Physarum Polycephalum changes polyaniline properties

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Abstract

Physarum polycephalum slime mould can modify polyaniline (PANI) features due to its internal activity. We created networks with different conductivity made by the slime mould on PANI substrates. Thus, Physarum’s growth results in changing the conductivity state of PANI layers, providing negative and positive patterning of the samples. A spectrophotometric scanner is here exploited to investigate and characterize the effects coming out from the interaction between Physarum polycephalum and PANI. The latter is an electro-chromic polymer that vary its colour and conductive properties according to its redox state.

Introduction

Physarum polycephalum slime mould is attracting interest of scientists, not only biologists, because of its incredible features and adapting capabilities. Indeed, since many years, it has been the object of studies in the field of unconventional computing, (Gale et al. 2013), networks modelling and development (Nakagaki et al. 2004), biorobotics (Mayne et al. 2013), and biochemistry (Romeo et al. 2015). Physarum polycephalum slime mould is a single cell organism with many nuclei dispersed in the cytoplasm. It belongs to the family of Myxomycetes, Physarales species, taxonomically classified as Protozoas or, simply, slime mould (Ratzel et al. 2013). In nature it feeds on bacteria and decaying organic materials and needs darkness, humidity and a temperature around 25-27°C. Therefore, the abovementioned characteristics have to be considered in order to properly culture a colony in a laboratory. Physarum polycephalum can be, thus, cultivated on agar gel or wet towels kept in a dark and humid chamber and fed with oat-flakes. Moreover, to ensure its safety, the colony has to be periodically replanted to a new fresh substrate (agar or towels). Physarum polycephalum life is characterized by different phases (Stephenson et al. 2000): sporulation, sclerotium and plasmodium. The first is the reproduction phase, exploited by meiosis and fructification; the second is the dormant phase, a sort of hibernation during which the mould protect itself from not proper environmental conditions. The latter, plasmodium, is the vegetative form of slime mould; during this phase the organism is more active and moves searching for food. It appears as a yellowish multinuclear mass of protoplasm, a single cell with a myriad of nuclei, able to regenerate autonomously, even cutting a part away. Therefore, it has not a fixed mass, but it continuously changes its shape as a function of the spatial food distribution; indeed, it has been also demonstrated its capability of choosing not only the closest but also the most nutritional food sources from a buffet (Latty et al. 2009). Figure 1 shows an example of the abovementioned feature. Its foraging behaviour consists in the formation of optimized networks (Bonifaci et al. 2012) of protoplasmic tubes branching towards nutrients. However, in the case of food abundance it tends to stasis creating “pancakes-like” structures that enwrapped the whole food allowing phagocytosis (Golderer et al. 2001). These are astonishing features for a simple amoeba. In addition, as its name suggest, it produces a certain amount of slime, an extracellular polysaccharide (Simon et al. 1970) working as a sort of “external brain”. The slime allows Physarum polycephalum to remember already trodden paths, in order to not retrace them when seeking for food or other attractants (Nakagaki et al. 2001). The protoplasmic veins of plasmodium’s network consist in a two phases sol-gel medium. The external, more rigid, gel is the ectoplasm and contains the sol, more fluid endoplasm (Wohlfarth-Bottermann, 1974) which is transported by means of proteins present in the ectoplasm, the actomiosins. The latter, by contracting, generates high-pressure gradients that push the endoplasm forward allowing locomotion of the whole organism, in the so called shuttle streaming mechanism (Matsumoto et al. 2008). The way by which Physarum feels attractors and food in a certain direction, as well as the individuation of the forces involved in the locomotion of this organism, are still unknown. In this interesting scenario, we exploited Physarum’s capability of creating networks (Shirikawa and Gunji, 2007) to pattern polyaniline (PANI) samples.

