Evaluation of Aging in Air of Poly (Ethylene Terephthalat) in Oxygen Plasma

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Polyester fabric samples – PET (poly (ethylene terephthalate)), were treated with oxygen plasma, in order to alter the hydrophilicity of such material. The process parameters: working pressure, current, tension and temperature were kept constant, varying only the treatment time. In order to evaluate the change caused on samples hydrophilicity, as well as the influence of the treatment time, the vertical wicking test was used. The samples were stored at standard temperature and pressure conditions, and their wettability were measured one day after the treatment and repeated after 30, 60 and 360 days, in order to check the stability of the effects produced. Scanning electron microscopy (SEM), Raman and ATR (Attenuated total reflection) spectroscopies were used to evaluate physical and chemical alterations on the samples surface. The results have shown a substantial improvement on the hydrophilicity of the treated samples compared to the non-treated sample.

Keywords: aging treatment, low-temperature plasma nitriding, vertical wicking, wettability

1. Introduction

Polymers are materials of large industrial application. However their properties like hydrophilicity; biocompatibility, adhesion, friction and dyeability do not serve, eventually, the industry desires. Therefore, additional surface modification of these materials is needed to alter their properties1-4.

Fabric water absorption capacity is an important characteristic that should be fulfilled by a textile materials used in clothing, which depend mainly on its fiber properties5.

The poly (ethylene terephthalate) (PET) is used by the textile clothing industry due to its versatility and durability, though it is uncomfortable when in contact with the skin mainly due to its low water absorption capacity (0.2 to 0.8%), which is due to its fiber smooth surface and absence of chemical hydrophilic groups, see Scheme 1, like –OH in its molecular structure6,7.

There are several techniques used to modify textile materials surface, and those involve the use of lots of water and chemical reagents, which in its majority are toxic and harmful to the environment. Low energy plasma interaction with the material occurs only on superficial level, so it only modifies a nanometric surface layer of the material and do not modify the bulk propriety of the material7,8. Therefore plasma technique is an important alternative to treat the surface of the textile materials, as it eliminates processing with water and use of chemical reagents9-12.

Low temperature plasma is produced by an electrical discharge in gases at low pressure. It consists of a mixture of highly reactive species like ions, radicals, electrons, and excited molecules, preserving the electrical neutrality. Its chemical composition and physical characteristics are determined by the process parameters, such as: composition of atmosphere, vacuum chamber geometry, gas pressure, gas flow rate and electrical power13,14.

Several phenomena may occur when the active plasma species reaches the material surface under treatment. The plasma process efficiency depends on the gas used, reactor working pressure, temperature and time of treatment, as well as the applied tension and also on the kind of material to be treated9. The main reactions produced by plasma treatment of polymers are surface etching and addition of new chemical groups to the surface. The etching reactions occur due to polymeric chain degradation, thus altering the surface topography. Addiction reactions occur due to recombination of carbon radicals on the polymer surface with the plasma active species, like atoms/ions of oxygen and nitrogen. Plasma of argon, oxygen, nitrogen and ammonia (NH3) are used to modify PET’s surface in order to make it hydrophilic15,16-17.

Others papers, reported that the hydrophilicity reduction of a polymeric material, after plasma treatment, is due to migration of added chemical polar groups from the polymer surface to its interior. This migration is difficult when, during the treatment, cross links are formed between the polymeric chains, resulting that the treated polymer may return to its hydrophobic original characteristic after same elapsed time17-22.
The purpose of this work was to study the increasing of the wettability of (100%) polyester fabrics by plasma treatment and to observe the aging of treated samples during one year. So, it was used 100% O\textsubscript{2} as the precursor gas altering only the time of treatment and all the others parameters like voltage, pressure, flow, current and temperature were kept constant. After treatment the wettability, topography and the chemical composition of the samples surface were analyzed. The wettability was measured through a capillarity test which was developed at the laboratory. The topography was observed through a scanning electron microscopy (SEM) and the chemical composition was studied using Attenuated Total Reflection (ATR-FTIR) Fourier Transform Infrared and Raman spectroscopies.

2. Material and Methods

The substrate used in this work was a 100% polyester (PET) fabric with 160 g/m\textsuperscript{2}, plain geometry and net form. Although clean, the fabric was washed in a bath of water and neutral detergent under agitation, at 100 °C for 10 minutes in order to remove any impurity from the production process and/or storage. Then, the fabric was rinsed to guarantee complete detergent removal, dried at room temperature and finally cut in rectangular shape measuring 25 mm × 200 mm.

The plasma apparatus (presented schematically in Figure 1) used in this work is constituted by a continuous power source of 1kW, connected to the reactor. This source had a maximum voltage of 900V.

The reactor was composed by a vacuum chamber with glass walls, about five liters in volume, and closed by two stainless steel flanges fitted with welded fluid and gas feedthroughs. The superior flange was grounded and contains the gas filling system whose flow was regulated by a flow controller. The inferior flange is free of electrical connections, assuming the floating potential condition during the process, and its central feedthrough is used to place the energized electrode which worked as a cathode. The chamber is evacuated to approximately 7×10\textsuperscript{–3} mbar, using a rotary vane vacuum pump. The chamber pressure was measured by a barometer, type capacitive membrane, and the cathode temperature is measured by a thermocouple of alumel-cromel, fitted inside the electrode. As samples were treated with oxygen plasma treatment rate varying from 10 to 60 minutes at a ratio of 10 minutes. Some process parameters were kept constant in all treatment as pressure 1.25 mbar current of 0.1 A, voltage of 470 V and 90 °C without thermocouple.

