Adsorption behavior of Eu(III) on partially Fe(III)- or Ti(IV)-coated silica

Hee-Jung Im*, Kyoung Kyun Park and Euo Chang Jung

Abstract
The adsorption behavior of Eu(III) onto silica surface, which was partially coated with Fe(III) or Ti(IV), was investigated to determine Fe(III) or Ti(IV) effects on the surface reaction of lanthanides on mineral surfaces in groundwater. Compared with a parallel uncoated silica, the Fe(III)-coated silica did not enhance the adsorption of Eu(III). However, enhanced adsorption of Eu(III) on the Ti(IV)-coated silica was observed by increasing the amount of Ti(IV) on the silica surface.

Keywords: partially coated silica, Ti(IV) coating effects, enhanced adsorption, surface complexation.

Introduction
There has been great interest in the immobilizations and adsorption mechanisms of various toxic ions in aqueous solutions by using silica-based sorbents [1-4]. The adsorption reaction of a metal ion onto a metal (hydr) oxide surface is explained in terms of surface complexation. Besides free metal ions, hydrolyzed or complexed species [5], or even the colloidal species can be adsorbed [6]. Surface precipitation may occur even in a concentration below the surface site saturation [7]. Frequently, experimental evidence indicates surface nucleation of metal hydroxides [8].

Due to their ubiquity in soils and sediments and high specific surface area, iron or titanium hydroxides (Fe or Ti oxides) around silicate minerals may play a role in the migration of actinides in groundwater. The interactions of actinides on immobile solid surfaces are important processes that determine retardation during transport. Usually, Eu(III) is considered to be an adequate chemical analogue of radiotoxic nuclides, Am(III) and Cm(III). These actinides consist mainly of long-lived nuclides that emit alpha radiation, and their radioactivity continues for several hundred thousands of years [9].

The following studies have been reported. Fe-modified silica gel was investigated as an adsorbent for humic acids employing an electrostatic binding to Fe and/or in coordination with Fe by direct substitution of OH and Cl on Fe sites [10]. TiO₂-coated SiO₂ synthesized by hydrolysis and condensation of various silicate and titane precursors has been actively studied as photocatalysts due to its photocatalytic and photovoltaic effects [11]. Eu(III) sorption onto clay minerals was quantitatively modeled with pH ranging from 3 to 10 using cation exchange reactions for Eu(III)/Na(I) and Eu(III)/Ca(II) [12].

However, Eu(III) sorption onto Fe(III)- or Ti(IV)-coated silica has not received as much attention as UO₂²⁺ sorption. The aim of this paper is to study the sorption of Eu(III) from an aqueous solution on Fe(III)- or Ti(IV)-coated silica to understand the trace radionuclide migration which occurs in groundwater.

Experimental section
The chemicals used in this study including silica (Sigma-Aldrich Corporation, St. Louis, MO, USA; particle size 40 to 63 μm; surface area 550 m²/g), ferric nitrate [Fe(NO₃)₃·9H₂O], titanium butoxide [Ti(OBu)₄], 1,10-phenanthroline [C₁₂H₈N₂], ethanol [C₂H₅OH], toluene [C₆H₅CH₃], and Europium(III) oxide [Eu₂O₃] were all of high purity and used as received. Perchloric acid [HClO₄], hydroxylamine hydrochloride [NH₂OH·HCl], sodium perchlorate [NaClO₄], sodium hydroxide [NaOH], sulfuric acid [H₂SO₄], and hydrofluoric acid [HF] were of analytical grade and used without further purification. NaOH solution was titrated with 0.1 M hydrochloric acid [HCl] standard solution (Merck &
Co., Inc., Whitehouse Station, NJ, USA) in the presence of a phenolphthalein indicator.

The dry silica was dispersed in 3.7 M HNO₃ for one day and washed with distilled water until the wet silica surface was neutral. Finally, the resulting silica was dried in an oven at 120°C for 6 h and stored in a capped bottle after cooling.

The 11.3 mM Eu₂O₃ in 20.62 mM HClO₄ was prepared as a stock solution for the Eu(III) adsorption tests. The metal-ion concentration of the stock solution was determined with inductively coupled plasma - atomic emission spectrometry [ICP-AES] before diluting for the adsorption experiments. All the solutions were handled under a nitrogen gas flow.

