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Continuous and rapid fabrication of photochromic fibers by facilely coating tungsten oxide/polyvinyl alcohol composites†

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Photochromic fibers have attracted great attention due to their wide use in areas of military camouflage, safety warnings, anti-counterfeiting, entertainment, etc. Compared with photochromic organic materials, inorganic photochromic tungsten trioxide (WO_3) materials have been extensively studied, because of their good stability and cost efficiency. In this work, we report the continuous fabrication of photochromic fibers in a simple and low-cost way by dip-coating WO_3 /PVA composites. The prepared photochromic fibers show fast and reversible color switch from light yellow to dark blue upon UV irradiation and infrared heating treatment. The obtained photochromic fibers can be produced on a large scale and be woven into various patterns with good mechanical strength and washability, showing great potential in developing photochromic textiles.

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1 Introduction

Modern clothing is currently moving towards more and more intelligent applications with the addition of some high-end functionalities, such as physiological signal acquisition,¹ performance monitoring,² flexible electronics and display,³ temperature control,⁴ energy storage,⁵ etc. However, most of the current so-called “smart clothing” is made of technologies being integrated into common garments rather than made of truly functional fibers. Nevertheless, many fibrous devices have been extensively developed in the past few years owing to their unique one dimensional structure with the ability to perfectly implant into textiles or clothes and with promising applications in portable and wearable devices, such as solar cells,⁶ supercapacitors,⁷ color variation,⁸ lithium ion batteries,⁹

electroluminescence,¹⁰ etc. These functional fibers represent a significant strategy towards the development of intelligent textiles.

Among the various functional fibers, smart color-changing fibers have become a hotspot for their potential applications, such as wearable displays, visual sensors, military camouflage and so on.^{11–15} So far, the existing smart chromatic fibers can change their colors upon encountering diverse stimuli like temperature,¹⁶ ray radiation,¹⁷ specific solvents and gas,^{18,19} stress tension²⁰ and electricity.^{21–23} Although electricity was usually thought to be an active and controllable approach to trigger the color change, the fabrication of electrochromic or electrothermal chromatic fibers is not easy due to their complex device structure with solid and liquid phase components. The multi-layer coating process, including conductive layer, photo-catalytic layer, thermochromic layer and encapsulation layer, leads to difficulties in continuous mass production. Power supply is essential for these chromatic fibers and current pass through the whole fibers, which raise obvious safety considerations for wearable applications. Compared with electrochromic devices, photochromic applications are free of complex device configurations and electric supply. Just upon photo-irradiation, coloration occurs.^{24,25}

Therefore, in this work, we designed a photochromic fiber with simple core–shell structure. Of the photochromic materials studied so far, tungsten trioxide (WO_3) has become one of the most extensively studied candidates. Integrating WO_3 nanomaterials into a polymer matrix is a feasible method for preparing composite materials with controllable photochromic properties. Several groups have studied the photochromism of the inorganic/organic composite photochromic materials.^{26–33}

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† Electronic supplementary information (ESI) available: The original pictures of WO_3 nanorods powder; UV-vis diffuse reflectance spectra of the WO_3 nanorods powder with different infrared heating time; the pictures of WO_3 /PVA composite films with different WO_3 nanorods contents; the pictures of three kinds of coating solutions with different WO_3 content after mixing and standing for 2 h at room temperature; the coating thickness as function of fiber drawing speed and SEM images of the surface of coated fiber with different drawing speeds; UV-vis diffuse reflectance spectra of one single photochromic fiber with different UV irradiation time; schematic diagram of the experimental process for studying the discoloration process of composite fiber by using various oxidation methods. See DOI: 10.1039/c8ra05170d





