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Sorption and redox speciation of plutonium at the illite surface under highly saline conditions

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Abstract. Natural groundwater may contain high salt concentrations, such as those occurring at several potential deep geological nuclear waste repository sites. Actinide sorption to clays (e.g. illite) under saline conditions has, however, been rarely studied. Furthermore, both illite surface and ionic strength may affect redox speciation of actinides like plutonium. In the present study, Pu sorption to illite is investigated under anaerobic conditions for $3 < \text{pH}_m (= - \log \text{mH}^+) < 10$ and $m_{\text{NaCl}} = 1.0$ and $3.2$ molal (m). Results are compared with previous data for $m_{\text{NaCl}} = 0.1$ m.

According to redox potential measurements and based on Eu(III)-illite sorption data (taken as analogue of Pu(III)), the strong effect of $m_{\text{NaCl}}$ on overall Pu uptake observed for $\text{pH}_m < 6$ is mainly attributed to the presence of Pu(III) and its competition with Na$^+$ for ion exchange sites. For $\text{pH}_m > 6$, overall Pu uptake is largely insensitive to $m_{\text{NaCl}}$ due to the prevalence of strongly adsorbed Pu(IV). By applying appropriate corrections to the activity coefficients of dissolved ions and using the 2-site protolysis non-electrostatic surface complexation and cation exchange (2 SPNE SC/CE) model, experimental data on Pu sorption to illite as a function of pH, Eh and $m_{\text{NaCl}}$ can be very well reproduced.

Keywords: Plutonium, illite, clay, saline, speciation, redox, sorption, surface complexation model, ion exchange.

Graphical Abstract
1. Introduction

Only few investigations of actinide uptake on clay minerals have been carried out at elevated ionic strength [1-3]. The majority of investigations pertain to I = 0.1 molar (m) and rarely up to 1.0 m (e.g. [4-6]; and references therein). However, clay rock pore waters as e.g. in the Jurassic and lower Cretaceous clay rock in Northern Germany, discussed as potentially appropriate host rock formations for a final nuclear waste repository, may contain salt contents as high as about 5 M [7]. Sedimentary rocks currently investigated in Canada are in contact with brine solutions up to 6.5 M [8]. Therefore, detailed sorption investigations of radionuclide onto clay materials under saline conditions become necessary. At high salt concentrations, activity coefficients of aqueous species change dramatically and actinide cations may form aqueous complexes with background anions, which can affect actinides speciation (including redox equilibria), but this can be predicted. By contrast, the effect of high ionic strength on mineral (i.e. including clay minerals) surface properties is elusive. Previous studies performed at high ionic strength show that non-electrostatic sorption models are quite suitable to simulate proton and metal ion sorption to naturally occurring surfaces, e.g. marine microalgae or bacteria [9-11]. Recently, Eu(III) sorption to illite and smectite was investigated in 0.1 < m_{NaCl} < 3.9 m [3]. The experimental results could be described by the 2 site protolysis non-electrostatic surface complexation and cation exchange (2 SPNE SC/CE) model [5,6] coupled to the specific ion interaction theory (SIT [12]) or the Pitzer formalism [13], to account for activity coefficients of solutes in concentrated media.

Compared to Eu(III), the complex redox chemistry of Pu adds another dimension to sorption studies. Pu occurs in the oxidation states +III, +IV, +V or +VI and its geochemical behavior, such as solubility and mobility, strongly depends on its redox state [14-15]. Under reducing conditions, Pu(IV) and Pu(III) prevail [16-20]. However, Pu(III) sorption to minerals has rarely
been studied separately as it is usually accompanied by Pu(IV). It has been shown, that the overall uptake of redox sensitive actinides and their redox speciation at mineral surfaces can be estimated by taking into account the uptake of the individual redox states and the measured redox potentials, i.e. the (apparent) electron activity, pe [21-22]. Using the 2 SPNE SC/CE model, previous work demonstrated the applicability of the approach to describe Pu uptake on illite under anaerobic conditions in 0.1 m NaCl, where the Pu(IV)/Pu(III) redox couple was involved [23]. Here, the approach is extended to Pu sorption and redox speciation in contact with illite under saline conditions (up to 3.2 m NaCl). While there is no study dedicated to tetravalent actinide sorption to illite at such high ionic strength, Eu(III) sorption to illite (often studied as chemical analogue of Pu(III)) is affected by NaCl concentration [3]. Therefore, Pu sorption and Pu(IV)/Pu(III) redox equilibria at the illite surface are expected to be affected by the ionic strength.

