


 Cite this: *RSC Adv.*, 2021, 11, 18131

Preparation of a double-network hydrogel based on wastepaper and its application in the treatment of wastewater containing copper(II) and methylene blue†

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To reclaim and utilize wastepaper (WP), a WP/acrylamide double-network hydrogel (WP/PAM) was prepared to transform WP into efficient adsorbent for heavy metals and dye wastewater treatment. The structure and properties of the WP/PAM were characterized systematically by scanning electron microscopy (SEM), thermogravimetric analysis (TGA), swelling performance (SR), Fourier transform infrared spectrum (FTIR), and X-ray photoelectron spectroscopy (XPS). Batch experiments showed that the adsorption process of Cu(II) and MB followed the pseudo-second-order kinetic model and the Langmuir model. The maximum adsorption capacities of the WP/PAM for Cu(II) and MB were 142.2 mg g⁻¹ and 1714.5 mg g⁻¹, respectively. The adsorption mechanism of Cu(II) on the WP/PAM was related to ion exchange and complexation, while MB adsorption was driven by hydrogen bonding and electrostatic interaction. Besides, the WP/PAM performed well in treating simulated wastewater. The regeneration test indicated that the WP/PAM could be successfully reused after 6 cycles. This work provided an alternative choice for the recycling of WP and produced a potential adsorbent for the dye and heavy metals wastewater treatment.

Received 24th March 2021

Accepted 13th May 2021

DOI: 10.1039/d1ra02321g

rsc.li/rsc-advances

1. Introduction

In the wake of development in urbanization and industrialization, wastewater with heavy metals or dye from different industrial processes, such as printing, electroplating, and steel manufacturing, has been released into the environment and resulted in increasingly serious water pollution.^{1–3} Take dye wastewater as an example, about 10–20% of dyes will be directly released into water to form organic dye wastewater in the process of dye production and use.⁴ Meanwhile, some metallic salts (e.g., Cu(II), Mn(II), Co(II), Ni(II)) are required as ingredients for the synthesis of some metal complexing dyes and mordants in the printing and dyeing processes, which is the reason that

printing and dyeing wastewater is usually accompanied by heavy metal ions.⁵ The heavy metal ions, which cannot be collected or degraded,⁶ would persist in the water environment, eventually endangering human health.^{7,8} Among various methods for treating heavy metal and dye wastewater, adsorption is widely used because of its simple operation and high tolerance to toxic chemicals in water.^{9,10} But the relatively high costs of adsorbents with high-efficiency cause an economic burden to the practical application of adsorption.¹¹ Hence, it is necessary to continuously develop more efficient, stable, low-cost, and environmentally compatible adsorbents.

Wastepaper (WP) as the main component of municipal waste had caused considerable environmental problems.¹² In the process of developing and utilizing WP, some scholars found the pure WP could be converted into adsorbents to remove pollutants, but the adsorption capacity was unsatisfactory.^{12,13} Furthermore, polyamides¹⁴ and chitosan¹⁵ were adopted to combine with WP to improve the performance for the elimination of heavy metal ions. Whereas, the application of these composites was still limited in water pollution remediation due to their relatively low adsorption capacity, long preparation time, and complex preparation methods. To overcome the drawback, WP-based composites with better adsorption ability should be further studied and improved.

Double-network polymer hydrogel with two independent network structures retains the advantage of traditional

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/d1ra02321g

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hydrogels that rich in buried water,¹⁶ and superior mechanical properties that cause less prone to collapse than conventional hydrogels.¹⁷ It is also easy to prepare, and its monomers are widely available. So far, there are some double-network hydrogels using natural polymer (*e.g.*, cotton cloth,¹⁸ rice husk,¹⁹ and straw²⁰) as monomers have been prepared. They possess good potential application value in the adsorption material, because of their low cost, simple production, and excellent removal effect for heavy metals. WP is the most common and easily available natural polymer in our daily life. Nevertheless, there are few double-network hydrogels prepared by WP as monomer, not to mention the report about its application in wastewater treatment.

In this study, acrylamide (AM), which has the advantages of rapid gelation through polymerization and cross-linking reaction, was selected as another layer of the network to prepare double-network hydrogel based on the WP cellulose network. It was expected that the WP/acrylamide composite (WP/PAM) could achieve the characteristics of rich active functional groups, three-dimensional porous structure, large exposure of adsorption sites, and high removal rate of heavy metals and dyes in wastewater. The specific objectives were as follows: (1) synthesizing and comparing the WP/PAM with different WP content; (2) evaluating the adsorption performance of the WP/PAM on Cu(II) and MB through batch experiments in the single and binary system; (3) characterizing the properties of the WP/PAM and clarifying the mechanism of the WP/PAM hydrogel adsorption for contaminants removal.

2. Material and methods

2.1 Materials

WP was mainly collected from common white print paper discarded in the laboratory. The WP was first immersed with glacial acetic acid (AR, 99.5%) for 0.5 h to remove impurities on the surface (environmental temperature: >20 °C), until the WP surface was clean and free of impurities. The WP pretreated by glacial acetic acid was took out and cleaned it with pure water for several times until the pH of the cleaning water was neutral. Then, the WP was torn into uniform pieces of about 5 mm × 5 mm after natural drying for 48 h, and put the WP pieces into a self-sealing bag and kept it in an electronic constant temperature moisture-proof box (AD-50S ANDBON, China) for further use. The reagents mainly include urea (AR), sodium hydroxide (NaOH, AR), acrylamide (AM, AR), epichlorohydrin (ECH, AR), potassium persulfate (KPS, AR), tetramethylethylenediamine (TEMED, AR), *N,N*-methylenebisacrylamide (MBA, AR), *etc.* All chemical reagents were supplied by Sinopharm Chemical Reagent Co, Ltd. (China). In addition, all the solutions involved in this work were prepared with deionized water (18.25 MΩ cm).

2.2 Preparation of WP/PAM

The preparation of cellulose solvent according to Zhang's work:²¹ NaOH/urea aqueous solution with a mass ratio of 7 : 12 was prepared and pre-cooled at 0 °C for 1 h.

The brief preparation process was shown in Fig. S1.† Here were the details: different quantities (1, 2, 3, and 4 g) of pre-treated WP were immediately added into the NaOH/urea aqueous solution with stirring strongly for 5 min to result viscous WP solution with mass fraction of 1 wt%, 2 wt%, 3 wt%, and 4 wt%. Afterwards, 6 mL cellulose solution was mixed with 2 mL AM solution (0.014 mol), then 0.0484 g MBA (crosslinking agent for AM), 0.3 mL ECH (crosslinking agent for WP), 0.0272 g KPS (initiator for AM), and 0.1 mL TEMED (catalyst) were added successively. After mixing evenly, the resulting mixture was poured into a glass culture dish immediately and covered with the 9 cm Petri dish cover, then reacted at 60 °C for 2 h to form hydrogel. The thickness of the product was about 1 mm. Finally, the resulting hydrogel was divided into uniform pieces with a knife, washed alternately with deionized water and anhydrous ethanol until the pH was neutral, then kept at 60 °C for 24 h in a vacuum oven to gain the dried WP/PAM. To be clear, WP/PAM prepared from 1 wt% WP solution was defined as the 1 wt% WP/PAM, the same is true for preparation of the 2, 3, and 4 wt% WP/PAM. The prepared WP/PAM were sealed into self-sealing bags and stored in an electronic moisture-proof box (AD-50S ANDBON, China) for further use. The cost analysis was given in ESI (ESI, Text S1).†

In order to obtain the hydrogel based on WP with the best adsorption performance for Cu(II) and MB, the effect of WP content (1, 2, 3, and 4 wt%) on the adsorption behavior was investigated by comparison.

2.3 Characterization of WP/PAM

Put a certain amount of dry hydrogel into deionized water until the swelling equilibrium was reached. Wiped the surface water of the wet hydrogel with filter paper, then weighed it. The swelling ratio (SR) of the WP/PAM was calculated by the following eqn (1):²²

$$SR = \frac{m_1 - m_0}{m_0} \quad (1)$$

where m_0 and m_1 (g) were the weight of dried and swollen samples, respectively.

