

A Polymer Blend Substrate for Skeletal Muscle Cells Alignment and Photostimulation

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Substrate engineering for steering cell growth is a wide and well-established area of research in the field of modern biotechnology. Here we introduce a micro-machining technique to pattern an inert and transparent polymer matrix blended with a photoactive polymer. We demonstrate that the obtained scaffold combines the capability to align with that to photostimulate living cells. This technology can open up new and promising applications, especially where cell alignment is required to trigger specific biological functions, e.g., generate powerful and efficient muscle contractions following an external stimulus.

1. Introduction

In tissue engineering, cell-substrate coupling has a crucial role. Its mechanical, chemical, physical and morphological features are decoded by cells as stimuli that strongly affect their behavior. Often tools such as electrodes are further added to the cell culture in order to probe or actively stimulate the cellular processes. The use of electrodes has large practical success but it does bear a number of downsides like like unnatural heating, Reactive

Oxygen Species (ROS) production,^[1] spreading of the signal in a tissue and inflammatory response.^[2] Alternatively, photostimulation provides advantages such as better spatial and temporal resolution, low toxicity and invasiveness. In addition, the stimulation map throughout the sample can be changed easily without the need of rewiring nor the limit of discrete points, owing to the wireless architecture of this optical strategy. The use of light to control living cells and organisms started indeed long ago, but it is regaining interest due

to the emerging of new techniques at the frontier between photonics and biology. Optogenetics,^[3] near infrared (NIR)^[4] and photoactive material based^[5] stimulation are the most popular and promising techniques. In particular, the use of non-genetic phototransducers has the highest translational potential. Phototransducers may come in a variety of shapes and structures, such as plasmonic nanodots,^[6,7] inorganic semiconductor devices,^[8–10] nanostructures,^[11–14] organic semiconductors^[15–19] and small molecules.^[20,21] Thanks to the similarity in composition and structure with biomolecules, organic semiconductors and molecular compounds express higher biocompatibility and multifunctional actuation. Their peculiar photophysical properties allow the employment of different photostimulation mechanisms (i.e., electrical, mechanical or thermal) and at the same time, their mechanical flexibility favors conformational matching that is crucial for *in vivo* applications.^[22,23] Furthermore, by using wet processes at room temperature, organic semiconductors are easily fabricated into a great number of forms. The surface morphology of these shapes can be patterned to mimic the extracellular matrix and improve both cell growth and cell coupling at the abiotic/biotic interface.^[24,25] In particular, in muscular tissue engineering, in order to efficiently reproduce the natural muscle organization and achieve contraction ability, cell alignment on a suitably stiff substrate is required. This can be done by depositing extra matrix proteins^[26–29] or polymer fibers that produce an aligned path to force cell's orientation.^[30–32] Alternatively, a simpler approach relies on the fabrication of microgrooves onto a substrate that physically confine the cell, steering the cells in a direction parallel to the realized pattern.^[33–38] There are no universal rules that define which pattern can lead to an optimal morphological stimulation, since it strongly depends on the target cell. Nevertheless, some guidelines can be identified, regarding both the depth and width of the pattern: The depth should be higher than the cell's height

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(at least of the order of micron);^[39] The width should be lower than the cell's length.

In this work, we present a polymer substrate designed to perform a double functionality: cell alignment and cell photostimulation. We selected poly-(3-hexylthiophene-2,5-diyl) (P3HT), a very well-known polymer of organic photovoltaics, as photoactive material. Upon blending it with high-density polyethylene (HDPE),^[40] we obtained a free standing polymer film with good mechanical properties that preserves P3HT optoelectronic features. Furthermore, by using a maskless laser ablation approach we were able to inscribe a customizable guiding pattern in the films suitable for achieving aligned cell growth. The combination of blending and micromachining is used as a novel approach to achieve a multifunctional substrate without the employment of complex and expensive fabrication methods such as photolithography. Moreover, the upscaling of this approach can be easily achieved in order to increase the complexity of the realized pattern. As a simple model system for preliminary biocompatibility and photostimulation studies we used Human Embryonic Kidney (HEK) cells. Once parameters had been optimized we focused on C2C12, a murine myoblast cell line characterized by myotube diameter ranging from 10 to 25 μm and length ranging from 130 to 520 μm .^[41] C2C12 show many peculiar properties of skeletal muscle cells (i.e., the ability to differentiate toward multinucleated myotubes that can display contraction) thus making them a widely accepted *in vitro* model to validate our purpose.

2. Result and Discussion

It has been widely reported in literature that P3HT can be blended with a large variety of insulating polymers preserving its optoelectronics properties.^[42–46] The blend strategy is

employed to introduce desired processing and mechanical properties of commodity polymers (i.e., common plastics) in organic electronics applications. Plastic “additives” are ductile, display high elongation at break, while in contrast semiconducting polymers are typically rather brittle owing to their low molecular weight.^[43] Moreover, insulating polymers have been reported to be beneficial in a number of applications leading for instance to improved charge carriers mobilities, increased device stability,^[47–50] tailored optical properties^[51,52] and decreased energetic disorder.^[53] We focus here on HDPE, a semi-crystalline insulating polymer that has been already employed in combination with P3HT to produce both thin and thick films for a large variety of applications. Remarkably, the blend system retains semiconducting optoelectronic properties even at very low concentrations of the photoactive material in the insulating matrix, while showing improved mechanical properties. Indeed, the improved viscoelastic properties of the blend system allow the production of robust self-standing films that can be used directly in tissue engineering, while films comprising neat semiconducting polymers would be too brittle to do so. Beside the combination of mechanical and optoelectronic properties, blending has an economic advantage for HDPE being much cheaper than P3HT, which is particularly useful when producing thick films (tens of microns).

