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Preparation and characterization of multilayered superfine fibrous mat with the function of directional water transport†

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Directional water transport in garment materials plays a pivotal role in maintaining human thermal and wet comfort. In the present work, a new type of multilayer fibrous mat with the specific function of directional water transport was prepared *via* the combination of melt-electrospinning and solution-electrospinning. The polypropylene (PP) fibrous layer prepared by melt-electrospinning technology was located in the inner layer (next to the skin), while the polyacrylonitrile-containing hydrophilic nano-silica particles (PAN-SiO₂) layer with remarkable hydrophilicity was located in the outer layer, which could effectively transport water to the outer surface of the composites. Treatment of the as-prepared PAN-SiO₂/PP with an alkaline aqueous solution of dopamine not only increased the wettability of the PP layer, but also further improved the hydrophilicity of PAN-SiO₂. A layer of cotton woven mesh was added between the TPP layer and TPAN-SiO₂ to form a sandwich structure in order to accelerate water transport in the bilayered fibrous mats. The directional water transport, mechanical flexibility, and permeability of the prepared multilayered superfine fibrous mat were characterized systematically. The experimental results exhibited that TPAN-SiO₂/cotton mesh/TPP exhibited an excellent accumulative one-way transport index (AOTI, 1071%), remarkable overall moisture management capacity (OMMC, 0.88), and reasonably high water vapor transport rate (WVT, 11.6 kg d⁻¹ m⁻²), indicating it is a promising candidate for the development of novel textile materials for use in the field of sportswear for fast sweat release applications.

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1. Introduction

The rapid transport of human sweat has become one of the required functions of textile materials, and thus has attracted increasing attention.^{1–3} If the sweat generated by the human body cannot be transported and released in time, it may remain in the micro-environment and affect the heat and humidity comfort of the human body, causing some related skin diseases. Thus, most commercially available jerseys are woven or knitted using the technique of hydrophilic modification, profiled fibers and post-finishing to endow textiles with the properties of sweat conduction and quick drying.^{26,32} However, these products often have multiple limitations, such as slow wicking rate, complex manufacturing processes and high cost, which restrict their wide application. Based on this, it is particularly significant to

develop new garment materials with the property of directional water transport. Water is transported to the outer surface of the garment through the fibers and capillary pores in the material.⁶ Janus wettability is an effective water transport mode in porous materials, which is abrupt conduction and relies mainly on the asymmetric water absorption properties of the inner and outer layers to accelerate the rate of transport.^{7,8} According to the above analysis, the inner layer has a relatively hydrophobic and large pore structure, and the outer layer possesses a small pore size with super-hydrophilic properties, which endow the material with remarkable directional water transport performances. For the construction of garment materials with ideal performances, the inner layer may be hydrophobic fibers such as polypropylene fiber (PP) and polyethylene fiber (PET), and polypropylene fine fiber (PAN), polyvinyl alcohol fiber (PVA) and cellulose fibers are ideal for the outer layers.⁹

Electrospinning is an efficient textile processing technology for most polymers for the manufacture of fibrous membranes of various sizes ranging from nanoscale to microscale.¹⁰ Previously, various types of fibrous materials with excellent directional water transport have been prepared by electrospinning technology (shown in Table below).

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Name	Characteristic
Trilayered fibrous membranes ⁵	Continuous, spontaneous, and directional water transport
Cellulose acetate-based nanofibrous membranes ²⁸	Breathable and colorful
Nonwoven/Polyacrylonitrile-SiO ₂ nanofiber composite membranes ¹⁰	Potential candidate for fast sweat release applications
Double-layered membranes constructed of poly(vinylidenefluoride) fibrous layer and nylon-6 nanofibrous layer ³³	Promising candidate for developing modern breathable textiles with enhanced moisture management properties such as sportswear
Unidirectional water-penetration composite of hydrophobic polyurethane and hydrophilic crosslinked poly (vinyl alcohol) fibrous layers ¹	An interesting “water diode” film is fabricated <i>via</i> a facile electrospinning technique
Dual-layer electrospun nanofibrous mats consisting of polyacrylonitrile layer and polystyrene layer with and without interpenetrating nanopores, respectively ²	Tailoring surface hydrophilicity of porous electrospun nanofibers to enhance capillary and push-pull effects for moisture wicking

Presently, electrospun fibrous membranes can be prepared *via* solution electrospinning and melt-electrospinning.^{11,12} The solution-electrospinning process is a method with ordinary operation, and fibers can reach the nano-scale. It is widely used for the development of filtration materials, biomedical materials, flexible sensors and other functional products.¹³ The melt-electrospinning technology uses a heating syringe to melt and eject the solid polymer, which is subjected to electrostatic force drawing, and then cooled and formed in air. Finally, micron-sized fibers are obtained.^{14,15} Also, due to its unique characteristic, melt-electrospinning can be used to spin polymers that are difficult to dissolve under normal conditions (PET, PP, PLA, *etc.*).^{16–18} In addition, the process of melt-electrospinning does not involve solvents, which prevents the volatilization of toxic solvents from causing negative effects on operators and the environment during the spinning process.⁴ It also can be widely used in air filtration, water filtration, high-efficiency oil absorption and other fields.¹⁹

Herein, a multilayered superfine fibrous mat spun by two-mode electrospinning is reported. The polypropylene (PP) fibrous layer inside the mat was spun *via* melt-electrospinning, which contacts with human skin directly due to its hydrophobicity. The outer surface was mixture of polypropylene (PAN) nanofibers and hydrophilic silicon dioxide (SiO₂) particles prepared by solution electrospinning, which is considered suitable as the outer layer material due to its natural hydrophilic property.^{20,21} A dopamine alkaline aqueous solution was used to treat the as-prepared bilayered fibrous mats to improve the wettability of the PP side, while the PAN-SiO₂ nanofiber membranes could partly be hydrolyzed and the silica particles in it fall off, promoting the water evaporation rate on the surface of the membranes. Meanwhile, cotton mesh was specially added between the inner and outer layers to accelerate water

transport and endow the material with outstanding mechanical properties.²² Schemes 1 and 2 illustrate the complete preparation process and performance mechanism of the treated PAN-SiO₂/PP (TPAN-SiO₂/TPP) and treated PAN-SiO₂/cotton mesh/PP (TPAN-SiO₂/cotton mesh/TPP), respectively.

