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# **Analysis of Membrane Fouling of nZVI@PVDF Using Infrared Spectroscopy-multivariable Curve Resolution Analysis**

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## *Authors' contributions*

*This work was carried out in collaboration among all authors. Author MZ wrote, reviewed and edited the manuscript. Authors MZ DC and ZR did data curation, formal analysis. Author YZ and XZ investigated the work. Authors MZ and XL did data visualization and wrote the original draft. Author SL conceptualized the study, did funding acquisition and searched for resources. Authors JM and HD did software analysis and supervised the study. Authors MZ and JM did data validation. Authors HD and JY did project administration. Author XZ performed the methodology. Author JY searched for resources and supervised the study. Author ZR did funding acquisition. All authors read and approved the final manuscript.*

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## **ABSTRACT**

Xanthan gum (XG) and humic acid (HA) belonging to dissolved organic matter are the main substances in membrane fouling, and their interaction affects the membrane fouling process. In this study, the fouling process of nZVI@PVDF membranes used to filter different concentrations of XG and HA were investigated by infrared spectroscopy-multivariable curve resolution analysis. The results showed that the interaction between polysaccharides and organic matter in water samples can affect membrane fouling. When the proportion of mixed pollutants in water samples changes, the degree of membrane fouling varied. The simulated water sample containing 25% XG caused the most serious membrane fouling, while the simulated water sample containing 60% XG caused the least membrane fouling. The results of ATR-IR combined with MCR-ALS showed that as the filtration time and the proportion of pollutants in water samples change, the dominant pollutants on the membrane surface will be different. For the simulated water sample with 25% XG and 75%HA, the main pollutant on the membrane surface was XG, while the main pollutant on the membrane surface of the simulated water sample with 60% XG and 40%HA was HA. In addition, the correlation between transparent extracellular polymers (TEP) and membrane fouling had also been verified in this study. The higher the TEP content, the more serious the membrane fouling.

*Keywords: Membrane fouling; nZVI@PVDF membrane; xanthan gum; humic acid.*

# **1. INTRODUCTION**

Facing the severe problems of water shortage and water pollution, membrane separation technology plays an important role in industrial wastewater purification and seawater desalination [1-4]. According to the pore size, the membrane can be divided into microfiltration membrane, ultrafiltration membrane (UF), nanofiltration membrane, and reverse osmosis membrane [5]. UF membrane technology has higher efficiency and lower energy costs because UF membrane's pore size ranges from 10 A to 1000 A and requires low-pressure filtration at 0.1 - 0.5 MPa [6]. Ultrafiltration (UF) membrane can remove bacteria and most viruses, which prevent water-related diseases and microbial death and improve effluent quality [7]. Wide-used polyvinylidene fluoride (PVDF) has excellent filmforming properties, stability, chemical stability, thermal stability, and mechanical strength, making it a suitable material for preparing porous separation membranes.

Recently, nanoscale zero-valent iron (nZVI) has been widely used in groundwater and industrial wastewater treatment due to its extremely small particle size, high specific surface area, and strong iron reducing power, showing excellent treatment effect [8]. However, nZVI particles have a strong aggregation trend, which is due to the strong magnetic attraction between particles [9]. Therefore, nZVI particles for water treatment will reduce the reactivity due to the reduced surface area. The combination of functional nanomaterials with PVDF membranes provides