2.1. Fabrication Process

The fabrication process is designed to obtain a photoexcitable substrate suitable for laser pattern ablation. Muscle cell have typically height of $\approx 15 \mu\text{m}$ and length of $\approx 200 \mu\text{m}$.^[41] According to the requirements discussed above, we selected a pattern geometry formed by parallel lines with a width of 60 μm and a depth of at least 15 μm , thus requiring a film

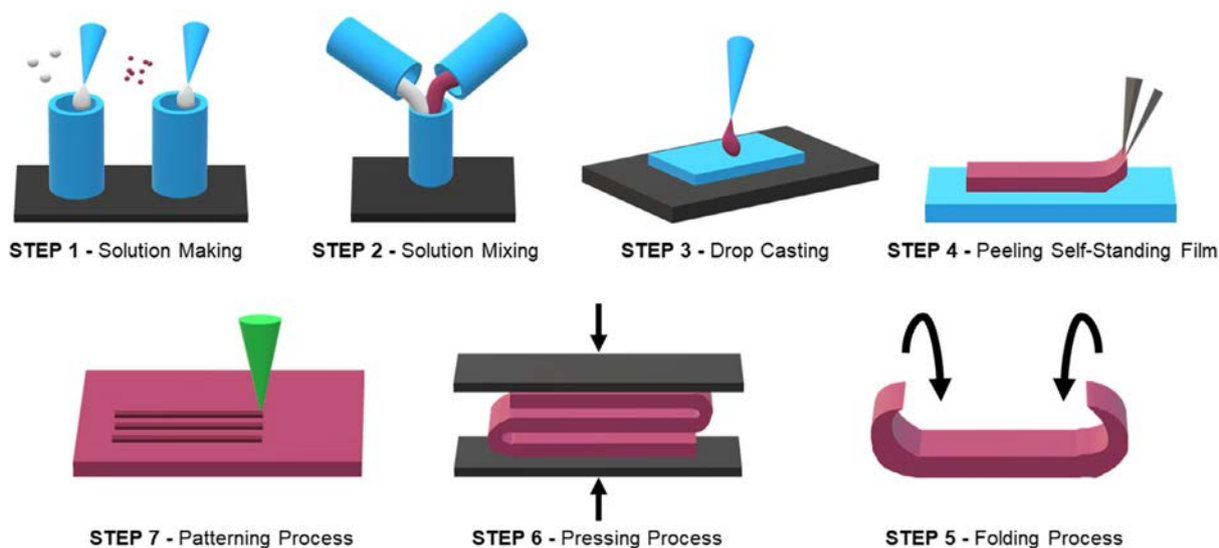


Figure 1. Schematic representation of the multistep fabrication process. Step 1 and 2 regards the solution making process: two different solutions are made for each neat polymer dissolved in the same solvent and at the same concentration and successively mixed with a specific ratio. From step 3 to step 6 the realization of a thick HDPE:P3HT film is illustrated. The process is based on drop casting (step 3), peeling and subsequent folding of the solid deposit (step 4–5) that precede the pressing step (step 6).^[56] Finally, a patterning process is performed upon laser ablation (step 7). During the deposition steps, glass slides and hot plate are used, they are shown in blue and dark grey in the sketch. The press plates, used in the pressing step, are depicted in dark grey.

thickness $>20\ \mu\text{m}$. The fabrication process (Figure 1) is based on a pre-compounding step performed in solution, followed by solid-state processing of the self-standing film and final laser patterning. In principle, the solution processing could be avoided and the polymer:polymer compounding could be achieved upon melt mixing P3HT and HDPE, leading to a solvent free process. However, the latter requires the employment of a large volume of materials and high temperature of processing owing to the high melting point of P3HT ($\approx T = 238\ ^\circ\text{C}$).^[54] Consequently, the solution-based method is more practical. Once a homogeneous solution is achieved, it was drop cast to obtain a solid state deposit. Despite this deposit exhibits the required self-standing and mechanical properties, it lacks the necessary thickness and homogeneity. In order to reach a suitable film thickness, around $30\ \mu\text{m}$, we delaminated, folded and pressed the polymer blend into a uniform film. Solid-state pressing is a versatile technique that is widely employed in the processing and embossing of polymer films.^[55] The blend strategy is particularly useful, not only to tune the mechanical properties of the films, but also to control the optical density. Indeed, HDPE does not absorb in the UV-vis range (see Figure S1, Supporting Information) and when intermixed with the semiconductor it enhances the transparency of the otherwise opaque P3HT thick film. Films obtained with a concentration of P3HT in the range 1–5 wt% show an optical density suitable for photostimulation experiments at the required thickness of $30\ \mu\text{m}$.