Kuboyama *et al.*²⁶ prepared polyethylene glycol (PEG)/WO₃ hybrids and studied their photochromic properties. The photochromism of WO₃ was dramatically enhanced in the presence of PEG. Yano *et al.*²⁷ produced poly(vinyl alcohol) (PVA)/WO₃ hybrid by using ion exchange process of Na₂WO₄·2H₂O. Good mechanical and photochromic properties were presented by the PVA/WO₃ hybrids. Chen *et al.*²⁸ successfully prepared WO₃-TiO₂-ZnO ternary composite films by a sol-gel method using PEG-400 as an organic additive. The addition of PEG-400 helped to improve the photochromism of composite film. Yamazaki *et al.*²⁹ fabricated photochromic film by incorporating WO₃ nanoparticles in methylcellulose matrix with addition of little amount agents. The biodegradable composite film had good photochromic properties with high contrast of the color change before and after UV irradiation. Recently, Zhou *et al.*³⁰ prepared photochromic inks by dispersing the WO₃ nanoparticles into aqueous PVP solution, and casted the photochromic films on flexible substrate. The composite film exhibited fast photochromic switching rate and good reversibility. Although these composite materials based WO₃ and polymers have been received much attention, and have been used in various applications, such as, photochromic films,³⁰ bio-electronic devices,³¹ rewritable media³² *et al.*, these researches are mainly focused on photochromic composite film, while the photochromic fiber based on WO₃ is rarely report. Recently, some hybrid WO₃/polymer nanofibers have been prepared *via* electrospinning,³⁴⁻³⁷ while these electrospun nanofibers are all applied in form of nonwoven fabric rather than the continuous monofilament because of low mechanical strength. Therefore, in this paper, the basic idea of this research is preparation of photochromic fiber with good mechanical strength by coating WO₃/PVA composites. The produced functional fiber has good photochromic properties, and can be woven into fabric blending with ordinary fiber, which may apply in some special fields like military, firefighting *etc.*

Herein, a simple process was employed to prepare photochromic fiber based on WO₃ nanomaterials. The first step was to prepare light-yellow WO₃ nanorods by a one-step hydrothermal method. The WO₃ nanorods were then uniformly dispersed into PVA solution by magnetic stirring, and finally coated on the surface of cotton fiber using a rapid and continuous dip-coating process. The WO₃-based photochromic fiber exhibited fast photochromic performance, good reversible color change. The produced photochromic fiber also displayed good mechanical strength and stability, and could be woven into various patterns and fabrics. This study proposes a feasible strategy for large-scale preparation of photochromic fiber, and the produced photochromic fiber may have potential applications in the areas of military, firefighting *et al.*

2 Experimental section

2.1 Materials

Sodium tungstate dehydrate (Na₂WO₄·2H₂O) used in this study was supplied by shanghai Titan Scientific Co., Ltd. Potassium sulfate (K₂SO₄) was purchased from Jiangsu Strong Functional Chemical Co., Ltd. Oxalic acid dihydrate (C₂H₂O₄·2H₂O) was

purchased from Admas Reagent Co., Ltd. Polyvinyl alcohol (PVA, 1799), hydrochloric acid and ethanol were purchased from Sinopharm Chemical Reagent Co., Ltd. Cotton fiber was purchased from the commercial market. All of the chemicals were used as received without further purification. Water used in all experiments was ultra-pure water.

2.2 Synthesis of WO₃ nanorods

WO₃ nanorods were synthetized as the following process.³³ 4 g Na₂WO₄·2H₂O and 3.3 g C₂H₂O₄·2H₂O were dissolved in 100 ml ultra-pure water to form a homogeneous solution. Next, the solution was acidified to a PH value of 1.68 by using 3 mol L⁻¹ hydrochloric acid. And then 6 g K₂SO₄ was added into the solution with stirring 1 h to form a transparent and homogeneous precursor solution. Next, the prepared precursor solution was transferred into a Teflon-lined autoclave with a stainless steel shell. After reaction at 180 °C for 36 h, the final naturally cooled products were collected by vacuum filtering and washed separately with ultra-pure water and ethanol to remove residual ions. Finally, the obtained products were dried at 60 °C in a vacuum oven.

