Effect of aluminum hydrolyte species on human enterovirus removal from water during the coagulation process

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Abstract

We prepared different types of aluminum-based coagulants, consisting of mainly monomeric aluminum species, polymeric aluminum species, or colloidal aluminum species, to investigate the effect of aluminum hydrolyte species on the removal of two types of human enteroviruses, poliovirus (PV) type 1 and the free-chlorine-resistant virus coxsackievirus (CV) B5, from lake and river water samples during the coagulation process. We found that differences in the distribution of the aluminum hydrolyte species in the coagulant affected the removal of these enteroviruses during coagulation: the removal ratios of PV and CV observed with polyaluminum chloride (PACl) with a high colloidal
aluminum content and a basicity of 2.1 (i.e., PACl-2.1c) were larger than those observed with high monomeric aluminum content coagulant (i.e., AlCl₃ solution) and with high polymeric aluminum content coagulant PACl (PACl-2.1b). Unlike AlCl₃ or PACl-2.1b, PACl-2.1c contains Al₃₀ species, indicating that Al₃₀ species probably play a major role in the removal of enteroviruses. The PV and CV removal ratios were almost identical, regardless of the coagulant type or viral quantification method used (plaque-forming unit method or real-time polymerase chain reaction method), suggesting that PV and CV behaved similarly during the coagulation process. We also experimentally confirmed that the main mechanism for virus removal was coprecipitation into growing aluminum hydroxide during charge neutralization; virus adsorption onto formed aluminum hydroxide flocs also contributed to virus removal, but played a limited role.

**Key words:** Aluminum hydrolyte species, Basicity, Charge neutralization, Coagulation, Coxsackievirus, Poliovirus
1. Introduction

Coagulation is an important process in water treatment to remove contaminants. Of the various coagulants used for the coagulation process, prepolymerized aluminum coagulants, such as polyaluminum chloride (PACl), have been widely used in drinking water treatment because of their superiority to traditional aluminum coagulants [e.g., aluminum chloride (AlCl₃) and alum]; PACls are more efficient, are less dependent on temperature and pH, can be used at a lower dosage, and produce less sludge [1–3].

The hydrolysis reactions of aluminum species are very complicated, depending on water quality and coagulant type, which contribute to the formation of various aluminum hydrolyte species [4, 5]. The ferron method has been employed extensively to categorize the aluminum hydrolyte species into the following three fractions on the basis of the kinetic differences between the reactions of the aluminum species and the ferron regent: monomeric species, fast-reacting polymeric species, and slow-reacting colloidal species, denoted as Ala, Alb, and Alc, respectively [6]. Ala is composed mainly of monomeric species such as Al³⁺, Al(OH)²⁺, and Al(OH)₃⁺; dimeric and trimeric species such as [Al₂(OH)₂]⁴⁺ and [Al₃(OH)₄]⁵⁺ are also classified as Ala [1]. Alb is the intermediate polymeric species, and many researchers have suggested that Alb could include the Al₁₃ species [AlO₄Al₁₂(OH)₂₄(H₂O)₁₂]⁷⁺, which was identified by using liquid ²⁷Al nuclear magnetic resonance (NMR) spectroscopy [1, 4, 7]. Alc is the large polymer or colloidal species, and the Al₃₀ species [Al₃₀O₈(OH)₅₆(H₂O)₂₄]¹⁸⁺, which was also identified by using liquid ²⁷Al NMR spectroscopy, is classified into Alc [7].
The distribution of aluminum hydrolyte species in PACls and the formation of specific aluminum hydrolyte species such as Al$_{13}$ and Al$_{30}$ might be controlled by basicity ([OH$^-$/[Al$^{3+}$]), aluminum concentration, rate and mode of neutralization, and reaction temperature and time during PACl preparation [7–9]. In addition, the aluminum species originally contained in the PACls or formed in situ during the coagulation process play an important role in determining the behavior and efficiency of coagulants [4, 5]. For these reasons, many researchers have paid particular attention to the effects of aluminum species on coagulation efficiency. For example, Yan et al. [1] reported that the removal efficiencies of turbidity and dissolved organic carbon (DOC) correlate well with the Al$_c$ content and Al$_b$ content in PACls, respectively, whereas both Al$_b$ and Al$_c$ content influence the removal of ultraviolet absorbance at 254 nm (UV254, an indicator of natural organic matter concentration). Our research group [10] and Duan et al. [11] reported that the residual aluminum concentration in treated water is closely related to the Al$_a$ content in the PACl: low Al$_a$ content in the PACl yields a low residual aluminum concentration. Moreover, among the identified aluminum hydrolyte species in PACls, Al$_{13}$ and Al$_{30}$ are believed to be effective coagulation species due to their strong charge neutralization capability and high structural stability [12]. In fact, Lin et al. [13] showed that DOC removal by PACl with a high Al$_{13}$ content is superior to that by commercially available PACl with a low Al$_{13}$ content at about pH 6, and that a lower dosage of high Al$_{13}$ content PACl than of commercially available PACl is required for optimal DOC removal. Zhang et al. [2] reported that compared to AlCl$_3$, PACl with a high Al$_{30}$ content exhibits greater UV254 removal efficiency and
leads to a lower residual aluminum concentration across a broad pH range and a wide coagulant dosage range. In addition, Hu et al. [4] showed the value of PACl with a high Al$\text{_{13}}$ content for the removal of arsenic (As) from As-spiked tap water over a broad pH range, and concluded that the removal efficiencies of As(V) correlated with the amount of Al$\text{_{13}}$ species present. Moreover, Mertens et al. [14] reported that PACl with a high Al$\text{_{30}}$ content contributed to the efficient removal of As(III) and As(V) from As-contaminated groundwater at pH 7–8.

