Flame Retardant Finish of Silk Fabric with Dimethyl Phosphonate Doped Silica Sol

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Abstract. In this paper, a dimethyl phosphonate doped (P-doped) silica sol was prepared from a tetraethyl silicate inorganic precursor and applied to silk fabric as a flame retardant finish through the sol-gel process. The X-ray photoelectron spectroscopy (XPS) analysis and scanning electron microscope (SEM) images indicated a phosphorus-containing silica network successfully formed on the silk fabric surface. The flammability of the treated silk fabric was investigated by the limiting oxygen index (LOI) and vertical combustion test. The results indicated that the treated silk fabric were flame retardant with a higher limiting oxygen index (LOI) and shorter damage length. Moreover, micro calorimeter combustion testing and thermal gravimetric analysis indicate 50% of P-doped silica sol treated silk fabric exhibited a lower heat release rate and higher char residue, which demonstrates a synergistic effect between the silica and phosphorus.

Introduction

The flame retardance of a textile is rather important for both researchers and consumers because textiles’ igniting during a fire can cause damage to property and life. According to Great Britain fire statistics from April 2012 to March 2013, 76% of fire-related fatalities occurred in dwelling fires, and textiles along with upholstery and furnishings accounted for nearly 60% of deaths[1]. Indeed, studying textile flame retardancy began very early, and most currently used flame retardant chemicals were developed during from 1950-1980. However, little work has occurred in the last decade due to increasing environmental sustainability, chemical toxicological acceptability, performance and cost concerns[2].

The sol-gel process, an emerging material surface modification technology, is mainly based on the hydrolysis and condensation of semi-metal alkoxides. The sol-gel method has been widely used to create functionalized textiles[3], such as antimicrobial, water repellent and super hydrophobic textiles because it yields new materials with a high degree of molecular homogeneity and potentially extraordinary physical and chemical properties[4]. In recent years, the sol-gel method has been increasingly used and reportedly exhibits great potential for flame retardancy[5].

The high nitrogen and sulfur content of silk fabric makes it less flammable than other fibers. However, it still cannot satisfy the strict demand for flame-retardant textiles. In this paper, silk fabric was treated with a P-doped silica sol using dimethyl phosphonate as a flame retardant additive via the sol-gel method. Flammability, combustion performance under thermal radiation and thermo stability tests were performed to comprehensively characterize the flame retardant properties. The surface morphologies and elemental composition for the finished fabrics were also presented.
Experimental

Material and Reagents

Silk fabric (weight: 36 g/m², density (ends/cm): 50×45 (warp×weft)) was supplied by Suzhou Huasi Silk Printing & Dyeing Co., LTD (Suzhou, China). Tetraethyl silicate (TEOS), the inorganic precursor, and dimethyl phosphonate (DP), the flame retardant additive, were both purchased from Sinopharm Chemical Regent Co., Ltd. Ethanol (EtOH) and hydrochloric acid (HCl) were purchased from China Sun Specialty Products Co., Ltd. All reagents were analytically pure.

Preparation of Flame Retardant Sol

The flame retardant sol was synthesized via the sol-gel method. The pure silica sol was prepared as follows: 3.6 ml distilled water was acidified with 1.4 ml hydrochloric acid and added dropwise to a 44.7 ml tetraethyl silicate and 35.0 ml ethanol solution while stirring at room temperature. This solution was stirred for 3 hours at 70°C and aged for 2 days. The molar ratio for the reagents was 1:3:1:0.085 for TEOS: EtOH: H₂O: HCl. For the P-doped silica sol, dimethyl phosphonate, used as a flame retardant, was added to the tetraethyl silicate before the hydrolysis. To investigate the synergistic effect between tetraethyl silicate and dimethyl phosphonate, different mole dimethyl phosphonate fractions (namely, 0, 10, 20, 30, 40 and 50% with respect to the sum of TEOS and DP) were employed.

