

SCIENTIFIC REPORTS



OPEN

Molecular Structures of Al/Si and Fe/Si Coprecipitates and the Implication for Selenite Removal

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Received: 25 October 2015

Accepted: 29 March 2016

Published: 20 April 2016

Aluminum and iron oxides have been often used in the coagulation processes during water purification due to their unique surface properties toward anions. In the presence of silica, the coprecipitation of Al/Si or Fe/Si might decrease the efficiency of wastewater purification and reuse. In this study, surface properties and molecular structures of Al/Si and Fe/Si coprecipitates were characterized using spectroscopic techniques. Also, the selenite removal efficiency of Al/Si and Fe/Si coprecipitates in relation to their surface and structural properties was investigated. While dissolved silicate increased with increasing pH from Fe/Si coprecipitates, less than 7% of silicate was discernible from Al/Si samples over the range from acidic to alkaline conditions. Our spectroscopic results showed that the associations between Al and Si were relatively stronger than that between Fe and Si in coprecipitates. In Al/Si coprecipitates, core-shell structures were developed with $\text{AlO}_6/\text{AlO}_4$ domains as the shells and Si frameworks polymerized from the SiO_2 as the cores. However, Si framework remained relatively unchanged upon coprecipitation with Fe hydroxides in Fe/Si samples. The Si core with Al shell structure of Al/Si coprecipitates shielded the negative charges from SiO_2 and thereby resulted in a higher adsorption capacity of selenite than Fe/Si coprecipitates.

Silica (SiO_2) has been widely found in the effluent discharged from the chemical-mechanical-planarization (CMP) process of semiconductor manufacturing, and it showed substantial stability in wastewater since it has a very low point of zero charge, suggesting that it could be the hardest one to remove in conventional wastewater treatment plants by aggregation-sedimentation^{1,2}. The residual SiO_2 in wastewater might decrease the efficiency of wastewater reuse as it would clog water pipes, decrease thermal conductivity of boilers, and especially block the reverse osmosis membranes³. As a conventional method, SiO_2 is often removed from wastewater by coagulating/flocculating (i.e., adsorption and/or coprecipitation) with aluminum (Al) or iron (Fe) salts^{4,5}. Due to the nano-scale particle size and the highly negative surface charges, however, the stability of Al/Si and Fe/Si coagulation is susceptible to change in various polymerization conditions⁵⁻⁸.

The stability of Al/Si and Fe/Si coagulation is in relation to the surface and structural properties of coprecipitates, which would vary with various environmental factors such as temperature, concentration of Si and Fe/Al, pH, and aging time⁸⁻¹². While coprecipitated with Fe hydroxides, for example, SiO_2 would decrease the surface charges of Fe/Si coprecipitates with increasing pH and aging time¹³. However, the surface charges of Al/Si coprecipitates tended to increase with increasing initial concentration of Al hydroxides¹². The promoted surface charges were derived from Al that hydrolyzed and then adsorbed or precipitated onto SiO_2 ⁹. The significant interaction between Al hydroxides and SiO_2 differentiated the sorption behaviors of Al/Si coprecipitates for nutrients, trace and heavy metals like Cd^{2+} and Ca^{2+} from that of pure Al hydroxides¹⁴. For Fe/Si coprecipitates, however, the Cd^{2+} removal efficiency was indistinguishable from that of pure Fe hydroxides^{14,15}. Collectively, the coagulation between Al/Fe with silica might substantially change the surface attributes of coprecipitates and further alter the mechanisms of metal and nutrient retention on Al and Fe hydroxides. However, the mechanistic mechanisms

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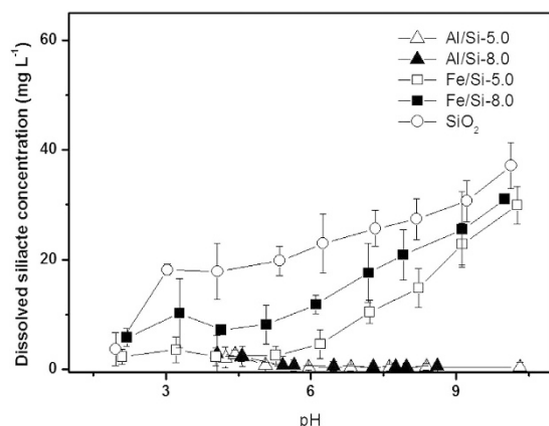


Figure 1. Trends of silicate dissolution from Al/Si and Fe/Si coprecipitates synthesized at pH 5.0 (Al/Si-5.0, Fe/Si-5.0) and 8.0 (Al/Si-8.0, Fe/Si-8.0) under an electrolyte concentration of 0.01 M NaNO₃.

at the molecular scale about how silica affects structural attributes of Al/Si and Fe/Si coprecipitates and the subsequent sorption behaviors have been still unclear.

The interactions of Al(III) or Fe(III) with SiO₂ on coprecipitate surfaces have been determined qualitatively and quantitatively using surface models, including complexation/precipitation and triple-layer models^{13,15,16}. However, the related mechanistic mechanisms, especially at molecular scale, could only be determined using spectroscopic methods. For example, the interaction between Al(III) and SiO₂ was classified as adsorption, surface-enhanced precipitation, and aluminosilicate precipitation according to the results of nuclear magnetic resonance (NMR) spectroscopy^{17,18}. The Fourier-transform infrared spectrometer (FT-IR) was employed to determine the elemental binding between Fe and SiO₂, and the results showed a replacement of Si by Fe on the Fe/Si coprecipitates¹⁹.

In this study, we aimed to determine the stability Al/Si and Fe/Si coprecipitates in relation to their surface and structural characteristics and further to characterize how such structural changes affect selenite removal. The reason to choose selenite as the target sorbate is that selenite is a common ingredient coming with SiO₄ in wastewater discharged from the semiconductor manufacturing²⁰. In addition, selenium is of environmental interest on account of the limited extent between biologically metabolically required and toxic effect concentrations in many organisms²¹. Wherein, selenite may threaten the environmental ecosystems seriously due to its low toxic doses and bioavailability in water²². In this study, we performed several spectroscopic analyses including powder X-ray diffraction (PXRD), transmission electron microscope (TEM), FT-IR, NMR, X-ray absorption spectroscopy (XAS), and X-ray photoelectron spectroscopy (XPS) to determine the molecular structural attributes for the Al/Si and Fe/Si coprecipitates. Subsequently, pH-dependent and isotherm sorption experiments of selenite on Al/Si and Fe/Si coprecipitates were performed to determine the selenite removal efficiency among individual coprecipitates. The combination of the structural characterization and sorption results was used to identify factors that can influence the stability of coprecipitate structures and the magnitude of selenite removal.

Results

Silicate Dissolution from Al/Si and Fe/Si coprecipitates. Trends of silicate dissolution from Al/Si and Fe/Si coprecipitates synthesized at pH 5.0 and 8.0 in 0.01 M NaNO₃ (Al or Fe/Si-5.0 and -8.0) were shown in Fig. 1. In raw silica solids (SiO₂), the silicate dissolution tended to increase with increasing pH, agreeing with the previous report⁹. There was a significant amount of dissolved Al or Fe under acidic conditions. However, only less than 0.15% of total added Al in Al/Si samples and 0.08% of total added Fe in Fe/Si samples was found at pH > 5.1 and 3.3, respectively (Figure S1). Such results indicated SiO₂ was surrounded by AlO₆ or FeO₆. While SiO₂ coprecipitated with Fe hydroxides, the amounts of silicate dissolution was general alleviated among all tested pH, especially for the Fe/Si samples coprecipitated at pH 5.0 (Fe/Si-5.0). Noteworthy, the silicate dissolution was essentially inhibited in Al/Si coprecipitates. For example, while 74%, 60%, and 62% of silicate was dissolved from SiO₂, Fe/Si-5.0, and Fe/Si-8.0 at pH 10, no discernable silicate dissolution was found for Al/Si samples at alkaline condition. Furthermore, only 5 to 7% of silicate was dissolved from Al/Si-5.0 and Al/Si-8.0 at pH < 4.5.