Human enteric viruses, which are frequently present in contaminated drinking water sources and do not settle from suspension by gravity, can also be removed by the coagulation process with PACl [15]. However, the relationship between the removal efficiencies of human enteric viruses and the aluminum hydrolyte species in PACl remains unclear, and the role of Al$\text{_{13}}$ and Al$\text{_{30}}$ species in the removal of human enteric viruses has not been investigated. We recently reported that laboratory-prepared PACl with a high Al$\text{_{c}}$ content removed bacteriophages (i.e., viruses that infect bacteria) more efficiently than did laboratory-prepared PACl with a high Al$\text{_{b}}$ content or commercially available PACls, suggesting that the Al$\text{_{30}}$ species in high Al$\text{_{c}}$ content PACl probably play a major role in bacteriophage removal during the coagulation process [16]. Because the removal ratios of human enteric viruses including poliovirus were different to and smaller than that of bacteriophage MS2 in the coagulation process with aluminum-based coagulant [17], the effect of aluminum hydrolyte species on virus removal, and specifically the role of Al$\text{_{13}}$ and Al$\text{_{30}}$, may also differ between human enteric virus and bacteriophage removal. In addition, because human enteric viruses have high
resistance to free-chlorine disinfection compared with human enteric bacteria [18], and the increase in free-chlorine dosage needed for sufficient disinfection of human enteric viruses sometimes results in the formation of high levels of toxic disinfection by-products [19], improvements in the coagulation efficiency of virus removal are highly desired. Therefore, the identification of aluminum hydrolyte species in aluminum-based coagulants that efficiently remove human enteric viruses from drinking water would improve coagulation efficiency and support the development of novel aluminum-based coagulants for the prevention and control of waterborne disease caused by exposure to such viruses through drinking water.

Here, we conducted batch coagulation experiments to investigate the effect of aluminum hydrolyte species on the removal of human enteric viruses, specifically human enteroviruses, by comparing five coagulants with different distributions of aluminum hydrolyte species. In addition, we investigated the mechanism of human enterovirus removal during the coagulation processes by examining the coprecipitation of enteroviruses into growing aluminum hydroxide during charge neutralization and the absorption of enteroviruses onto preformed aluminum floc particles. Because poliovirus (PV) type 1 is commonly used as representative of human enteric viruses [17, 20], and coxsackievirus (CV) B5 has high resistance to free-chlorine disinfection compared with other types of CVs (e.g., CV B3 and CV B4) and other human enteric viruses including PVs (types 1, 2 and 3), echoviruses (EV types 1 and 11), and adenoviruses (AdV types 40 and 41) [21, 22], we chose these two human enteroviruses for use in our study.
2. Materials and Methods

2.1. Source water and coagulants

Lake water and river water were sampled from Lake Imba-numa in Chiba, Japan, and the Toyohira River in Sapporo, Japan, on 17 November 2014 (water quality data are shown in Table 1). The source water samples were stored at 4 °C until use, and the temperature was adjusted to 20 °C prior to use.

Five aluminum-based coagulants were used for the coagulation experiments (Table 2). AlCl₃ solution was prepared by dilution of reagent-grade aluminum(III) chloride hexahydrate (AlCl₃·6H₂O, Wako Pure Chemical Industries, Osaka, Japan) dissolved in Milli-Q water (Milli-Q Advantage, Millipore Corp., Billerica, MA, USA). Four nonsulfated high-basicity PACls (PACl-2.1b, PACl-2.4b, PACl-2.1c, and PACl-2.4c, where 2.1 and 2.4 are basicity values, and “b” and “c” indicate high Al₃⁺ and Al₅⁻ content, respectively, as measured by the ferron method; see below) were prepared by using a base titration method in our laboratory. A certain amount of 0.25 M AlCl₃ solution was transferred to a glass beaker, and then the solution was titrated with 0.15 M NaOH at a constant rate (4 mL/min) by use of a peristaltic pump to achieve the targeted basicity value to prepare PACl-2.1b and PACl-2.4b. The solution in the beaker was stirred at 630 rpm and the temperature was maintained at 90–95 °C by a hot plate stirrer during the titration. PACl-2.1c and PACl-2.4c were prepared from 1.5 M AlCl₃ solution and 0.9 M NaOH by using the same method just described, and then the solution was
continuously stirred at 630 rpm and kept at 90–95 °C for 24 h after the titration. All coagulants were used in batch coagulation experiments immediately after dilution with Milli-Q water.