PANI is a redox electro-chromic polymer widely studied, also by authors for organic memristive device realizations (Erokhin and Fontana, 2011). Moreover, a spectrophotometer device has also been used to get evidence of the modifications induced by the mould on the PANI samples. In this work, we found a method by which it is possible to characterize Physarum’s networks by electrical and spectrophotometric measurements. The latter, built by the mould onto PANI substrate, were transferred, as a lithographic process, on PANI itself.

Therefore, we finally project *Physarum* done networks on the PANI layer.

![Figure 1: Optical microscope image of Physarum polycephalum on Agar nutrient gel, in this case it is clearly visible that the mould is creating a sort of "pancake-like" structure growing in all directions homogeneously. Thus demonstrating its shape is a function of spatial food distribution.](image)

**Materials and methods**

**Physarum culture**

*Physarum polycephalum* slime mould, as mentioned in the introduction section, was cultured in dark and humid chamber at room temperature. The colony was maintained in Petri dishes with agar 1.5% non nutrient gel, fed with oat-flakes and periodically replanted to a new fresh agar.

**Spectrophotometer**

The device (Dimonte et al. 2015), a spectrophotometric scanner, is generally exploited in the art and restoration fields, being able to appreciate chromatic differences with high spatial resolution between points. The instrument produces an image consisting in an ordered collection of the spectral reflectance factors of each pixel.

The spectrophotometer here used is composed of a transmission spectrometer (Inspector V8 manufactured by Specim, Finland) designed for a 2/3 inch CCD sensor equipped with a 25 μm entrance slit and covering the 400÷780 nm spectral range with a spectral resolution of about 2 nm. The spectrometer is coupled to a monochromer 2/3 inch CCD matrix chill digital camera (Hamamatsu C4742-12bit, 1280×1024 pixels, 9 f/sec) while a collecting lens (Computar TEC-M55 designed for a 2/3 inch sensor) focalizes the painting on the plane of the entrance slit. The illumination is obtained by means of two 150 W halogen lamps whose light is filtered preventing the illumination of the painting. The digital camera is interfaced to a PC by means of a 12 bit frame grabber (Mutech MV1000). A software program drives the scanner, acquires data of a strip of the scene and allows to save its image as a spectral image. The program is implemented to reproduce the colours on a calibrated CRT monitor. The light is dispersed by the spectrometer and focalized in the plane, containing the sensor of the camera. The spectrometer has a 1:1 image magnification then, the image of the input is focalized on the pixel rows of the sensor, while its position along the vertical axis of the sensor, depends on the light wavelength. White light in the range 400÷780 nm enterig into the device fills the whole sensor. The acquisition of one frame of the digital image can be thought as the acquisition of the reflectance spectra of each pixels of the strip of the painting.

**Polyaniline samples**

Polyaniline is a redox electrochromic polymer, discovered in 1985 (Kang et al. 1998). Its main feature is represented by a high conductivity difference between the conductive, green, oxidized state and the insulating, blue reduced one (Bredas et al. 1985). The transition depends on the doping degree. It can be achieved by chemical treatment (i.e. water or basic solutions for reducing and acid chloride for oxidizing) or by proper voltage application (Berzina et al. 2007). Emeraldine base polyaniline was purchased from Sigma (Mn ca. 100,000). The deposition of the active PANI layer was carried out with a KSV 4000 LB trough, using a Langmuir–Schaefer technique. Pure water, purified with a Milli-Q system, with a resistivity of 18.2 MΩcm, serves as the subphase (Dimonte et al. 2014). Polyaniline powder was dissolved in 1-methyl-2-pyrrolidones/NMPd and carefully filtered. The real concentration of the solution was determined and then NMP was added to achieve the final concentration of 0.2 mg/ml.

**Oxygen-Containing Plasma Treatment**

The treatment has been performed with a Plasma Matrix bdiscom machine with a 10 min exposure at 99 W. This allows us to remove PANI from the areas of the samples not covered by mould’s network.