2. Experimental

2.1 Materials

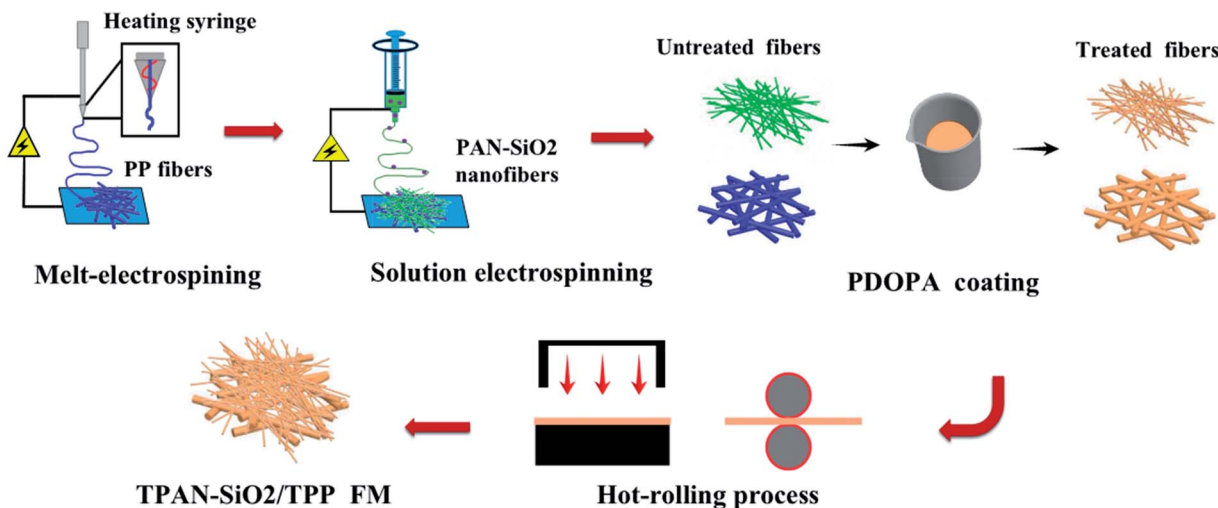
Polyacrylonitrile (PAN) ($M_w = 90\,000$) was purchased from Aladdin Chemicals Co., China. Silica nanoparticles (SiO₂ NP, 10–50 nm), dopamine hydrochloride ($M_w = 189.64$), ammonia water, and dimethylformamide (DMF) were purchased from Aladdin Chemicals Co., China. Distilled water was obtained from a Heal-Force water purifying system. Polypropylene (PP, melt index = 16) was purchased from Yousuo Chemical Technology Co., Ltd. cotton mesh (58.16 g m⁻²) was purchased from Jinhai Textile Co., Ltd. The shirt textile, which was only used for the draping and blending test, was a 100% cotton single jersey (148.64 g m⁻²). All analytical grade chemicals were used as received with no further purification.

2.2 Preparation of fibrous mats

Single-layer fibrous mats from polypropylene (PP) were fabricated *via* melt-electrospinning using a melt-electrospinning machine (MES-001 Laizhou Electronics Co., Ltd.). Before electrospinning, the PP masterbatches were processed into strips by an extruding machine as the as-prepared substance. The as-prepared substance was formed by transporting it in a syringe (diameter: 0.4 mm) heated to 340 °C at a speed of 5 cm min⁻¹ and fixing it on a slide moving at a speed of 30 cm min⁻¹. A high direct current voltage of 36 kV was applied between the syringe and the collector, with a distance of 7 cm maintained between them. The PP fibers were collected on a grounded metallic rotating roller or metallic rotating roller covered cotton mesh at a rotation rate of 45 rpm. The relative ambient temperature during the electrospinning was 45 °C. The syringing was in this mode for 0.5 h, 1 h, 1.5 h and 2 h to prepare PP layers with difference thicknesses, which were denoted as PP-0.5, PP-1, PP-1.5, and PP-2, respectively.

The spinning solution (100 g) comprised of silica nanoparticles (SiO₂ NP) concentration (10 wt% relative to polymer) and polyacrylonitrile (PAN, 12 wt%) was obtained by dissolving the powder reactants in dimethylformamide (DMF) solvent following by magnetic stirring for 8 h. The mixed and uniform solution was electrospun on the surface of PP and cotton mesh-PP by the needle (diameter: 0.3 mm) on an RES-001 Rotary Dynamic Electrospinning Device at a fixed feeding rate of 0.002 mm s⁻¹. A high direct current voltage of 20 kV was applied to the needle tip. The fibrous nanofibers were collected on a grounded metallic rotating roller at the same rotation rate (45 rpm) covered by PP, cotton mesh-PP, and the needle tip-to-collector distance was maintained at 15 cm. The syringing was conducted in a fixed mode for 3 h at room temperature and 45 ± 3% relative humidity.





Scheme 1 Illustration of the preparation method of TPAN-SiO₂/TPP composite FM.

2.3 Treatment with alkaline aqueous solution of dopamine

Poly-dopamine (PDOPA) was coated on the composite fibrous mats *via* *in situ* polymerization. In a typical experiment, ammonia water was dripped into 500 mL distilled water until the pH became 8.5 (measured by a pH tester). A suitable amount of dopamine hydrochloride was weighed to prepared 2 g L⁻¹ dopamine alkaline aqueous solution with 8.5 pH ammonia aqueous solution as the solvent, and the mixed solution was magnetically stirred for 15 min finally. The treatment on the surface of PAN-SiO₂ was conducted for 0.5 h, 1 h, and 1.5 h, and the resulting samples were denoted as TPAN-SiO₂-0.5 h, TPAN-SiO₂-1 h, and TPAN-SiO₂-1.5 h, respectively. Moreover, the surfaces of PAN-SiO₂/PP and PAN-SiO₂/cotton mesh/PP were treated for 1.5 h under continuous stirring, and denoted as TPAN-SiO₂/TPP and TPAN-SiO₂/cotton mesh/TPP, respectively. All the treated mats were washed with distilled water three times and dried under vacuum at 60 °C for 24 h.

2.4 Characterization

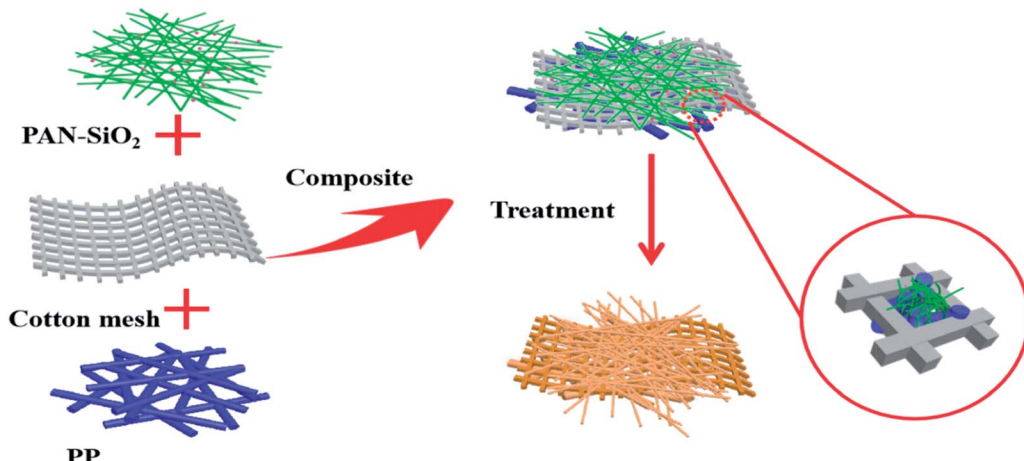
The morphology of the composite fibrous mats was characterized *via* scanning electron microscopy (SEM, S-3400N, Japan).

Fourier transform infrared spectroscopy (FTIR) was conducted on a Spectrum Two spectrometer to examine the molecular structure and chemical information of the different treatment time PAN-SiO₂ and PP samples before and after treatment (Perkin Elmer Inc., USA).

The pore size and pore size distribution of the fibrous membranes were analyzed using a capillary flow porometer (POROLUX 100 FM, Promet Co., Ltd., GER).

The thermal mass loss was investigated using a thermogravimetric analyzer (TGA4000, Perkin Elmer Inc., USA) from 30 °C to 800 °C at a heating rate of 30 °C min⁻¹, which showed the amount of SiO₂ in the treated PAN-SiO₂ fibrous membranes.

The contact angles of droplets (5 µL) were measured using a contact angle tester (Beijing Jinshengxin Testing Instrument Ltd., China).



Scheme 2 Illustration of the preparation of TPAN-SiO₂/cotton mesh/TPP with a "sandwich" structure.



The thickness of each layer of the composite mat was measured using a fabric thickness gauge by applying a certain pressure (500 g).