2.2. Characterization of coagulants

The aluminum hydrolyte species in the coagulants were analyzed by using the ferron method and liquid $^{27}$Al NMR spectroscopy after the coagulants were diluted with Milli-Q water to a concentration of 2.7 g-Al/L, that is, 0.1 M-Al (for AlCl$_3$ solution, PACl-2.1c, and PACl-2.4c), or 1.35 g-Al/L, that is, 0.05 M-Al (for PACl-2.1b and PACl-2.4b), depending on the aluminum concentration in the coagulant (Table 2). The positive colloid charges of the coagulants (diluted with Milli-Q water to 1–3 mg-Al/L) were determined by using a colloid titration method. The details of the ferron method, liquid $^{27}$Al NMR analysis, and the colloid titration method were described in our previous report [16].

2.3. Human enteroviruses

The Sabin attenuated LSc/2ab strain of PV type 1 was kindly provided by Dr. Hiroyuki Shimizu, National Institute of Infectious Diseases, Tokyo, Japan. The Faulkner strain of CV B5 (ATCC VR-185) was obtained from American Type Culture Collection (ATCC, Manassas, VA, USA). PV and CV were propagated in buffalo green monkey kidney epithelial cells (BGM cells), kindly supplied by Dr. Daisuke Sano, Hokkaido University, Sapporo, Japan, and were maintained in 1X Eagle’s minimum essential medium (EMEM, with phenol red, Nissui Pharmaceutical Co. Ltd., Tokyo, Japan)
supplemented with 10% (v/v) heat-inactivated fetal bovine serum (Life Technologies, Carlsbad, CA, USA), 2 mM L-glutamine (Life Technologies), 100 units/mL penicillin, 100 µg/mL streptomycin (Pen Strep, Life Technologies), and 1.125 g/L NaHCO₃. One milliliter of 10-fold diluted PV or CV stock solution [diluted with Dulbecco’s phosphate-buffered saline (without Ca and Mg ions, Nissui Pharmaceutical Co. Ltd.)] was inoculated onto a monolayer of approximately 90% confluent BGM cells in a 75 cm² flask. The flask was then incubated in a humidified incubator at 37 °C in 5% CO₂ for 30 min. After this incubation, 20 mL of 1X EMEM was added to the flask, and then the flask was incubated at 37 °C in 5% CO₂ for 2–3 days until 100% cytopathic effects of BGM cells were confirmed, at which point the viruses were harvested. Following incubation, viruses were released from the infected BGM cells by freezing at −83 °C and thawing at 37 °C, three times. The PV or CV culture solution was passed through a membrane filter (nominal pore size 0.2 µm, hydrophilic cellulose acetate; Dismic-25cs, Toyo Roshi Kaisha, Ltd., Tokyo, Japan) to prepare the PV or CV stock solution. The filtrate was further purified by using a tangential-flow filtration cassette (nominal molecular weight cutoff 1,000,000, regenerated cellulose (RC); Pellicon XL, Millipore Corp.), and by using a centrifugal filter device (nominal molecular weight cutoff 100,000, RC; Amicon Ultra-15, Millipore Corp.) twice to ensure that only a minimal amount of DOC components from the culture medium was introduced into the virus-spiked source water (see Section 2.4). The concentrations of the purified PV and CV solutions were approximately 10⁶ and 10⁷ PFU/mL, respectively.
2.4. Coagulation experiments

Batch coagulation experiments were conducted with 300 mL of virus-spiked source water in square plastic beakers at 20 °C. The purified PV or CV solution (see Section 2.3) was added to the beaker at a concentration of approximately $10^3$ PFU/mL ($C_0$). Because the purified virus solution was diluted during spiking of the source water, virus addition contributed less than 0.2–0.3 mg/L of unintentional DOC carryover. After enough HCl or NaOH was added to the spiked water to bring the final pH to a target value of 6, 7, or 8, coagulant was injected into the water. Because the dosages used in the actual drinking water treatment plants for the treatments of Lake Imba-numa water (Kashiwai drinking water treatment plant, Chiba, Japan) and Toyohira River water (Moiwa drinking water treatment plant, Sapporo, Japan) were 2.70 and 1.08 mg-Al/L, respectively, on the day we sampled the source water, those coagulant dosages were used in the present study. The water was stirred rapidly for 1 min ($G = 200$ s$^{-1}$, 94 rpm) and then slowly for 10 min ($G = 20$ s$^{-1}$, 20 rpm) with an impeller stirrer. The water was then left at rest for 60 min to allow the generated aluminum floc particles to settle. Supernatants were taken from the beaker after settling for quantification of the PV or CV concentration ($C_s$).