2. Materials and methods

Chemicals (all pro analytical quality or better) were obtained from Merck (Darmstadt, Germany) or Riedel de Haen (Seelze, Germany). Solutions were prepared with de-ionized “MilliQ” water (specific resistivity, 18.2 MΩ cm⁻¹). The purified Na-illite was provided within the EC project CP CatClay. The source material derives from lacustrine continental sediments deposited at the Upper Eocene (~ -35 Ma) in the basin of Le Puy en Velay (Massif Central, France). The purification procedures and the characterization of the purified illite (<63 μm) were previously detailed [21], and will not be repeated here. Note that in the last step of the purification, the clay suspension was freeze dried, to exclude bacterial activity.
2.1. Plutonium and Europium stock solutions

A $^{238}$Pu stock solution was prepared from Pu solution dissolved in nitric acid, which was fumed three times by 0.1 M HClO$_4$, in order to remove all impurities and organic traces. The concentration of the Pu stock solution was $3.9 \times 10^{-5}$ M in 0.1 M HClO$_4$. From this, a more dilute solution ($[\text{Pu}] = 1.9 \times 10^{-6}$ M) in 0.1 M HClO$_4$ was prepared for experiments at low Pu(IV) concentration. The diluted $^{238}$Pu stock solution contained 85% Pu(IV), 11% Pu(V) and 4% Pu(III), as determined by liquid-extraction methods [23]. Aqueous $^{238}$Pu concentrations were determined by liquid scintillation counting (LSC) using the scintillation cocktail Ultima Gold XR with a liquid scintillation analyzer (Tri-Carb 3110TR). In addition, the stock solution of $^{238}$Pu was checked by ICP-MS and the results were in excellent agreement with LSC measurements.

Eu(III) was used in some experiments as a chemical analogue of Pu(III). A radiotracer solution was purchased from Amersham International (total Eu concentration: $6.0 \times 10^{-4}$ M) with isotopic composition $^{151}$Eu (83%), $^{152}$Eu (13%, $t_{1/2} = 13.33a$) and $^{153}$Eu (4%). $^{152}$Eu is a $\beta$, $\gamma$-emitter and can be conveniently analyzed by $\gamma$-counting. In the present study, precise determination of dissolved $^{152}$Eu was performed using a Perkin Elmer Wallac gamma counter (Wizard 1480).

2.2. Determination of pH and Eh

The pH in the clay suspensions was measured by an Orion 525A (pH meter) and a Ross electrode calibrated with 4 standard buffers (pH 3, 5, 7, and 9; Merck). The error in pH measurements is ± 0.05. For pH measurements in highly saline conditions ($I > 0.1$ m) a correction term is applied to the measured operational pH-values ($pH_{\text{exp}}$). The molal proton concentration, i.e. $-\log m_{H^+}$ ($pH_m$), was obtained involving an empirical correction coefficient ($A_{\text{NaCl}}$) according to equations 1 and 2:
\[ pH_m = pH_{exp} + A_{NaCl} \]  

\[ A_{NaCl} = 0.0013 \times (m_{NaCl})^2 + 0.1715 \times m_{NaCl} - 0.0988 \]

\( A_{NaCl} \) depends on background electrolyte composition and concentration and has been accurately determined for NaCl solutions for the electrodes we use [24]. \( m_{NaCl} \) is the molality (mol kg\(^{-1}\)) of the background electrolyte.

