Structural coloration of chitosan coated cellulose fabrics by electrostatic self-assembled poly (styrene-methyl methacrylate-acrylic acid) photonic crystals

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ARTICLE INFO

Keywords: Photonic crystal Chitosan Textile Cotton Structural color Iridescence

ABSTRACT

The structural coloration of a chitosan-coated woven cotton fabric obtained by glutaraldehyde-stabilized deposition of electrostatic self-assembled monodisperse and spherically uniform (250 nm) poly (styrene-methyl methacrylate-acrylic acid) photonic crystal nanospheres (P(St-MMA-AA)) was investigated. Bright iridescent coatings displaying different colors in function of the viewing angle were obtained. The SEM, diffuse reflectance spectroscopy, TGA, DSC and FTIR analyses confirm the presence of structural color and the glutaraldehyde and chitosan ability to provide durable chemical bonding between cotton fabric and photonic crystal (PCs) coating with the highest degradation temperature and the lowest enthalpy. The coatings are characterized by a mixture of face-centered cubic and hexagonal close-packed arrays alternating random packing regions. For the first time a cost-efficient structural coloration with high washing and light fastness using self-assembled P(St-MMA-AA) photonic crystals was successfully developed onto woven cotton fabric using chitosan and/or glutaraldehyde as stabilizing agent opening new strategies for the development of dye-free coloration of textiles.

1. Introduction

Colors can be produced by different methods like emission, interference, scattering or optical absorption. A pigment mainly relies on the selective absorption of light by chemical chromophores to achieve coloration (Zhao, Xie, Gu, Zhu, & Gu, 2012). Differently, structural colors are originating from micro structural variation at a length scale comparable to the optical wavelength (Saito, 2016). Structural colors are responsible for the most vivid and brightest colors in nature. Numerous examples of structural coloration can be found in insects, birds, mollusks, sea mice, fish, plants and mineral gems (Liu, Shao et al., 2015). They are mainly used by animals and plants for signaling, mimicry and mate (Vignolini et al., 2012). Research currently focus in studying structural color properties in the wavelength ranges of ultraviolet (UV) light, visible light, and infrared (IR) light using both inorganic and organic materials (Diao & Liu, 2012). Structural colors have a multiplicity of potential applications due to their high-resistance to discoloration and inimitability. Moreover, structural colors are a green technology due to the lack of chemical pigments, high efficiency in energy consumption and in the use of light (Saito, 2016; Zhao et al., 2012). Most of the structural colors found in nature are originated from five fundamental optical processes and their combinations: thin-film interference, multilayer interference, diffraction grating effect, photonic crystals, and light scattering (Kinoshita & Yoshioka, 2005). Photonic crystals (PCs) are highly ordered materials that possess a periodically modulated dielectric constant, having the properties of confining and controlling the propagation of light due to the existence of photonic band gap, a band of frequencies where light propagation in the photonic crystal is forbidden (Liu, Shao et al., 2015). They can manipulate pathways of certain light waves with periodic array structures allowing structural color formation. This unique phenomenon is attributed to the light inside the array interacting with the crystal structures (Liu & Huang, 2006). PCs are important for applications in paints, paper, cosmetics, modulators, sensors, filters, laser and light emitting diodes (LEDs) (Jones, Burdett, Ryan, & Walsh, 2016; Sun & Williams, 2012). The most simplest configuration of PCs are monodisperse highly charged micro or nanoparticles colloidal that spontaneously self-assemble into face-centered cubic or body-centered cubic...
crystalline colloidal arrays in low ionic strength aqueous solutions (Xu, Friedman, Humfeld, Majetic, & Asher, 2002). Colloidal particles are of particular interest because of their phononic band gaps and strong interaction with light and have long been used as the major components of industrial products (Zhao, Shang, Cheng, & Gu, 2014). The properties of these monodispersed colloidal spheres can be closely controlled by varying different parameters such as diameter, crystallinity, functional group, bulk substructure, chemical composition, interfacial free energy and surface charge density (Xia, Gates, Yin, & Lu, 2000). Moreover, colloidal phononic structures can be easily fabricated using different methods such as auto-cloning process (Turbierfeld, Campbell, Sharp, Harrison, & Denning, 2000), holographic lithography (Kuramochi et al., 2002), colloidal and block-copolymer self-assembly (Kang, Walsh, Gorishnyy, & Thomas, 2007; Kim, Lim, Lee, Heo, & Yang, 2010). Colloidal self-assembly is the most reliable method due to the low production costs of PCs over large areas, and the wide range of shapes that can be obtained (Gallisteo-López et al., 2011). Electrostatic self-assembly is based on the alternating adsorption of multiply charged cationic and anionic species and it becomes one of the most promising methods for thin-film fabrication (Uğur & Sarsıklı, 2011). Its simplicity and high coating quality combined with the achievable uniform distribution of nanoparticles in wide areas opens broad perspectives for this technique both in textile research and in industry (Ostrander, Mamedov, & Kotov, 2001). Conventional dyeing and printing processes in textile industry face many challenges such as the use of huge amounts of water and chemicals as well as wastewater generation and low color fastness. Structural coloration applications can provide enhanced properties to textile coloration industry since they have special low color fastness. Structural coloration using self-assembled poly (styrene-methyl methacrylate-acrylic acid) (P(St-MMA-AA)) were synthesized by a modified soap-free emulsion polymerization method in a three-necked flask equipped with a reflux condenser and a mechanical stirrer as previously described by Cong and Cao (2003). Briefly, the monodispersed P(St-MMA-AA) colloidal particles were synthesized as follows: 120 mL of aqueous solution (A), containing 0.4 g of Na2S2O3 and 0.8 g of NaHCO3 in a funnel, and 25 mL of monomer mixture (B), consisting of St/MMMA/AA (90:5:5 v/v/v) in another funnel, were added at the same time into a 250 mL three-necked flask. The mixture was stirred at 70 °C in N2 atmosphere for 5 h to obtain a homogeneous dispersion of particles with uniform size distribution.