an excellent opportunity to degrade pollutants, improve osmotic flux, and change hydrophilicity [10]. Pd/Fe nanoparticles (NPs) were immobilized in hydrophilically polyacrylic acid (PAA)/PVDF membrane to decrease the aggregation of Pd/Fe nanoparticles [11]. Nano Pd/Fe-based PVDF membrane had been successfully used in treating trichloroacetic acid [11], trichloroethylene [12], polychlorinated biphenyls [13], etc. In Our previous study, nZVImodified PVDF membranes (nZVI@PVDF membranes) were functional successfully prepared by a simple and easy-to-follow filterpress coating method and was applied in the ultrafiltration process to treat 2-chlorophenol (2- CP) wastewater [14]. Due to the hydrophobicity of the PVDF membrane, serious membrane fouling is more likely to occur, which shortens the service life of the membrane, resulting in higher costs [15]. Simple surface coating with nZVI can improve both hydrophilicity and have little effect on the internal structure of the membrane. However, this method is often easy to plug the membrane pores, resulting in the decrease of the average pore size and porosity of the separation membrane, which makes it difficult for water molecules to penetrate the separation membrane and the water flux decreases sharply [16]. However, there is little literature on the fouling performance of the nanomaterial-modified ultrafiltration membrane.

Membrane contamination is due to the accumulation of organic matter, inorganic matter, or organisms on the membrane surface and inside the membrane pores [17]. Among these dissolved organic matters (DOMs) in wastewater have been considered as a key factor in membrane fouling [18]. In general, humic substances, proteins, and polysaccharides that belong to DOMs have been identified as the principal foulants that could block the membrane pores, and reduce permeate flux [19]. Different properties of DOMs make them more difficult to solve the membrane fouling problem. Humic substances are the main component of DOMs and are considered to be one of the most serious membrane contaminants. Xanthan gum (XG), with unique rheological properties and good water solubility, is an extracellular polysaccharide of microorganisms with extensive functions [20]. XG molecules in a water environment are easy to combine with dissolved organic matter in water to form copolymers. The membrane contamination caused by the mixing of different organic pollutants is more complex due to its synergistic effect [21]. For example, the interaction between polysaccharides and proteins, such as humic acid (HA), may also increase membrane fouling during membrane filtration. In the process of studying the formation mechanism of membrane fouling, the mechanism of membrane fouling caused by the interaction among HA and XG is rarely reported. Notably, the release of iron ions will be accompanied by the use of nZVI@PVDF modified membranes, and the effect of the interaction between iron ions and organic matter on membrane contamination is worth studying.

Transparent exopolymer particles (TEP) is a gellike acidic polysaccharide widely distributed in the ocean and wastewater, whose role in membrane fouling has attracted wide attention [22]. TEP is closely related to membrane fouling for the following reasons. Firstly, TEP has a high viscosity, which makes TEP easy to attach to the membrane surface and form a hydrogel layer, resulting in large membrane resistance [23]. Secondly, TEP has a three-dimensional structure and a large specific surface area, which is colonized by microorganisms [22]. Finally, the deformable TEP can under pressure go through pores smaller than its particle size by bending, breaking, and dispersing [24]. To analyze the components of pollutants and explore the interaction between pollutants. IR-ATR (infrared attenuated total reflection) technology is adopted, which can reflect the structural information of pollutant functional groups in the process of membrane fouling, and its characterization of pore space distribution on the membrane surface is effective [25]. The main drawback of this method is that the spectrum on each pixel is

overlap different pollution components, so it is difficult to distinguish different components [26]. Multivariable curve resolution-alternating leastsquares analysis (MCR-ALS) assumes that the spectrum of each pixel in the image is a linear combination of a series of pure spectral signals. The pixel spectrum is preprocessed and decomposed into pure components of its contribution [27]. The analysis method considered the complete spectral profile to identify compounds and visualize their distribution in the samples, to classify different chemical regions [28]. MCR-ALS was used to characterize the characteristics of membrane fouling, and the results were rapid and easy to explain and evaluate membrane fouling from chemical and biological perspectives.

In this study, the analytical methods of IR-ATR and MCR-ALS were combined to build an infrared imaging platform for membrane fouling. The fouling of nZVI@PVDF modified membrane by HA and XG during ultrafiltration was studied for the first time. The identification of pollutants, the evaluation of membrane pore blockage, and the visualization of membrane surface pollution were carried out. The mechanism of membrane fouling caused by the interaction between HA and XG during the formation of membrane fouling was studied. Understanding TEP-induced modified membrane fouling mechanisms during ultrafiltration is of primary importance for developing effective fouling control and prevention strategies, and for improving membrane filtration performance.

