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### Transformation of polyvinyl chloride (PVC) into a versatile and efficient adsorbent of Cu(II) cations and Cr(VI) anions through hydrothermal treatment and sulfonation

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22 The reuse of waste polyvinyl chloride (PVC) has drawn much attention as it can reduce 23 plastic waste and associated pollution, and provide valuable raw materials and products. In 24 this study, sulfonated PVC-derived hydrochar (HS-PVC) was synthesized by two-stage hydrothermal treatment (HT) and sulfonation, and shown to be a versatile adsorbent. The 25 26 reoval of Cu(II) cations and Cr(VI) anions using HS-PVC reached  $81.2 \pm 1.6\%$  and  $60.3 \pm$ 27 3.8%, respectively. The first stage of HT was crucial for the dichlorination of PVC and the 28 formation of an aromatic structure. This stage guaranteed the introduction of -SO<sub>3</sub>H onto 29 PVC-derived hydrochar through subsequent sulfonation. HT intensities (i.e., temperature and time) and sulfonation intensity strongly determined the adsorption capacity of HS-PVC. 30 31 Competitive adsorption between Cu(II) and Cr(VI) onto HS-PVC was demonstrated by 32 binary and preloading adsorption. The proposed Cu(II) cations adsorption mechanism was 33 electrostatic adsorption, while Cr(VI) were possibly complexed by the phenolic -OH and 34 reduced to Cr(III) cations by C=C groups in HS-PVC. In addition, HS-PVC derived from PVC waste pipes performed better than PVC powder for Cu(II) and Cr(VI) removal (>90%). 35 36 This study provides an efficient method for recycling waste PVC and production of efficient 37 adsorbents.

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Keywords: Polyvinyl chloride; Hydrothermal treatment; Sulfonation; Adsorption;
Cu(II) cations and Cr(VI) anions

#### 42 **1. Introduction**

Polyvinyl chloride (PVC) is irreplaceable in the infrastructure of modern society (Cui et al., 2021). However, large quantities of plastic waste continuously enter aquatic and terrestrial ecosystems, leading to ubiquitous pollution worldwide (Kawecki and Nowack 2019, Miao et al., 2020). For example, the accumulated PVC waste in China will exceed 500 million tons by the end of 2050, posing a daunting challenge for the safe and effective disposal of this halogenated waste (Liu et al., 2020).

49 Mechanical recycling (25.5%), incineration (9.3%), and landfill disposal (36.0%) are 50 currently the main solutions for solving the problem of PVC waste (Yu et al., 2016, Ye et al., 2017, Liu et al., 2020). However, mechanical recycling is still rudimentary and only suitable 51 52 for source-separated PVC waste. Incineration and landfill disposal of PVC waste normally 53 release hazardous compounds, such polychlorinated dibenzo-p-dioxins as and 54 polychlorinated dibenzofurans. Furthermore, the land requirements of landfills must be considered (Wiedinmyer et al., 2014). Thus, reliable alternative methods to treat PVC waste 55 56 safely and efficiently in an environmentally sound and high value-added way are worth 57 pursuing (Lau et al., 2020). Such an approach is in line with the European Union's strategy 58 for plastics in a circular economy (Matthews et al., 2021).

Hydrothermal treatment (HT), conducted at relatively low temperatures of 160 to 300°C,
can pretreat PVC waste for dehydrochlorination (Ma et al., 2019, Ning et al., 2020),
permitting subsequent pyrolysis to produce chars and liquid hydrocarbons with a low Cl
content, such as alkene and polyene (Poerschmann et al., 2015, Jiang et al., 2021). However,

cost- and energy-intensive conditions, such as noble metal catalysts and high temperature 63 (500°C), are typically adopted for lighter oil production by catalytic pyrolysis of plastic 64 65 (Kunwar et al., 2016, Maity et al., 2020). On the other hand, PVC waste can be converted into coal-alternative fuels by HT processes amended with cellulose (Shen et al., 2017), 66 pinewood sawdust (Huang et al., 2019), and corncob (Lu et al., 2020). Yet, another important 67 recycling method has been undervalued and ignored, i.e., the potential transformation of 68 69 PVC-derived hydrochar (H-PVC) into an effective adsorbent for the removal of contaminants, 70 such as heavy metals.

71 Unfortunately, the adsorption capacity of organic compounds by H-PVC is very low, primarily because surface-related adsorption prevails over hydrophobic partitioning 72 73 (Poerschmann et al., 2015). Several studies showed that the adsorption of hydrochar derived from natural biomass sources was enhanced by introducing functional groups, such as -OH, -COOH, and -NH<sub>2</sub>, which could trap heavy metal ions from aqueous solutions (Azzaz et al., 2020, Yang et al., 2021). Sulfonation is an attractive modifications method for various carbon materials due to the unique surface chemistry and low production costs. Large amounts of oxygen functionalities, such as -OH, -COOH, and lactones, along with -SO<sub>3</sub>H groups, are introduced into the surface of the carbon materials, improving their hydrophilic/oxyphilic properties after sulfonation. Sulfonated carbon materials have been widely applied in the catalysis of esterifications, acetalization, etherifications, and transesterifications (Konwar et al., 2019). However, sulfonated PVC-derived hydrochar (HS-PVC) has rarely been reported, albeit potentially being a cost-effective adsorbent for heavy metals.

84	The novelty of the present work is the development of two-stage HT and sulfonation to
85	synthesize PVC-derived adsorbents for the removal of both Cu(II) cations and Cr(VI) anions.
86	A systematic investigation was conducted to discern the mechanisms underlying functional
87	group modifications of PVC through different strategies and treatment intensities. The
88	adsorption mechanisms of Cu(II) and Cr(VI) were clarified to understand the adsorption
89	behavior onto HS-PVC. The performance of HS-PVC to adsorb Cu(II) and Cr(VI) from
90	natural water was evaluated. PVC waste pipes were used to synthesize HS-PVC and test its
91	applications. Furthermore, the versatility of HS-PVC for the adsorption of low concentrations
92	of Cu(II) and Cr(VI) (20 mg/L) and other metals, i.e., Mn(II), Cd(II), Ni(II), and Co(II) was
93	investigated. This study provides a practical approach for the sustainable reuse of waste PVC
94	as an efficient adsorbent to remove heavy metal cations and anions.