2.2. Photophysical Properties

The optical properties and elementary excitation dynamics of the P3HT:HDPE films were investigated by measuring absorbance, photoluminescence (Figure S1, Supporting Information) and transient photoinduced absorption (Figure 2) in a spin-cast P3HT thin film (P3HT-TF) and for the three different HDPE: P3HT pressed blends (with P3HT concentration equal to 1, 3, and 5 wt%). Note that HDPE is virtually transparent in the considered spectral range (400–800 nm) as shown in Figure S1, Supporting Information, thus spectroscopy essentially addresses the P3HT properties.^[57–61] The optical features of the three blends are very similar to that of the neat film suggesting that: a) the fabrication process has a minor effect onto the P3HT photophysics and b) P3HT successfully phase separates from HDPE in the blend. We selected a deposition temperature of $100\ ^\circ\text{C}$, in a range in which P3HT is known to phase separate and crystallize^[62–64] prior the HDPE crystallization during the solvent evaporation,^[40] leading to an unperturbed UV-Vis absorption of the semiconductor.

The presence of ordered P3HT domains is highlighted by the structured absorption spectra, which are typical of aggregates in this polymer.^[63,65] By and large the transient absorption data confirm this picture, even though quantitative differences are present depending on the blend composition. In particular, in the 1% blend the photoinduced absorption (PA) band

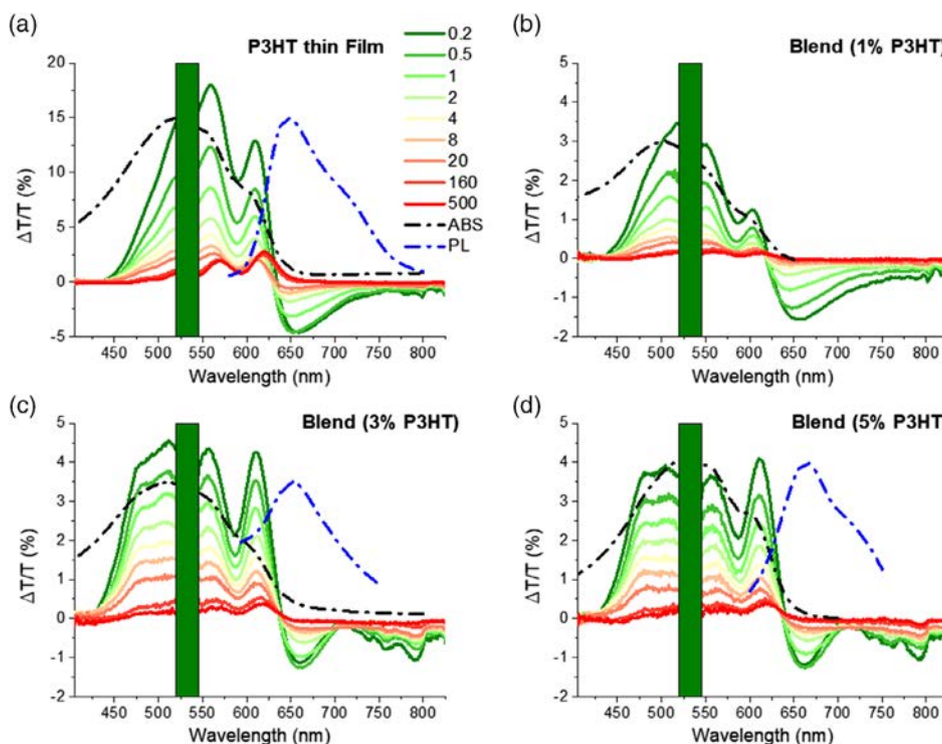


Figure 2. Comparison between spectroscopic behavior of P3HT-TF and the blended film obtained with different percentages of P3HT. (a) P3HT-TF, (b) blend (1% P3HT), (c) blend (3% P3HT), (d) blend (5% P3HT). In each panel the absorbance spectrum (dash-dot black line), the photoluminescence spectrum (dash-dot blue line) and the transient absorption spectra (solid lines) are reported. The time evolution is indicated by the color gradient of the line (from dark green = 0.2 ps to dark red = 500 ps). In 2-b the photoluminescence spectrum is not reported because it is overwhelmed by the sample scattering (see Supplementary Information for more details).

($\Delta T < 0$) appears much broader than in the other spectra, suggesting a more efficient charge separation process. In the 3% and 5% samples the first vibronic resonance of the bleaching band ($\Delta T > 0$) is enhanced suggesting a higher degree of order in the P3HT domains. This hypothesis is enforced by the red shift of the emission spectra as a function of the P3HT concentration that suggests an enhanced aggregation as well as a planarization of the P3HT molecules (Figure S1, Supporting Information). Despite a more detailed analysis is beyond the scope of this work, we can speculate that the crystal packing of P3HT chains secluded in the blend might favor singlet fission into triplets. The transient absorption analysis corroborates the steady state data, confirming that the P3HT mostly retains its excited state dynamics upon blending in the optically inert matrix. The enhanced vibronic replica, as well as the appearance of the SE convoluted with the PA at 720 nm observed at the highest P3HT loadings might be related to the planarization of P3HT chains.

