Hydrothermal Synthesis of Zirconia/Silica Hybrid Materials and their Application on Cotton Fabrics

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Abstract — A series of novel zirconia/silica hybrid materials were prepared via the hydrothermal process from zirconium n-propoxide (ZNP), tetraethoxysilane (TEOS), and n-octyltrietoxysilane (OTES). The structures of the zirconia/silica hybrid materials were characterized using Fourier transform infrared (FTIR) analysis and $^{29}$Si nuclear magnetic resonance (NMR) imaging. Grey cotton fabrics were first dyed and then coated with the zirconia/silica hybrid materials through a padding–drying–thermosol process. The bonding structures of the treated cotton fabrics were evaluated using scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDS) analyses. The evenness of the coated layer on the treated cotton fabrics was confirmed by SEM images, and the interaction of the hybrid materials with the treated cotton fabrics was verified. Moreover, water repellency and warmth retention analyses of the treated fabrics were conducted. Our data show that the warmth retention of the cotton fabrics treated with the zirconia/silica hybrid materials was improved with increasing concentrations of zirconium n-propoxide in the reaction mixture. After modification with zirconia/silica hybrid materials, all the treated cotton fabrics revealed good water repellency abilities.

Keywords — zirconia, silicacotton fabrics, hydrothermal, warmth retention, water repellency.

I. INTRODUCTION

Most new products are designed to have multiple functions and a form of intelligence. Thus, hybrid materials have recently become one of the main trends in materials science research. Compared to traditional materials, the most important feature of a hybrid material is its designability. The mechanical, physical, and chemical properties of a hybrid material can be designed and controlled by changing its composition or using interface control, compounding technology, or molding technology to meet the requirements of maximum usability and environmental compatibility [1–3]. Organic–inorganic hybrid materials contain both organic and inorganic functionalities and provide functional compensation and optimization [4–8]. Consequently, they are widely used in numerous fields including optics, catalysis, and in other biomaterials.

Zirconia (ZrO$_2$) is an interesting oxide material that has been developed recently. Zirconia offers ablation resistance, chemical stability, and a high refractive index; it also possesses good dielectric, insulating, and mechanical properties [9–12]. It is applied in thermal-barrier-, bioactive-, ultrafiltration-, and corrosion-protection coatings [13–17], as well as during the manufacture of photoelectric components and electrochemical biosensors [18–21]. It is frequently used in various industries, including aviation, aerospace, ferrous metallurgy, machine manufacture, and optics. In addition, it offers strong potential for other applications. There are many methods used to prepare the coating material; these methods can be classified into two major types, namely physical processes, such as vapor-deposition and sputtering methods, and chemical processes, such as chemical-vapor deposition, spray pyrolysis, and sol-gel methods. The sol–gel method is considered extremely practical for the fabrication of mesoporous ceramic membranes [22, 23].

Furthermore, silica is a natural material derived from common materials such as quartz, sand, and flint. It has high chemical stability, a low thermal expansion coefficient, and high heat-resistance characteristics. The relatively high chemical stability of the silica phase can be advantageous in some cases [24–27]. Khatib’s use of silica membranes for a hydrogen separation application reveals that silica-based membranes have emerged as promising materials at high temperatures owing to their high permeation rates, high selectivity, hydrothermal stability, resistance to poisons, and mechanical strength [28].

Hydrothermal method provides advantages such as direct crystallization under hydrothermal conditions without sintering, regular morphology, uniform grain size, lower agglomeration, etc., it is widely used for the preparation of nanoparticles. It is also used in the preparation of hybrid materials that possess both the flexible performance of organic materials and the anti-friction, resistance to aging, and weathering characteristics of inorganic materials [29–32]. The
modification of inorganic–organic materials greatly improves the surface properties of the organic material and expands its application [33-36].

This study adopted the hydrothermal method to synthesize a series of composite materials of zirconium n-propoxide and tetraethoxysilane (TEOS) and n-octytriethoxysilane (OTES) as chemical precursors. Various molar ratios of the zirconia/silica hybrid materials were used to generate the network structures, and the chemical and physical properties of the treated cotton fabrics were evaluated. Water repellence and warmth retention were analyzed to evaluate the physical properties of the treated cotton fabrics and to identify multifunctional benefits.