A moisture management tester (MMT, SDL ATLAS, Ltd., China) was used to measure the directional water transport performance of the mats. During the test, the sample was placed between two arrays of electric resistance sensors. The transfusion tube in the sensors dripped a fixed concentration of saline water onto the fibrous mats. The probe of the sensors tested the resistance between different rings to reflect the absorption and diffusion of the liquid in the fibrous mats by the change in resistance. The accumulative one-way transport index (AOTI) and over moisture management capacity (OMMC) were calculated as follows:

$$AOTI = \frac{\int(U_b) - \int(U_t)}{T} \quad (1)$$

where, U_b is moisture content in the bottom layer, U_t is the moisture content in the top layer and T is the total testing time.

$$OMMC = 0.25BAR + 0.5AOTI + 0.25BSS \quad (2)$$

where, BAR is the bottom absorption rate ($\% \text{ s}^{-1}$), AOTC is the accumulative one-way transport index and BSS is the bottom spreading speed ($\% \text{ s}^{-1}$).

The tensile strength test was based on GB/T3923.1-2013, with modification. The samples were examined using a testing machine (YG006, China) under a fixed stretching speed of 100 mm min^{-1} . The dimensions are fixed parameters ($50 \times 150 \text{ mm}$) and the distance gauge was set to 100 mm.

The water vapor transmission rate test was based on GB/21655-2008, with modification. Samples were cut into fixed sizes ($100 \text{ mm} \times 100 \text{ mm}$), and suspended naturally after wetting with water droplets (0.2 g). The mass of the samples was measured every 5 min.

The wicking height test was based on GB/21655-2008, with modification. The samples were cut into strips (20 mm wide) and dipped into distilled water, and the wicking height was recorded every 2 min.

The drape coefficient and blending length were measured using a YG811 fabric drape tester and FAST-2 bending tester, respectively, followed a reported procedure.^{23,30} The drape coefficient was calculated using the following formula:

$$DC = \frac{S_d}{S_o} \times 100\% \quad (3)$$

where, DC is the drape coefficient, S_d is the area of dropping part and S_o the original area.

Also, the drape stiffness was calculated using the following formula:

$$BS = 9.81 \times 10^{-8} WL_B^3 \quad (4)$$

where, BS is blending stiffness, and W and L_B the sample of unit area quality (g m^{-2}) and blending length (mm), respectively.

The air permeability test was conducted using an automatic permeability meter (YG461E-III, Ningbo Textile Instruments Ltd., China).

The moisture permeability was tested according to GB/21655-2008, and the moisture permeability was calculated using the following formula:

$$WVT = \frac{24 \times \Delta m}{S} \quad (5)$$

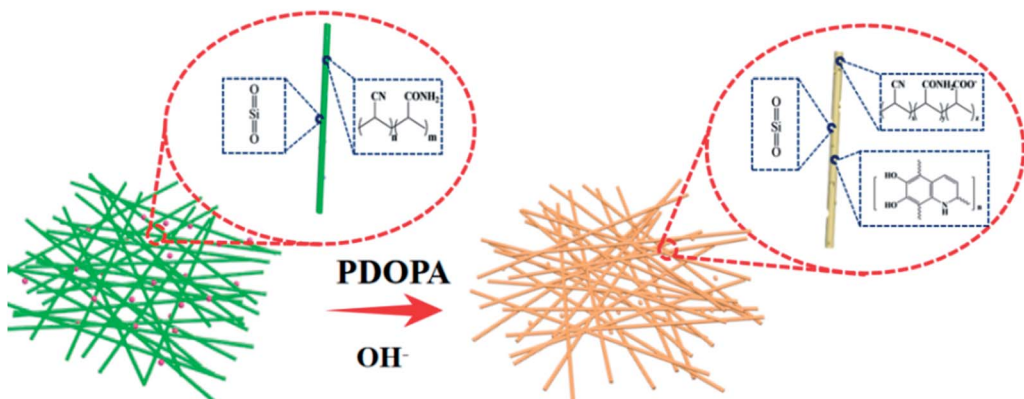
where, WVT is the moisture permeability, Δm is the difference in the mass of water and S is the experimental area of the sample.

3. Results and discussion

3.1 Effect of treatment time for PAN-SiO₂ nanofiber membrane on its hydrophilic property

PAN-SiO₂ nanofibers were modified by hydrolysis and dopamine treatment in alkaline aqueous solution, and subsequently used to prepare nanofiber membranes with superhydrophilic properties (Scheme 3). Fig. 1a-d show the scanning electron microscopy (SEM) images of the PAN-SiO₂ nanofibers membrane, which illustrate the detailed surface morphology obtained by treatment with dopamine alkaline aqueous solution for different times (0 h, 0.5 h, 1 h, and 1.5 h, respectively). It can be seen from the images that the surface of the PAN-SiO₂ fibers became rougher with an increase in treatment time. After treatment for 1.5 h, obvious grooves appeared on the surface of the PAN-SiO₂ fibers, which were formed *via* the partial hydrolysis of the fibers under the action of hydroxyl radicals (OH⁻) in the treatment solution. Moreover, the average pore size of the PAN-SiO₂ membranes treated with dopamine alkaline aqueous solution was less than that of the untreated nanofibers (1.64 μm). When the treatment time was 0.5 h and 1 h, the average pore size was 0.97 μm and 1.1 μm , respectively. When the treatment time was extended to 1.5 h, the average pore size became 1.54 μm (Fig. 2). Although the nanofibers tended to be thin with the prolongation of the treatment time (FS 1↑), which led to an increase in the pore size of the membranes, during the dipping process, the tiny dopamine particles adhered to the surface of the fibrous membranes, making the pore size partially sheltered and reduced. However, when the dipping treatment was 1.5 h, the pore size became large due to the decrease in the diameter of the fibers, which was a more significant factor than the effect of dopamine particles. Fourier transform infrared spectroscopy (FTIR) was used to observe the surface chemical structure (Fig. 3a) of the PAN-SiO₂ nanofibers treated in alkaline dopamine solution for different times. All the spectra showed obvious characteristic peaks at 2246 cm^{-1} , which correspond to the stretching vibration of the nitrile group (C≡N). After treatment with the alkaline dopamine solution, the characteristic peaks were observed at 1616 cm^{-1} , while the intensity of the peak at 1659 cm^{-1} gradually decreased. This indicated that hydroxyl (COO⁻) was present in the PAN-SiO₂ nanofiber membranes treated with alkaline dopamine solution. Although the intensity of the peak at 1659 cm^{-1} gradually weakened with an increase in treatment time, it still showed that amide bonds (CONH₂) were present in all the samples. The characteristic peaks at 1088 cm^{-1} indicated that SiO₂ still existed on the surface of the material after treatment. Table 1





Scheme 3 Illustration of the hydrolysis of the PAN-SiO₂ membrane in alkaline aqueous solution of dopamine.

shows the parameters of PAN-SiO₂ with different treatments time during the thermal decomposition. According to the thermogravimetric analysis (TGA) results and the differential TGA diagram (FS 2†), the weight residue of the nanofiber membranes tended to decrease with an increase in treatment time, indicating some of the SiO₂ nanoparticles fell off the nanofibers. This is because the thermal stability of SiO₂ was superior to that of the polymer,³⁴ and its surface became rougher. Moreover, after 1.5 h of treatment, the characteristic peak appeared at around 3350 cm⁻¹, which may be attributed to the stretching vibration of -OH and -NH, indicating that a certain amount of dopamine was polymerized on the surface of the nanofibers.

The existence of nitrile groups and amide bonds in PAN-SiO₂ nanofibers caused the nanofibers with a rough surface have certain hydrophilic properties. With an increase in the

treatment time of the nanofibers in alkaline aqueous solution of dopamine, the wicking height (Fig. 4a) of the nanofibers increased correspondingly, which indicates that the hydrophilicity of the nanofibers was further improved under the action of the stronger hydrophilic chemical groups on the surface and hierarchical micro-nanoscale roughness.^{5,20,26} When the treatment time was extended to 2 h, the wicking height tended to drop, which is attributed to the increase in the pore size as the fibers became thinner, weakening the wicking effect of the materials (FS 3†).