The porosity of the WP/PAM was measured by liquid replacement method.²³ First, the dried WP/PAM was weighed, then immersed in ethanol solution for 24 h, and the saturated WP/PAM was weighed again after the removal of the excess ethanol. The porosity (%) of the WP/PAM was calculated by the following eqn (2):

$$\text{Porosity} = \frac{m_0 - m_2}{\rho V} \times 100\% \quad (2)$$

where m_2 (g) were the weight of the WP/PAM after immersion in ethanol, ρ was the density of ethanol and V was the volume of the WP/PAM.

The morphology was analyzed by scanning electron microscope (SEM, JEOL JEM-2100F, Japan). The thermogravimetric analysis (TGA, TGA5500, USA) was carried out using a thermogravimetric analyzer in a nitrogen atmosphere, with temperatures ranging from room temperature to 600 °C and a heating rate of 10 °C min⁻¹. The isoelectric point of the WP/



PAM pH value (pH_{PZC}) was analyzed *via* ΔpH drift (see in Text S2†).¹⁸ The WP/PAM were analyzed before and after Cu(II) and MB adsorption by the Fourier transform infrared spectrum (FTIR, Bruker Vertex 70, Germany) in the wavenumber range of 400 to 4000 cm^{-1} and an X-ray photoelectron spectroscopy (XPS, ThermoFischer ESCALAB Xi+, USA). The change in performance of the WP/PAM after storage was measured by comparing the removal efficiency of Cu(II) and MB by the WP/PAM after storing 16 months.

2.4 Batch adsorption experiments

Dissolving 3.8020 g $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ (AR) in a 1000 mL volumetric flask to obtain the Cu(II) stock solution (1000 mg L^{-1}) and the required Cu(II) concentration through appropriate dilution of the stock solution. MB was directly dissolved in different concentrations of deionized water to prepare MB solution. In this work, the concentration of Cu(II) and MB were measured by an atomic absorbance spectrometer (ASAP 2020 Plus, Micromeritics) and a UV-2550 spectrophotometer (AOE, China) at 664 nm, respectively. The adsorption capacity at equilibrium (Q_e , mg g^{-1}) was calculated by the following eqn (3):

$$Q_e = V(C_0 - C_e)/m \quad (3)$$

where V (L) and m (g) represented the volume of the adsorbate solution and the dosage of the WP/PAM, respectively. C_e and C_0 (mg L^{-1}) were the equilibrium concentration and initial concentration of adsorbate, respectively.

In batch experiments, conical flask containing 50 mL Cu(II) or MB solution oscillated at 170 rpm under the required conditions. The adsorbent dose (0.25–1.5 g L^{-1}), pH (2–6 for Cu(II); 2–9 for MB), contact time (0–180 min), and initial concentration (0–2000 mg L^{-1} for Cu(II);²⁴ 400–2000 mg L^{-1} for MB) on the adsorption process were investigated. The pH was adjusted by 0.1 M HNO_3 and 0.1 M NaOH.

The competition effect between Cu(II) and MB was tested in two systems. In one of the systems, the effect of MB concentration (0–200 mg L^{-1}) on the adsorption capacity of the WP/PAM on Cu(II) was investigated when the concentration of Cu(II) was 40–100 mg L^{-1} . In the other system, the effect of Cu(II) concentration (0–100 mg L^{-1}) on the adsorption capacity of the WP/PAM on MB was investigated when the concentration of MB was 400–700 mg L^{-1} .

The amount of adsorbent used in this work represented the amount of dry hydrogel. The dry hydrogel was weighed quantitatively and put into deionized water for full swelling before adsorption. The thickness of the swollen hydrogel was about 2 mm. All the experiments were performed in triplicate.

2.5 Desorption and regeneration of the WP/PAM

To assess the regeneration of the WP/PAM in Cu(II) and MB adsorption–desorption process, the used WP/PAM were eluted and regenerated with 1.0 M HCl solution and 0.1 M NaOH solution, respectively. Firstly, the WP/PAM after adsorption of Cu(II) and MB was separated from the supernatant by filtration, then 50 mL 1.0 M HCl solution was added and the desorption

was completed by shaking at 170 rpm for 3 h. Afterwards, the WP/PAM was placed in 50 mL 0.1 M NaOH solution for 3 h for activation treatment. Finally, the material was washed with deionized water to neutral and left to use. The cyclic adsorption of Cu(II) and MB on the regenerated material was investigated under the original experimental conditions.

The desorption efficiency (η_d , eqn (4)) and the cyclic adsorption capacity (η_c , eqn (5)) were calculated by follows:

$$\eta_d (\%) = (Q_d/Q_e) \times 100\% \quad (4)$$

$$\eta_c (\%) = Q_{e(n)}/Q_{e(n-1)} \times 100\% \quad (5)$$

where Q_d (mg g^{-1}) was the maximum amount of Cu(II) or MB desorbed by the WP/PAM, n was number of cycles.

2.6 Simulated printing and dyeing wastewater treatment

A series of simulated wastewater samples were prepared respectively by the deionized water, the tap water from the laboratory, and the Xiangjiang River water (Fig. S2,† Changsha, Hunan province, China) with the addition of MB and some common metallic salt mordants (CuSO_4 (AR), CoCl_2 (AR), MnCl_2 (AR), and NiCl_2 (AR)). Captured by 3 g L^{-1} WP/PAM, the MB concentration was monitored on a UV-2550 spectrophotometer by full-spectrum detection, and the heavy metals concentrations were analyzed by an inductively coupled plasma-mass spectrometry (ICP-MS7700, Agilent Technologies, USA) after digestion.

3. Results and discussion

3.1 Effect of WP content

The adsorption capacities of the WP/PAM prepared with different WP contents on Cu(II) and MB were shown in Fig. S3.† When the WP content increased from 1 wt% to 2 wt%, the Q_e value for Cu(II) on the WP/PAM increased from 107.1 mg g^{-1} to 121.6 mg g^{-1} and the Q_e value for MB increased from 942.9 mg g^{-1} to 958.4 mg g^{-1} . The increase of WP content brought more hydroxyl groups resulted in the enhancement of the interaction between the WP/PAM and the adsorbate molecules.²⁵ However, as higher WP contents the decreasing on the adsorption was observed, by reason that larger number of WP fiber blocked part of the pore network and limited the molecular diffusion and transfer of pollutants,²⁶ in agreement with what was shown in Fig. S4† and the measured WP/PAM porosity (Table S1†). As shown in Fig. S4(a)–(d),† with the increase of WP content, it was obvious that the cellulose composition in the prepared WP/PAM increased in the prepared WP/PAM. Under the irradiation of SEM (Fig. S4(e)–(h)†), the changes of pores in the WP/PAM was more visually seen. When the WP content increased, the pore size became smaller and the laminated waste paper fiber structure began to appear at a WP content of 3 wt%. It can also be seen from Table S1† that with the increase of WP content, the porosity of the WP/PAM increased at first and then decreased. Among them, the porosity of 2 wt% WP/PAM was the highest (98.1%), which further explained why 2 wt% WP/PAM had the best adsorption performance. Moreover, the p -values (1.51 ×



10^{-5} for Cu(II) and 0.00508 for MB) < 0.05 also revealed that the WP content in the WP/PAM had a significant difference in the adsorption effect of Cu(II) and MB. Consequently, the 2 wt% WP/PAM was taken as the optimal ratio to carry out the following experiment.