To investigate the importance of charge neutralization on virus removal during the coagulation process, 300 mL of source water without virus addition was coagulated rapidly for 1 min at pH 7, as described above. Following charge neutralization, the purified PV solution was added to the beaker at approximately $10^3$ PFU/mL ($C_0$). The water was stirred slowly for 10 min and then left at rest for 60 min. Supernatants were taken from the beaker after settling for quantification of the PV
concentration ($C_s$). To assess virus adsorption onto preformed aluminum floc particles, 300 mL of source water without virus addition was coagulated rapidly for 1 min and then slowly for 10 min at pH 7, as described above, to generate aluminum floc particles. Following the formation of the aluminum floc particles, the purified PV solution was added to the beaker at approximately $10^3$ PFU/mL ($C_0$). The water was stirred slowly for 10 min and then left at rest for 60 min. Supernatants were taken from the beaker after settling for quantification of the PV concentration ($C_s$).

2.5. Enterovirus assay

Infectious PV or CV was quantified by use of a plaque assay. Approximately 90% confluent BGM cells in a 75 cm$^2$ flask were seeded in 6-well (8.96 cm$^2$/well) tissue culture plates. One milliliter of 10-fold serially diluted sample (diluted with 1X EMEM supplemented with 2 mM L-glutamine, 100 units/mL penicillin, 100 μg/mL streptomycin, and 1.125 g/L NaHCO$_3$) or of 2-fold diluted sample (diluted with 2X EMEM containing 4 mM L-glutamine, 200 units/mL penicillin, 200 μg/mL streptomycin, and 2.25 g/L NaHCO$_3$) was inoculated onto a monolayer of 100% confluent BGM cells in a 6-well plate, which was then incubated in a humidified incubator at 37 °C in 5% CO$_2$ for 90 min. After this incubation, the inoculum was removed by inverting the plate, and then 3 mL of agar overlay was applied to the monolayer. The overlay was prepared by combining equal volumes of 2.5% (w/v) agarose (Agar-EPI, Nacalai Tesque, Inc., Kyoto, Japan) and 2X EMEM (without phenol red, Nissui Pharmaceutical Co. Ltd.) supplemented with heat-inactivated fetal bovine serum (2% for PV, 20%
for CV), 4 mM L-glutamine, 200 units/mL penicillin, 200 μg/mL streptomycin, and 2.25 g/L NaHCO₃.

After the addition of the agar overlay, the plate was incubated at 37 °C in 5% CO₂ for 2 days. After this incubation, the cell monolayer was stained with 1 mL of 0.15 g/L neutral red at 37 °C in 5% CO₂ for 3 h, and then excess stain was discarded by inverting the plate. Plaques of each well were counted for 1–3 days (1 day for PV, 3 days for CV) after discarding the neutral red until no new plaques appeared. The average plaque count of triplicate wells or twelve wells prepared from one sample was considered as the infectious PV or CV concentration for that sample.

The real-time polymerase chain reaction (PCR) method, which detects all viruses regardless of their infectivity or the presence of aggregates, was used to quantify viral RNA. PV and CV RNA was specifically quantified by using the real-time reverse transcription-PCR (RT-PCR) method. Viral RNA was extracted from 200 μL of sample with a QIAamp MinElute Virus Spin Kit (Qiagen K. K., Tokyo, Japan) to obtain a final volume of 20 μL. The extracted RNA solution was added to a High-Capacity cDNA Reverse Transcription Kit with RNase Inhibitor (Applied Biosystems Japan, Tokyo, Japan) for the RT reaction, which was conducted at 25 °C for 10 min, 37 °C for 120 min, and 85 °C for 5 s, with subsequent cooling to 4 °C in the thermal cycler (Thermal Cycler Dice Model TP600, Takara Bio Inc., Otsu, Japan). The cDNA solution was then amplified with a TaqMan Universal PCR Master Mix, No AmpErase UNG (Applied Biosystems Japan) with 400 nM primers (HQ-SEQ grade, Takara Bio Inc.) and 250 nM TaqMan probe (Applied Biosystems Japan). The oligonucleotide sequences of the primers and the probe used for the PV and CV quantifications were taken from
previous reports [23, 24]. Amplification was conducted at 50 °C for 2 min, 95 °C for 10 min, and then 40 cycles of 95 °C for 15 s and 60 °C for 1 min in an Applied Biosystems 7300 Real-Time PCR System (Applied Biosystems Japan). The standard curve for the real-time PCR method was based on the relationship between the infectious PV or CV concentration of a freshly prepared purified PV or CV solution measured by the plaque assay described above, and the number of cycles (Ct value) of PCR amplification.