II. EXPERIMENT

A. Materials

Milled, scoured, and bleached plain weave Cotton fabrics [ends (100)*picks (56)/(32s/1)*(32s/1)] were supplied by Everest Textile Industry Co., Ltd., Tainan, Taiwan. Tetraethoxysilane, zirconium n-propoxide, and OTES were respectively prepared with 50 mL of deionized water and 5 mL of 99% ethanol. To disaggregate the agglomerated particles into well-dispersed colloidal particles, Fumed zirconia was hydrolyzed in water using nitric acid, and the mixture was then stirred until it became clear. Then, constant ratios of TEOS and OTES were used to make clear. The pH value was adjusted to approximately 3 using nitric acid, and the mixture was then stirred until it became clear. Then, constant ratios of TEOS and OTES were respectively prepared with 50 mL of deionized water and 5 mL of 99% ethanol. To

c. Treating Cotton fabrics

The treatment of the cotton fabrics was performed using the “two dips, two nips” padding method with a pickup of 80%. A were dipped for 3 min in a so solution containing the required weight percent of hybrid materials. A padding–drying–curing finishing procedure was used to disaggregate the agglomerated particles into well-dispersed colloidal particles. Fumed zirconia and silica sol-treated fabric samples were padded and nipped to remove excess liquid and to obtain a percent wet pickup of 80% using a paddler (Rapid Labortex Co., Ltd., Taoyuan, Taiwan) with a set nipping pressure. The treated fabric was pre-dried in an oven at 70°C for 5 min. Then, the treated fabrics were cured at 100°C for 90 s in a preheated curing oven (R3, Chang Yang Machinery Co., Taiwan). The treated fabrics were rinsed with water several times to thoroughly remove any hybrid material residue and dried at 70°C for 5 min prior to analysis. The labels CT1–CT4 and CZ1–CZ4 denote the dyed cotton fabrics treated with hybrid materials T1–T4 and Z1–Z4, respectively. Figure 1 shows a schematic describing the interaction between the hybrid materials and fabric.
III. RESULTS AND DISCUSSION

A. FTIR analysis

In the IR spectra of the Z<sub>i</sub>-Z<sub>i</sub> and T<sub>i</sub>-T<sub>i</sub> hybrid materials, a strong absorption band in the range from 3456 to 3444 cm<sup>-1</sup> and 3452 to 3441 cm<sup>-1</sup>, respectively, showed that the hybrid materials had N–H groups. Absorption peaks at approximately 451–447 cm<sup>-1</sup> and 459–455 cm<sup>-1</sup>, respectively, revealed that the hybrid materials had Zr–O groups, which enabled the formation of Zr–O–Zr networks. A second typical absorption region for the Si–O stretching vibration of Z<sub>i</sub>–Z<sub>i</sub> and T<sub>i</sub>–T<sub>i</sub> at 1072–1068 cm<sup>-1</sup> and 1072–1070 cm<sup>-1</sup>, respectively, has been reported. In the absorption region at 1188–1184 cm<sup>-1</sup>, a band due to Si–C bond stretching was found, and the absorption region at 798–794 cm<sup>-1</sup> was due to deformation of the Si–C bond. Asymmetric stretching vibrations of the C–H bonds in Z<sub>i</sub>–Z<sub>i</sub> and T<sub>i</sub>–T<sub>i</sub> were found from 2927 to 2925 cm<sup>-1</sup> and from 2928 to 2927 cm<sup>-1</sup>, respectively. Symmetric stretching vibrations of the C–H bonds of Z<sub>i</sub>–Z<sub>i</sub> and T<sub>i</sub>–T<sub>i</sub> were found from 2858 to 2854 cm<sup>-1</sup> and from 2858 to 2856 cm<sup>-1</sup>, respectively. Thus, it is reasonable to assume that reactions occurred between some of the zirconia and silica. Figure 2 shows the FTIR spectra of Z<sub>i</sub>–Z<sub>i</sub>. Increasing the zirconia concentration increased the strength of the Zr–O absorption and also strengthened the bonding. Figure 3 shows the FTIR spectra of T<sub>i</sub>–T<sub>i</sub>. Increasing the OTES concentration increased the strength of the Si–O absorption and strengthened the bonding. The structures of Si–O–Si and Zr–O–Zr were also analyzed using <sup>29</sup>Si NMR.

