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Secondary particle size determining sedimentation and adsorption kinetics of titanate-based materials for ammonia nitrogen and methylene blue removal

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13 Abstract

The effect of the main size distribution of particles on the adsorption process for adsorbent materials has been well-recognized; however, the impact of secondary particles size (agglomerated, aggregated or hydrated ones) on adsorbent properties and performance was rarely reported so far. In this study, a series of sodium titanates (STs) and peroxide modified sodium titanates (PSTs) with different primary particle sizes, and secondary sizes are synthesized by controlling synthesis conditions and

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20	subsequently applied to batch adsorption experiment. By employing scanning electron
21	microscopes and Laser particle size analyzers, the particle size of STs and PSTs are
22	found to be closely correlated with synthesis conditions. The surface morphology and
23	specific surface area of titanates are size-dependent, while the components of all the
24	samples maintained constant. The sedimentation experiment and CFD simulation
25	demonstrated that particles with larger secondary sizes tended to settle more quickly
26	than those with a bigger sizes. PSTs or STs particles with smaller secondary sizes could
27	reach equilibrium more rapidly than those with the bigger size. The fitting results from
28	Elovich and Weber-Morris models demonstrated that the particles size affect kinetics
29	mainly through the liquid film diffusion process within the initial stages.
30	Keywords: Titanate; Secondary size; Sedimentation; Adsorption; Water treatment
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44 **1. Introduction**

Recently, there has been considerable concerns about emerging threats from the 45 total nitrogen (TN) and chemical oxygen demand (COD) in water and other ecosystems 46 [1, 2]. The removal of ammonia nitrogen (NH_4^+) or methylene blue (MB) from 47 wastewater has drawn much attention in water purification due to their great 48 contribution to TN or COD pollution, respectively [3, 4]. Adsorption, one of the 49 50 practical and popular physicochemical techniques used in water treatment, has been frequently employed for removing NH4⁺ and MB because of its high efficiency, cost-51 effective and easy operation [5, 6]. Numerous studies have been carried out to 52 53 understand the adsorption process [7]. It has been widely demonstrated that solution pH, environment temperature or contact time, and the properties of an adsorbent such 54 as surface charge, functional groups, or pore structure can significantly impact the 55 adsorption performance[8-12]. However, the particle size effect on the adsorbent 56 properties and adsorption process is less still unclear, even though most adsorbent 57 materials for NH₄⁺ or MB are powders or particles in nature [13, 14]. It has been shown 58 that many fundamental properties of powder materials like mechanical and electrical 59 performance are size-dependent when the diameter of particles are nanoscale [15]. 60 Indeed, particle size change could impact the movement of particles in solution 61 according to the Brownian motion. In addition, the particle size of materials could also 62

affect the value of surface free energy and then might have effects on the adsorptioninteraction [16].

It has been found that the adsorption of MB onto diatomite is size-dependent when 65 the size scale is larger than 250 µm, and the maximum uptake capacity decreases with 66 the reduction of size. The same phenomenon was found in the adsorption of bisphenol 67 A onto montmorillonite [16, 17]. However, the particle size value discussed in most 68 reports was determined directly from SEM image or sieve action, which could not show 69 the actual scale of particles in solution because of aggregation [18]. The size of solid 70 71 materials in an aqueous environment could be described through two different approaches: the primary size for individual particles and the secondary size for 72 agglomerated, aggregated or hydrated particles [19]. Therefore, the secondary size 73 74 value is closer to the actual conditions and appropriate for the size-dependent discussion. Titanate, one of the common inorganic ion-exchange adsorbents or photocatalysts 75 [20, 21], was chosen as the main adsorbent for size dependence experiment according 76 to the following two merits: 1) amorphous sodium titanate (ST) and peroxide modified 77 sodium titanate (PST) particles can be facilely synthesized through only mild 78 hydrolysis method. The secondary particle size can be easily controlled for ST and PST 79 by changing synthesis conditions such as the ratio and concentration of reagents. 2) 80 Titanate based materials exhibit excellent adsorption efficiency for cationic 81 contaminants due to their ion-exchange ability demonstrated by literature [22-25]. 82