# **2. MATERIALS AND METHODS**

# **2.1 Preparation of nZVI@PVDF Modified Membrane**

The nZVI preparation method used in this study was the liquid phase transformation method [13]. A certain amount of dried nZVI was weighed, transferred to anhydrous and anaerobic ethanol, and then put into a bottle. Ultrasonic was carried out in an ultrasonic cleaner to make nZVI dispersed evenly. Then the anhydrous ethanol containing nZVI was poured into the ultrafiltration cup, and the nZVI was uniformly coated on the PVDF membrane (Zhongke Ruiyang Membrane Technology Co., Beijing, China) surface by nitrogen pressure. The membrane was washed three times with anaerobic ultrapure water aeration for 30 min in advance, and finally, the nZVI@PVDF modified ultrafiltration membrane coated with nZVI was obtained. The coating

amount of nZVI used in this study was 2 mg/cm<sup>2</sup>, and the filtration area of the membrane was 36.32 cm<sup>2</sup>.

# **2.2 nZVI@PVDF Modified Membrane for Ultrafiltration Treatment of Simulated Wastewater**

1.0 g of humic acid powder and 0.4 g of NaOH powder were added to approximately 800 mL of deionized water. The mixture was stirred rapidly on a magnetic stirrer for about 1 hour until the humic acid powder was completely dissolved. The solution was then adjusted to a final volume of 1 L to obtain a stock solution of humic acid at a concentration of 1 g/L. 0.5 g of xanthan gum powder was added to approximately 400 mL of deionized water. The mixture was stirred rapidly on a magnetic stirrer until the large lumps of gum were dispersed. The solution was then placed in an ultrasonic cleaner to ensure complete dissolution, and the final volume was adjusted to 500 mL to obtain a stock solution of xanthan gum at a concentration of 1 g/L. The total concentration of HA and XG in the mixed solution was 100 mg / L, and the proportions of XG were 0 %, 25 %, 50 %, 60 %, 75 % and 100 %, respectively (Table 1). The ultrafiltration experiment was performed using a cup filter for dead-end filtration, and the specific experimental device was shown in Fig.1. Firstly, the nZVI@PVDF modified membrane was fixed in the cup filter, and then the simulated water sample was poured into the cup filter. The magnetic stirrer was used to continuously and slowly stir water samples. The upper end of the cup filter was connected with a nitrogen bottle, and the pressure was adjusted to 0.1 MPa when filtering. The mass of the filtrate was recorded at a time interval of 10 s through the data acquisition software provided by the electronic balance. The PVDF ultrafiltration membrane was used in the experiment and immersed in ultrapure water for 24 h before use.

#### **Table 1. Concentrations of substances in mixed water samples (total pollutant concentration of 100 mg/L)**



#### **2.3 TEP Characterization**

Alcian blue 8GX (0.02 %) was dissolved in 0.06 % acetic acid to prepare Alcian blue solution. The solution was filtered with a 0.05 μm pore water filter membrane to remove the influence of substances in Alcian blue on TEP test results. A total of 50 mL mixed solution of membrane pollution source was passed through 0.22 μm aperture of water system filter membrane under 0.2 bar pressure (keeping low pressure during filtration to protect TEP from destruction) and then dyed with 1 mL alcian blue solution for 5 s. The filter was washed with 1 mL ultrapure water. The filter membrane after dyeing was taken out and immersed in 5 mL of 80 % sulfuric acid for 2 h. During the immersion, the sample tube was shaken 3 – 5 times to ensure that alcian blue was completely dissolved in sulfuric acid. Finally, the absorbance of alcian blue in sulfuric acid solution was measured with a spectrophotometer at the wavelength of 612 nm using a 1 cm colorimetric dish. The average value of each sample was obtained after three repeated measurements. The concentration of TEP was finally expressed by the equivalent of xanthan gum per liter of water, namely mgXeq / L.