B. <sup>29</sup>Si NMR spectra analysis

<sup>29</sup>Si NMR was used to observe the structure formed by the hydrolysis of Si. While the FTIR results indicate the formation of Si–O–Si by a hydrothermal reaction, <sup>29</sup>Si solid-state NMR provides data on the structure of the silica and the extent of the Si–OH condensation reaction. The <sup>29</sup>Si NMR spectra of T<sub>i</sub>–T<sub>i</sub>, shown in Figure 4, indicated that the increase in the molar ratio of OTES under constant molar amounts of ZNP and TEOS made the hybrid materials react more completely, so that the absorption peaks of hybrid materials tend to be smooth, and so the main absorption peaks of the hybrid materials T<sub>i</sub>–T<sub>i</sub> tend to be strong. The high-resolution solid-state NMR spectra of T<sub>i</sub>–T<sub>i</sub>, respectively, illustrated absorption peaks at T<sub>2</sub>, corresponding to Si–OR, following the hydrolysis of TEOS with OTES, indicating that the absorption structure tended to be of the type (R–O)Si(–OSi≡)₃. The structure of the Si–OR absorption peak indicated that some Si quadruple bonds had an unreacted Si–OH functional group, (H–O)Si(–OSi≡)₃, and some Q<sup>3</sup> quadruple bonds had an unreacted Si–O functional group, Si(–OSi≡)₃. The structure of Q<sup>4</sup> indicates that the Si quadruple bond reacted completely with the Si–O functional group: Si(–OSi≡)₄. In the spectrum of hybrid T<sub>j</sub> and T<sub>k</sub>, the strength of the Q<sup>4</sup> absorption peak (at approximately -100.42 and -100.40 ppm) increased, and the Q<sup>4</sup> absorption peak appeared at a δ value of ~108.14 and ~109.09 ppm. In addition, the weak absorption peak T<sub>1</sub> appeared at approximately 63.91 and 63.83 ppm. The results for hybrid T<sub>j</sub>
indicated a Q^3 absorption peak at a δ value of ~100.06 ppm; the Q^4 absorption peak decreased, and the T^2 absorption peak increased significantly. Regarding hybrid T4, the Q^3 absorption intensity increased substantially, and the T^2 absorption intensity increased. The Q^4 absorption peak for hybrids T1–T4 gradually shifted into a Q^3 absorption peak, as shown in Figure 4. Therefore, with increasing OTES proportion, Si–O replaced Zr–O, and Si–O–Zr was formed. The noise peak reduction indicated that increasing the TEOS proportion enhanced the network structure purity.

When additional zirconia gel was added to hybrids Z4, the quantity of zirconium increased, which caused the quantity of Si to decrease. It is possible that the carbon proportion decreased and the oxygen increased in the hybrid material because of the formation of additional Zr–O–Zr bonds in zirconia thin films of hybrids Z1–Z4 and the gradually increasing zirconium proportion.

Table 2 shows the results of the EDS analyses of T1–T4. The molar ratio of the ZNP/TEOS/OTES hybrid material increased with increasing amounts of OTES because the Si(OH)4 structure (formed by hydrolysis–condensation of OTES and TEOS) combined with the zirconia. The Si peaks revealed that the intensity of the ZNP/TEOS/OTES hybrid material increased as the amount of OTES increased. The amounts of carbon and zirconium decreased for the T4 hybrid material. This was because the T4 ratio formed more Si–O–Si bonds, which is shown by the high Si content in Fig. 6. Furthermore, the amount of zirconium increased in the Z4-treated Cotton fabric. Therefore, a larger hybrid material ratio resulted in greater Zr–O–Si bonding. According to Table 2, when the TEOS and OTES amounts were fixed, the silica content decreased with increasing zirconium content. The maximum Z4 via ZNP addition level in the treated cotton fabrics resulted in the maximum zirconium content.

C. X-Ray diffraction analysis

In this study, we performed an X-ray diffraction analysis of hybrid materials in the Z series with various proportions developed by a sol–gel process. Because neither the series of hybrid materials nor the Zr powder was sintered, all of these materials exhibited amorphous structures without clear crystal phase peaks, as shown in Figures 5. Figure 5 shows that the diffraction peaks of Z1, Z2, Z3, and Z4 occur at 2θ = 21.36°, 20.89°, 21.12°, and 21.50°, respectively. These results indicate that with the exception of the Zr powder, the morphology of amorphous-phase OTES and TEOS affected the diffraction peaks of the hybrid materials Z1–Z4.