2.3. Substrate Biocompatibility

The biocompatibility of the substrate and of the patterning process were tested with two distinct assays. For substrate biocompatibility, the alamarBlue proliferation test was performed with HEK cells (Figure 3b). These cells were preferred for this kind of methodology due their dimension, which is lower than C2C12 and therefore more suitable for acquiring data for longer proliferation times and for choosing a higher seeding density, a crucial aspect for this test. The biocompatibility of the patterning process was also tested, to be sure that laser ablation did not produce toxic fabrication debris. Conversely, C2C12 viability was evaluated with the HOECHST 33342/NucGreen Dead 488 ReadyProbes assay (Figure 3c). Both the film and the patterning process highlight very good biocompatibility showing proliferation and viability rates similar to those obtained seeding the cells on glass, that represents our standard control condition.

2.4. Cell Photostimulation

Having established that the blends retain their optical properties and have suitable biocompatibility, we went on studying the performance in cell photo-stimulation. The electrophysiological properties were measured by carrying out patch clamp experiments in whole-cell configuration and current clamp mode ($I = 0$). Using this approach, changes of the cell membrane potential are recorded during light excitation. As we showed in previous works,^[66,67] the cell membrane undergoes a depolarization and a subsequent hyperpolarization upon illumination when cells are plated on P3HT-TF. We measured the cell membrane variation of both HEK and C2C12 cells seeded on thick solid state pressed P3HT:HDPE films when short (20 ms) and long (200 ms) light pulses ($\lambda = 530$ nm) are applied (Figure 4) at a fixed power density (40 mW mm^{-2}).

We detected a transient depolarization upon illumination with short pulses, quickly evolving into hyperpolarization when the light was switched off. The membrane potential recovered its resting value approximately after 300 ms (Figure 4a). By using longer pulses the initial depolarization turned into hyperpolarization during illumination. At the light offset there was again an overshoot increase of the hyperpolarization, followed by a recovery to the initial resting potential in about 300 ms (Figure 4b). A very good reproducibility of the data is observed, as it can be noticed from the low dispersion of the data in both the cell lines and the pulse length shown in Figure 4b–d. We performed measurements on a control sample (black line Figure 4) without observing any signal during the measurement, ruling out a direct interaction between cell or HDPE with light. Once the photoinduced effect has been demonstrated, we performed further studies on the biophysical mechanisms. In previous works, the photoinduced membrane potential variation, measured in cells seeded on bare P3HT films and subjected to light stimulation protocol similar to the one adopted here, was assigned to a thermal effect. In particular, the initial depolarization is reported to be due to the increase of capacitance of the cell membrane, while the following hyperpolarization, still under light exposure,

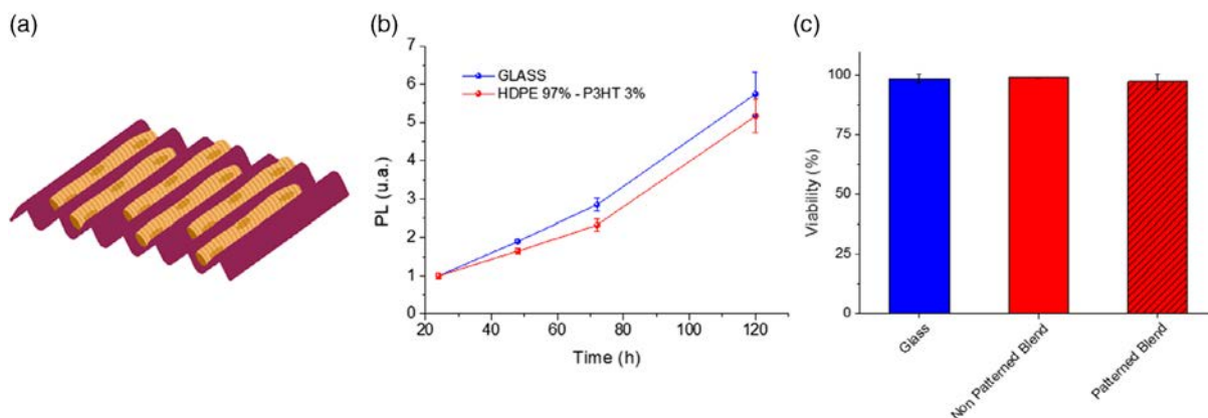


Figure 3. Results of biocompatibility tests. a) Sketch of cells seeded on the patterned blend film. b) Proliferation assay (alamarBlue) performed on HEK cells. Measurements were performed after 1, 2, 3, and 5 days after seeding. c) Viability test for C2C12 cells cultured on bare glass, blend and patterned film (3% P3HT). Data are presented as mean \pm s.e.m., $n = 3$ wells measured 5 times for proliferation and $n = 80$ cells for viability.