#### 96 2. Materials and methods

#### 97 2.1. Preparation of PVC-derived adsorbents

Raw PVC powder (sieved through a 100 mesh) was purchased from a plastic chemical company (Guangdong, China). Hydrothermal batch reactions were conducted by loading 10 g of Raw PVC with 40 mL distilled water in 100-mL airtight stainless-steel tubular reactors with internal polyphenylene containers. The reactors were heated to 250 °C for 6 h in a homothermic oven (DHG-9013A, China). The reactors were then cooled to room temperature in a water bath to prepare the H-PVC. Ten grams of Raw PVC was sulfonated in flasks (250 ml) with concentrated sulfuric acid (98%) at a sulfonation intensity of 1:10 (g/mL) in a shaking 105 bath (70 °C and 110 rpm for 12 h) to produce the sulfonated PVC (S-PVC). Unless otherwise 106 specified, sulfonation intensity refers to the ratio of adsorbent mass to the volume of sulfuric 107 acid in this paper. Specifically, a relatively low temperature of sulfonation (70 °C) was chosen 108 according to the literature (Xiao and Hill 2020). The HS-PVC was synthesized by sulfonation 109 of H-PVC with the same process as described for S-PVC. The S-PVC and HS-PVC were 110 washed three times with distilled water to eliminate the unreacted sulfuric acid. The Raw PVC, 111 H-PVC, S-PVC, and HS-PVC were dried in a homothermic oven at 105 °C for 24 h. The dried 112 PVC-derived adsorbents were then ground and sieved through a 100 mesh (147 um) before 113 adsorption of Cu(II) and Cr(VI).

#### 114 2.2. PVC-derived adsorbents for highly efficient removal of Cu(II) and Cr(VI)

#### 115 2.2.1. Cu(II) and Cr(VI) removal using Raw PVC, H-PVC, S-PVC, and HS-PVC

116 A stock solution of 500 mg Cu(II)/L (7.87 mM) was prepared by dissolving 1965 mg of 117 copper sulfate pentahydrate (CuSO<sub>4</sub>·5H<sub>2</sub>O) in 1 L of ultrapure water. A stock solution of 250 mg Cr(VI)/L (4.81 mM) was prepared by dissolving 770 mg of potassium dichromate 118 119 (K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub>) in 1 L of ultrapure water. Batch experiments of Cu(II) and Cr(VI) removal were 120 performed by adding 10 g/L of Raw PVC, H-PVC, S-PVC, and HS-PVC to flasks (250 mL) in a water bath shaker (25 °C and 100 rpm) for 12 h. All samples were withdrawn from the 121 122 flasks and filtered through membrane filters with a pore size of 0.22 µm for subsequent Cu(II) and Cr(VI) determination. 123

Removal efficiency = 
$$\frac{C_0 - C_t}{C_0} \times 100\%$$
 (1)

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124

where  $C_0$  (mg/L) and  $C_t$  (mg/L) were the concentrations of Cu(II) and Cr(VI) at the start of the experiment and time t (h). Unless otherwise specified, all calculations of Cu(II) and Cr(VI) removal efficiencies by PVC-derived adsorbents followed this equation. The concentrations of Cu(II) and Cr(VI) were determined by inductively coupled plasma mass spectrometry (ICP-MS) (Thermo Scientific iCAP Q, USA).

# 131 2.2.2. Cu(II) and Cr(VI) removal using HS-PVC under different intensities of HT 132 temperature, HT time, and sulfonation and correlation analyses

HS-PVC was synthesized under different intensities of HT (temperature and time) and 133 134 sulfonation as follows: HT temperatures from 220 °C to 260 °C (in 10 °C intervals, HT time of 6 h and sulfonation intensity of 1:10), HTC times from 2 h to 6 h (in 1 h intervals, HT 135 136 temperature of 250 °C, and sulfonation intensity of 1:10), and sulfonation intensities from 1:1 137 to 1:10 (in 1:10 intervals, HT temperature of 250 °C, and HT time of 6 h). 10 g/L HS-PVC 138 fabricated with different HT intensities and sulfonation were used for Cu(II) and Cr(VI) 139 removal at 25 °C and 100 rpm. To investigate the individual effects of the intensities of HT 140 temperature, HT time, and sulfonation on Cu(II) and Cr(VI) removal, enhancements of Cu(II) 141 and Cr(VI) removal were calculated by comparing the lowest value with the highest one (250 °C 142 vs. 220°C, 6 h vs. 2 h, and 1:10 vs. 1:1, respectively).

#### 143 2.2.3. Adsorption kinetics and isotherms of Cu(II) and Cr(VI) removal using HS-PVC

The adsorption kinetics of Cu(II) and Cr(VI) removal were studied based on the removal efficiencies of HS-PVC as described in Section 2.2.1. Adsorption isotherm experiments were carried out with different initial concentrations of Cu(II) (i.e., 100, 200, 300, 400, 500, 600,

- and 800 mg/L) and Cr(VI) (50, 100, 150, 200, 250, and 300 mg/L) at three different
  temperatures (25, 35, and 45 °C) using 10 g/L HS-PVC under optimum operational
  conditions (250°C, 6 h, and a sulfonation intensity of 1:10).
- Pseudo-first-order (PFO) and pseudo-second-order (PSO) models were used to evaluate
  Cu(II) and Cr(VI) removal kinetics of HS-PVC, and their linearized equations are given in *Eqs.* (2) and (3), respectively.
- 153  $\ln(q_e q_t) = \ln q_e k_1 t$  (2)

154 
$$\frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \frac{t}{q_e}$$
(3)

where  $k_1$  and  $k_2$  are the rate constants of PFO and PSO, respectively, and  $q_e$  and  $q_t$  (mg/g) are the amounts of adsorbed Cu(II) and Cr(VI) (mg/g) at equilibrium and at time t, respectively. Values of  $k_1$ ,  $k_2$ , and  $q_e$  were calculated based on the Levenberg-Marquardt method at the 95% confidence level.

