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1	Minimizing residual black particles in sand filtrate when applying super-
2	fine powdered activated carbon: coagulants and coagulation conditions
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18 Abstract

19 Because of the eminent adsorptive capacity and rate for dissolved organic molecules 20 compared to conventionally-sized powdered activated carbon (PAC), super-fine powdered 21 activated carbon (SPAC) is gathering momentum for use in not only the pretreatment for 22 membrane filtration for drinking water purification but also the conventional water 23 purification process consisting of coagulation-flocculation, sedimentation, and rapid sand-24 filtration (CSF). However, the probability of SPAC particles to leak through a sand bed is higher than that of PAC, and their strict leakage control is an issue to be challenged when 25 26 applying SPAC to CSF. However, study focusing on very high particle removal, which yield 27 residual concentrations down to around 100 particles/mL, has been very limited. A previous 28 study mentioned that the tendency of SPAC leakage is related to its low destabilization. In 29 response to this, the present study focused on the two key components of coagulation (mixing 30 intensity and coagulants) and investigated how to effectively reduce the residual SPAC after 31 CSF.

Astonishingly, the flash mixing (the first process of CSF), especially its G (velocity gradient) value, played the most important role in determining the residual SPAC in the filtrate of sand filter (the fourth process). Even if the slow mixing time was short, a sufficiently large G value but short T (mixing time) value in flash mixing effectively reduced the residual SPAC. When the total GT value of flash and slow mixing was fixed at a constant, priority should be given to flash mixing to reduce the residual SPAC.

Among 23 PACl (poly-aluminum chloride) coagulants, PACl with a high-basicity (basicity 70%) and with sulfate ion (0.14 of sulfate/aluminum in molar ratio), produced by Al(OH)₃dissolution, were the most effective to reduce the residual SPAC after CSF. PACls produced by base-titration, which have been intensively investigated in previous researches, were not effective due to lack of floc-formation ability. However, their Al species composition

43	determined by the ferron method were almost the same as those of PACl by Al(OH)3-
44	dissolution, and their charge-neutralization capacities were higher. PACls produced by
45	Al(OH)3-dissolution possessed both charge-neutralization and floc-formation abilities, but
46	the former ability was more important to minimize the residual of SPAC.
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49	Keywords
- 0	

50 SPAC; filtration; mixing; floc; basicity

52 **1. Introduction**

53

54 When removing dissolved organic compounds, in particular those of low molecular size, in 55 water treatment processes, adsorption by activated carbon (AC) has been a common practice 56 for many years. Activated carbon is used as an adsorbent either in powdered or granular form. 57 While granular activated carbon is used in a bed adsorber, powdered activated carbon (PAC) 58 can be applied at any point in the treatment process if PAC is sufficiently removed before the 59 treated water enters the distribution system. The main concern for the adsorption processes 60 is the removal efficiencies of target adsorbate compounds to be treated, but plant operators 61 also pay attention to avoid fine black carbon particles from remaining in treated water and 62 entering the distribution system. It is actually not uncommon that PAC, which is applied 63 upstream of a separation process such as rapid sand filtration, sometimes passes through the 64 filters and enters the distribution system, provoking black or grey water complaints from 65 consumers (American Society of Civil Engineers and American Water Works Association 66 1998). Black water is usually caused by inadequate coagulation and sedimentation or high 67 doses of PAC (American Water Works Association and American Society of Civil Engineers 68 2012). Even trace concentrations of black carbon particles remaining in treated water, which 69 do not make the water black or grey and do not violate drinking water quality standards for 70 turbidity, could deteriorate the quality of food, such as tofu, that is produced from the water, 71 etc., which causes complains from food manufacturers. Plant operators pay closer attention 72 to the turbidity or particle count in treated water when PAC is dosed than when PAC is not 73 dosed (Bureau of Waterworks Tokyo Metropolitan Government 2014). Control of fine, black 74 carbon remaining in treated water should be a critical issue, in particular, under the situation 75 in which the application of super-fine PAC (SPAC), carbon particles with a size smaller than

76 the conventional size, attracts attention (Amaral et al. 2016, Ando et al. 2010, Bonvin et al.

77 2016, Ellerie et al. 2013, Matsui et al. 2004, Matsui et al. 2012, Partlan et al. 2016).

78

79 Plenty of studies have been conducted for the effect of coagulants and coagulation conditions 80 on floc formation and turbidly removal (Letterman and Yiacoumi 2011). Some studies have 81 been conducted for the effect of PAC on coagulation-sedimentation; however, studies 82 regarding the control of carbon particles at trace concentration levels are very limited. Aguilar 83 et al. (2003) reported that the use of PAC considerably increased the elimination efficiency 84 of particulate matters by coagulation and sedimentation, but there was a diminution in the 85 elimination efficiency for particles with diameters in the range 5.5-8.5 µm. They attributed 86 this diminution to the carbon particles remaining in the treated water. In contrast, Younker 87 and Walsh (2016) reported that the addition of PAC into FeCl₃ flocs reduced floc size but 88 had little impact on final turbidity after sedimentation. Therefore, the researches of the two 89 reports are not completely consistent with regard to the effect of PAC, but several reasons 90 can be considered for this apparent discrepancy: raw water characteristics, coagulation 91 conditions employed, etc. The effects of SPAC and PAC on floc formation were investigated 92 in a membrane filtration study (Matsui et al. 2009). The authors reported that the floc particles 93 that formed during coagulation preceded by SPAC pretreatment were larger and more porous 94 than the floc particles formed during coagulation preceded by PAC pretreatment and those 95 formed during coagulation without any pretreatment. The authors explained this result due 96 to increased particle-particle collision frequency and better removal of natural organic 97 matter, which inhibits coagulation by consuming coagulant, before the coagulation reaction. 98 These previous researches focused on the effect of adsorbent on the formation of floc 99 properties, settling velocity, permeability etc., and carbon particles have been investigated as

a ballasting agent for settling, nuclei for coagulation, or a pretreatment agent of coagulation-hindering compounds.

102

103 However, researches to investigate the effect of coagulation conditions and coagulants on the 104 remaining carbon particles in finished water have, to the best of our knowledge, never been 105 conducted except for that of our research group, although their control at a trace concentration 106 level is an important issue in terms of customer satisfaction. Our research group recently 107 revealed that at equivalent carbon doses to achieve the same adsorptive removal performance 108 (30 mg/L for PAC and 7.5 mg/L for SPAC), the residual particle number concentrations after 109 coagulation-flocculation, sedimentation, and sand filtration (CSF) were similarly low (100-110 200 particles/mL) between SPAC and PAC and the particle sizes were similarly small 111 (Nakazawa et al. 2018). There is no drinking water quality standard or guideline in the world 112 for black carbon particles, to the best of our knowledge. When applying SPAC instead of 113 PAC, however, a criterion from plant operator sites, in particular those with a safety 114 orientation, would be that the level of carbon particles remaining in treated water when SPAC 115 is applied should be at a similar or lower level than when PAC is applied.

116

On the basis of these consideration, this study set its objectives to find better coagulation conditions and find a better coagulant (poly-aluminum chloride: PACl) for the removal of SPAC. In actual water treatments, SPAC/PAC is applied for the removal of dissolved organic compounds, and the removal of SPAC/PAC by CSF occurs in the presence of other substances, such as clay particles and natural organic matter, which may influence the removal of SPAC/PAC. Moreover, coagulation conditions better for SPAC removal may necessarily not bring a good performance for natural organic matter removal. However, the

124	CSF experiments in this study were conducted mainly with water made from filtered tap
125	water supplemented with carbon slurry in order to gain basic information for carbon particle
126	removal without any influence of other substances.
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129	2. Materials and Methods
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131	2.1. Activated carbon particles
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A commercially available wood-based PAC (Taiko W; Futamura Chemical Co., Ltd., Nagoya, Japan) was prepared as a slurry in pure water (Milli-Q water; Merck KGaA, Darmstadt, Germany) at a concentration of ~15% (w/w). The PAC in the slurry were milled in a closed-chamber ball mill (Nikkato, Osaka, Japan) with 5- and 10-mm-diameter Al₂O₃ balls at 45-50 rpm for 4–5 h to afford the particles with D₉₅ (the diameter larger than 95% of the entire distribution) < 30 μ m (Pan et al. 2017). The slurry was taken out from the chamber of the ball mill and then milled with a bead mill (LMZ015, Ashizawa Finetech, Ltd., Chiba, Japan) with 0.3-mm-diameter ZrO₂ beads at a rotational speed of 8 m/s (2590 rpm) in recirculation mode for 20 min to produce a SPAC slurry (Table 1).

The true particle size distributions of the carbons were determined using a laser light
diffraction and scattering method (Microtrac MT3300EXII, Nikkiso Co., Inc., Tokyo, Japan),
which followed addition of a dispersant (Triton X-100; Kanto Chemical Co., Inc., Tokyo,
Japan; final concentration, 0.08% w/v) and ultrasonic dispersion with 150 W for 1 min (Fig.
138 1S, Supplementary Information of SI).