D. Energy-dispersive X-ray spectra analysis

The energy-dispersive X-ray spectroscopy (EDS) analysis results are presented in Table 2 and Figure 6.

![Fig. 4 Si-NMR spectra of hybrid material T1–T4](image)

![Fig. 5 XRD spectra of Z1–Z4](image)

![Fig. 6 The EDS diagram of hybrid materials Z4](image)

Table 2: EDS analysis of hybrid materials Z1–Z4 and T1–T4

<table>
<thead>
<tr>
<th>Samples</th>
<th>Elemental composition (%)</th>
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<tbody>
<tr>
<td></td>
<td>C</td>
</tr>
<tr>
<td>Z1</td>
<td>35.5</td>
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<tr>
<td>Z2</td>
<td>33.23</td>
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<tr>
<td>Z3</td>
<td>32.51</td>
</tr>
<tr>
<td>Z4</td>
<td>30.31</td>
</tr>
<tr>
<td>T1</td>
<td>35.89</td>
</tr>
<tr>
<td>T2</td>
<td>35.17</td>
</tr>
<tr>
<td>T3</td>
<td>34.23</td>
</tr>
<tr>
<td>T4</td>
<td>34.01</td>
</tr>
</tbody>
</table>
E. Surface morphology of finished cotton fabric

SEM was used to study the morphology of the cotton fabrics. The untreated cotton fabric shows a smooth morphology (Fig. 7a). Figures 7b–7e show SEM images of fabrics treated with varying ZNP molar ratios. An increase in the molar ratio of ZNP with constant amounts of OTES and TEOS revealed irregularly sized particles with a flake-like morphology. As the amount of zirconia increased, the pore size of the treated surface texture decreased as a result of the Zr–O–Zr network structure, which effectively filled the pores of the fabrics, thus improving the water repellency characteristics of the treated fabric. SEM micrographs of fabrics treated with varying OTES molar ratios (Figs. 8a–8d) show irregularly sized particles with a flake-like morphology. Figure 8a shows small particle blocks and uneven distribution of zirconia and silica particles on the surface. As shown in Figs. 8b–8d, the particle distribution was much more uniform with increasing amounts of silica, indicating that the hybrid materials were more evenly dispersed across the fabric, resulting in the formation of a protective layer. Surface-bound particles were more homogeneous in terms of size and distribution and rough regions appeared on the surface. Fabric pores were marginally filled, resulting in a slight increase in the water repellency characteristics and conductivity. Through the above analysis confirms that the Z and T series hybrid materials can be universally processed in cotton dyed fabrics. It also echoes the preamble of using the hydrothermal method to prepare the hybrid materials to uniformly disperse the cotton fabrics and make the surface of materials appear adhesion.

F. Contact angle analysis of treated cotton fabrics

Table 3 shows the results of the contact angle analysis for the treated cotton fabrics. The contact angle of untreated cotton fabrics was 0°. The average contact angle for CT1 before washing was 130.9° and this value increased with increasing OTES molar ratio. The increase of OTES molar ratio, in turn, increased the number of siloxane groups in the hybrid materials, which enhanced crosslinking and improved the water repellency characteristics of the cotton fabrics. Thus, CT1 attained the highest contact angle, 138°. The results presented in Figure 9 for the CZ1–CZ4 fabrics show trends similar to those of the CT1–CT4 fabrics as illustrated in Figure 10. However, the contact angles slightly decreased after 10 washing cycles. The reason for these results may be that the hybrid compounds that adhered on the Cotton fabrics were removed from the surface of the fabrics because of mechanical stirring, thereby negatively affecting the contact angle.

By comparing the contact angles of the CZ1–CZ4 fabrics, it can be seen that the fabrics treated with CZ4 exhibited the best contact angle of 134°. Further, the dyeing showed little effect on the contact angle of the cotton fabrics, as shown in Table 3. The contact angles of CZ4 and CT4 after washing were 126.4° and 131.8°.