2. Materials and methods

2.1. Materials

Commercial black dyed cotton fabric with a warp density of 34 threads cm⁻¹, a weft density of 30 threads cm⁻¹ and weight per unit area of 140 g m⁻² was used in this study. The samples were pre-washed with a 1 g L⁻¹ of non-ionic detergent solution at 30 °C for 30 min and then rinsed with water for another 15 min in order to minimize contaminations. Chitosan (DD 85%, ChitoClear hq95-43000, Mw = 350 kDa) was purchased from Primex (Iceland). Styrene (St), methyl methacrylate (MMA), and acrylic acid (AA) were distilled before use. All the other reagents were analytical grade purchased from Sigma–Aldrich, St. Louis, MO, USA and used without further purification.

2.2. Preparation of monodispersed P(St-MMA-AA) nanospheres

Monodispersed latex spheres of poly (styrene-methyl methacrylate-acrylic acid) (P(St-MMA-AA)) were synthesized by a modified soap-free emulsion polymerization method in a three-necked flask equipped with a reflux condenser and a mechanical stirrer as previously described by Cong and Cao (2003). Briefly, the monodispersed P(St-MMA-AA) colloidal particles were synthesized as follows: 120 mL of aqueous solution (A), containing 0.4 g of Na2S2O3 and 0.8 g of NaHCO3 in a funnel, and 25 mL of monomer mixture (B), consisting of St/MMMA/AA (90:5:5 v/v/v) in another funnel, were added at the same time into a 250 mL three-necked flask. The mixture was stirred at 70 °C in N2 atmosphere for 5 h to obtain a homogeneous dispersion of particles with uniform size distribution.

2.3. Cationization process

Cationic cotton fabrics (7 × 7 cm) were prepared by padding chitosan (CH) solutions of 1.0, 0.5 and 0.3 wt% dissolved in 1% acetic acid aqueous solution. The CH solutions were stirred at 300 rpm for 30 min at 70 °C. The heating was kept until the chitosan was completely dissolved. The mixture was stirred until room temperature was reached. The samples were padded in a mini-foulard through the CH solution at 1.5 bar of pressure and 4 rpm (80% pick-up). The excess coating was then removed by gently rinsing with distilled water and the samples dried in an oven for 12 h at 50 °C. Cationic cotton fabrics (7 × 7 cm) used for all the subsequent experiments were prepared only with a chitosan (CH) solutions of 1.0 wt%.

2.4. Coating of cotton fabrics with photonic crystals

Three types of samples were tested: (1) Pristine cotton sample dipped in a 14% photonic colloid solution for 5 min and then dried at 60 °C; (2) Cationized cotton sample treated as sample 1; (3) Sample 2 followed by a second impregnation in a crosslinked chitosan solution and dried at 60 °C. Crosslinked chitosan solution was prepared by dissolving 1 g of chitosan in 100 mL of 1% (v/v) aqueous acetic acid mixed with 50 mL of 1% (m/m) glutaraldehyde solution for 30 min.

2.5. Scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDS)

Morphology analyses of the coatings were carried out with an Ultra-High Resolution Field Emission Gun SEM, NOVA 200 Nano SEM, FEI Company. Secondary electron images were acquired with an acceleration voltage of 5 kV, while backscattering electron images were
obtained with an acceleration voltage of 15 kV. To improve conductivity, the tested surfaces were covered with a film of Au-Pd (80–20 wt%) using a high-resolution sputter coater, 208HR Cressington Company, coupled to a MTT-20 Cressington high resolution thickness controller.