The redox potentials in the clay suspensions were measured using an Orion 525A (\( E_h \) meter) and a Pt electrode combined with a Ag/AgCl reference system (Metrohm). Raw data were converted into \( E_h \) vs. standard hydrogen electrode (SHE) by correcting for the potential of the reference electrode. \( E_h \) was converted to the apparent electron activity, 
\[ p_e = -\log a_e = 16.9 \times E_h (V) \] at 25°C. A commercial redox-buffer (220 mV, Schott instruments) was used for calibration. An equilibration time of 15 min was allowed for all \( E_h \) measurements, after having stirred the suspension. Uncertainties in \( E_h \) measurements are ±50 mV (±0.8 for \( p_e \)-scale) [20, 25]. Unlike for the measurement of pH, to our knowledge, there is no ionic strength dependent correction to apply to the experimental \( E_h \) with the presently used set-up.

### 2.3. Batch sorption experiments

All sorption studies were performed as batch type experiments. The procedure is the same as in our previous work dedicated to \( m_{NaCl} = 0.1 \) m [23]. The effect of \( pH_m \) was investigated at an initial Pu concentration ([Pu]\(_{tot}\)) of \( 8 \times 10^{-11} \) M in 3.2 m NaCl. In addition, the effect of [Pu]\(_{tot}\) was investigated for \( m_{NaCl} = 1.0 \) and 3.2 m, \( 8 \times 10^{-11} < [Pu]_{tot} < 10^{-8} \) M and \( pH_m \approx 4.5, 6 \) and 9.5.
Batch experiments were carried out in 40 mL polypropylene centrifuge tubes at room temperature in an argon glove box (< 1 ppm O₂, absence of CO₂). The suspension volume was 25 mL. At a solid to liquid ratio of 2 g L⁻¹, the suspensions were preconditioned in 1 or 3.2 M NaCl under continuous shaking for 4-5 days to achieve a given target pHₘ value by adding 0.1 M HCl or 0.1 M NaOH. After adding Pu to the illite suspension, pHₘ was adjusted again to the respective target pHₘ. Neither pH nor Eh buffers were used. The vials were then closed and shaken end-over-end. According to previous studies [23, 26-29], potentially occurring redox processes might be rather slow. Consequently, an equilibration time of one year was chosen to make sure that equilibrium has established [23, 28]. After one year, pHₑₓᵖ and Eh were measured in the suspension and an aliquot of each sample was transferred to centrifuge tubes (Beckmann, Recorder No.: 356562) and centrifuged (Beckmann Coulter XL-90 K) at 90,000 rpm (~700,000 g max) for one hour. The supernatant was analyzed for dissolved Pu by LSC.

Results from the batch experiments will be expressed throughout as distribution coefficients (Rₖ in L kg⁻¹), calculated by the following equation:

\[
R_d = ([\text{Pu}]_{\text{tot}}/\text{[Pu]}_{\text{aq}} - 1) \times V/m
\]  

where [Pu]ₐq and [Pu]ₜₜ (M) are the dissolved (final) equilibrium and total (initial) concentrations of Pu, respectively. The term V/m corresponds to the aqueous solution volume to illite mass ratio (L kg⁻¹). According to a previous experimental study [23], an uncertainty of ±0.2 is assigned to log Rₖ, although it could be larger according to comparable studies where more than 99% uptake is obtained [6]. Under such conditions, larger uncertainties are induced by analytical constraints.
Batch Eu sorption experiments were performed for [Eu]_{tot} = 3 \times 10^{-9} \text{ M} applying the same protocol as for Pu, except that Eh was not recorded because Eu is not redox sensitive under our experimental conditions. After one week contact time and subsequent ultracentrifugation, the supernatant was analyzed for dissolved Eu by γ-spectrometry. The $m_{NaCl}$ values for Eu(III) (0.1, 0.9 and 3.9 m) slightly differ from those for Pu (0.1, 1.0 and 3.2 m). This has no impact on our conclusions. Previous work suggested no significant effect of incorporation reactions or sorption by secondary phases on Eu/Pu uptake between 1 week and 1 year [23].