Modification of Silk Fabric with Prepared Sols

According to the padding & curing modification process, the silk fabric was first impregnated with the prepared sols and squeezed for a 100% pickup via a padder with a 0.4 kg/cm² nip pressure. The samples were then dried at 80°C for 3 min and immediately cured at 120°C for 6 min. All samples were conditioned under standard atmospheric conditions (25± 2°C; 65% humidity) for 48 h before testing.

Characterization and Measurements

Morphology of the samples was observed by a Hitachi TM3030 Desktop Scanning Electron Microscope (SEM) at an acceleration voltage of 3kV under vacuum condition. The elemental binding state was measured via X-ray photoelectron spectroscopy (XPS, Axis Ultra HAS, Kratos). The step size for the high-resolution scan was 0.1eV, and the pressure in the analysis chamber was maintained at 4.0×10⁻⁹pa.

The limiting oxygen index (LOI) was measured by LCK-09 Oxygen index measuring instrument according to ASTM D2863. The vertical combustion test was measured by LYF-26 vertical combustion instrument, and the fabric bottom (300mm×80mm) was exposed to a 40 ± 2mm high flame for 12s to calculate the damage length, repeating 4times for each sample. Micro calorimeter combustion (MCC) instrument was used to test the sample combustion properties. The samples (~5mg) were heated from room temperature to 700°C under nitrogen flow at 80cm³/min with a linear heating rate of 1°C. The gaseous pyrolysate mixture in the pyrolyser was mixed with a 20cm³/min stream of oxygen before combusting in a 900°C furnace for 10s. The thermo stability the silk fabrics were performed with a 2960 SDT 290 TA instruments. The temperature was set from 50°C to 700°C at a rate of 10°C/min and nitrogen atmosphere was selected.

In addition, the add-on values of the samples (Table 1) were calculated according to equation 1:

\[
\text{Add-on} = \frac{w_2 - w_1}{w_1} \times 100\%
\] (1)
where $w_1$ and $w_2$ is the weight of silk fabric before and after modification, respectively. The treated fabrics were washed for 60min using a 5g/l soap-flake at 60°C before testing to understand the modified silk fabric durability.

**Results and Discussions**

**Surface Morphology and Chemical Composition Characterization**

Figure 1 shows the surface morphology of the treated (pure and 50% P-doped) and control silk fabrics. It can be seen from the figure that the control sample surface was clean and smooth with clear boundaries between each filament (a), while the treated fabric surface were fuzzy and covered by the coatings (b and c). The char residue after heating the treated samples (e and f) to 700°C retained a complete and stable structure in contrast to the control (d). This result indicates the silicon network successfully formed on the silk fabric surface after the treatment and could guarantee the structural integrality at high temperatures.

![Fig.1 SEM Images of Silk Fabric for Control (a), Pure Silica Sol Treated (b) and 50% P-doped Silica Sol Treated Sample (c) and Char Residue for the Control (d), Pure Silica Sol Treated (e) and 50% P-doped Silica Sol Treated Sample (f), respectively](image)

The surface composition for the samples were investigated by XPS. Figure 2 shows a typical survey for the three samples. The distinctive silica peak was present for the treated silk fabrics (b and c) but not the control (a). In particular, according to the literature [6,7], the Si 2p peaks from the -Si-O-Si- chemical bond appeared with a binding energy of 98 eV for pure silica sol treated silk fabric and 100eV for the 50% P-doped silica sol treated silk fabric. The SiO$_2$ films formed on the surface of the treated silk fabrics via the sol-gel method. The Si 2s peaks from the -Si-O-C-chemical bond exhibited a binding energy of 149eV for the pure silica sol treated silk fabric and 151eV for the P-doped silica sol treated silk fabric, which confirmed the silica films were grafted to the sample surface. Furthermore, the P 2p peak with a binding energy of 130eV for the 50% P-doped silica sol treated silk fabric indicated phosphorus was deposited on the silk fabric via silica sol doping [7].
Flammability Test

The LOI and vertical combustion experimental results are shown in Table 1.