3.2 Characterization of 2 wt% WP/PAM

The SEM of 2 wt% WP/PAM was depicted in Fig. 1(a). It was obvious that the freeze-dried WP/PAM presented a 3D porous network structure cross-linked by bundles of paper fibers and

PAM. The chemical crosslinking network effectively exposed a large number of active sites, promoting the mass transfer of Cu(II) and MB and enabled the removal of contaminants. The EDS result (Fig. 1(b)) further revealed that the synthetic material contained not only the characteristic elements of WP solution such as C, O, and Na, but also N elements peculiar to PAM, which confirmed the successful composite of WP and AM. The specific surface area and porous ratio of the WP/PAM were given in Table S2† and discussed in ESI (Text S3).†

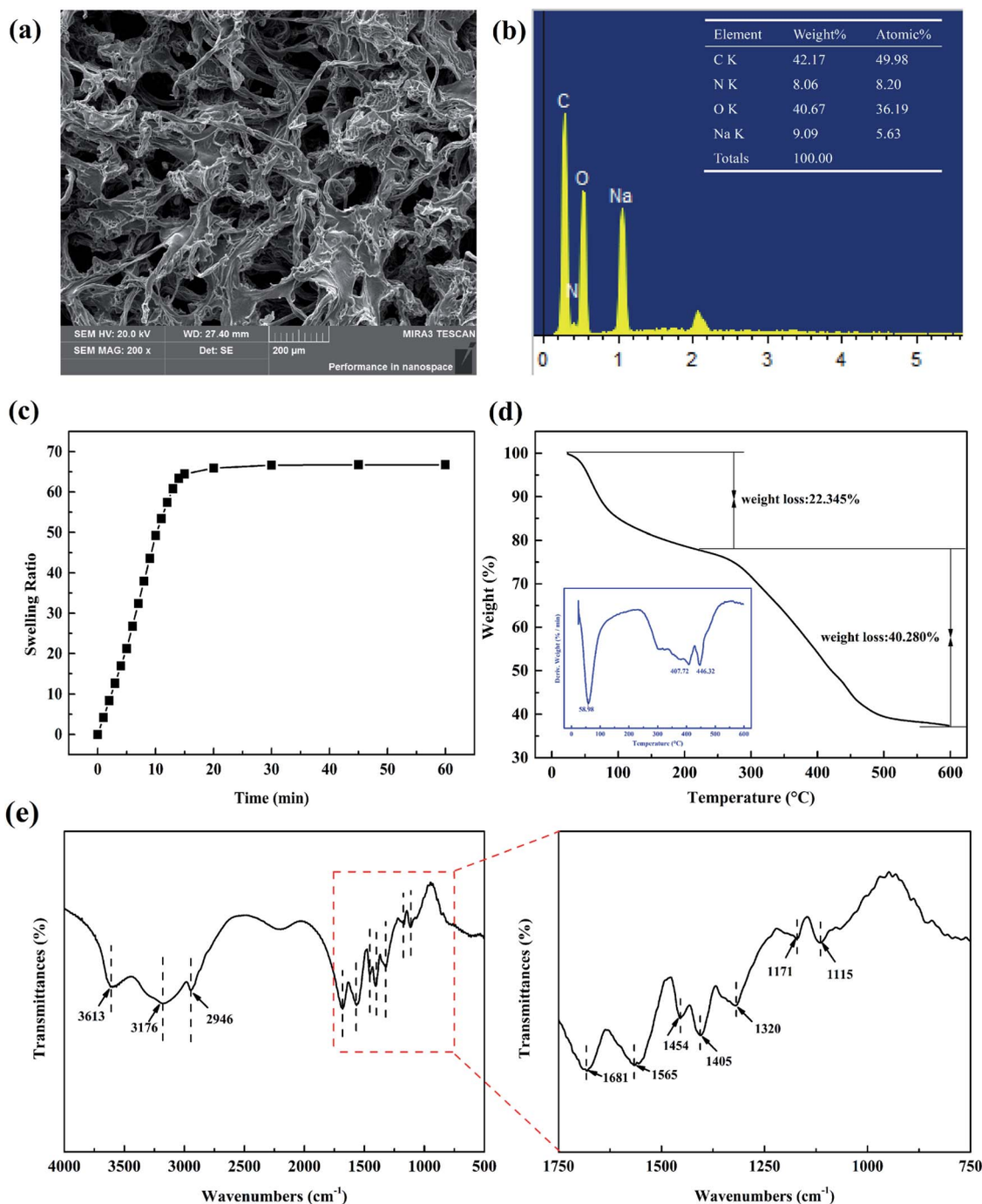


Fig. 1 The (a) SEM, (b) EDS, (c) SR, (d) TGA, (e) FTIR of the 2 wt% WP/PAM.



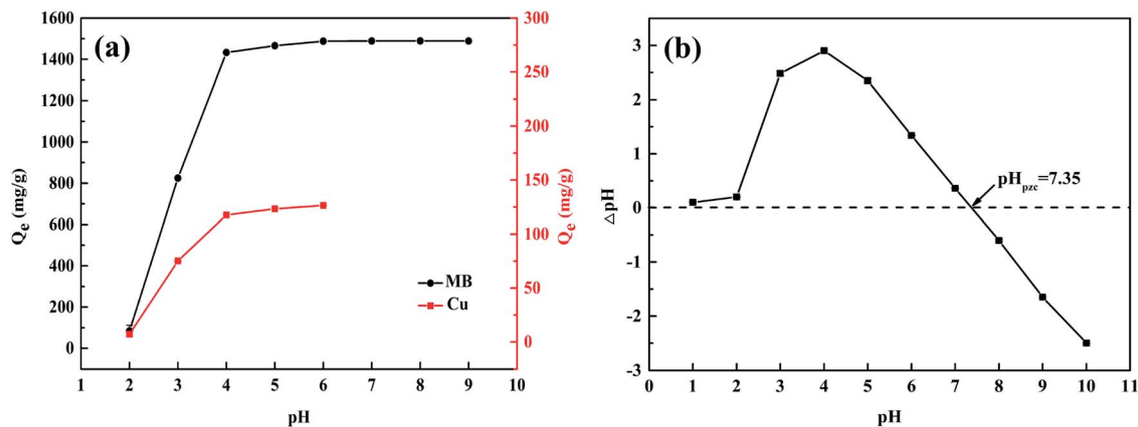


Fig. 2 (a) Effect of pH (b) zeta potential of the WP/PAM at different pH values. ($\text{dosage}_{\text{Cu(II)}}$: 0.7 g L^{-1} , $\text{dosage}_{\text{MB}}$: 0.4 g L^{-1} ; $C_{0(\text{Cu(II)})}$: 100 mg L^{-1} , $C_{0(\text{MB})}$: 600 mg L^{-1} ; 180 min; temperature: $25 \text{ }^\circ\text{C}$).

The swelling curve of the WP/PAM was observed in Fig. 1(c). The swelling equilibrium was reached within 30 min and SR was 66.75, which evinced excellent water absorption performance compared with other hydrogels.^{27,28} The excellent swelling properties of the WP/PAM would account for the loose double-network structure and rich hydrophilic groups such as hydroxyl groups.²⁹ Besides, amine groups on the PAM network were also reported as potential hydrogen bonds, which made hydrogel more likely to absorb abundant water in a short time, could facilitate the contaminants to enter into the hydrogel and be adsorbed.³⁰

Thermal characteristics of the prepared 2 wt% WP/PAM were presented in Fig. 1(d). Based on the TG-DTG curve, the degradation of the WP/PAM could be divided roughly into two main stages. At first, when the temperature was below $200 \text{ }^\circ\text{C}$, there was a 22.345% weight loss because of the loss of water in the WP/PAM.³¹ Then the WP/PAM lost 40.280% weight in the second stage, part of which corresponded to the pyrolysis of oxygen-containing groups (e.g., carboxyl and epoxy groups) and the release of CO_2 , CO, and steam when the temperature rose from $200 \text{ }^\circ\text{C}$ to $407.72 \text{ }^\circ\text{C}$.²⁷ when the temperature was greater

than $407.72 \text{ }^\circ\text{C}$, the thermal cracking of cellulose and PAM network structure of the WP/PAM would account for the further weight loss.³² In addition, it could be seen from Fig. 1(d) that the WP/PAM behaved steadily at the temperature required in this experiment ($15\text{--}35 \text{ }^\circ\text{C}$).