2.4. Thermodynamic modeling

pH-pe diagrams for Pu were constructed using PhreePlot [30], which contains an embedded version of the geochemical speciation program PHREEQC [31]. Thermodynamic constants for Pu and Eu aqueous speciation were taken from the NEA thermodynamic database [32]. Lanthanides and actinides exhibit similar chemical behavior for the same redox state and chemical analogues are often used to estimate the complex geochemical behavior of Pu [14, 33]. Therefore, in case of gaps in the Pu database, data for analogues were chosen (i.e. Eu(III), Np(IV), Np(V) and U(VI) for the respective Pu redox states). The specific ion interaction theory (SIT [12]) was used to calculate activity coefficients of aqueous species and the activity of water. These calculations are required in order to correct thermodynamic constants (including for redox equilibria) for ionic strength effects. SIT is generally considered valid for ionic strengths up to 3 - 4 m. Pitzer model [13], which is valid for a larger range of ionic strengths but more complex than SIT, can also be applied to calculate activity coefficients for aqueous ions in concentrated media (see e.g. [3]). SIT and Pitzer equations are implemented in PHREEQC but only SIT was used in
this study. At 25 °C, activity coefficients for an aqueous species \( i \) (\( \gamma_i \)) with a charge \( z_i \) are calculated as follows:

\[
\log \gamma_i = -z_i^2 \frac{0.509 \times \sqrt{I}}{1 + 1.5\sqrt{I}} + \sum_k \varepsilon(i,k) \times m_k
\]  

(4)

where \( m_k \) is the molality of the aqueous species \( k \) (mol kg\(^{-1}\)), and \( \varepsilon(i,k) \) is the specific ion interaction coefficient between species \( i \) and \( k \) (kg mol\(^{-1}\)). Auxiliary reactions and constants are from the SIT database provided with PHREEQC (sit.dat file). All parameters involved in Pu and Eu aqueous speciation calculations are listed in Tables S1 and S2 in the supporting information.

The 2 SPNE SC/CE model was used to simulate Pu and Eu sorption to illite. The cation exchange capacity (CEC) of the illite was set to 0.225 eq kg\(^{-1}\) [34]. Only the strong sites of the 2 SPNE SC/CE model are considered in the adsorption calculations with a site density of 2×10\(^{-3}\) mol kg\(^{-1}\) [34]. The weak sites play a negligible role in our experiments, since a maximum loading of only 5×10\(^{-6}\) mol kg\(^{-1}\) is investigated (i.e. for \([\text{Pu}]_{\text{tot}} = 10^{-8} \text{ M}\)). Surface complexation constants for Eu(III) and Pu(IV) extrapolated to zero ionic strength are available from previous work [23, 34]. These constants accurately predict Eu(III), Pu(III) and Pu(IV) uptake on the present illite in 0.1 m NaCl [23]. A summary of the parameters for the 2SPNE SC/CE model is given in supporting information (Table S3).

3. Results and discussion

3.1. Plutonium sorption to illite in solutions of different ionic strength

Results of log \( R_d \) for the Pu-illite system (8×10\(^{-11}\) < \([\text{Pu}]_{\text{tot}} < 10^{-8} \text{ M}\)) plotted versus pH\(_{\text{m}}\) for 0.1, 1 and 3.2 m NaCl are shown in Fig. 1a, 1b and 1c, respectively. Data for 0.1 m NaCl are taken from our previous work [23]. Experimental Pu sorption data in 0.1 < \( m_{\text{NaCl}} < 3.2 \) m are
plotted together for comparison on Fig. 1d. Generally, Pu uptake on illite increases with pH\_m and
remains constant for pH\_m > 6. For pH\_m < 6, Pu sorption decreases with increasing m\textsubscript{NaCl},
whereas it is not affected by m\textsubscript{NaCl} for pH\_m > 6. Pu uptake data at 0.1 m NaCl is explained by
accounting for the presence of both Pu(III) and Pu(IV) for pH\_m < 6, where Pu(III) uptake is
weaker than that of Pu(IV), whereas Pu(IV) prevails for pH\_m > 6 [23]. Experimental data for
Eu(III) (taken as a chemical analogue for Pu(III)) sorption to illite (for m\textsubscript{NaCl} = 0.1, 0.9 and 3.9
m) are included in Fig. 1. In agreement with a recent study on Eu(III) sorption to illite in saline
solutions [3], log R\textsubscript{d}(Eu(III)) decreases with increasing m\textsubscript{NaCl} for pH\_m < 6, and is not affected by
m\textsubscript{NaCl} for pH\_m > 6. The ionic strength dependence of log R\textsubscript{d}(Pu) and log R\textsubscript{d}(Eu) at pH\_m < 6 is
consistent with the assumption of a cation exchange mechanism being at least partly responsible
for the uptake of Pu(III). Note, however, that our Pu uptake data for pH\_m < 6 scatter significantly,
which might be due to slight variations in redox conditions in individual batch experiments, as
discussed later.