Surface and structural properties of Al/Si and Fe/Si coprecipitates. BET surface area, PXRD, FE-SEM of Al/Si and Fe/Si coprecipitates. The BET surface area (S_{BET}) for Fe hydroxides (283 and 284 m²g⁻¹ for Fe-5.0 and Fe-8.0) are greater than that of Al hydroxides (54 and 158 m²g⁻¹ for Al-5.0 and Al-8.0), wherein the Al-5.0 sample showed the lowest S_{BET} (Table S1). While coprecipitated with silica, the S_{BET} for the Al/Si samples were increased substantially. Especially for the Al/Si-5.0 sample, its S_{BET} (155 m²g⁻¹) was almost three times greater than that of pure Al hydroxides. In Fe systems, however, the S_{BET} was slightly decreased while coprecipitated with SiO₂. Figure 2(a) showed the PXRD patterns for Al/Si and Fe/Si samples, which indicated that all samples were non- or poorly crystalline. For Al/Si and SiO₂ samples, the broad peak centered at 4.1 Å ($2\theta = 23^\circ$) demonstrated the presence of SiO₂ structures^{23,24}. However, no discernable peak was found in Fe/Si samples. While no discrete phase was found in Al/Si samples (Fig. 2c,d), there were significant precipitates (dark spots in

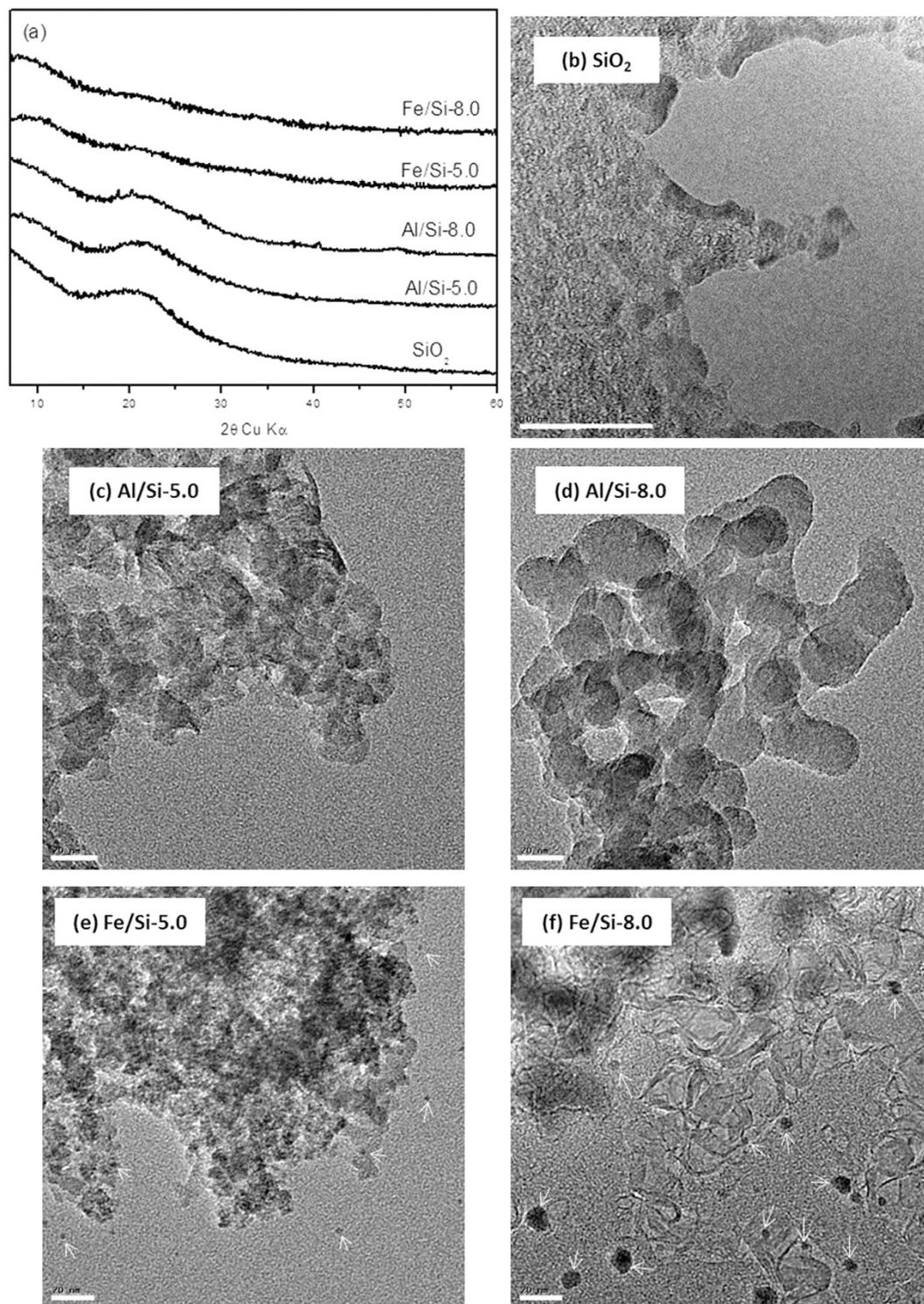


Figure 2. PXRD patterns and TEM images of raw SiO_2 and Al/Si, Fe/Si coprecipitates synthesized at pH 5.0 and pH 8.0 (Al/Si-5.0, -8.0, Fe/Si-5.0 and -8.0). The white bars represent 20-nm scales and the white arrows pointed out the plausible Fe precipitates.

Fig. 2e,f) in Fe/Si samples. The single phase in Al/Si sample implied a strong association between Al and SiO_2 surfaces. On the other hand, Fe precipitates and SiO_2 might form separate phases in Fe/Si systems.