2.6. Aluminum species distribution analysis during the coagulation process

The effect of pH on the aluminum hydrolyte species transfer of coagulants during the coagulation process was analyzed mostly by following the method of Yan et al. [25]. Batch coagulation experiments were conducted with 1,000 mL of prepared Milli-Q water containing 0.5 mM NaHCO₃ and NaNO₃ in square plastic beakers at 20 °C. After enough HCl or NaOH was added to the prepared Milli-Q water to bring the final pH to a target value of 6, 7, or 8, 2.7 mg-Al/L of coagulant was injected in the water. The water was stirred rapidly for 1 min (G = 339 s⁻¹, 200 rpm), and then the solution was analyzed by using the ferron method.

3. Results and Discussion

3.1. Characterization of coagulants
To investigate the influence of the type of aluminum hydrolyte species present in coagulants on virus removal, five aluminum-based coagulants containing different aluminum hydrolyte species were prepared and used in this study. Application of the ferron method showed that the major aluminum species in AlCl₃ was a monomeric aluminum species (Alₐ) (Table 2); ²⁷Al NMR spectroscopy confirmed that the major aluminum species in AlCl₃ was monomeric because only two peaks, (the aluminum monomer peak at 0 ppm and the internal standard substance peak at 80 ppm) were observed in the AlCl₃ spectrum (Figure 1). In contrast, the major aluminum species in PACl-2.1b and PACl-2.4b was polymeric aluminum species (Alₐ) (Table 2). Moreover, unlike AlCl₃, these PACls contained Al₁₃ species as shown by the central tetrahedral Al peak in the Al₁₃ species at 63 ppm in the ²⁷Al NMR spectra of PACl-2.1b and PACl-2.4b (Figure 1). In contrast, PACl-2.1c and PACl-2.4c contained a high proportion of colloidal aluminum species (Alₐ) (Table 2), and these PACls contained not only Al₁₃ species but also Al₃₀ species, which was not the case for PACl-2.1b and PACl-2.4b, because the broad peaks of the octahedral Al of the external shells in the Al₁₃ and Al₃₀ species at 10–12 ppm and the central tetrahedral Al in the Al₃₀ species at 70 ppm were observed in the ²⁷Al NMR spectra of PACl-2.1c and PACl-2.4c, in addition to the peak of the Al₁₃ species (Figure 1). These results concur with those of other researchers who have reported that aluminum monomer can be classified as Alₐ, Al₁₃ species can be classified as Alₐ, and Al₃₀ species can be classified as Alₐ [1, 4, 7].
3.2. Effect of aluminum species in coagulants on virus removal

The effect of aluminum species in the coagulants on the infectious PV removal ratio (log\(_{10}[C_0/C_s]\)) during the coagulation process was evaluated by using the PFU method after settling (Figure 2a). Because of the small size of PV, no PV removal (0.1-log\(_{10}\)) was observed in the absence of coagulant at approximately pH 7. In contrast, the coagulation process removed infectious PV over the pH range of 6–8 no matter what type of coagulant was used. The addition of the coagulant destabilized the dispersion of the virus particles in the source water. The particles were then entrapped in or adsorbed onto the aluminum floc particles generated during coagulation, and then the floc particles with the entrapped or adsorbed viruses settled from suspension under the influence of gravity during the settling process. The removal ratios of the infectious PV depended on the coagulant type: whereas coagulation with AlCl\(_3\) and PACl-2.4b resulted in approximately 2-log\(_{10}\) removal over a pH range of 6–7, approximately 3-log\(_{10}\) removal was achieved with PACl-2.1b, PACl-2.1c, and PACl-2.4c. In addition, the removal ratios observed with PACl-2.1c were somewhat larger than those observed with the other aluminum-based coagulants, and approximately 2-log\(_{10}\) removal was obtained with PACl-2.1c even at pH 8. Taken together, these data indicate that the ability of the coagulants to remove infectious PV from virus-spiked lake water followed the order: PACl-2.1c > PACl-2.4c ≥ PACl-2.1b > PACl-2.4b = AlCl\(_3\). The removal ratios of total PV, evaluated by using the PCR method, were also observed to be somewhat larger with PACl-2.1c than those observed with the other aluminum-based coagulants tested (Figure 2b). The ability of PACl-2.1c to remove infectious PV from virus-spiked
river water was also higher than that of the other aluminum-based coagulants, in agreement with the result from the virus-spiked lake water experiments (Figure 3a), although the removal ratios obtained with the lake water and the river water were different, due to differences in water quality (Table 1). Moreover, the total PV removal ratios were also found to be somewhat larger with PACl-2.1c than those with the other aluminum-based coagulants tested in the river water sample (Figure 3b). These results indicate that differences in the distribution of the aluminum species in the coagulants affected virus removal performance during coagulation and that PACl-2.1c effectively removed PV, particularly at pH 6–7.

