0.99$) yielding the hole mobility $\mu = (6.6 \pm 0.2) \cdot 10^{-2} \frac{cm^2}{V \cdot s}$. b) Gate current response to a constant gate voltage pulse of 15 s with $V_{ds} = 0$ V. The charge Q accumulated in the channel has been obtained by integration of I_{gs} curves, the linear fit of Q vs V_{gs} ($r^2 = 0.98$) yielded a capacitance $C = (890 \pm 60) \, \mu F$. c) I_{ds} response to a constant V_{gs} pulse of 15 s at $V_{ds} = -600$ mV (central panel) and $V_{ds} = -10$ mV (lower panel). The red dotted line in the central panel is an exponential fit of the switching behavior of the device, yielding a rise time $\tau_r = (1.13 \pm 0.01) \, s$ and a fall time $\tau_f = (4.45 \pm 0.04) \, s$.

Figure 5c shows the I_{ds} response to a V_{gs} pulse of 0.5 V at V_{ds} = -600 mV (central panel) and V_{ds} = -10 mV (lower panel). From Bernards model³⁵ the transient drain current can be modeled as:

$$I_{ds}(t) = I_{SS}(V_{gs}) + \Delta I_{SS} \left[1 - f \frac{\tau_h}{\tau_i} \right] \exp\left(-\frac{t}{\tau_i} \right)$$
 (2)

Where I_{SS} is the steady state drain current, ΔI_{SS} is the difference between initial and final steady state current, f takes in account the spatial non-uniformity in the de-doping process, τ_h is the hole time of flight and τ_i is the ionic RC time constant

At low drain voltage (lower panel) a spike and recovery behavior is obtained as expected from Bernards model³⁵ for slow hole transport. At high drain voltage the response is dominated by a monotonic decay of the drain current, the exponential fit yielding a rise time $\tau_r = (1.13 \pm 0.01) s$ and a fall time $\tau_f = (4.45 \pm 0.04) s$. The greater fall time is consistent with the slow recovery obtained during the TOF measurements. The presence of a small spike in the drain current upon application/removal of the gate voltage pulse even at high V_{ds} is attributed to the large source/drain spacing of 2 mm, from the ratio of this spike in drain current versus the corresponding spike in gate current a non-uniformity factor $f \simeq 0.26$ has been calculated⁴¹.

Dopamine biosensing

We made an analysis dedicated to the check of performances by the composite resin used in OECTs architectures. This analysis is aimed at giving a tangible example of 3D OECTs suitability in biosensing applications. We hence explored the 3D OECTs efficiency in sensing mode of operation detecting biomolecules of interest in medicine. In the specific case, we implemented the detection, by the 3D OECT, of the dopamine (DA), which is a neurotransmitter regulating the correct functioning of several organs whose loss in some areas of the brain causes, for instance, the Parkinson's disease. The choice of dopamine is also motivated by the fact that, starting from the work by Tang et al. ⁴², it is a prototyping molecule in OECTs biosensing ^{43–46}.

The calibration curve for DA detection implemented using a PEGDA:PEDOT OECT and calculated from transfer curves is reported in Figure 6a. Some selected (averaged) transfer curves at lower and higher DA concentrations ([DA]= 0, 10 μ M, 100 μ M, 1 mM and 5 mM in PBS 100 mM) and the related universal curve are reported as an example in Figures 6b and 6c, respectively. Quasi-reversibility of electrochemical processes in the composite resin is indicated by the large error bars upon averaging repeated measurements, especially at lower V_{gs} (the baseline current, I_{ds} for $V_{gs} \rightarrow 0$, shifts towards lower values and the largest shift is systematically recorded after the first measurement). The basic idea is to exploit the reactivity of catecholamines with a bare Pt electrode, whereas dopamine is expected to