Point of zero charge (PZC) of Al/Si and Fe/Si coprecipitates. Trends of zeta potential as a function of pH for pure hydroxides and that coprecipitated with silica are showed in Fig. 3. In general, the coprecipitation with silica tended to decrease the PZC for both Al and Fe hydroxides. While hydrolyzed at pH 5.0, for example, the PZC of Fe hydroxide was 7.4, and that was decreased to 6.4 for the Fe/Si-5.0 sample. Similar trend was also found in

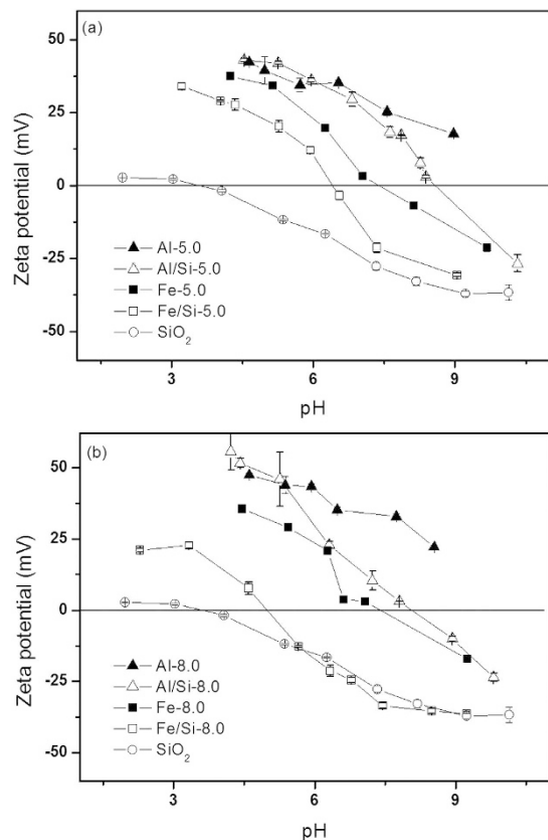


Figure 3. Results of zeta potential for SiO_2 , Al-, Fe-hydroxides, and Al/Si, Fe/Si coprecipitates synthesized at (a) pH 5.0 and (b) pH 8.0 under an electrolyte concentration of 0.01 M NaNO_3 .

the systems that hydrolyzed and coprecipitated at pH 8.0 (Fig. 3b). From the perspective of coprecipitation pH, however, the PZC for samples prepared at pH 8.0 were lower than that prepared at pH 5.0. For example, the PZC of Fe/Si-8.0 is 2.4 units less than that of Fe/Si-5.0.

FT-IR spectra of Al/Si and Fe/Si coprecipitates. The FT-IR signals at 1300–1000, 807, and 478 cm^{-1} was found in both pure silica and coprecipitated samples (Fig. 4a), which could be assigned to asymmetric Si–O–Si stretching of isolated SiO_4 units, symmetric stretching vibration of Si–O–Si network, and bending Si–O–Si/ O–Si–O, respectively^{25–31}.

As shown in Fig. 4b,c without stack increments in the y axis, the intensities at 1300–1000, 807, and 478 cm^{-1} of the Al/Si samples were generally greater than that of Fe/Si samples. Compared with Fe/Si samples, the shoulders at 907, 728, 629, or 554 cm^{-1} of the Al/Si coprecipitates intensified with increasing pH. Noteworthy, the intensity of Si-related signals (1300–1000, 807, and 478 cm^{-1}) for Al/Si samples, especially the Al/Si-8.0 one, was relatively greater than that of Fe/Si samples. In Al/Si samples, we also found the emerging shoulders at 907, 728, 629, or 554 cm^{-1} (Fig. 4b). The 907 cm^{-1} mode may derive from the Al–OH deformation vibration³². For the ridge at 728 cm^{-1} , it may be caused by stretching vibration of Si–O–Al with AlO_6 coordination³⁰ or the adsorbed tetrahedral AlO_4 ^{33–35}. For the shoulder of 629 and 554 cm^{-1} that was found in Al/Si-5.0 and Al/Si-8.0 respectively, the signals represented the symmetric stretching Si–O–Al³⁶ and/or the AlO_6 coordination^{35,37,38}. In fact that the intensity of an absorption band is related to the change of the dipole moment of the bond and the amount of the specific bond present, the greater intensity at such Si-related signals for Al/Si samples suggested more or stronger Si–O or Si–O–Si interaction. The changes in the quantity or intensity for such Si–O or Si–O–Si bonding might be resulted from the interaction between Si and Al as suggested by emerging shoulder showed in the Fig. 4b. While Al was demonstrated to be closely associated with silica in the coprecipitates, the interaction between Fe and Si in Fe/Si coprecipitates was not found due to the similar FT-IR spectra to that of SiO_2 and the absence of the significant Si–O–Fe vibration at 656 cm^{-1} (Fig. 4)³⁹.

Solid-state ^{27}Al and ^{29}Si MAS NMR spectra of Al/Si and Fe/Si coprecipitates. Solid-state ^{27}Al and ^{29}Si MAS NMR spectra for the Al/Si and Fe/Si coprecipitates were shown in Fig. 5. The ^{27}Al NMR spectra for the Al/Si coprecipitates synthesized at various pH (Fig. 5a) showed a discernible crest at 58 ppm and a significant peak around 5 ppm. The feature at around 58 and 5 ppm suggested tetrahedral (AlO_4) and octahedral (AlO_6) Al coordination environments, respectively^{40,41}. Regarding the peak at 58 ppm, the decreasing intensity with increasing pH indicated a fewer amount of AlO_4 at alkaline condition, agreed with what Houston *et al.* reported¹⁷, wherein the amount of AlO_4 tended to decrease and transform to the precipitated AlO_6 on the surface of silica as pH

