Synthesis of Polyphosphate Polymer Employing the Bisphenol (BHBF) and the Dichloride of Phenylphosphonic (PPDC): Evaluation of the Thermal Characteristics

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Abstract

This work presents studies about the synthesis and thermal characterization of a new polymer synthesized from the chloride phenylphosphonic molecule containing. The interfacial polycondensation method was utilized for the synthesis at low temperature. As monomer were used the phenylphosphonic dichloride (PPDC) and the 4,4'-dihydroxybenzophenone (DHBP). The synthesis reactions were conducted at room temperature with heterogeneous medium and constant stirring. The polymer was characterized by FTIR, NMR 31P and MALDI-TOF, which proved the formation of a compound polyphosphonate. DSC and TGA showed that the vitreous transition occurs at temperature of 66.0 °C and that the value of 220.0 °C is the degradation temperature of the polymer. After thermal degradation process the percentage of residual material is around 22.0 wt%. These results suggest the possibility of application of this material as a flame retardant in polymer materials.

Keywords

Synthesis of polyphosphonated polymer, Polyphosphate polymer, Flame-retardants polymers, Thermal properties of polyphosphate polymer and Flame retardant compound

Introduction

Polymeric materials are used in many industrial segments. These materials are easily burned and release toxic compounds during the combustion reactions. Scientific communities have manifested high interest in this area. Due the broad application of these materials in many products, several studies have been realized with polymeric materials containing compounds with flame retardant properties [1-5]. The flame retardant compounds are studied and applied in the polymeric materials to delay its combustion reactions. Normally, these materials are referred as flame retardant additives [1].

Some halogenated molecules have the flame retardant property. These materials are incorporated into polymeric materials for retard the spread of flame. But, the products from these reactions combustion are toxic gases and cause damages to health. Thus, an efficient alternative is the application of compounds containing phosphorus in structures. The chemical groups containing phosphorus can have the flame retardant property and to present low toxicity [2,3,6]. Molecules containing red phosphorus in its structure, inorganic phosphate and organophosphorus generally are good flames retardants. Red phosphorus structures have higher concentration of phosphorus elements, so it can be used in minor amounts compared to the others. However,
present great disadvantage because during the combustion reaction releases the gas phosphine (PH3) that is highly toxic.

Polyphosphates compounds are a special class of the inorganic phosphates. Its performance as flame retardant is based in an effective mechanisms of release of hydroxyl radicals of the type HO• during the combustions reactions. This dehydration reaction favors the formation of a protective layer on the polymeric molecules and acts as a protection barrier to combustion reactions [8,9].

Some works have been realized to synthesize polymers or oligomers containing phosphorus into its structure for these applications. The insertion of flame retardants elements directly in the polymeric chain has been a good method and the results has been good. The phosphorus elements impregnated in the polymer chain inhibits the ignition process and favors the formation of a protective layer between the polymer chain and the oxygen molecules [4]. Also, processes of combustions involving these polymers generates gases with low toxicity when compared with the polymers containing flame retardants additives based in halogens elements [2].

One way of obtaining a phosphorous polymer is by polycondensation synthesis methods that at low temperature the polymeric chains have a good linearity [5]. This is a condensation reaction where two molecules react and generate a complex polymeric molecule and water, for example [10]. Studies show that the flame retardant ability of these polymers depends of the amount of phosphorus groups present in the polymeric chain [5,11,12].

In this context, the present work has the purpose describing the synthesis and characterization of a new phosphorated polymer obtained by phenylphosphonic compound as monomers. The synthetized molecules needs subsequently have detailed studies about their properties as flame retardants, but shows important and interesting results that indicates possible flame retardant ability.

**Experimental**

For the synthesis of the polyphenyl phosphonate containing 4,4'-dihydroxy-benzophenone(DHBP-polyphenyl phosphonate) were utilized the chemical reagents showed on the Table 1.

In first step of the synthesis was prepared a solution of NaOH with 0.70 M in deionized water. In this solution was added a solution of Benzylltriphenyl-phosphonium chloride (BTPPC) with 0.44 M solubilized in Dichloromethane (DCM). The resultant solution was agitated for 40 minutes at room temperature.