**Optical measurements**

The samples have been characterized by means of an optical microscope Leica D300. In addition, it was also fundamental, at the final state of the work, to make proper contacts of the PANI networks by means of micro-manipulating tips.

**Electrical charaterizations**

The electrical conductivity of the PANI networks was recorded by means of a Keithley 2400 SourceMeters and two micromanipulators ending with tungsten tips.

**Experiments and Results**

Polyaniline samples were prepared by depositing 40 molecular layers of PANI on glass substrates by Langmuir–Schaefer technique (see material and methods); afterward the layer was doped by HCl treatment.

We put 2 µl blob of *Physarum polycephalum* in the centre of a glass with deposited PANI film and attractors (oat flakes) at the external boundaries, to stimulate the mould in exploring the sample. Therefore, *Physarum polycephalum*, spanning towards the food sources in 8-12 hours, created networks on the polymer substrate as it is shown in Figure 2a).

The sample was then transferred to open air and light conditions to let *Physarum* enter in the sclerotic phase and, consequently, maintain unvaried the designed network. Subsequently, we applied Oxygen plasma to remove PANI in all samples areas not covered by the mould.

The result is shown in Figure 2b). The reported picture allows to conclude that the performed treatment did not destroy the network and the sclerotized mould stays attached to the surface without obvious damages. Successively, the mould was removed from the support surface by dipping the sample in water and than washing it gently. As expected, underneath the sclerotium we have found the PANI network. The PANI was doped to transfer it into conductive state by acid treatment (see materials and methods), we then contacted the channels of the network by means of micro-manipulating tips connected to a measurement station and verified that the most of channels were rather conductive with a calculated resistance in the range of 24-70 MOhm depending on the tested areas. Figure 3 shows the analysis scheme developed on the typical patterned samples. In particular it is clearly visible the green polyaniline network designed by *Physarum*.

Figure 2 Optical microscope photograph of a network created by *Physarum polycephalum* on PANI sample before (part a) and after (part b) oxygen plasma treatment.

Figure 3: The part above shows an optical microscope photograph of an analyzed sample after Oxygen Plasma treatment and removal of the mold. The 2 black tips are the contact points for electrical characterization and the resulting value is 24 MΩ. In the part below, the spectrum refers to the point “a” signed in the image. Since the picture is a collection of spectra, it is possible to obtain spectra from lines or areas by selecting them directly on the image.

The latter can be considered as a type of lithography. The pattern, created during the slime mold growth can be transferred on the PANI layer, developed by the oxygen plasma and realizing the conductivity patterning after the acid treatment doping. Moreover, micromanipulating tips allow contacting the channels, one at a time, and checking the conductivity in desirable channels. The point marked on the image corresponds to the spectrum below. Indeed, after electrical characterizations, samples were measured with spectrophotometer and data have been elaborated considering pure polyaniline film as a reference.

Conclusions

The study here presented shows interesting results and opens many possible developments. Considering the non-invasiveness of the spectral analysis method (no influence on the conductivity variations), it will be possible to perform measurements in real time, characterizing the network during the motion of Physarum polycephalum. Therefore, we connected the growth of the slime mould with a variation of optical and electrical properties of the sub-layer (Dimonte et al. 2014). Here, putting a step forward, we demonstrated not only that Physarum polycephalum induces redox reactions, but also we exploited Physarum polycephalum’s network as a lithographic mask. Therefore, we found and elaborated a technique by which it is possible to design conductive channels of polyaniline. Moreover, we applied the spectral imaging method for characterizing the patterned polyaniline surfaces formed during the Physarum polycephalum growth. Considering such kind of images, it is possible to appreciate chromatic differences with high spatial resolution between space-separated points. This is an important feature for the characterization of the grown networks of Physarum polycephalum. Therefore, the spectrophotometric scanner is here exploited in a non-conventional context, but it helps us to visualize the interactions between the growing mould and polyaniline layer.

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References
