Removal of Se(IV) and Se(VI) from drinking water by coagulation

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Selenium (Se) is one of contaminants required to be regulated during drinking water treatment, however, little information has been collected to date regarding Se removal by coagulation. In this study, the performance of Se removal by coagulation has been evaluated with respect to the dependence on Se species, coagulant type, water pH and interfering ions. The results showed that a Fe-based coagulant was much more efficient than Al-based coagulants in Se removal. The removal of selenite (Se(IV)) by coagulation was much more pronounced than that of selenate (Se(VI)). With an FC dosage of more than 0.4 mM Fe/L, Se(IV) removal efficiency of more than 98% could be achieved when the initial Se(IV) concentration was 250 µg/L. For Al-based coagulants (AlCl3 (AC) and polyaluminum chloride (PACl)) Se removal efficiency was positively correlated with the content of Al13 species during the coagulation process. Adsorption onto hydroxide flocs was the most active coagulation mechanism for Se removal and precipitation also played specific roles at low dosage, especially for Se(IV) removal and with Fe coagulant. High coagulant dosage and weakly acidic pH could enhance the formation of hydroxide flocs having more active adsorption sites and high zeta potential, and thus favored Se removal. These findings are important to understand the efficiency and mechanisms of Se removal by coagulation.

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1. Introduction

Selenium (Se) is an essential trace element for both animals and humans, but it can be toxic if taken in excess [1]. Excessive amounts of Se can cause lower reproduction rate and fetal deformity [2,3]. Its toxicity led the World Health Organization (WHO) to recommend a maximum Se concentration in drinking water of 10 µg/L. Se is introduced in the environment from both natural and anthropogenic sources. Agriculture and combustion of fossil fuels are the primary sources of Se from human activities [4]. It is reported that the surface and ground water in some areas of China, India and America contain much higher levels of Se (45–341 µg/L) than required by the drinking water standard [4–6]. In the environment, Se can exist in different oxidation states: elemental selenium (Se0), selenite (SeO3^2^-), selenide (Se^-2), selenate (SeO4^2-) and organic Se [7]. Se(IV) and Se(VI) are found in most aqueous media and are the predominant chemical forms [1]. The toxicity of Se depends on its oxidation state. Se(IV) is considered to be more toxic than Se(VI) [8].

A variety of treatment methods have been studied for Se removal from contaminated waters, including ion-exchange [9], reverse osmosis [10], zero-valent iron [11], microbial reduction [12], chemical precipitation [13], a hybrid electro-coagulation membrane process [14] and adsorption [15–23]. Among these, adsorption is believed to be the most promising process and has been studied extensively. Metal oxides such as various iron (Fe) oxides and aluminum (Al) oxide have been demonstrated to be effective adsorbents for Se removal [15,18,20,22–24]. Coagulation by Al and Fe salts is an important water treatment process and has been widely used in drinking water plants. For Al salt that is widely used as coagulant, it has been believed that the Al species distribution during coagulation can determine the coagulation behavior and efficiency [25,26]. After coagulant addition, Al or Fe cations hydrolyze quickly and in-situ form abundant precipitates of Al or Fe hydroxide oxides. Contaminant adsorption to Al or Fe hydroxide oxide can play a very important role in the coagulation process [27,28]. Therefore, coagulation may be an effective treatment to remove Se through adsorption by Al or Fe hydroxide oxide. Coagulation is an existing water treatment process in drinking water plants. However, so far no study has been conducted on the removal of Se from drinking water by coagulation.

The objective of this paper is to evaluate the coagulation performance in Se removal. The efficiency of Se(IV) and Se(VI) removal by Al and Fe salts as coagulants was examined using jar tests. The effect of water pH and coexisting anions on Se removal by coagulation was clarified. The effect of Al speciation on Se removal was also investigated from the viewpoint of Al species transformation.

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The present study also provided insight into the mechanisms of Se removal by Al- and Fe-based coagulants. We hope this work contributes to effective Se removal during coagulation process and makes drinking water supply more safely.