In the second steps was added a solution of 4,4'-dihydroxy-benzophenone (DHBP) 0.46 M, solubilized in DCM, and 1.40 g of Phenylphosphonic dichloride (PPDC). The new emulsion was agitated in a jacketed reactor during one hour, at temperature of 24.0 °C.

After these procedures the DHBP-polyphenyl phosphonate was precipitated with hexane. The precipitated compound was dried and purified by solvent evaporation in a vacuum oven during 24 h. All procedures were realized at temperature of 24.0 °C.

DHBP-polyphenyl phosphonate was characterized by Fourier transform infrared spectroscopy (FTIR), Nuclear Magnetic Resonance for the Phosphorus (NMR 31P) and by Matrix-assisted laser desorption/Ionization spectroscopy (MALDI), Differential scanning calorimetric (DSC) and Thermo gravimetric analysis (TGA).

FTIR analyses were realized with a Spectrophotometer Shimadzu (IR Prestige-21) with the transmittance mode and the samples were prepared in circular tablets of KBr. The RMM 31P of solid states was realized with a Spectrometer Bruker (AC200), with the samples solubilized in protonated DMSO and the MALDI-TOF MS spectra were obtained with an AB Sciex4800plus MALDI-TOF/TOF MS apparatus, equipped with Nd:YAG Laser (operated at 100 Hz). The spectra were recorded in the reflection positive ion mode and externally calibrated with “TOF/TOF calibration” standard solution. The acceleration voltage was of 100 mV and the spectra of 400 shots were averaged to improve the signal-to-noise ratio.

TGA analyses were realized with a Systems Shimadzu (TGA-50), at temperature range 25.0 to 900.0 °C and with heating rate of 10.0 °C/min in nitrogenous atmosphere. The thermograms were obtained in a Differential Scanning Calorimeter Perkin Elmer (Jade DSC). For the DSC measurements the samples were prepared with a polymeric mass of 5.0 mg and the temperature range studied were of -20.0 to 230.0 °C. The heating/cooling rate was a 10 °C.min⁻¹ in nitrogenous atmosphere.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Chemical Formula</th>
<th>Physical Aspects</th>
<th>Purity (%)</th>
<th>Molar Mass (g)</th>
<th>Furnisher</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenylphosphonic dichloride</td>
<td>C6H5Cl2OP</td>
<td>Liquid</td>
<td>90.0</td>
<td>194.9</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>4,4'-dihydroxy-benzophenone</td>
<td>(HOC6H4)2CO</td>
<td>DHBP</td>
<td>99.0</td>
<td>214.0</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>Benzyltriphenyl-phosphonium chloride</td>
<td>C25H24ClP</td>
<td>BTPPC</td>
<td>99.0</td>
<td>491.9</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>2,6-Dimethylphenol</td>
<td>C6H10O</td>
<td>DMF</td>
<td>99.0</td>
<td>122.0</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>Hexane</td>
<td>C6H14</td>
<td>C6H14</td>
<td>99.0</td>
<td>86.18</td>
<td>Lafan</td>
</tr>
<tr>
<td>Sodium Hydroxide</td>
<td>NaOH</td>
<td>NaOH</td>
<td>99.0</td>
<td>86.18</td>
<td>Lafan</td>
</tr>
<tr>
<td>Dichloromethane</td>
<td>CH2Cl2</td>
<td>DCM</td>
<td>99.5</td>
<td>84.93</td>
<td>F.MAIA</td>
</tr>
</tbody>
</table>
Results and Discussion

Figure 1 shows the global reactions proposed for the polymerizations of the DHBP-Polyphosphonate polymer. The reaction occurs by interactions between 4,4'-dihydroxybenzophenone (DHBP) and a Phenylphosphonic dichloride (PPDC) molecule, at room temperature (25 °C) for two hours under constant agitation.

![Figure 1: Global reaction proposed for the polymerization of the DHBP-Polyphosphonate.](image1)

Figure 2: Reaction proposed for substitution of the chlorate ion of the BTPPC by the hydroxide ion and the generating of the BTPP-OH.

![Figure 2: Reaction proposed for substitution of the chlorate ion of the BTPPC by the hydroxide ion and the generating of the BTPP-OH.](image2)

Figure 3: Mechanism proposed for the polymerization of the DHBP-Polyphosphonate.

