Hydrophobization of cotton fabric with silanes with different substituents

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Abstract In this paper hydrophobization of cotton fabric with a range of silanes, differing in number of chlorine atoms and in other substituents, was explored. The fabric modification was carried out in silane solutions in anhydrous toluene and in n-hexane. Water contact angles were measured to characterize hydrophilic/hydrophobicity of the modified fabrics. Surfaces of the fibers were analyzed using SEM, EDS and ATR-FTIR. Treatment with dichlorodimethylsilane, dichloro(methyl)phenylsilane, 3-chloropropylmethyldichlorosilane and (3,3,3-trifluoropropyl)dichloromethylsilane resulted in hydrophobization, whereas trimethylchlorosilane and 3-aminopropyl(diethoxy)methylsilane were inefficient. Mechanisms of coating with the silanes were proposed. Dielectric spectroscopy of the modified fabric demonstrated that resistance was sensitive to efficiency of reactions removing water from the cotton fibers.

Keywords Cotton fabric · Superhydrophobicity · Silanes · Dielectric spectroscopy · ATR-FTIR
Introduction

Cotton fibers are the most popular natural fibers used to produce a broad range of textiles. Being lightweight, porous and flexible, cotton textiles are widely used. Cotton is hydrophilic, which is an advantage in many applications. However, engineering of superhydrophobic textile surfaces gains significant scientific and industrial interest because of potential applications in outdoor wear and protective textiles. Different methods of the cotton modification were investigated (Vasiljevic et al. 2013; Xu et al. 2012; Xue et al. 2009; Yang et al. 2018). Superhydrophobicity of soft woven fabrics can be achieved by coating; the most desirable from the practical point of view are coatings that are thin, transparent and durable (Brzezinski et al. 2012; Ma et al. 2018). To impart superhydrophobicity, modification of surfaces can be carried out, which changes surface energy or increases surface roughness by deposition of particles (Darmanin and Guittard 2015). The latter method was applied recently by (Makowski et al. 2014, 2019) to achieve superhydrophobicity of surfaces of fibrous materials coated with carbon nanoparticles. Trichloromethylsilane (TCMS) in an anhydrous environment was used to form polymethylsilsesquioxane globular particles on the modified surfaces, which resulted in the lotus effect. It was demonstrated that the shape and size of the particles formed on fiber surfaces depended on relative humidity during pre-treatment conditioning of cotton fibers (Artus and Seeger 2014; Makowski et al. 2014). It is worth noting that TCMS hydrophobization of cotton fabric coated with reduced graphene oxide resulted in filament structures on the fiber surfaces (Shateri-Khalilabad and Yazdanshenas 2013). Similar structures were also found on TCMS hydrophobized neat cotton fabric (Shirgholami et al. 2011). Other authors superhydrophobized cotton fabric through treatment with silica nanoparticles and water-repellent agent (Bae et al. 2009), polyphenol (Gu et al. 2017) or industrial waterproof reagent—potassium methyl silicone (Li et al. 2008). The possibility to hydrophobize cotton fabric in the gas phase using fluoromonomers was also explored (Maity et al. 2010).

It is worth noting that silanes other than TCMS, for instance dichloromethylsilane (DCMS), were used to hydrophobize silicon wafers, glass and paper by chemical vapor deposition (Cech et al. 2001) although it is a high-energy method and not always economically viable.

Preparation of hydrophobic or hydrophilic microspheres by a sol–gel method from the functional alkoxyisilanes and by cross-linking of polyhydroxysilanes with low molar mass or oligomeric divinyl compounds was recently reviewed (Slomkowski et al. 2017). Based on the described mechanisms it can be envisaged that not only TCMS, but also other silanes can be applied to hydrophobization of a cotton fabric.

It should be noted that the presence of hydroxyl groups on the cellulose surface makes a condensation reaction with silanes possible (Cunha and Gandini 2010; Fadeev and McCarthy 2000). However only one bond in monofunctional organosilane (R3SiX) is hydrolysable; usually X = Cl or N(CH3)2. Combinations of silica nanoparticles and multifunctional silanes allowed to obtained hydrophobic cotton surfaces (Roe et al. 2012). Recently, bacterial cellulose was hydrophobized with vinyl triethoxysilane or 3-aminopropyl triethoxysilane, and also by acylation and acrylation (Frone et al. 2018).

In this work hydrophobization of cotton fabric using a range of silanes differing in number of chlorine atoms and in other substituents was explored. These compounds were not used previously for such purpose. Water contact angles were measured to characterize hydrophilicity of the modified fabrics and surfaces of the fibers were analyzed using scanning electron microscopy (SEM), SEM with energy dispersive spectroscopy (EDS), attenuated total reflection—Fourier transform infrared spectroscopy (ATR-FTIR) and dielectric spectroscopy.