2. Materials and methods

2.1. Materials

All of the chemical solutions used in this study were prepared using de-ionized water and analytical grade chemicals. FeCl₃ (FC) was used as the Fe-based coagulant. AlCl₃ (AC) and polyaluminum chloride (PACl) were used as a traditional Al-based coagulant and inorganic Al polymer flocculant, respectively. PACl was prepared by an electrochemical process [29]. The predominant Al species of AC and PACl were Al monomer (Al₃⁺) and Al₁₁₃⁺ polymer, respectively. Al₁₁₃⁺ polymer in PACl used in this study accounted for 87.1% of the total Al concentration, which was analyzed using the method of ²⁷Al nuclear magnetic resonance [30]. 1000 mg/L Se(IV) and Se(VI) stock solutions were prepared by dissolving Na₂SeO₃ and Na₂SeO₄·10H₂O (both from Aldrich) into de-ionized water, respectively. Water samples were synthesized by spiking Se(IV) or Se(VI) into tap water for coagulation and adsorption experiments. The initial Se concentration was 250 μg/L for all experiments.

2.2. Determination of Se concentration

Se concentration was analyzed by a spectrometer (AF-610, Beijing Rull Analytical Instrument Co., Ltd., China) on the basis of hydride generation–atomic fluorescence spectrometry [31]. The operation conditions were as follows: PMT-voltage of 220 mV, lamp current of 50 mA, argon carrier gas flow rate of 800 mL/min, and atomization at room temperature. All samples used in our analysis were analyzed within 24 h of collection. Total Se (Se(IV) and Se(VI)) concentration analysis was performed as follows: 10 mL of filtered supernatant was transferred into a 15 mL test tube, then 1.5 mL HCl (6 mol/L) was added. The solution was heated to 90 °C for 30 min, then was diluted to 15 mL with 1.2 mol/L HCl for final analysis. HCl (1.2 mol/L) and potassium borohydride (15 g/L, containing 2 g/L NaOH) were used as carrying fluid.

2.3. Jar test

Jar tests were performed using a six-paddle stirrer. The procedures consisted of a 2 min rapid mix (200 rpm), 30 min slow mix (40 rpm), and a 30 min settling period, after which the supernatants were sampled and filtered by a 0.45 μm pore size membrane filter. The filtrates were tested for Se concentration. A small amount of sample was taken immediately to measure Zeta potential (Zetasizer 2000, Malvern, UK) during the slow mix period. Before the coagulant dose a predetermined amount of 0.2 mol/L NaOH or 0.05 mol/L HCl solution was added into water samples to approximately get an expected pH value. After dosing with coagulants, water pH was accurately adjusted to the expected value during the rapid mix period by adding HCl or NaOH solution, after which the water pH was constant during the subsequent flocculation process.

2.4. Adsorption experiments

Adsorption experiments were initiated to evaluate Se(IV) and Se(VI) removal by adsorption on Al or Fe hydroxide flocs. In the water samples without Se, hydroxide flocs preformed during the slow mix periods. After this, a certain amount of Se was added into the preformed floc suspension and then the slow mixing (40 rpm) was maintained for 30 min to allow the Se to be adsorbed. After the adsorption step, the settling step was applied. The experimental conditions for adsorption were same as those for coagulation. The Se removal performance by adsorption was compared with that by coagulation.

2.5. Effect of competing anions on Se removal

To test the effects of co-existing anions on Se(IV) removal by coagulation, four types of oxyanions (SO₄²⁻, CO₃²⁻, SiO₃²⁻ and PO₄³⁻) of similar molecular structures were evaluated individually. The experiments were conducted at pH 6.0. To avoid the interference of matrix constituents in tap water, the experiments were carried out with de-ionized water containing 0.5 mmol/L NaNO₃. Water samples were synthesized by spiking co-existing anions into de-ionized water for the experiments of co-existing anions effect. The concentrations of the four oxyanions were controlled at three levels (0.1, 1 and 10 mM).

3. Results and discussion

3.1. Se removal by Al- and Fe-based coagulants

Fig. 1 shows the results of Se(IV) and Se(VI) removal by FC, AC and PACl. With the increase of coagulant dosage, Se(IV) and Se(VI) concentration gradually decreased, except at the high dosages of AC. FC was more effective for Se removal than the Al-based coagulants. The removal efficiency of Se(IV) by FC could reach above 98% at dosages of more than 0.4 mmol Fe/L, while the maximum removal efficiency of Al-based coagulants was about 80% at the dosage of 1.2 mmol Al/L. The removal efficiency of Se(VI) by FC was also notably higher than that by the two Al salts at the same coagulant dosages. Fig. 1 shows that the removal of Se(IV) by coagulation was much more effective than that of Se(VI). For example, at the dosage of 0.4 mmol/L Fe or Al, the removal efficiencies of Se(IV) were twice as high as those of Se(VI). Se(IV) concentrations...
in the product water after treatment by more than 0.4 mmol Fe/L. FC could meet the required maximum contaminant level (10 μg Se/L).