Tab.1 Flammability Test for Treated and Untreated Silk Fabric

<table>
<thead>
<tr>
<th>Code</th>
<th>Concentration of dimethyl phosphonate (%)</th>
<th>Add-on (%)</th>
<th>LOI (%)</th>
<th>Damage length (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Before washing</td>
<td>After washing</td>
</tr>
<tr>
<td>CONTROL</td>
<td>---</td>
<td>---</td>
<td>24.2</td>
<td>24.2</td>
</tr>
<tr>
<td>TEOS</td>
<td>0</td>
<td>23.17</td>
<td>26.5</td>
<td>26.3</td>
</tr>
<tr>
<td>TEOS 10P</td>
<td>10</td>
<td>22.56</td>
<td>26.9</td>
<td>26.3</td>
</tr>
<tr>
<td>TEOS 20P</td>
<td>20</td>
<td>26.74</td>
<td>27.4</td>
<td>26.9</td>
</tr>
<tr>
<td>TEOS 30P</td>
<td>30</td>
<td>24.41</td>
<td>27.4</td>
<td>26.5</td>
</tr>
<tr>
<td>TEOS 40P</td>
<td>40</td>
<td>26.07</td>
<td>28.2</td>
<td>26.8</td>
</tr>
<tr>
<td>TEOS 50P</td>
<td>50</td>
<td>30.02</td>
<td>33.1</td>
<td>27.7</td>
</tr>
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</table>

The LOI values were higher for the treated silk fabrics than the control sample, which means the flame retardance of silk fabric was improved after treating with the prepared sols. More specifically, the LOI value for the pure silica sol treated silk fabric was 26.5%, a 9.5% increase over the control sample (24.2%). The LOI value for the P-doped silica sol treated silk fabric increased relatively slowly for a P concentration below 30%. However, increasing the dimethyl phosphonate concentration to 50% gave the treated silk fabric an optimal flame retardance with a 33.1% LOI because both silica and phosphorus are flame retardant elements. The sol-gel treatment forms a nonflammable silica network on the treated silk fabric surface, which prevented the flames from spreading. The dimethyl phosphonate, a phosphorus-based flame retardant, exhibits a significant flame retardant effect on the fabric by generating phosphorous acid that dilutes the flammable gas from combustion and improves the flame retardance of the treated fabric. Notably, the LOI values for all of the silk fabrics treated with a P-doped silica sol remained above 26.0% after home laundering once, which indicates the flame retardance has a certain durability. Furthermore, the damage length was much shorter for all of the treated silk fabrics than the control sample, which indicates reduced vertical combustion.

Combustion Performance

Table 2 shows the heat release data for the silica sol treated and control silk fabrics during MCC test and Figure 3 shows the heat release curves versus temperature for the test samples.
Table 2 shows the peak heat release rate (pHRR) decreased with increasing P concentration in the prepared sol. Compared to the pure silica sol treated silk fabric, the pHRR for the silk fabric treated using a silica sol with 50% P content decreased from 111.9 to 71.5 W/g. However, compared to the control sample, the pHRR of the silica sol treated silk fabric did not decrease using pure silica or a lower P content (below 20%). The same pHRR trend was observed for the heat release capacity (HRC) value, which was obtained by dividing the maximum specific heat release rate (SHRR) by the heating rate for the test \(^8\). The silica layer could only act as heat insulation on the treated fabric surface and prevent heat release to some extent because a low P content only weakly prevents the thermal degradation of the sample. When the heat under the insulation layer reached a limit, all of the accumulated heat broke through the layer and was released, which makes the maximum heat release rate and heat release capacity larger than the control sample. However, when the P content increased from 20% to 50%, the thermal degradation of the silica sol treated silk fabric changed significantly, which made the heat insulation from the silica layer stable enough to endure the heat release, which decreased the pHRR and HRC. As depicted in Figure 6, the treated (pure and 50% P-doped) silk fabric began decomposing at 250°C (b) and 230°C (c), which is higher than for the control sample, 200°C (a), as shown by the rapid increase in HRR and dramatic decomposition at lower temperatures. Furthermore, the treated silk fabric exhibited a lower total heat release (THR) and higher percent char residue than the control sample. These results indicated improved combustion properties for the treated silk.