It was observed that the wicking height reached about 10 cm when the PAN-SiO₂ nanofiber membrane was treated in the alkaline aqueous solution of dopamine for 1.5 h; thus, the optimum treatment time for the PAN-SiO₂ nanofiber membrane was fixed at 1.5 h in subsequent experiments. Simultaneously, the water evaporation rate (Fig. 4b) increased significantly with

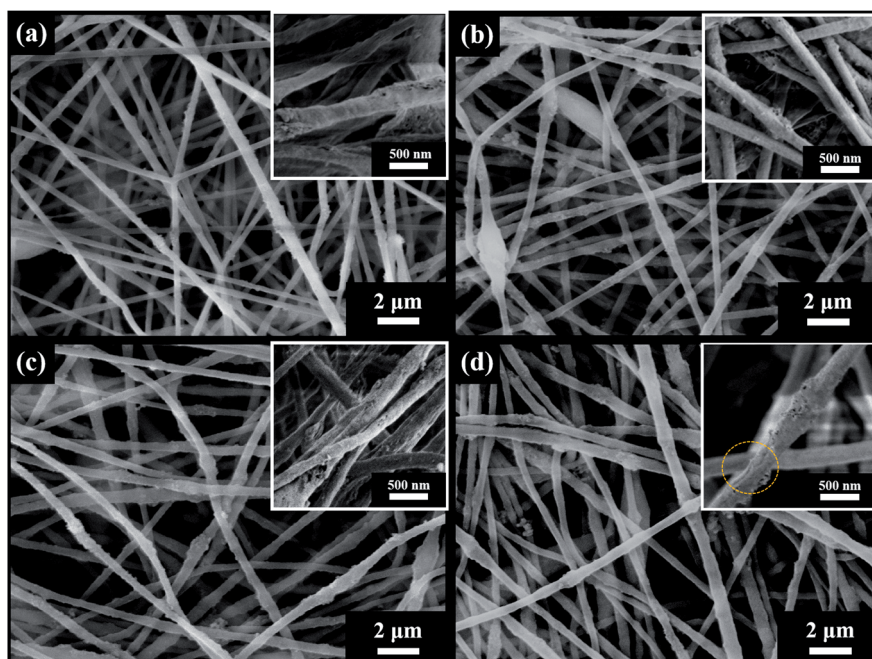


Fig. 1 Microscopic structure of PAN-SiO₂ fiber membranes with different treatment times: (a) 0 h, (b) 0.5 h, (c) 1 h, and (d) 1.5 h.

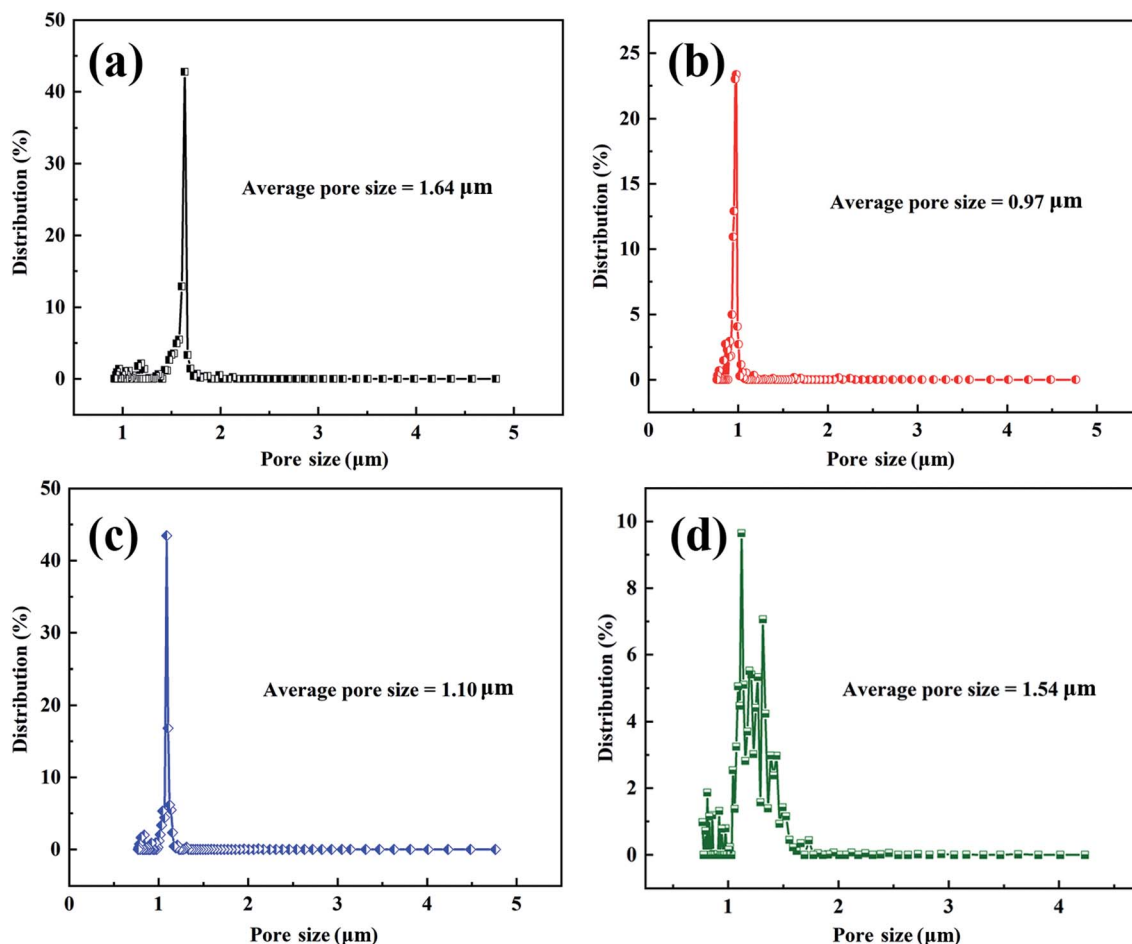


Fig. 2 Pore size distribution of the PAN-SiO₂ fibrous membranes after various alkali treatment times of (a) 0, (b) 0.5, (c) 1 and (d) 1.5 h.

an increase in the treatment time, indicating the quick drying property of the membranes, which is attributed to the wider wetting area of the hydrophilic nanofiber membrane.^{25,27}

Fig. 4c depicts the apparent water contact angle of a droplet on the surface of PAN-SiO₂ nanofibers with different treatment times. The water absorption times for TPAN-SiO₂-0.5, TPAN-SiO₂-1, TPAN-SiO₂-1.5 had no significant difference.