## **2.4 Characterization of Membrane Fouling**

#### **2.4.1 Membrane flux**

The calculation formula of membrane flux (J) is:

$$
J = V/T \times A \tag{2.1}
$$

In the formula, J is the membrane flux  $(m / s)$ , T is the sampling time (s), V is the volume of filtrate in the sampling time  $(m<sup>3</sup>)$ , A is the membrane filtration area  $(m^2)$ . The degree of membrane fouling in this study is characterized by the attenuation of specific flux  $(J/J_0)$ , where J represents the membrane flux at any time in the filtration process, and  $J_0$  represents the initial membrane flux in the filtration process. With  $J/J_0$ as the ordinate axis and the filtration time as the abscissa axis, the change of specific flux in the filtration process was plotted to reflect the change of membrane fouling.

#### **2.4.2 Infrared imaging platform for surface contamination of modified membrane**

The nZVI@PVDF membrane after filtration of simulated wastewater was dried at room temperature. The membrane surface was divided into 136 points (5 mm  $\times$  5 mm), and then the IR-ATR measurement (Nicolet iS10, Germany) was



**Fig.1. Diagram of ultrafiltration experimental setup.**

performed on each point. Test conditions: scan times: 32; resolution: 4; the wavenumber ranged from  $4000 \, \text{cm}^{-1}$ ~550 cm<sup>-1</sup>. The infrared data obtained from the test were aggregated into the spectral matrix and imported into Matlab R2018b software for analysis by using the MCR-ALS script program. After several iterations of the ALS algorithm, the data of 136 points can be decomposed into the infrared spectrum and concentration distribution of pollutants on the membrane surface. Finally, the infrared image of the polluted film was obtained by drawing with software Origin 2019b.

#### **2.4.3 SEM characterization**

A scanning electron microscope (FE-SEM-EDS, Quant 250 FEG, FEI Corporation). was used to observe and characterize the surface morphology of the film. The test conditions were as follows: 220 V  $(\pm 10\%)$ , 50 Hz. Working temperature:  $15 \sim 20$  °C. Working humidity: < 80%.

## **3. RESULTS AND DISCUSSION**

# **3.1 Effects of Different Mixing Ratios of HA and XG on Flux Attenuation**

In this study, the effect of the composition of HA and XG in simulated water samples (total concentration of 100 mg/L) on membrane pollution was explored by changing the content of XG (0%, 25%, 50%, 60%, 75%, 100%) within 1h filtration cycle. Fig. 2 showed the relationship between membrane-specific flux and filtration time of nZVI@PVDF modified membrane in the process of filtering simulated water samples under 0.1 MPa. The membrane flux of the simulated water sample decreased rapidly in the

first 5 min of filtration and then decreased slowly after the first 5 min. When the filtration time reached 30 min, the membrane flux remained basically unchanged. When the content of XG was 60%, the specific flux of the membrane decreased the least. This means that the membrane fouling was the lightest, and the specific flux after 30 min was 30.7%. When the content of XG was 25%, the specific flux of the membrane decreased most. The membrane fouling was the most serious obviously, and the specific flux after 30 min was only 14.5% When both XG and HA were present in the water samples, the membrane flux was lower than that of pure HA or XG, except for the simulated water sample with 60% XG. Moreover, the membrane flux decreased with increasing XG concentration. The experimental results indicated that there was an interaction between XG and HA, which affects the formation of membrane fouling.

In order to determine the interaction between organic matter and polysaccharides, the correlation between TEP of HA and XG mixed solution and membrane fouling was worth studying. The TEP values for different XG contents are shown in Table 2. The measured value of TEP decreased first and then increased with the increase of XG content in the mixed solution. When the content of XG in the mixed solution is 25%, the TEP measurement value is the highest, reaching 84.65 mg Xeq/L. When the XG content was 60%, the measured value of TEP was the lowest, only 31.91 mg Xeq/L. This result corresponds to the attenuation of membrane flux in Fig. 2, indicating that membrane pollution was positively correlated with TEP in water samples. Therefore, the TEP value can reflect membrane fouling to some extent.



