1. Introduction

With the rapid development of global industrialization, haze weather conditions are becoming increasingly serious worldwide, which has attracted significant attention from the public. According to a series of reports released by the UN Environment Programme, haze causes seven million deaths every year worldwide; among them, 4.3 million people die of indoor air pollution [1]. Among air pollutants, particulate matter (PM), especially PM2.5 (particles with a diameter less than or equal to 2.5 μm), is the most harmful to humans. These particles can combine with other substances in the air and remain airborne for extended periods. They can be easily inhaled and may cause allergies or respiratory infectious diseases [2-4]. Considering the long-term indoor activities of people, reducing indoor PM2.5 is an effective method to protect human health. At present, the most widely used air purification method is to remove PM2.5 particles using air-cleaners. However, air-cleaners only circulate and filter indoor air and cannot supply fresh air; thus, opening windows for ventilation is still required [5, 6]. Mashes mode of glass fiber, metal, etc. are commonly used window screen materials. However, the mesh of these window screens is on the millimeter scale, which can block mosquitoes and other small insects but cannot block PM2.5 [7]. The application of an ultra-fine fiber film as a window screen enables the appropriate filtering of PM2.5 while maintaining sufficient air permeability and light transmittance [8, 9].

Electrospinning is the most common and effective method for preparing nanofibers. Electrospun nanofiber membranes have the advantages of small pore size, high porosity, and good channel connectivity [10, 11]. This enables the reduction of air permeability of materials; thus, it is considered to have great application potential in air filtration [12, 13]. However, the strength of pure
nanofiber membranes cannot meet the realistic requirements; thus, nanofiber membranes need to be combined with other substrate materials using composite technology.

In this study, a composite window screen with a sandwich structure was prepared, which consisted of a polyacrylonitrile (PAN) nanofiber membrane as the filter layer, a 30 mesh glass fiber mesh as the supporting layer, and a 120 mesh polyester fiber mesh as the protective layer. We studied the effect of different composite methods and composite parameters on the performance of anti-haze window screens and determined the optimal composite technology and parameters. This work aims to provide a solid foundation for the industrial production of nanofiber anti-haze window screens.

2. Experimental Section

2.1 Materials

Polyacrylonitrile (Mr = 150,000) was provided by Tianjin Kemiou Chemical Reagent Co., Ltd., China, lithium chloride (LiCl) was provided by Tianjin Aopusheng Chemical Co., Ltd., China; N,N-dimethylformamide (DMF) was obtained from Xilong Scientific Co., Ltd., China, and polyurethane hot melt adhesive (PUR, particle size 75 µm) was supplied by the Shenzhen Renshan New Material Technology Co., Ltd., China. Both the 30 mesh glass fiber gauze and the 120 mesh polyester gauze were obtained from Shanghai Zhengxi New Material Technology Co., Ltd., China. All chemicals were of analytical grade and used without further purification.

2.2 Electrospinning

In the early stage of the study, we found that the optimal spinning concentration of PAN was 12% [14]. First, different LiCl amounts were dissolved in DMF and stirred for 12 h at 60 ℃. Then, quantitative PAN was added to the LiCl/DMF solution, heated, and stirred for 24 h until the PAN was completely dissolved. Finally, a series of solutions with LiCl contents of 0 wt%, 0.01 wt%, 0.02 wt%, and 0.03 wt% were prepared.

Nanofiber membranes were prepared by linear electrode electrospinning using a 120 mesh polyester fiber mesh as the receiving base fabric, as shown in Fig. 1. The mesh cord speed was set to 150, 200, 250, 300, and 350 mm/min to obtain nanofiber membranes with different thicknesses. The spinning parameters were the same as those in our previous work: spinning voltage of 35 kV, spinning pitch of 18 cm, temperature of 25 ℃ (±3 ℃), and humidity of 30 %

![Fig. 1 Schematic diagram of the electrospinning setup](image)

![Fig. 2 Schematic diagram of the ultrasonic bonding machine](image)
(±5 %).

2.3 Ultrasonic composite process

To composite the polyethylene terephthalate (PET) fiber mesh covered by nanofibers and the glass fiber mesh (Fig. 2), an ultrasonic bonding machine was used when the feeding speed of the compound reached the set value. The effects of different composite pressures (0.5, 1.5, 2.6, 3.9, and 6 MPa) on the peel strength and filtration efficiency of the window screen were studied. The compounding time was 0.4 s and the holding pressure time was 0.6 s.

2.4 Hot melt adhesive composite process

The PUR was coated on the surface of the glass fiber mesh using hot melt adhesive equipment and then bonded with the PET-fiber mesh with a nanofiber membrane, as shown in Fig. 3. Using this method, we prepared anti-haze window screens with different amounts of glue (0.4, 0.8, 1.2, 1.6, and 2 g/m²), and studied the effects of the amount of glue on the peel strength and filtration performance. In addition, we studied the effect of PUR curing times (4, 8, 12, 24, 36, and 48 h) on the peel strength. The recombination time and pressure were 13 s and 0.3 kPa, respectively, and the compound velocity was 5 m/min.