141

142 Twenty-three kinds of PACls were used as coagulants in the present study. Among them, 143 PACI-70 (basicity 2.1; the number, 70, in the name indicates the % basicity; sulfate ion 2% 144 w/w) and PACI-50 (basicity 1.5, sulfate ion 3% w/w) are commercial products provided by 145 Taki Chemical Co., Ltd. (Hyogo, Japan), which are produced by dissolving Al(OH)₃ into 146 HCl and H₂SO₄ [as described by, e.g., Itoh and Sato (1995) and Sato and Matsuda (2009)]. 147 On the other hand, B70ns, B70s0.11, B70s0.12, B70s0.13, B70s0.14, B70s0.15, and 148 B70s0.17 (the numbers after "B" and "s" represent the "% basicity" and "mole ratio of sulfate 149 ion to aluminum: SO₄/Al" respectively, and "ns" stands for "non-sulfated") were produced 150 in the authors' laboratory by the following method named Al(OH)₃-dissolution.

151

152 Predetermined amounts of AlCl₃ · 6H₂O (FUJIFILM Wako Pure Chemical Corporation, 153 Osaka, Japan) and Al₂(SO₄)₃·14~18H₂O (FUJIFILM Wako Pure Chemical Corporation) 154 were dissolved into pure water (Milli-Q water) to get aluminum aqueous solution. This 155 solution was mixed with Al(OH)₃ powder (FUJIFILM Wako Pure Chemical Corporation) to 156 form a milk-like slurry and then heated with microwaves (ETHOS TC, Milestone S.r.l., 157 Sorisole (BG), Italy) at 150°C for 3 h. After that, the slurry was centrifuged by a centrifugal 158 separator (H-9R, Kokusan Co., Ltd., Tokyo, Japan) at a rotation speed of 3,000 rpm (6,000 159 g) for 10 min at 20°C, and then the clear top of the liquid was filtered by a membrane filter 160 (pore size 0.45 µm, PTFE, Toyo Roshi Kaisha, Ltd., Tokyo, Japan) to obtain a solution of 161 PACl with basicity of 40~50%. The solution was poured into a beaker, stirred with a rotation 162 speed of >600 rpm, and heated at 50°C using a hot plate/magnetic stirrer, and then a 163 predetermined amount of aqueous solution of sodium carbonate (FUJIFILM Wako Pure

164 Chemical Corporation) was dripped into the solution to adjust the basicity up to 70%. The165 basicity was calculated by the following formula.

166

167 Basicity (%) =
$$\frac{[OH^{-}]}{3[Al_t]} \times 100$$

168

where [Alt] is the total aluminum concentration determined by an inductively coupled plasma
mass spectrometer (ICPMS, 7700x, Agilent Technologies, Inc., Santa Clara, CA, USA),
[OH⁻] is the concentration of hydroxyl groups from the ingredients Al(OH)₃ and sodium
carbonate. The total mass of OH⁻ introduced to the final product was calculated by taking a
mass balance.

174

PACls produced by the method named base-titration were used in the CSF experimentsdescribed in supplementary experiments. The method is described in SI.

177

178 The distributions of aluminum species in the coagulants were analyzed by the ferron method. 179 On the basis of their reaction rates with ferron reagent (8-hydroxy-7-iodo-5-quinoline 180 sulfonic acid, FUJIFILM Wako Pure Chemical Corporation), the aluminum species were 181 divided into three categories: Ala, Alb, and Alc. Ala denotes aluminum species that reacted 182 with ferron instantaneously (within 30 s). Alb denotes species that reacted with ferron within 183 120 min. Alc denotes species that did not react with ferron. These species were assumed to 184 be monomeric, polymeric, and colloidal aluminum species, respectively (Wang et al. 2004). 185 Ferron analyses of the PACls were conducted immediately after dilution with Milli-Q water 186 to 2.7 g-Al/L (0.1 mol-Al/L) (Jia et al. 2004, Wang et al. 2004). Dilution reportedly has little 187 effect on the ferron speciation distribution of PACI (Kimura et al. 2013, Wang et al. 2002).

188

189 2.3. Coagulation–flocculation, sedimentation, and rapid sand filtration

190

191 Tap water in Sapporo city was filtered through a membrane filter (nominal pore diameter, 192 0.1 µm; Toyo Roshi Kaisha, Ltd.) to remove suspended matter, and then the water was 193 adjusted for M alkalinity to 30 mg/L as CaCO₃ by adding NaHCO₃ (FUJIFILM Wako Pure 194 Chemical Corporation). The water was supplemented with an activated carbon slurry to make 195 a raw water for CSF experiments: activated carbon concentrations were 30 mg/L for PAC; 6 196 mg/L, 7.5 mg/L, 10 mg/L, or 30 mg/L for SPAC. Most CSF experiments were conducted with these waters, but some CSF experiments were additionally conducted with water from 197 198 the Toyohira River (Sapporo, Japan) after supplementing it with SPAC at 10 mg/L. The river 199 water was sampled at the location where it becomes the raw water source for the Moiwa 200 Water Purification Plant (Sapporo).

201

202 The experimental setup and procedure were basically the same as those of Nakazawa et al. 203 (2018) but are shown in Fig. 2S (SI). The coagulation-flocculation and sedimentation steps 204 were conducted in a 4-L rectangular beaker. After a predetermined volume of HCl or NaOH 205 (0.1 N) was added to adjust the coagulation pH to 7.0, the coagulant (PACl) was injected into 206 the beaker to a final concentration of 4 mg-Al/L, unless otherwise noted. Then the water was 207 mixed rapidly followed by slow mixing with a three-stage mixing intensity. The mixing 208 intensities (G value: velocity gradient, see Fig. 3S and Table 1S of SI for the G value 209 calculation) and times (T value: mixing time) of the flash (indicated by subscript F) and slow 210 mixing (indicated by subscript S, with the order: 1, 2, and 3) were varied depending on each 211 experiment, and these values are described in each case in the Results and Discussion section 212 (see also Table 2S, SI). The water was then left at rest for 1 h. Next, the top three liters of the 213 water (supernatant) were transferred to another beaker for determination of the turbidity 214 (2100Q portable turbidimeter; Hach Company, Loveland, CO, USA) and for rapid sand filtration. Sand filtration was conducted for 40 min at a rate of 90 m d⁻¹ in the down-flow 215 216 direction using a column (Φ 4 cm) filled to a depth of 50 cm with sand (effective diameter, 217 0.6 mm; uniformity, 1.3). The sand filtrate was collected from 13 to 40 min after the start of 218 filtration, and the turbidity and carbon particle count of the filtrate were determined. After 219 each filtration run, the sand filter was backwashed with tap water, then washed forward with 220 pure water (Milli-Q water) and membrane-filtered tap water.

221

222 2.4. Membrane filtration and microscopic image analysis

223

224 To sample the carbon particles in the water, the water was filtered through a PTFE membrane 225 filter (nominal pore diameter, 0.1 µm; Φ25 mm; Merck KGaA) supported by a glass filter 226 holder (KG-25; Toyo Roshi Kaisha, Ltd.) (Fig. 4S, SI). After drying the filter, color digital 227 photomicrographs were captured for nine or eighteen predetermined observation zones 228 (microscope view area, 247×330 µm) per filter (Fig. 5S, SI) with a digital microscope (VHX-229 2000; Keyence corporation, Osaka, Japan) at 1000× magnification. The photomicrographs 230 were analyzed using the image analysis software supplied with the microscope. Details of 231 the analytical procedures are reported elsewhere (Nakazawa et al. 2018).

- 233
- 234 **3. Results and Discussion**
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236 3.1. Comparison of commercially-available PACI-50 and PACI-70

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Fig. 1 shows the number concentrations of residual carbon particles in sand filtrate after CSF when water containing SPAC or PAC of various mass concentrations was coagulated using PAC1-70 (a high-basicity PAC1) or PAC1-50 (a conventional PAC1 having normal basicity). When PAC at the initial concentration 30 mg/L was treated with PAC1-70, the residual concentrations were around 200 particles/mL, and PAC1-70 resulted in a slightly higher residual concentration than PAC1-50, but the difference was small.

245 When SPAC of the same initial concentration (30 mg/L) was treated using conventional 246 PACI-50, the residual concentration was much higher (800 particles/mL). When PACI-70 247 was used to treat the SPAC, the residual concentration was reduced to 400 particles/mL. The 248 lower the initial concentration of SPAC was, the lower the residual concentration, both for 249 PACI-50 and PACI-70; and at the initial concentration of 7.5 mg/L or less, the difference of 250 residual concentrations between the two cases of using PACI-50 or PACI-70 became smaller. 251 Overall, high-basicity PACI-70 was superior to PACI-50 to remove SPAC. With PACI-70, 252 the removal rates in terms of particle number concentrations depended on the concentration 253 of SPAC: 5.3 log removal was lowest at an SPAC concentration of 10 mg/L. Nevertheless, 254 the removal rates with PACI-70 were slightly higher than with PACI-50 (Fig. 6S, SI). In 255 contrast, the turbidity of the supernatant after sedimentation following coagulation by PACI-256 50 was slightly lower than that by PACI-70, and the result was opposite to the residual 257 number concentration mentioned above (Fig. 7S, SI).

258

Higher-basicity PACls (e.g. PACl-70) are known to neutralize negative charges more
according to the results of colloid titration (Fig. 8S, SI) (Matsui et al. 2017). Accordingly,

although PACI-70 is not superior compared with PACI-50 in terms of forming large flocs
bringing a lower turbidity of supernatant water, PACI-70 is inferred to neutralize the negative
charges of carbon particles efficiently due to its high capacity in charge neutralization. The
high capacity in charge neutralization seemed to contribute to the removal of carbon particles.