2.3 Preparation of composite solution of WO₃ and PVA

Firstly, 10 wt% transparent PVA solution was prepared by dissolving PVA in DI water at 85 °C under stirring for 4 h. Then, the as-synthesized WO₃ nanorods powders were added to the prepared PVA solution with three different WO₃ solid contents: 1 wt%, 3 wt%, 5 wt%. The mixtures were pre-stirred at 1000 rpm for 10 min, and then 1 wt% no-ionic surfactant (TX-100) related to amount of WO₃ were added to improve dispersion of WO₃ nanorods in PVA solution.^{38,39} Finally, the mixtures were agitated strongly using stirring bar at 1000 rpm for 1 h to gain the homogeneous WO₃/PVA gel solution.

2.4 Continuous fabrication of photochromic fiber

As shown in Fig. 1, the photochromic fiber was prepared continuously by three steps: (1) surface coating; (2) fast coagulation; (3) dry collection. The detailed process was described as follows: firstly, the homogeneous blend solution was placed in a 5 ml lab-made conical vessel with aperture inner diameter of 0.8 mm, which was fixed vertically on an iron stand. The cleaned and oxygen plasma (100 W, 5 min) pre-treated cotton fiber, was drawn slowly at constant speed (10 mm min⁻¹) through the blend solution by a computer controlled motor. Secondly, the coated fiber was immersed into alcohol bath subsequently to coagulate the WO₃/PVA composite coating layer quickly. Finally, the cotton fiber with coagulated coating was dried at 70 °C with infrared lamp and collected using a stainless roller with diameter of 2 cm.

2.5 Characterization

The transmission electron microscopy (TEM) images, high resolution transmission electron microscopy (HRTEM) images and the electron diffraction pattern were collected with a transmission electron microscope (FEI Tecnai G2 F20). The

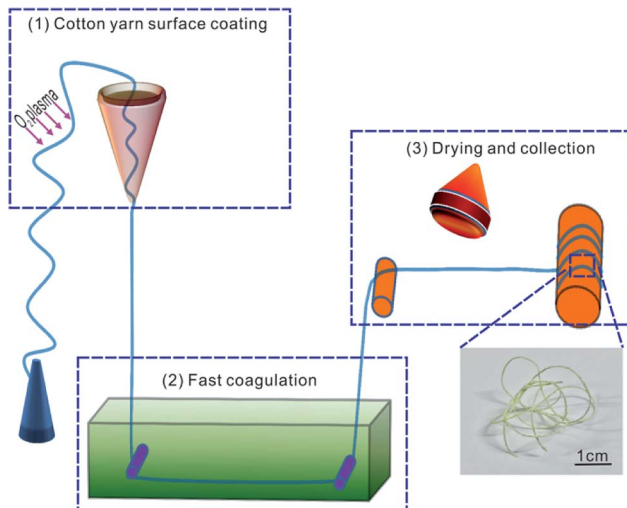


Fig. 1 Schematic illustration of the continuous fabrication process of photochromic fiber.

scanning electron microscopy (SEM) images, the selected area elemental analysis and the energy dispersive spectroscopy (EDX) were performed by a Field Emission-SEM system (Hitachi S-4800). X-ray diffraction (XRD) patterns were collected by a Philips X'Pert-Pro MRD diffractometer with a Cu K α radiation. X-ray photoelectron spectroscopy (XPS) was measured by a Kratos AXIS ULTRA DLD system. All the optical pictures were taken by Nikon D750 digital camera, and the photochromic coloration process was carried out by using a handle UV lamp with 254 nm emission. The UV-vis diffuse reflectance spectroscopies of solid WO_3 samples and photochromic fiber samples were captured by Ideaoptics PG4000 spectrometer system.