![Figure 3: Mechanism proposed for the polymerization of the DHBP-Polyphosphonate.](image3)
Global reaction proposed is constituted by a series of chemical reactions. In first step occurs the anionic substitution of the chloride ions in the BTPC by the hydroxides ions originated of the dissociation of the NaOH compound, according Figure 2. For that reactions of anionic substitutions occur is necessary that the organic and aqueous phases are miscible. The ions of Benzyltriphenyl-phosphonium chloride are soluble in the organic solvents and not offer restrictions for the anionic substitutions and the BTPP-OH produced during reactions contains the hydroxide ion and is the responsible by the change of the electrical charges across the interface between the organic and inorganic phases [12].

In the second step of the global synthesis the DHBP is solubilized in DCM is added to BTPP-OH. When the ion BTPP-OH is transferred to the organic phase the hydroxide ion keeps the neutrality of the solution. So, the hydroxide ion is transferred from the aqueous phases to the organic phase, promoting the reaction of deprotonating of the DHBP, according reaction proposed in the Figure 3.

In the sequences the PPDC is added in the reaction medium containing DHBP-deprotonated. The PPDC have the P = O chemical bond and characterizes the phosphorus as an electropositive agent and can interact easy with a nucleophilic center. The group P = O have two leaving groups of Cl, that allows a nucleophilic substitution by reaction SN2. So, the ion produced in the protonation process of the DHBP acts as a nucleophile phase and reacts with the PPDC to originate the DHBP-Polyphosphonate. So, a polyphenyl phosphonate is formed by successively reactions of the DHBP deprotonated with the two actives positive center of the PPDC.

A sequence of results proves the formation of a polyphenyl phosphonated compound proposed by this work. The Figure 4 shows the FTIR spectrums in range 500 to 1750 cm\(^{-1}\) obtained for the samples of polyphenyl phosphonate polymer and in the Figure 5 with the range 2000 to 4000 cm\(^{-1}\).

FTIR Spectrums shows transmittance bands typical of a polyphenyl phosphonated compound. These results are strong evidences of the occurrence of the synthesis of a polyphenyl phosphonate polymer, according proposed by global synthesis reaction. But, characteristic bands are observed too due the presence of the residual monomers and of the reagents.

In details it’s possible to observe the presence of the bands in 1157 cm\(^{-1}\), related to the vibrational mode of the bonds type P-O-C aromatic, and the band in 989 cm\(^{-1}\) characteristics of the bonds type O = P-C aromatic [13]. The absence of the peaks in 601 cm\(^{-1}\) and in 524 cm\(^{-1}\) is a strongly evidence of the absence of the P-Cl bonds in molecular structure and the low intensity peak in 3410 cm\(^{-1}\) is related the presence of the hydroxyls of the bisphenol [14].

FTIR band in 1436 cm\(^{-1}\) confirms the presence of bonds type P-C aromatic in the structure of polyphenyl phosphonate [15]. The P = O bonds in the polymer are confirmed by the bands in 1315 cm\(^{-1}\) and 1273 cm\(^{-1}\) [12,13,16]. The bands in 989 cm\(^{-1}\) is related to bonds type O = P-C aromatic of the polyphenyl phosphonate [13].

Presence of others FTIR bands can be associated with others vibrational modes exists in the polyphenyl phosphonate or with the residual molecules present due the incomplete synthesis reactions. For example, the FTIR bands in 1597, 1504, 1116 e 719 cm\(^{-1}\) are associated with C = C aromatic bonds and the bands in 2918 cm\(^{-1}\) and in 848 cm\(^{-1}\) with the H-Caromatic, while the bands in 1647 cm\(^{-1}\) are associated with the C = O bonds, existents in the bisphenol [17].

Figure 6 shows details of the FTIR spectrum in the range of 1500 to 500 cm\(^{-1}\) obtained from the monomer.
Phenylphosphonic Dichloride. It’s possible to observe vibrational modes related to bonds of P-Cl in 601 cm\(^{-1}\) and 524 cm\(^{-1}\). These transmittance bands not exists in the FTIR spectrum obtained for the polyphenyl phosphonate polymer and indicate that the bonds of P-Cl were substituted by the bonds type P-O. Presence of the bonds P = O in the monomer is confirmed by the presence of the bands in 1315 cm\(^{-1}\) and 1273 cm\(^{-1}\) [12,13,16].