2.6. Fourier transform infrared spectroscopy (FTIR)

A Shimadzu FTIR spectrophotometer (IR-Affinity 1) with an attenuated total reflectance accessory (ATR) was used to record the FTIR spectra of the fabric samples. Spectra were collected in the region of 4000–400 cm\(^{-1}\) and at a resolution of 4 cm\(^{-1}\) with 45 scans at room temperature. A background scan with no samples and no pressure was acquired before the spectra collection.

2.7. Photographs

Optical photos of the fabrics coated with PCs were taken with a Nikon CoolPix4300 digital camera. The pictures were acquired under natural light, at the same time and environmental conditions, perpendicularly to the fabrics and at the distance of 15 cm. In the case of the multi-angle photographs the camera was disposed at different angles (0º, 45º and 90º) using a goniometer maintaining the same distance and using the same sample.

2.8. Thermogravimetric analysis (TGA)

TGA was carried on a Pyris 1 TGA (Perkin Elmer, USA) according to the standard ISO 11358:1997(E). The TGA trace was obtained in the range 40–700 °C under nitrogen atmosphere with a flow rate of 20 mL/min and heating rate of 10 °C/min. Samples were dried at 60 °C for 1 h and placed into a porcelain sample pan before analysis. Graph was plotted with weight (percentage) vs. temperature.

2.9. Differential scanning calorimeter (DSC) analysis

DSC was carried on a Power compensation Diamond DSC (Perkin Elmer, USA) with an Intracooler ILP, based on the standards ISO 11357-1:1997, ISO 11357-2:1999 and ISO 11357-3:1999. Samples were dried at 60 °C for 1 h and placed in an aluminum sample pan before testing. The analysis was carried out in nitrogen atmosphere with a flow rate of 20 mL/min and heating rate of 10 °C/min. The thermogram was obtained in the range of 30 °C–500 °C. Graph was plotted with heat flow vs. temperature.

2.10. Diffuse reflectance spectroscopy

The color of the fabrics was evaluated using a Spectraflash 600 (Datacolor) diffuse reflectance spectrophotometer at standard illuminant D65 (LAV/Spec. Inc., d/8, D65/10°). Five areas on each sample were measured in various positions, and the results represent average values with up to 1% variation. All measurements were performed in triplicate. The responses analyzed were the color characteristics: K/S, L*, a*, b*. K/S is the color strength calculated using Kubelka-Munk’s equation (K/S = (1-R)/2R, where R is the reflectance). L*, a*, and b* are the coordinates of the color in the cylindrical color space, based on the theory that color is perceived by black-white (L*), lightness, red-green (a*), and yellow-blue (b*) sensations. The lightness, L*, represents the darkest black at L* = 0 and the brightest white at L* = 100. The color channels, a* and b*, will represent true neutral gray values at a* = 0 and b* = 0. The red/green opponent colors are represented along the a* axis, with green at negative a* values and red at positive a* values. The yellow/blue opponent colors are represented along the b* axis, with blue at negative b* values and yellow at positive b* values. The results were also summarized by the overall color difference (ΔE*) value using the black dyed cotton as control.

2.11. Light fastness tests

The light fastness was applied in an Atlas XenoTest 150 S chamber with a filtered Xenon lamp with an intensity of 60 Wm-2 according to ISO 4892-2 Method A without water spraying. The light of the Xenon lamp was filtered under λ > 300 nm with an UV window combined with six IR filter glasses. The XenoTest creates an accelerated environment of the natural weathering conditions, simulating materials behavior during its lifetime.

2.12. Washing fastness

The washing fastness was evaluated according to the standard ISO 105 C06, A1S method at a temperature of 40 °C.

3. Results and discussion

In order to determine the most effective chitosan concentration for cotton cationization, three different chitosan solutions (1.0%, 0.5%, 0.3%) were padded onto the dyed cotton before the nanophotonic crystals addition by dipping method. The binding efficiency was monitored using the change in color coordinates (Table 1) according to CIELAB system, where L* characterizes the lightness of a color and can range between 0 (dark) and 100 (light); a* defines a color on a red-green axis; and b* describes the blue part of the color (Okubo, Kanawati, Richards, & Childress, 1998). The color difference (ΔE*) and color coordinate values of the fabrics coated with the nanophotonic crystals are directly proportional to the change in chitosan concentration. Higher is the amount of chitosan, higher are ΔE* and color coordinates values. A chitosan concentration higher than 0.5% does not show significant improvements in color differences. However, the study was conducted at the higher concentration of 1 wt% due to the even coating and highest cationization effect obtained.