As depicted in Fig. 1(e), there were the characteristic peaks of paper fiber network^{14,33} and PAM network:^{24,34} the bands at 3613 and 3176 cm^{-1} was defined as the stretching vibration of $-\text{OH}$; the $\text{C}-\text{O}-\text{H}$ and $\text{C}-\text{O}-\text{C}$ stretching vibration appeared at 1454 and 1320 cm^{-1} , respectively; the bands at 1171 and 1115 cm^{-1} related to sugar structure was attributed to $\text{C}-\text{O}$ stretching vibration. The absorption peaks at 1681 , 1565 , and 1405 cm^{-1} were ascribed to the $\text{C}=\text{O}$ of amide stretching vibration peak, the $\text{N}-\text{H}$ bending vibration, and the $\text{C}-\text{N}$ stretching vibration, respectively. For the WP/PAM, all the above characteristic absorption peaks of paper fiber and PAM could be clearly observed, proving that the grafting copolymerization reaction was successful.

XPS analysis about the WP/PAM was discussed in Section 3.4.1 XPS spectra.

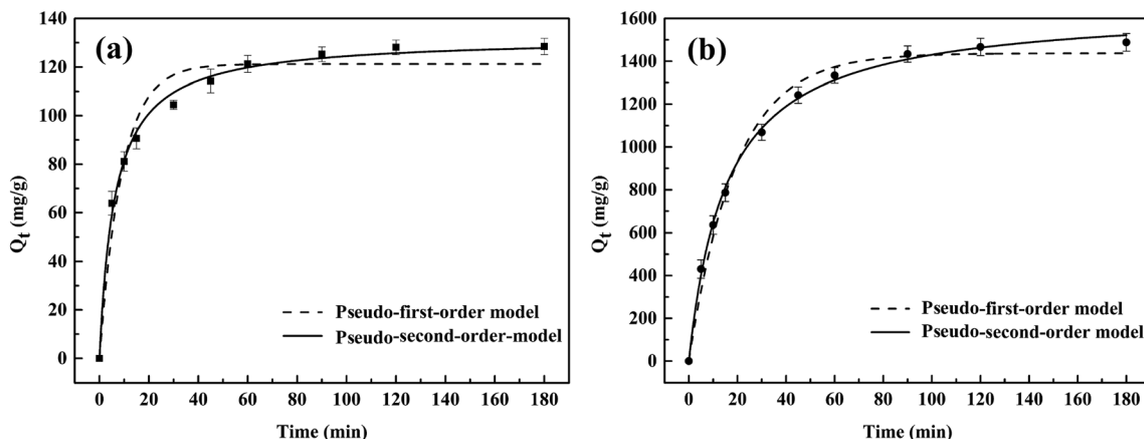


Fig. 3 The pseudo-first-order (a), and the pseudo-second-order (b) kinetic models of Cu(II) and MB adsorption on the WP/PAM ($\text{dosage}_{\text{Cu(II)}}$: 0.7 g L^{-1} , $\text{dosage}_{\text{MB}}$: 0.4 g L^{-1} ; $C_{0(\text{Cu(II)})}$: 100 mg L^{-1} , $C_{0(\text{MB})}$: 600 mg L^{-1} ; 180 min; temperature: $25 \text{ }^\circ\text{C}$).



In order to verify the performance of the WP/PAM stored in an electronic moisture-proof box, the adsorption capacity of the WP/PAM stored after 16 months on Cu(II) and MB was compared, and the results were shown in Table S3.† As could be seen from Table S3,† the adsorption property of the WP/PAM for Cu(II) and MB still retained more than 98.5% after 16 months of storage.

3.3 Adsorption of Cu(II) and MB by 2 wt% WP/PAM

3.3.1 Effect of adsorbent dosage. It was conducive to obtaining the balance between removal efficiency and material cost to study the effect of adsorbent dosage. As shown in Fig. S5,† the removal rate of the WP/PAM on 100 mg L⁻¹ Cu(II) gradually increased to 100% with the increase of the WP/PAM dosage from 0.25 to 0.75 g L⁻¹. Moreover, the 500 mg L⁻¹ MB had been completely removed even when the dosage was only 0.5 g L⁻¹. These proved the WP/PAM had an excellent and efficient ability to remove pollutants. However, as the WP/PAM dosage continued to increase, there will be more idle adsorption sites which reduced the adsorption capacity of per unit mass of the WP/PAM for Cu(II) and MB.³⁵ Based on revenue maximization and convenience, 0.7 g L⁻¹ and 0.4 g L⁻¹ were selected as the optimal dosage for Cu(II) and MB, respectively.

3.3.2 Effect of pH. As reported, the pH of most wastewater containing heavy metals is extremely acid, and the pH of the solution could affect the form of pollutants in the solution and the charge transferred on the interface of the pollutant/adsorbent, thus affecting the removal efficiency.^{36,37} To facilitate the better function of the WP/PAM, the wastewater needs to be adjusted to an appropriate pH before the adsorption reaction. Therefore, it is necessary to explore the optimal pH value for the adsorption reaction. Fig. 2(a) illustrated that the adsorption efficiency of the WP/PAM on Cu(II) and MB depended on pH.

When the solution pH was 2, the removal of Cu(II) and MB by the WP/PAM was not ideal. It might be due to the positive charge on the surface of WP/PAM (Fig. 2(b)), which enhanced the positive charge characteristics of the surface, resulting in the electrostatic repulsion to Cu(II) and cationic dyes (MB) with positive charge.³⁸ Besides, the solution contained abundant H⁺ could compete with pollutants for adsorption sites, resulting in weak adsorption of pollutants by the WP/PAM at this pH.³⁹ Simultaneously, low pH condition reduced the swelling ability of the WP/PAM, led to less removal of pollutants by the hydrogel.²⁴ When pH increased to 3–6 < p*H*_{pzc} = 7.35, the surface of the WP/PAM was still positively charged (Fig. 2(b)), but the adsorption efficiency increased rapidly until it reached saturation state. It indicated that electrostatic interaction was not the main adsorption mechanism. As the pH continued to rise, the removal of MB by the WP/PAM would not be affected, and reversely, the precipitation of Cu(II) would affect the adsorption.⁴⁰ Therefore, the WP/PAM had acceptable removal capacities on Cu(II) and MB under pH 3–6 and pH 3–9, respectively. For convenience, most experiments in this work were performed without adjusting the pH of the pollutant solution.

Table 1 The kinetic constants of Cu(II) and MB adsorption on the WP/PAM

Model	Parameters	Cu(II)	MB
Pseudo-first-order	K_1 (min ⁻¹)	0.1103	0.0521
	Q_e cal (mg g ⁻¹)	121.2	1437.1
	R^2	0.9600	0.9873
Pseudo-second-order	K_2 (g mg ⁻¹ min ⁻¹)	1.217×10^{-3}	3.875×10^{-5}
	Q_e cal (mg g ⁻¹)	132.2	1653.6
	R^2	0.9944	0.9980

3.3.3 Adsorption kinetic. Adsorption kinetics was often employed to describe the adsorption process and to assess the efficiency of the WP/PAM. As depicted in Fig. 3(a) and (b), the WP/PAM showed fast adsorption efficiency on Cu(II) and MB, which all reached 50% in just 15 min and then reached equilibrium in about 60 min. The high water-absorption and permeability of the WP/PAM were conducive to the diffusion and adsorption of pollutants, which would account for the rapid adsorption phenomenon.⁴⁸ In order to further explore the adsorption behavior of the WP/PAM for Cu(II) and MB, pseudo-first-order (eqn (6)), and pseudo-second-order (eqn (7))^{40,41} were respectively employed to fit the adsorption process of the WP/PAM on Cu(II) and MB. The fitting curves were exhibited in Fig. 3(a), (b), and (c), the correlation coefficient (R^2) and fitting parameters were given in Table 1.

$$Q_t = Q_e(1 - e^{-K_1 t}) \quad (6)$$

$$Q_t = \frac{K_2 Q_e^2 t}{1 + K_2 Q_e t} \quad (7)$$

where Q_t (mg g⁻¹) was the adsorption capacity of the WP/PAM on Cu(II) and MB at reaction time t . The K_1 (min⁻¹), and K_2 (g mg⁻¹ min⁻¹) were the adsorption equilibrium rate constants of the pseudo-first-order, and pseudo-second-order model, respectively.