266 We previously reported that the particles remaining after CSF were smaller in size than the 267 particles before CSF (Nakazawa et al. 2018). Although the tendency of small carbon particles 268 to remain after CSF is overall attributable to their size, the size effect is comprised of the 269 following three reasons: 1) low destabilization rate during the coagulation process, 2) low 270 frequency of particle-particle collisions during flocculation, and 3) low probability of the 271 particles coming into contact with sand grains during the sand filtration process. Therefore, 272 particle destabilization is a key to reduce remaining carbon particles after CSF. When 273 comparing the zeta potentials of residual carbon particles after CSF treatments with PACI-274 70 and PACI-50, they were not different (Fig. 2). Therefore, particles with a certain zeta 275 potential value or lower regardless of PACI-70 or 50 passed through the filter. Accordingly, 276 the merit of PACI-70 in reducing carbon particle concentration after CSF was due to its high 277 capacity to minimize the number of non-neutralized carbon particles (with a certain zeta 278 potential value or lower) rather than due to its high positive charge to neutralize the negative 279 charge of particles to nearly zero level.

280

When the SPAC suspension was coagulated using PACI-70, the formation of micro floc particles was not confirmed in the photograph taken soon after flash mixing (Fig. 3). When the PAC suspension was coagulated with PACI-70, the formation of micro floc particles was confirmed marginally. In contrast, when PACI-50 was used in the coagulation process, the formation of micro floc particles was clearly observed in the experiments of all experimental conditions. However, floc size at the end of slow mixing was not different between each pairof experiments using PACI-50 and PACI-70.

288

289 Therefore, it can be said that the rapid formation of large-size floc particles and the lowered 290 turbidity of supernatant water, although such information can be obtained by a conventional 291 jar test, are not indicators for a lower residual carbon particle concentration after CSF. PACls 292 are rich in diversity in terms of two different aspects: charge-neutralization capacity to 293 destabilize particles and the bridge-formation ability to form large floc particles, but the 294 charge-neutralization capacity is not related to the bridge-formation ability (Zhao et al. 2010). 295 Consequently our study revealed that PACIs having a feature of the charge-neutralization 296 capacity to destabilize particles rather than the bridge-formation ability to form large-size 297 floc particles are beneficial for lowering residual carbon particle concentration after CSF.

298

299 3.2. Effect of mixing intensity in coagulation and flocculation

300

301 As mentioned in the preceding paragraph, the high-basicity PACl (PACl-70) formed flocs 302 slowly, but it had a high charge neutralizing capacity. Hence, it was hypothesized that 303 optimizing the mixing intensity and time would lead to further reduction of residual carbon 304 particle concentration in sand filtrate, and experiments were conducted. Fig. 4 illustrates the 305 residual SPAC concentrations in sand filtrates when coagulation-flocculation were conducted 306 with three different G_TT_T values ranging from 19,500 to 78,000 by adjusting T_T. (The G_TT_T 307 value means total GT, which is the sum of flash-mixing GT (G_FT_F) and slow-mixing GT 308 (G_ST_S) .) In the case with the small G_TT_T value of 19,500, the residual carbon particle 309 concentration was 800-particles/mL after CSF using PACI-50, whereas a much higher 310 residual concentration of 2,200 particles/mL was observed using PACI-70. In such a low

311 mixing intensity, large floc particles were seen in the photographs at the end of slow mixing, 312 but many small floc particles were also observed in the same photographs, in both cases of 313 PACI-50 and PACI-70 (Fig. 5); the mixing intensity was obviously insufficient. In contrast, 314 as the G_TT_T value increased from 39,000 to 78,000, small floc particles diminished, seen in 315 the photographs taken at the end of slow mixing (Fig. 5). At the same time, the turbidity of 316 supernatant water (Fig. 9S, SI) and the residual carbon particle concentration of sand filtrate 317 also decreased substantially. As shown in Fig. 4 and Fig. 9S (SI), furthermore, the effect of 318 G_TT_T on the turbidity of supernatant water and the residual carbon particle concentration of 319 sand filtrate was more pronounced for PACI-70, which had a higher capacity for charge 320 neutralization, than PACI-50.

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322 *3.3. Effect of flash mixing intensity in coagulation*

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324 Because charge neutralization occurs mostly during the flash mixing stage, we hypothesized 325 that the residual carbon particle concentration of sand filtrate is influenced more by the flash 326 mixing conditions than by the slow mixing conditions, so we conducted other series of CSF 327 experiments. In the first series of experiments, the G_TT_T was fixed at 39,000, but its allocation 328 to flash mixing was changed; i.e. the larger the G_FT_F value of flash-mixing was, the smaller 329 the G_ST_S value of slow-mixing. The results are shown in Fig. 6. When the flash-mixing G_FT_F 330 was small (6,000), the residual carbon particle concentration was high at 900 particles/mL. 331 The residual concentration, however, decreased as the flash-mixing G_FT_F increased. The 332 effect of the flash-mixing G_FT_F was more prominent with PACI-70 than with PACI-50. At 333 the largest flash-mixing G_FT_F (24,000), the residual carbon particle concentration was 334 reduced to <200 particles/mL.

336 In a series of experiments for which flash-mixing G_FT_F was changed by adjusting T_F but slow 337 mixing intensity was fixed at a constant value (G_ST_S=15,100), a G_FT_F>24,000 resulted in a 338 low residual carbon particle concentration (Fig. 7), indicating that there is a certain minimal 339 value for G_FT_F required for lowering the carbon particle concentration in sand filtrate. Next, 340 the effect of G_F value on residual carbon particles was examined in the condition where GT 341 values of both flash and slow mixing intensities were fixed (G_FT_F=24,000 and G_ST_S=15,100). 342 Residual carbon particle concentration became smaller with a larger G_F value (smaller T_F value) (Fig. 8). It was minimal at $G_F=600 \text{ s}^{-1}$ (experiments at G value>600 s⁻¹ could not be 343 344 conducted because the mixing caused the water to splash out of the container. Kan et al. 345 (2002) conducted jar tests by using alum and commercially-available PACl to investigate the 346 effect of flash-mixing time on the removal of clay particle. They report that the residual 347 turbidity decreased sharply with increasing duration of flash-mixing. Lin et al. (2013) also 348 conducted jar tests by using turbid surface-water and commercially-available PACl to 349 investigate the effect of flash-mixing intensity on the turbidity and DOC removal. They 350 reported that the removal performances of turbidity and DOC were improved as flash-mixing 351 intensity increased. We used fine carbon particles as the target matter in the present study, 352 but the tendency that the turbidity after settling and residual carbon particles remaining in 353 sand filtrate reduced as flash-mixing intensity or time increased was much the same as those 354 previous studies.

355

It is of great interest that the trace residual-carbon-particle concentration as an outcome after the fourth step of sand-filtration in the CSF was largely affected by the mixing intensity of the first step of coagulation in the CSF, although the flocculation and sedimentation processes occurred in between the coagulation and the sand-filtration. The flocculation and sedimentation processes had long detention time, and most of the carbon particles were 361 removed by the sedimentation process. During the flash mixing process for coagulation, the 362 PACls hydrolyzed in water and formed a hydrolyzing aluminum polymer, which efficiently 363 neutralized the negative charge of carbon particles. Thus, providing both rapid dispersion of 364 dosed PACls and high-frequent contact between the hydrolyzing aluminum polymer and the 365 carbon particles, in particular very fine carbon particles, was necessary to decrease charge-366 un-neutralized particles. Finally, achieving these conditions produced low residual carbon 367 particles, which penetrated through the sand bed situated in the last stage in the CSF. A high 368 G_F value created such conditions of the rapid dispersion and the high-frequent contact.

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370 *3.4. Effect of coagulant dosage*

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372 It is generally known that turbidity removal by coagulation-sedimentation improves as the 373 dosage of coagulant increases when the coagulation processes are mainly due to sweep 374 flocculation mechanism, which operates at near neutral pH, but not due to charge-375 neutralization mechanism, which operates at pH < 6 (Amirtharaja and O'Melia 1990, 376 Hendricks 2010, Letterman and Yiacoumi 2011). The present study of CSF, which was 377 conducted at pH 7, also confirmed the relationship between the dosage of coagulant and the 378 residual concentration of SPAC or the turbidity of supernatant water in CSF (Fig. 10S and 379 Fig. 11S, SI); increasing coagulant dosage resulted in an increase in the removal of carbon 380 particles both after sedimentation and sand filtration. Additionally, the superiority of PACI-381 70 to PACI-50 was held in the entire range of coagulant dosages tested in terms of residual 382 carbon particle concentration. The very low residual concentration of 6 particles/mL was 383 attained with the highest coagulant dosage of PACI-70.