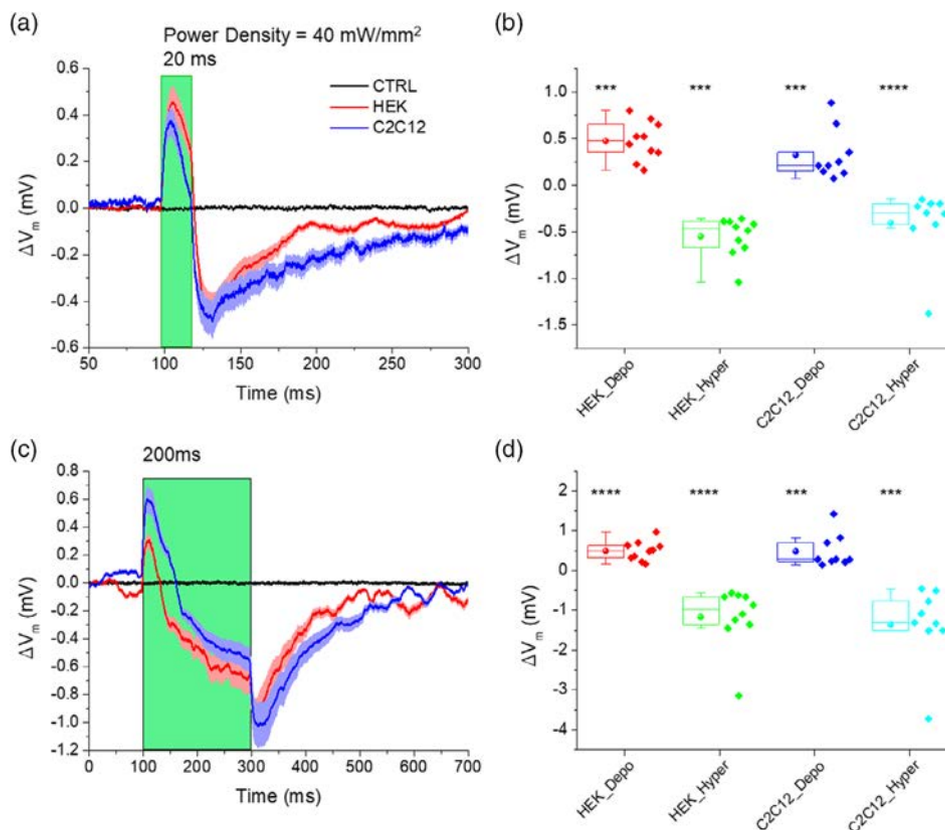


Figure 4. Electrophysiological response of HEK and C2C12 cells. Representative curves of the membrane potential during light excitation for 20 ms (a) and for 200 ms (c) light pulses. The black curves are the control measurement from cells grown on HDPE films. Box plots of depolarization (b) and hyperpolarization (d) changes obtained for both HEK ($n = 10$) and C2C12 ($n = 10$) cells with a photoexcitation density of 40 mW mm^{-2} at 530 nm. Statistical comparison is evaluated between resting and light induced membrane potential value.

is assigned to the change in the cell baseline.^[66] The dark evolution towards equilibrium is ascribed to the recovery of capacitance and cooling off. In principle our blend could have a different thermodynamic behavior under illumination when compared with the neat thin film.

We therefore measured the temperature variation in close proximity to the substrate, using the calibrated pipette method (Figure 5a,b) described by Martino et al.^[66] Effectively, upon

irradiation with a 200 ms light pulse, the temperature rose up to 33°C , with $\Delta T_{\text{Blend}} = +9^\circ\text{C}$, which is comparable to that observed on neat P3HT thin films, $\Delta T_{\text{P3HT-TF}} = +7^\circ\text{C}$ with a comparable power density ($\approx 40 \text{ mW mm}^{-2}$). Effectively, absorption in the blend is large enough to give rise to a change in temperature that slightly overcomes that observed in the neat film, despite the low P3HT density in the blend. This could probably be due to the different thickness and the different thermal

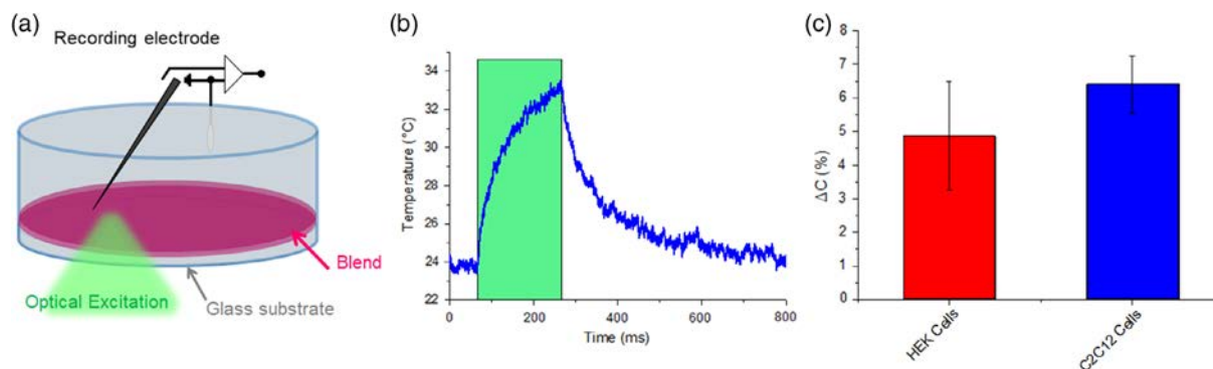


Figure 5. Proposed photothermal mechanism. a) Sketch representation of the calibrated pipette method. b) Temperature variation of the extracellular bath upon 200 ms photoexcitation as a function of time. The experiment is performed at room temperature (24°C) with a photoexcitation density of 40 mW mm^{-2} at 530 nm. c) Light induced variation (mean \pm s.e.m.) of the whole cell capacitance for HEK ($n = 10$) and C2C12 ($n = 10$) cells.