In this study, NH₄⁺ and MB were chosen as the targeted contaminants not only
because of their contribution to TN and COD but also due to their large difference in

molecular weight and size, which might help uncover the relation between particle size 85 dependence and adsorbate molecular properties. Several challenges in secondary 86 particle size-dependent sedimentation and adsorption kinetics of titanate-based 87 materials for ammonia nitrogen and methylene blue removal have been systematically 88 studied: 1) a series of titanates named sodium titanates (STs) and peroxide modified 89 sodium titanates (PSTs) with different particle sizes were facilely synthesized by 90 changing synthesis conditions. 2) The effect of titanates particle secondary size on 91 adsorbent properties and adsorption performance in removal of NH4⁺ or MB were 92 conducted and discussed. 3) The potential mechanisms of the effect of particle size were 93 also revealed. 94

95 2. Experimental procedures

96 2.1. Materials

TIPT (titanium isopropoxide, Ti(OC₃H₇)₄), H₂O₂ (hydrogen peroxide, 30 wt%), 97 NaOH (sodium hydroxide), HCl (hydrochloric acid), anhydrous ethanol(99.7%), HDA 98 99 (hexadecylamine, 90 wt%) and isopropanol were of analytical reagent grades and used without any further purification. The simulated wastewater used in this study were 100 101 obtained by dissolving ammonium chloride (NH₄Cl, GR) or methylene blue (MB, BS, Fig. S1) (both from Sinopharm Chemical Reagent Co. Ltd, China) into deionized water. 102 The deionized water used for all the experiments was prepared by an EPED-40TF Super 103 pure Water System, China. 104

105 **2.2. Synthesis of titanate samples**

A series of sodium titanates (STs) with different particle sizes were prepared based 106 on our previous study [22]. The particle size of STs powders was adjusted by controlling 107 the hydrolysis rate of TIPT, where the volume of H₂O was changed according to the 108 molar ratio of TIPT: H₂O (1:1, 1:10, 1:20). HDA was served as a structure-directing 109 agent [26]. In a typical synthesis procedure, 0.8 g of NaOH and 2 g of HDA were 110 dissolved in a mixed solution of 200 mL ethanol and a certain amount of H_{2O} (0.61, 111 6.10 and 12.2 mL) with agitation at ambient temperature. 10 mL of TIPT was added 112 dropwise into the above solution under stirring. After 24 h, the white suspension was 113 recovered by centrifugation. The solid sample was washed with ethanol five times and 114 dried at 60°C for 12 h [27]. The as-prepared white samples were labelled as ST (1:1), 115 ST(1:10) and ST(1:20) according to the TIPT: H₂O molar ratio. All the samples were 116 117 further purified by the sieve with 200 mesh.

A series of peroxide modified sodium titanate (PSTs) with different particle sizes 118 were synthesized as follows: a mixture of TIPT (10 mL) and isopropanol with the 119 volume ratio of 5:2 was added into 200 mL NaOH solution (0.1 mol·L⁻¹) with magnetic 120 stirring at 60°C [28]. Afterwards, an appropriate amount of H₂O₂ (5 mL, 10 mL and 20 121 mL respectively) was dropwise added into the white suspension in 1 h, accompanied 122 by stirring. The solution gradually became transparent and bright yellow, and then the 123 yellow suspension was stirred for another 30 min at ambient temperature. The yellow 124 solid was filtered out and subsequently washed with deionized water until the filtrate 125 pH reached approximately 7.0, then dried at 60 °C for 12 h. The as-prepared PST 126 samples were denoted as PST-5, PST-10 and PST-20 according to the volume of H₂O₂ 127

used in the synthesis process, respectively. All the samples were sieved by the sievewith 200 mesh.

130 2.3. Analysis and characterization methods

The morphology of samples was investigated by scanning electron microscopy 131 (SEM, MAIA3 LMH, US). The imaging and elemental mapping were performed using 132 acceleration voltages of 15 kV. The element's contents of powder samples were 133 measured by X-ray Fluorescence (XRF, Bruker S8 Tiger, Germany) without any 134 pretreatment. The sample's secondary size in solution was determined by a laser particle 135 136 size analyzer (LS-909, China). The actual density of samples was tested by a Gas Displacement Pycnometer (AccuPyc II 1340, US). The surface functional groups of 137 adsorbents were identified by a Fourier transform infrared spectrometer (FTIR, Bruker, 138 Germany) with the KBr pellet method at the wavenumber ranging from 400 to 4000 139 cm^{-1} . The Brunauer-Emmett-Teller specific surface area (S_{BET}) and pore structure 140 characteristics of samples were determined by the Builder SSA-4300 (Beijing, China) 141 142 at 77 K using the Barrett-Joyner-Halenda (BJH) method. The zeta potentials of the samples were measured with Brookhaven 90plus Zeta, samples of which (1.0 mg) were 143 added into 10 ml NaCl solution (10⁻³ mol·L⁻¹) at different pH values (adjusted by 0.5 144 mol·L⁻¹ HCl and NaOH solution). 145