Table 3. Surface hydrophobicity evaluation of CZ1–CZ4 and CT1–CT4

<table>
<thead>
<tr>
<th>Compd.</th>
<th>Contact angle (°)</th>
<th>Compd.</th>
<th>Contact angle (°)</th>
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<tbody>
<tr>
<td>Before washing</td>
<td>After washing</td>
<td>Before washing</td>
<td>After washing</td>
</tr>
<tr>
<td>C</td>
<td>0</td>
<td>C</td>
<td>0</td>
</tr>
<tr>
<td>CZ1</td>
<td>127.4</td>
<td>CT1</td>
<td>130.9</td>
</tr>
<tr>
<td>CZ2</td>
<td>130.1</td>
<td>CT2</td>
<td>132.0</td>
</tr>
<tr>
<td>CZ3</td>
<td>132.4</td>
<td>CT3</td>
<td>135.0</td>
</tr>
<tr>
<td>CZ4</td>
<td>134.0</td>
<td>CT4</td>
<td>138.0</td>
</tr>
<tr>
<td></td>
<td>126.4</td>
<td></td>
<td>131.8</td>
</tr>
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</table>
respectively, slightly lower than those of CZ₄ and CT₄ before washing.

G. Warmth retention analysis of processed cotton fabrics CZ₁–CZ₄ and CT₁–CT₄

We used a four-point temperature probe to measure the temperature variation of the finished cloths to determine their thermal insulation properties. The analysis results, shown in Figures 11–12, indicate that the grey cotton cloth exhibited the smallest temperature difference following 600 s of irradiation with a halogen lamp. In the case of the grey finished cloth, CZ₄, which had the greatest zirconia sol content, the temperature difference was approximately 31.72 °C, which indicates that the temperature difference increases as the zirconia sol content increases. Thus, the enhanced endothermic effects of the hybrid materials in the finished cloth provided the finished fabric with a thermal storage effect. The insulation properties of the materials improved as the zirconia/TEOS/OTES sol concentrations were increased. Among finished cloths CT₁–CT₄, the largest temperature difference, 28.77 °C, was determined between finished cloth CT₄ and the cotton grey cloth. Thus, the Z and T series of the finished cloth, with additional TEOS/zirconia sol, exhibited comparatively improved thermal storage capacities. An analysis of the insulation properties of finished cloths CT₁–CT₄ revealed a temperature difference of 7.35 °C between finished cloth CT₄ and the cotton grey cloth, which was the optimal result obtained. An analysis of the insulation properties of finished cloths CZ₁–CZ₄ indicated that there was temperature difference of 8.15 °C between finished cloth CZ₄ and the polyester grey cloth, which was the optimal result obtained. Therefore, we determined that the zirconia sol alone causes the inorganic zirconium to agglomerate, which reduces the size and number of gaps in the finished cloth and increases the air content. In addition, the properties of the zirconium and agglomerated inorganic zirconium prevented the material from cooling, contributing to the superior thermal insulation properties of the material.

IV. CONCLUSIONS

Zirconia/silica hybrid materials were synthesized using a hydrothermal process with various molar ratios of the aluminum isopropoxide and OTES chemical precursors. The properties of cotton fabrics coated with the ZNP/TEOS/OTES hybrid materials were examined, and FTIR, ²⁹Si NMR, and EDS analyses confirmed that the zirconia and silica in the hybrid materials bonded to form a network. The FTIR spectra indicated increasing numbers of Si–O–Si and Zr–O–Zr linkages as the molar ratios of ZNP and OTES increased. The diffraction peak of the Zr powder, inoculated Zr-series hybrid material Z₄ appeared at 2θ = 21.05°. Thus, the addition of adequate amounts of ZNP/TEOS/OTES can improve the water repellency properties of treated cotton fabrics, as indicated by the contact angle results for untreated cotton fabrics (0°) and treated fabrics after washing (120.1–131.8°). The thermal storage analysis indicated that the effects became stronger as the zirconia sol content was increased. There was a temperature difference of 31.72 °C between finished cloth CZ₄ and the cotton grey cloth. We can infer that the endothermic effects of the zirconia sol in the hybrid materials enhanced the thermal storage capacity of the treated fabrics. The insulation effect that occurred in the case of finished cloth CZ₄, in which only zirconia sol was added, resulted in a
temperature increase of 8.15 °C compared with that of the cotton grey cloth.

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