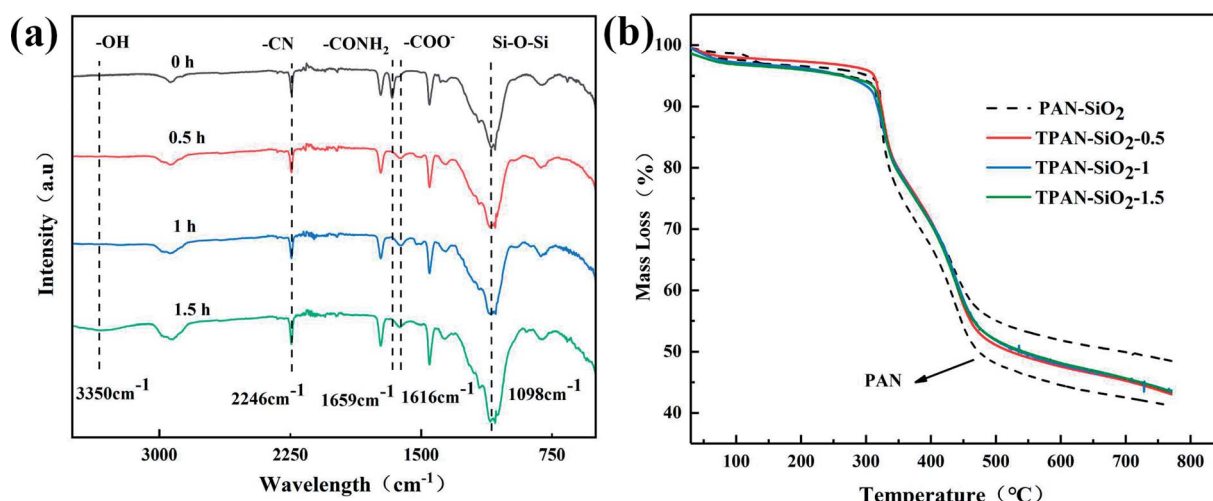


Fig. 3 FTIR spectra (a) and TGA curves (b) of the fibrous membranes.

Table 1 Parameters of the different samples during the thermal decomposition^a

Sample	T_d (°C)	α (%)
PAN-SiO ₂	321.6	48.5
PAN-SiO ₂ -0.5	326.1	43.5
PAN-SiO ₂ -1	326.3	43.4
PAN-SiO ₂ -1.5	326.5	43.3
PAN	324.3	39.6

^a T_d the thermal decomposition temperature at the maximum weight loss rate of material. α weight residue at 800 °C.

Nevertheless, the WCA of PAN-SiO₂ was different from that of the other samples, and the water droplets could be absorbed and diffused completely by the nanofiber membrane after 13 s. This indicated that the hydrophilicity of the PAN nanofibers was enhanced due to the hydrophilic SiO₂ nanoparticles (PAN-SiO₂). Thus, the nanofiber membranes treated by dopamine alkaline solution contained hydrophilic groups on their surface, and the original hydrophilic SiO₂ particles did not completely fall off

and still existed, leading to a further improvement in the water absorption on the surface of the membrane.²⁴ At the same time, the wettability of the PP layer was also improved in alkali solution of dopamine (FS 4†).

3.2 Effect of treatment on water transport behavior in the bilayered fibrous mats

To investigate the effect of the treatment on the water transport behavior in the bilayered fibrous mats, the change in contact angle with time for the PAN-SiO₂/PP bilayered fibrous mats treated at different times (0 and 1.5 h) was examined (Fig. 5). The initial WCA of PAN-SiO₂/PP-0 h was 130.5°, which was almost the same as that of the single PP layer, and the droplet could not diffuse after 300 s. After the bilayered fibrous mat was treated for 1.5 h with dopamine alkaline aqueous solution, the initial contact angle of the water droplet decreased to 57.9° and it was completely absorbed in about 3 s. This phenomenon is attributed to the wettability of the PP layer and the significantly weakened hydrophobicity after dipping in dopamine alkaline aqueous solution. The hydrostatic pressure of the water droplets was greater than the hydrophobicity of the PP layer; thus,

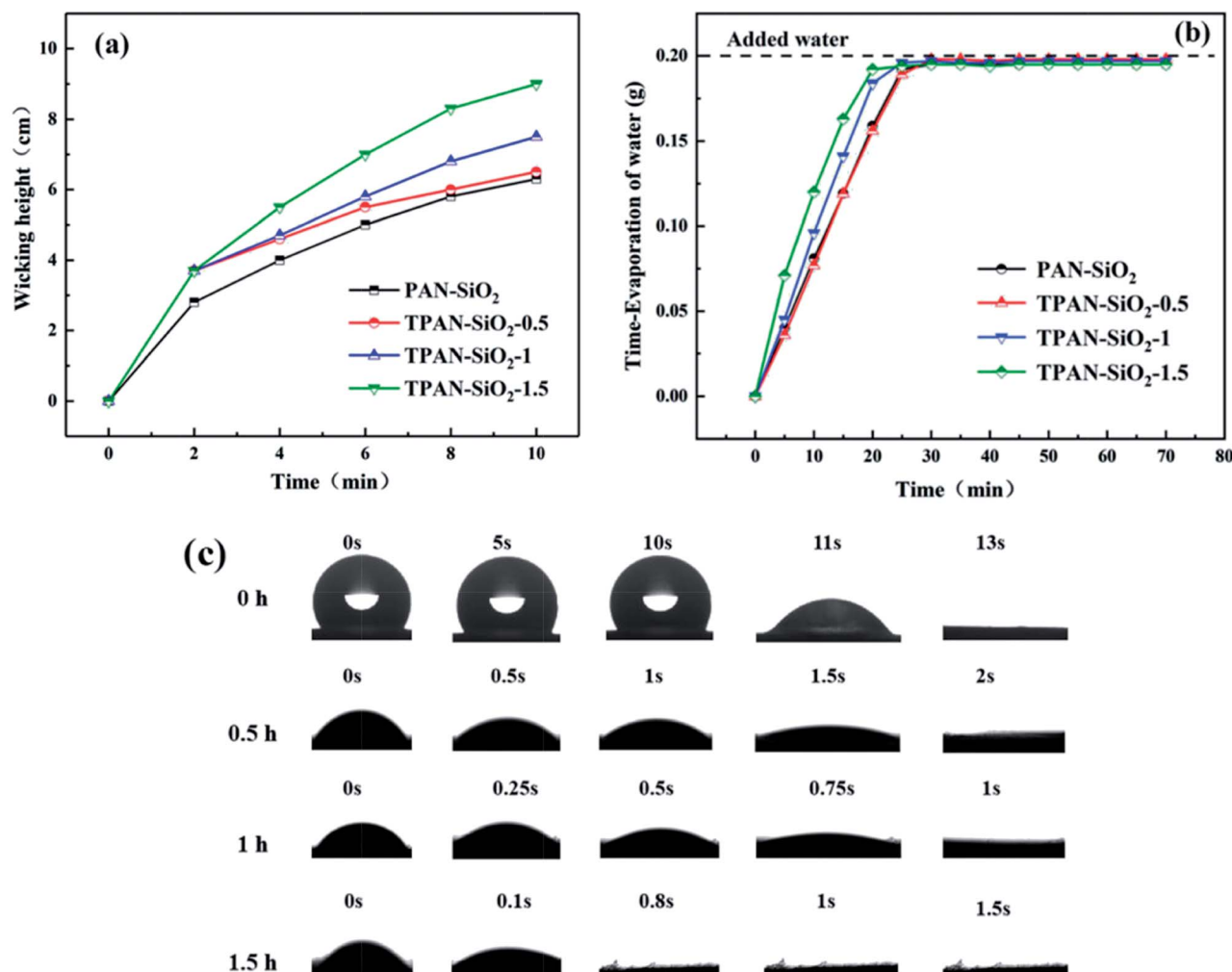


Fig. 4 Wicking height (a), water evaporation rate (b), and apparent WCA of PAN-SiO₂ (c) with different treatment times.