3. Results and discussion

The morphology and microstructure of as-synthesized samples were shown in Fig. 1a, which indicates the diameter of obtained WO_3 nanorods is about 500 nm. Phase purity and crystallographic structure of the samples were determined by XRD. As shown in Fig. 2b, all the diffraction peaks of XRD patterns can be indexed to the hexagonal phase of WO_3 (JCPDS no. 75-2187). The TEM image of Fig. 2c further indicates the rod-like microstructure of the synthesized WO_3 . The HR-TEM image shown in Fig. 2d illustrates that the spacing between adjacent lattice planes is 0.38 nm, which is indexed as (002) plane of hexagonal WO_3 and in accordance with XRD data. All the above measured results indicate that the WO_3 nanorods with hexagonal structure have been successfully produced.

The reversible photochromic process of WO_3 nanorods was examined with UV irradiation for coloration and the infrared lamp heating treatment for bleaching. The UV irradiation was implemented using a 36 W commercial UV lamp, and the heat treatment was performed by a 50 W infrared lamp. Fig. 3a shows the gradually changing photographs of WO_3 nanorods powder during UV coloration and heat bleaching cycle with various exposure times. Under UV irradiation process, the color of WO_3

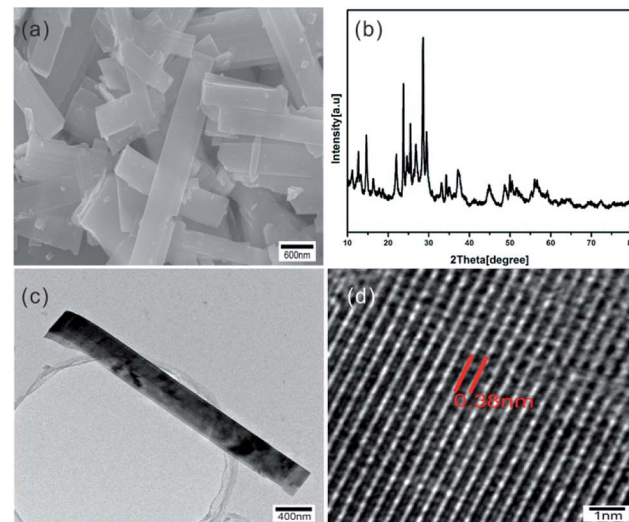


Fig. 2 (a) SEM image, (b) XRD pattern, (c) TEM image and (d) HR-TEM image of as-synthesized WO_3 nanorods.