To predict the adsorption capacity of the adsorbents, two widely used isotherms (i.e., Langmuir and Freundlich) were selected among several models. *Eqs.* (4) and (5) express the Langmuir isotherm and the Freundlich isotherm, respectively.

$$q_e = \frac{bq_m c_e}{1 + bc_e} \tag{4}$$

$$q_e = K_F C_e^{1/n} \tag{5}$$

where  $q_m$  (mg/g) is the maximum adsorption capacity of the adsorbent and b (L/mg) is the Langmuir constant related to the adsorption energy.  $K_F$  (mg/g) is the Freundlich adsorption isotherm constant related to the extent of adsorption and n is the heterogeneity factor of the Freundlich model. Values of b,  $q_m$ ,  $K_F$ , and n were also calculated based on the

162

168 Levenberg-Marquardt method at the 95% confidence level.

# 169 2.2.4. Competitive adsorption of Cu(II) and Cr(VI) onto HS-PVC in binary solution and 170 preloading adsorption

Binary adsorption of Cu(II) and Cr(VI) was carried out by adding 10 g/L of HS-PVC to a binary solution with 500 mg Cu(II)/L and 250 mg Cr(VI)/L in the same solution to investigate the competitive adsorption between Cu(II) and Cr(VI). A separation factor was utilized to investigate the adsorption selectivity in the binary solution, defined as:

175 
$$\alpha_2^1 = \frac{q_{e1} \times c_{e2}}{c_{e1} \times q_{e2}}$$
 (6)

where  $q_{e1}$  (mmol/g) and  $C_{e1}$  (mmol/L) were the equilibrium adsorption capacity and equilibrium concentration toward constituent 1, and  $q_{e2}$  (mmol/g) and  $C_{e2}$  (mmol/L) were the equilibrium adsorption capacity and equilibrium concentration toward constituent 2, respectively.

Preloading adsorption tests for Cr(VI) were performed by incubating 2.5 g HS-PVC in 250 mg/L Cr(VI) solution for 12 h. Then, the HS-PVC preloaded with Cr(VI) was used for the adsorption of Cu(II) for 12 h, as described in Section 2.2.1. The preloading adsorption tests for Cu(II) were performed under the same experimental conditions as those for Cr(VI). The removal efficiencies of Cu(II) and Cr(VI) were calculated to evaluate the competitive adsorption between Cu(II) and Cr(VI) onto the HS-PVC.

#### 186 2.2.5. Effect of K(I) on Cr(VI) removal

187 The cations K(I) was present in the experimental Cr(VI) solutions prepared from
188 K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub>. To assess the impact of K(I) on the adsorption of Cr(VI) by HS-PVC, HClO<sub>4</sub> (36.62)

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mL, calculated as described in Text S1) was added to precipitate K(I) (*Ksp* of KClO<sub>4</sub>  $1.5 \times 10^{-2}$ )

- 190 in the Cr(VI) solution (total volume 100 mL). Then, HS-PVC (10 g/L) was added to the
- 191 treated Cr(VI) solution for adsorption for 12 h. Finally, the removal efficiency of Cr(VI) was
- 192 calculated to evaluate the impact of K(I) on the adsorption of Cr(VI) using HS-PVC.

#### 193 2.2.6. Low concentration Cu(II) and Cr(VI) removal using HS-PVC

The concentrations of Cu(II) and Cr(VI) in industrial applications, such as electroplating, leather tanning, and alloy manufacturing, range from 20 to 600 mg/L. Thus, 10 g/L of HS-PVC was added to 20 mg/L Cu(II) and Cr(VI) solutions to evaluate its adsorption performance for low concentrations of these heavy metals. The adsorption procedure was the same as in Section 2.2.1.

#### 199 2.2.7. Cu(II) and Cr(VI) removal in natural water

To test the potential interference of natural water regarding Cu(II) and Cr(VI) adsorption on HS-PVC, natural water (untreated surface water) was sampled from the Jingyue Lake in Donghua University, Shanghai, China. This water was used to prepare Cu(II) (500 mg/L) and Cr(VI) (250 mg/L) solutions. 10 g/L of HS-PVC was added to these Cu(II) and Cr(VI) solutions and incubated for 12 h at 25°C and 100 rpm.

#### 205 2.2.8. Cu(II) and Cr(VI) removal using HS-PVC derived from PVC waste pipes

To test the applications of HS-PVC in Cu(II) and Cr(VI) removal, real PVC waste pipes were used to prepare HS-PVC (real waste PVC). First, the PVC waste pipes were fragmented with a crusher and then washed with tap water. Subsequently, 10 g/L of the pretreated HS-PVC from pipes was added to Cu(II) (500 mg/L) and Cr(VI) (250 mg/L) solutions for 12

#### 210 h adsorption (at 25°C and 100 rpm).

#### 211 **2.3.** Characterizations of PVC-derived adsorbents

#### 212 **2.3.1.** Characterizations of Raw PVC, H-PVC, S-PVC, and HS-PVC

213 The surface properties of Raw PVC, H-PVC, S-PVC, and HS-PVC were characterized by 214 scanning electron microscopy (SEM) (HITACHI, S-4800, Japan) and contact angle 215 measurements ( $\theta$ ) (Kino, SL200KS, USA). The specific surface area and pore distribution of 216 Raw PVC, H-PVC, S-PVC, and HS-PVC were determined by the Brunauer-Emmett-Teller 217 (BET) (Quantachrome, Autosorb-iQ, USA) method and mercury intrusion porosimetry (MIP) 218 (Micromeritics, Auto pore lv, USA), respectively. Particle size distributions were calculated based on the SEM images (Table S1). The elemental composition of Raw PVC, H-PVC, 219 220 S-PVC, and HS-PVC was analyzed by elemental analysis (Elementar, EL III, Germany) and 221 energy dispersive spectroscopic (EDS) (IE 300 X, England) element mapping. The variation 222 of functional groups of Raw PVC, H-PVC, S-PVC, and HS-PVC was characterized by Fourier 223 transform infrared (FTIR) spectroscopy (Thermo Fisher, NEX US-670, USA). The 224 thermostability of Raw PVC and HS-PVC was evaluated by thermogravimetric analysis (TGA, 225 TA, Netzsch, Germany).