conductivity of the two films. To confirm the mechanism reported above we also measured the capacitance in dark condition and upon light excitation by applying the double sinusoidal technique. Figure 5c reports the results. As expected, the capacitance increases due to light excitation in both the investigated cell lines. The percentage increase, ΔC , is $4.87 \pm 1.62\%$ for the HEK cells and $6.40 \pm 0.85\%$ for C2C12 cells.

2.5. Laser Ablation and Cell Orientation

Once the cell photostimulation ability of the plain pressed blend was assessed, we further aimed at micromachining its surface in order to force cell alignment. The manufacturing process creates a grooved line pattern on our blend substrate taking advantage from the ultra-short pulse laser direct surface ablation technique. This maskless fabrication technique gives a high degree of freedom in designing both the shape and the wall dimension of such patterns, allowing geometries that are more complex. By changing the repetition rate, the pulse energy or the position of the laser beam focal spot it is possible to tune the wall depth, the

geometry (which can also be curvilinear) and control the spatial resolution of the chosen pattern (as shown in Figure S3, Supporting Information). Furthermore, direct laser ablation provides a quicker and cheaper manufacturing technique, compared to classical lithographic methods, that well matches the cost-effectiveness of the blended substrates presented in this work. We selected a simple square cross-section target pattern with a wall depth greater than $20\ \mu\text{m}$ and a width of $60\ \mu\text{m}$ (Figure 6). We then assessed the ability of the realized pattern to align cells with Scanning electron microscopy (SEM), by acquiring images of C2C12 cells seeded directly on the substrates. To better visualize the effect of the pattern we first selected the experimental conditions that allowed the growth of isolated cells. A representative image acquired in this condition is shown in Figure 7a, where it is possible to appreciate how the substrate pattern effectively steers the cells orientation. In Figure 7b we show the polar plot of the alignment angles, where 0° corresponds to the pattern direction.

The mean orientation angle is $12.95^\circ \pm 1.52^\circ$. Moreover, a good alignment efficiency is observed, indeed 77% of the cell's

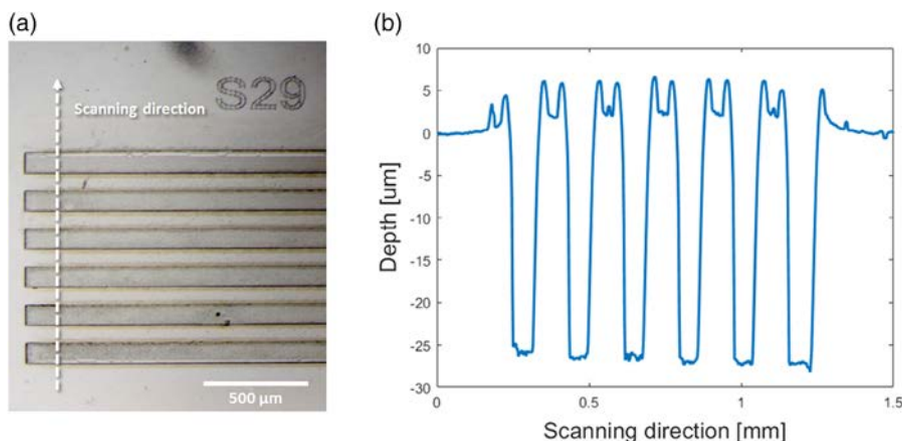


Figure 6. Obtained pattern by laser ablation. a) Microscope image of the pattern obtained with the ultra-short pulse laser direct ablation process. b) A profile of the obtained pattern shows its characteristic dimensions, a depth of $25\ \mu\text{m}$ and a width of $65\ \mu\text{m}$. The scanning direction is represented with a dashed arrow in panel a.

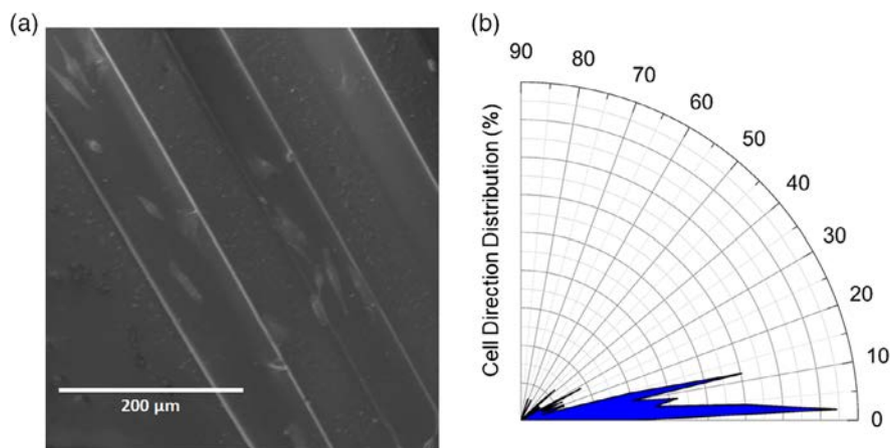


Figure 7. Pattern induced cell orientation. a) SEM image of C2C12 seeded on patterned substrate in isolated cells condition. b) Polar plot of the cell directionality distribution normalized on the pattern direction ($n = 83$ cells).