Fig. 2 shows all sorption data measured for pH\_m > 6, 8×10\textsuperscript{-11} < [Pu]\textsubscript{tot} < 10\textsuperscript{-8} M and m\textsubscript{NaCl} = 0.1, 1 and 3.2 m as log R\textsubscript{d} versus the logarithm of the final Pu concentration in solution after
phase separation (log [Pu]\textsubscript{aq}). Fig. 2 suggests that Pu uptake is not significantly influenced by
[Pu]\textsubscript{tot} or ionic strength within the range of concentrations investigated and for pH\_m > 6. [Pu]\textsubscript{aq} in
presence of illite is close to or below the solubility limit of Pu(IV) in equilibrium with
PuO\textsubscript{2}(am,hydr) (10\textsuperscript{-10.4±0.5} M [35]). But the more or less constant R\textsubscript{d} values even at the highest
investigated Pu concentrations suggest the absence of significant precipitation or surface
precipitation of PuO\textsubscript{2}(am,hydr). On average for all m\textsubscript{NaCl} investigated, log R\textsubscript{d} = 5.3 ± 0.3 (1\sigma) for
pH\_m > 6, which is in excellent agreement with previous work in m\textsubscript{NaCl} = 0.1 m (log R\textsubscript{d} = 5.2 ±
0.2) [23].
Fig. 3 shows pe values of all our experiments plotted versus pH\textsubscript{m} together with the predominance diagram for Pu in solution for \(m_{\text{NaCl}} = 0.1\) m. The corresponding predominance fields in 1 and 3.2 m NaCl shift because the activity coefficients of aqueous species vary with the ionic strength. Note that formation of aqueous Eu(III)- and Pu(IV)-Cl complexes is almost insignificant (< 5%) for 0.1 < \(m_{\text{NaCl}}\) < 3.2 m (preliminary calculations not shown). However, the ionic strength dependence of the Pu(III)/Pu(IV) redox borderline is minor relative to the uncertainty of measured pe (see Fig. S1). In view of the scattered experimental pe data we are not able to clearly identify an ionic strength dependence of pH\textsubscript{m}-pe values, and the present results are in excellent agreement with previous studies in \(m_{\text{NaCl}} = 0.1\) m [21, 23]. The calculated predominance diagram and measured pH\textsubscript{m}-pe values suggest that Pu(III) could predominate or exist in significant fractions at pH\textsubscript{m} < 6 and Pu(IV) becomes dominant above this pH\textsubscript{m}. The pronounced ionic strength dependence of Pu uptake (comparable to that of Eu) at low pH can thus easily be explained by the ion exchange competition of Pu\(^{3+}\) and Na\(^{+}\) [3,23,34]. Based on our pH/pe data, we can also conclude that the ionic strength independent experimental Pu-illite sorption data at pH\textsubscript{m} > 6 are due to the exclusive presence of Pu(IV) undergoing inner-sphere surface complexation only [6, 31].