#### 226 2.3.2. Characterizations of HS-PVC before and after adsorption of Cu(II) and Cr(VI)

The electric potentials of HS-PVC, HS-PVC after adsorption of Cu(II) (HS-PVC-Cu), and HS-PVC after adsorption of Cr(VI) (HS-PVC-Cr) were determined by zeta potential measurements (Brook Haven, 90 PALS, USA). In addition, the functional groups of HS-PVC and HS-PVC-Cr were analyzed by X-ray photoelectron spectroscopy (XPS) (Thermo

231	Scientific, K-Alpha, USA), FIIR, solid-state "C nuclear magnetic resonance spectroscopy
232	(NMR) (Agilent, 600M, USA), and Raman spectroscopy (Thermo Fisher, DXR, USA). The
233	C1s XPS spectra were separated and characterized by C=C (284.8 eV), C-C (285.1 eV), C-O
234	(286.1 eV), and C=O groups (288.7 eV). The regions of functional groups derived from the
235	solid-state <sup>13</sup> C NMR spectra were divided into Carboxylic (220-161), Aromatic (161-113),
236	Anomeric (113-93), Carbohydrate (93-44), and Alkyl (44-0) (Wang et al., 2016, Li et al.,
237	2019).
238	2.4. Statistical analysis
220	

All adsorption experiments were run in triplicate, and the results were expressed as mean values  $\pm$  standard deviation. The triplicate PVC-derived adsorbents were pooled to minimize the potential variation before its characterizations (SEM, MIP, elemental analysis, FTIR, TGA, XPS, solid-state <sup>13</sup>C NMR, and Raman spectroscopy). Analyses of variance (ANOVA) were used to test for differences between treatments, and *p*<0.05 was considered statistically significant.

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001

#### 246 **3. Results and discussion**

### 247 **3.1.** Highly efficient Cu(II) cations and Cr(VI) anions removal using HS-PVC

Regarding Cu(II) removal, HS-PVC was significantly more efficient  $(81.2 \pm 1.6\%)$ compared to Raw PVC  $(28.5 \pm 1.7\%)$  and PVC solely pretreated by either HT  $(14.5 \pm 3.2\%)$ or sulfonation  $(20.6 \pm 4.8\%)$  within 12 h (Fig. 1A). In addition, HS-PVC was also highly efficient in removing Cr(VI)  $(60.3 \pm 3.8\%)$  compared to Raw PVC  $(6.1 \pm 1.3\%)$  and PVC 252 solely pretreated by either HT (14.0  $\pm$  4.6%) or sulfonation (7.0  $\pm$  4.8%) within 12 h (Fig. 1B). The adsorption of Cu(II) reached equilibrium within 30 min while the adsorption 253 254 efficiency of Cr(VI) was only  $49.3 \pm 1.0\%$  at 30 min and then slowly increased to equilibrium 255 within 3 h (Fig. S1). This suggested different adsorption mechanisms for Cu(II) and Cr(VI) 256 onto HS-PVC. Biomass-derived hydrochar was also widely employed in the adsorption of 257 metal cations, such as Cu(II), Zn(II), and Pb(II) (Deng et al., 2020, Kazak and Tor 2020, Song et al., 2020) or metal anions, such as Cr(VI) ( $Cr_2O_7^{2-}$ ) (Li et al., 2020). Cu(II) and Cr(VI) 258 259 could be simultaneously removed by hydrochar derived from chitosan with Fe<sub>2</sub>O<sub>3</sub> 260 modifications. However, the main adsorption mechanisms were chelation adsorption of Cu(II) and electrostatic adsorption-coupled Cr(VI) reduction due to the specific dual-core Fe<sub>2</sub>O<sub>3</sub>@C 261 262 structure (Wang et al., 2017). The present study firstly provided a metal-free modification of hydrochar derived from waste PVC for simultaneous Cu(II) cations and Cr(VI) anions 263 264 removal.

The adsorption capacity of hydrochar was highly dependent on the functional groups 265 formed during HT (Tong et al., 2021). However, the structure and component of PVC 266 267 ([-CH<sub>2</sub>-CHCl-]n) were simpler compared to natural biomass, suggesting fewer functional 268 groups anchored in H-PVC than in natural biomass. This may explain why H-PVC is of little significance for the adsorption of Cu(II) and Cr(VI) (Fig. 1), which was in line with poor 269 270 adsorption in wastewater remediation (Poerschmann et al., 2015). Sulfonation was also an 271 efficient modification of adsorbents derived from tea waste and cellulose for Cu(II) or Cr(VI) removal (Dong et al., 2016, Ahsan et al., 2018). However, the adsorption of Cu(II) and Cr(VI) 272

using S-PVC was still low in this study. This paper showed a possible synergistic effect of
HT and sulfonation in the modifications of the methodology to produce a highly efficient
adsorbent.