The control experiment was conducted to evaluate the property of as-prepared materials in solution as follows: 0.5 g of powders were added into a measuring cylinder with 25 mL deionized water and dispersed under ultrasound sonication for 30 min. Afterwards, the cylinder with PSTs or STs solution was put on the surface of a flat desk,respectively. The photos were taken at different time intervals.

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2.4. Numerical simulations

In the present study, Computational Fluid Dynamics (CFD) was also conducted to 152 reveal the relationship between settlement property and particles size. Considering the 153 symmetry of the test tube, a 2D axisymmetric domain with a height of 200 mm and a 154 width of 20 mm was employed. Structured grids were firstly generated for the 155 computational domain through software ICEM. Then, based on the finite volume 156 157 method, numerical simulation was carried out using the software ANSYS-Fluent to analyze unsteady flow behavior. The Eulerian multiphase model was utilized to model 158 granular flow, allowing two phases to share a single pressure and momentum, and 159 160 continuity equations are solved for each phase [29]. The phase coupled SIMPLE algorithm was used for the pressure-velocity coupling scheme. The velocities were 161 calculated coupled by phases in a segregated method, and the pressure correction 162 equations were established based on total continuity. According to ANSYS-Fluent 163 theory guide, the k- ε dispersed turbulence model was applicable when there is one 164 primary continuous phase, and the rest are dispersed dilute secondary phases. Thus, the 165 k- ε dispersed turbulence model with standard wall function was adopted to close the 166 turbulence terms in the simulation. Moreover, the gravitational acceleration was 167 specified as 9.81 m/s² to consider the gravity, and the Gidaspow method [30] was 168 implemented for the drag law. The thickness of the initial particle layer was set as 5 mm 169

with a volume fraction of 0.2. The models with different particle diameters and densitieswere constructed to study their effect on the setting ability.

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2.5. Adsorption experiments

Unless otherwise stated, NH_4^+ and MB removal experiments were performed at 25°C in a temperature-controlled shaker with a 200 rpm stirring rate. The initial concentration of NH_4^+ and MB were 45 and 275 mg·L⁻¹, respectively. The suspensions after adsorption were filtered in syringes equipped with 0.45 µm cellulose acetate membrane filters.

To investigate the effect of the solution pH on the adsorption of pollutants, 0.4 g of as-prepared STs or PSTs separately mixed with 20 mL individual NH4⁺ or MB solution at different pH values ranging from 2.0-12.0 (adjusted by either HCl or NaOH) in 50 mL centrifugal tube. The above solution was shaken for 120 min.

Adsorption kinetic experiments were carried out in a conical flask by adding 0.2 182 g of adsorbent into a 100 mL solution containing NH4⁺ or MB with initial pH of 7.0. At 183 various time intervals, 1.5 mL suspension was withdrawn and filtered for determination. 184 Adsorption isotherms experiments were performed as follows: 0.04 g of each 185 adsorbent (PSTs and STs) was added into a 50 mL centrifuge tube containing 20 mL 186 solution with different initial concentrations varying from 10 to 160 mg \cdot L⁻¹ for NH₄⁺, 187 50-300 mg·L⁻¹ for MB at pH=7.0. The mixture was shaken at ambient temperature for 188 120 min. 189