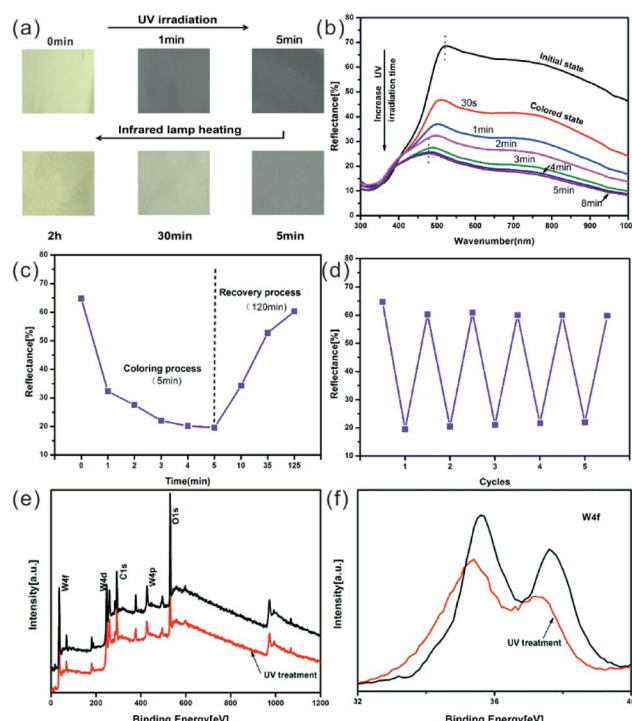


Fig. 3 (a) Photographs of the WO_3 nanorods powder show gradual color change under UV irradiation and infrared heating, these pictures were cut and rebuilt from the original pictures as shown in Fig. S1;† (b) UV-vis diffuse reflectance spectra of the WO_3 nanorods powder with different UV irradiation time; (c) the reflectance intensity of the WO_3 nanorods powder at 600 nm monitored for one coloration–bleaching cycle; (d) the reflectance intensity of the WO_3 nanorods powder at 600 nm after 5 min UV irradiation (coloration) and 2 h infrared heating treatment (bleaching), monitored for continuous 5 cycles; (e) XPS survey spectra of WO_3 nanorods powder before (black) and after (red) UV irradiation; (f) W 4f XPS for before (black) and after (red) UV irradiation states respectively.

nanorods changed from initial pale yellow into dusty blue in one minute and gradually changed into dark blue color within 5 min. During the following infrared lamp heating treatment, the sample's color reversed back to its initial status within about 2 hours. The corresponding reflectance spectrums of color variation were measured as shown in Fig. 3b. The intensity of reflective peak dramatically decreased with UV irradiation, and finally stabilized at 27.6% (initial 67.9%) after 5 min. When extending the exposure time to 8 min, the curve of spectrum almost unchanged. The reflectance spectrum can be recovered to its initial stage after 2 h heating treatment (Fig. S2†), which is in consistent with the discoloration of WO_3 nanorods powder from blue to yellow (Fig. 3a). The intensities of spectrum curves at 600 nm were collected during one coloration–bleaching cycle as shown in Fig. 3c. In coloration process, the intensity decreased from 64.8% to 32.2% within 1 min and the coloration was saturated (19.5%) with UV irradiation for 5 min. In bleaching process, the intensity increased to 60.3% with 2 h heating treatment, which was slight lower than the initial value. If prolonging the thermal treatment time to 5 h, the intensity of reflectance spectrum can fully recover. To evaluate the reversibility of color turning, the WO_3 sample was processed with sequential UV irradiation for 5 min and heating treatment for 2 h repeatedly, and the corresponding reflectance intensities at 600 nm were recorded as 5 coloration–bleaching cycles as shown in Fig. 3d, which demonstrated a good reversibility of color switching.

XPS was employed to study the evolutional structural changes in the coloration process. The wide-scan XPS in Fig. 3e indicates that the W and O are the primary elements in WO_3 nanorods without any other impurities. In order to further investigate binding state, the narrow-scan XPS spectra for W element was carried out before and after UV irradiation. In Fig. 3f, the W 4f two peaks with binding energies of 35.6 eV and 37.7 eV corresponds to W^{6+} state in WO_3 powder before UV treatment. And the energy separation between the main peaks of W 4f_{7/2} and W 4f_{5/2} was 2.1 eV, which further agreed with the tungsten in the W^{6+} valence state. After UV treatment, the W 4f shoulder peaks appear shift towards lower binding energy and the intensity also decreased. The newly formed shoulder peaks can be assigned as W^{5+} species, which were reduced upon UV irradiation.

The as-synthesized WO_3 nanorods were mixed with PVA solution to form homogeneous blend coating solution as shown in Fig. 4a. With the addition of a non-ionic surfactant, the WO_3 nanorods were uniformly dispersed in PVA solution with light yellow color. Three kinds of coating solutions were prepared with different weight content of WO_3 nanorods (1 wt%, 3 wt%, 5 wt%). Fig. S3† shows these blend solutions all have good film forming abilities when they were spin-coated on glass substrate, and the films' color gradually deepens with the increase of WO_3 content both before and after UV irradiation. However, as WO_3 content increases, some precipitation occurs. Three kinds of coating solutions were observed after mixing and standing for 2 h. A small amount precipitation appears in the blend solution with 5 wt% WO_3 content, but there's no obvious aggregation of WO_3 in the blend solution with 3 wt% content, as shown in