Obviously, the higher R^2 value ($R^2 > 0.99$) in the pseudo-second-order model evinced that the adsorption behavior of the WP/PAM on Cu(II) and MB was related to chemical adsorption.⁴² The $K_2 < 1$ indicated that the reaction between the WP/PAM and Cu(II) and MB were rapid.⁴³ Besides, the Q_e values obtained by the pseudo-second-order model were 132.2 mg g⁻¹ for Cu(II) and 1653.6 mg g⁻¹ for MB, respectively, which were near to the experimental values (139.5 mg g⁻¹ for Cu(II) and 1705.5 mg g⁻¹ for MB).

3.3.4 Adsorption isotherm. Studying the adsorption isotherm could evaluate the surface properties and adsorption properties of the WP/PAM and understand the adsorption mechanism better. The curves fitted according to isothermal models were exhibited in Fig. 4,¹⁰ and relevant parameters were provided in Table 2.

The Langmuir model:

$$Q_e = Q_m K_L C_e / (1 + K_L C_e) \quad (8)$$

The Freundlich model:



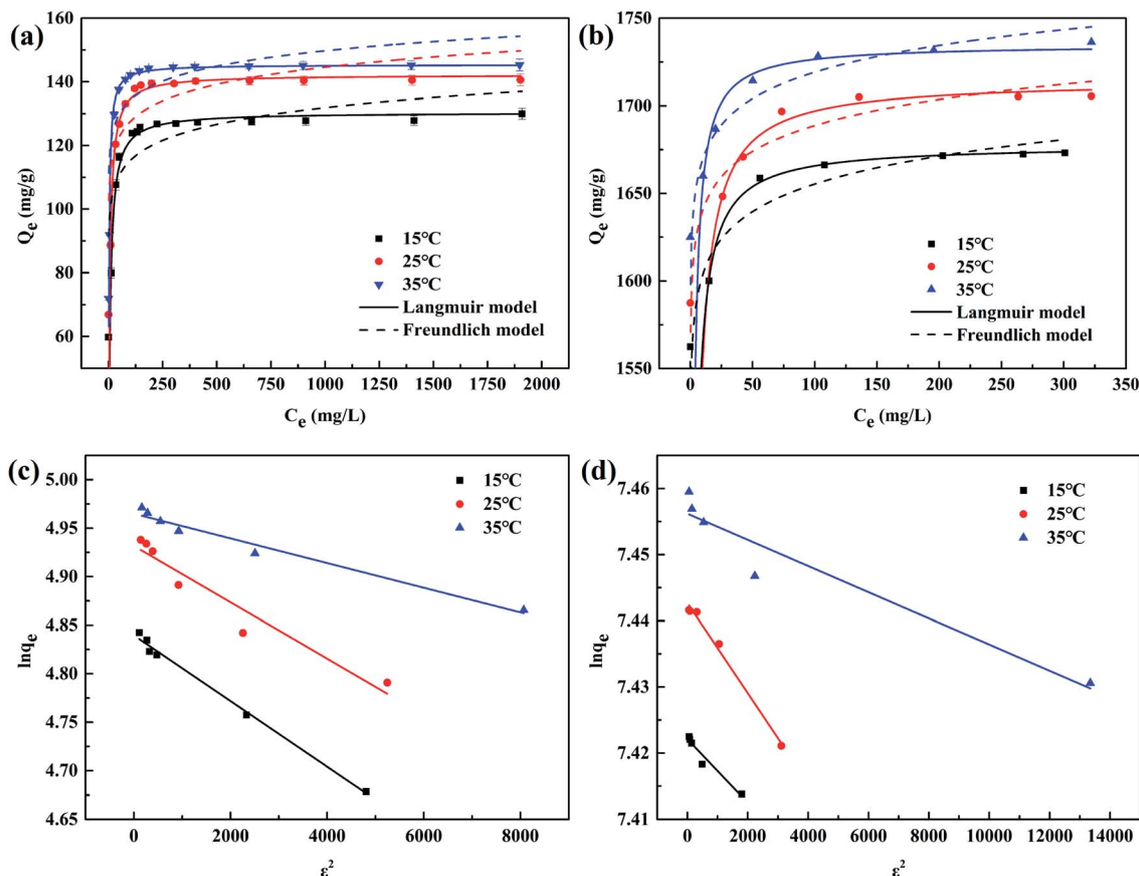


Fig. 4 The Langmuir model and the Freundlich model of (a) Cu(II) and (b) MB, and the D–R model of (c) Cu(II) and (d) MB adsorption on the WP/PAM (dosage_{Cu(II)}: 0.7 g L⁻¹, dosage_{MB}: 0.4 g L⁻¹, C₀(Cu(II)): 100 mg L⁻¹, C₀(MB): 600 mg L⁻¹, 180 min).

Table 2 The isotherm constants of Cu(II) and MB adsorption on the WP/PAM

Model	Parameters	Cu(II)			MB		
		15 °C	25 °C	35 °C	15 °C	25 °C	35 °C
Langmuir parameters	Q _m (mg g ⁻¹)	130.4	142.2	145.3	1677.7	1714.5	1734.9
	K _L (L mg ⁻¹)	0.1356	0.1767	0.4046	1.3847	0.9662	2.0530
	R _L ²	0.9849	0.9935	0.9949	0.9979	0.9662	0.9708
Freundlich parameters	K _F (mg g ⁻¹)	88.1	99.2	108.1	1533.3	1593.4	1621.3
	n	17.11	18.32	21.16	72.31	79.24	78.56
	R _F ²	0.5807	0.5816	0.5761	0.8141	0.7617	0.9014
Dubinin–Radushkevich parameters	Q _m (mg g ⁻¹)	126.4	138.6	143.3	1672.6	1707.4	1730.5
	E (kJ mol ⁻¹)	20.88	24.35	25.98	10.20	8.573	15.90
	R ²	0.9923	0.9197	0.9649	0.9300	0.9891	0.9064

$$Q_e = K_F C_e^{1/n} \quad (9)$$

The D–R model:

$$\ln Q_e = \ln Q_m - K_{DR} \varepsilon^2 \quad (10)$$

$$\varepsilon = RT \ln(1 + 1/C_e) \quad (11)$$

$$E = 1 / \sqrt{2K_{DR}} \quad (12)$$

where Q_m (mg g⁻¹) and C_e (mg L⁻¹) were the maximum capacity and the equilibrium concentration of pollutants, respectively. K_L (L mg⁻¹), K_F (mg g⁻¹), n, and K_{DR} (mol² kJ⁻²) were the Langmuir constant, the Freundlich constants and the Dubinin–Radushkevich constant, respectively. R = 8.314 (J mol⁻¹ K⁻¹), T (K) represented the temperature and E (kJ mol⁻¹) was the free energy of the reaction.