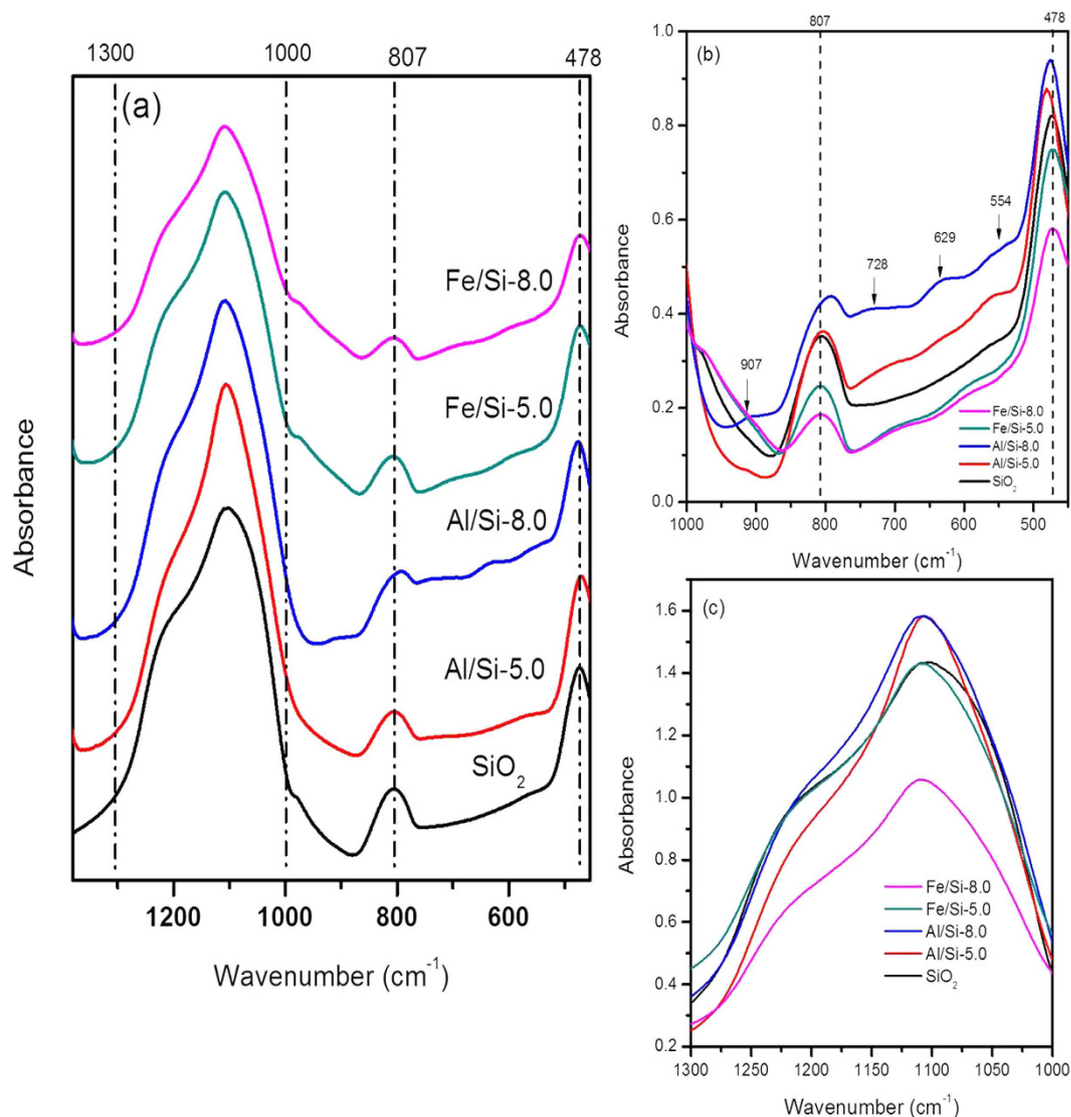


Figure 4. The FT-IR spectra ranged from (a) 450–1380 cm^{-1} and the inlet from (b) 450–1000 cm^{-1} as well as (c) 1000–1300 cm^{-1} for SiO_2 and Al/Si or Fe/Si coprecipitates synthesized at pH 5.0 or 8.0 under an electrolyte concentration of 0.01 M NaNO_3 , respectively.

increased. Such result was in line with the shift of the 5 ppm peak to the higher magnetic field with increasing pH, which suggested more bonding was formed between Al and Si at higher pH⁴².

The ^{29}Si MAS NMR spectra of Al/Si samples in Fig. 5b showed a broad peak at -109 ppm and a shoulder at around -101 ppm, which are contributed by the siloxanes with four bridging oxygen (Q_4) and silanol sites with three bridging oxygen (Q_3), respectively⁴². Compared with pure silica, the indiscernible signal at -101 ppm for Al/Si samples might indicate the polymerization of silicate, which transformed the silanol sites on silica to siloxane bonds⁴³. Besides the -109 ppm peak, there was a shoulder occurred between -101 to -109 ppm in Al/Si coprecipitates, and the position of such shoulder tended to shift to the low-field chemical shift (i.e., more negative shift) with increasing pH. Two possible explanations may account for the signal between -101 to -109 ppm in Al/Si samples: (1) the Q_3 polymerization to Q_4 coordination that caused a signal with chemical shift less than -101 ppm; and (2) the bond formation between Si and Al that shifted the Q_4 signal to the high-field chemical shift, as one Al bonding to Si would increase the ^{29}Si NMR signal by 5 ppm⁴². For Fe/Si coprecipitates, the ^{29}Si NMR spectra showed a main peak at -109 ppm with a minor signal at -101 ppm regardless of the changes at pH (Fig. 5c), indicating the predominance of Si polymerization⁴³. Given that the shoulders at -101 ppm in Fe/Si coprecipitates remained at similar intensity and position at various pH conditions, the absence of Fe/Si bonding in the coprecipitates was suggested⁴⁴.

Si K-edge XAS analysis of Al/Si and Fe/Si coprecipitates. Normalized Si and Fe K-edge X-ray absorption near-edge structure (XANES) data for the Al/Si or Fe/Si coprecipitates synthesized at various pH were shown in

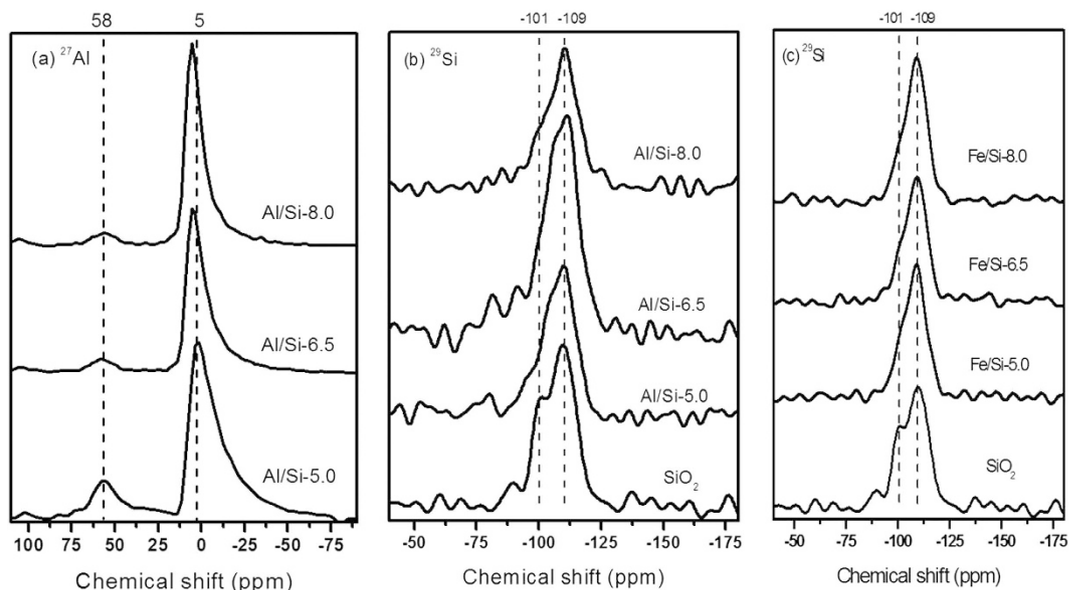


Figure 5. (a) ^{27}Al , (b) ^{29}Si MAS NMR spectra for Al/Si coprecipitates synthesized at pH 5.0, 6.5, and 8.0 (Al/Si-5.0, -6.5, and -8.0), and (c) ^{29}Si MAS NMR spectra for the Fe/Si coprecipitates synthesized at pH 5.0, 6.5, and 8.0 (Fe/Si-5.0, -6.5, and -8.0) under an electrolyte concentration of 0.01 M NaNO_3 , respectively.

Fig. 6. For the Si K-edge spectra, all samples showed a sharp white line (WL) centered at 1847.5 eV, wherein the WL intensity for both Al/Si and Fe/Si samples tended to increase with increasing coprecipitation pH (Fig. 6a,b).

While comparing the WL intensity between samples synthesized at the same pH, we found a higher WL peak for Al/Si-6.5 than that for Fe/Si-6.5. Similar trends were also observed between Al/Si-8.0 and Fe/Si-8.0 samples, although the WL intensity for the Al/Si-5.0 sample was lower than that of Fe/Si-5.0. Given that the WL peak for Si K-edge XANES is caused by the electron transition from the 1s to 3p orbitals⁴⁵, whose intensity is related to orbital hybridization⁴⁵ and/or the Si polymerization⁴⁶. Hence, such enhanced WL peaks in Al/Si samples, especially the one coprecipitated at alkaline conditions, suggested a higher degree of orbital hybridization between central Si and the surrounding Al and/or the enhanced polymerization of SiO_2 . In the matter of Fe/Si samples, however, we suspected the changes in WL intensity were mainly due to the silica polymerization as no significant difference in the Fe coordination environments was found in the Fe K-edge XANES (Fig. 6c) and extended X-ray absorption fine structure (EXAFS) (Fig. 6d) data for the Fe/Si samples.