Termogravimetry (TGA) analysis provides information about fiber degradation indicating the temperature at which mass loss starts and also gives unique features at the molecular level of the analyzed material. TGA showed that the degradation of untreated and chitosan treated cotton fabric takes place in two well-differentiated steps (Fig. 1). In the first step the polymer showed a small initial weight loss between 35 and 100 °C. This first mass loss is associated with the removal of physically absorbed water and pore liquor that were not removed during the drying process (Kumar, Joo, Choi, Koo, & Chang, 2004). Above this temperature the weight remains constant until a second phase attributed to cellulose decomposition that starts at around 290 °C and continues till 376 °C with a major weight loss of about 70% (Luduena, Fasce, Alvarez, & Stefani, 2011). The highest loss occurs at the maximum decomposition temperature (Tmax) of 334 °C as shown in derivative thermal gravimetric (DTG) peak (Table 2). Slowly, a progressive carbonized structure was formed as the temperature increased further. The high solid residue content (18%) at 700 °C might be due to the presence of impurities such as cations (Na+, K+, Ca2+), which acted as a bridge between cellulose polymer chains, or due to a

Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>K/S</th>
<th>ΔE*</th>
<th>L*</th>
<th>a*</th>
<th>b*</th>
</tr>
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<tbody>
<tr>
<td>CO + CH</td>
<td>584.5</td>
<td>30.3</td>
<td>0.3</td>
<td>0.4</td>
<td>3.3</td>
</tr>
<tr>
<td>CO + CH (0.3%) + pH</td>
<td>214.31</td>
<td>14.4</td>
<td>13.8</td>
<td>1.5</td>
<td>3.6</td>
</tr>
<tr>
<td>CO + CH (0.5%) + pH</td>
<td>141.85</td>
<td>20.5</td>
<td>20.1</td>
<td>1.2</td>
<td>3.6</td>
</tr>
<tr>
<td>CO + CH (1.0%) + pH</td>
<td>132.78</td>
<td>22.2</td>
<td>21.7</td>
<td>1.5</td>
<td>4.8</td>
</tr>
</tbody>
</table>

a: CO: Black dyed cotton; CH: 0.3, 0.5 or 1% Chitosan solution; pH: 14% Photonic colloid.
low carbonization degree (Pooja & Chandra, 2009). No noticeable degradation peaks for chitosan are visible in the Tg spectra of cotton fabric. The introduction of the photonic crystals in the coating show a third mass drop with a DTG peak around 420 °C. This peak is attributed to the degradation of (P(St-MMA-AA)) nanospheres. The absence of Tg features is frequently detected in cellulose due to its rigid polymer backbone having strong inter- and/or intra-molecular hydrogen bonding and to the rigid amorphous phase because of its heterocyclic units (Lee, Kim, & Lee, 2000). The addition of chitosan alone to the fabrics does not show significant difference compared to the control, only a slight increase in the degradation peak temperature and a decrease of the enthalpy due to the lower crystallinity of the coating. However, the addition of the photonic crystal onto the fabric show the appearance of a second endothermic peak at around 430 °C due to the decomposition of the (P(St-MMA-AA)) as confirmed in the TG thermogram. The position of the endothermic peak of the cotton fabric does not change significantly. However, the enthalpy of the cellulose showed a significant reduction after the coating especially in the photonic crystals deposited in the chitosan-modified surface because of its amorphous structure and lowest crystallinity. In particular, the photonic sample pretreated with chitosan and submitted to cross-linking treatment by glutaraldehyde shows the highest degradation temperature (350 °C) and the lowest enthalpy (30 J g⁻¹) suggesting that the cross-linker improves fabric stability and affects the cellulose crystallinity (Park, Choi, & Oh, 2014).

The most important factors to create durable and uniform structural color coatings on textiles are their washing fastness and reflectance properties. The diffuse reflectance spectra of the unwashed and washed cotton fabrics are shown in Fig. 3 and the color data are shown in Table 3. Before washing (Fig. 3a), the uncoated and chitosan coated black cotton almost absorb across the whole visible range (650–400 nm) with an increasing reflectance in the red region (650–700 nm) as expected for this type of dye. The choice of a black dyed fabric as substrate is due to the fact that compared to other base colors the black fabric is able to absorb almost all of incident light, regardless of frequency or angle of incidence. The deposition of the photonic crystal onto the cotton surface increases significantly the surface reflectance (15%) showing a peak in the region between 500 and 600 nm in agreement with its predominant orange color. The other configurations display a similar behavior but with a higher reflectance (20%). The sample coated with chitosan followed by a photonic layer