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385 3.5. Development and performance of coagulants by base-titration

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The above described investigation suggests the importance of decreasing the number of 388 charge-un-neutralized particles in order to enhance the separation of SPAC particles and 389 lower the number of residual SPAC particles after CSF. Available techniques for decreasing 390 the number of charge-un-neutralized particles were enumerated using a coagulant with high 391 a charge-neutralization capacity, such as high-basicity PACl with the optimized flash mixing 392 intensity, as well as increasing the dosage of coagulant. We further searched for a better PACl 393 that lowered the number of residual SPAC particles. We produced a series of PACIs with 394 high basicity (Table 3S, SI) and conducted jar tests (Table 4S, SI). 395 396 It is widely known that the Al₁₃ species in PACl have a high charge-neutralizing ability (Gao 397 et al. 2005, Lin et al. 2008, Parthasarathy and Buffle 1985, Wu et al. 2007, Zhao et al. 2009), 398 and Alb analyzed by the ferron method corresponds to the Al_{13} species (Parker and Bertsch 399 1992, Parthasarathy and Buffle 1985). In the present study, some PACls with a high content 400 of Alb (Alb-type PACl) were produced by base-titration (Fig. 12S, SI), and experiments of 401 coagulation-flocculation and sedimentation were conducted. The zeta potential of carbon 402 particles became zero at a lower dosage of Alb-type PACl (B80ns-1) than of PACl-70 (Fig. 403 13S, SI), and Alb-type PACls (B65ns, B80ns-1, B80ns-2) supplied a higher positive charge 404 to carbon particles than PACI-70 at the same coagulant dosage (Fig. 14S, SI). These results 405 indicate that Alb-type PACls actually had a higher charge-neutralizing ability than PACl-70. 406 However, these Alb-type PACls could not flocculate carbon particles, and consequently 407 resulted in a high turbidity after sedimentation, even given large mixing intensity (Fig. 14S, 408 SI). Even at the coagulant dose, which brought the zeta potentials of carbon particles to

409 around zero, the turbidities after sedimentation were still high, compared with those obtained

410 by PACI-70 (Fig. 13S, SI). 411

It was reported that sulfate ions in PACl play a role in enhancing floc formation (Wang et al. 2002). Actually, the commercial PACl products, PACl-50 and PACl-70, both of which produce larger floc particles than the Alb-type PACl, contain sulfate ion. Accordingly, we supplemented sulfate ion and prepared three Alb-type PACls (Fig. 15S, SI). One PACl (B82s0.11) among the sulfated Alb-type PACls slightly flocculated carbon particles at a large mixing intensity (as a GT value of 57,000). However, turbidity-removal performance was considerably inferior to PACl-50 and PACl-70.

419

420 Gao and Yue (2005) produced sulfated PACls of different SO₄/Al ratios by base-titration at 421 ambient temperature and reported that a sulfated high-basicity PACl (basicity 67%) with 422 SO₄/Al=0.066 in the range from 0 to 0.15 reduced residual turbidity the most when an 423 experiment consisting of coagulation-flocculation and sedimentation was conducted with 424 lake water. In our experiments, however, the production of sulfated high-basicity PACIs at 425 ambient temperature by base-titration was unsuccessful due to the occurrence of aluminum 426 hydroxide precipitation during titration, and we were not able to apply the sulfated high-427 basicity PACl. Shi et al. (2007) also mentioned that room temperature was not suitable for 428 preparation of PACl because a large amount of aluminum precipitate could form during 429 titration.

430

431 Yan et al. (2008) investigated the relationship between Al species and their behavior in 432 coagulation targeting NOM. They reported that Ala (monomeric species) combined with 433 NOM but most of the flocs formed by Ala were small and difficult to settle. Alb (polymeric 434 species) had high charge-neutralizing ability but flocs formed by Alb were not larger and did 435 not settle faster than those formed by Alc (colloidal species), and Alc adsorbs and removes 436 the target matter most efficiently. In the present study, in addition, given that the major 437 aluminum fraction in PACI-50 and PACI-70 was not Alb but Alc, Alc-type PACIs (rich in 438 content of Alc fraction) were also produced (see SI for the preparation method). Those Alc-439 type PACls showed high charge-neutralization ability, however their flocculation 440 performances were almost the same as the Alb-type PACls and had a low removal efficiency 441 (Fig. 16S, SI). In summary, Alb or Alc in PACIs prepared by base-titration had a high 442 capacity for charge-neutralization, but the bridge-formation ability to form floc particles was 443 poor and resulted in high turbidity after sedimentation.

444

445 3.6. Development and performance of coagulants by Al(OH)₃-dissolution

446

447 Subsequently, we applied another method to produce PACls: Al(OH)₃-dissolution (Table 2). 448 Eleven PACls with a basicity of 70% and with a variety of sulfate ion contents were produced 449 (Table 2): it was impossible to produce PACls with a basicity >70% because of the gelation 450 of Al solution or the presence of too much undissolved residue. As shown in Fig. 9, the 451 performance on reducing residual SPAC particles increased as sulfate content increased from 452 SO₄/Al=0 to SO₄/Al=0.14 but decreased for SO₄/Al>0.14. The preparation of PACls with 453 SO₄/Al>0.14 required rising the temperature from 50°C to 80°C when adjusting the basicity. 454 In terms of high temperature, this process was similar to the process of base-titration PACls, 455 which do not have sufficient bridge-formation ability. The loss of the bridge-formation ability 456 might be related with a high temperature in production process.

457

Among the PACls prepared by Al(OH)₃-dissolution, B70s0.14 with the SO₄/Al=0.14 showed
the highest performance on reducing residual SPAC particles after CSF. B70s0.14

460 outperformed commercially available PACI-70, which was used in the experiments of the 461 paragraph 3.1, 3.2, and 3.3 and effectively removed SPAC. As shown in Fig. 10 (A), 462 B70s0.14 highly reduced residual carbon particles at a higher mixing intensity, and was superior to PACI-70 for all mixing conditions tested. The superiority of B70s0.14 to PACI-463 464 70 was also confirmed in the experiments where natural water supplemented with SPAC was 465 used (Fig. 10 (B)). Moreover, the trend that high G_FT_F results in low concentration of residual 466 carbon particles, which described in the section 3.3, was also observed in the data of natural 467 water. In case of very high $G_F T_F$ (60,000) applied, very low concentrations of residual carbon 468 particles were attained with both B70s0.14 and PACI-70 and their difference was small. 469 Residual carbon particle concentrations obtained in the experiments using river water (Fig. 470 10 (B)) were somewhat lower than those using filtered tap water (Fig. 10 (A)). However the 471 effect of organic matter concentrations and turbidity should not be inferred by the comparison 472 of these two data because the two waters differed in ionic composition as well as organic 473 matter concentrations and turbidity, all of which affect coagulation performance.

474

475 The PACls produced by Al(OH)₃-dissolution, including B70s0.14, which showed the high 476 performance on the reduction of residual SPAC after CSF, contained Alc as the main 477 component. However, the percentages of Ala, Alb, and Alc were almost the same as those of the Alc-type PACls prepared by base-titration (Fig. 17S, SI), which were not effective 478 479 coagulants. This means that the Al species distribution determined by the ferron method is 480 not informative in evaluating the ability for coagulation treatment targeting SPAC, in 481 particular the ability of floc formation. Base-titration required a high temperature when 482 adjusting the basicity and resulted in a high salt (Na and Cl) content in comparison to 483 Al(OH)3-dissolution. These differences were inferred to influence the ability of floc 484 formation. Studies on PACl produced by Al(OH)3-dissolution are scarcely seen, and further

485 investigations are expected.

486

487

488 4. Conclusions

489

The addition of SPAC in water treatment is a promising technology to remove organic contaminants, but efficient removal of the loaded SPAC is needed. In this study, key control points to attain the high log removal rates of SPAC and to reduce residual SPAC particles at trace concertation levels of 100 to 1000 particles/mL in treated water were examined, and their background mechanisms were discussed, focusing on the process points of coagulationflocculation and coagulants in CSF. The following conclusions were obtained.

496

497 1) A sufficiently large G value in flash mixing for the first step of coagulation was clearly a 498 dominant factor for minimizing residual carbon particle concentration in the treated water 499 after the fourth step of sand filtration. Adequate mixing for coagulation would enable 500 complete dispersion of coagulant thereby reducing the number of particles that were not 501 charge-neutralized. Such particles would pass through the sand filter into the treated water.

502

2) Commercially available PACls were compared with each other: high-basicity PACl (basicity 70%) was more effective in removing residual carbon particles. It was because the high-basicity PACl had a high charge neutralizing capacity to destabilize many particles, though it formed flocs slowly. However, this superiority of high-basicity PACl was provided by the sufficient flash mixing. Without sufficient flash mixing, not only the removal rate decreased but also the high-basicity PACl became inferior to conventional-basicity PACl.

510 3) High-basicity PACl with SO₄/Al=0.14 reduced residual carbon particles down to the

511 minimum concentration. Such a high performance PACl coagulant was produced by using

512 Al(OH)₃-dissolution, and the dominant Al species was Alc determined by the ferron method.

513 In contrast, Alb and Alc type PACls produced by base-titration had a high positive charge

- 514 capacity and were poor at forming floc particles due to lack of bridge-forming ability; this
- 515 was true even if they were used at a large dosage or the optimal dosage in terms of charge
- 516 neutralization.
- 517
- 518

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- 522

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- 629
- 630
- 631

Table 1. Activated carbon particle size. The median diameters were determined by laser light diffraction and scattering.