Experimental

Materials

Commercial plain weave cotton fabric (145 g/m²) 0.36 mm thick, with 205 threads/10 cm and 295 threads/10 cm in the warp and weft directions, respectively, was used in this study. Before the modification the cotton fabric was cleaned, as previously described (Makowski et al. 2014). The fabric samples were purified by extraction for 2 h with diethyl ether (98% p.a. purity) and for next 2 h in anhydrous ethanol (p.a. purity), both from Chempur, Poland. The samples were then additionally purified.
for 2 h in boiling ethanol under a reflux condenser, rinsed with doubly distilled water and dried at 120 °C for 30 min.

The following silanes were used for hydrophobization: trimethylchlorosilane C3H9SiCl (TMCS), dichlorodiethylsilane C3H6Cl2Si (DMDCS), dichloro(methylphenyl)silane C7H5Cl2Si (DCMPhS), 3-chloropropylmethyldichlorosilane C4H9Cl2Si (CPTCS), (3,3,3-trifluoropropyl)dichloromethylsilane C4H7Cl2F3Si (TFDCMS) and 3-aminopropyl(diethoxy)methylsilane C8H21NO2Si (AMDES).

TMCS with 98% purity, was purchased from Fluka Sigma-Aldrich (Germany) whereas DMDCS with 99% purity, DCMPhS, CPTCS, TFDCMS and AMDES, all four with 97% purity, were delivered by ABCR (Germany). The structures of the silanes used are shown in Fig. 1.

In addition, hydrochloric acid, 35%, from Chempur (Poland), toluene p.a. purity from StanLab (Poland), n-hexane, 95%, from HiPerSolv Chromanorm (Poland), 1,4-dioxane, 99.8%, from Chem-Lab NV (Belgium) and ethanol, 96%, from Avantor (Poland) were used.

Hydrophobization of cotton fabric

Before hydrophobization all fabric samples were conditioned at room temperature (RT) in dessicators under vapors of saturated salt aqueous solutions at relative humidity (RH) of 30% and 70% (Makowski et al. 2014).

To impart hydrophobicity, each sample was placed in a polypropylene vessel containing 0.5 M solution of the silane in anhydrous toluene or n-hexane, with exception of 3-aminopropyl(diethoxy)methylsilane, which was dissolved in 1,4-dioxane. 100 ml of solution per 1 g of fabric was used. The process was carried out at RT for 1 h. The samples were then rinsed sequentially in: toluene, ethanol, ethanol-distilled water mixture (1:1) and distilled water, and then dried in air at RT.

Characterization

All cotton fabric samples were coated with 10 nm thick gold layer by ion sputtering using a coater Quorum EMS150R ES (UK) and then analyzed with SEM JSM-6010LA JEOL (Japan) with EDS.

ATR-FTIR spectra of the neat and modified fabric were recorded on a FT/IR-6200 Jasco (Germany) spectrometer equipped with ATR PRO610P-S device.

To determine water contact angles (WCA), 5 µl distilled water droplets were placed on surfaces of the materials at 25 °C. WCA values were measured using a RameHart NRL Goniometer, 100-00-230 (Rame-Hart, NJ) coupled with a camera and optical system. Determination of WCA values was carried out using Drop Analysis program. To obtain average values, the measurements were repeated five times for three samples of each material and the results were averaged.

The impedance spectra in the frequency range of 20 Hz–1 MHz were determined using a HP4284A precision RLC meter, with sinusoidal signal with 20 V amplitude. The samples were sandwiched between

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Fig. 1 Structures of siloxanes: (1) trimethylchlorosilane; (2) dichlorodiethylsilane; (3) dichloro(methylphenyl)silane; (4) 3-chloropropylmethyldichlorosilane; (5) (3,3,3-trifluoropropyl)dichloromethylsilane; (6) 3-aminopropyl(diethoxy)methylsilane
two copper electrodes in the form of disks with 10 mm in diameter, and connected to the measurement unit with 1 m cable. The distance between the two electrodes was constant and equal to 301 μm.

**Results and discussion**

Photographs of water droplets on surfaces of the modified fabric samples are shown in Table 1 together with corresponding WCA values. The untreated cotton samples as well as those treated with AMDES and TMCS were hydrophilic and water droplets soaked in. In the case of DMDCS, DCMPhS, CPTCS and TFDCMS, superhydrophobicity was achieved, reflected in WCA angles 157–165° and 163–174°, after conditioning at RH of 30% and 70%, respectively. The higher RH during the conditioning increased the WCA values, similarly as reported previously (Makowski et al. 2014). In turn, the type of solvent used had no significant effect on the WCA values, except for CPTCS and TFDCMS treatment of fabric conditioned at RH of 30%, for which the use of n-hexane as a solvent increased somewhat these values. Most possibly, due to a smaller content of residual water in n-hexane, the reactions leading the hydrophobization (described below) were more localized on fiber surfaces. It is worth noting that the effect vanished for the fabric samples conditioned at RH of 70% due to higher water content in the fabric.