In order to avoid samples damage that could be attributed to the thermal energy dissipated by the cathode, the samples were placed at a distance of 70 mm from it, where the temperature was approximately 90 °C.

3. Samples Characterization

The vertical wicking test, Scanning Electron Microscopy – SEM, Raman and Attenuated Total Reflection (ATR-FTIR) Spectroscopy were used to characterization and evaluation of the plasma treatment applied to the samples.

3.1. The vertical wicking test

The vertical wicking test was used to evaluate changes on samples hydrophilicity. It is a simple test of capillarity, on which a sample measuring 25mm × 200mm, has one of its end suspended, while 20 mm on the other end is immersed in a colored solution (Figure 2). A chronometer is used to measure the solution ascending time. The dye used to facilitate the visualization of the ascending liquid is a solution prepared dissolving 1g of reactive yellow dye in one liter of distilled water at room temperature.\textsuperscript{23}
3.2. Scanning Electron Microscopy – SEM

A scanning electron microscope (Phillips model XL-30-ESEM) was used to carry out the analysis to identify morphological alterations caused by the plasma treatment on samples surface. Some fibers were removed from the samples, and then fixed on the equipment stage with a carbon adhesive tape. The fibers were coated by a thin gold film to work as the conducting coating since the samples are dielectric.

3.3. Attenuated Total Reflectance (ATR) and Raman spectroscopy

A non-treated (reference sample) and the plasma treated samples were analyzed using ATR-FTIR and Raman spectroscopies in order to identify any chemical alteration occurred caused by the treatments.

4. Results and Discussions

4.1. The vertical wicking test

Figure 3 presents the results of the vertical wicking test for the treated samples as well as the standard one. The standard sample behaved as expected presenting a constant hydrophilicity, almost equal to zero. The results presented on Figure 3a, were conducted immediately after the samples treatment. It can be observed that in the 30 first seconds, the behavior of the ascending colored solution was nearly the same for all samples, becoming different afterwards. The maximum height of the wicking (19 cm) was reached by the sample treated for 60 min, and the minimum height of the wicking (16 cm) by the samples treated for 10 and 30 min.

According to Rodrigues24, a fabric sample is considered hydrophilic if in the vertical wicking test the solution reaches a minimum height of 5 cm in 5 minutes.23,25,26 Thus, all treated samples presented a substantial improvement on its hydrophilic character, since the minimum height reached was 16 cm. Furthermore, it also became evident that the sample hydrophilic capacity increases with the increasing of the plasma treatment time.

In order to investigate the ageing effect of the applied plasma treatment, and consequently of the hydrophilicity of the samples, these were submitted again to the vertical wicking test after periods of 30, 60 and 360 days of storage. The results of these tests are presented in Figure 3b-d, respectively. As can be seen from these figures, the samples still present a hydrophilic capacity superior to the reference
sample. However, when compared to Figure 3a, it can be observed a drop in the hydrophilic capacity of all aged samples.

The degradation of the effect produced by the plasma treatment as a function of the storage time becomes evident in the analysis of the results presented in Figure 4, where the colored solution maximum height values for each sample are plotted as a function of the time elapsed after the treatment. It can be observed that the ageing effect is accentuated in the 60 first days and then it gradually tends to stabilization. Although a substantial reduction on the solution maximum height occurred, all the treated samples, except that one treated for 10 min, can be still classified as hydrophilic.

4.2. Scanning Electron Microscopy – SEM

Figure 5a-d are SEM micrographs of the fibers removed from the samples untreated and treated by 10, 30 and 60 minutes, respectively. Figure 5a presents the image of a fiber with uniform and smooth surface, as it was expected for the non treated polyester fabric sample. Figure 5b does not reveal any significant physical surface alteration, looking pretty much the same as Figure 5a. On Figure 5c it can be seen a discrete physical surface alteration (marked region). Figure 5d shows that a violent attack occurred on the fiber surface, promoting substantial physical alteration on its surface. There were formations of deep erosions, which can be attributed to the break of molecular chains, due to reactions with plasma excited species on the polymer surface, as well as to the ultraviolet radiation present in the plasma.

![Figure 4. The effect produced by the plasma treatment as a function of the storage time.](image1)

![Figure 5. Fibers microscopic image of samples (a) untreated; (b) treated by 10 minutes; (c) treated by 30 minutes (d) treated by 60 minutes.](image2)
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5. Conclusion

The plasma treatment applied to the samples in this study was proved adequate to enhance the hydrophilicity of PET hydrophobic fabric. Although the results have shown an accentuated reduction of the samples hydrophilicity on the first 60 days, the hydrophilicity became stable afterwards. It became evident that increasing the samples treatment time conducts to more and more intense physical alterations on their surfaces. This gradually provokes surface erosion, increasing the contact area of the samples and improving the hydrophilicity. Although some chemical modifications might occur on samples surface, it was not possible to characterize them using ATR and Raman spectroscopy.

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