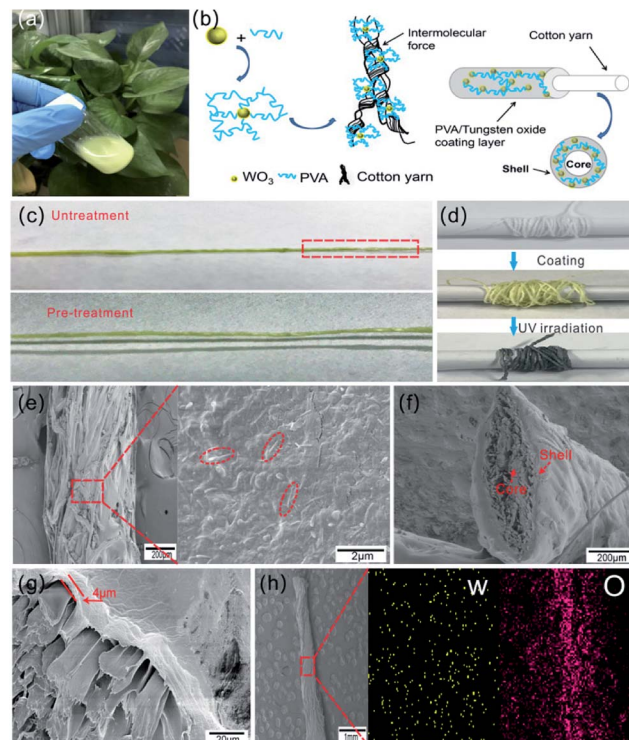


Fig. 4 (a) Picture of WO_3 /PVA blend coating solution with WO_3 content of 3 wt%; (b) the schematic diagram of the coating process; (c) pictures of photochromic fiber separately fabricated with pre-treated and untreated cotton fiber, the area marked with red dotted line shows the uneven coating layer; (d) pictures of pure cotton fiber (upper), photochromic fiber before (middle) and after (below) UV irradiation; (e) SEM images of photochromic fiber's surface with low and high magnification, the area marked with red dotted line indicates the embedded WO_3 nanorods; (f) SEM image of photochromic fiber's cross-section; (g) SEM image of WO_3 /PVA coating layer's cross-section, the cotton fiber drawing speed was 10 mm min^{-1} ; (h) the elemental distribution maps of W and O elements of the composite coating layer.

Fig. S4.† Note that the precipitation can block the nozzle in the coating process, which may cause difficulties in continuous production. Therefore, the blend solution with 3 wt% WO_3 content was selected as coating solution for fabrication of photochromic fibers. The schematic diagram of Fig. 4b shows the mechanism of coating process: WO_3 nanorods were dispersed in PVA solution, when the cotton fiber was drawn through the blend solution, PVA/ WO_3 composite layer was uniformly coated on its surface with core–shell structure. WO_3 nanorods were embedded in PVA polymer to form the functional coating layer with photochromic property. Note that cotton fiber must be pre-treated with O_2 plasma to improve the hydrophilic property. Fig. 4c shows the picture of coated fibers with and without pre-treatment at the same drawing speed. It's obvious that the coating layer is not continuous and has bad binding force with the untreated cotton fiber. In contrast, the composite layer was continuously coated on the surface of pre-treated cotton fiber with strong adhesion force due to hydrogen bonding. This manufacturing process is suitable for mass production, as indicated in Fig. 4d, the photochromic fiber in

length of two meters was collected by a roller, and after UV irradiation, the light yellow color changed into dark blue.