As displayed in the Fig. 4(a) and (b), when C₀(Cu(II)) and C₀(MB) raised, the Q_e increased and moved towards a plateau eventually, illustrating that increasing the C₀ of the pollutant was



Table 3 Comparison of removal of Cu(II) and MB by cellulose based hydrogels

Cellulose based hydrogel	Solution conditions	Q_m (mg g ⁻¹)		Ref.
		Cu(II)	MB	
Cotton	25 °C, 6 h	138.9		18
Microcrystalline cellulose powder	Room temperature, 8 h	52.3		30
Eucalyptus wood	30 °C, 4 h	16.7		46
Carboxymethyl cellulose	25 °C, 24 h		357–526	35
Bagasse	25 °C		37.2	47
Sugarcane bagasse	30 °C, pH = 9		632.9	48
Pineapple peel	30 °C		172.4	27
WP	25 °C, 3 h	142.2	1714.5	This study

conductive to improving the driving force of the solid–liquid interface (*i.e.*, ion exchange, electrostatic interaction, and chemical association) until the adsorption reached saturation.⁴¹ Comparing the R^2 under three temperatures, it was noticeable that the systems of the WP/PAM adsorption for Cu(II) and MB followed the Langmuir model. It meant that only monolayer adsorption occurred on the WP/PAM, once a molecule was adsorbed, the adsorption site became unusable.⁴⁴ What's more, the E value of Cu(II) adsorption process calculated in D–R model was 20.88–25.98 kJ mol⁻¹ beyond the E range of chemical interaction (8–16 kJ mol⁻¹) (Fig. 4(c) and (d)), suggesting that the WP/PAM had physical interaction with Cu(II) in addition to chemical interaction.⁴⁵

The Q_m values obtained by the Langmuir model at 25 °C were 142.2 mg g⁻¹ for Cu(II) and 1714.5 mg g⁻¹ for MB, respectively, which were near to the experimental values (139.5 mg g⁻¹ for Cu(II) and 1705.5 mg g⁻¹ for MB). Furthermore, higher temperature was conducive to the improvement of Q_m . It might illustrate that the adsorption of the WP/PAM with Cu(II) and MB were endothermic reaction and the affinity of the WP/PAM for Cu(II) and MB were reinforced at higher temperatures. To revealing excellent adsorption capacity of the WP/PAM more

intuitively, the comparison of the adsorption capacity of the WP/PAM and other cellulose-based hydrogels on Cu(II) and MB was listed in Table 3.

3.3.5 Competitive effect between Cu(II) and MB. Cu(II) usually existed together with MB as a mordant in printing and dyeing wastewater,⁵ so it was necessary to discuss the competition effect between Cu(II) and MB in a single and binary system. As depicted in Fig. 5(a) and (b), it was noticeable that when the concentration of one adsorbate increased in binary solution, the Q_e of another adsorbate gradually decreased compared with that in the single system which show in Fig. 5(a) and (b) for Cu(II) and MB, respectively. The results verified that Cu(II) and MB would compete for the same active site of the WP/PAM in the binary system, resulting in insufficient effective adsorption sites for Cu(II) and MB.⁴⁹ From the details in the Fig. 5(a), the Q_e of Cu(II) decreased slightly when MB concentration increased. As the $C_{0(\text{Cu(II)})}$ was 100 mg L⁻¹, the Q_e value of Cu(II) decreased from 142.9 to 110.5 and then decreased to 102.1 mg g⁻¹ with $C_{0(\text{MB})}$ increased from 0 mg L⁻¹ to 100 mg L⁻¹ and then increased to 200 mg L⁻¹, respectively. Whereas, while the $C_{0(\text{Cu(II)})}$ raised from 0 to 50 mg L⁻¹ and then raised from 50 to 100 mg L⁻¹ in the solution containing 700 mg L⁻¹ MB, the

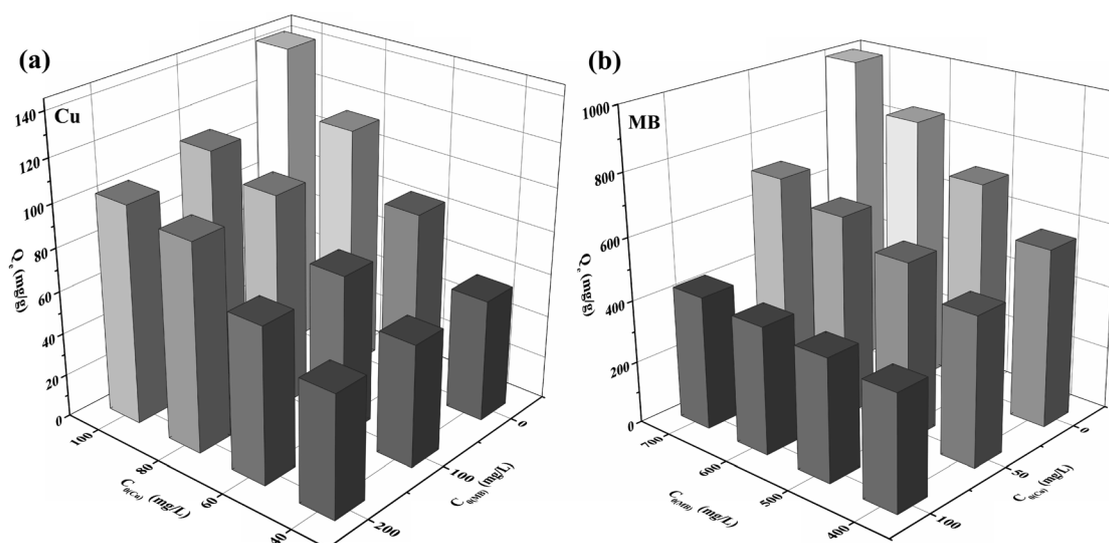


Fig. 5 The adsorption capacity of (a) Cu(II) and (b) MB in single and binary systems (dosage: 0.7 g L⁻¹; 180 min; temperature: 25 °C).