3.2. Modeling results

Pu(IV) uptake on illite is simulated for \(m_{\text{NaCl}} = 0.1, 1.0\) and 3.2 m with the 2 SPNE SC/CE model. The same exercise is made for Eu(III) and \(m_{\text{NaCl}} = 0.1, 0.9\) and 3.9 m. Previously reported surface complexation constants for Pu(IV) and Eu(III) are used for this purpose, without any parameter adjustment [23,34]. Results are shown in Fig. 1. In agreement with a previous study [3], the 2SPNE SC/CE model accurately predicts the effect of high \(m_{\text{NaCl}}\) on Eu(III) uptake.
on the whole investigated pHm range, when appropriate ionic strength corrections for the activity coefficients of aqueous species are applied with SIT. For pHm < 6, Eu(III) sorption to illite decreases due to its competition with Na⁺ for ion exchange sites whereas, for pHm > 6, Eu(III) uptake is weakly affected by the ionic strength because of the formation of innersphere Eu-illite surface complex [3]. Pu(IV) uptake on illite is predicted to be only weakly affected by the ionic strength in the range 0.1 < m_NaCl < 3.2 m. Model calculations agree with the observed absence of significant ionic strength effects on experimental uptake data of Th(IV) (taken as actinide analogue of Pu(IV)) on montmorillonite between 0.1 and 1.0 M NaClO₄ [4], and the sorption data for Pu obtained in the present study for pHm > 6. These results support the assumption that tetravalent actinides undergo innersphere complexation rather than ion exchange at illite surface.

As previously inferred for actinide uptake on clays, the redox speciation of actinides is influenced by the formation of surface complexes [21-23]. The Pu(IV)/Pu(III) borderline at the illite surface (denoted as \{Pu(IV)/Pu(III)\}_{surf}) is expected to depend on the individual Pu(IV) and Pu(III) sorption behavior [22-23], and can be calculated for m_NaCl = 0.1, 1.0 and 3.2 m (Fig. 3) according to the following equation:

\[
\{\text{Pu(IV)/Pu(III)}\}_{\text{surf}} = \{\text{Pu(IV)/Pu(III)}\}_{aq} + (\log R_d(\text{Pu(III)}) - \log R_d(\text{Pu(IV)}))
\]

\{Pu(IV)/Pu(III)\}_{aq} is the Pu(IV)/Pu(III) borderline in solution (i.e. plotted as a black line in Fig. 3), which is calculated using the Nernst equation. Values of log R_d(Pu(III)) and log R_d(Pu(IV)) in eq. 5 represent the respective individual predicted uptake of the two Pu oxidation states under the same physico-chemical conditions (same pH, ionic strength, etc), which can be seen in Figure 1a,b,c (“Pu(IV) only” and “Pu(III) only”). For all investigated m_NaCl and pHm > 6 log R_d(Pu(III))
\[ \approx \log R_d(\text{Pu(IV)}) \] and, as a consequence, \( \{\text{Pu(IV)/Pu(III)}\}_{aq} \) and \( \{\text{Pu(IV)/Pu(III)}\}_{surf} \) coincide (Fig. 3). For pH < 6, \( \{\text{Pu(IV)/Pu(III)}\}_{aq} > \{\text{Pu(IV)/Pu(III)}\}_{surf} \) because of the stronger sorption of Pu(IV) as compared to that of Pu(III). With increasing \( m_{NaCl} \), \( \log R_d(\text{Pu(IV)}) \) remains constant whereas \( \log R_d(\text{Pu(III)}) \) decreases. Hence, \( \{\text{Pu(IV)/Pu(III)}\}_{surf} \) decreases with increasing \( m_{NaCl} \) (i.e. the Pu(IV)/Pu(III) borderline at the illite surface is shifted to lower pe). According to this model, when the redox conditions (pH/pe) fall between \( \{\text{Pu(IV)/Pu(III)}\}_{aq} \) and \( \{\text{Pu(IV)/Pu(III)}\}_{surf} \), Pu(III) prevails in solution whereas Pu(IV) prevails at the illite surface, and the overall Pu uptake by illite is intermediate to the individual uptake of Pu in the two distinct oxidation states [22,23].