Both AC and PACl could remove some Se, but the coagulation behaviors of AC and PACl were different. Fig. 1 shows that the performance of PACl in Se removal was more stable than that of AC. The Se removal efficiency of AC at the middle dosage (0.4 mmol/L) was higher than that of PACl, while lower at the low (0.05–0.2 mmol/L) and high (more than 0.8 mmol/L) dosages. Fig. 2 shows that water pH was depressed by coagulant addition. The most significant decline of water pH was observed with FC coagulation. For Al-based coagulants, the extent of the drop was inverse to the coagulant basicity. From the lowest dosage to highest dosage, the pH value with AC addition decreased from 7.5 to 4.2, which was larger than the pH range with PACl. At the optimal dosage (about 0.6 mmol/L) of AC for Se removal, the final pH was weakly acidic (about pH 5), which has been proven to be the optimal pH condition for AC coagulation. It has been demonstrated that water pH plays an important role in Al species distribution during the coagulation process [25,32], which may contribute to the differences between AC and PACl in Se removal. The effect of pH on Se removal by coagulation was further investigated and discussed in the following text.

3.2. Effect of pH on Se removal

The effect of pH on Se removal was evaluated in the range of pH 2–9 for FC and 4–9 for the two Al-based coagulants, respectively. As shown in Figs. 3 and 4, the effect of pH on Se(IV) removal by coagulation was similar to that on Se(VI) removal. In the range of pH 4–9, the Se(IV) and Se(VI) removal efficiencies with 0.1 mmol Fe/L FC were clearly higher than those using the two Al-based coagulants at 0.4 mmol Al/L. Compared with Al-based coagulants, the Se removal by FC was less dependent on pH. The optimal pH for Se removal by FC was lower than for the Al-based coagulants. The optimal pH values for Se(IV) and Se(VI) removal by FC, AC and PACl were about 4, 5 and 6, respectively. Figs. 3 and 4 shows that the trend of Se removal as a function of pH resembled that of the zeta potential, which indicated that Se removal by coagulation positively correlated with the zeta potential of the hydroxide flocs. The optimal pH for Se removal was approximately consistent with the pH values for the maximal zeta potential. The result revealed that the surface potential of flocs might play an important role in coagulation to remove Se. The zeta potential of flocs was positive at weakly acidic conditions, where the Se(IV) and Se(VI) removal efficiency reached the maximal value. Since both Se(IV) and Se(VI) are anions in water, the high positive charge of flocs facilitated the Se removal during the coagulation process. Generally, Fe and Al hydroxide flocs readily and largely form in weakly acidic pH conditions due to the minimum solubility of Al and Fe in water in this pH region. Abundant hydroxide flocs may favor Se removal through coprecipitation into growing hydroxide and adsorption on formed hydroxide flocs, which are believed to be the main mechanisms for removal of anionic contaminants by the coagulation process. Hydrous Fe oxides are less soluble than hydrous Al oxides, thus producing more flocs for Se removal.
copolymers and adsorption during the coagulation process [33], which may contribute to the better performance of FC in Se removal.

Figs. 3 and 4 show that both Se(IV) and Se(VI) removals by AC were higher than those by PACl in the pH range 5–6.5, which was the optimal pH range for Se removal by Al-based coagulants. At acidic and alkaline pH conditions, except Se(IV) removal at pH 4, Se removal by AC was inferior to that by PACl. The optimal pH value (pH 5) for Se removal by AC was lower than that (pH 6) by PACl. Al monomer and Al\(_{13}\) polymer were the dominant Al species for AC and PACl, respectively. Al\(_{13}\) polymer has been believed to be the most active species in Al coagulants responsible for coagulation [25,26,34]. It has been demonstrated that Al\(_{13}\) polymer could largely form in-situ through the transformation of Al monomer in a weakly acidic pH environment, while preformed Al\(_{13}\) polymer was quite stable even in acidic and alkaline pH conditions [25,32]. Overall, PACl performance in Se removal was less dependent on pH than AC. This may be a result of the stable character of preformed Al\(_{13}\) polymer during the PACl coagulation process. By comparison, AC showed a relatively labile performance with pH change. The higher removal efficiency at middle dosage (Fig. 1) and weakly acidic pH condition (Figs. 3 and 4) may be ascribed to the function of in-situ formed Al\(_{13}\) polymer during the AC coagulation process. This indicated that for Al-based coagulants, Se removal efficiency was positively correlated with the content of Al\(_{13}\) species during the coagulation process.