2.5 Scanning electron microscopy test

Scanning electron microscopy (SEM, Pw-100-515, Shanghai Funa Scientific Instruments Co., Ltd., China) was used to observe the morphology of the fiber membrane. The nanofiber membrane was applied to the sample table by a spraying current of 10 mA and scanning voltage of 10 kV. Fifty samples were randomly tested using a nanofiber measurer, and the average values were obtained.

2.6 Filter performance test

The filter performance of the nanofiber membrane was analyzed using an automatic filter material tester (8130A, TSI Incorporated, USA) at an air flow rate of 32 l/min. The sample was cut into a circle with a diameter of 15 cm, and the average value of ten tests for each sample was determined.

2.7 Peel strength test

A laminated ultra-thin fiber membrane automatic peel strength tester (QJ210A, Jinan Languang Electromechanical Technology Co., Ltd., China) was used to measure the peel strength, referring to the Chinese Standard GB/T 2790-1995. The specific peel strength test method was flexible versus rigid material. The sample was cut to a length of 260 mm and width of 50 mm, the stretching speed was set to 500 mm/min, and the stretching distance was 250 mm [15]. In this experiment, the peel force at the maximum peak value was used as the value for calculating the peel strength, and the average value was obtained after five tests. The peel strength was calculated as

\[ \sigma_{180^\circ} = F / B, \]

where \( \sigma_{180^\circ} = 180^\circ \) is the peel strength, \( F \) is the maximum peak peel force, and \( B \) is the width of the sample.

3. Results and discussion

The grid size of an ordinary window screen is approximately 1-20 mm (Fig. 4(a)), which can only block dust and mosquitoes; however, it provides no filtration for haze compounds with PM2.5. The core layer of a nanofiber composite window screen is a nanofiber non-woven membrane with pore sizes in the range of 0.5-2 µm and with a three-dimensional structure (Fig. 4(b)), which has a good filtering effect for PM2.5.

3.1 The effect of LiCl concentration on the diameter of PAN nanofibers

The pore size of the nanofiber membrane is directly related to the fiber diameter, and the conductivity of the solution is one of the most
important process parameters affecting the diameter of the electrospun fibers. Therefore, the diameter of the electrospun nanofibers is typically adjusted by the addition of an electrolyte [16]. In this study, LiCl was chosen for the adjustment of the conductivity of the spinning solution, and the effect of the fiber morphology was further investigated. As shown in Fig. 5, with the addition of LiCl, the content of the finer fibers increased, the diameter distribution was concentrated at around 150-200 nm, and the average diameter decreased significantly. At the LiCl content of 0.02 wt%, the diameter distribution of nanofibers was the narrowest, and the concentration was approximately 150 nm, as shown in Fig. 5(c). However, as the amount of LiCl was gradually increased, the fiber diameter and dispersion increased, and there was a bonding phenomenon between adjacent fibers, forming a rod-like structure, as shown in Fig. 5(d). However, with the excessive addition of LiCl, the higher salt content increased the viscosity of the solution, resulting in incomplete fiber splitting and twinning, which increased the dispersion of the fiber diameter. Therefore, in this study, the optimum LiCl content was determined to be 0.02 wt%.

3.2 Effect of receiving net speed on the filtration efficiency of the nanofiber

To effectively block fine particles, the three-dimensional gap between fibers needs to be sufficiently small, while the bulk density of the fibers in the non-woven membrane directly affects the pore size. In this study, the bulk density of the nanofibers was adjusted by the running speed of the electrospinning receiving mesh, and its effect on the filtration efficiency of the fiber membrane was analyzed [17]. As can be seen in Fig. 6, the filtration resistance decreased gradually with the increase in feeding speed, and their relationship was nearly linear. However, the change in the filtration efficiency was completely different. At a running speed less than 250 mm/min, the filtration efficiency of the nanofiber membrane was maintained above 90%. When the running speed was greater than 250 mm/min, the filtration efficiency decreased rapidly. The bulk density of the fiber membrane at different feeding speeds could clearly be observed by SEM. At a running speed of 150 mm/min, the nanofibers were densely packed and the pore sizes of the fiber membrane were in the range of 1-5 µm, as shown in Fig. 7(a). Thus, the filtration efficiency was maintained at approximately 94%. At a feeding speed of 350 mm/
The pore size between the nanofibers increased to nearly 40 µm, and the filtration efficiency was decreased to approximately 72%. Considering realistic demand and production efficiency, the running speed of the curtain was set to 250 mm/min.