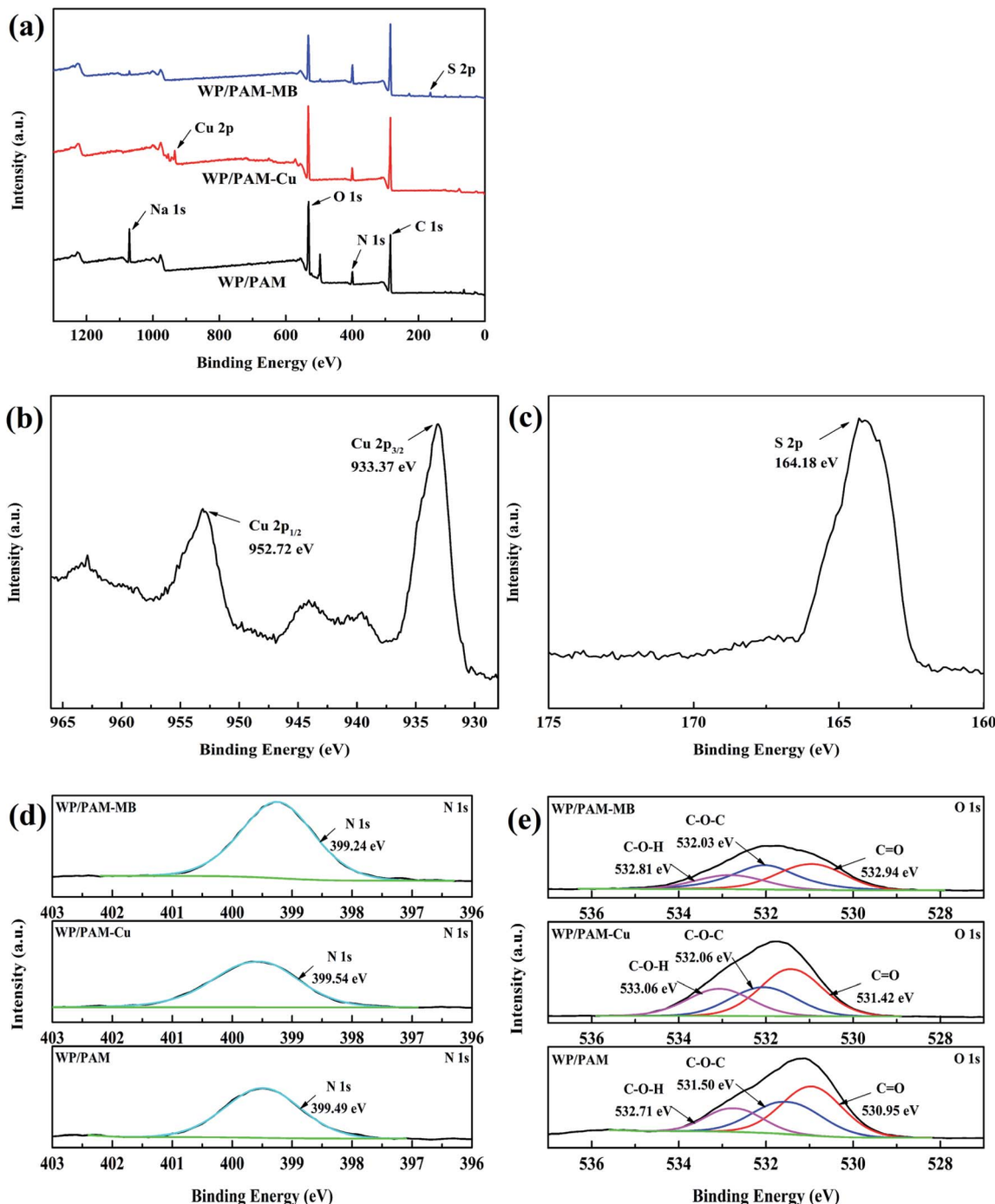


Fig. 6 XPS spectra of (a) the WP/PAM before and after Cu(II) and MB adsorption, (b) Cu 2p, (c) S 2p, (d) N 1s and (e) O 1s.

loss of Q_e (for MB) on the WP/PAM were 29% and 52%, respectively (Fig. 5(b)). Compared with the fact that there was a slight loss of Cu(II) removal when the concentration of MB increased, the 29% and 52% loss of MB removal indicated that Cu(II) was more inclined to adsorb on the WP/PAM than MB in the binary system.

3.4 Adsorption mechanism

3.4.1 XPS spectra. XPS characterization of the WP/PAM was performed before and after Cu(II) and MB adsorption to further

clarify the interaction mechanism between adsorbate and the WP/PAM and the results were placed in Fig. 6. After adsorption, Cu 2p (Fig. 6(b), Cu 2p_{1/2}: 952.72 eV and Cu 2p_{3/2}: 933.37 eV) and S 2p (Fig. 6(c), 164.18 eV) characteristic peaks appeared on the wide scan XPS of the WP/PAM (Fig. 6(a)),^{37,50} indicating Cu(II) and MB were successfully captured by the WP/PAM. It could be further seen from Table 4 that the Na element could not be detected on the WP/PAM after Cu(II) adsorption, certified that the WP/PAM had ion exchange with Cu(II).⁵¹ Besides, the increased relative content of the N element in the WP/PAM after

Table 4 The element content ratio (%) of the WP/PAM before and after Cu(II) and MB adsorption

	C	O	N	Na	Cu	S
WP/PAM	60.39	27.98	6.95	4.68	—	—
WP/PAM-Cu	66.65	25.05	6.57	—	1.73	—
WP/PAM-MB	69.24	18.51	9.44	0.80	—	2.01

the adsorption of MB was observed because of the N in the MB constituent element, which equally confirmed the successful adsorption of MB on the WP/PAM.^{52,53}

Refer to N 1s (Fig. 6(d)), there was no significant change in the binding energy between the WP/PAM (399.49 eV) and the WP/PAM-Cu (399.54 eV), showed that the nitrogen containing functional group did not participate in the Cu(II) adsorption process.⁵⁴ This was due to the fact that the lone pair electrons of nitrogen atoms in the amide group were occupied and could not be contributed to adsorbate.¹⁹ While the binding energy of the WP/PAM-MB decreased slightly from 399.49 eV to 399.24 eV, indicating that the WP/PAM had electrostatic interaction with N element in MB, increasing the density of the nearby electron cloud.^{42,50}

The O 1s of the WP/PAM (Fig. 6(e)) could fit three peaks with binding energies of 532.71, 531.50, and 530.95 eV, contributed to C-O-H, C-O-C, and C=O, respectively.^{55,56} After reacted with Cu(II), the binding energy of C-O-H and C=O had an increase from 532.71 eV to 533.06 and 530.95 to 531.42 eV, respectively. It was an indication that the density of the electron cloud around the O atom in C-O-H and C=O reduced. The phenomenon enabled Cu(II) and oxygen-containing functional groups to form Cu-O through complexation reaction.⁴³ As for the WP/PAM-MB, the C-O-H peak originally located at 532.71 eV had moved to a higher binding energy (532.81 eV), demonstrating that C-O-H was involved in the MB adsorption, which was in accordance with the results reported previously.⁵⁷ Combined with the analysis of N 1s, it was reasonable to conclude that it was the C-O-H in the WP/PAM had electrostatic interaction with N in MB.⁵⁸ As a result, XPS results revealed that the adsorption

mechanism of the WP/PAM for Cu(II) mainly involved complexation and ion exchange, and the adsorption process for MB was mainly controlled by electrostatic interaction, which were consistent with the results^{59,60}

3.4.2 FTIR spectra. Compared with the WP/PAM (Fig. 7(a)), the C-O-H, C-O-C, and C-O stretching vibration were weakened after Cu(II) adsorption, indicating that the cellulose network was involved in the adsorption process.⁵⁸ Moreover, a new absorption band was observed at 1603 cm⁻¹, suggested that the oxygen-containing functional group on the WP/PAM formed a bridge carboxylate with Cu(II).⁶¹ The C=O at 1681 cm⁻¹ was shifted to 1669 cm⁻¹, which could be attributed to the formation of complexes between carboxyl groups in surface and Cu(II) in the solution.⁴⁹ These revealed that the Cu(II) adsorption process on the WP/PAM involved complexation,⁶² which was in accordance with the XPS analysis.

In the case of MB, new peaks appeared at 1338, 1219, 1173, 1133, and 881 cm⁻¹, pertain to the C-N, AR-N, C=S, C-S connected to the benzene ring, and C-H near the benzene ring, respectively.^{63,64} After reacting with MB, the C-N on the WP/PAM was shifted from 1405 to 1396 cm⁻¹, which was consistent with previous studies.⁶⁵ It was reported that MB is a cationic dye, which could utilize the N atom of the C-N group and S atom of the C-S group as hydrogen bond acceptor to form an intermolecular hydrogen bond with H atom of the hydroxyl in the WP/PAM.^{64,66} The C=O at 1681 cm⁻¹ was shifted to 1669 cm⁻¹ after adsorption, which was caused by the reaction between negatively charged groups in the composite matrix with the unsaturated dimethylamine groups of MB and the formation of hydrogen bonds according to Gomes *et al.*²⁹ and Abdel *et al.*⁶⁷

Fig. 8 provided a brief illustration of the adsorption mechanism.

3.5 Desorption and regeneration of the WP/PAM

The reusability of the WP/PAM was also an important index to evaluate the material practicability. It could be seen in Fig. S6 and Table S4† that the WP/PAM hydrogel presented good stable adsorption performance and reusability. After 6 adsorption-desorption cycles, the loss of removal rate of the WP/PAM for

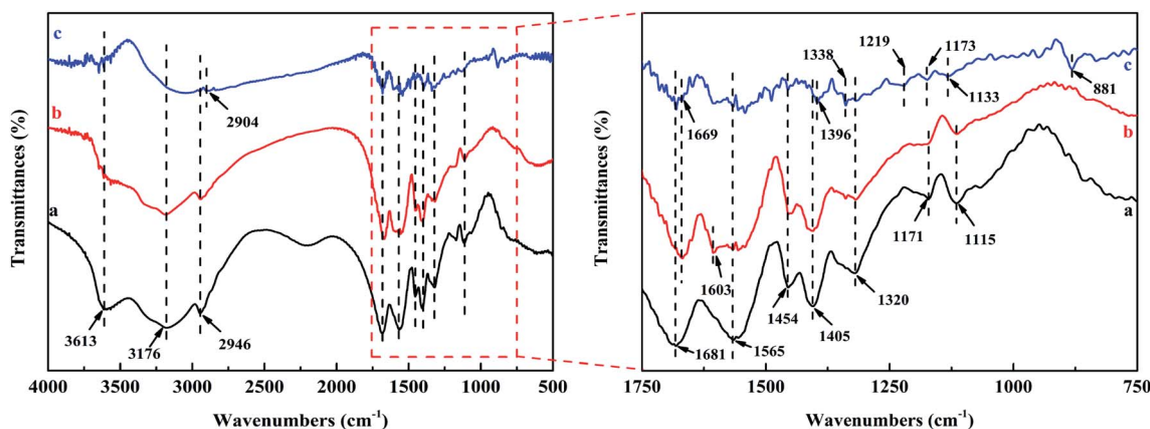


Fig. 7 FTIR spectra of the WP/PAM (a) before, (b) after Cu(II) and (c) after MB adsorption.