3.3. The mechanism of Se removal by coagulation

Three mechanisms for the removal of anionic contaminants (e.g., P, As and Sb) by Al and Fe-based coagulants have been suggested [35–37]: (i) Precipitation and formation of insoluble compounds; (ii) Coprecipitation into growing hydroxide; (iii) Adsorption onto formed hydroxide flocs. In the present study, special experiments were performed to discriminate between these mechanisms and determine which one is predominant, and the differences between PACl, AC and FC during Se(IV) and Se(VI) coagulation removal. The results of Se coagulation by AC and PACl and adsorption by the corresponding hydroxide flocs at pH 6 are shown together in Fig. 5. Similar trends were observed in adsorption and coagulation processes with AC and PACl. The Se removal efficiency of both coagulation and adsorption increased with the increase of coagulant dose. Although Se removal by coagulation was more efficient than that by hydroxide floc adsorption, the gap in Se removal efficiency between adsorption and coagulation was small. The gap in Se(IV) removal was larger than that of Se(VI). The difference was more pronounced at low dosages than high dosages. Se(IV) removal by adsorption and coagulation both for AC and PACl were almost the same at high dosages, where the removal efficiency was close to 100% (Fig. 5a). Se removal by FC coagulation and FC floc adsorption was investigated at pH 6 and 4, respectively. As shown in Fig. 6, similar trends were also observed in adsorption and coagulation processes with FC. The gap in Se(IV) removal efficiency between FC coagulation and FC floc adsorption, except at the high dosages, was significantly larger than those of Al-based coagulants and that of Se(VI). In addition, the difference was more pronounced at pH 6 than at pH 4. The results from these experiments indicated that adsorption onto hydroxide precipitations was the most important mechanism for Se removal by Al and Fe coagulants and precipitation/coprecipitation also played specific roles at low dosage, especially for Se(IV) removal and with Fe coagulant. Adsorption onto hydroxide flocs played a more important role in Se(IV) removal, and it was almost the sole coagulation mechanism when Se(IV) removal efficiency reached more than 95%. Previous studies [36,38,39] obtained similar results when they studied the coagulation behavior of Al- or Fe-based coagulants in the removal
of anionic contaminants. Since fewer hydroxide flocs occur at low coagulant dosages, coprecipitation of Se(IV) into growing hydroxide flocs via inclusion or occlusion seems to be difficult to occur as the predominant mechanism. Therefore, a precipitation mechanism, with formation of insoluble $\text{Al}_3(\text{SeO}_3)_2$ or $\text{Fe}_3(\text{SeO}_3)_2$ [40], may play an important role in Se(IV) coagulation at low dosage.

Fig. 5a shows that Se(IV) removal by AC coagulation was higher than that by PACI coagulation, but Se(IV) removal by AC floc adsorption was lower than that by PACI floc adsorption. Fig. 5b shows that Se(VI) removal by both coagulation and adsorption on PACI was higher than those for AC. These results indicated that PACI flocs had stronger adsorption affinity for Se than AC flocs did. As shown in Figs. 3b and 4b, the surface potential of PACI flocs at pH 6 was higher than that of AC flocs, which might lead to a greater attractive force between PACI flocs and Se ions. Adsorption onto formed hydroxide flocs was more active during PACI coagulation to remove Se. Precipitation was relatively more active during AC coagulation to remove Se. The predominant Al species of AC is Al monomer, while Al$_{13}$ polymer is the dominant Al species in PACI. The monomer may more easily complex with Se ions compared with Al$_{13}$ polymer since the binding sites of Al in Al$_{13}$ polymer are occupied by hydroxyl.

During adsorption onto preformed flocs, Se removal depended on the coagulant dose and consequently on the number of active sites on the hydroxide flocs’ surfaces. As mentioned above, FC could produce more flocs due to the lower solubility of hydrous Fe oxides. Moreover, it is believed that the structure of Fe hydroxide flocs is more open and loose than that of Al hydroxide flocs, ensuring a large available surface area for adsorption [41]. Consequently, Fe hydroxide flocs had a greater affinity toward Se(IV) and Se(VI) than Al hydroxide flocs, thereby bringing about higher Se removal efficiency for FC coagulation. Figs. 3 and 4 show that the zeta potential of FC hydroxide flocs at pH 4 was higher than that at pH 6, thus Fe hydroxide flocs generated at pH 4 may have a greater affinity toward Se(IV) and Se(VI) than those at pH 6. This may contribute to the higher Se removal at pH 4 during the FC coagulation process.