The surface morphology of the photochromic fiber was observed by SEM, as shown in Fig. 4e. The images with low and high magnification indicate a dense layer of PVA/WO₃ composite was coated on the surface of cotton fiber, and the WO₃ nanorods were uniformly embedded in the composite layer, as shown in the area marked with red dotted line. The cross section image in Fig. 4f shows the complete core-shell structure of the photochromic fiber, and the thickness of coating shell was about 4 μm (Fig. 4g). Note that the coating thickness negatively correlate with the fiber drawing speed. In order to form continuous and uniform coating layer, various drawing speeds were tested and the detailed relationship between thickness and drawing speed was shown in Fig. S5.† Finally, the speed of 10 mm min^{-1} was performed as the drawing speed of cotton fiber in the continuous process. The elemental distribution maps of W and O elements of the composite coating layer (Fig. 4h) show uniform distributions of the WO₃ nanorods, which further indicates the inorganic and organic parts are uniformly mixed.

The reversible photochromic response of the prepared composite fibers was also investigated upon continuous UV irradiation and infrared heating process as shown in Fig. 5a. The color of composite fiber changes from initial light yellow to dark blue under UV exposure within 7 min, and gradually turns back to its initial status by using infrared lamp heating within about 2 h. Fig. 5b shows the reflectance spectrum of composite fiber as function of UV irradiation time. It's obvious that the reflective intensity remarkably decreased with continuous UV exposure, and reached saturation within 7 min. The reflectance spectra of one single composite fiber was also recorded, as shown in Fig. 5b, and the original unsmoothed spectral data was

shown in Fig. S6.† The discoloration process of composite fiber was also investigated by using various oxidation methods. Fig. S7† shows the detailed experimental process, one single composite fiber colored with UV irradiation was evenly cut into four pieces, and each piece fiber was bleached with different method. As shown in Fig. 5c, under ambient conditions, the colored composite fiber can be oxidized slowly and gradually get back to original status in 2–3 days, while heating in oven or by infrared lamp at 100 °C, the dark blue color can transfer to initial color in about 2 h. If the sample is put in an ozone generator which can provide a large amount of oxygen radicals, the color will be bleached quickly in only 9–10 min. After comparing these methods, we chose the heating way *via* an infrared lamp as oxidation method to bleach the composite fiber's color because of its non-contact heating process and ozone free pollution. The color switching repeatability is essential for practical applications of photochromic fiber. The UV irradiation (within 7 min) and infrared lamp heating (within 2 h) were applied to study the photochromic reversibility and repeatability of composite fiber, and the corresponding reflectance intensities at 600 nm were recorded as 5 coloration–bleaching cycles as shown in Fig. 5d. After 5 cycles, the color intensity only decreased slightly, which indicated the good reversibility of color turning.

The prepared photochromic fiber can be woven into some interesting shapes, such as grass ring, bow-tie, Chinese knot and even knitted fabric, as shown in Fig. 6a. And after UV irradiation, the color can change from initial light yellow to dark blue. The photochromic reversibility of the composite fiber may have a good prospect application in smart and functional textiles, which may be used in some special fields like military, firefighting, *etc.* Fig. 6b shows the tensile strength–