190 The NH4⁺ concentration remaining in the solution was determined by the 191 conventional salicylate spectrophotometric method. The MB concentration in solution 192 was determined by a direct UV-Vis spectrophotometry method at a wavelength of 665

nm. The amounts of adsorbates adsorbed on the as-prepared STs or PSTs at a certain 193 time Q_t (mg·g⁻¹) and equilibrium Q_e (mg·g⁻¹) were calculated by Eqs. (1)-(2). The 194 adsorption efficiency R (%) was calculated by Eq. (3). Where $C_0 (\text{mg} \cdot \text{L}^{-1})$ is the initial 195 concentration of adsorbates, C_t (mg·L⁻¹) and C_e (mg·L⁻¹) are the residual adsorbate 196 concentration at time t (min) and equilibrium; m (g) is the mass of adsorbent and V(L) 197 is the solution volume. As shown in Eqs. (S1)-(S6), four kinetic models (pseudo-first 198 199 order, pseudo-second order, elovich and web-morris models) and two isotherms models (Langmuir and Freundlich models) were used to analyze the adsorption data [31-33]. 200

201
$$Q_t = \frac{C_0 - C_t}{m} \times V \tag{1}$$

$$202 \qquad Q_e = \frac{C_0 - C_e}{m} \times V \tag{2}$$

$$R = \frac{C_0 - C_t}{C_0} \times 100 \tag{3}$$

203

204 **3. Results and discussion**

3.1. Structure and morphology analyses



Figure 1. SEM images of ST (1:1) (a), ST (1:10) (b), ST (1:20) (c), PST-5 (d), PST-10
(e) and PST-20 (f).

The morphology of the primary size samples was analyzed by SEM, as shown in 209 Fig. 1. It is observable that the diameter of granules in STs (Fig. 1a-c) decrease with the 210 increase of H₂O volume in the synthesis procedure. The reduction of the Ti/H₂O molar 211 ratio increased the hydrolysis rate of Ti precursor and cut down the production of large 212 particles [34]. The average primary size of ST(1:1), ST(1:10) and ST(1:20) is about 500 213 nm, 300 nm and 150 nm, respectively. Agglomeration occurs primarily on the surface 214 of ST(1:20) because the size of nano-particles is small [35]. The change of particles 215 size and morphology in STs might affect the sample properties like surface area and the 216 adsorption behavior in the aqueous phase [16]. The effect of synthesis condition on the 217 morphology of PSTs is more significant than that of STs, as shown in Fig. 1d-f. The 218 complexation reaction between H₂O₂ and Ti⁴⁺ can be affected by increasing H₂O₂ 219 content during synthesis and result in different morphology and particles size [36]. It 220 can be seen that the images that PST-5 (prepared with 5 mL H₂O₂) still maintains a 221 rough surface consisting of particles like STs. However, PST-10 and PST-20 exhibit a 222 well-organized morphology with tiny drapes and fewer particles, which might have a 223 negative effect on the pore volume and specific surface area of samples. All the above 224 results suggest that the modification process succeeded in changing the primary size of 225 samples, favouring the modulation of secondary size in the aqueous phase. 226

Though the primary size is influential on the property of the sample, the secondary size of powder samples is possibly more significant when served as adsorbents in the aqueous phase [19]. The actual particles size (secondary size) of STs and PSTs owing

230	to agglomeration or hydration in agitating solution was measured by a Laser particle
231	size analyzer and shown in Table 1. It is illustrated that the secondary size of samples
232	is greatly different from the primary size of SEM images. The results also indicate that
233	secondary particle size of STs and PSTs are in the order of ST(1:1)> ST(1:10)> ST(1:20)
234	and PST-20> PST-10> PST-5, respectively. Controlling the hydrolysis rate (for STs)
235	and complexation degree (for PSTs) successfully caused the difference in secondary
236	particle size. It is also indicated from Table 1 and Fig. S2 that the surface area and pore
237	volume of STs vary with the particle size value. The reduction of particle diameter
238	resulted in an order of specific surface area as ST(1:20)> ST(1:10)> ST(1:1) and PST-
239	5> PST-10> PST-20. However, the surface area value of PSTs is relatively low because
240	the addition of H ₂ O ₂ could effectively break the morphology of samples and reduce the
241	number of pores. In addition, the XRF result (Table 1) demonstrated that the distribution
242	of the element of STs or PSTs was hardly affected by the synthesis procedure.

Table 1. Physical-chemical properties of STs and PSTs by BET, XRF and Laser particle

size analyzer.