3.4. Effect of co-existing anions on Se removal

At a fixed pH of 6, the effects of four common oxyanions at three concentration levels (0.1, 1.0 and 10 mM) on Se(IV) removal are illustrated in Fig. 7. In the presence of SO$_4^{2-}$, CO$_3^{2-}$, SiO$_3^{2-}$ and PO$_4^{3-}$, the Se(IV) removal decreased to varying degrees. The effect of co-existing anions on Se(IV) removal by AC was more pronounced than that by FC. PO$_4^{3-}$ caused the greatest percentage decrease in Se(IV) removal among the anions at each concentration level. Although sulfur and Se are located in the same main group of the periodic table, the influence of SO$_3^{2-}$ on Se removal was lowest among the four oxyanions. The negative influence on Se(IV) removal among the four oxyanions followed the order: PO$_4^{3-}$ > SiO$_3^{2-}$ > CO$_3^{2-}$ > SO$_3^{2-}$, which is similar to the result of adsorptive Se(IV) removal using iron-coated GAC adsorbents [17]. Since adsorption onto hydroxide flocs is the main coagulation mechanism for Se(IV) removal by Al- and Fe-based coagulants, the results can be explained by the triple layer model, which is widely used to describe solid–liquid interface adsorption [42]. PO$_4^{3-}$ can bind strongly with the surface sites of metal hydroxides, forming inner-sphere (o-plane) surface complexes that are barely affected by ionic strength. In contrast, sulfate is weakly bound to the surface sites. The complex formation usually takes place on outer-sphere (o-plane) and is significantly affected by ionic strength. This can explain why PO$_4^{3-}$ has a relatively larger negative effect than SO$_3^{2-}$. For SiO$_3^{2-}$ and CO$_3^{2-}$, the competition between the two anions and Se(IV) may involve another reaction mechanism, surface accumulation/precipitation [16,43].

In addition, the competing anions may have accumulated or precipitated on the metal hydroxide surface, which in turn reduced the surface potential of the hydroxide flocs. The resulting electrostatic repulsion between Se(IV) and the hydroxide surface sites increased, which reduced Se(IV) adsorption and thus the Se(IV) removal by coagulation decreased. As mentioned above, compared with Al-based coagulants, the hydroxide flocs of FC have more surface adsorption sites for Se(IV) removal. Consequently, the negative interference of co-existing anions on Se(IV) removal by FC was less than that by AC.

4. Conclusions

The principal conclusions of this study are as follows:

1. The Fe-based coagulant was much more efficient than the Al-based coagulants. Compared with Se(VI), Se(IV) could be more easily removed by coagulation. With an FC dosage of more than 0.4 mM Fe/L, Se(IV) removal efficiency of more than 98% could be achieved when the initial Se(IV) concentration was 250 mg/L. Our results indicated that coagulation with iron salt is an effective treatment technique for Se(IV) removal.

2. Adsorption onto hydroxide flocs was the most active coagulation mechanism for Se removal and precipitation also played specific roles at low dosage, especially for Se(IV) removal and with the Fe coagulant. The surface character of hydroxide flocs, especially the zeta potential and active adsorption sites, determined the coagulation performance in Se removal. For Al-based coagulants, Se removal efficiency was positively correlated with the content of Al$_{13}$ during the coagulation process.

3. High coagulant dosage and weakly acidic pH conditions favored Se removal. AC performed better than PACI for Se removal under weakly acidic pH conditions. Compared to the performance of Al-based coagulants, Se removal by FC was relatively independent on water pH and insensitive to the presence of oxyanions. The negative influence on Se(IV) removal among the four oxyanions tested followed the order: PO$_4^{3-}$ > SiO$_3^{2-}$ > CO$_3^{2-}$ > SO$_3^{2-}$.
Pre-reduction of Se(VI) to Se(IV) seems to be necessary to achieve effective Se removal. In addition, some coagulant aids, chitosan for example, may have the potential to enhance Se(VI) removal by coagulation. Whether the pre-reduction process or the coagulant aids are used, their efficiency in enhancing Se(IV) removal by coagulation needs to be evaluated.

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