Activate	Median diameter (µm)		
PA	13.7		
SPAC	SPAC ₁	0.96	
SIAC	SPAC ₂	0.90	

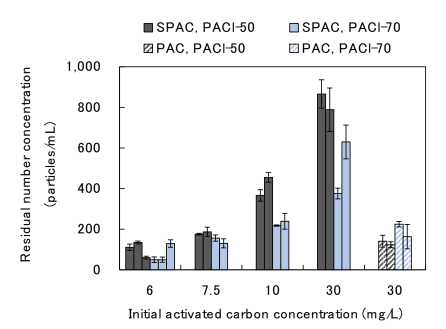
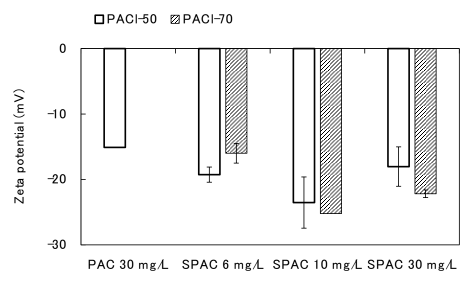


Figure 1. Particle number concentration of residual activated carbon vs. initial activated carbon concentration at a fixed mixing intensity (G_TT_T: 39,000, G_F: 600 s⁻¹, T_F: 20 s; G_{S1}: 50 s⁻¹, T_{S1}: 300 s, G_{S2}: 20 s⁻¹, T_{S2}: 300 s, G_{S3}: 10 s⁻¹, T_{S3}: 600 s). PAC and SPAC₁ were used. Error bars indicate standard deviations of measurements.



Initial activated carbon concentration

Figure 2. Zeta potentials of residual carbon particles in sand-filtrate vs. initial activated carbon concentration. Experimental conditions were the same as those described for Fig. 1. The number of measurements was described as follows: with PACI-70, two measurements for SPAC 6 mg/L and 30 mg/L, and one measurement for SPAC 10 mg/L; with PACI-50, three measurements for SPAC 30 mg/L, two measurements for SPAC 6 mg/L and 10 mg/L, and one measurement for PAC 30 mg/L. Error bars indicate standard deviations of measurements.

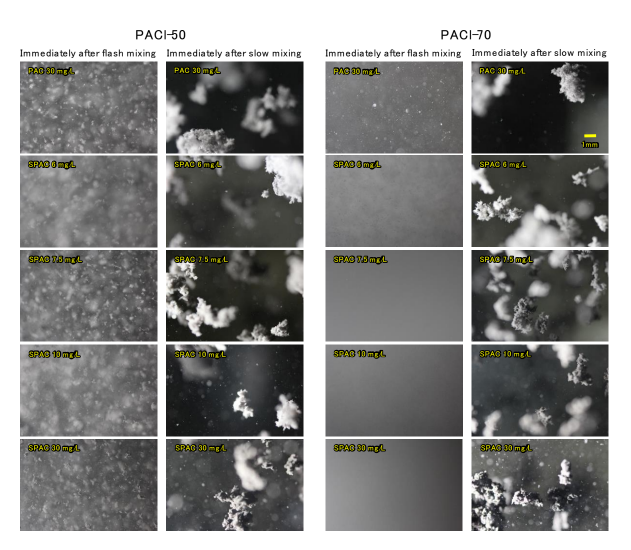


Figure 3. Photos of floc particles during the experiments of Fig. 1. The two columns of the left side are experiments using PACI-50. The two columns of the right side are experiments using PACI-70. The photos in the first column in each set of two rows were taken immediately after flash mixing was finished. The photos in the second column in each set of two rows were taken immediately after slow mixing was finished.

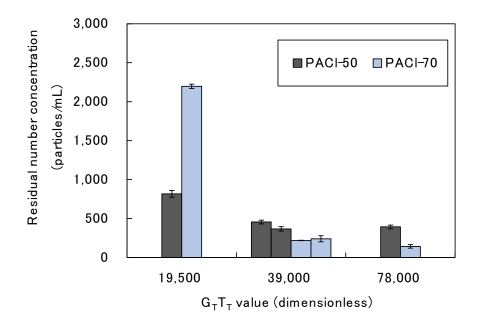


Figure 4. Effect of total mixing intensity (G_TT_T value= G_FT_F value of flash mixing+ G_ST_S value of slow mixing) on the particle number concentration of residual carbon. SPAC₁ was used. Initial SPAC₁ concentration was 10 mg/L. Details of the mixing conditions are shown in Table 2S of SI. The number of experiments for each mixing condition was described as follows: with total GT 39,000, two experiments; with total GT 19,500 and 78,000, one experiment each. Error bars indicate standard deviations of measurements.

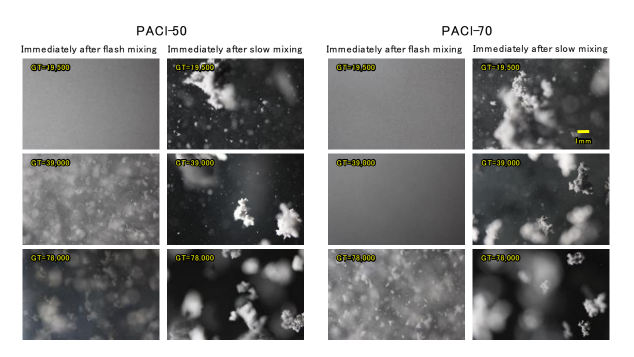


Figure 5. Photos of floc particles during the experiments of Fig. 4. The two columns of the left side are experiments using PACI-50. The two columns of the right side are experiments using PACI-70. The photos in the first column in each set of two rows were taken immediately after flash mixing was finished. The photos in the second columns in each set of two rows were taken immediately after slow mixing was finished.

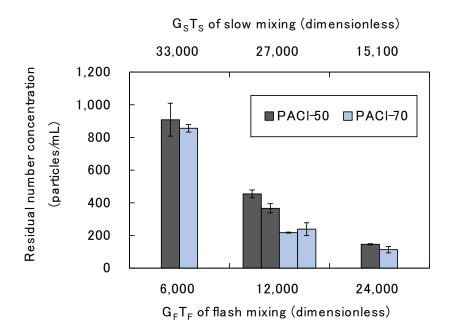
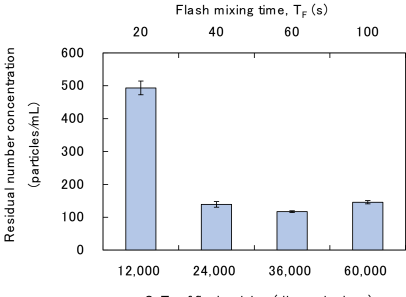


Figure 6. Effect of mixing intensity in flash and slow mixing on the particle number concentration of residual carbon at a fixed total mixing intensity. SPAC₁ was used. Initial SPAC₁ concentration was 10 mg/L. G_TT_T value (= G_FT_F of flash mixing+ G_ST_S of slow mixing) was fixed at about 39,000. Details of the mixing conditions are shown in Table 2S of SI. The number of experiments for each mixing condition was described as follows: with G_TT_T 39,000 (G_FT_F =12,000), two experiments; with G_TT_T 39,000 (G_FT_F =6,000) and 39,100 (G_FT_F =24,000), one experiment. Error bars indicate standard deviations of measurements.



G_FT_F of flash mixing (dimensionless)

Figure 7. Effect of flash mixing time (T_F) on the particle number concentration of residual carbon. SPAC₂ was used. Initial SPAC₂ concentration was 10 mg/L. PACI-70 was used. G_F value was fixed at 600 s⁻¹. G_ST_S of slow mixing=15,100. Details of the mixing conditions are shown in Table 2S of SI. Error bars indicate standard deviations of measurements.

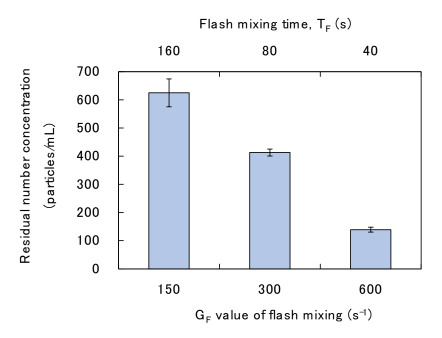


Figure 8. Effect of mixing speed (G_F) and time (T_F) in flash mixing on the particle number concentration of residual carbon at a fixed flash mixing intensity ($G_FT_F=24,000$) and a fixed slow mixing intensity ($G_ST_S=15,100$). SPAC₂ was used. Initial SPAC₂ concentration was 10 mg/L. PAC1-70 was used. Error bars indicate standard deviations of measurements.

Name	Basicity	SO ₄ /Al	Cl/Al	Na/Al	Al content	Titrating process	
						Time	Temperature
	%	mole ratio	mole ratio	mole ratio	mol/L	h	°C
B70ns	70	0.00	1.68	0.78	2.0	1.0	50
B70s0.11	70	0.11	1.46	0.79	2.0	1.0	50
B70s0.12	70	0.12	1.36	0.69	2.2	1.0	50
B70s0.13	70	0.13	1.36	0.72	2.0	1.3	50
B70s0.14-1	70	0.14	1.43	0.79	2.1	1.3	50
B70s0.14-2	70	0.14	1.44	0.81	2.1	1.4	50
B70s0.14-3	70	0.14	1.37	0.75	1.9	1.4	50
B70s0.14-4	70	0.14	1.37	0.75	1.9	1.4	50
B70s0.15-1	70	0.15	1.31	0.71	2.2	1.4	50~77
B70s0.15-2	70	0.15	1.29	0.68	2.1	1.3	50~77
B70s0.17	70	0.17	1.34	0.78	2.1	1.0	50~80

Table 2. Coagulants made by Al(OH)₃-dissolution.