In the present study, similar removal ratios of PV were observed with AlCl₃ and PACl-2.4b, even though the original compositions of the aluminum species in these coagulants were quite different (Figure 4, the analytical pH condition was approximately 4–5, these are the same data as those shown in Table 2). Because further transformation of aluminum species after coagulant dosing is known to occur, and the distribution of aluminum species depends largely on the original composition of the coagulants and the pH of the treated water [4, 25, 26], we investigated the effect of pH on the transformation of the aluminum species after coagulant dosing across a pH range of 6–8 (Figure 4). The distribution of the aluminum species of the four PACls during coagulation tended to be maintained (i.e., the original composition of the aluminum species was retained): the Al₆ fraction, which represented a large proportion of PACl-2.1b and PACl-2.4b, and the Al₅ fraction, which represented a large proportion of PACl-2.1c and PACl-2.4c, were almost unchanged between the pH
ranges of 4–5 and 6–8 (Figure 4), although the Alc fraction, which represented a small proportion of PACl-2.1b and PACl-2.4b, was smaller when the pH was increased from 4–5 to 6. This result indicates that most of the original Alb and Alc content of the PACls was relatively stable over the pH range of 4–8, in agreement with previous studies [4, 26]. In contrast, the distribution of the aluminum species of AlCl3 changed markedly during coagulation compared with the original composition: the Ala fraction, which represented a large proportion of AlCl3 decreased and the Alb fraction increased when the pH was increased from 4–5 to 6–8 (Figure 4). This result indicates that a large proportion of the original Ala content of AlCl3 transformed into Alb during coagulation, and that the formation of Alb in situ probably led to the similar removal ratios of PV that were observed with PACl-2.4b.

The removal ratios of PV observed with high Alc content PACls tended to be larger than those observed with high Alb content PACls (Figures 2 and 3). Because PACl-2.1c and PACl-2.4c contained not only Al13 species but also Al30 species, which was not the case for PACl-2.1b and PACl-2.4b, we hypothesize that the Al30 species in PACl played a major role in PV removal during the coagulation process. These results, taken together with our previous findings [16], suggest that Al30 species in aluminum-based coagulants are the important species to achieve efficient removal of not only bacteriophages but also human enteric viruses during the coagulation process.

When the basicity was increased from 2.1 to 2.4, the Alc content in the high Alc content PACls increased slightly and the Ala content in those PACls decreased slightly (Figure 4, the analytical pH condition was approximately 4–5). A similar trend was observed for the distribution of the aluminum
species of PACls during coagulation (Figure 4, the analytical pH condition was approximately 6–8). However, the removal ratios of PV observed with PACl-2.4c tended to be lower than those with PACl-2.1c (Figures 2 and 3). This result suggests that the coagulation efficiency of virus during the coagulation process is not determined simply by the amount of Al₃ in the coagulant.

To elucidate why PACl-2.1c effectively removed PV, we used the colloid titration method to determine the positive colloid charges of the coagulants (Figure 5). The colloid charge densities of AlCl₃ were almost zero and constant, regardless of the aluminum concentration. In contrast, the colloid charge densities of the four PACls increased with increasing aluminum concentration. However, we observed no large differences in the colloid charge densities of the PACls, whereas the removal ratios observed with PACl-2.1c were somewhat larger than those observed with the other aluminum-based coagulants, as described above. Therefore, the virus coagulation efficiency was not solely dependent on either the amount of Al₃ in the coagulant or the positive colloid charge of the coagulant. Ye et al. [12] recently reported that the amount of Al₃₀ species in the coagulant increases with increasing basicity from 1.0 to 2.0, but decreases when the basicity increases from 2.0 to 2.5. Therefore, the difference in the amount of Al₃₀ species between PACl-2.1c and PACl-2.4c due to the difference in basicity probably led to the difference in virus removal performance during coagulation. Further investigation is needed to determine why the removal ratios of PV observed with PACl-2.4c tended to be lower than those with PACl-2.1c, even though the Al₃ in PACl-2.4c was slightly higher than that of PACl-2.1c.
3.3. Comparison of PV and CV removal ratios during coagulation

As described above, PACl-2.1c removed PV more efficiently than did the other aluminum-based coagulants tested, especially at approximately pH 7. To confirm that PACl-2.1c actually removed viruses more effectively than did AlCl₃ or PACl-2.1b, we also evaluated the removal ratios of a CV that has high resistance to free-chlorine disinfection relative to other human enteric viruses, including PV [21, 22]. We evaluated the CV removal ratios by using the PFU and PCR methods after settling during the coagulation process, and then compared the results with those for PV (Figure 6). For both enteroviruses, the removal ratios observed with PACl-2.1c were larger than those observed with AlCl₃ and PACl-2.1b at about pH 7: approximately 3-log₁₀ removals were achieved with PACl-2.1c. This result indicates that coagulation with PACl-2.1c more effectively removes human enteroviruses, including a virus with high resistance to free-chlorine disinfection, relative to coagulation with AlCl₃ and PACl-2.1b.