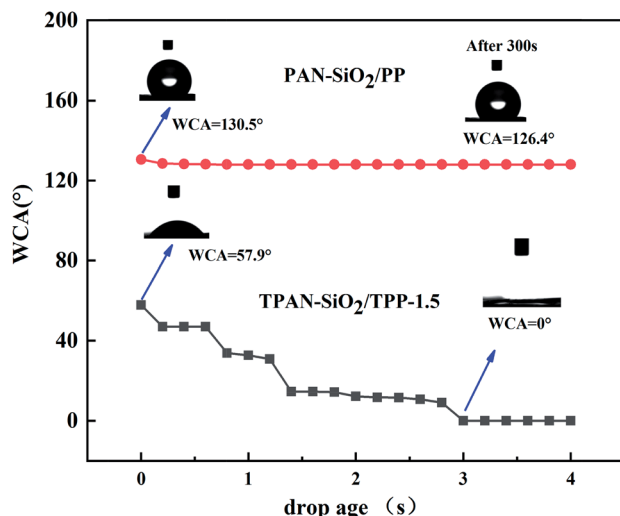


Fig. 5 Apparent WCA of the PAN-SiO₂/PP bilayered fibrous mats with 0 and 1.5 h treatment times.

the water droplet could penetrate the PP layer vertically. Meanwhile, the wicking property of the PAN-SiO₂ layer improved, and the water droplet was pulled by the hydrophilic layer quickly.

Fig. 6a and b demonstrates the directional water transport of PAN-SiO₂/PP and TPAN-SiO₂/TPP, which was quantitatively measured using a liquid moisture management tester (MMT). 0.2 g of saltwater was continuously dropped on the surface of the composite fibrous mats for 120 s. The blue line in the figure represents the relative water content at the top of the mat, while the red line represents the water content at the bottom.

The resultant AOTI value for PAN-SiO₂/PP and TPAN-SiO₂/TPP were -852% and 330% , respectively. The water transport behavior was promoted after treatment in alkaline aqueous solution of dopamine due to the improvement in the wettability of the treated PP fibers. The observed images show that when

a saltwater droplet was dropped on the PAN-SiO₂/PP, the majority of water stayed on the surface of the PP layer and hardly transported to the opposite side. This is because the surface energy was so small that the PP layer was endowed with strong hydrophobic property; hence, the water droplet could not contact the hydrophilic layer (PAN-SiO₂) and pulled out by capillary. However, the water content at the bottom was significantly larger than that at the top when a certain amount of water was dropped on TPAN-SiO₂/TPP. Initially, the water content at the top increased faster than that at the bottom, and reached the maximum at the point of 10 s. This is attributed to the relatively strong initial hydrophobic force on the TPP side, which slowed the downward transport of water droplets. The water content at the bottom of the layer increased steadily with time and exceeded that at the top at the end of 37 s owing to the water being pulled out by the TPAN-SiO₂ layer under the remarkable wicking action, which accelerated the rate of wetting in the bottom area. The bottom water content did not increase significantly after 80 s, and remained at a stable level. Nevertheless, when the composite mat was reversed and the saltwater droplet was added to the TPP-SiO₂ side, all the water accumulated at the top without transporting downwards, making the bottom hardly wetted (FS 5a†).

When a suitable amount transparent blue ink droplets (35 μL) was on the TPP side, it was observed that the water was quickly transported from the TPP layer to the opposite side (Fig. 7a and b), showing the excellent wettability on the TPP side and capillary force on the TPAN-SiO₂ side. Also, it was observed that the water spreading area on the TPP side was small, because of a large amount of water was absorbed by TPAN-SiO₂ and diffused along the transverse direction. Thus, the experimental results prove that TPP has decent water transport ability and did not absorb the water, which meets the requirements of quick drying of inner layer to ensure thermal and humid comfort for the human body. TPAN-SiO₂ had remarkable wicking ability, and it could pull water out quickly and accelerate the rate of water transport. However, when water was

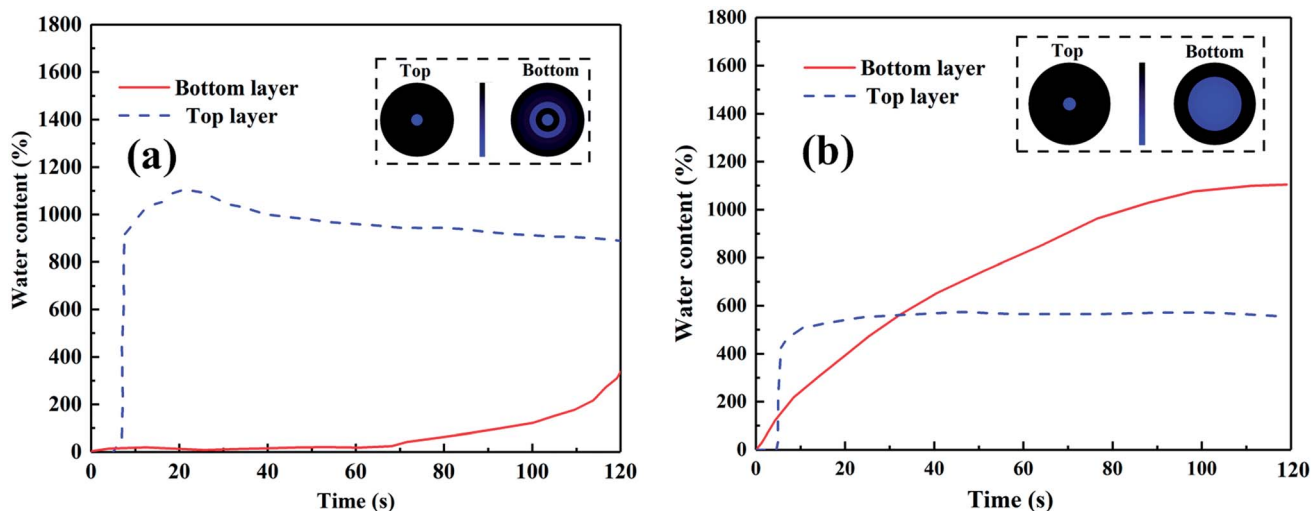


Fig. 6 MMT results of the PAN-SiO₂/PP bilayered fibrous mats with (a) 0 and (b) 1.5 h treatment time.

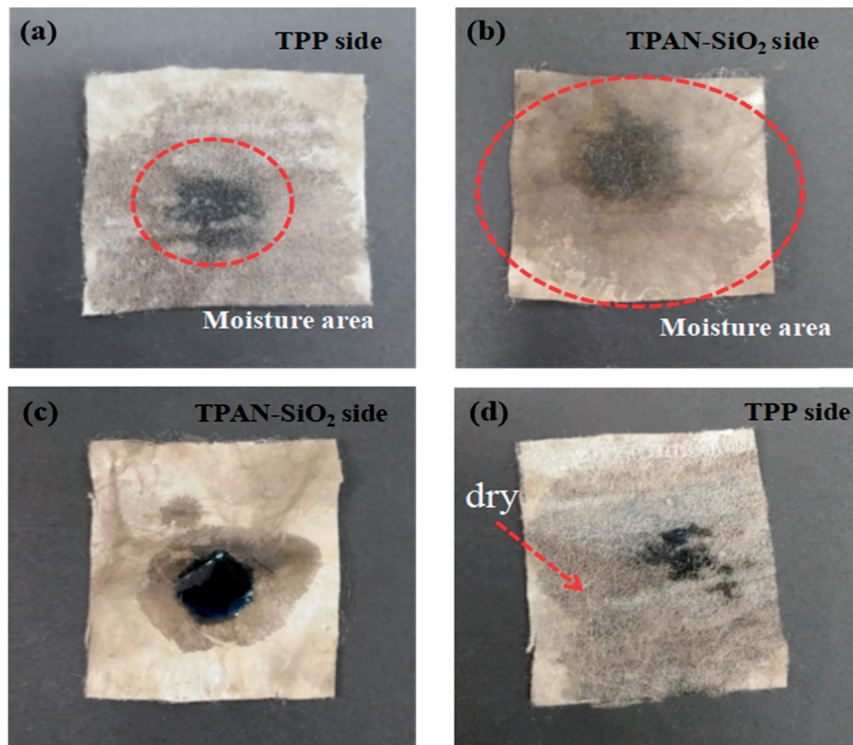


Fig. 7 Photographs showing the water spreading and transport abilities of the bilayered fibrous mats. Illustration of the directional water transport mechanism: Water is dropped on the TPP side (a and b) and TPAN-SiO₂ side (c and d).

dropped in the opposite direction (Fig. 7c and d), water accumulated on the surface of TPAN-SiO₂ and diffused rapidly along the transverse plane, while only a small amount of water was transported to the other side; thus, most areas of the TPP side remained dry. This phenomenon intuitively demonstrates that TPAN-SiO₂/TPP has outstanding directional water transport.