Figure 7 shows results of RMN 31P (phosphorus 31) obtained for the samples of polyphenyl phosphonate. The peak in 12.6 ppm is related to the phosphorus elements bonded in the polymer chain and the peak in 12.1 ppm refers to the phosphorus elements exists in the ends of the polymer.
Figure 7: Spectrum of RMN 31P for the DHBP-Polyphosponate.

Figure 8: MALDI spectrum for the DHBP-Polyphosponate polymer.
The endings of the polyphenyl phosphonate chain presents P-Cl or P-OH groups, being the chlorine and the oxygen the electron receptors. The extreme group of the polymer chain presents a RMN peak with smaller magnitude than the phosphonated group of the chain of the polymer [18]. Peaks between 10.0 and 11.0 ppm can be attributed to the residues of the reagents present in the reaction medium after the synthesis process. These evidences are showed too by FTIR results.

So, the RMN analysis is associated with the FTIR results confirms the synthesis of a polyphenyl phosphonated polymer from the PPDC molecules.

Figure 8 shows the MALDI spectrum for the polymer synthesized. The result proves the formation of a DHBP-Polyphenylphosphonate polymer with molecular weight above 2000 atomic mass unit. The spectrum shows patterns for the repetitive units consistent for a polyphenyl phosphonate polymer containing its functional groups of finalization.

Pattern type A is a set of peaks relative to a representative molecular unit present in the structure of the synthesized polymer. The MALDI spectrum shows configurations for these molecules containing amounts above 2000 units of measurement.

Pattern type B and C are the sets of peaks relative to the bonds between fundamental units of the polymer with the phosphonated groups or with a bisphenol group (benzophenone), respectively. The difference between peaks of the pattern A and B is of 140 m/z and between the peaks of the pattern A and C is of 196 m/z and coincides with the molecular weight of these molecules. This information proves that a phosphonated group and a bisphenol groups can bond with the polyphenyl phosphonated molecule in the end of its chain.

Analysis of the MALDI spectrum suggest three possible configurations for the polymers synthesized: i) Polyphenyl phosphonate containing 4,4’-dihydroxy-benzophenone in the chain - Pattern A; ii) Polyphenyl phosphonate with terminations of phosphonated groups - Pattern B and iii) Polyphenyl phosphonate with terminations of benzophenone group - Pattern C. Figure 9 shows these possible molecular formulas.

MALDI results prove the occurrence of the polymerization reactions with the formation of polymeric molecules containing phosphorated structures, in accordance with FTIR and NMR analysis.

Figure 9: Possible configurations for the three types of the polyphenyl Phosphonate polymeric molecules.
Figure 10a: Thermogram obtained by the Thermal Differential Analysis (DSC) for the DHBP-Polyphosphonate polymer.

Figure 10b: Respective first order differential diagram for the signal of the DSC thermogram.
Polyphenyl phosphonate molecules are formed by aromatic functional groups and is expected that its crystalline structures are amorphous. The Figure 10a shows the DSC thermograms obtained for a polymeric sample synthesized and the Figure 10b its respectively first order differential diagram. It’s possible to observe only vitreous transitions for the polyphenyl phosphate molecules that occur between at temperatures 60.0 °C to 70.0 °C. For temperatures above 140.0 °C occurs the thermal decomposition of the molecules with lower molecular weight and of the residual compounds due incomplete synthesis reactions.

DSC thermogram is typical of an amorphous compound and not shows signals of phase transformations. The first order differential diagram shows that the average temperature associated with the vitreous transitions of the polyphenyl phosphonate polymer is 66.0 °C and above of 147.0 °C the reactions of thermal degradations are accentuated.

Figure 11 shows the Thermal analyses (TGA) for the polyphenyl phosphonate. The results indicate the mass loss in function of the temperature. The mass loss at temperature of 147.0 °C is due the decomposition of the residual compounds from the incomplete synthesis reaction and confirms the results of the Differential Thermal Analysis. The greatest mass loss is observed above the temperature 220.0 °C that indicates the start of the thermal decomposition of the phosphorated polymer.