Coated, adsorbed, and desorbed metal concentrations were determined with an ultraviolet and visible [UV-vis] absorption spectrophotometer (Cary 3, Varian, Inc., Santa Clara, CA, USA) and an ICP-AES (ULTIMA 2C, HORIBA Jobin Yvon, HORIBA, Ltd., Minami-ku, Kyoto, Japan). A spectrofluorometer (FS-900CD, Edinburgh Instruments, Livingston, U.K.) was used to obtain appropriate fluorescence spectra.

Partial Fe(III) coating on silica surface

Fe(NO₃)₃·9H₂O (0, 35, 70, 140, 280, and 420 mg) was added with stirring to each silica (50 g) in 500 mL of distilled water. The pH was adjusted to 4.5 with 0.1 M HClO₄ or 0.1 M NaOH, and each mixture was stirred for 2 h. The partially Fe(III)-coated silica was glass filtered, washed with a pH 4.5 HClO₄ solution and distilled water three times each, and dried in sequence.

Partial Ti(IV) coating on silica surface

Ti(OBu)₄ was slowly added with stirring to each silica (15 g) in C₂H₅OH and C₆H₅CH₃ (1:1) mixed solution until 0, 5, 10, 20, 100, and 200 mM of Ti(IV) was added in the 50 mL total solution. Then, each mixture was stirred for 2 h. The partially Ti(IV)-coated silica was washed with C₂H₅OH and C₆H₅CH₃ (1:1) mixed solution three times each, and dried at 120°C for 6 h sequentially.

Eu(III) adsorption onto Fe(III)- or Ti(IV)-coated silica

In each test, 500 mg of dissimilar Fe(III)- or Ti(IV)-coated silica was placed in a 60-mL beaker, and 20 mL of distilled water was added in the beaker. The total volume of each mixture was adjusted to 50 mL, and the final concentration was 0.1 mM Eu₂O₃ in 0.18 mM HClO₄ with a controlling ionic strength with 0.1 M NaClO₄. At this point, for the observation of a pH-dependent adsorption, 0.1 M NaOH under the N₂ gas flow in order to eliminate the remaining CO₂ in the solution, was properly added to each mixture, and the pH went up to 8 for Eu(III) adsorption tests. The mixture in each polyethylene beaker was stirred for more than 30 min until the pH equilibrium was achieved. The mixture was then analyzed by ICP-AES after being filtered through a 0.1-μm pore-sized membrane filter. Fluorescence of Eu ions on Fe(III)- or Ti(IV)-coated silica was obtained from the sediments.

Results and discussion

It has been known that Si-OH in silica (SiO₂) is dissociated into surface ≡Si-O and free H⁺ at pH > 3, and as the result, the surface is negatively charged, which is appropriate to incorporate electron-deficient metals to the silica surfaces. Here, Fe(III) or Ti(IV) was primarily fixed on the silica surface through the ≡Si-O-Fe or ≡Si-O-Ti structure. In contrast, Eu(III) ion is easily hydrolyzed [13] and forms insoluble trihydroxide precipitates [14] and polynuclear hydroxo complexes [15]. The hydrolyzed Eu-OH is assumed to be sorbed into Fe(III)- or Ti(IV)-coated silica in aqueous Eu(III) solutions.

Each and every coated Fe ion on the silica surface was stripped using 5 M HCl, and the amount was measured using a UV-vis spectrophotometer by reducing all Fe (III) to Fe(II) with NH₂OH·HCl for the production of colored Fe(II)-orthophenanthroline complexes (ferroin, (Phen)₃Fe²⁺), which are sensitive to UV-vis absorption [16]. From the UV-vis absorption spectra, 0, 2.05, 3.81, 7.38, 13.8, and 21.3 μmol/g (Fe/silica) Fe-coated silica was obtained when 0, 35, 70, 140, 280, and 420 mg of Fe(NO₃)₃·9H₂O were added in 50 g of each silica. During this process, the following product is expected:

≡Si - OH + Fe³⁺ + H₂O → ≡Si - O - Fe - OH.

The adsorption of Eu(III) onto the silica at various pH shows no influence of a surface coating by Fe(III) (Figure 1). In other words, Fe(III) coating on silica surfaces did not enhance adsorption of Eu(III) compared with the uncoated silica. The paper written by Pokrovski et al. [17] states, 'at pH > 2.5 in the presence of aqueous silica (mₑ approximately at 0.01 mol/kg), Fe-Fe dimers and trimers shared one or two edges of FeO₆-octahedra, and silicon tetrahedra linked to two neighboring Fe octahedral via corners’. Due to linkages to the free corners of FeO₆-octahedra, the number of available sorption sites for Eu(III) in Fe(III)-OH had decreased, and thus, it did not show a significant Fe(III)-coating effect compared with the uncoated silica.