**Thermal Stability**

The TG curves for the treated (pure and 50% P-doped) and control silk fabrics are shown in Figure 4.

Tab.2 MCC Data for Treated and Untreated Silk Fabric

<table>
<thead>
<tr>
<th>Sample</th>
<th>HRC (J/g*K)</th>
<th>pHRR (W/g)</th>
<th>THR (kJ/g)</th>
<th>Tmax (°C)</th>
<th>Char residue (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CONTROL</td>
<td>109</td>
<td>97.3</td>
<td>12.3</td>
<td>296.4</td>
<td>20.60</td>
</tr>
<tr>
<td>TEOS</td>
<td>122</td>
<td>111.9</td>
<td>10.5</td>
<td>313.7</td>
<td>33.96</td>
</tr>
<tr>
<td>TEOS 10P</td>
<td>115</td>
<td>104.6</td>
<td>8.7</td>
<td>309.7</td>
<td>32.20</td>
</tr>
<tr>
<td>TEOS 20P</td>
<td>103</td>
<td>93.8</td>
<td>7.6</td>
<td>306.5</td>
<td>33.60</td>
</tr>
<tr>
<td>TEOS 30P</td>
<td>101</td>
<td>92.7</td>
<td>7.4</td>
<td>305.2</td>
<td>34.93</td>
</tr>
<tr>
<td>TEOS 40P</td>
<td>99</td>
<td>90.8</td>
<td>7.5</td>
<td>304.2</td>
<td>36.40</td>
</tr>
<tr>
<td>TEOS 50P</td>
<td>78</td>
<td>71.5</td>
<td>6.3</td>
<td>293.5</td>
<td>45.60</td>
</tr>
</tbody>
</table>

Fig.3 HRR Curves of (a) Control (b) Pure Silica Sol Treated and (c) 50% P-doped Silica Sol Treated Sample
Fig. 4 TG Curves of (a) Control (b) Pure Silica Sol Treated and (c) 50% P-doped Silica Sol Treated Sample

The three curves exhibited similar weight loss behavior below 275°C due to water loss. The silk fabric began to seriously degrade with a major weight loss from 280°C to 500°C. Notably, the temperature for the treated silk fabric for any given weight was slightly higher than for the control sample. For example, when the weight residue decreased to 81.6%, the temperature of the treated silk fabric was nearly 314°C while the control sample was 303°C. This result indicated the thermal-degradation temperature was improved for the treated silk fabric. Additionally, increasing the temperature stabilized the weight loss rate for the silk fabric and a steady char residue was obtained at nearly 700°C. As shown in the figure, the char residue weight was much lower for the control than the treated samples, i.e., 33.30% for the control sample and 45.10% and 49.65% for the pure and 50% P doped silica sol treated silk fabrics, respectively. Indeed, according to the report, individual phosphorus flame retardants could not improve the thermal stability. Phosphorus flame retardants act as a condensed form of phosphoric acid and polyphosphoric acid during thermal decomposition and cannot gain char residue but only dehydrates carbon-containing materials, which accelerates oxidation[9]. Therefore, the increased char residue for the 50% P-doped sample could evidence a synergistic effect between the silica and phosphorus, that is, the phosphorus-containing component could promote the material dehydrating to carbon at high temperatures, and the silica layer could then improve the produced carbon stability.

Summary

A nano-scaled silica sol system was prepared and applied as a flame retardant finish on silk fabric. The results indicated the flame retardance of the silk fabric was improved after treating with pure and dimethyl phosphonate doped silica sols. The treated silk fabric reached the optimal flame retardant effect (33.1% of the LOI compared to 24.2% for the control sample) and superior thermal stability when treated with a 50% P doped silica sol.

Acknowledgement

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References