![Fig. 1. Thermogravimetric (TGA) curve of the cotton fabrics with different coating procedures from 20 °C to 700 °C performed at a heating rate of 10 °C/min under nitrogen atmosphere with a flow rate of 20 mL/min. CO: Black dyed cotton; CH: 1% Chitosan solution; pH : 14% Photonic colloid; CL: 1% cross-linked chitosan solution.](image-url)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Tm (°C)</th>
<th>ΔH (J g⁻¹)</th>
<th>T peaks of 1st derivative (°C)</th>
<th>Residual weight at 700 °C (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>307.1</td>
<td>148.2</td>
<td>333.9</td>
<td>18.1</td>
</tr>
<tr>
<td>CO + CH</td>
<td>312.5</td>
<td>123.5</td>
<td>326.4</td>
<td>19.9</td>
</tr>
<tr>
<td>CO + pH</td>
<td>308.8</td>
<td>72.0</td>
<td>97.5</td>
<td>420.2</td>
</tr>
<tr>
<td>CO + CH + pH</td>
<td>300.8</td>
<td>31.9</td>
<td>126.8</td>
<td>323.4</td>
</tr>
<tr>
<td>CO + pH + CL</td>
<td>399.3</td>
<td>67.6</td>
<td>96.0</td>
<td>343.0</td>
</tr>
<tr>
<td>CO + CH + CL</td>
<td>327.7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO + CH + pH + CL</td>
<td>351.1</td>
<td>30.5</td>
<td>77.8</td>
<td>360.5</td>
</tr>
<tr>
<td>+ CL</td>
<td>434.6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO + CH + pH + + CH</td>
<td>395.1</td>
<td>80.8</td>
<td>59.9</td>
<td>327.1</td>
</tr>
<tr>
<td>+ CH</td>
<td>432.6</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* CO: Black dyed cotton; CH: 1% Chitosan solution; pH : 14% Photonic colloid; CL: 1% cross-linked chitosan solution.

The differential scanning calorimetry (DSC) allows rapid detection and measurement of the physical and chemical transformations of a material when subjected to heating. DSC analysis (Fig. 2) shows one major peak at around 310 °C attributed to the partial pyrolysis with fragmentation of carbonyl and carboxylic bonds from anhydrous glucose units of cellulose producing carbon and/or carbon monoxide (Santos et al., 2014). The enthalpy of this decomposition was 148 J g⁻¹ (Table 2). No crystallization or fusion peaks were observed below the range of thermal degradation as previously observed in others unmodified polysaccharides (Alves, Caridade, Mano, Sousa, & Reis, 2010). The absence of Tg features is frequently detected in cellulose due to its rigid polymer backbone having strong inter- and/or intra-molecular hydrogen bonding and to the rigid amorphous phase because of its heterocyclic units (Lee, Kim, & Lee, 2000). The addition of chitosan alone to the fabrics does not show significant difference compared to the control, only a slight increase in the degradation peak temperature and a decrease of the enthalpy due to the lower crystallinity of the coating. However, the addition of the photonic crystal onto the fabric show the appearance of a second endothermic peak at around 430 °C due to the decomposition of the (P(St-MMA-AA)) as confirmed in the TG thermogram. The position of the endothermic peak of the cotton fabric does not change significantly. However, the enthalpy of the cellulose showed a significant reduction after the coating especially in the photonic crystals deposited in the chitosan-modified surface because of its amorphous structure and lowest crystallinity. In particular, the photonic sample pretreated with chitosan and submitted to cross-linking treatment by glutaraldehyde shows the highest degradation temperature (350 °C) and the lowest enthalpy (30 J g⁻¹) suggesting that the cross-linker improves fabric stability and affects the cellulose crystallinity (Park, Choi, & Oh, 2014).
that was stabilized with glutaraldehyde showed the highest reflectance (30%). A strong reflectance in the photonic band gap indicates a stable colloidal crystals structure onto the black fabric surface. Moreover, the unwanted scattered wavelengths outside the photonic band gap are reduced due to the ability of the black dyed fabric to absorb the transmitted and scattered light outside the photonic band gap enhancing the chroma of structural colors (Liu et al., 2015b). After washing only the sample cross-linked with glutaraldehyde and the sample with an extra layer of chitosan preserve some coating losing in average 5% of the initial reflectance (Fig. 3b). It is clear that glutaraldehyde acting as a bridging agent between its aldehyde groups and the amino groups of chitosan, the acid groups of photonic nanosphere and the hydroxyl groups of cotton. The observed result of the sample with the P(St-MMA-AA) layer imbibed between two layers of chitosan can be explained by the protective action of the outmost layer of chitosan that during washing is removed exposing, without removing, the photonic crystal layer below. The L*, a*, b*, ΔE* and K/S cm\(^{-1}\) values listed in Table 3 show in detail the structural color changes. The reference sample used in this work was the unwashed black dyed cotton (CO, K/S = 697.93, L* = 0, a* = 0, b* = 0). As can be noticed in Table 3 the CO + pH sample after washing is very similar to the control (ΔE* = 0.7, L* = 0.3) due to the very low adhesion of the photonic crystals to the cotton surface. The CO + CH sample is also similar to the control sample but due to the high chemical affinity of chitosan with the cellulose structure some lightness is present on the cotton dyed surface even after washing. The sample pre-impregnated with chitosan and with deposited cross-linked nanophotonic crystals (before and after washing) shows the lowest K/S and the highest L*, a*, b*, and ΔE* values indicating the presence of a high amount of stable photonics absorbed onto the fabric surface. The ΔE* values before and after washing indicate the color changing and can be considered as a direct measure of the degree of stability of the photonic crystals onto the fabric surface providing further evidence that cross-linked photonics deposited onto chitosan offer high washing durability.