3.3 Effect of PP layer thickness on water transport behavior in the multilayered superfine fibrous mats

Multilayered superfine fibrous mats were prepared with different PP layer thicknesses by controlling the melt-electrospinning time, as shown in Table 2, and the effect of thickness on their directional water transport behavior was investigated.

Fig. 8a–d present the water transport behavior and quantitative analysis of the amount of water retained in the top and bottom layers of TPAN-SiO₂/cotton mesh/TPP with various thicknesses. The resultant AOTI values for the obtained multilayered superfine fibrous mats were 861%, 1071%, 841%, and 352%. The water content in the TPP layer in TPAN-SiO₂/cotton mesh/TPP-1 was almost zero and the hydrophilic layers of all the samples were completely wetted by water after the cotton mesh was sandwiched, which indicate the cotton mesh acts as a transfer layer to accelerate water transport.

When the time of melt-spinning was 0.5 h, the thickness of the PP layer was 0.11 ± 0.01 mm. At this time, the water droplet experienced a weaker hydrophobic force because the PP layer was relatively thin. Thus, the water droplet could easily pass through the hydrophobic layer and be absorbed by the

Table 2 The thickness of every single layer and multilayered superfine fibrous mats with different melt-electrospinning times

Sample	Sample ID	Average thickness (mm)
Single-layer sample		
TPAN-SiO ₂		0.074
Cotton mesh		0.33 ± 0.02
TPP-0.5		0.11 ± 0.01
TPP-1		0.34 ± 0.03
TPP-1.5		0.54 ± 0.03
TPP-2		0.71 ± 0.04
Multi-layer sample		
TPAN-SiO ₂ /cotton mesh/TPP-0.5		0.52 ± 0.03
TPAN-SiO ₂ /cotton mesh/TPP-1		0.74 ± 0.01
TPAN-SiO ₂ /cotton mesh/TPP-1.5		0.93 ± 0.01
TPAN-SiO ₂ /cotton mesh/TPP-2		1.15 ± 0.03



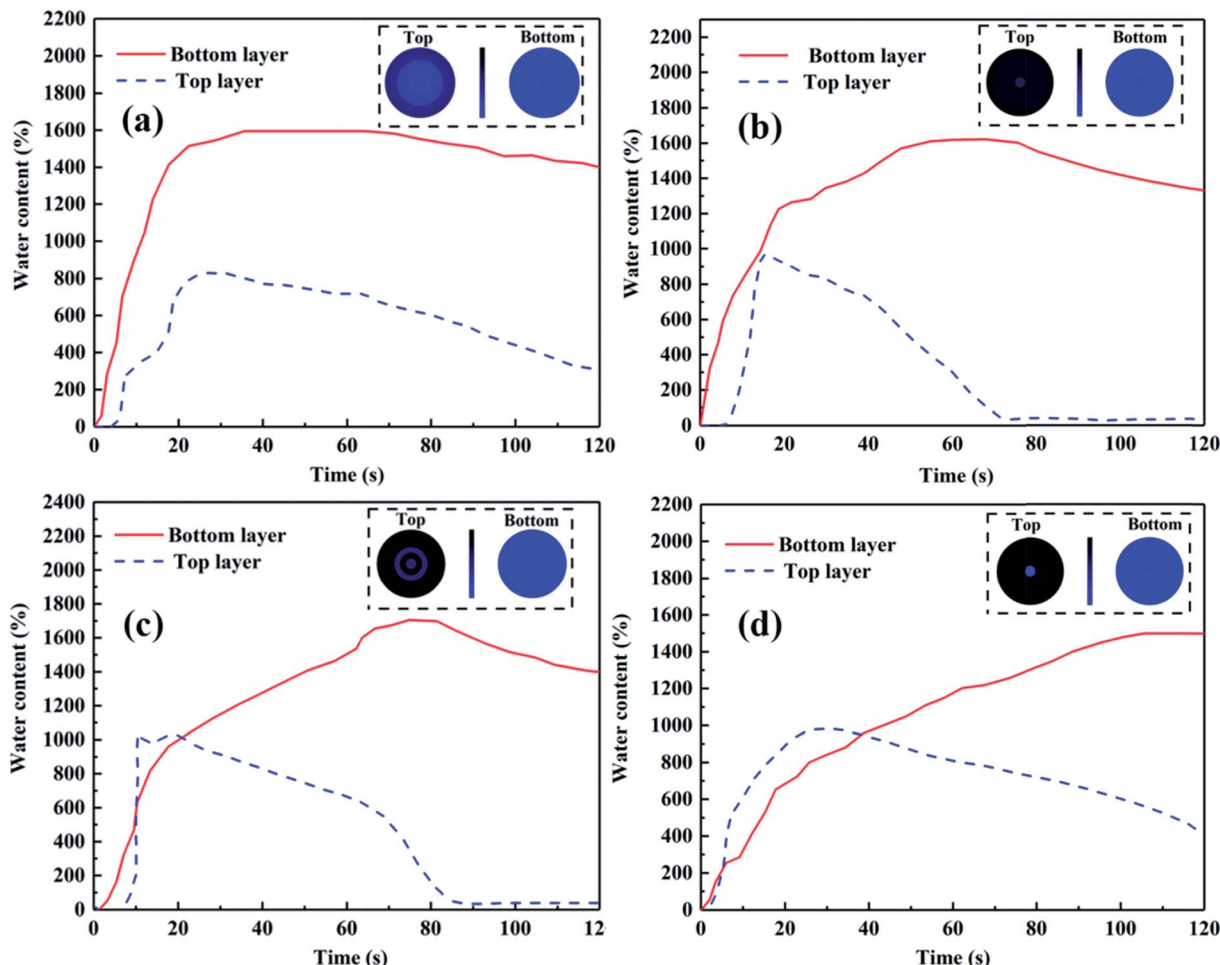


Fig. 8 MMT results for the TPAN-SiO₂/cotton mesh/TPP with the 0.5 h (a), 1 h (b), 1.5 h (c), 2 h (d) melt-electrospinning time.

hydrophilic side. However, the water droplet easily diffused transversely under the action of the cotton mesh in this case, making the PP layer wetted and affecting the directional water transport of the material, as shown in Fig. 8a. When the spinning time was extended to 1 h, the thickness of the PP layer reached 0.34 ± 0.03 mm. The water droplets could be absorbed by the outer layer and the inner layer kept the dry state for 120 s, showing an excellent water transport effect, as shown in Fig. 8b. Moreover, the composites still exhibited an excellent wearability performance (FS 6 and FS 8[†]).³⁵ Continuous prolongation of the spinning time to 1.5 h and 2 h caused the thickness of the PP layer to increase constantly. The water droplet was subjected to a greater hydrophobic force in the inner layer, which limited the transport of water in the material, thus affecting the overall performance of the directional water transport (Fig. 8c and d, respectively).