Surfaces of neat cotton fibers are presented in Fig. 2 showing surface roughness on a microscale, resulting from the presence of microfibrils (Brzezinski et al. 2012). The silane treated surfaces are shown in Fig. 3. Except for Fig. 3a, which shows fiber surface very similar to that of neat cotton fiber, thin layers and particles resulting from silane treatment are visible. Small cracks in the layers, occasionally seen, resulted most probably from damage caused by the electron beam. However, the surface topography is different than that described previously (Makowski et al. 2014). No evenly distributed spherical objects or filament

| Reagents     | RH 30% WCA(°) | WCAh(°) | RH 70% WCA(°) | WCAh(°) |
|--------------|---------------|---------|---------------|---------|
| Untreated cotton | Soak          |         | Soak          |         |
| AMDES*       | Soak          |         | Soak          |         |
| TMCS         | Soak          |         | Soak          |         |
| DMDCS        | 163 164       |         | 168 169       |         |
| DCMPhS       | 164 165       |         | 167 169       |         |
| CPTCS        | 154 158       |         | 174 174       |         |
| TFDCMS       | 157 163       |         | 172 172       |         |

*1,4-Dioxane
structures were clearly discernible on the surfaces studied unlike in (Makowski et al. 2014; Shirgholami et al. 2011). Exemplary EDS spectra for silane treated fabric samples, conditioned previously at RH of 70%, are shown in Fig. 4. Very similar results were obtained for samples conditioned at RH of 30% (not shown).
Fig. 4  EDS spectra of cotton fabric samples conditioned at RH of 70%, and treated with silanes dissolved in toluene: a TMCS, b DMDCS, c DCMPhS, d CPTCS, e TFDCMS, f AMDES
All spectra demonstrated the presence of Si, which evidences the deposition of reaction products on the fiber surfaces.

The probable mechanisms of hydrolysis and condensation reactions are shown Fig. 5. The first step was reaction with water molecules loosely associated with cotton or residual water molecules in solvents, which resulted in formation of silanol groups, capable of condensation reaction. In all cases linking of reaction products to cotton was possible through condensation involving cotton hydroxyl groups. Hydrolysis and condensation reactions of silanes with two Cl atoms bound to Si atom can result in the formation of linear oligomers/polymers, that form a hydrophobic coating on the surface of the fiber. This explains superhydrophobicity of the fiber surfaces treated with DMDCS, DCMPhS, CPTCS and TFDCMS. It is worth noting that only on the surface of CPTCS treated fibers the presence of Cl was shown by EDS, most probably because Cl in the chloropropyl group remained unreacted, being less reactive than Cl atoms bonded directly to Si (Rozga-Wijas et al. 2010). In the case of AMDES the condensation reactions are also possible but the presence of –NH2 end groups makes the resulting coating hydrophilic.

![Fig. 5 Schemes of reactions during treatment of cotton fibers with: a TMCS, b DMDCS, DCMPhS, CPTCS, TFDCMS, c AMDES](image)

![Fig. 6 ATR-FTIR spectra of neat and modified cotton fabric](image)

ATR-FTIR spectra of the neat and modified cotton fabric are shown in Fig. 6. Bands characteristic of polysiloxanes are indicated with arrows: at 789–796 cm\(^{-1}\) (–CH\(_3\) rocking and Si–C stretching in Si–CH\(_3\)), at 1074 cm\(^{-1}\) (Si–O–Si stretching), at 1260–1259 cm\(^{-1}\) (CH\(_3\) deformation in Si–CH\(_3\)) (Johnson et al. 2013). Bands at 730 cm\(^{-1}\) and 1430 cm\(^{-1}\) are characteristic of Si–Ph, the latter very narrow (Launer and Arkles 2013). It must be added that those bands are weak, because the coatings are
very thin and the spectra are dominated by signals of cotton.

Figure 7 shows frequency dependence of resistance (real part of impedance), R, of the modified and neat cotton and the value of R at 500 Hz.

It can be seen that after the modification with TCMS, DCMPhS, CPTCS and TFDCMS R increased, especially at low frequency. R is most possible related to the presence of water associated with cotton and its increase reflects efficiency of reactions leading to removal of water from the fibers.

Conclusions

Treatment of cotton fabric with a range of commercially available silanes, with different numbers of chlorine atoms and different substituents, was explored. Superhydrophobicity of the fabric was achieved after 1 h treatment with DMDCS, DCMPhS, CPTCS and TFDCMS dissolved in anhydrous solvents. As a result of the treatment, the surfaces were coated although globular or filament structures, like those observed previously in the case of TCMS treatment (Makowski et al. 2014; Shirgholami et al. 2011), were absent. AMDES and TMCS treatment was inefficient, and did not impart hydrophobicity to the cotton fabric. Hydrolysis and condensation reactions of silanes with two Cl atoms bound to Si atom can result in the formation of linear oligomeric structures that form a hydrophobic coating on the surface of the fibers. The ATR-FTIR spectra confirmed the modification of the cotton fabric with the siloxane compounds. The results obtained can contribute to optimization of technological processes of hydrophobization of cotton textiles.

Acknowledgments

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