The size, shape and distribution of the colloidal nanospheres are of great importance for the resulting optical properties of the photonic crystals, thus a SEM analysis was performed to evaluate the synthesized nanospheres and the coating morphologies (Liu et al., 2016). Mono-dispersed composite latex spheres of P(St-MMA-AA) were successfully synthesized with spherical shape, smooth surface and uniform size distribution by soap-free emulsion polymerization. The P(St-MMA-AA) nanospheres are spherically uniform with an average diameter of 250 ± 10 nm (Fig. 4a). The P(St-MMA-AA) nanospheres deposited on black cotton fabrics show a close-packed array with some cracks usually
in proximity of the fibers interspace (Fig. 4b). Spheres of equal size in order to occupy the minimum volume can display two different packing arrangement: i) face-centered cubic (FCC) or ii) hexagonal close-packed (HPC). The SEM micrographs confirm a regular close-packed sphere arrangement on cotton fabric but they also show a mixture of FCC and HPC regions with some random packing area. Obtain a specific packing type of P(St-MMA-AA) nanospheres is important because it have a direct effect on the photonic properties. Energetically the FCC structure is preferred over the HCP structure, although the energy difference is small (Zhou, Liu, Wu, Fan, & Shao, 2014).

The fabrics with deposited PCs on the untreated cotton and chitosan pre-impregnated cotton before and after washing show similar results, however chitosan provide a smoother surface with less cracks on the PCs coating (Fig. 5a and c). After washing, the PCs are significantly reduced on both fabrics exposing the cotton fiber surface (Fig. 5b and d). Nevertheless, chitosan treated fabric display higher washing durability. Chitosan has active hydroxyl and amino groups on its macromolecule chain that strongly interact with the cotton surface producing cationized cotton able to bound the negatively charged PCs (Chattopadhyay, 2001).

The introduction of the cross-linker to the system clearly improves the deposition of the photonic crystal onto the fabric. The untreated and chitosan treated cotton fabric show similar results due to the glutaraldehyde action that promote the formation of an even surface encaising thick layers of photonic structures (Fig. 6a and c). Sometimes thin layers of photonic structures are also present on film surface. It is important to point out that this superficial film does not interfere with the iridescence of the coated fabric. After washing the sample without chitosan cationization step display delamination of the photonic structures exposing in some extent the dyed cotton surface (Fig. 6b). On the contrary, the cationized cross-linked cotton does not show significant changes after washing confirming that the glutaraldehyde action is not limited to the P(St-MMA-AA) nanospheres but it is extended to the chitosan layer providing enhanced stability and washing fastness (Fig. 6d). Glutaraldehyde and chitosan act as a cross-linked matrix filling up the gaps between P(St-MMA-AA) nanospheres in a complex process forming random N-C\(\text{CH}_2\)-C< and O-C\(\text{CH}_2\)-C< bond types (Kildeeva, Perminov, Vladimirov, Novikov, & Mikhailov, 2009). In this way, the anionic \(-\text{COO}\) groups of the photonic crystal nanospheres by glutaraldehyde action can link to the amine of the chitosan macromolecule chain forming a very stable and resistant coating. The sample without glutaraldehyde but with the additional chitosan surface layer shows a very close packed structure in which the polysaccharide filling the gaps among the P(St-MMA-AA) nanospheres (Fig. 6e). This peculiar configuration display a different opalescence and color appearance due to the new geometry provided by the chitosan surrounding the nanospheres but also due to the change in the lattice structure of the fabric surface induced by the differences in light adsorption between P(St-MMA-AA) and chitosan (Xin-Yuan, 2004). After washing most of the chitosan in removed together with a first layer of nanospheres exposing an untouched second layer of photonic crystals. However, only part of the first layer is removed maintaining a regular coating of photonic crystal with large gaps among nanospheres that alter the opalescence effect without reducing its intensity. This result suggests that the use of a layer-by-layer approach using chitosan for the construction of effective structural color coatings can avoid the utilization of harmful chemicals such as glutaraldehyde.