Samples	ST(1:1)	ST(1:10)	ST(1:20)	PST-5	PST-10	PST-20
<i>D50</i> /µm	89.13	72.78	53.56	62.81	76.59	92.40
<i>D90</i> /µm	200.21	166.92	131.99	149.99	174.27	211.18
$ ho/{ m g}\cdot m cm^{-3}$	2.48	2.77	2.60	3.05	2.98	3.06
$S_{BET}/m^2 \cdot g^{-1}$	55.2	73.56	139.2	6.9	1.36	0.85
$V/\mathrm{cm}^3\cdot\mathrm{g}^{-1}$	0.30	0.22	0.52	0.06	0.03	0.03
<i>R</i> _{pore} /nm	10.7	5.9	7.5	16.7	40.7	72.5
Ti/wt%	64.6	63.9	63.2	66.3	64.5	63.6
O/wt%	26.3	28.7	29.7	27.7	28.8	30.2
Na/wt%	7.1	7.4	7.1	6.0	6.7	6.2

245	It is evident from FTIR spectra of STs (Fig. S3a) and PSTs (Fig. S3b) that the
246	typical peak responding to Ti-O-Na ⁺ (Ti-OH destroyed in the alkaline environment [37])
247	in 1340 cm ⁻¹ remains unchanged in these samples. Meanwhile, the peak responding to
248	the Ti-O bond at about 900 cm ⁻¹ appears in PSTs and becomes stronger in the PST-10
249	and PST-20, which accounts for the coordination between the Ti-O framework and Na^+
250	instead of bridge connection[38]. During the PSTs synthesis, Na ⁺ might enter into the
251	interior structure and made the sample's surface more negative in solution [36], which
252	is meaningful for the adsorption of positive pollutants like cationic dyes. In addition, it
253	is observed that Ti-O-O bonds (responding to 680 cm ⁻¹ peak [39]) appears in PSTs after
254	the modification by H ₂ O ₂ , which might increase the acidity of the surface and promote
255	attracting cations. In addition, the zeta potential results (Fig. S4a-b) show that the pH_{iep}
256	(pH at isoelectric point) of ST(1:1), ST(1:10), ST(1:20), PST-20, PST-10 and PST-5 are
257	1.58, 2.01, 2.23, 1.15, 1.52 and 2.33, all of which are in low pH range. It is acceptable
258	that the change of synthesis conditions only brought about a slight difference in the
259	surface charge distribution. Therefore, the surface of STs and PSTs samples in solution
260	would be negatively charged in a wide range of pH, which is beneficial for the cationic
261	contaminants adsorption behavior [40].

262

3.2. Sedimentation property/adsorbent separation

The sedimentation property of adsorbent powders can reflect their separation tendency or affinity from the water after the adsorption process [41]. It is well known that some theoretical formulas are describing the sedimentation velocity of a spherical particle in a static solution, like the Stokes equation for the laminar region and Newton

equation for the turbulent region [42]. A meaningful conclusion can be made from these 267 equations that the diameter and density of sphere particles could affect the 268 sedimentation process. However, these ideal or empirical equations sometimes can't 269 perfectly predict the sedimentation of powders because the actual environment is 270 complicated. Therefore, the sedimentation experiment and CFD simulation are 271 necessary for analyzing the role of particle size in sedimentation. The density and 272 secondary size of as-prepared ST and PST samples have been changed through 273 controlling the synthesis conditions and are showed in Table 1. It is evident that the 274 275 density of PST samples is different from STs though the density value among PSTs or STs is similar. The setting experiment evaluated all the samples' setting properties, and 276 the resulting photos are shown in Fig. S5. It can be seen that the settling velocity of 277 278 PSTs or STs particles changes significantly with the change of synthesis conditions. The setting ability of these samples are in the order of ST(1:1) > ST(1:10) > ST(1:20)279 and PST-20> PST-10> PST-5, which is consistent with the order of secondary size. 280





(a) ST(1:1)



(d) PST-5



Figure 2. The simulation results of the setting process of STs and PSTs samples byAnsys-Fluent.