However, as shown in Figure 2, the fluorescence decreased with an increased amount of coated Fe(III) even though the shape of the fluorescence spectra did not change. In Figure 2a, the Eu(III) fluorescence spectra excited at 394 nm were scanned from 525 to 650 nm.
nm. This range covers the wavelengths corresponding to $^5\!D_0 \rightarrow ^7\!F_J$ ($J = 0, 1,$ and 2) transitions, and the peaks at 588 nm and 613 nm correspond to $^5\!D_0 \rightarrow ^7\!F_1$ and $^5\!D_0 \rightarrow ^7\!F_2$, respectively. Since the $^5\!D_0 \rightarrow ^7\!F_1$ transition is allowed in magnetic dipole and its strength is not sensitive to coordination environment, the decrease in the peak intensity can be explained in terms of a quenching effect. The fluorescence intensities of Eu(III) at 613 nm were quantitatively expressed in Figure 2b according to the amount of coated Fe(III). The $^5\!D_0 \rightarrow ^7\!F_2$ transition should also be affected by the quenching effect as with the $^5\!D_0 \rightarrow ^7\!F_1$ transition. In contrast, Ti(IV)-coated
silica showed higher Eu(III) adsorption capacities while increasing the amount of Ti(IV) on silica surface as shown in Figure 3.

The amount of every coated Ti ion on the silica surface was measured using ICP-AES after a pretreatment process with H₂SO₄-HF solution to remove SiO₂. From the ICP-AES analysis, 0, 16.1, 40.5, 63.9, 324, and 407 μmol/g (Ti/silica) Ti-coated silica was obtained when 0, 5, 10, 20, 100, and 200 mM (in other words, 0, 20, 40, 80, 350, and 700 μmol/g (Ti/silica)) of Ti(OBu)₄ were added in 15 g of each silica. In this process, the following product is expected:

≡Si - OH + Ti⁴⁺ + H₂O → ≡Si - O - Ti - OH.

The Ti(IV)-coated silica exhibited a stronger binding toward Eu(III) than the uncoated silica, and the preferential binding is considered due to a higher metal Lewis acidity of Ti than Si. The hard Lewis acid, Eu(III), forms more stable complexes with hydroxyl ligands on relatively harder TiO₂ than those on SiO₂. The Eu(III) adsorption processes onto the partially Ti(IV)-coated silica involve the combination of Eu(III) hydrolysis and the adsorption of the hydrolysis product, Eu³⁺-OH, to produce ≡Si-O-Ti-OH-Eu³⁺ and/or ≡Si-O-Ti-O-Eu³⁺ in addition to ≡Si-OH-Eu³⁺ and/or ≡Si-O-Eu³⁺, depending on the pH of the prepared solutions [12].

The enhanced adsorption of Eu(III) onto the silica coated by Ti(IV) is partially confirmed by observing the increase in fluorescence intensities as increasing the amount of Ti(IV) on the silica surface as shown in Figure 4[18]. The fluorescence intensities of Eu(III) at 613 nm were quantitatively expressed in Figure 4b according to the amount of coated Ti(IV). In the case of Eu(III), the maximum fluorescence peaks corresponding to [⁵D₀ → ⁷F₂ transition at 613 nm were red-shifted in a wavelength nearly 618 nm with an increased coated Ti(IV) (Figure 4a). It suggests the formation of other species with a reduced hydration number such as ≡Si-O-Ti-O-Eu³⁺ or ≡Si-O-Ti-O-Eu³⁺-OH. No fluorescence-quenching effect of the increased amount of coated Ti(IV) indicates a similar chemical environment between the species reacting with the titanol and silanol functional groups.

Conclusions
This study shows an example of foreign ion effects on the adsorption of actinide onto a mineral surface. In the case of the Ti(IV) ion for Eu(III) adsorption onto a silica surface, Ti(IV) enhances the adsorptivity as far as it exists as a surface hydroxide. The enhancement in
adsorptivity decreases when the surface hydroxide converts to oxide prior to Eu(III) adsorption. In contrast, Fe (III) coating on silica surfaces did not enhance adsorption of Eu(III), nor were there any changes in fluorescence properties compared with uncoated silica.