Element distribution for Al/Si and Fe/Si coprecipitates. The proportions of Al or Fe relative to Si on near surfaces of Al/Si or Fe/Si coprecipitates detected by XPS analysis were tabulated in Table S2. The XPS analysis was used to determine the elemental distribution on near surfaces of coprecipitates rather than the elemental composition for coprecipitated particles. XPS is the most widely used surface analysis technique because it can be applied to a broad range of materials and provides valuable quantitative and chemical state information from the surface of the material being studied⁴⁷. The average depth of analysis for an XPS measurement is approximately 5 nm. On the near surfaces of Al/Si samples, the Al proportions increased from 3.9%–7.2% as pH increased from 5.0–8.0; however, the atomic ratio of Si remained at around 22% regardless of the changes at pH. Such trends resulted in the increasing Al/Si molar ratios from 0.17–0.34 with increasing pH. In contrast with Al/Si samples, the Fe proportion on Fe/Si samples showed insignificant variations as a function of pH, leading to a narrow range in Fe/Si molar ratios from 0.06–0.07. Although the element analysis of XPS was not highly accurate, the present modification in sensitivity factors of XPS rendered it as a reliable quantitative method^{48–51}. The merit of XPS in this study was that it demonstrated the trend that Al tended to precipitate on near surfaces of particles with increasing pH, but the Fe/Si ratios on near surfaces of Fe/Si coprecipitates stayed relatively consistent.

Selenite removal by Al/Si and Fe/Si coprecipitates. Given that the pH values for industry and domestic wastewater effluent are generally in the wide range between 1.5–8.5^{52,53}, it is worthy to examine the pH-dependent removal efficiency of selenite by Al/Si and Fe/Si coprecipitates. As shown in Fig. 7a, for both Al/Si and Fe/Si coprecipitates, the proportion of sorbed selenite decreased as suspension pH was artificially increased. With the exception of Al/Si-8.0 samples, more than 80% of selenite was fixed at pH 4. At pH > 8, however, less than 20% of selenite was fixed on Al/Fe-Si coprecipitates. This observed trend in decreasing sorbed Se with increasing sorption pH could be attributed to the fewer protonated surface sites on coprecipitates that serve as binding sites for selenite^{54,55}. Given that the Al/Si-5.0 and Fe/Si-5.0 showed relatively greater efficiency for selenite removal than that of Al/Si- and Fe/Si-8.0 among all tested pH, the samples coprecipitated at pH 5.0 were further used to conduct the isotherm experiments for selenite. As shown in Fig. 7b, the sorption capacity for Al/Si-5.0 (0.40 mmol g^{-1}) was 81% higher than that of Fe/Si-5.0 (0.22 mmol g^{-1}). As shown in Figure S2, the maximum adsorption capacities of selenite on pure $\text{Al}(\text{OH})_3$ and $\text{Fe}(\text{OH})_3$ are 0.41 and 0.28 mmol g^{-1} by Langmuir

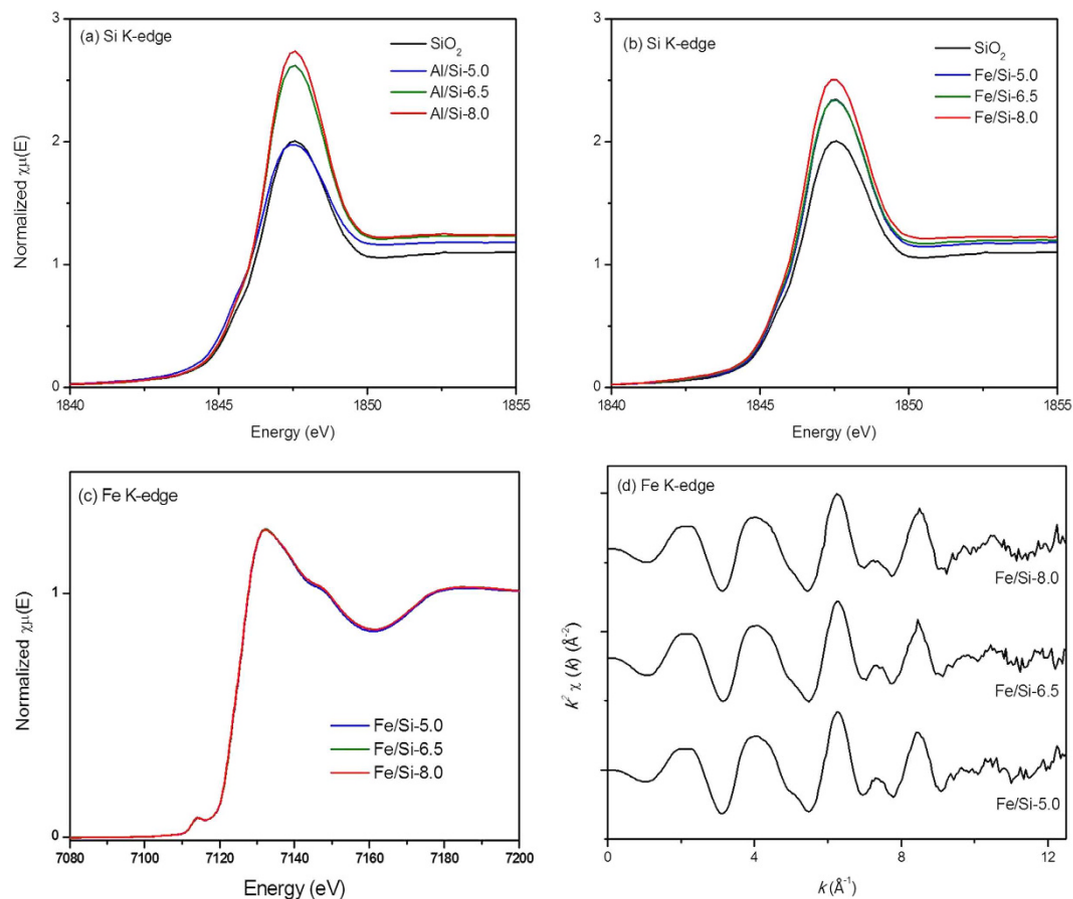


Figure 6. Normalized XANES spectra at Si K-edge for (a) SiO₂, Al/Si coprecipitates (Al/Si-5.0, -6.5, and -8.0), and (b) Fe/Si coprecipitates (Fe/Si-5.0, -6.5, and -8.0). The Fe K-edge XANES and k^2 -weighted EXAFS data for Fe/Si coprecipitates (Fe/Si-5.0, -6.5, and -8.0) were shown in (c,d).

isothermal model, slightly greater than Al/Si (0.40 mmol g⁻¹) and Fe/Si sample (0.22 mmol g⁻¹). Given that SiO₂ particles have almost no sorption capacity for selenite (data not shown), the association of SiO₂ on Al/Fe hydroxides might decrease the selenite sorption capacity for the Al/Fe-Si coprecipitates. Wherein silica might occupy active sorption sites on Al hydroxides by forming the core-shell structures in Al/Si coprecipitates. In Fe/Si samples, the dissolved silicate might compete for sorption sites on Fe hydroxides with selenite.