# 276 3.2. Characterization during the fabrications of HS-PVC from Raw PVC in 277 combination with HT and sulfonation

278 The microstructure of the PVC-derived adsorbents was characterized by SEM (Fig. 2A to 2D). A rough surface and non-porous structures were observed in Raw PVC (Fig. 2A). 279 After HT treatment, the surface of H-PVC became smoother with an internal porous structure 280 281 (Fig. 2B) mainly due to dehydrochlorination, aromatization, and depolymerization reactions 282 during HT (Ning et al., 2020). There was no apparent difference between Raw PVC and 283 S-PVC in the SEM images (Fig. 2A and 2C), indicating that sulfonation possessed a marginal 284 impact on the surface structure. In contrast, the HS-PVC was fragmented with an internal 285 porous structure due to HT and sulfonation (Fig. 2D), which was consistent with a significant 286 decrease of particle size in HS-PVC compared to the Raw PVC (Fig. 3A and Table S2). In 287 addition, the hydrophobicity and hydrophilicity of adsorbents may influence the performance 288 of the adsorption process (Adam et al., 2020). The hydrophobicity of S-PVC was similar to that of Raw PVC (Fig. 2E and 2G). However, the hydrophobicity was higher after HT (Fig. 289 290 2E and 2F), resulting in hydrophobic partitioning as described in the literature (Poerschmann 291 et al., 2015), which was consistent with the low adsorption efficiency of Cu(II) and Cr(VI) by 292 H-PVC (Fig. 1). Furthermore, the hydrophilicity of HS-PVC was significantly higher than 293 that of H-PVC (Fig. 2F and 2H), indicating that sulfonation improved the hydrophilicity of HS-PVC, which is a prerequisite for the adsorption of Cu(II) and Cr(VI).

The BET surface areas of Raw PVC (2.59 m<sup>2</sup>/g), H-PVC (4.95 m<sup>2</sup>/g), S-PVC (2.01  $m^2/g)$ , and HS-PVC (0.59 m<sup>2</sup>/g) (Table S3) were relatively low compared to those of biomass-derived hydrochar (118 m<sup>2</sup>/g) in the literature (Sheng et al., 2019). According to the analysis of MIP (Fig. S2), the pore size in HS-PVC was mainly in the mesoporous (2-50 nm) and microporous (>50 nm) regions (Erdem et al., 2020, Rueda-Marquez et al., 2021). The results revealed that the Cu(II) and Cr(VI) adsorption mostly occurred in the mesoporous and microporous regions of the HS-PVC.

302 Elemental analysis (Fig. 3B) showed that the Cl/C molar ratio (0.5) of Raw PVC was 303 similar to the stoichiometric value of ([-CH<sub>2</sub>-CHCl-]<sub>n</sub>). However, the molar ratio of Cl/C on 304 the surface of Raw PVC ( $0.85 \pm 0.37$ ) was higher than 0.5 according to EDS element 305 mapping (Fig. 3C), suggesting that most of the -Cl functional groups were on the surface of 306 Raw PVC. The Cl/C ratio of H-PVC significantly decreased (Fig. 3B and 3C) compared to 307 the Raw PVC after pretreatment by HT due to the dehydrochlorination (Poerschmann et al., 308 2015). The elemental composition of the entire Raw PVC and S-PVC was similar (Fig. 3B), 309 but surface Cl was lower, and surface O slightly higher in S-PVC than in Raw PVC (Fig. 3C). 310 This indicated that sole sulfonation only impacted the chemical composition on the surface of Raw PVC (Fig. 3C), causing no obvious chemical modifications to the entire Raw PVC (Fig. 311 3B). Additionally, the O/C and S/C ratios of HS-PVC increased, resulting in an O/S ratio of 312 3.04 (elemental analysis in Fig. 3B) or 3.16 (EDS element mapping in Fig. 3C). Thus, it 313 314 could be hypothesized that the introduction of S-containing functional groups such as -SO<sub>3</sub>H

315	(O/S=3) into HS-PVC occurred. The absorption peaks of HS-PVC from the FTIR at 1030
316	$cm^{-1}$ (O=S=O) and 1169 $cm^{-1}$ (-SO <sub>3</sub> H) verified this assumption (Fig. 3D) (Chen et al., 2019).
317	The functional group -SO <sub>3</sub> H was only detected in HS-PVC, indicating that the introduction of
318	-SO <sub>3</sub> H onto HS-PVC was attributed to the synergy effect of HT and sulfonation. In addition,
319	the thermostability of HS-PVC was higher than that of Raw PVC according to the TGA
320	patterns (Fig. S3), which was mainly due to aromatization during HT (Park et al., 2016). It
321	has been reported that the presence of aromatic groups was essential for the sulfonation of
322	petroleum coke by concentrated H <sub>2</sub> SO <sub>4</sub> (Xiao and Hill 2020). In this study, HT was crucial to
323	forming an aromatic structure (Fig. 3D) for the introduction of -SO <sub>3</sub> H during sulfonation to
324	fabricate HS-PVC for highly efficient Cu(II) and Cr(VI) removal. Moreover, the functional
325	group C=C in HS-PVC (Fig. 3D) was crucial for removing Cr(VI), which was confirmed in
326	the following section.

#### 327 3.3. Correlations of Cu(II) and Cr(VI) removal with HT intensities and sulfonation

The adsorption properties of HS-PVC were highly dependent on HT temperature and 328 time (see time curves in Fig. S4 and Fig. S5). The adsorption efficiencies of Cu(II) and Cr(VI) 329 330 increased from 36.9  $\pm$  3.0% and 46.0  $\pm$  3.8% at 220 °C to 81.1  $\pm$  2.1% and 61.7  $\pm$  2.7% at 250 °C, but declined to  $63.7 \pm 3.2\%$  and  $57.2 \pm 1.3\%$  at 260 °C, respectively (Fig. 4A). In 331 332 addition, the adsorption efficiencies of Cu(II) and Cr(VI) also increased from  $56.5 \pm 1.7\%$ and 9.2  $\pm$  1.1% at 2 h to 80.0  $\pm$  2.8% and 58.5  $\pm$  1.1% at 6 h, followed by declines to 54.5  $\pm$ 333 334 2.0% and 2.2  $\pm$  1.2% at 8 h, respectively (Fig. 4B). Higher HT intensities benefited PVC dehydrochlorination (Zhang et al., 2020), providing more sites for subsequent SO<sub>3</sub>H 335

introduction during sulfonation. It has been reported that proper HT intensities (HT temperature =  $250 \,^{\circ}$ C and HT time = 6 h in this study) benefited the formation of the aromatic structure (Ning et al., 2020). In contrast, higher HT intensities affected the stability of the carbon structure (Ning et al., 2020), which explained the declines in the adsorptions of Cu(II) and Cr(VI) onto HS-PVC at 260 °C (Fig. 4A) and 8 h (Fig. 4B).