The basic idea is to exploit the reactivity of catecholamines with a bare Pt electrode, whereas dopamine is expected to be involved in a 2-electrons exchange mechanism with the platinum electrode 47 . The measurement protocol is defined by the scheme used by Tang et al. 42 . The proposed scheme relies on the fact that the reaction of dopamine at the platinum gate electrode is able to modulate the effective gate voltage ($V_{gs,eff}$) due to an offset voltage (V_{offset}) strictly connected to the analyte concentration and expressed as 48 :

connected to the analyte concentration and expressed as ⁴⁸:
$$V_{offset} = V_{gs,eff} - V_{gs} = \Delta V_{gs} = 2.30 (1 + \gamma) \frac{K_B T}{nq} log[DA] + cost \qquad (2)$$

where K_B is the Boltzmann's constant, T is temperature, n is the number of electrons transferred at the gate, [DA] is the dopamine concentration and γ is the ratio between the capacitances at the gate electrode/electrolyte and electrolyte/active channel interfaces. This equation predicts that an increase of dopamine concentration causes the enhancement of V_{offset} that, in turn, implies a reduction of the channel current.

The device performance can be rationalized by extracting two parameters from the calibration curve, i.e. the Limit of Detection (LoD) and the sensitivity, the former representing the lowest concentration of the analyte that can be reliably

distinguished if compared to a blank measurement. Sensitivity, indeed, describes the minimum input parameter able to create a detectable output response by the sensor. The assessment of ΔV_{gs} by evaluating the shift of transfer curves is expected to be less effective in determining lower LoDs if compared to its evaluation based on real-time measurements, such as the recording of the step-like device response upon subsequent injections of the analyte to be detected in an electrolytic reservoir ^{49,50}. Nevertheless, both devices show a LoD falling in the micromolar range and a dynamic window varying between the micromolar and the millimolar range of [DA]. Assessed LoD is two orders of magnitude higher than that experimentally achievable by analytical tools, such as Cyclic Voltammetry in the case of SPEs having a Pt working electrodes (50 nm⁴²), but is still in line with LoD values obtained using OECTs with unmodified Pt gate electrodes ^{42,45}. However, the OECT operation consists of an electronic transduction (by the active channel) of ionic signals associated to some reactions at the gate electrode/electrolyte interface (as in the case of DA and Pt electrodes) and/or capacitive effects (Electrical Double layer formation). In the former case, the efficiency of such transduction often requires a specific functionalization aimed at amplifying the charge transfer mechanisms at the metallic gate/liquid electrolyte interface, hence LoDs comparable to the lowest DA concentrations determined by CV measurements can only be achieved upon functionalizing the Pt gate electrode^{43,51}

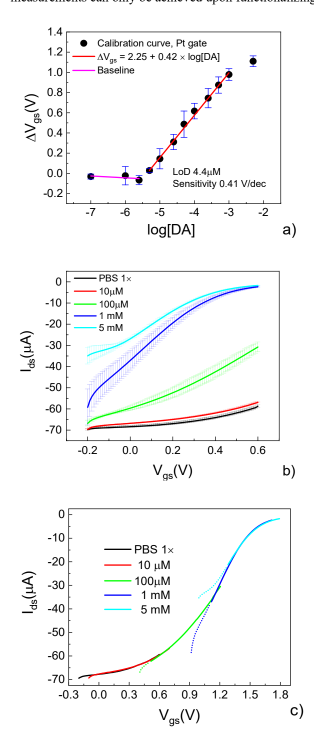


Figure 6: calibration curve for dopamine sensing by (a) a PEGDA/PEDOT 3D OECT; (b) typical transfer curves recorded at different dopamine concentrations ([DA]= 0, $10~\mu$ M, $100~\mu$ M, 1~mM and 5~mM, error bars are standard deviations) and (c) universal curve obtained by merging transfer curves in (c) upon shifting them along the V_{ss} axis;

On the contrary, calibration curves show a larger sensitivity if compared to that assessed using different experimental approaches ⁵⁰. Specifically, PEGDA:PEDOT OECTs show a markedly enhanced sensitivity of 0.41 V/dec. This enhanced sensitivity may be ascribed to the PEGDA:PEDOT microstructure. In fact, the composite resin microstructure (Figure 3e) shows a homogeneous dispersion of the PEDOT phase, whose exposure to the electrolyte is not mediated by the PSS phase. This is difficult to achieve with standard PEDOT:PSS, where both solvent ⁵² and thermal ³⁴ annealing cause the PEDOT clusters to be segregated in the bottom part of PEDOT:PSS films and the PSS phase interfaced to the gate electrolyte.