directions are included in the angular interval 0° – 15° and 90.3% in 0° – 30° . Having established that cell growth anisotropy is achieved in the low-density cell regime, we carried out also high cell density experiments trying to achieve a muscular tissue (see supporting materials). Despite it is hard to obtain a quantitative evaluation of the alignment; nevertheless it is still possible to observe that the cell growth follows the pattern direction (see Figure S5, Supporting Information). The effect is particularly evident at the edge of the pattern where it is possible to appreciate an evolution of the cells from a random to an aligned orientation condition canalized by the pattern.

3. Conclusion and Perspective

We have developed a substrate that combine, for the first time, the ability to align and photostimulate cells. We have achieved this by fs-micromachining a polymer blend comprising the semiconducting polymer P3HT and the insulating polymer HDPE. This is an enabling technology that can open up new applications, especially when cell alignment is mandatory to carry out specific functions, e.g., muscle contraction. For instance, this novel platform can lead to the fabrication of systems in which fully developed skeletal or cardiac muscle cells can grow in an organized array that allows carrying out specific functions under light control. The shape of the array can be easily tuned with our fs-micromachining approach, given its capability of easily patterning any shapes and sizes, without the need of using any masks. The same setup also allows a high-resolution soft-cut of the material to give a macro customized shape to the patterned areas. The P3HT:HDPE blend is a proof-of-concept system, however the absorption spectra of our substrates can be tailored upon selecting a different semiconducting polymer, so that light can penetrate deeper into the tissue in order to broaden the spectrum of possible biological applications. Moreover, HDPE can be replaced with hydrogel or elastomeric materials to well reproduce biological stiffness and to achieve ink-properties compatible with 3D-printing techniques, thus aiming to reproduce the structural complexity of living tissue.

4. Experimental Section

Sample Preparation: Regio-regular P3HT (99.995% purity, $M_w = 20\text{--}45\text{ kg mol}^{-1}$) was purchased from Sigma Aldrich and used without any further purification. High-density Polyethylene ($M_w = 125\text{ kg mol}^{-1}$) was purchased from Alfa Aesar and used without any further purification. Solutions of the two material (P3HT and HDPE) were prepared in di-chlorobenzene at concentration of 15 g L^{-1} , mixed in a specific ratio (1–99, 3–97, and 5–95 wt%) and stirred for at least 30 min. The obtained mixing solution was drop casted on a hot plate at 100°C . Once the solvent was evaporated the obtained film was folded and pressed with a hydraulic press with several step to arrive to the maximum pressure of 7 MPa held for 8 min at 140°C . Although the solution step is not required and the exact amount of powder can be directly pressed to obtain the film, in this work we decided to go through solution to be more precise in the mixing step. The obtained film was circular with a diameter of 18 mm and thickness of 30 μm .

UV-Vis Absorption: The UV-Vis absorption spectra were acquired with a Perkin Elmer Lambda 1050 spectrophotometer equipped with deuterium (280–320 nm) and tungsten (320–3300 nm) lamps. The signal is acquired

with three detectors acting in different spectral region (photomultiplier [180, 860] nm, InGaAs [860, 1300] nm and PbS [1300, 3300] nm).

PL Measurements: The PL measurements were carried out with a Horiba Nanolog Fluorometer equipped with two detectors (photomultiplier and InGaAs). The sample was excited at 530 nm using the same LED lamp and power employed for transmission measurements (see above).

Ultrafast Time-Resolved Spectroscopy: For the femtosecond TA measurements, we used a Ti:Sapphire laser with a repetition rate of 1 kHz and a pulse width of 100–150 fs. We excited the films at 530 nm. The beam was generated by means of an optical parameter amplifier and probed with a white light beam generated by a CaF₂ plate. The excitation energy was of $\approx 30\text{ }\mu\text{J}$ and the beam spot size of $\approx 200\text{ }\mu\text{m}$ in diameter.

Laser Pattern: The micromachining setup consists of an amplified Yb:KGW femtosecond laser system (Pharos, Light Conversion) with 230 fs pulse duration, 515 nm wavelength (frequency doubled), 500 kHz repetition rate focused with a 20X–0.42 NA microscope objective (Mitutoyo). Computer-controlled, 3-axis motion stages (ABL-1000, Aerotech) interfaced by CAD-based software (ScaBase, Altechna) with an integrated acousto-optic modulator are used to translate the sample relative to the laser irradiation desiderate patch. An average power (pulse energy) of 15 mW (30 nJ) and a scan speed of 1 mm s^{-1} are used to laser pattern the substrates.