The removal ratios of infectious PV and CV as evaluated by use of the PFU method were almost identical during coagulation regardless of the coagulant type used (Figure 6). A similar trend was observed for the removal ratios of total PV and CV, as evaluated by using the PCR method. In contrast, Mayer et al. [20] reported that the removal ratio of CV B6, evaluated by using the 50% tissue culture infectious dose (TCID₅₀) method, was consistently larger than that of PV type 1 evaluated by the same method during enhanced coagulation with FeCl₃: for example, the optimal enhanced
coagulation conditions of 40 mg/L FeCl$_3$ and a pH of between 5.5 and 6.5 resulted in 1.0–2.5-log$_{10}$ and 1.5–3.0-log$_{10}$ removals of infectious PV and CV, respectively. This result is not consistent with our result, although we did observe similar ranges of removal ratios for infectious PV and CV. In general, the removal ratios of microorganisms including viruses during coagulation are influenced by several factors such as source water quality, coagulant type, coagulant dosage, and coagulation pH [27]. Several of these factors, in particular coagulant type (aluminum-based coagulants vs. FeCl$_3$) and coagulation pH (7 vs. 5.5–6.5), were different between our study and that of Mayer et al., which may explain the differences in the removal ratios between the two studies. Nevertheless, our findings clearly show that there was no large difference between the PV and CV removal ratios when aluminum-based coagulants were used as the coagulant at about pH 7.

The removal ratios of PV and CV determined by using the PFU and PCR methods were almost the same during coagulation with AlCl$_3$ and PACl-2.1c; however, when PACl-2.1b was used as the coagulant, the removal ratios determined by using these two methods differed slightly (Figure 6). This result suggests that the PCR method, which allows for rapid, highly sensitive, and highly specific quantification of viruses, is an acceptable alternative to the PFU method, which is time-consuming and labor-intensive, for the evaluation of enterovirus removal by coagulation with aluminum-based coagulants. However, if the removal ratios determined by using these two methods showed a greater difference, cell culture–based infectivity assays, including the PFU method, would be required to evaluate the removal of infectious viruses in order to predict the human health risk. Ryu et al. [28]
reported that the removal ratios of AdV, CV, and EV evaluated by using a cell culture–based infectivity assay (i.e., the PFU or TCID\textsubscript{50} method) and by using the PCR method were almost identical during coagulation with FeCl\textsubscript{3}, which is in agreement with our result even though the coagulation conditions were quite different between the two studies.

3.4. Mechanisms of virus removal

Coprecipitation into growing aluminum hydroxide during charge neutralization and adsorption onto formed aluminum hydroxide flocs are two known mechanisms for removal of natural organic matter and As(V) during the coagulation processes with aluminum-based coagulants [4, 5, 29]. To investigate whether these mechanisms contribute to virus removal, and if so, whether differences exist in the contributions of these mechanisms for virus removal among AlCl\textsubscript{3}, PACl-2.1b, and PACl-2.1c, two types of batch coagulation experiments were conducted at approximately pH 7 (Figure 7).

Although the removal ratios of infectious PV, as evaluated by using the PFU method, during coagulation with AlCl\textsubscript{3}, PACl-2.1b, and PACl-2.1c were 1.6-log\textsubscript{10}, 2.7-log\textsubscript{10}, and 3.0-log\textsubscript{10}, respectively, the effectiveness of these coagulants to remove virus decreased to ≤0.5-log\textsubscript{10} when the PV was added to the beaker after charge neutralization (i.e., after rapid mixing) (Figure 7a). A similar trend was observed for the removal ratios of total PV evaluated by using the PCR method (Figure 7b). These results indicate that coprecipitation into growing aluminum hydroxide during charge neutralization is the main mechanism for virus removal by aluminum-based coagulants, and that the
difference in the charge neutralization capabilities of the coagulants, which we could not evaluate simply by the colloid titration method, probably led to the difference in virus removal efficiency during coagulation. In contrast, the removal ratios of infectious PV in the adsorption process were \leq 0.5-\log_{10} no matter what type of coagulant was used (Figure 7a). A similar trend was observed for the removal ratios of total PV evaluated by using the PCR method (Figure 7b). Several researchers have reported that the properties of aluminum floc particles generated during coagulation differ depending on the distribution of the aluminum species in the coagulants [1, 11, 30]. Our results suggest that there was no difference in the adsorption of PV onto the preformed aluminum floc particles produced by AlCl₃, PACl-2.1b, or PACl-2.1c. Therefore, virus adsorption on formed aluminum hydroxide flocs is not the main mechanism for virus removal by aluminum-based coagulants, although this mechanism made a partial contribution to virus removal during coagulation.