Fig. 7 shows the photographs under natural light at different angles of a cross-linked cotton fabric coated with the self-assembled colloidal PCs after a washing cycle. Some gaps exposing the black cotton have appeared however the fabric still maintains a reasonable degree of uniformity and iridescence. This mean that an homogeneous

<table>
<thead>
<tr>
<th>Sample</th>
<th>K/S</th>
<th>(\Delta K/S)</th>
<th>(L^*)</th>
<th>(a^*)</th>
<th>(b^*)</th>
<th>K/S</th>
<th>(\Delta K/S)</th>
<th>(L^*)</th>
<th>(a^*)</th>
<th>(b^*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO + CH</td>
<td>584.5</td>
<td>3.4</td>
<td>0.3</td>
<td>0.4</td>
<td>3.3</td>
<td>597.2</td>
<td>2.6</td>
<td>-0.1</td>
<td>0.4</td>
<td>2.6</td>
</tr>
<tr>
<td>CO + pH</td>
<td>107.0</td>
<td>23.6</td>
<td>23.5</td>
<td>0.5</td>
<td>1.9</td>
<td>581.8</td>
<td>0.7</td>
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<td>-0.6</td>
</tr>
<tr>
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<td>29.0</td>
<td>0.9</td>
<td>2.0</td>
<td>121.5</td>
<td>21.0</td>
<td>20.9</td>
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<td>30.0</td>
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<td>5.0</td>
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<td>1.6</td>
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<td>23.4</td>
<td>23.4</td>
<td>1.1</td>
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* CO: Black dyed cotton; CH: 1% Chitosan solution; pH: 14% Photonic colloid; CL: 1% cross-linked chitosan solution.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Unwashed</th>
<th>Washed</th>
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</table>
| Fig. 4. P(St-MMA-AA) nanospheres size (a) and distribution (b) on black cotton fabric.
distribution of nanospheres are maintained allowing large domains of crystalline arrays that are necessary for the occurrence of structural colors in the visible region (Zhou et al., 2015). The fabric displays different structural colors at different viewing angles. It is clear that the orientation-dependent Bragg diffraction change to shorter wavelength increasing the viewing angles from 0° to 90°. The iridescence of structural colors can be also tuned changing the nanospheres size (data not shown). An increase in the PCs sizes result in a red shift of the photonic band gap while a decrease on their size result in a blue shift in the visible spectrum (Liu et al., 2015a). Light fastness test on the fabric shows no significant changes in color compared to a conventional dyed fabric.