Discussion

Interactions between silica and Al or Fe controlled the surface and structural properties of coprecipitates that further affected the stability of coprecipitates and led to unique selenite sorption results. Compared with Fe/Si coprecipitates, silica in Al/Si coprecipitates seemed to be fixed more rigidly. There was less than 7% of silicate dissolved from Al/Si samples over the range from acidic to alkaline conditions (Fig. 1). That is, coprecipitation with Al hydroxides showed the promise to inhibit the silica reactivity with OH in the solution^{16,56}. The TEM images (Fig. 2b–f) demonstrated that the single phase in Al/Si sample implied a strong association between Al and SiO₂ surfaces. On the other hand, Fe precipitates and SiO₂ might form separate phases in Fe/Si systems. Based on the solubility product constant of 10⁻³⁹ for Fe(OH)₃ and 10⁻³⁴ for Al(OH)₃^{57,58}, Fe hydroxide domains plausibly formed faster than Al hydroxide domains under the same hydrolysis condition. For Al(III) that precipitated slower, the free Al³⁺ could directly bond with Si, resulting in the single phase in Al/Si systems. According to our spectral results (FT-IR, NMR, and XAS in Figs 4,5 and 6), such relatively more stable structure of Al/Si coprecipitates may derive from the bonding formation between Al and Si atoms.

For Al/Si samples, Al³⁺ rapidly hydrolyzed as suspension pH was raised, and the precipitated Al hydroxides may adhere on silica surfaces, wherein the silica served as the active heteronuclei in the coprecipitation processes⁵⁹. While precipitated on silica surfaces, the AlO₆ domains with a minor amount of AlO₄ units (Fig. 5a) may interact with the Si framework that polymerized from the SiO₂ (Figs 4 and 5b), which was evidenced by the stretching vibration of Si–O–Al in FT-IR spectra of Al/Si samples (Fig. 4). Given that the indiscernible interaction between Fe and Si was evidenced by spectral analyses, we proposed Fe hydroxides might form discrete phases with SiO₂ during coprecipitation. If any bonding between Fe and Si formed, a tetrahedral Fe coordination should be found, which would show the fingerprints of 7131.4, 7137.4, and 7148.2 eV at Fe K-edge XANES spectrum⁶⁰. For our Fe/Si samples, however, the Fe K-edge XANES and EXAFS data (Fig. 6c,d) suggested that the Fe structures were more like poorly crystalline Fe hydroxides⁶¹. The possible explanation for the discrete Fe phases in our

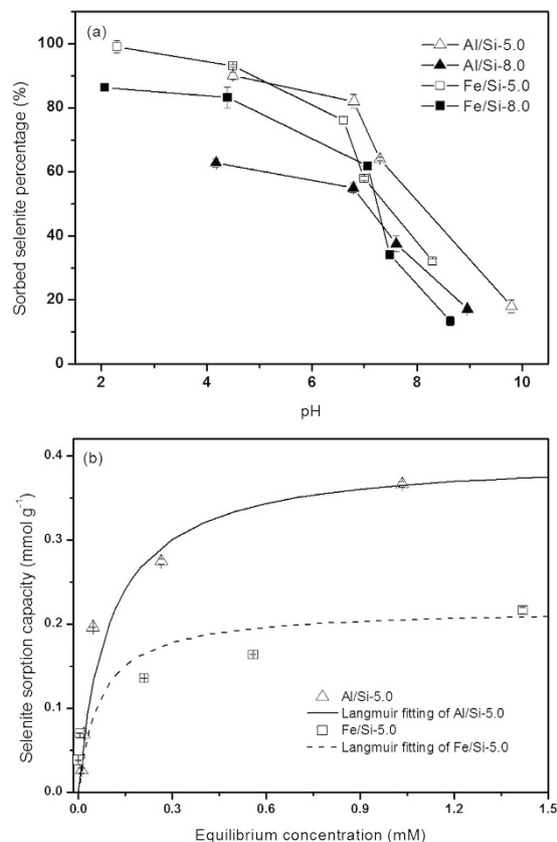


Figure 7. Results of (a) pH-dependent and (b) isotherm sorption of selenite on Al/Si and Fe/Si coprecipitates synthesized at pH 5.0 and 8.0 (Al/Si-5.0, -8.0 and Fe/Si-5.0, -8.0) at 25 °C under a 0.01 M NaNO₃ background. The Langmuir fitting results showed the maximum adsorption capacities for Al/Si and Fe/Si coprecipitates were 0.40 and 0.22 mmol g⁻¹. The coefficients of determination for the fitting are 0.993 and 0.980 ($P < 0.05$, $n = 5$), respectively.

Fe/Si samples is that the coprecipitation process was not performed under thermal condition. Previous studies^{44,60} suggested that the Fe-Si interaction might occur when coprecipitates are subjected to thermal treatments.

Taken together, our findings allow us to conceptualize the structural development for Al/Si and Fe/Si coprecipitates in Fig. 6. Here, AlO₆ with minor amounts of AlO₄ that hydrolyzed from Al³⁺ adhered onto SiO₂ surfaces by forming the direct bonding of Si-O-Al (Fig. 8a). Such robust structural development suppressed the silicate dissolution. On the contrary, the octahedral Fe domains that grew from Fe³⁺ hydrolysis were loosely associated with SiO₂ colloids. The relatively isolated SiO₂ in Fe/Si samples, therefore, led to the more pronounced silicate release (Fig. 8b). The forgoing mechanisms of structural developments were also supported by our XPS data. The Al/Si ratios on the near surfaces of Al/Si coprecipitates were significantly greater than the Fe/Si ratios for Fe/Si samples. The enrichment of Al relative to Si implied that the SiO₂ might serve as a core with Al attached at SiO₂ surfaces. This was in line with the core-shell structure that plausibly formed during the heterogeneous nucleation of Al hydroxide with SiO₂ serving as a template^{59,62}. For Fe/Si samples, however, the trifling amount of Fe relative to Si on the near surfaces of coprecipitates implied the relatively independent phases formed between SiO₂ and Fe hydroxides.

The trend in decreasing sorbed selenite with increasing sorption pH could be attributed to the fewer protonated surface sites on coprecipitates that serve as binding sites for selenite (Fig. 7a)^{54,55}. In relation to the coprecipitation pH condition, both Al/Si and Fe/Si samples prepared at pH 5.0 showed the greater removal efficiency for selenite over the tested pH range (Fig. 7a). Such results might be ascribed to the higher PZC values for samples prepared at pH 5.0 than that prepared at pH 8.0 (Fig. 3a). In terms of coprecipitated elements, the Al/Si-5.0 sample showed a relatively superior sorption capacity of selenite than Fe/Si-5.0 (Fig. 7b), which also agreed with the much higher PZC at Al/Si-5.0 than at Fe/Si-5.0 (Fig. 3a). Due to the core-shell structures, the negative charges on silica might be sterically shielded by Al hydroxides. Such steric effect that decreased the exposure of negative charges of silica was rarely achieved in the independent phases between SiO₂ and Fe hydroxides (Fig. 8). Moreover, while silica occupied the active sorption sites on Al hydroxides by forming the core-shell structures in Al/Si coprecipitates, the dissolved silicate competed for active sites on Fe hydroxides with selenite (Fig. 8)⁶³. On the basis of the differences between Al/Si and Fe/Si samples, surface properties that plausibly deduced from structural attributes for coprecipitates showed the promise to influence the extent of selenite removal. Although the association between Al and Si in Al/Si-8.0 was stronger than that in Al/Si-5.0, the surface charges of Al/Si-8.0 derived from OH groups, resulting in less selenite removal than Al/Si-5.0 (Fig. 7a).