The simulation by CFD was also conducted to figure out the effect of secondary 295 size and density on the setting ability of adsorbent particles. The numerical method is 296 showed in section 2.4, and the parameters of particles were set according to the results 297 in Table 1. For convenience, diameter D_{50} and average true density ρ were used to 298 represent the chaotic state of actual particles in solution. On this basis, the simulation 299 of the settlement process of these samples is showed in Fig. 2. The simulated settling 300 conditions of STs or PSTs are consistent with the actual state, as shown in Fig. S5, 301 which demonstrates the rationality of the numerical and simplification method. In 302

addition, we found that smaller particles tend to interact with others in a longer time 303 than the larger ones in the initial phase of the settling process. The phenomenon can be 304 attributed to the fact that the diffusion movement of particles with smaller sizes is 305 stronger than particles with larger sizes due to the Brownian movement [43]. From 306 Table 1 and Fig. 2, sample particles with larger density (PSTs) settle more easily and 307 quickly than the small ones (STs). However, for the same kind of samples whose 308 synthesis procedures are identical, the true density value of particles are similar. Thus, 309 the main factor that affects the settling process is only secondary particle size. Therefore, 310 311 controlling the diameter of particles through finitely changing synthesis conditions can effectively enhance or reduce the settlement ability of samples, which is meaningful for 312 regulating the separation ability of saturated adsorbent materials. 313

314 **3.3.** Adsorption results

315 **3.3.1.** Effect of pH

Usually, the initial solution pH in adsorption experiments is one of the remarkable 316 317 influence factors since the surface property of adsorbents and the form of adsorbate are pH-dependent [44, 45]. It is illustrated that the adsorption capacity of NH₄⁺ onto 318 STs or PSTs reached the maximum when the initial solution pH is 3~4 (Fig. S6a-b). 319 The adsorption capacity of MB onto these adsorbents increased along with the increase 320 321 of initial solution pH (Fig. S6c-d). The above results can be explained with the following two reasons: firstly, MB is cationic in all the tested pH ranges, and ammonia 322 nitrogen is mainly cationic in the solution pH < 7 (NH₄⁺); secondly, the STs and PSTs 323 samples could be negatively charged at a large pH range according to the zeta potential 324

results [46]. In addition, considering the conclusion from the above characterization, 325 the only particle size of samples is highly affected by synthesis conditions, but the 326 particles' size has little effect on the adsorption capacity of NH4⁺ and MB by STs or 327 PSTs. The adsorbent samples with smaller particle sizes possess a slight better 328 adsorption performance at the same pH value. However, it is also indicated that the 329 optimal pH value for the adsorption behavior of STs or PSTs is kept constant among 330 different samples, which means that the particle size of the adsorbent sample is 331 irrelative to the pH adaption. 332

333

3.3.2. Adsorption isotherms

The adsorption isotherm experiments of NH4⁺ and MB onto samples were 334 conducted to find the effect of particle size on adsorption behavior with a series of initial 335 336 solution concentrations. It is observable that the difference in particle size has a more powerful impact on MB adsorption than NH4⁺ for the same adsorbent material (please 337 see Fig. S7). In addition, the difference in adsorption capacity for the same contaminant 338 among the three PST samples is higher than that among the STs. The above results 339 demonstrate that the effect degree of particle size on adsorption performance is relative 340 to the property of both adsorbent and adsorbate. The diagram also indicates that smaller 341 particle size is favorable for adsorption when other conditions remain constant. For 342 instance, the adsorption of MB on STs is in order of ST(1:20)> ST(1:10)> ST(1:1). The 343 increasing specific surface area and more intensive diffusion movement for smaller size 344 345 samples might provide more active adsorption sites and contacts.

346	Two typical isotherm models (Langmuir and Freundlich model) were employed to
347	fit the experimental data to understand the effect of particle size on the adsorption
348	process and evaluate the adsorption performance. The nonlinear fitting curves were
349	plotted in Fig. S7a-d, and a list of corresponding parameters was showed in Table 2.
350	The correlation coefficients (R^2) value reveals that the Langmuir model is more
351	reasonable than the Freundlich model in describing the process, which also indicates
352	that the change of particle size doesn't bring effects on the monolayer adsorption nature
353	of MB or NH_4^+ onto STs and PSTs [47, 48].