3.4 Directional water transport mechanism in the multilayered superfine fibrous mat

Based on the above results, we discuss the mechanism of directional water transport. As illustrated in Fig. 9a, when a water droplet was dropped on the hydrophobic layer side (TPP

side), the droplet was subjected to two opposite forces, namely, hydrostatic pressure (HP) and hydrophobic force (HF). HF impeded the downward transport of the droplet to wet the fibrous mat, and the effect of HF was related to the nature of the water itself, being proportional to the weight of the droplet. When the HP was larger than the HF, the droplet would penetrate the hydrophobic layer and contact the hydrophilic layer (TPAN-SiO₂ side), which would be accelerated by the capillary force provided by the hydrophilic layer (TPAN-SiO₂ side). When the droplet was dropped on the other layer (TPAN-SiO₂ side), the water droplet would absorb and transverse diffuse quickly on its surface. After the water droplet reached the contact plane between the two layers, the hydrophobic force of the opposite hydrophobic layer (TPP side) would prevent the water from continuing to penetrate the other side, thus effectively reducing the wetting area of the hydrophobic layer (next to the skin).^{29,31}

Fig. 9b depicts the mechanism of directional water transport in the multilayered superfine fibrous mat. When the water droplet was dropped on the hydrophobic side, it still retained a Wenzel–Cassie state subjected to two forces in opposite directions, hydrostatic pressure (HP) and hydrophobic force (HF), which is the same as mechanism in the bilayered fibrous



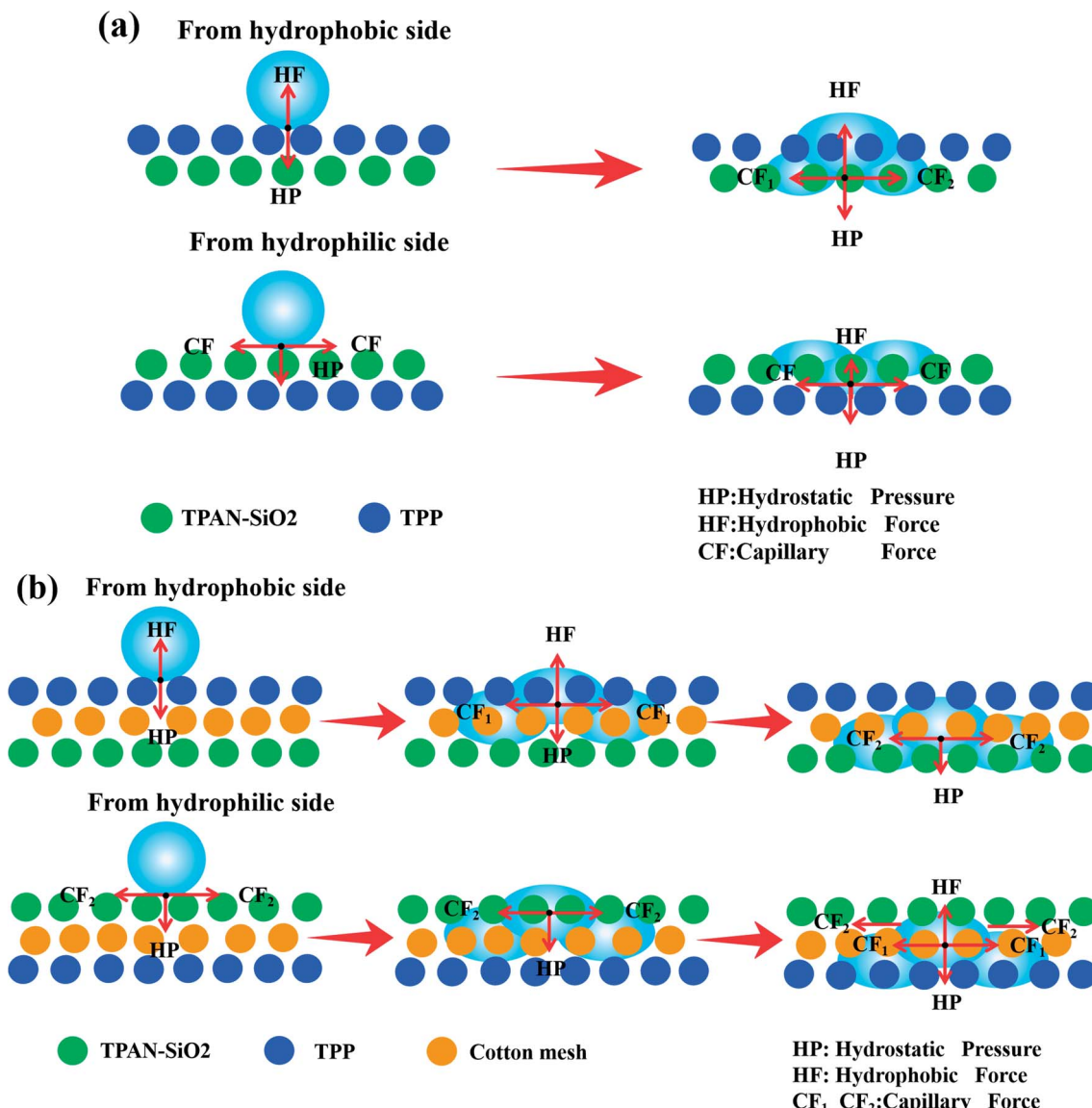


Fig. 9 Illustration of the directional water transport mechanism in the bilayered fibrous mats (a) and multilayered superfine fibrous mats (b): water is dropped on the upward hydrophobic TPP side and TPAN-SiO₂ side, respectively.

mat. When the water droplet moved downward, it would be continuously attracted by the capillary force (CF_1 and CF_2) provided by the cotton mesh and TPAN-SiO₂ successively, resulting in progressive wettability to speed up the water droplet transport to the opposite side, thus increasing the diffusion area of water on the hydrophilic side, as shown in the inserted images in Fig. 6 and 8. On the other hand, when the water droplet was on the hydrophilic layer side, it would be absorbed rapidly by the cotton mesh and TPAN-SiO₂ and diffuse horizontally. When water droplets contacted the TPP layer, they suffered from hydrophobic force in the upward direction, preventing the block inner layer from being wetted. Interestingly, insignificant areas of the inner layer in TPAN-SiO₂/cotton mesh/TPP were wetted by water due to the close combination of PP fibers and cotton mesh under the action of electrostatic force (FS 5b†), but large areas remained dry in the TPP layer.

4. Conclusion

In summary, the TPAN-SiO₂/cotton mesh/TPP multilayered superfine fibrous mat prepared by dual-mode electrospinning technology and subsequent treatment meets two types of water transport requirements for materials used in the field of functional textiles. The wettability of its inner layer and hydrophilicity of its outer layer were improved by dipping it in a dopamine alkaline aqueous solution, which weakened the initial hydrophobic force for water transport. Through MMT test, the multilayered superfine fibrous mats exhibited obvious directional water transport ability, exhibiting outstanding AOTI (1071%) and excellent OMMC (0.88) values, and when they were flipped, the AOTI value changed to -1332%, exhibited a reasonable performance of water directional water transport. The TPAN-SiO₂/cotton mesh/TPP multilayered superfine fibrous

mat exhibited decent strength and excellent permeability while remaining soft, which give it further practical application value.

Interestingly, in during the experiment, we found that when spinning the PP fiber layer by melt-electrospinning with cotton mesh as the substrate, the PP fibers and cotton yarn were attached relatively close due to the electrostatic force interaction between them. The cellulose fiber also had good hydrophilicity, which meets the requirement of directional water transport. Thus, we will subsequently conduct in-depth research on this material.

Conflicts of interest

There are no conflicts to declare.

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