The sulfonation intensity also affected the introduction of functional groups (Konwar et al., 2019), thus affecting the adsorptive performance of HS-PVC. The adsorption time curves of sulfonation intensity regarding Cu(II) and Cr(VI) removal by HS-PVC were shown in Fig. S6. With the increasing sulfonation intensity, the adsorption efficiencies of Cu(II) and Cr(VI) increased from  $38.1 \pm 3.2\%$  and  $28.4 \pm 3.5\%$  at the sulfonation intensity of 1:1 to  $81.0 \pm 1.3\%$ and  $59.6 \pm 2.4\%$  at the sulfonation intensity of 1:10 (Fig. 4C). The adsorption of Cu(II) and Cr(VI) was positively correlated with the increase in sulfonation intensity from 1:1 to 1:10.

348 As summarized in Fig. 4D, the adsorption efficiency of Cu(II) increased by 117.4% 349 while that of Cr(VI) only by 34.1%, when increasing the temperature from 220 °C to 250 °C. 350 In contrast, the adsorption efficiency of Cu(II) improved only by 41.6%, while that of Cr(VI) 351 increased by 534%, with HT time increasing from 2 to 8 h. These results indicated that the 352 adsorption of Cu(II) was sensitive to HT temperature while the adsorption of Cr(VI) was rather sensitive to HT time. However, the impact of sulfonation intensity on the adsorption of 353 Cu(II) (increasing 112.5% from 1:1 to 1:10) was similar to the adsorption of Cr(VI) (increasing 354 110.3% from 1:1 to 1:10). 355

#### 3.4. Kinetics and isotherms of the adsorption of Cu(II) and Cr(VI) onto HS-PVC

357 Adsorption kinetics were essential to investigate the adsorption process (Song et al., 358 2020, Wang and Guo 2020). The adsorption of Cu(II) and Cr(VI) onto the HS-PVC (Fig. 5A 359 and 5B) increased rapidly in the first 30 min and then decreased as the reaction approached 360 equilibrium due to the saturation of active adsorbent sites. The values of  $q_e$  and rate constants 361  $(k_1 \text{ and } k_2)$  obtained from the kinetic models (Fig. 5A and 5B) were shown in Table 1, along with the correlation coefficients ( $R^2$ ). The  $R^2$  value of the PSO model was higher than that of 362 the PFO model for both Cu(II) and Cr(VI) adsorption. In addition, the  $q_e$  calculated by the 363 364 PSO model was more consistent with the experimental value compared to that of the PFO model (Table 1). Hence, the adsorption process of Cu(II) and Cr(VI) onto HS-PVC could be 365 366 well described by the PSO model, indicating that chemisorption might be the rate-limiting step. In addition, more active sites were reported with the PSO model compared to the PFO 367 368 model (Wang and Guo 2020), which was crucial to the adsorption of Cu(II) and Cr(VI) onto 369 HS-PVC.

Adsorption isotherms play a crucial role in investigating the chemical interactions between adsorbates and adsorbents (Yang et al., 2019, Wang and Guo 2020). The adsorption isotherms of the adsorption of Cu(II) and Cr(VI) onto HS-PVC at three different temperatures (25, 35, and 45°C) were shown in Fig. 5C-5F. During the adsorption of Cu(II), the  $R^2$  values of the Langmuir model (0.94–0.96) were higher than those of the Freundlich model (0.82– 0.89) at all tested temperatures (25, 35, and 45 °C) (Fig. 5C, 5D, and Table S4), indicating that the adsorption of Cu(II) by HS-PVC was a monolayer process. Similarly, the adsorption of Cr(VI) was also a monolayer process due to the higher  $R^2$  values of the Langmuir model (0.94 at 25 °C and 0.95 at 35 °C) compared to those of the Freundlich model (0.90 at 25 °C and 0.89 at 35 °C) (Fig. 5E, 5F, and Table S4). However, the adsorption of Cr(VI) was fitted by the Langmuir and Freundlich model equally well according to  $R^2$  values of 0.96 at 45°C. In addition, the values of  $K_F$  and  $q_m$  increased with the increasing temperature (Table S4), indicating an endothermic adsorption process (Xu et al., 2019).

#### 383 **3.5.** Mechanisms for the adsorption of Cu(II) and Cr(VI) onto HS-PVC

The removal efficiency of Cu(II) decreased (p < 0.01) by 19.9 % with the co-adsorption 384 385 of Cr(VI), and that of Cr(VI) also significantly decreased (p < 0.01) by 48.8% with the co-adsorption of Cu(II) (Fig. 6A). The selectivity of Cu(II) ( $\alpha_{Cr(VI)}^{Cu(II)} = 4.11$ ) was higher than that 386 of Cr(VI) ( $\alpha_{Cr(VI)}^{Cu(II)} = 0.24$ ) in binary-component solution, indicating that Cu(II) can occupy 387 388 adsorption sites more easily than Cr(VI). However, during the preloading test, the removal efficiency of Cu(II) decreased by 70.5% using Cr(VI) loaded HS-PVC, which was similar to 389 390 the decrease in Cr(VI) removal (65.8%) using Cu(II) loaded HS-PVC (Fig. 6A). These results 391 suggested that the Cu(II) and Cr(VI) adsorption might share similar adsorption sites on 392 HS-PVC. The zeta potential, an important index for the electric potential at the mineral surface (Ramasamy et al., 2017, Gu et al., 2021), significantly decreased after HT and 393 394 sulfonation (Fig. S7). The value increased after the adsorption of Cu(II) (Fig. 6B), indicating 395 electrostatic adsorption (Fig. 7). However, only a slight increase in zeta potential was 396 observed after the Cr(VI) adsorption, implying that other mechanisms govern Cr(VI) 397 adsorption onto HS-PVC.