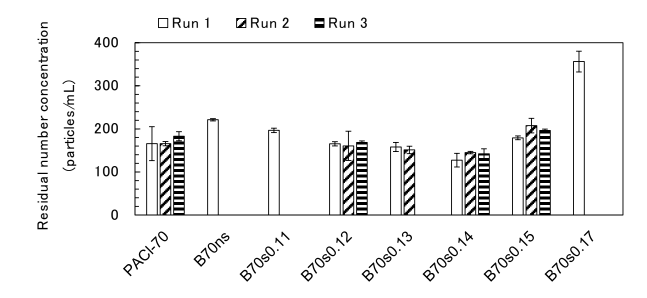


Figure 9. Particle number concentration of residual carbon after CSFs with various PACls. SPAC₁ was used. Initial SPAC₁ concentration was 10 mg/L. G_TT_T value was fixed at 39,000. Three runs were conducted for PACl-70, B70s0.12, B70s0.14 (B70s0.14-1 was used for all Runs), and B70s0.15 (B70s0.15-1 was used for Run 1, and B70s0.15-2 was used for Run 2 and 3). Two runs were for B70s0.13. One run was conducted for B70ns, B70s0.11, and B70s0.17. Error bars indicate standard deviations of three measurements.

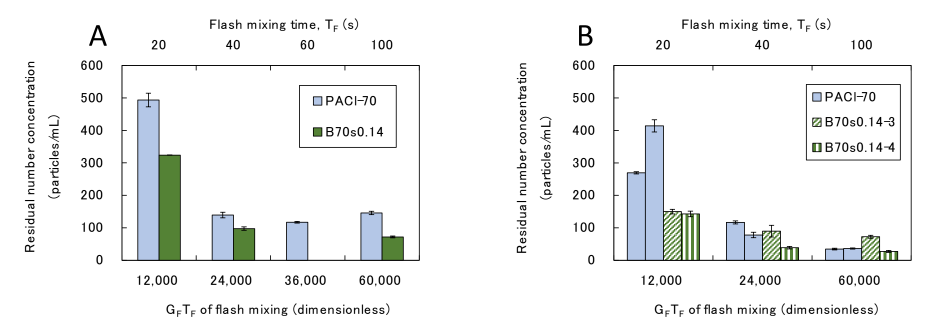


Figure 10. Effect of flash mixing time on the particle number concentration of residual carbon. SPAC₂ was used. Initial SPAC₂ concentration was 10 mg/L. G_F value was fixed at 600 s⁻¹. G_ST_S of slow mixing was 15,100. Details of the mixing conditions are shown in Table 2S of SI. Error bars indicate standard deviations of measurements. (A) Filtered tap water supplemented with SPAC₂ was used as raw water; PACl-70 and B70s0.14-2 were used as coagulant. (B) River water (turbidity 2.5 NTU and TOC 0.6 mg/L) supplemented with SPAC₂ was used as raw water; PACl-70, B70s0.14-3, and B70s0.14-4 were used.

Supplementary Information

Minimizing residual black particles in sand filtrate when applying super-fine powdered activated carbon: coagulants and coagulation conditions

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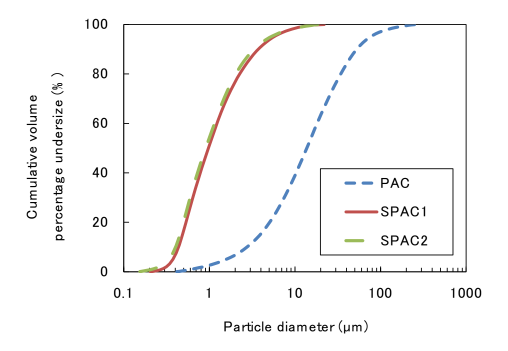


Figure 1S. Particle size distributions of PAC, SPAC₁, and SPAC₂.

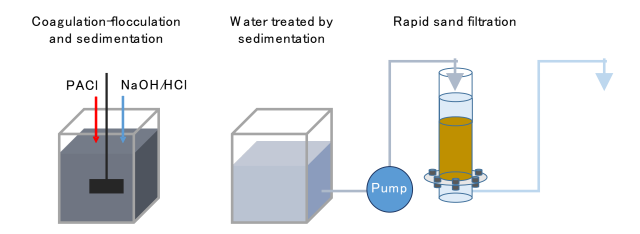


Figure 2S. Schematic diagram of the experimental setup for the coagulation-flocculation, sedimentation, and sand filtration experiment.

Calculation of G (velocity gradient) value

The mean velocity gradient $G(s^{-1})$ is defined as a function of P, μ , and V as follows.

$$G = \sqrt{\frac{P}{\mu V}} \tag{1}$$

where P is power consumed by the agitating impeller in a mixing vessel (W), μ is the viscosity of water (Pa s), and V (m³) is the volume of the water in the mixing vessel.

Power consumption is given by the product of force acting on an impeller blade and its moving velocity. Velocity and force acting on the blade of the impeller varies radially such that the tip of the blade moves fastest (tip speed) and the force acting on the tip is strongest, but the root end of the blade moves slowest and its force is smallest. Therefore, power consumed by a rotating impeller blade is given by:

$$P_B = \int v \, p dA \qquad (2)$$
$$p = \frac{1}{2} \rho_w C_d v^2 \qquad (3)$$

where, P_B is the power required to move a blade (W), v is the velocity of the moving blade (m/s), p is the pressure acting vertically on the blade (Pa), A is the area of blade (m²), ρ_w is the density of fluid (kg/m³), and C_d is drag coefficient (dimensionless).

For a flat rectangular blade shown in Fig. 3S,

$$v = 2\pi (1 - K_r) x N_s$$
(4)
$$dA = h dx$$
(5)

where, K_r is (dimensionless), x is the radial distance from the impeller center (m), N_s is rotational speed (rps), and h is the height of a impeller blade (m).

By substituting equations (3), (4) and (5) into (2):

$$P_B = \int_{r_1}^{r_2} \frac{1}{2} \rho_w C_d [2\pi (1 - K_r) x N_s]^3 h dx = \rho_w C_d [\pi (1 - K_r) N_s]^3 h (r_2^4 - r_1^4)$$
(6)

The total power input to the mixing tank having multiple impeller blades is given by:

$$P = nP_B = n\rho_w C_d [\pi N_s]^3 h(r_2^4 - r_1^4)$$
(7)

where, n is the number of impeller blades in a mixer.

Substituting Eq. (10) into Eq. (1) gives,

$$G = \sqrt{\frac{n\rho_w C_d \pi^3 (1 - K_r)^3 N_s^3 h(r_2^4 - r_1^4)}{\mu V}}.$$
 (8)

The values used for the calculation of G values by equation (8) were summarized in Table 1S.

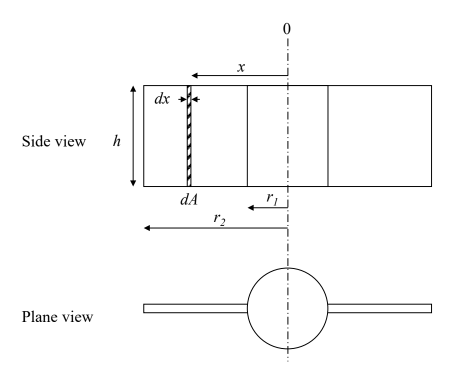


Figure 3S. Schematic diagram of mixing blade for the calculation of G value.

Parameter symbol	Unit	Value				
V	m ³	0.004				
μ	Pa s	1.005×10^{-3}				
h	m	0.035				
r_{1}	m	0.014				
r_2	m	0.05				
$ ho_w$	kg m ⁻³	9.982×10^{2}				
C_d	dimensionless	1.5				
K_r	dimensionless	0				
п	dimensionless	4				
N_S	rps	from 0.22 to 3.28				

Table1S. Parameter values used for G value calculation.

	Activ	ated carbon	Coagulant		Flash mixing		Slow mixing						Flash + slow mixing
	Name	Initial concentration (mg L ⁻¹)	Name	Dosage (mg-Al L ⁻¹)	G_F (s ⁻¹)	T _F (s)	G _{S1} (s ⁻¹)	T _{S1} (s)	G _{S2} (s ⁻¹)	T _{S2} (s)	G _{S3} (s ⁻¹)	T _{S3} (s)	G _T T _T (dimensionless)
Fig. 1 Fig. 6S Fig. 7S	PAC and SPAC ₁	30 (PAC) and 6~30 (SPAC ₁)	PACI-50 and PACI-70	4	600	20	50	300	20	300	10	600	39,000
Fig. 4 Fig. 9S	SPAC ₁	10	PACI-50 and PACI-70	4	600 600 600	10 20 40	50 50 50	150 300 600	20 20 20	150 300 600	10 10 10	300 600 1,200	19,500 39,000 78,000
Fig. 6	SPAC ₁	10	PACI-50 and PACI-70	4	600 600 600	10 20 40	50 50 50	375 300 170	20 20 20	375 300 170	10 10 10	675 600 320	39,000 39,000 39,100
Fig. 7	SPAC ₂	10	PACI-70	4	600 600 600 600	20 40 60 100	50 50 50 50	170 170 170 170	20 20 20 20	170 170 170 170	10 10 10 10	320 320 320 320	27,100 39,100 51,100 75,100
Fig. 8	SPAC ₂	10	PACI-70	4	150 300 600	160 80 40	50 50 50	170 170 170	20 20 20	170 170 170	10 10 10	320 320 320	39,100 39,100 39,100
Fig. 9	SPAC ₁	10	PACI-70 and PACIs by Al(OH)3-solution	4	600	20	50	300	20	300	10	600	39,000
Fig. 10	SPAC ₂	10	PACI-70 and B70s0.14	4	600 600 600 600	20 40 60 100	50 50 50 50	170 170 170 170	20 20 20 20	170 170 170 170	10 10 10 10	320 320 320 320 320	27,100 39,100 51,100 75,100
Fig. 10S Fig. 11S	SPAC ₁	10	PACI-50 and PACI-70	1.5~8	600	40	50	600	20	600	10	1,200	78,000

Table 2S. The experimental conditions of coagulation in CSF experiments.