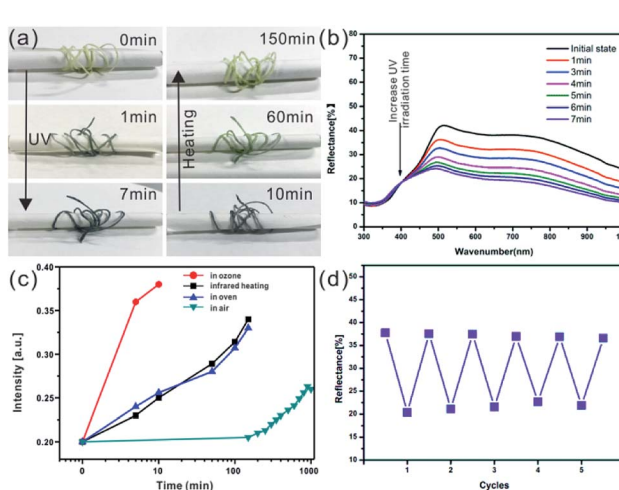


Fig. 5 (a) Optical pictures of the prepared photochromic fiber show gradual color change under UV irradiation and infrared heating; (b) UV-vis diffuse reflectance spectra of one single photochromic fiber with different UV irradiation time; (c) plots of the discoloration process for composite fiber in different environments by monitoring the reflection at 600 nm as a function of time; (d) the cyclic stability of the coloration–bleaching process of the prepared photochromic fiber by recording the reflectance intensity evolution at 600 nm for 5 cycles.

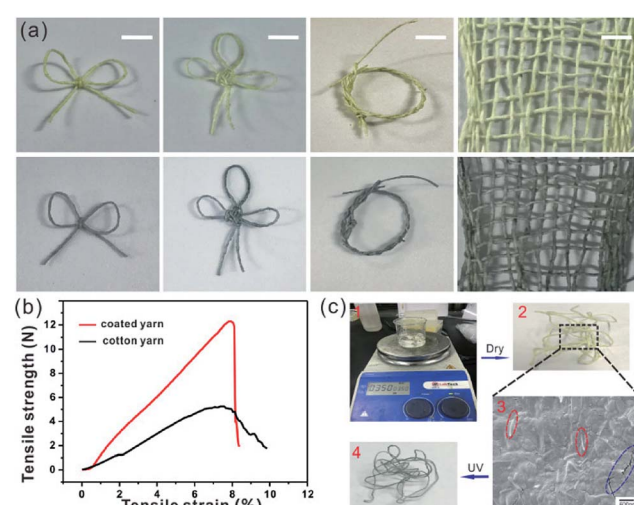


Fig. 6 (a) Pictures of the various shapes woven by prepared photochromic fiber before (upper) and after (below) UV irradiation, the scale bar is 1 cm; (b) tensile strength–elongation curves for cotton and photochromic fiber; (c) the washability of prepared photochromic fiber, (1) picture of the simulated washing process by using a magnetic stirrer; (2) optical image of the washed photochromic fiber after drying; (3) SEM image of the surface of washed photochromic fiber; (4) the picture of the washed photochromic fiber after UV irradiation.



elongation curves of prepared photochromic fiber and pure cotton fiber. Due to the composite coating layer with thickness of 4 μm (drawing speed is 10 mm min^{-1}), the photochromic fiber has a higher breaking strength of 12.2 N, which is almost three times than that of cotton fiber. In order to investigate the washability of prepared photochromic fiber, a simulated washing process was performed by using magnetic stirrer as shown in Fig. 6c1. After washing at room temperature with 350 rpm for 10 min, the photochromic fiber's color (Fig. 6c2) was unchanged, which means the composite coating was not stripped from cotton fiber during the washing process. The SEM image in Fig. 6c3 further indicates the composite coating was not destroyed except for some cracks (area marked with blue dotted line). The washed composite fiber still has the photochromic property, as shown in Fig. 6c4, the fiber's color changed from light yellow into duck blue after UV irradiation. The wonderful mechanical property and washability indicate the prepared photochromic fiber may have good practicability.

4. Conclusions

In summary, WO_3 nanorods were successfully synthesized by a simple one-step hydrothermal method, and the as-synthesized WO_3 nanorods were further uniformly dispersed in PVA solution to form homogeneous composite coating solution. The photochromic fiber was fabricated by coating the as-obtained composite solution onto the surface of cotton fiber. The prepared photochromic fiber shows fast photochromic rate and good reversible color change performance, wonderful mechanical strength and good washability, and the photochromic fiber can be woven into various patterns and fabrics. The WO_3 -based photochromic solution coating method may provide a step toward the development of photochromic fiber for functional textiles, which may have potential application in the areas of military camouflage, safety warning and entertainment, *et al.*

Conflicts of interest

There are no conflicts to declare.

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