**Fig. 2. Membrane flux attenuation of nZVI/PVDF modified membrane for filtering simulated water samples with different xanthan gum content**





# **3.2 Different HA and XG Mixing Ratios Correspond to Changes in the IR-ATR Spectrum**

Fig. 3 showed the infrared spectra of HA and XG pure components. It can be seen from Fig.3(a) that HA had a broad absorption peak at 3350 cm-1, representing O-H stretching vibration; the peak at 2970 cm-1~2850 cm-1 was related to aliphatic C-H stretching vibration; The peak at 1724 cm<sup>-1</sup> was caused by the stretching vibration of the C=O bond of the carbonyl group; the absorption peak at 1605 cm-1 was the aromatic C=C, COO-; the absorption at 1244 cm-1 was the C-O stretching of the carboxyl group Caused by vibration; 1027 cm-1 was related to carbohydrate CO stretching vibration [29]. It can be seen from Fig. 3(b) that XG had a broad absorption peak at 3275 cm-1 , which represented the -OH group; 1602 cm-1 and 1398 cm-1 belonged to the COOgroup.

The MCR-ALS method was used to decompose the original data measured by IR-ATR into component spectra and their concentration data. Four components can be obtained, namely XG, HA, PVDF, and nZVI. The correlation coefficient R2 obtained by iteration of each infrared spectrum data matrix can reach more than 99.99%, and the relevant characteristic peaks in the infrared spectrum of each component obtained can also correspond to the infrared spectrum of the pure component, indicating the MCR-ALS analysis The results obtained are very reliable.

The concentration data obtained from the analysis are mapped to 136 points on the membrane surface one by one. The HA and XG pollution distribution diagrams on the membrane surface after filtering the simulated water sample for 1 h under different mixing ratios are shown in Fig. 4 and Fig. 5. When filtering pure HA and XG solutions of the same concentration, as shown in Fig. 4 (a) and Fig. 5 (e), the residue of XG on the membrane surface was significantly greater than that of HA. EPS, composed of polysaccharides, proteins, and humic substances, had been considered to cause serious membrane fouling. While polysaccharides usually have the most serious effect on the decrease of membrane flux [30]. Therefore, XG had a higher pollution potential than HA. When the XG content was 25% and 75%, the main residual contaminant on the film surface was XG, which was consistent



**Fig. 3. Infrared spectra of HA (a) and XG (b)**



**Fig. 4. The surface of the membrane is contaminated by HA after filtering the simulated water sample for 1 hour under different mixing ratios of HA and XG. (XG content: a-0%, b-25%, c-50%, d-60%, e-75%)**



**Fig. 5. The surface of the membrane is contaminated by XG after filtering the simulated water sample for 1 hour under different mixing ratios of HA and XG. (XG content: a-25%, b-50%, c-60%, d-75%, e-100%)**

with the large TEP value in Table 1. When the XG content was 60%, the main contaminant on the membrane surface was HA, which was consistent with the result of the smallest TEP measurement. When the XG content was 50%, the main contaminants on the membrane surface were HA and XG. The results also showed that TEP has a certain connection with membrane fouling.

### **3.3 The Dynamic Change of the Film Surface Pollution Corresponds to the IR-ATR Spectrum**

To investigate the dynamic changes of the membrane surface pollution during the filtration process, two special XG contents (25%, 60%) were selected during the membrane flux attenuation process. The membrane surface with

different filtration times was characterized by infrared, combined with MCR-ALS analysis of the changes of the components on the surface of the membrane. Since the membrane flux tends to be flat or even basically unchanged after the filtration time reaches 20 min, the filtration time selected for the study of dynamic changes is 5 min, 10 min, and 20 min.