398	To further explore the mechanism of the adsorption of Cr(VI) onto HS-PVC, XPS
399	analysis was used to investigate the adsorption behavior of hydrochar. The C1s XPS spectra
400	were separated, and the characteristic C=C, C-C, C-O, and C=O bonds were quantified in
401	HS-PVC before and after Cr(VI) adsorption. The intensity of C=C in HS-PVC decreased
402	from 35.5% (Fig. 6C) to 28.5% (Fig. 6D) after adsorption of Cr(VI) (Fig. 6C), suggesting the
403	involvement of this bond in Cr(VI) removal. According to solid-state <sup>13</sup> C NMR, aromatic
404	functional groups decreased from 55.0% (Fig. 6 E) to 51.4% (Fig. 6F) after Cr(VI) adsorption,
405	also supporting a decrease of C=C intensity after the adsorption of Cr(VI). In addition, the
406	degree of graphitization of HS-PVC decreased after Cr(VI) adsorption according to the
407	decrease in the ratio of $I_G/I_D$ from 0.77 to 0.73 (Fig. 6G), which might be related to the
408	reduction of Cr(VI). Interestingly, Cr(III) was detected in HS-PVC after the adsorption of
409	Cr(VI) (Fig. 6H). These results indicated that the functional group C=C and aromatic groups
410	might reduce Cr(VI) to Cr(III). In addition, there were shifts of the ring vibration peak in the
411	FTIR spectra after adsorption of Cr(VI) (arrows in Fig. 6I). Since a ligand aligns itself to a
412	metal, the energy of the ligand material will most likely be perturbed, leading ultimately to
413	subtle shifts in the absorption peaks (Sergios et al., 2010). Thus, HS-PVC might also bind
414	Cr(VI), which was consistent with Cr(III) and Cr(VI) in Fig. 6H. It has been reported that
415	Cr(VI) was complexed with phenolic -OH using hydrochar derived from leaves and wood
416	sticks. The adsorbed Cr(VI) was subsequently reduced to Cr(III) by the hydrochar along with
417	the decrease of the functional group C=C (Chen et al., 2017, Chen et al., 2021). In this study,
418	the Cr(VI) was supposed to be complexed by the phenolic -OH (Fig. 6I) existed in HS-PVC

419 and reduced by the C=C in the aromatic structure formed by HT (Fig. 7).

Note that the removal efficiency of Cr(VI) significantly decreased from 63.8  $\pm$  2.0% in the presence of K(I) to 28.2  $\pm$  5.1% after K(I) was precipitated by HClO<sub>4</sub> (Fig. S8). The introduction of H<sup>+</sup> ions increased the proportion of H<sub>2</sub>CrO<sub>4</sub>, thereby breaking the equilibrium balance of HCrO<sub>4</sub><sup>-</sup> and Cr<sub>2</sub>O<sub>7</sub><sup>2-</sup> due to negative charges. This, in turn, suggested that K(I) ions might neutralize the negatively charged HS-PVC surface, thereby facilitating the binding of dichromate ions on HS-PVC (Fig. 7).

#### 426 **3.6. Versatility of HS-PVC**

427 Natural water contains ions and organic compounds (Jeon et al., 2020), which might hinder Cu(II) and Cr(VI) removal by HS-PVC. Due to the competitive adsorption of 428 co-existing ions, the efficient removal of Cu(II) and Cr(VI) decreased only by 19.2% and 25.0% 429 430 (Fig. 8), respectively, indicating low interference. In addition, HS-PVC was particularly 431 suitable for the removal efficiencies of Cu(II) and Cr(VI) at low concentrations (20 mg/L) 432  $(99.6 \pm 0.1\%$  and  $94.3 \pm 2.5\%$ , respectively) (Fig. 8). Further, the removal efficiencies of Cu(II) 433 and Cr(VI) using HS-PVC from waste pipes increased by 13.7% and 43.9% (Fig. 8), which 434 were higher than those of HS-PVC derived from PVC powder (Fig. 1). The better performance 435 of HS-PVC (pipes) might be attributed to plasticizers and mineral fillers in the PVC (Correa et al., 2019). The removal efficiencies of other metals such as Mn(II), Cd(II), Ni(II), and Co(II) 436 amounted to  $81.2 \pm 2.3\%$ ,  $80.8 \pm 1.7\%$ ,  $70.8 \pm 1.8\%$ , and  $70.1 \pm 3.2\%$  (Fig. S9), respectively, 437 438 indicating that HS-PVC is suitable for the treatment of complex industrial wastewater. 439 Moreover, the adsorption of Cu(II) and Cr(VI) decreased only by 10.0% and 31.0% (Fig. S10)

440 after three adsorption-desorption cycles (1 mol/L HCl as eluent), indicating excellent441 recyclability.