		Langmuir model			Freundli	Freundlich model		
Adsorbates	Samples	$Q_{max}/mg \cdot g^{-1}$	$K_L/L \cdot mg^-$	R^2	$K_{F}/(\mathrm{mg}^{(\mathrm{n-n})})^{1/\mathrm{n}}\cdot\mathrm{g}^{-1}$	1/n	R^2	
	ST(1:1)	52.81	0.012	0.9978	3.30	0.46	0.9526	
	ST(1:10)	51.76	0.017	0.9875	4.48	0.42	0.9963	
MD	ST(1:20)	58.94	0.013	0.9780	3.76	0.47	0.9750	
MB	PST-5	98.52	0.010	0.9904	3.59	0.56	0.9474	
	PST-10	94.47	0.007	0.9943	2.61	0.59	0.9743	
	PST-20	98.23	0.007	0.9942	2.13	0.62	0.9602	
	ST(1:1)	44.82	0.029	0.9921	3.61	0.50	0.9806	
	ST(1:10)	44.48	0.031	0.9888	3.90	0.48	0.9725	
NILL.+	ST(1:20)	45.97	0.032	0.9926	3.93	0.49	0.9776	
1114	PST-5	45.26	0.034	0.9886	4.22	0.48	0.9713	
	PST-10	46.18	0.027	0.9916	3.40	0.51	0.9773	
	PST-20	42.91	0.028	0.9942	3.26	0.51	0.9747	

Table 2. Isotherm parameters of NH₄⁺ or MB adsorption onto STs and PSTs.

355 **3.3.3.** Adsorption kinetics

The kinetic experiment was conducted to find the relationship between particle size and adsorption kinetics, and the results of the experiments are shown in Fig. 3. It is illustrated that samples with smaller particle sizes can reach the adsorption equilibrium of either NH₄⁺ or MB more quickly than those with a bigger size. For

example, the adsorption of MB onto ST(1:20) (D_{50} = 53.56 µm) can be finished within 360 10 min, which is shorter than the 60 min of ST(1:10) (D_{50} = 72.78 µm) and 90 min of 361 ST(1:1) (D_{50} = 89.13 µm) at the same conditions. Thus, the effect of particle size on the 362 equilibrium time was independent of the adsorbate species. In addition, two typical 363 kinetic models (pseudo-first-order and pseudo-second-order models) were used to fit 364 the experimental data to find the effect of particle size on the kinetic nature of the 365 adsorption process. The fitting curves are shown in Fig. 3, and the related parameters 366 are listed in Table S1. The value of kinetic constants (K_1 and K_2) of the two models 367 indicate the order of adsorption rate values as ST(1:20)> ST(1:10)> ST(1:1) and PST-368 5 PST-10> PST-20. The value of correlation coefficient R^2 indicates that the pseudo-369 second-order model is more suitable for describing the kinetic adsorption process of 370 371 both NH4⁺ and MB onto the as-prepared samples, which further demonstrates that the difference in particle size doesn't have a pronounced effect on the chemisorption nature 372 of the kinetic process. 373





Figure 3. Adsorption kinetics of NH_{4^+} onto STs (a), PSTs (b) and MB onto STs (c), PST(d). (Experimental conditions: $C_0=45 \text{ mg}\cdot\text{L}^{-1}$ for NH_{4^+} and 275 mg $\cdot\text{L}^{-1}$ for MB; initial solution pH=7)

375

It is also illustrated from the experimental data that samples with smaller sizes 379 could adsorb pollutants more quickly than those with bigger sizes at the beginning short 380 time, which reveals that the size value of particles can significantly affect the adsorption 381 rate at the primary stage. Therefore, the Elovich model was also employed to fit the 382 experimental data, as shown in Fig. 4 and Table S2. It is evident that the value of α 383 (initial adsorption rate) of STs and PSTs in adsorption of MB and NH4⁺ are all in the 384 order of PST-5> PST-10> PST-20 and ST(1:20)> ST(1:10)> ST(1:1), which is opposite 385 to the order of secondary particle size. The above results demonstrate that the effect of 386 particle size on the difference of adsorption rate mainly occurs at the very early time 387 388 when samples just in contact with the solution.





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Figure 4. The fitting results of adsorption of NH4⁺ onto STs (a), PSTs (b) and adsorption
of MB onto STs (c), PSTs (d) by Elovich model.