Here we have shown that PACl-2.1c, which contains Al₃₀ species, removed human enteroviruses, specifically PV and CV, more efficiently than did the other aluminum-based coagulants tested. Ye et al. [12] recently reported that Al₁₃ aggregates, which are precursors for Al₃₀ species, induced by an aging process promoted higher yields and faster production of Al₃₀ species, whereas Al₁₃ aggregates prepared by increasing the basicity hampered the production of Al₃₀ species. In the present study, PACl-2.1c was prepared by using the base titration method without any aging process. Therefore, further optimization of the preparation of PACls for the effective formation of Al₃₀ species should further improve virus removal performance during coagulation and lead to the development of novel
4. Conclusions

We found that differences in the distribution of aluminum hydrolyte species in the coagulants affected the removal of the enteroviruses during the coagulation process: PV and CV removal ratios were larger with high Alc content PACl (PACL-2.1c) than with high Ala content coagulant (AlCl₃ solution) and high Alb content PACl (PACL-2.1b). Unlike AlCl₃ or PACl-2.1b, PACl-2.1c contains Al₃₀ species, indicating that Al₃₀ species probably play a major role in enterovirus removal during the coagulation process. The PV and CV removal ratios were almost identical during the coagulation process, regardless of the coagulant type or virus quantification method used, suggesting that PV and CV behaved similarly during the coagulation process.

The main mechanism for virus removal during the coagulation process was coprecipitation into growing aluminum hydroxide during charge neutralization. Virus adsorption onto formed aluminum hydroxide flocs made a limited contribution to virus removal.

Acknowledgements

We thank Dr. Hiroyuki Shimizu (National Institute of Infectious Diseases) for providing the Sabin
attenuated LSc/2ab strain of PV type 1. We also thank Dr. Daisuke Sano and Ms. Rie Nomachi
(Hokkaido University) for providing buffalo green monkey kidney epithelial cells and for teaching
us how to cultivate and quantify infectious PV. We thank the staff of the drinking water treatment
plants for providing source water samples. This research was supported in part by a Grant-in-Aid for
Young Scientists A (no. 25709044, 2013), a Grant-in-Aid for Scientific Research S (no. 24226012,
2012), and a Grant-in-Aid for Scientific Research B (no. 15H04064, 2015) from the Japan Society
for the Promotion of Science; by a Health and Labor Sciences Research Grant (Research on Health
Security Control) from the Ministry of Health, Labor and Welfare of Japan; and by a grant from the
Kurita Water and Environment Foundation (no. 14A007, 2014).

References

  species (Alₐ, Alₐ, and Alₐ) during coagulation with polyaluminum chloride: a case study with the

  Coagulation characteristics of polyaluminum chlorides PAC-Al₃0 on humic acid removal from water,


coagulation efficiency, residual Al and floc properties in surface water treatment, Colloid Surface A 459 (2014) 14-21.


Table 1. Water quality data for the source water samples.

<table>
<thead>
<tr>
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<th>Lake water</th>
<th>River water</th>
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<td>pH</td>
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<tr>
<td>Turbidity (NTU)</td>
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<td>DOC (mg/L)</td>
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<td>UV260 (cm⁻¹)</td>
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<td>Alkalinity (mg-CaCO₃/L)</td>
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Table 2. Characteristics of the aluminum-based coagulants used in the present study.

<table>
<thead>
<tr>
<th>Coagulants</th>
<th>Basicity</th>
<th>Aluminum concentration (μg-Al/L)</th>
<th>Relative density at 20 ºC</th>
<th>Aluminum species distribution</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Al⁺⁺⁺ (%)</td>
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<tr>
<td>AK⁺⁺⁺b</td>
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<tr>
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<tr>
<td>PACl-2.4c</td>
<td>2.4</td>
<td>10.3</td>
<td>1.1</td>
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</table>
Figure 1. $^{27}$Al NMR spectra of the coagulants used in the present study.
Figure 2. Effect of aluminum hydrolyte species in coagulants on (a) infectious PV removal as evaluated by using the PFU method and on (b) total PV removal as evaluated by using the PCR method after settling during the coagulation process. Source water, lake water; coagulant dosage, 2.70 mg-Al/L.
Figure 3. Effect of aluminum hydrolyte species in coagulants on (a) infectious PV removal as evaluated by using the PFU method and on (b) total PV removal as evaluated by using the PCR method after settling during the coagulation process. Source water, river water; coagulant dosage, 1.08 mg-Al/L.
Figure 4. Effect of pH on the transformation of aluminum species after coagulant dosing, as evaluated by using the ferron method.
Figure 5. Positive colloid charges of the coagulants used in the present study, as evaluated by using the colloid titration method.
Figure 6. Comparison of removal ratios of PV (white) and CV (gray) after settling during the coagulation process at approximately pH 7. Source water, lake water; coagulant dosage, 2.70 mg-Al/L. Values were determined from one experiment or are the means of duplicate experiments; error bars indicate the data range.
Figure 7. Effects of charge neutralization and adsorption on preformed aluminum floc particles on (a) infectious PV removal as evaluated by using the PFU method and on (b) total PV removal as evaluated by using the PCR method during the coagulation process at approximately pH 7. Source water, lake water; coagulant dosage, 2.70 mg-Al/L. Values were determined from one experiment or are the means of duplicate experiments; error bars indicate the data range.