Fig. 6 showed the temporal and spatial distribution of pollutants on the membrane surface when the XG content was 25%. It could be seen from Fig. 6 that the pollutant distribution on the membrane surface was generally uniform. The concentration of HA on the membrane surface decreased with time, while the concentration of XG increased with time. When the filtration time was 5 minutes, the HA concentration on the membrane surface was slightly higher than that of XG. Therefore, the rapid decline of membrane flux in the first 5 minutes may be caused by the blockage of membrane pores by HA. When the filtration time reached 10 minutes, the membrane surface was obviously dominated by XG. When the filtration time reached 20 min, the XG concentration on the membrane surface was much higher than that of HA, which meant the influence of the pollution layer formed by XG on membrane fouling was significantly greater than that of HA.

Fig. 7 showed the dynamic changes of pollutants on the membrane surface when the XG content

was 60%. The fouling degree of the membrane surface with 60% XG content was lighter than that with 25% XG content, which corresponded to the attenuation of membrane flux. In the first 10 minutes of the filtration process, the concentration of HA or XG on the membrane surface changed little. When the filtration time reached 20 min, the concentration of pollutants on the membrane surface was higher than that before, and the concentration of XG was slightly higher than that of HA, indicating that XG had begun to form a pollution layer on the membrane surface.



**Fig. 6. HA concentration distribution on the membrane surface when the XG content is 25% and the filtration time is 5 min (a), 10 min (b), and 20 min (c). XG concentration distribution on the membrane surface when the XG content is 25% and the filtration time is 5 min (d), 10 min (e) and 20 min (f)**



**Fig. 7. The HA concentration distribution on the membrane surface when the XG content is 60% and the filtration time is 5 min (a), 10 min (b), and 20 min (c). the XG concentration distribution on the membrane surface when the XG content is 60% and the filtration time is 5 min (d), 10 min (e), and 20 min (f)**

# **3.4 Characterization of Fouling Morphology on the Membrane Surface**

To understand the dynamic changes of membrane surface pollution more intuitively in the filtration process, the simulated water samples containing 25% XG with the most serious attenuation of membrane flux were selected. The membrane surface was characterized by SEM after 5 min, 10 min, and 20 min of modified membrane filtration. The morphology of the membrane surface under different filtration durations is shown in Fig. 8. When the filtration time was 5min, the fouling layer on the membrane surface could be observed under 1000 times magnification. However, when the filtration time was10 min, there was no pore on the membrane surface, indicating that with the increase of filtration time, the fouling layer on the membrane surface would become denser. As shown in Fig. 8 (f), when the magnification was 20000 times and the filtration time was 20 min, the gel material could be obviously observed in the lower-left corner of SEM. This confirmed that XG could promote the formation of a gel layer on the membrane surface due to its gelation. Combined with the accumulation of HA on the membrane surface, the filtration resistance was increased, resulting in the decrease of membrane flux.

This study showed that the mixture of HA and XG could affect nZVI@PVDF membrane fouling. A surprising finding was that the most serious pollution occurred when 75% HA and 25% XG

are mixed, and the main pollutant on the membrane surface was XG after 1 h of the filtration cycle. From the fifth minute to twenty minutes, the concentration of XG was higher than HA. Similarly, the concentration of XG on the membrane surface was higher when pure XG and pure HA are filtered at the same concentration. These results showed that in the fouling process of the modified nZVI@PVDF membrane, the fouling layer formed mainly by XG on the surface, while the membrane pore blockage caused by HA had little effect on membrane fouling. XG has a high molecular weight and high viscosity at low concentration, which affected the filtration flux [31]. In addition, XG has gel properties and bridges between molecules, forming a three-dimensional gel network on the membrane surface and accumulating on the membrane surface [31]. Therefore, the macromolecular XG caused more serious pollution by blocking the membrane surface. In addition, due to the use of nZVI@PVDF modified membrane, the nZVI coated on the membrane surface will be beaded due to its magnetism, which will also promote the accumulation of XG on the membrane surface. The SEM images of the membrane surface in Fig. 8 confirmed that XG could promote the formation of a gel layer on the membrane surface due to its gelation. Combined with the accumulation of HA on the membrane surface, the filtration resistance was increased, resulting in the decrease of membrane flux and high TEP (84.65 mg Xeq/L). The concentration of pollutants on the membrane surface increased with time,