Acknowledgements
This work was supported by the nuclear research and development program through the National Research Foundation of Korea funded by the Ministry of Education, Science and Technology.

Authors’ contributions
H-JI drafted the manuscript, prepared samples, and acquired various adsorption and spectrographic data. KKP conducted the preparation of sample and analysis of data. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Received: 10 September 2011 Accepted: 5 January 2012 Published: 5 January 2012

References
1. Im H-J, Barnes CE, Dai S, Xue Z. Functionalized sol-gels for selective copper(II) separation. Environ Sci Technol 2000, 34:2209-2214.
2. Im H-J, Barnes CE, Dai S, Xue Z. Functionalized sol-gels for mercury(II) separation. A comparison of mesoporous materials prepared with and without surfactant templates. Micropor Mesop Mater 2004, 70:57-62.
3. Im H-J, Barnes CE, Dai S, Xue Z. Mechanistic investigation of hydrolysis reactions of diethacetal derivatives grafted on silica gel. Talanta 2004, 63:259-264.
4. Im H-J, Yost TL, Yang Y, Bramlett JM, Yu X, Fagan BC, Allain LR, Chen T, Barnes CE, Dai S, Roecker LE, Sepaniak MJ, Xue Z-L. Reactions of dithioacetal derivatives grafted on silica gels. J Environ Radioact 1993, 20:151-167.
5. Degueulde C, Wemli B. Association behavior of NAm(III) on SO4(2-)/amorphous silica and SO4(2-)/quartz colloids. J Environ Radioact 1993, 20:151-167.
6. Cujic-Andjan D, Cujic-Andjan N, Pravic M. Sorption of europium(III) and iron (III) in ionic and colloidal states on silica gel. J Chromatogr 1894, 586:180-186.
7. Farley KL, Dzombak DA, Marel FMM. A surface precipitation model for the sorption of cations on metal oxides. J Colloid Interface Sci 1985, 106:226-242.
8. Charlet L, Manceau A. X-ray absorption spectroscopic study of the sorption of Cr(III) at the oxide-water interface. J Colloid Interface Sci 1992, 148:443-458.
9. Kim J-H. Significance of actinide chemistry for the long-term safety of waste disposal. Nucl Eng Technol 2006, 38:459-482.
10. Monguchi T, Yano K, Tahara M, Yaguchi K. Metal-modified silica adsorbents for removal of humic substances in water. J Colloid Interface Sci 2005, 283:300-310.
11. Shi A, Cheng A. The effect of different preparation conditions on the photocatalytic activity of TiO2·SiO2 beads. Surf Coat Technol 2006, 200:3637-3643.
12. Bradbury MH, Baeyens B, Geckeis H, Rabung TH. Sorption of Eu(III)/Cm(III) on Ca-montmorillonite and Na-illite. Part 2: surface complexation modeling. Geochim Cosmochim Acta 2005, 69:5403-5412.
13. Rocha E, Choppin GR. In Handbook on the Physics and Chemistry of Rare Earths. Volume 15. Edited by: Gschneidner KA, Jr, Eyring L. Amsterdam: Elsevier Sci. Pub. B.V; 1991:393-442.
14. Smith RM, Martel AE. In Critical Stability Constants. Volume 4. New York: Plenum Press; 1976.
15. Cauvery L, Lukano M, Porto R. The hydrolysis of the La(III) ion in aqueous perchlorate solution at 60°C. Polyhedron 1987, 6:1283-1290.
16. Nestorides A. Rapid method analysis of Portland cement. Analyst 1970, 95:51.
17. Pokrovski GS, Schott J, Farges F, Hazemann J-L. Iron (III)-silica interactions in aqueous solution: Insights from x-ray absorption fine structure spectroscopy. Geochim Cosmochim Acta 2003, 67:3559-3573.
18. You H, Nagami M. Optical properties and local structure of Eu(III) ions in sol-gel TiO2·SiO2 glasses. J Phys Chem B 2004, 108:12003-12008.

Figure 4 Fluorescence spectra and intensities of Eu(III) on partially Ti(IV)-coated silica. (a) Fluorescence spectra (λexc = 394 nm), and (b) fluorescence intensities (λexc = 394 nm) of Eu(III) on partially Ti(IV)-coated silica at 613 nm, according to the amount of coated Ti(IV) at pH 7.