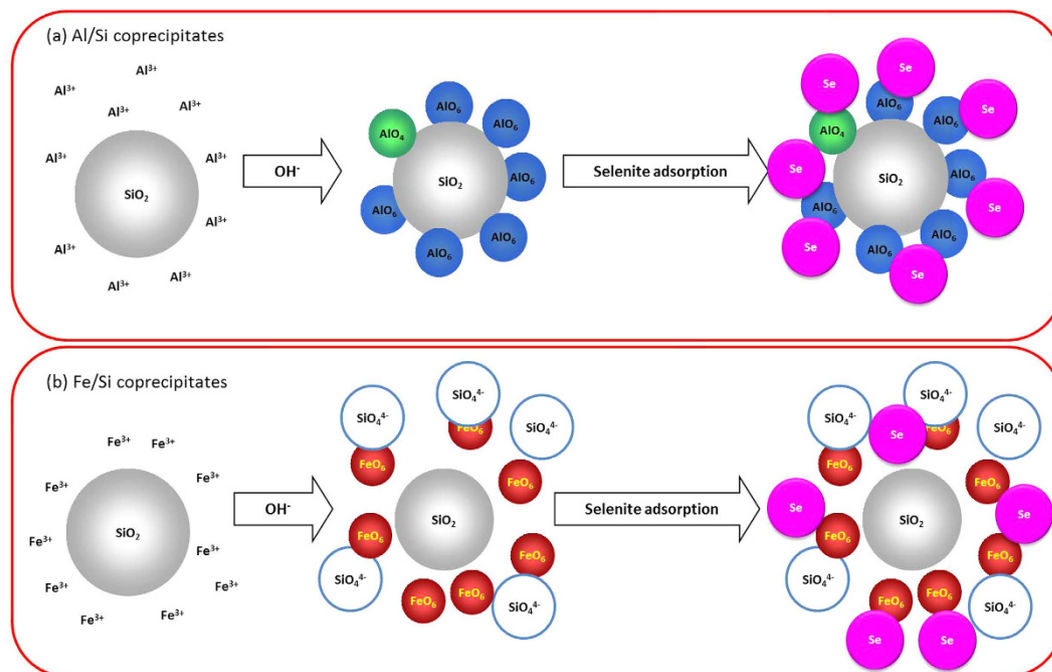


Figure 8. Conceptualized structural development and removal mechanisms of selenite for (a) Al/Si and (b) Fe/Si coprecipitates.

Our spectroscopic results for Al/Si and Fe/Si coprecipitates indicated that Al tended to interact with silica and precipitate as AlO_6 and/or AlO_4 polyhedra on surfaces of silica particles. Such structural development was more pronounced with increasing coprecipitation pH. However, no discernible interaction between Fe and Si was found in Fe/Si coprecipitates. Silica and Fe hydroxides seemed to occur as independent phases. While the formation of Si–O–Al bonds on Al/Si coprecipitates modified their surface properties from the pure SiO_2 , the independent phases in Fe/Si coprecipitates reserved the original SiO_2 properties, including silicate dissolution, surface charges, and structures. The selenite removal by Al/Si and Fe/Si coprecipitates could generally be estimated using their PZC values, however the differences in PZC between Al/Si and Fe/Si samples were a consequence of the various structural development. The Al/Si samples showed a relatively rigid coprecipitate structures and a relatively higher selenite sorption capacity than Fe/Si. Thus, we would suggest using Al rather than Fe to remove Si and selenite from the wastewater treatment discharged from semiconductor manufacturing.

Methods

Preparation and stability of Al/Si and Fe/Si coprecipitates. Briefly, Al/Si and Fe/Si coprecipitates were synthesized by mixing Al(III) or Fe(III) solutions with suspensions of noncrystalline SiO_2 (Cab-O-Sil[®]M5 from Cabot Corp., Tuscolca, IL) under a 0.01 M NaNO_3 electrolyte. Prior to coprecipitation, SiO_2 suspension in a concentration of 10 g L^{-1} 0.01 M NaNO_3 was aged at 25°C under an N_2 atmosphere for 24 h. Subsequently, an appropriate amount of 0.5 M $\text{Al}(\text{NO}_3)_3 \cdot 9 \text{ H}_2\text{O}$ or $\text{Fe}(\text{NO}_3)_3 \cdot 9 \text{ H}_2\text{O}$ was added into the SiO_2 suspension to achieve a final concentration of 5 mM Al or Fe in 1 g L^{-1} SiO_2 . The mixture was agitated at 300 rpm and pH 5.0 using a pH-stat (TIM865 Titration Manager, Radiometer Analytical) with 0.01 M NaOH or HNO_3 . After 24 h, the suspension was centrifuged (Hitachi, 18 PR-52) at $21,400 \times g$ for 15 min. The collected precipitates were washed three times using double deionized water ($\text{DDW} > 18 \text{ M}\text{-cm}$) to remove excessive salts and then freeze-dried (Millitorr Elemech FD-101) for further analyses. Identical coprecipitation procedures, but with different coprecipitation pH at 6.5 and 8.0, was also conducted. Al/Si and Fe/Si coprecipitates synthesized at pH 5.0, 6.5, and 8.0 were hereafter entitled as Al/Si- or Fe/Si-5.0, 6.5, 8.0. Samples of pure Al- and Fe-hydroxides were also hydrolyzed using identical procedures at pH 5.0 and 8.0 (Al- or Fe-5.0, 8.0), but in the absence of SiO_2 .

The degree of stability of Al/Si or Fe/Si coprecipitates was determined based on the silicate dissolution from the solids. The pH-dependent leaching experiment of silicate from Al/Si or Fe/Si coprecipitates was performed in a suspension at a concentration of 3 g L^{-1} DDW. The suspensions were continuously mixed end-over-end for 24 h at room temperature ($25\text{--}27^\circ\text{C}$) across the pH range from 2 to 10 (adjusted using 1 M NaOH or HNO_3). The pH was measured after incubation and prior to the centrifugation. Separation of supernatant from the residual coprecipitates was conducted using the centrifugation at $21,400 \times g$ for 15 min. Amounts of silicate in aqueous samples were analyzed using inductively coupled plasma atomic emission (ICP-AES) spectroscopy (Perkin Elmer, Optima 2000DV).

Surface analyses of Al/Si and Fe/Si coprecipitates. Freeze-dried subsamples of Al/Si and Fe/Si coprecipitates were used to determine the single-point Brunauer-Emmett-Teller (BET) surface areas with a Micromeritics Tristar 3000 gas adsorption analyzer. To understand the morphology of Al/Si and Fe/Si

coprecipitates, a drop of the suspension was spread on a thin cellulose film supported by a copper grid. The specimen was dried for a few minutes at room temperature. The TEM images were investigated with a Jeol JEM-2100 ultrahigh resolution transmission electron microscope (accelerating voltage: 200 KV, stability: $2 \times 10^{-6} \text{ min}^{-1}$). Suspensions of individual coprecipitates in the concentration of 3 g L^{-1} 0.01 M NaNO_3 were prepared, shaken for 24 h, and used for zeta potentials analysis with Malvern Zetasizer 3000 HS.