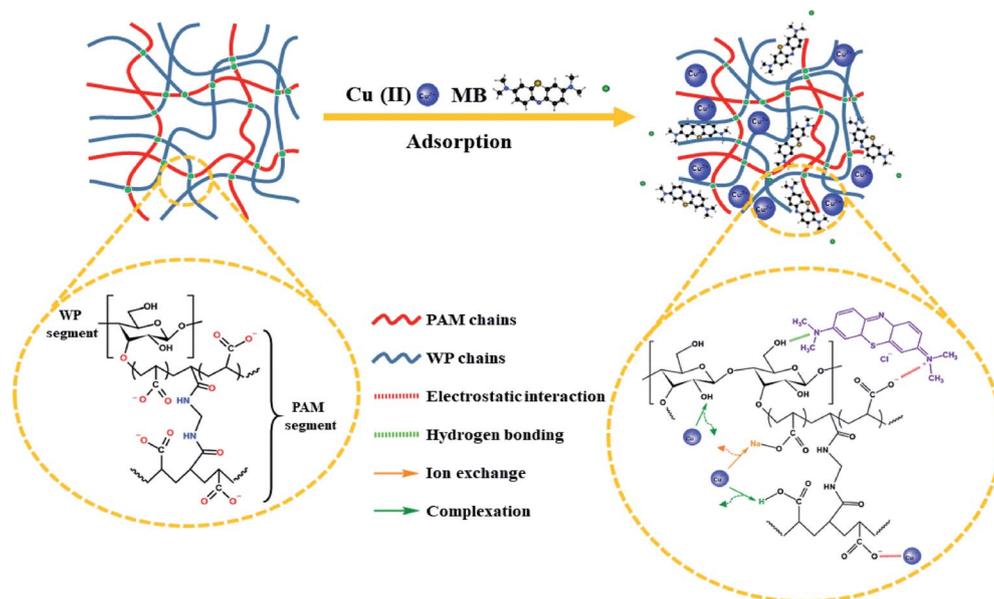


Fig. 8 Adsorption mechanism of the WP/PAM on Cu(II) and MB.

Table 5 Parameters before and after manipulation of printing and dyeing wastewater prepared by deionized water, tap water, Xiangjiang river

Parameters	Sample prepared by deionized water		Sample prepared by tap water		Sample prepared by Xiangjiang river	
	Before	After	Before	After	Before	After
pH	4.64	7.46	5.51	7.63	5.40	7.18
Pollutants (mg L^{-1})	MB	505.35	3.85	495.45	5.40	491.28
	Cu(II)	125.60	0.45	114.78	0.45	115.55
	Co(II)	54.67	0.16	51.92	0.17	57.36
	Mn(II)	35.84	0.33	35.09	0.27	34.84
	Ni(II)	20.07	0.73	22.89	0.74	22.35

Cu(II) could be negligible, while the adsorption and removal rate of MB could still reach more than 82% with high concentration (600 mg L^{-1}). Additionally, the WP/PAM could be separated directly from wastewater by natural settling. The recovery method with easy to operation and low loss presented a great potential of the WP/PAM in wastewater treatment.

3.6 Treatment of simulated printing and dyeing wastewater

The parameters of simulated wastewater before and after treatment were summarized in Table 5. The pH of the simulated printing and dyeing wastewater were acidic, because the metal salts (*i.e.*, CuSO_4 , CoCl_2 , MnCl_2 and NiCl_2) added to the simulated wastewater belong to strong acid and weak base salts. When they dissolved in water, hydrolysis occurred to generate weak electrolyte hydroxides and H^+ , resulting in acidic solution. The reason why the pH of the treated simulated wastewater reached neutral was that due to the adsorption of WP/PAM, and the corresponding hydrolytic equilibrium shifted, resulting in the decrease of H^+ in the solution. Therefore, the pH of the solution increased after the WP/PAM reacted with metal ions.

Furthermore, captured by 3 g L^{-1} WP/PAM, the ion concentrations of wastewater were basically lower than the limit of industrial wastewater discharged in China (Cu(II): 0.5 mg L^{-1} , Co(II): 1.0 mg L^{-1} , Ni(II): 1.0 mg L^{-1} and Mn(II): 2.0 mg L^{-1}). After the treatment, incineration and landfill were considered as the disposal methods of the adsorbed WP/PAM. In general, the WP/PAM showed good potential for treating MB industrial wastewater containing heavy metal ions.

4. Conclusions and outlook

In this study, the double-network hydrogel based on WP cellulose was successfully synthesized, which could effectively remove Cu(II) and MB. The WP/PAM presented a 3D porous network structure cross-linked by bundles of paper fibers and PAM, which effectively exposed a large number of active adsorption sites. In addition, the WP/PAM had excellent swelling properties ($\text{SR} = 66.75$), which promoted the mass transfer of Cu(II) and MB on the WP/PAM. The excellent structural properties of the WP/PAM played a favorable role in the



adsorption of Cu(II) and MB. Batch experiments showed that the adsorption process of Cu(II) and MB was more consistent with the pseudo-second-order kinetic model and the Langmuir model. At 25 °C, the maximum adsorption capacities of the WP/PAM for Cu(II) and MB obtained by Langmuir model fitting were 142.2 mg g⁻¹ and 1714.5 mg g⁻¹, respectively. The competitive adsorption experiments between Cu(II) and MB showed that Cu(II) was more inclined to adsorb on the WP/PAM than MB. The adsorption mechanism analysis results indicated that the adsorption mechanism mainly included ion exchange, complexation for Cu(II), while electrostatic interaction and hydrogen bonding for MB. Besides, the WP/PAM can effectively handle simulated wastewater containing Cu(II) and MB. This study provides a new way for recycling of WP and a valuable reference for its application in wastewater treatment.

However, there were still shortcomings and areas for improvement in our work. In order to apply the WP/PAM better into practical wastewater treatment, it is necessary to study the effect of different sources of WP on the adsorption performance of WP/PAM, more simplified synthesis steps and more economical amounts of regeneration solution to ensure lower costs. The adsorption behavior and the adsorption mechanism of the WP/PAM on other heavy metal mordants also need to be explored in more depth. In the future, an adsorption tower will be prepared for research on the practical application. The equipment will include a storage tower (filled with water to ensure the expansion effect), adsorption tower, regeneration device, etc. Some important details, such as how these devices can be properly assembled and placed in the adsorption tower to make the treatment process simple and efficient, the replacement period of the WP/PAM, and the more economical way of waste disposal, will be the direction and focus of our future work.

Author contributions

Yaoning Chen: supervision, validation, funding acquisition, writing – review & editing. Linshenzhang Li: data curation, investigation, methodology, visualization, writing – original draft. Yuanping Li: supervision, writing – review. Yihuan Liu: writing – review & editing. Yanrong Chen: writing – review & editing. Hui Li: supervision, funding acquisition & writing – review. Meiling Li: methodology, writing – review & editing. Fangting Xu: software, writing – review & editing. Yuqing Liu: writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This study was financially supported by the National Natural Science Foundation of China (51979104), Training Program for Excellent Young Innovators of Changsha (kq1802010, kq1802040), Science and Technology Planning Project of Hunan

Province (2018RS3109), and Natural Science Foundation of Hunan Province, China (2020JJ5019).

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