Thermograms show too the presence of a significant quantities of residual mass after thermal decomposition of the phosphorated polymer. After heating until 700.0 °C the residual mass is approximately of 24.0 wt%. This characteristic is very important if the future application of this polymer is as flame retardant additive in polymeric matrices. In these applications the residues after thermal decomposition can favor an intumescent effect in the polymeric matrix and generate a physical barrier for the oxygen into the polymeric matrices. This process inhibits with good efficiency the combustions processes of the polymers.

Phosphorated compounds are applied as flame retardant by having phosphorus elements in its structure [7]. The phosphorus elements favor the formation of an inhibitors layer on the surface of the polymer matrices during the combustions processes. These mechanisms decreases the heat generation during the combustion and reduces the contact area between polymeric molecules and the oxidizing agents, particularly with the oxygen. The predominant mechanisms in this processes depends of the characteristics of the polymer matrix and of the structure of the phosphorated compound applied [20].

Figure 11: Thermogram obtained by the Thermogravimetry Analysis (TGA) for the polyphenyl Phosphonate in nitrogen atmosphere and heating rate of 10.0 °C/min.
In general, the mass of the residues generated after the thermal decomposition of the polyphenyl phosphonate compounds are directly related by the amounts of phosphorus elements in the polymeric chain [21]. During the combustion reactions occurs the formation of a carbonaceous structure, that is an inhibitors layer (solid chair) originated by the pyrolysis reaction. The decomposition of the polyphosphate polymer releases radicals of PO•, P• and P2• that have the ability of capturer radicals of H•, •O• and HO•. These phenomena allow the formation of high amounts of the residual mass after the thermal decomposition, at high temperatures even in inert atmosphere [22].

Table 2: Relationship of phosphorus quantities in the structure of the DHBP-Polyphosphonate and in others phosphorated compounds and the amount of residual mass generated after their thermal decomposition.

<table>
<thead>
<tr>
<th>Molecular structure</th>
<th>P (wt%)</th>
<th>T (°C)</th>
<th>Residual Mass (%)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="Molecular structure 1" /></td>
<td>8.8</td>
<td>700</td>
<td>24.2</td>
<td>Polyphenyl phosphonate</td>
</tr>
<tr>
<td><img src="image2" alt="Molecular structure 2" /></td>
<td>6.4</td>
<td>600</td>
<td>27.0</td>
<td>[5]</td>
</tr>
<tr>
<td><img src="image3" alt="Molecular structure 3" /></td>
<td>6.6</td>
<td>700</td>
<td>20.0-30.0</td>
<td>[19]</td>
</tr>
<tr>
<td><img src="image4" alt="Molecular structure 4" /></td>
<td>8.9</td>
<td>800</td>
<td>52.0</td>
<td>[22-23]</td>
</tr>
<tr>
<td><img src="image5" alt="Molecular structure 5" /></td>
<td>8.9</td>
<td>800</td>
<td>23.0</td>
<td>[12]</td>
</tr>
<tr>
<td><img src="image6" alt="Molecular structure 6" /></td>
<td>5.2</td>
<td>850</td>
<td>17.2</td>
<td>[13]</td>
</tr>
<tr>
<td><img src="image7" alt="Molecular structure 7" /></td>
<td>13.9</td>
<td>700</td>
<td>40.9</td>
<td>[24]</td>
</tr>
</tbody>
</table>
Mass loss is significant after thermal decomposition of the polymer synthesized and the quantities is similar to others flame retardants applied in polymers. The Table 2 relate the results obtained to polyphenyl phosphonate synthesized in this work with others phosphorated materials obtained by others scientific works [23,24].

Conclusions

Results showed that by the interfacial polycondensation synthesis methods and utilizing a phenylphosphonic as monomer is possible to obtain a polymer containing phosphorous in its principal chain and with molecular weight higher than 2000 units. FTIR, RMN 31P and MALDI spectrocopies proved the formation of the DHBP-Polyphosphonate polymer employing the Bisphenol (BHBF) and the Dichlorate of phenylphosphonic (PPDC).

Thermal analysis showed that the DHBP-Polyphosponate has good thermal stability until temperatures of 220.0 °C and after its thermal degradation the residual mass is approximately of 24.0%. These results potentiates the application of the DHBP-Polyphosponate for the future as good flame retardant additive for the polymer materials.

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References


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