Conclusion

We demonstrated that even devices as complex as OECTs can be well fabricated by standard SL technique. RP of 3D OECTs was achieved by optimizing a new composite material: the PEGDA:PEDOT resin. The latter is a photocurable composite that we demonstrate to maintain performance comparable to that of PEDOT:PSS in terms of electrical conductance, bulk ionic diffusion mechanism, gating response in OECTs architectures and, hence, showing quite good sensing response. The optimization of this characteristics depends not only on the PEDOT content inside the resin, but also on the active material pre-treatment and the printability, which ultimately affects the OECTs behavior. Finally, 3D OECTs showed an enhanced sensitivity of 0.41 V/dec towards dopamine detection, thus demonstrating a precise assessment of concentrations when used as biosensors and, hence, in a perspective view, the freeform fabrication enabled by 3D printing paves the way for their integration in IoT and smart objects.

Methods

PEGDA:PEDOT resin

The highly conductive PEDOT is prepared starting from the commercial solution Clevios™ PH1000 (Heraeus) eluted with 0.5 M H₂SO₄ at 1:10 ratio (Clevios:0.5 M H₂SO₄) for 12 h. Then the PEDOT agglomerates were separated from the initial solution by centrifugation at at 4000 rpm for 15 min with Ohaus Frontiers Centrifuge. The agglomerates were dispersed in ethanol and a 20 minutes fractionation was performed with IKA Ultraturrax. A further separation by centrifugation at 4000 rpm for 15 min allowed to obtain the final PEDOT agglomerates, which were added to PEGDA-575:IRGACURE 819 (100:1 wt%) resin (Merk) in a ratio PEGDA:PEDOT (55:45 wt%).

CV measurements

Voltammograms (potential range -0.6 to 0.6 vs. Ag/AgCl) were acquired using commercial SPEs (C220BT Dropsens, Metroohm: Au Working Electrode, WE, surface area $0.126~\text{cm}^2$, Au Counter Electrode, CE, Ag(AgCl) Reference Electrode, RE) with and without modification of the WE by the resin composite (drop casted, $3\mu\text{L}$), using a Palmsens4 potentiostat/impedance analyzer. Curves were acquired at different scan rates (v = 25, 50, 10, 150, 200 and 300~mV/s) in the case of a potassium ferricyanide salt ($K_3[\text{Fe}(\text{CN})_6]$) in PBS solution (25 mM in PBS solution, 100 mM, pH 7.4) as electrolyte

3D printing of the device

Two resins were used to print the device: SpotHT resin from SpotA materials and PEGDA:PEDOT resin prepared as reported in the previous section. The customized SL printer was set to build up the object by printing a 100 µm thick layer each time and alternating the materials. In the top-down printing configuration, the printer platform moves inside a vat filled with liquid resin, then it comes out and a recoater blade slides on it (at a distance almost equal to the layer thickness). In such step, the resin is made more homogeneous at the polymerization site and is ready to be crosslinked by the exposure through of a UV laser (405 nm wavelength in this case) moved by a galvanometric head. For the 3D printed OECTs, the printing process was slightly modified to allow for the use of different materials. First of all, the insulating base was printed with SpotHT resin setting the following parameters: 10 mW laser nominal output power, 1000 mm·s⁻¹ laser hatch speed, 50 µm hatch spacing. Then, the printing was paused and the printer vat filled with PEGDA:PEDOT resin. The printing was resumed to print the OECT channels and pads, setting the following parameters: 20 mW laser nominal output power, 1000 mm·s⁻¹ laser hatch speed, 50 µm hatch spacing. Finally, the insulating support, the insulating frame and the wells were printed after the PEGDA:PEDOT resin was replaced with the SpotHT resin. The printed devices were carefully rinsed with isopropyl alcohol and dried. A final post-curing step was performed in a thermal chamber (Binder VDL): starting from room temperature, the devices were heated in vacuum up to 120°C for one hour. Then, they were left cooling in vacuum, until the room temperature was reached again. A Zeiss Supra 40 Field Emission Scanning Electron Microscopy (FESEM), electron high tension of 5 KV, was used to investigate the PEDOT agglomerates in the 3D OECTs.