To determine elemental compositions on near surfaces of Al/Si and Fe/Si coprecipitates, XPS analyses were performed using a ULVAC-PHI (PHI 5000 VersaProbe/Scanning ESCA Microprobe) with monochromatic Al $K\alpha$ radiation ($h\nu = 1486.6 \text{ eV}$) for plane-view samples at 45° angle of emission. The X-ray spot size was approximately $100 \mu\text{m} \times 100 \mu\text{m}$. The pressure in the analytical chamber during spectral acquisition was less than $6.7 \times 10^{-8} \text{ Pa}$. The analyzer pass energy was 187.85 eV. The energy step size for high resolution scans was 0.1 eV. Surface concentrations of elements in atomic percent were obtained from spectral deconvolution and fitting of O 1s, Si 2p, Fe 2p $_{3/2}$, and Al 2p signals.

Structural analyses of Al/Si and Fe/Si coprecipitates. We further used PXRD, FT-IR, ^{27}Al and ^{29}Si solid-state magic angle spinning nuclear magnetic resonance spectroscopy (MAS NMR), and XAS to characterize the structural properties of Al/Si and Fe/Si coprecipitates.

For structural analysis, PXRD patterns were obtained using a diffractometer (PHILIPS X'PERT Pro MPD) with monochromatized and Ni-filtered Cu $K\alpha$ radiation ($\lambda = 1.5405 \text{ \AA}$). The diffractometer was operated at 45 kV and 40 mA.

For FT-IR analysis, spectra were collected using a Bomem, DA8.3 spectrometer with an optical resolution of 2 cm^{-1} . Powers for FT-IR pellets were prepared by mixing oven-dried (110°C) coprecipitates with KBr at a concentration of 5 mg g^{-1} . Spectra were acquired in the range from $4000\text{--}450 \text{ cm}^{-1}$.

The ^{27}Al and ^{29}Si MAS NMR analysis of coprecipitates was performed in a Bruker DSSX-400 NMR solid state high resolution spectrometer, which was operated at 9.4 T with spinning rate varied between 6.4 kHz (7 mm rotor) and 10 kHz (4 mm rotor). The ^{29}Si spectra were calibrated using $\text{Si}(\text{CH}_3)_4$ and acquired at 79.4 MHz with 2.5 μs pluses (45° pulse width) and 7 s recycling times. The ^{27}Al spectra were calibrated using $\text{AlCl}_3(\text{aq})$ and collected at 104.1 MHz with 1 μs pluses (45° pulse width) and 0.5 s recycling times.

The local coordination of Si and/or Fe in Al/Si and Fe/Si samples was characterized using XAS, including X-ray absorption near-edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) spectroscopy. The XAS spectra at Si and/or Fe K-edge were collected at National Synchrotron Radiation Research Center (NSRRC), Hsin-Chu, Taiwan, where the storage ring was operated at 1.5 GeV with a fixed current of 250 mA. An aliquot of freeze-dried sample was mounted in an acrylic sample holder at a thickness calculated to yield unit edge step across the Si and/or Fe K-edge near 1839 and 7112 eV, respectively⁶⁴.

For Si XAS, the wet paste samples were mounted on the holder without cover. The spectra were collected at beamline BL-16A1. The synchrotron radiation that passes through focusing mirrors was detuned by 50% at 50 eV above the Si K-edge. The InSb(111) monochromator was calibrated to 1839 eV based on the first inflection point in the first derivative spectra of elemental Si. Spectra were collected in fluorescence mode with a $\text{He}(\text{g})$ -filled Lytle detector between -70 to $+610 \text{ eV}$ relative to the Si K-edge at 1839 eV, using a step size of 0.2 eV across the absorption edge region.

For Fe XAS, the samples were sealed with Kapton tape, and spectra were collected at Wiggler 20 beamline BL-17C1. The Si(111) monochromator energy of a Fe foil was calibrated to 7112 eV monitored during data collection. Samples were analyzed in fluorescence mode using an $\text{Ar}(\text{g})$ -filled Lytle detector between -200 to $+800 \text{ eV}$ relative to 7112 eV, using a step size of 0.2 eV in the near edge region ($-30\text{--}50 \text{ eV}$) and a step size of $k = 0.06 \text{ \AA}^{-1}$ at higher energies⁶¹.

Multiple XAS scans on each sample were aligned, merged, and processed using the Athena program, an interface to IFEFFIT (version 1.2.10)^{65–67}. Self-absorption effects, if they existed, were corrected in the Athena program. Spectra were baseline corrected using a linear pre-edge function between -75 to -10 eV for Si-XAS and -150 to -30 eV for Fe-XAS. The spectral normalization was conducted using a linear or quadratic function between $20\text{--}600 \text{ eV}$ for Si-XAS and $50\text{--}750 \text{ eV}$ for Fe-XAS, including a flattening function in the post-edge region⁶⁴. For Fe K-edge EXAFS spectra, backgrounds were removed using a cubic spline fit with nodes defined by the AUTOBKG function in IFEFFIT, and the EXAFS data were extracted from the normalized XAS spectra and converted to their (k) function as the k^2 -weighted data⁶¹.

pH-dependent and isotherm sorption of selenite on Al/Si and Fe/Si coprecipitates. The pH-dependent sorption experiments were conducted by mixing 1 mM of selenite reacted with 50 mL the Al/Si-5.0, 8.0 or Fe/Si-5.0, 8.0 coprecipitates at a solid concentration of 3 g L^{-1} . The incubations were carried out at 25°C for 24 h in a 0.01 M NaNO_3 background as a function of pH from 2.0–10.0. Besides, isotherm sorption experiments of selenite (2 to 0.1 mM) were conducted at pH 5.0 with same other incubation conditions as pH-dependent experiments. After the 24-h incubation, suspensions for both pH-dependent and isotherm experiments were passed through a $0.22 \mu\text{m}$ pore-size membrane filter (Millipore filter). Selenite in filtrates was analyzed using ICP-AES, and the isothermal results were fitted by Langmuir isotherm model. In the end of sorption experiments, the suspension samples were collected and analyzed using the Se-XAS. The Se-XANES spectra (data not shown) indicated that no oxidation or reduction of selenite was found during the experiments. The differences between the initial and final selenite concentrations were attributed to sorption onto the coprecipitates.

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Acknowledgements

This study was supported, in part, by a grant from the Ministry of Science and Technology, Taiwan (NSC 102-2313-B-029-005-MY2 and MOST 104-2313-B-005-014-MY3) and also a grant from the Ministry of Economic Affairs, Taiwan (103-EC-17-A-21-S1-229). We also thank Dr. Ling-Yun Jang and Jyh-Fu Lee for providing the beamtime of BL16A1 and BL17C1 at NSRRC, Taiwan. Thanks to Prof. C.S. Lin and Ms. Y.T. Lee of Instrumentation Center, National Taiwan University for FE-SEM experiments.

Author Contributions

Y.-T.C., W.-H.K., Y.-M.T., T.-Y.C. and Y.-T.L. conceived the experiments, Y.-T.C. conducted the experiments, M.-K.W. and H.-Y.T. conducted the TEM experiments, and we add them to the co-authors, and Y.-T.C. and Y.-T.L. analyzed the results. All authors reviewed the manuscript.

Additional Information

Supplementary information accompanies this paper at <http://www.nature.com/srep>

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Chan, Y.-T. *et al.* Molecular Structures of Al/Si and Fe/Si Coprecipitates and the Implication for Selenite Removal. *Sci. Rep.* **6**, 24716; doi: 10.1038/srep24716 (2016).



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