**Fig. 8. The SEM image of the membrane surface after the nZVI/PVDF membrane filters 25%XG simulated wastewater for different filtration times. (A) 5 min, (b) 10 min, (c) 20 min magnification is 1000 times; (d) 5 min, (e) 10 min, (f) 20 min magnification is 20000 times**

and the main pollutants on the membrane surface changed due to the composition of water samples and filtration time. Overall, in the process of nZVI@PVDF membrane fouling in the mixed solution of XG and HA, the main contributor to the pollution was the pollution layer formed by XG on the surface of the membrane, while the pore blockage caused by HA had little effect on the membrane fouling.

More importantly, lighter membrane fouling occurred at 60 % XG concentration. The main pollutant on the membrane surface was HA when the filtration time was 1 h. In contrast, after 1 hour of pure HA filtration, the membrane flux decayed seriously, but the concentration of pollutants on the membrane surface was low. The membrane fouling was related to the TEP content in water samples which the higher the TEP content was accompanied the more serious the membrane fouling and the higher the membrane flux attenuation. However, the concentration of TEP was not proportional to the amount of XG when XG is 60%. This indicated that the interaction between HA and XG had an impact on the formation process of TEP in water samples, resulting in a new change in membrane surface pollution. This was because HA may enter the membrane pore and block the membrane pore. In our experiment, nZVI was chain-like on the membrane surface, so the average pore size of the nZVI layer was much larger than that of the HA molecule, resulting in many HA molecules not being intercepted by nZVI, and HA molecules could still reach the membrane surface, blocking the membrane pores in the filtration process. When 60% XG was mixed with HA, the XG concentration attached to the membrane surface was higher in the fifth to twentieth minutes. Subsequently, the gel network formed by XG captured HA, which made HA stay in the fouling layer on the membrane surface. HA was intercepted by the pollutants on the membrane surface, so the main pollutants on the membrane surface became HA. The mixed solution of HA and XG will basically aggravate the membrane fouling of the nZVI@PVDF membrane, but there may be a certain proportion range, which can reduce the membrane fouling, and even there is an optimal proportion, which can greatly reduce the membrane fouling.

# **4. CONCLUSION**

In this study, we investigated the membrane fouling behavior of nZVI@PVDF membranes

when exposed to varying concentrations of xanthan gum and humic acid, two key dissolved organic matter components. Our findings demonstrate that the interaction between XG and HA significantly impacts the formation of transparent exopolymer particles, which directly correlates with the degree of membrane fouling. The results indicate that XG plays a predominant role in membrane fouling, with the most severe fouling occurring at lower XG concentrations (25%) and the lightest fouling at 60% XG. This suggests that an optimal XG-to-HA ratio can mitigate fouling, offering valuable insights for enhancing the performance of modified membranes in water filtration processes. Furthermore, the use of advanced techniques such as infrared spectroscopy and multivariable curve resolution (MCR-ALS) enabled us to accurately characterize the fouling mechanisms and pollutant distribution on the membrane surface. These findings provide a foundation for developing more effective fouling control strategies in ultrafiltration systems, particularly those involving<br>
nanomaterial-modified membranes. Future nanomaterial-modified membranes. Future research could explore the influence of other organic pollutants and further refine the optimal composition of dissolved organic matter to improve membrane longevity and filtration efficiency.

#### **DISCLAIMER (ARTIFICIAL INTELLIGENCE)**

Author(s) hereby declare that NO generative AI technologies such as Large Language Models (ChatGPT, COPILOT, etc) and text-to-image generators have been used during writing or editing of this manuscript.

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#### **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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