OECT characterization

Electrical characterization has been performed using a Keysight B2912A Source Meter precision unit connected to the micromanipulators of a probe station through triax cables and controlled by the proprietary Keysight software. A solution of 100 mM NaCl in DI water (pH = 7) was used as gate electrolyte during output and transfer-characteristic measurements and during transient response measurements. An Ag/AgCl electrode (1 mm of diameter) was used as gate electrode. Transfer characteristics, i.e., the channel current (I_{ds}) as a function of the applied gate voltage applied (V_{gs}) at fixed drain-source voltage (V_{ds}), were measured at fixed V_{ds} = -100 mV, -200 mV, -300 mV, -400 mV, -500 mV, -600 mV and -700 mV, by sweeping V_{gs} from -750 mV to +1 V and back with a scan rate of 5 mV/s. Output characteristics were measured by sweeping the V_{ds} from 0 V to -600 mV with a scan rate of 200 mV/s at different fixed V_{gs} of -1 V, -600 mV, -200 mV, +200 mV, +600 mV and +1 V.

Time of flight measurements have been performed driving the device with a 15 s constant gate current pulse of 1 μ A, 3 μ A, 7 μ A, 10 μ A and 30 μ A at fixed V_{ds} = -100 mV.

In order to extract the volumetric capacitance, the gate current has been measured at fixed $V_{ds} = 0 \text{ V}$ while V_{gs} pulses of 15 s were applied at 50 mV, 100 mV, 200 mV, 500 mV and 800 mV.

The response behavior of I_{ds} to a fixed V_{gs} pulse has been investigated by applying a gate voltage pulse of +500 mV for 15 s while fixing $V_{ds} = -600$ mV and $V_{ds} = -10$ mV. All the transient response measurements have been performed with a sampling rate of 33.33 Hz.

Dopamine biosensing

The PEGDA-PEDOT OECT response was evaluated by acquiring transfer characteristics in presence of different concentrations of dopamine (MW= 189.54 mg/mol, Alfa Aeser) in 100 mM PBS solution. All the electrical measurements were performed by varying the gate voltage from -0.2 V to 1 V at fixed V_{ds} =-0.6 V, with a scan rate of 10 mV/s. The PDMS well was filled with 150 μ L of PBS/DA solutions with dopamine diluted at concentrations ranging from 100 nM to 5 mM. A vertical Pt wire gate (0.6 mm diameter, Franco Corradi) was used as the gate electrode. Each measurement was repeated three times and the channel was washed with H₂O Milli-Q and PBS solution after each measurement step. ΔV_{gs} was calculated by merging transfer curves at different concentrations (three runs per device for each DA concentration, repeated using three different devices) upon shifting them along the V_{gs} axis in order to define a universal curve ⁴⁹. Error bars have been calculated as the statistical error of the mean (σ_{dev}/N -1, where σ_{dev} are the standard deviations and N is the number of measured devices).

The device LoD against DA sensing was assessed by estimating the intersection between the slope of the linear fit in the dynamic window (sensor transfer curve) and the ideal baseline for concentrations falling below such window. Sensitivity was calculated as the slope of the sensor transfer curve.