G and T stand for velocity gradient and mixing time, respectively. Subscripts F, S, and the numbers after S mean flash mixing, slow mixing, and the order of the slow mixing part, respectively.

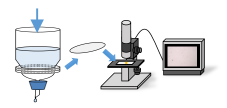


Figure 4S. Schematic diagram of the experimental setup for membrane filtration and microscopic image analysis.

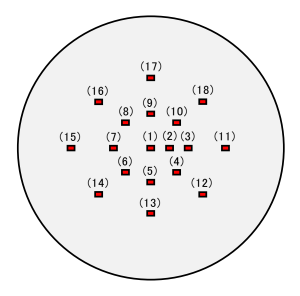


Figure 5S. The position of observation zones on a membrane filter. The observation was conducted with 18 zones in the case that the particle number concentration was expected to be lower than 200 particles/mL. In the other case, the observation was conducted with 9 zones (zone numbers 1 and from 3 to 10).

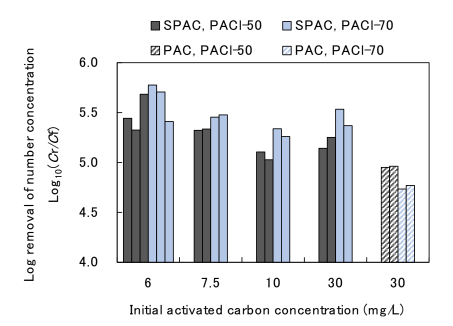


Figure 6S. Removal rate in terms of particle number concentration after CSF in the experiments of Fig. 1. *C*r and *C*f indicate the particle number concentration of carbon in raw water and sand filtrate, respectively.

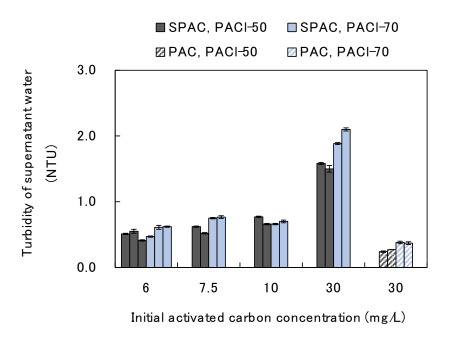


Figure 7S. Turbidity of supernatant after coagulation, flocculation, and sedimentation in the experiments of Fig. 1. Error bars indicate standard deviations of measurements.

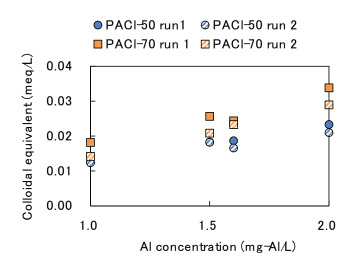


Figure 8S. Colloid charge capacity of PACI-50 and PACI-70 (Matsui et al. 2017).

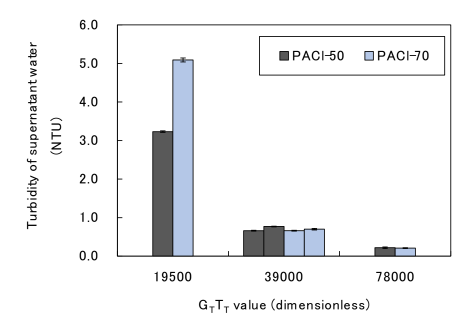


Figure 9S. Turbidity of supernatant after coagulation, flocculation, and sedimentation in the experiments of Fig. 4. Error bars indicate standard deviations of measurements.

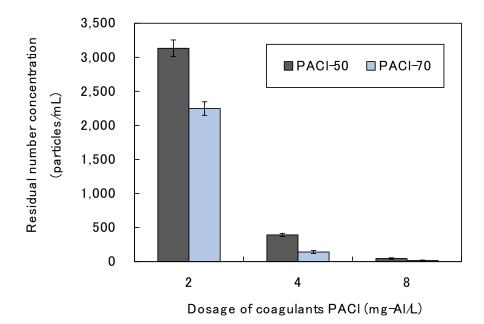


Figure 10S. Effect of coagulant dosage on the particle number concentration of residual carbon in sand filtrate. SPAC₁ was used. Initial SPAC₁ concentration was 10 mg/L. G_TT_T value was fixed at 78,000 (G_FT_F=24,000, G_ST_S=54,000). Error bars indicate standard deviations of measurements.

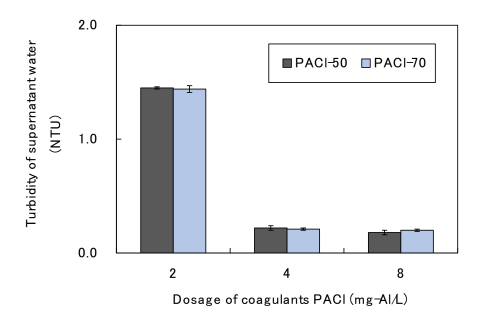


Figure 11S. Turbidity of supernatant after coagulation, flocculation, and sedimentation in the experiments of Fig. 10S. Error bars indicate standard deviations of measurements.

Name	Basicit y	SO4/Al	Cl/Al	Na/Al	Al conten t	Heating process		
						Time	Temperatur e	
	%	molar ratio	molar ratio	molar ratio	mol/L	h	C°	
B65ns	65	0	3.00	1.95	0.1			
B65ns-heat	65	0	3.00	1.95	0.1	5.0	121	
B79s0.10-1	79	0.10	2.80	2.38	0.04			
B79s0.10-2-heat	79	0.10	2.80	2.38	0.09	(5.0)	(121)	
B80ns-1	80	0	3.00	2.40	0.05			
B80ns-2	80	0	3.00	2.40	0.1			
B80ns-2-heat	80	0	3.00	2.40	0.1	0.33	120	
B82s0.11	82	0.11	2.79	2.45	0.1			
B82s0.11-heat	82	0.11	2.79	2.45	0.1	72	90~95	
B88s0.10	88	0.10	2.80	2.64	0.05			

Table 3S. Coagulants made by base-titration.

Preparation of coagulants by base-titration

The 10 poly-aluminum chlorides (PACls) were produced by base-titration. These PACls are designated by the following rules: the number after "B" represents "% basicity", "s" indicates "sulfated", "ns" indicates "non-sulfate", "heat" indicates "heated", the number after "s" represents "mole ratio of sulfate ion to aluminum: SO_4/Al ", and the number after the hyphen indicates serial number. $AlCl_3 \cdot 6H_2O$, $Al_2(SO_4)_3 \cdot 14 \sim 18H_2O$, and Na_2CO_3 were provided by FUJIFILM Wako Pure Chemical Corporation (Osaka, Japan), and NaOH was by Kanto Chemical Co., Inc., (Tokyo, Japan).

B65ns was prepared as follows: an AlCl₃ solution (0.5 M as Al, 80 mL) in a 500 mL Erlenmeyer flask was titrated with NaOH (0.24 M, 320 mL) by means of a peristaltic pump at a rate of 4 mL/min. During the titration, a combined hot plate/magnetic stirrer was used to agitate the solution in the flask and maintain the temperature at 85–90°C. B80ns-1 was prepared with the same materials and procedure for B65ns, except that the concentrations of AlCl₃ solution and NaOH were 0.25 M as Al and 0.15 M. The same materials and procedure were used to prepare B80ns-2, except that the concentration of AlCl₃ solution was 0.3 M. B82s0.11 was prepared as follows: an Al solution (0.49 M as Al, SO₄/Al=0.11 in mole ratio, 80 mL) was prepared with AlCl₃ and Al₂(SO₄)₃ in a 500 mL Erlenmeyer flask and titrated with NaOH (0.3 M, 320 mL) as the same procedure described above.

B65ns-heat was prepared by heating B65ns at 121°C for 5 h with an autoclave (MLS-3781, PHC Holdings Corporation, Tokyo, Japan). B80ns-2-heat was prepared by heating B80ns-2 at 120°C for 20 min with the same autoclave. B80s0.05-heat was prepared by heating B82s0.11 at 90~95°C for 72 h with a combined hot plate/magnetic stirrer.

B79s0.10-1 was prepared by adding a Na₂CO₃ solution and an Al₂(SO₄)₃ solution into B65ns at ambient temperature until the basicity became 79% (calculation of the basicity is described in paragraph 2.2 of the main paper) and SO₄/Al became 0.10. B79s0.10-2-heat was prepared by adding a Na₂CO₃ solution and an Al₂(SO₄)₃ solution into B65ns-heat at ambient temperature. B88s0.10 was also prepared by adding a Na₂CO₃ solution and an Al₂(SO₄)₃ solution into B65ns-heat at ambient temperature.