The ATR-FTIR analysis (Fig. 8) of the uncoated black cotton control fabric (CO) displays the typical absorption peaks of cellulose structure. The two broad bands of the OH—stretching vibrations arising from hydrogen bonding in cellulose at 3330 and 3270 cm\(^{-1}\) are assigned to intra- and inter-molecular hydrogen bonding of cellulose, respectively (Abidi, Cabrales, & Haigler, 2014). The intensities of methylene peaks at 2920 and 2850 cm\(^{-1}\) are attributed to CH\(_2\) asymmetrical and symmetrical stretching indicating the amount of noncellulosic materials such as proteins, pectin and wax remaining in the fibers after the washing out of the soluble sugars moeites (Souza et al., 2014). The strong peaks at 1150, 1100 and 1020 cm\(^{-1}\) are from the vibrations of the C—O—C bond of the glycosidic bridges of the cellulose structure (Algar et al., 2015). The chitosan-coated fabric (CO + CH) did not show significant differences compared to the control because the characteristic bands of amide I and amide II of the chitosan at 1640 and 1420 cm\(^{-1}\), respectively, are also attributable to the black dye structure (Ibrahim, Osman, & Mahmoud, 2011). Moreover, the peaks between 990 and 1100 cm\(^{-1}\) attributed to the C—O stretching of free and condensed C—OH groups are the same of the cotton structure (Goodarzi, Jafari, Khonakdar, Ghalei, & Mortazavi, 2013). In the photonic crystal coated cotton (CO + pH) the main bands of cellulose disappeared indicating an effective coating of the fabric surface. The bands at 3027, 2922 and 1541 cm\(^{-1}\) can be assigned to sp\(^3\) and sp\(^2\) CH stretching and to the bending of CH\(_2\) groups of the nanophotonic crystals, respectively. The peaks in the region from 1400 to 1490 cm\(^{-1}\) belong to the deformation of CH bonds in the methyl groups of the P(St-MMA-AA) structure (Yap, Que, & Chia, 1991). The absorption band at 750 cm\(^{-1}\) is attributed to the out-of-plane C—H deformation of the monosubstituted or 1,2 disubstituted aromatic rings of the crystals (Costa, Avataneo, Bracco, & Brunella, 2002). The chitosan-pretreated fabric with deposited photonic crystals (CO + CH + pH) and the sample with the additional chitosan layer (CO + CH + pH + CH) show the previously observed peaks of the P(St-MMA-AA) and the bands attributed to the chitosan structure. The peak at 750 cm\(^{-1}\) is more intense result from the aromatic ring out-of-plane C—H deformation and the in-plane NH deformation vibration of chitosan (Oliveira, Zille, & Souto, 2014). The new band around 1590 cm\(^{-1}\) can be assigned to C—N stretching vibration and refers to the amide group because of the NH\(_2\) bending vibration (Souza et al., 2014). The other new peak observed at 1720 cm\(^{-1}\) is attributed to the −COO\(^{-}\) anion of the carbonyl group suggesting the existence of a
polyelectrolyte complex formed by interaction between the chitosan amino groups and the carboxylic group of P(St-MMA-AA) (Davidenko, Peniche, Diaz, San Roman, & Sastre, 2007). The addition of the glutaraldehyde to the system (CO + pH + CL and CO + CH + pH + CL) shows the appearance of two new peaks at 1640 and 1550 cm$^{-1}$ attributable to the carbonyl and ionized carbonyl groups, respectively (Peppas & Wright, 1996). The presence of this two bands and the disappearance of the peaks of the methyl groups from 1400 to 1490 cm$^{-1}$ prove the effective crosslinking of the nanophotonic P(St-MMA-AA) crystal structures. After washing, the chitosan and the photonic coated fabrics show the characteristic spectrum of the untreated black cotton. The CO + CH + pH sample shows a decrease of the peak at 750 cm$^{-1}$ assigned to aromatic group of nanospheres. The cross-linked coatings (CO + pH + CL and CO + CH + pH + CL) do not

![Image](image-url)
show significant differences as well as the sample with the additional chitosan layer (CO + CH + pH + CH). These results confirm the previous observation by SEM that not only the stabilization ability of glutaraldehyde confer washing durability to the photonic crystal coating but also the additional chitosan layer is able to partially stabilize the nanophotonic P(St-MMA-AA) coating.

4. Conclusion

This work demonstrated the structural coloration on woven cotton fabric surface using self-assembled colloidal photonic crystals. The fabric is able to display different iridescence at different viewing angles changing to shorter wavelength as the viewing angle increase from 0° to 90°. Monodisperse and spherically uniform nanospheres of poly (styrene-methyl methacrylate-acrylic acid) with an average diameter of 250 ± 10 nm were successfully synthesized by soap-free emulsion polymerization and efficiently deposited by electrostatic self-assembly technique onto a chitosan-cationized woven cotton fabric. The glutaraldehyde-promoted crosslinking allows durable chemical bonding and higher washing durability between cotton fabric, chitosan and photonic crystals showing the highest degradation temperature (350 °C) and the lowest enthalpy (30 J g⁻¹) due to rearrangement of the crystalline structure of the coated cellulose fabric. The color coordinates, K/S and ΔE* values show the structural color changes among different coating indicating the elevated amount of stable photonics absorbed onto the cross-linked fabric surface and its high washing durability. The SEM micrographs show coatings formed by a mixture of face-centered cubic and hexagonal close-packed regions with some random packing area. FTIR analysis confirms that not only the stabilization ability of glutaraldehyde confers high washing and light fastness but also the additional chitosan layer is able to partially stabilize the nanophotonic coatings. For the first time, structural coloration using self-assembled poly (styrene-methyl methacrylate-acrylic acid) photonic crystals was successfully developed onto woven cotton fabric using chitosan and/or glutaraldehyde as stabilizing agent. This efficient and cost-efficient technique opens new strategies for the development of dye-free coloration of textiles allowing the production of garments for special applications and innovative design with a good degree of washing and light fastness.

Acknowledgments

This work is supported by FEDER funding on the COMPETE program and by national funds through Portuguese Foundation for Science and Technology (FCT) within the scope of the project POCI-01-0145-FEDER-007136 and UID/CTM/00264. A. Zille also acknowledge funding from FCT within the scope of the contract IF/00071/2015.

References