References

- 1. Falco, A., Petrelli, M., Bezzeccheri, E., Abdelhalim, A. & Lugli, P. Towards 3D-printed organic electronics: Planarization and spray-deposition of functional layers onto 3D-printed objects. *Org. Electron.* **39**, 340–347 (2016)
- 2. Aoyama, T., Tanaka, H. & Tajima, Y. Three-dimensional (3D) concave OLEDs fabricated by a multilayer coating technique of electrospray deposition (ESD). in *Proceedings of SPIE The International Society for Optical Engineering* **10529**, (2018).
- 3. Huang, Y. *et al.* Assembly and applications of 3D conformal electronics on curvilinear surfaces. *Mater. Horizons* **6**, 642–683 (2019).
- 4. Marasso, S. L. *et al.* PLA conductive filament for 3D printed smart sensing applications. *Rapid Prototyp. J.* (2018). doi:10.1108/RPJ-09-2016-0150
- 5. Perrucci, F. *et al.* Optimization of a suspended two photon polymerized microfluidic filtration system. *Microelectron. Eng.* **195**, 95–100 (2018).
- 6. Massaccesi, A. *et al.* 3D-Printable Dielectric Transmitarray With Enhanced Bandwidth at Millimeter-Waves. *IEEE Access* **6**, 46407–46418 (2018).
- 7. Bertana, V. *et al.* 3D-printed microfluidics on thin poly(methyl methacrylate) substrates for genetic applications. *J. Vac. Sci. Technol. B, Nanotechnol. Microelectron. Mater. Process. Meas. Phenom.* **36**, 01A106 (2018).
- 8. Muth, J. T. *et al.* Embedded 3D printing of strain sensors within highly stretchable elastomers. *Adv. Mater.* **26**, 6307–6312 (2014).
- 9. Baig, H., Sellami, N., Chemisana, D., Rosell, J. & Mallick, T. K. Performance analysis of a dielectric based 3D building integrated concentrating photovoltaic system. *Sol. Energy* (2014). doi:10.1016/j.solener.2014.03.002
- 10. Derakhshanfar, S. *et al.* 3D bioprinting for biomedical devices and tissue engineering: A review of recent trends and advances. *Bioactive Materials* (2018). doi:10.1016/j.bioactmat.2017.11.008
- 11. Parmeggiani, M. *et al.* P3HT Processing Study for In-Liquid EGOFET Biosensors: Effects of the Solvent and the Surface. *Proceedings* **15**, 39 (2019).
- 12. Preziosi, V. *et al.* Monitoring emulsion microstructure by Organic Electrochemical Transistors. *J. Mater. Chem. C* **5**, 2056–206 (2017).