When different Pu redox states coexist, the overall uptake of actinides on clays highly depends on pe [21-23]. As pointed out before, \( R_d \) data scatter at acidic pH\textsubscript{m}, which might be due to slight variations in redox conditions in individual batch experiments. The latter finding can be explained by the fact that the concentration of redox active components in the system is obviously low as discussed in a previous study on the interaction of Np with the same illite [21].

The exact nature of the redox partners determining measured pe values is not quite clear, but is most likely related to structurally bound Fe(II)/Fe(III). By using measured pH\textsubscript{m} and pe values as model input parameters, overall Pu uptake onto illite is calculated using the 2 SPNE SC/CE model and SIT without further parameter adjustment. Model results are plotted as empty symbols on Fig. 1a,b,c. In addition, Fig. 4 shows predicted versus experimental \( \log R_d \) values. Both figures show good agreement of predicted and experimental Pu uptake onto illite. The average deviation between predicted and experimental results equals 0.3 \( \log R_d \) units (shown as bold dashed lines), which corresponds to the experimental uncertainty. Larger deviation (maximum 1 \( \log R_d \) units, shown as thin dashed lines) can be observed for some data, especially for lowest \( \log R_d \) values, which might be attributed to uncertainties in pe measurements (±0.8). Indeed, these
data were obtained in redox conditions (pH/pe) between \{\text{Pu(IV)/Pu(III)}\}_{aq} and \{\text{Pu(IV)/Pu(III)}\}_{surf} (3 < \text{pH}_m < 6; \text{Fig. 3}), for which predicted log R_d values highly depend on pe \cite{21-23}. Therefore, it is possible to conclude that the present model can accurately predict Pu sorption to illite under a large range of pH_m, pe and ionic strength conditions.

4. Conclusions

In this study, we provided experimental data for Pu sorption to illite under slightly reducing conditions (2 < pe < 10), where Pu(IV)/Pu(III) couple is involved, under saline conditions (up to 3.2 m NaCl), which are relevant for several potential deep geological nuclear waste repository sites. We observed that Pu sorption to illite was ionic strength dependent for pH_m < 6 because of the presence of Pu(III) and its competition with Na^+ for ion exchange sites. By contrast, Pu sorption to illite did not depend on the ionic strength for pH_m > 6, due to the predominance of Pu(IV), which undergoes innersphere surface complexation. According to our calculations, although Pu redox speciation in solution is only slightly influenced by NaCl concentration, Pu(IV)/Pu(III) redox equilibria at the illite surface can significantly be affected by the ionic strength when pH_m < 6. Overall Pu uptake could be accurately predicted by the 2 SPNE SC/CE model coupled to SIT. Results of the present work together with the outcome of previous investigations \cite{3,21-23} demonstrate that the proposed modeling approach is robust and applicable, at least for the investigated systems, and might be a reliable predictive tool for performance safety assessment for clays under highly saline conditions. This recently developed approach now covers effects of pH, redox reactions and high background electrolyte concentrations, and allows estimating actinide sorption to clays at various redox conditions. Even
at relatively low concentrations of redox governing components, measured PE still appears to be a meaningful parameter, which can be used to estimate retention of redox sensitive actinide ions and their redox speciation in solution and at the mineral surface with a quasi-mechanistic sorption model.

Nevertheless, the applicability of the approach to natural systems characterized by a high degree of heterogeneity and complexity still has to be demonstrated. Further studies dedicated to the uptake of actinides on natural soil- or clay rock formation-\textit{in situ} porewater systems are therefore required to validate the present modeling approach. Our approach might also be tested for redox sensitive elements other than actinides (e.g. cerium, selenium, arsenic, chromium, iron) and adsorbing phases other than clays (e.g. oxides, natural organic matter).

Finally, non-electrostatic models were recently shown to be particularly suitable for the prediction of metal ion sorption to various types of surfaces in brines [3, 9-11], which is further demonstrated in this study, and would find applications not only in the field of nuclear waste disposal but also, for instance, in marine chemistry.