### 3.3 Effect of ultrasonic composite parameters on the performance of the window screen

The ultrasonic compound process involves the changing of the mechanical energy of the high-frequency vibration into heat energy [18, 19], which results in the thermal adhesion of the thermoplastic polymer. Owing to its advantages including high efficiency, pollution-free operation, and simple equipment, it is one of the most widespread composite methods in the field of non-woven industry [20].

Fig. 5 SEM images and diameter distribution diagrams of different LiCl contents: (a) 0 wt%, (b) 0.01 wt%, (c) 0.02 wt%, and (d) 0.03 wt%

Fig. 6 Effect of receiving wire speed on the filtration performance of the PAN nanofiber membrane

Fig. 7 SEM image of the PAN nanofiber membrane with a receiving network line speed of (a) 150 mm/min and (b) 350 mm/min
anti-haze window screen designed in this study consists of three layers: a glass fiber mesh, a nanofiber membrane, and a polyester fine mesh. First, ultrasonic composite technology was used to bond window screens, and we studied the effect of ultrasonic composite parameters on its performance to assess the applicability of ultrasonic composites in this study. As shown in Fig. 8(a), the peel strength of the composite window screen gradually increased with the increase in the composite pressure. When the composite pressure increased to 2.6 MPa, the peel strength reached 211.04 N/m that satisfied the requirements of practical applications. However, under this combined pressure, the filtration efficiency of the window screen decreased sharply. The SEM morphology showed that in this case, the front and back screens in the bonding point area were damaged, and the nanofiber membrane also had holes; thus, the filtration efficiency decreased rapidly. This suggests that the ultrasonic composite energy was too high to bond the nanofiber membrane materials.

### 3.4 Effect of hot melt adhesive composite parameters on the performance of the window screen

As discussed in the previous section, the ultrasonic process was not suitable for the adhesion of nanofiber membranes. Therefore, hot melt adhesive composite technology was employed to complete the bonding of the window screen. An environmentally friendly and high-performance adhesive resin, PUR [21–23], was chosen for this purpose. We studied the effect of the PUR amount on the performance of the composite window screen. The results showed that the peel strength increased with the increase of the coating amount. When the coating amount of PUR reached 1.2 g/m², the peel strength was 307.46 N/m. As the coating amount continued to increase, the peel strength remained nearly unchanged (Fig. 10(a)). The filter resistance of the composite window screen increased with the increase in the coating amount, but the filter efficiency remained constant. As the coating surface was a net structure, when all grid lines were coated with PUR, the bonding area saturated and the strength of the glass also became maximal. Therefore, the increase in the coating amount had little effect on the adhesion, as shown in Fig. 11(b). However, if the coating amount was too high, the glue flowed over the screen, resulting in a constant increase in the filter resistance, as shown in Fig. 11(d). Nevertheless, the...
entire bonding process did not damage the nanofiber membrane; thus, it had little effect on the filtration efficiency.

During the bonding process of the PUR adhesive, the –NCO group reacts with H2O in the air to form a cross-linking structure, which is a very slow process [24, 25]. Therefore, after the multi-layer mesh is coated with PUR and the layers are stacked together, they need to be held for a certain time to bond and solidify. In this study, the change in the peel strength of the composite window screen was analyzed as a function of storage time. As can be seen in Fig. 12, the peel strength of the window screen increased rapidly in 24 h with the increase in the storage time, the growth rate decreased sharply after 24 h, and after 36 h the strength was constant. Considering this, the optimal storage time of the window screen after the hot melt adhesive compound was set to 24 h.

4. Conclusions

In this study, a sandwich structure of a nanofiber anti-haze window screen was designed, which was composed of a 120 mesh polyester screen, a nanofiber membrane as the middle layer, and a 30 mesh glass fiber screen as the protective layer. The analysis of the spinning process showed that at a LiCl content of 0.02 wt%, the average diameter of the nanofibers was the smallest, and the distribution was higher.

In this study, we analyzed the effect of ultrasonic composite technology and hot adhesive compounding technology on the performance of an anti-haze window screen. The results show that glass fiber mesh, nanofiber membrane, and polyester fiber mesh can be bonded together by ultrasonic compounding. However, ultrasonic energy can destroy the nanofiber membrane at the bonding point, which decreases the efficiency of the window screen. In the hot adhesive
compounding process, with an increase in the coating amount, the peel strength of the window screen made of hot melt adhesive was improved, and the filtration efficiency was nearly constant and maintained at approximately 95%, but the filtration resistance increased. In this study, the optimal amount of glue was 1.2 g/m² and the bonding time was 24 h.

It can be seen that hot melt adhesive composite technology is a suitable composite process for fabricating nanofiber window screens, with simple technology and controllable parameters, which are beneficial to the industrial manufacturing of anti-haze window screens.

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