- 13. Gentili, D. *et al.* Integration of organic electrochemical transistors and immuno-affinity membranes for label-free detection of interleukin-6 in the physiological concentration range through antibody-antigen recognition. *J. Mater. Chem. B* **6**, 5400–5406 (2018).
- 14. D'Angelo, P. *et al.* Monitoring the adaptive cell response to hyperosmotic stress by organic devices. *MRS Commun.* 7, 1–7 (2017).
- 15. Tarabella, G. *et al.* A hybrid living/organic electrochemical transistor based on the Physarum polycephalum cell endowed with both sensing and memristive properties. *Chem. Sci.* **6**, 2859–2868 (2015).
- 16. Battistoni, S. *et al.* Synaptic response in organic electrochemical transistor gated by a graphene electrode. *Flex. Print. Electron.* (2019). doi:10.1088/2058-8585/ab4dce
- 17. Scordo, G. *et al.* A novel highly electrically conductive composite resin for stereolithography. *Mater. Today Commun.* **19**, 12–17 (2019).
- 18. Bihar, E. *et al.* Fully printed all-polymer tattoo/textile electronics for electromyography. *Flex. Print. Electron.* **3**, 034004 (2018).
- 19. Tarabella, G. *et al.* Aerosol jet printing of PEDOT:PSS for large area flexible electronics. *Flex. Print. Electron.* (2019).
- 20. Tarabella, G. *et al.* Multifunctional operation of an organic device with three-dimensional architecture. *Materials (Basel).* **12**, (2019).
- 21. Fan, J., Montemagno, C. & Gupta, M. 3D printed high transconductance organic electrochemical transistors on flexible substrates. *Org. Electron.* (2019). doi:10.1016/j.orgel.2019.06.012
- 22. Majak, D., Fan, J. & Gupta, M. Fully 3D printed OECT based logic gate for detection of cation type and concentration. *Sensors Actuators, B Chem.* **286**, 111–118 (2019).
- 23. Havener, R. *et al.* Freeform fabrication of organic electrochemical transistors. *18th Solid Free. Fabr. Symp. SFF 2007* **1**, 60–73 (2007).
- 24. Kim, Y. H. *et al.* Highly conductive PEDOT:PSS electrode with optimized solvent and thermal post-treatment for ITO-free organic solar cells. *Adv. Funct. Mater.* **21**, 1076–1081 (2011).
- 25. Kim, N. *et al.* Highly conductive PEDOT:PSS nanofibrils induced by solution-processed crystallization. *Adv. Mater.* **26**, 2268–2272 (2014).
- 26. Xiong, Z. & Liu, C. Optimization of inkjet printed PEDOT:PSS thin films through annealing processes. *Org. Electron.* **13**, 1532–1540 (2012).
- 27. Baccar, Z. M. & Hafaiedh, I. Immobilization of HRP Enzyme on Layered Double Hydroxides for Biosensor Application. *Int. J. Electrochem.* **2011**, 1–5 (2011).
- 28. Pandurangachar, M. *et al.* Electrochemical investigations of potassium ferricyanide and dopamine by 1-butyl-4-methylpyridinium tetrafluoro borate modified carbon paste electrode: A cyclic voltammetric study. *Int. J. Electrochem. Sci.* **5**, 1187–1202 (2010).
- 29. Lyons, M. E. G. & Keeley, G. P. The redox behaviour of randomly dispersed single walled carbon nanotubes both in the absence and in the presence of adsorbed glucose oxidase. *Sensors* **6**, 1791–1826 (2006).
- 30. Moldenhauer, J., Meier, M. & Paul, D. W. Rapid and direct determination of diffusion coefficients using microelectrode arrays. *J. Electrochem. Soc.* **163**, H672–H678 (2016).
- 31. Gualandi, I. *et al.* Textile Organic Electrochemical Transistors as a Platform for Wearable Biosensors. *Sci. Rep.* **6**, 1–10 (2016).
- 32. Taormina, G., Sciancalepore, C., Messori, M. & Bondioli, F. 3D printing processes for photocurable polymeric materials: technologies, materials, and future trends. *J. Appl. Biomater. Funct. Mater.* **16**, 151–160 (2018).
- 33. Friedel, B. *et al.* Effects of Layer Thickness and Annealing of PEDOT:PSS Layers in Organic Photodetectors. *Macromolecules* **42**, 6741–6747 (2009).
- 34. D'Angelo, P. *et al.* PEDOT:PSS Morphostructure and Ion-To-Electron Transduction and Amplification Mechanisms in Organic Electrochemical Transistors. *Materials (Basel)*. **12**, 9 (2018).
- 35. Bernards, D. A. & Malliaras, G. G. Steady-State and Transient Behavior of Organic Electrochemical Transistors. *Adv. Funct. Mater.* 17, 3538–3544 (2007).
- 36. Rivnay, J. *et al.* Organic electrochemical transistors with maximum transconductance at zero gate bias. *Adv. Mater.* (2013). doi:10.1002/adma.201303080
- 37. Hofmann, A. I. *et al.* All-Polymer Conducting Fibers and 3D Prints via Melt Processing and Templated Polymerization. *ACS Appl. Mater. Interfaces* **12**, 8713–8721 (2020).
- 38. Khodagholy, D. *et al.* High transconductance organic electrochemical transistors. *Nat. Commun.* **4**, 2133 (2013).
- 39. Inal, S. *et al.* Organic electrochemical transistors based on PEDOT with different anionic polyelectrolyte dopants. *J. Polym. Sci. Part B Polym. Phys.* (2016). doi:10.1002/polb.23938
- 40. Rivnay, J. *et al.* High-performance transistors for bioelectronics through tuning of channel thickness. *Sci. Adv.* 1, 1–6 (2015).
- 41. Friedlein, J. T., McLeod, R. R. & Rivnay, J. Device physics of organic electrochemical transistors. *Org. Electron.* **63**, 398–414 (2018).
- 42. Tang, H., Lin, P., Chan, H. L. W. & Yan, F. Highly sensitive dopamine biosensors based on organic electrochemical transistors. *Biosens. Bioelectron.* **26**, 4559–4563 (2011).