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\section*{References}
Figure Caption

Fig. 1. Pu sorption to illite (log $R_d$ in L kg$^{-1}$) versus pH$_m$ (= - log $m_{H^+}$) in (a) 0.1 (taken from our previous work [23]), (b) 1.0 and (c) 3.2 m NaCl. (d) Experimental Pu sorption data in 0.1 < $m_{NaCl}$ < 3.2 m are plotted together for comparison. Experimental Eu(III)-illite sorption data for similar $m_{NaCl}$ ((a) 0.1, (b) 0.9 and (c) 3.9 m NaCl) are shown for comparison. Solid and dashed lines show the predicted uptake of Pu(IV) (taken as independent component) and Eu/Pu(III), respectively. Empty symbols are model overall Pu sorption results using measured pH/pe values as input parameters.

Fig. 2. Experimental data for Pu sorption onto illite (log $R_d$, in L kg$^{-1}$) versus the final Pu concentration in solution after phase separation (log [Pu]$_{aq}$, in M) for pH$_m$ > 6 and 0.1 < $m_{NaCl}$ < 3.2 m (data for 0.1 m NaCl are taken from our previous work [23]). The average log $R_d$ (= 5.3) is shown as a line, associated with an uncertainty of ± 0.3 (1σ; dashed lines).

Fig. 3. The pe values measured in the illite suspensions in $m_{NaCl}$ = 0.1, 1 and 3.2 m are plotted versus pH$_m$ (= - log $m_{H^+}$). Data for 0.1 m NaCl are taken from our previous work [23]. The black lines correspond to the predominance pH$_m$-pe diagram for Pu in solution calculated for $m_{NaCl}$ = 0.1 m and [Pu]$_{tot}$ = 10$^{-11}$ M. The thick colored lines correspond to the Pu(IV)/Pu(III) borderline at the illite surface for $m_{NaCl}$ = 0.1 m (blue), 1.0 m (green) and 3.2 m (red).

Fig. 4. Modeled versus experimental log $R_d$ values for the whole dataset (present data for $m_{NaCl}$ = 1.0 and 3.2 m and from our previous work for $m_{NaCl}$ = 0.1 m [23]). The 1:1 (solid) line, associated with an uncertainty of ± 0.3 log $R_d$ units (bold dashed lines), which corresponds to both the experimental uncertainty on log $R_d$ and the average deviation between experimental and modeled results, and an uncertainty of ± 1 log $R_d$ units (thin dashed lines), which corresponds to the largest deviation between experimental and modeled results.
Fig. 1

(a) $m_{NaCl} = 0.1$ m

(b) $m_{NaCl} = 1.0$ m

(c) $m_{NaCl} = 3.2$ m

(d) $m_{NaCl} = 0.1$ m, $m_{NaCl} = 1.0$ m, $m_{NaCl} = 3.2$ m
Fig. 2

\[ \log R_d (L \text{ kg}^{-1}) = 5.3 \pm 0.3 \]

- \( m_{NaCl} = 0.1 \text{ m} \)
- \( m_{NaCl} = 1.0 \text{ m} \)
- \( m_{NaCl} = 3.2 \text{ m} \)

\[ \log [ Pu]_{aq} (M) \]
Fig. 3

- $\text{Pu}(\text{III})_{(aq)}$
- $\text{Pu}(\text{IV})_{(aq)}$
- $\text{Pu}(\text{V})_{(aq)}$

$[\text{NaCl}] = 0.1 \text{ m}$
$[\text{NaCl}] = 1.0 \text{ m}$
$[\text{NaCl}] = 3.2 \text{ m}$

$O_{2(g)} > 0.2 \text{ atm}$
$H_{2(g)} > 1 \text{ atm}$
Fig. 4

![Graph showing log R_d (model) versus log R_d (experiment) for different NaCl concentrations. The graph includes data points for m_{NaCl} = 0.1 m, m_{NaCl} = 1.0 m, and m_{NaCl} = 3.2 m.](image-url)