Cell Culture Maintenance: HEK and C2C12 cells were cultured in T-75 flasks containing Dulbecco's modified Eagle's medium (DMEM) supplemented with Fetal Bovine Serum (10%), glutamine (2 mM L^{-1}), Penicillin (100 IU mL^{-1}) and Streptomycin ($100\text{ }\mu\text{g mL}^{-1}$). Culture flasks were maintained in a humidified incubator at 37°C with 5% CO₂ and, when at confluence, plated at a density of 15,000 cells cm^2 and cultured for 48 h on samples. Prior to cell plating a layer of fibronectin ($2\text{ }\mu\text{g mL}^{-1}$ in PBS buffer solution) was deposited on the samples surface and incubated for 1 h at 37°C , to promote cellular adhesion. Excess fibronectin was then removed by rinses with PBS.

Viability Assay: To preliminarily evaluate the substrate cytotoxicity, alamarBlue proliferation assay, that allows a continuous monitoring of cells in culture, was performed on HEK cells. Briefly, the alamarBlue Reagent (Invitrogen DAL 1100) was diluted 11:1 with DMEM without phenol red. 500 μL of the obtained solution were added on each well. The solution was incubated for 3 h at physiological condition. Once removed from the well, the emission at 590 nm of three aliquots per each well was measured. The emission value was acquired 5 times per each aliquot to obtain a reliable measure. Furthermore, the biocompatibility of the patterning process was accomplished on the more relevant biological model C2C12 via HOECHST 33342/NucGreen Dead 488 ReadyProbes assay. The substrates were incubated in extracellular containing the two dyes (HOECHST 33342 ($10\text{ }\mu\text{g mL}^{-1}$) and NucGreen Dead 488 ReadyProbes Reagent (2 drops mL^{-1})) for 5 min protected from ambient light. The samples were then washed with extracellular solution and multiple images were acquired with a Nikon Eclipse Ti-S epifluorescence inverted microscope. Standard DAPI and FITC filter sets were employed for HOECHST and NucGreen respectively. The percentage of viable cells was estimated by counting the total number of cells nuclei (stained by HOECHST) and the total number of dead cells nuclei (stained by NucGreen).

Scanning Electron Microscopy (SEM): C2C12 cells were plated on Patterned HDPE-P3HT (97–3% in volume) substrates. These substrates were prepared for SEM following three steps. First, fixation in glutaraldehyde 4% in PBS for 20 min at room temperature; Second, immersion in increasing concentrations of ethanol (20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, and 100%, 20 min for each concentration) followed by air-drying at room temperature; Three, evaporation of a thin gold layer on top of samples surface (thickness 5 nm, 1.5 Cr adhesion layer). All SEM images were acquired by using a TESCAN MIRA III scanning electron microscope (operating voltage 3 kV, working distance 7 mm, stage tilt angle 0°).

Electrophysiology: Standard patch clamp recordings were performed using an Axopatch 200B (Axon Instruments) coupled to an inverted microscope (Nikon Eclipse Ti). HEK and C2C12 cells seeded on bare glass/HDPE/HDPE:P3HT blend were measured in whole-cell configuration with freshly pulled glass pipettes (4–7 M Ω), filled with the following intracellular

solution [mM]: 12 KCl, 125 K-Gluconate, 1 MgCl₂, 0.1 CaCl₂, 10 EGTA, 10 HEPES, 10 ATP-Na₂. The extracellular solution contained [mM]: 135 NaCl, 5.4 KCl, 5 HEPES, 10 Glucose, 1.8 CaCl₂, 1 MgCl₂. Only single cells were selected for recordings. Acquisition was performed with pClamp-10 software (Axon Instruments). Membrane currents were low pass filtered at 2 kHz and digitized with a sampling rate of 10 kHz (Digidata 1440 A, Molecular Devices). Data were analyzed with Origin 9.0 (OriginLab Corporation) and with Matlab software. The light source for excitation was provided by a green LED coupled to the fluorescence port of the microscope and characterized by maximum emission wavelength at 530 nm; the illuminated spot on the sample has an area of 0.23 mm² and a photoexcitation density of 40 mW mm⁻², as measured at the output of the microscope objective (Pobj).

Cell Capacitance Measurement: A double sinusoidal voltage-clamp signal is applied to the cell in whole-cell configuration. The response current signal is acquired and membrane capacitance, membrane resistance and access resistance are then extracted fitting the current with a custom Matlab program.

Statistical Analysis: Data are represented as mean ± standard error of the mean (s.e.m.). Normal distribution was assessed using D'Agostino-Pearson's normality test. To compare two sample groups, either the Student's t-test or the Mann-Whitney U-test was used. In all experimental settings, **P* < 0.05, ***P* < 0.01, ****P* < 0.001, *****P* < 0.0001. Statistical analysis was performed using GraphPad Prism 7 software (GraphPad software, Inc, La Jolla, CA).

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Authors Contributions

V.V. and A.D.S. contributed equally to this work. V.V. and G.L. planned the experiments. V.V. and A.D.S. developed the fabrication process of the blend. V.V. and F.L. carried out photostimulation experiment. V.V. took care for cell cultures, and S.E.M. sample preparation. A.M.R. acquired the transient absorption data. F.S. design, developed and performed the laser ablation. F.M. acquired the S.E.M. images. V.V., A.D.S., F.L., and G.L. wrote the manuscript. All authors contributed to data interpretation and approved the final version of the manuscript.

Keywords

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