442 Previous studies commonly focused on the conversion of waste PVC into coal-alternative fuel (Lu et al., 2020) or lighter oil (Maity et al., 2020). Using HT and 443 444 sulfonation modifications, the present study provided a novel strategy for resource recovery 445 of PVC waste by producing efficient adsorbents for Cu(II) cations and Cr(VI) anions removal. 446 The sulfonation intensity was mild in this study (98% sulfuric acid at 70°C). Furning H<sub>2</sub>SO<sub>4</sub>, 447 ClSO<sub>3</sub>H, aryldiazonium sulfonates, and higher temperatures (80–150°C) could be considered 448 for additional optimization. In addition, as it is a "sulfonated carbon" (Konwar et al., 2019), HS-PVC may show promising catalytic performances in various organic transformations, 449 450 including esterifications, ester exchange (transesterifications), acetalization, etherifications, condensation, and multi-component organic reactions that should be studied in the future. 451

#### 452 **4. Conclusion**

This study provided a novel method to fabricate efficient adsorbent (HS-PVC) from 453 454 waste PVC for both Cu(II) cations and Cr(VI) anions removal. The removal efficiencies of 455 Cu(II) and Cr(VI) using HS-PVC reached  $81.2 \pm 1.6\%$  and  $60.3 \pm 3.8\%$ , respectively. The first stage of HT was crucial for the dichlorination of PVC and the formation of an aromatic 456 structure, guaranteeing the introduction of -SO<sub>3</sub>H onto PVC-derived hydrochar through 457 subsequent sulfonation. In addition, sulfonation also significantly increased the hydrophilicity 458 459 of HS-PVC. The adsorption of Cu(II) was sensitive to HT temperature, while that of Cr(VI) 460 was more sensitive to HT time. Adsorption isotherm data were accurately fitted by the

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Langmuir isotherm model with a maximum adsorption capacity of 48.9 mg/g for the 461 adsorption of Cu(II) and 20.2 mg/g for the adsorption of Cr(VI) at 45°C. Competitive 462 463 adsorption of Cu(II) and Cr(VI) occurred, and the selectivity of HS-PVC for the removal of Cu(II) was higher than that of Cr(VI). The adsorption mechanism of Cu(II) was electrostatic 464 adsorption, while Cr(VI) were possibly complexed by the phenolic -OH and reduced to Cr(III) 465 466 cations by C=C groups in HS-PVC. In addition, HS-PVC was particularly suitable for the adsorption of both metals at low concentrations (>94% at 20 mg/L). HS-PVC derived from 467 468 PVC waste pipes also showed high absorption efficiencies for Cu(II) and Cr(VI) (>90%).

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### 649 **Table 1**

	Metals	$q_{ m exp}$	PFO			PSO		
		(mg/g)	$q_1$ (mg/g)	$R_l^2$	$k_1$ (min <sup>-1</sup> )	$q_2 (\mathrm{mg/g})$	$R_2^2$	$k_2$ (g/(mg·min))
	Cu(II)	40.34	38.18	0.86	17.92	40.24	0.95	0.70
	Cr(VI)	17.75	15.41	0.88	2.39	16.23	0.94	0.25
651								
652 653								

650 PFO and PSO model parameters for Cu(II) and Cr(VI) adsorption onto HS-PVC.





656 Fig. 1. Cu(II) and Cr(VI) removal efficiencies using Raw PVC, H-PVC, S-PVC, and

657 HS-PVC after adsorption for 12 hours.



660 Fig. 2. Surface properties of PVC derived adsorbents: SEM images of Raw PVC (A), H-PVC

661 (B), S-PVC (C), and HS-PVC (D); contact angle ( $\theta$ ) of Raw PVC (E), H-PVC (F), S-PVC

662 (G), and HS-PVC (I).



665 Fig. 3. Characterization of Raw PVC, H-PVC, S-PVC, and HS-PVC: particle size distribution

666 (A), elemental analysis (B), EDS element mapping (C), and FTIR spectra (D).



668

Fig. 4. Effects of HS-PVC preparation intensities on Cu(II) and Cr(VI) removal: HT temperature (A), HT time (B), sulfonation intensity (C), and comparison of Cu(II) and Cr(VI) removal efficiencies between the optimal and the lowest efficiency (D) (HT temperature: 250 °C vs. 220 °C, HT time: 6 h vs. 2 h, and sulfonation intensity: 1:10 vs. 1:1). The initial concentrations of Cu(II) and Cr(VI) were 500 and 250 mg/L. The adsorbent dose was 10 g/L and the temperature 25 °C.

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Fig. 5. Adsorption kinetics of Cu(II) (A) and Cr(VI) (B) onto HS-PVC. Equilibrium isotherms of Cu(II) fitted by the Langmuir (C) and Freundlich models (D) at different temperatures (25, 35, and 45 °C). Equilibrium isotherms of Cr(VI) fitted by the Langmuir (E) and Freundlich models (F) at different temperatures (25, 35, and 45 °C). The colored area covers the 95% confidence intervals.



Fig. 6. Mechanisms for the adsorptions of Cu(II) and Cr(VI) by HS-PVC. Competitive 683 684 adsorption between Cu(II) and Cr(VI) by HS-PVC (A). Zeta potential of HS-PVC, HS-PVC after the adsorption of Cu(II) (HS-PVC-Cu), and HS-PVC after the adsorption of Cr(VI) 685 (HS-PVC-Cr) (B). C 1s XPS spectra of HS-PVC before the adsorption of Cr(VI) (C) and 686 687 after the adsorption of Cr(VI) (D). Solid-state <sup>13</sup>C NMR of HS-PVC before the adsorption of Cr(VI) (E) and after the adsorption of Cr(VI) (F). Raman spectra of HS-PVC before and after 688 689 adsorption of Cr(VI) (a higher I<sub>G</sub>/I<sub>D</sub> ratio indicates a higher degree of graphitization) (G). Cr 2p XPS spectra of HS-PVC after the adsorption of Cr(VI) (H). FTIR spectra of HS-PVC 690 691 before and after the adsorption of Cr(VI) (I).







Fig. 8. Versatility of HS-PVC for Cu(II) and Cr(VI) removal efficiencies. Removal of
Cu(II) and Cr(VI) by HS-PVC from natural water, at low concentration (20 mg/L), and by
HS-PVC derived from waste PVC pipes (blue and red lines represent the removal efficiency
of Cu(II) and Cr(VI) by PVC powder in Fig. 1).