- 43. Liao, C., Zhang, M., Niu, L., Zheng, Z. & Yan, F. Organic electrochemical transistors with graphene-modified gate electrodes for highly sensitive and selective dopamine sensors. *J. Mater. Chem. B* **2**, 191–200 (2014).
- 44. Gualandi, I. *et al.* Selective detection of dopamine with an all PEDOT:PSS Organic Electrochemical Transistor. *Sci. Rep.* **6**, 35419 (2016).
- 45. Tybrandt, K., Kollipara, S. B. & Berggren, M. Organic electrochemical transistors for signal amplification in fast scan cyclic voltammetry. *Sensors Actuators, B Chem.* **195**, 651–656 (2014).
- 46. Ji, W. *et al.* Carbonized silk fabric-based flexible organic electrochemical transistors for highly sensitive and selective dopamine detection. *Sensors Actuators B Chem.* **304**, 127414 (2019).
- 47. Chumillas, S., Figueiredo, M. C., Climent, V. & Feliu, J. M. Study of dopamine reactivity on platinum single crystalelectrode surfaces. *Electrochim. Acta* **109**, 577–586 (2013).
- 48. Bernards, D. A. *et al.* Enzymatic sensing with organic electrochemical transistors. *J. Mater. Chem.* **18**, 116 (2008).
- 49. D'Angelo, P. *et al.* Nanomolar detection of the antitumor drug tamoxifen by flexible organic electrochemical devices. *AIP Conf. Proc.* **1990**, (2018).
- 50. Wang, N., Liu, Y., Fu, Y. & Yan, F. AC Measurements Using Organic Electrochemical Transistors for Accurate Sensing. *ACS Appl. Mater. Interfaces* **10**, 25834–25840 (2018).
- 51. Qing, X. *et al.* Wearable Fiber-Based Organic Electrochemical Transistors as a Platform for Highly Sensitive Dopamine Monitoring. *ACS Appl. Mater. Interfaces* (2019). doi:10.1021/acsami.9b00115
- 52. Timpanaro, S., Kemerink, M., Touwslager, F. J., De Kok, M. M. & Schrader, S. Morphology and conductivity of PEDOT/PSS films studied by scanning-tunneling microscopy. *Chem. Phys. Lett.* **394**, 339–343 (2004).

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Author contributions statement

V.B., M.P. and S.L.M. conceived the device architecture; V.B., G.S. processed the materials and optimized the device fabrication; L.S., M.C. and S.F. performed the material analysis; M.G.G and V.B. optimized the printing parameters and analyzed the printing performance; M.P. conducted the electrical characterization and analyzed the related results; D.V. and P.D.A. performed the electrochemical characterization and the dopamine sensing, analyzing the related results; S.I. and C.F.P. supervised the study and validate the analysis: G.S., V.B., M.P., D.V., P.D.A. and S.L.M. wrote the original draft; all authors reviewed the manuscript.