The entire aqueous adsorption process usually contains three stages: liquid film 393 diffusion, intraparticle diffusion and internal adsorption reaction [32]. In the early time 394 of the adsorption process, adsorbate molecules can move to the surface of sample 395 particles and enter into the internal framework through liquid film diffusion. The 396 decrease of particle size would make adsorbent powders diffuse more easily and quickly 397 in the solution and reduce the liquid film thickness, which successfully accelerated the 398 initial adsorption stage [49]. To make sure that the liquid film diffusion process is one 399 of the rate-limiting steps in the adsorption of NH4⁺ or MB onto STs and PSTs, the 400 Weber-Morris model was used to fit the experimental data as shown in Fig. 5 and Table 401 S3. The fitting curves don't cross the origin, and the value of C are all bigger than zero, 402 which indicates that intraparticle diffusion is not the only rate-limiting step. The whole 403 adsorption rate is controlled by multiple diffusion modes [50]. Therefore, the effect of 404 particle size on the liquid film diffusion process can affect the adsorption rate to some 405 extent, especially at the early contact time. 406



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Figure 5. Adsorption kinetics of NH4⁺ (a, b) and MB (c, d) onto samples with the fitting of the Weber-Morris model.

Therefore, the regulation of secondary size value can be carried out to improve the 411 separation ability and adsorption kinetics for the powder-shape adsorbent. The 412 enhancement of sedimentation ability can increase the separation efficiency and reduce 413 the relative cost for recovering the exhausted adsorbents. However, the larger secondary 414 size would mean more prolonged adsorption kinetics and finally reduce the adsorption 415 efficiency. A balance or a key point should be found between these two opposite results. 416 Therefore, an entire cost calculation is suggested to help determine the secondary size 417 value when designing adsorbent materials in the actual application. The investigation 418 of this issue will be focused on in future studies. 419

420 4. Conclusions

In this study, a series of sodium titanates (STs) and peroxide modified sodium 421

titanates (PSTs) particles with different sizes were synthesized through the control of 422 hydrolysis and complexation by changing the ratio of reagents. The as-prepared 423 samples were applied in the adsorption of NH4⁺ and MB from an aqueous solution to 424 figure out the effect of particle size on the adsorbent properties and adsorption 425 performance. The sedimentation experiment and CFD simulation indicated that the 426 change of secondary size (particles size after agglomeration, aggregation or hydration 427 in solution) could effectively affect the settling ability of adsorbent powders, which is 428 meaningful for regulating the separation ability of adsorbent materials. In addition, the 429 effect of secondary size on adsorption kinetics is also non-negligible. The PSTs or STs 430 particles with smaller secondary sizes can reach adsorption equilibrium for NH4⁺ and 431 MB more quickly than those with a bigger size. The fitting results from Elovich and 432 433 Weber-Morris models demonstrate that the particles size affect kinetics mainly in the liquid film diffusion process at the early contact time. Therefore, the secondary size 434 significantly affects the sedimentation ability and adsorption kinetics of titanate-based 435 adsorbents in the opposite results, which is helpful for adsorbent design and mechanism 436 analysis. 437

438

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442 Appendix A. Supplementary data

443 Eqs. (S1)- (S2): Langmuir and Freundlich models

- 444 Eqs. (S3)- (S4): Pseudo-first order and Pseudo-second models.
- Eqs. (S5)- (S6): Elovich model and Web-morris model
- 446

- 447 Table S1. Parameters of the kinetics models for the adsorption of NH_{4^+} and MB onto STs and PSTs.
- 448 Table S2. Parameters of the Elovich model for the adsorption of NH_{4^+} and MB onto STs and PSTs.
- 449 Table S3. Parameters of the Weber-Morris model for the adsorption of NH_{4^+} and MB onto STs and
- 450 PSTs.
- 451
- 452 Figure S1. The molecular structures of methylene blue.
- $\label{eq:453} Figure \ S2. \ N_2 \ gas \ adsorption-desorption \ isotherm \ of \ STs \ and \ PSTs. \ The \ inset \ is \ the \ corresponding$
- 454 pore size distribution.
- 455 Figure S3. FTIR spectra of (a) STs and (b) PSTs.
- 456 Figure S4. Zeta potential of (a) STs and (b) PSTs at different solution pH.
- 457 Figure S5. The setting state of PSTs at (a) 0s, (b) 60s, (c) 10 min, (d) 30 min and STs at (e) 0s, (f)
- 458 60s, (g) 10 min, (h) 30 min.
- Figure S6. Effect of initial solution pH on the adsorption capacity of NH_{4^+} onto STs (a), PSTs (b) and adsorption capacity of MB onto STs (c), PSTs (d).
- $\label{eq:461} {\mbox{Figure S7. Adsorption isotherm results of NH_4^+ onto STs (a), $PSTs$ (b) and MB onto STs (c), $PSTs$}$
- 462
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464 **References**

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