SPAC	Coagulant	Fla	sh	Slow						Total
Median diameter	Dosage	G _F	$T_{\rm F}$	G _{S1}	$T_{\rm S1}$	G_{S2}	$T_{S2} \\$	G_{S3}	T_{S3}	G _T T _T
μm	mg-Al/L	s ⁻¹	S	s^{-1}	S	s^{-1}	S	s^{-1}	S	dimensionless
1.0	0~1.5	190	60	19	600					22,800
1.0	0~1.5	200	60	20	600	15	600	10	1,200	45,000
1.1	1.5	200	120	20	600	15	600	10	1,200	57,000

Table 4S. The mixing conditions of coagulation in the jar-test (Figures 13S to 16S).

Jar-tests using the coagulants made by base-titration

Filtered tap water was obtained by the method described in the main text of this paper (paragraph 2.3) except that its alkalinity was not adjusted. The water was supplemented with an SPAC slurry (median diameter is described in Table 4S) at concentration 10 mg/L to make raw water for the experiment consisting of coagulation-flocculation and sedimentation (jar test). Jar-tests were conducted in a 1-L rectangular beaker. After a predetermined volume of HCl or NaOH (0.1 N) was added to order to achieve the coagulation pH at 7.0, the coagulant (PACl) was injected into the beaker to a final concentration between 0 and 1.5 mg-Al/L (see also Table 4S). Then, the water was mixed rapidly and slowly. The mixing speed (G value) and times (T value) of the flash and slow mixing were varied depending on each experiment (see also Table 4S). The water was then left at rest for 1 h. A supernatant of 50 mL was sampled and determined for turbidity (2100Q portable turbidimeter; Hach Company, Loveland, CO, USA).

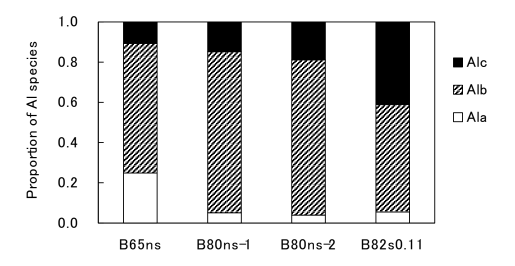


Figure 12S. Proportions of Al species in PACl coagulants made by base-titration.

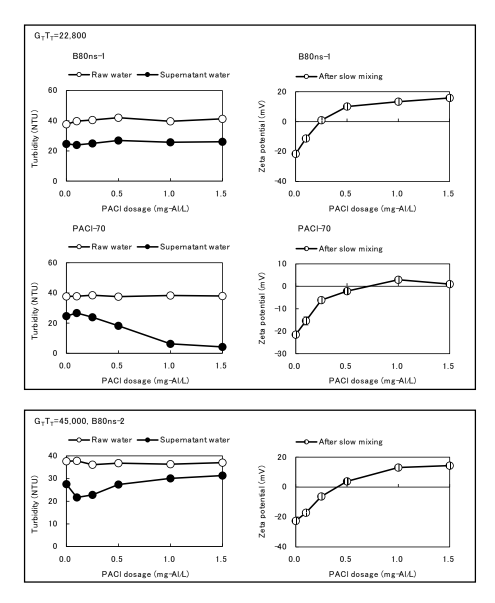


Figure 13S. Turbidity removal of and zeta potential of carbon particles. Initial SPAC concentration was 10 mg/L. Alb type base-titration PACls (B80ns-1, B80ns-2) and PACl-70 were used as coagulant with a dosage of 0 to 1.5 mg-Al/L. Coagulation pH was 7.0. Settling time was 60 min. Error bars of turbidity and zeta potential indicate standard deviations of three measurements and five measurements, respectively. Details of the mixing conditions are shown in Table 4S.

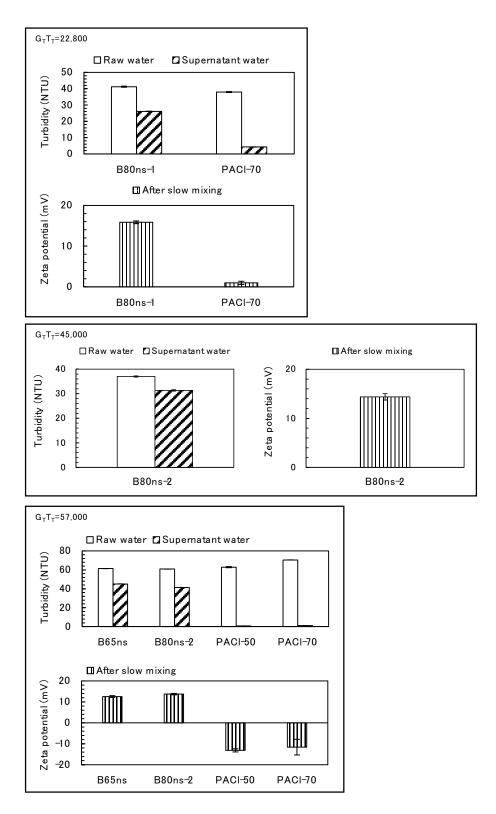


Figure 14S. Turbidity removal of and zeta potential of carbon particles. Initial SPAC concentration was 10 mg/L. Alb type base-titration PACls (B65ns, B80ns-1 and B80ns-2), PACl-50, and PACl-70 were used as coagulant with a dosage of 1.5 mg-Al/L. Coagulation pH was 7.0. Settling time was 60 min. Error bars of turbidity and zeta potential indicate standard deviations of three measurements and five measurements, respectively. Details of the mixing conditions are shown in Table 4S.

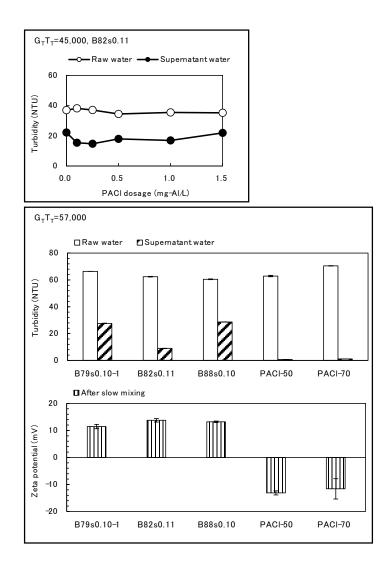


Figure 15S. Turbidity removal and zeta potential of carbon particles. Initial SPAC concentration was 10 mg/L. Base-titration PACls that contained sulfate ion (B79s0.10-1, B82s0.11, B88s0.10), PACl-50, and PACl-70 were used as coagulant with a dosage of 0 to 1.5 mg-Al/L. Coagulation pH was 7.0. Settling time was 60 min. Error bars of turbidity and zeta potential indicate standard deviations of three measurements and five measurements, respectively. Details of the mixing conditions are shown in Table 4S.

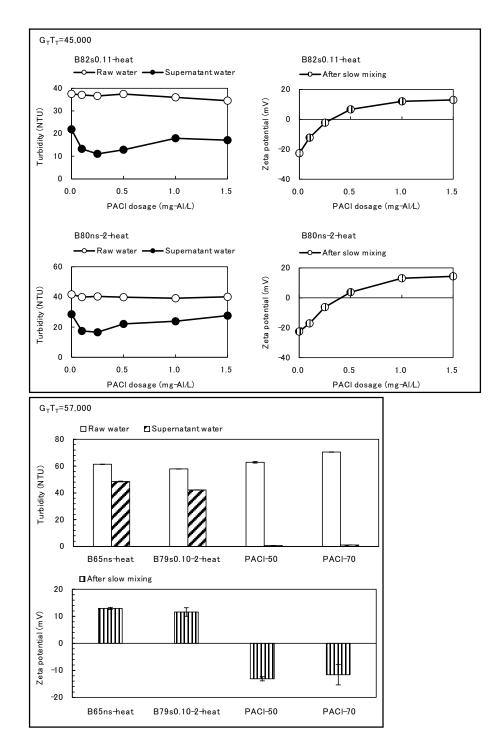


Figure 16S. Turbidity removal and zeta potential of carbon particles. Initial SPAC concentration was 10 mg/L. Alc-type base-titration PACls (B82s0.11-heat, B80ns-2-heat, B65ns-heat, and B79s0.10-2-heat), PACl-50, and PACl-70 were used as coagulant with a dosage of 0 to 1.5 mg-Al/L. Coagulation pH was 7.0. Settling time was 60 min. Error bars indicate standard deviations of measurements. Details of the mixing conditions are shown in Table 4S.

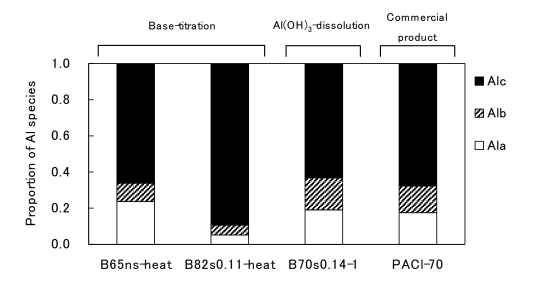


Figure 17S. Proportions of Al species in PACl coagulants made by base-titration, Al(OH)₃-dissolution, and commercial product.

Referene

Matsui, Y., Shirasaki, N., Yamaguchi, T., Kondo, K., Machida, K., Fukuura, T. and Matsushita, T. (2017) Characteristics and components of poly-aluminum chloride coagulants that enhance arsenate removal by coagulation: Detailed analysis of aluminum species. Water Research 118, 177-186.