THE INFLUENCE OF URANYL HYDROLYSIS AND MULTIPLE SITE-BINDING REACTIONS ON ADSORPTION OF U(VI) TO MONTMORILLONITE

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Abstract - Adsorption of uranyl to SWy-1 montmorillonite was evaluated experimentally and results were modeled to identify likely surface complexation reactions responsible for removal of uranyl from solution. Uranyl was contacted with SWy-1 montmorillonite in a NaClO₄ electrolyte solution at three ionic strengths (I = 0.001, 0.01, 0.1), at pH 4 to 8.5, in a $N_{2(g)}$ atmosphere. At low ionic strength, adsorption decreased from 95% at pH 4 to 75% at pH 6.8. At higher ionic strength, adsorption increased with pH from initial values less than 75%; adsorption edges for all ionic strengths coalesced above a pH of 7. A site-binding model was applied that treated SWy-1 as an aggregate of fixed-charge sites and edge sites analogous to gibbsite and silica. The concentration of fixed-charge sites was estimated as the cation exchange capacity, and non-preference exchange was assumed in calculating the contribution of fixed-charge sites to total uranyl adsorption. The concentration of edge sites was estimated by image analysis of transmission electron photomicrographs. Adsorption constants for uranyl binding to gibbsite and silica were determined by fitting to experimental data, and these adsorption constants were then used to simulate SWy-1 adsorption results. The best simulations were obtained with an ionization model in which AlOH₂⁺ was the dominant aluminol surface species throughout the experimental range in pH. The pH-dependent aqueous speciation of uranyl was an important factor determining the magnitude of uranyl adsorption. At low ionic strength and low pH, adsorption by fixed-charge sites was predominant. The decrease in adsorption with increasing pH was caused by the formation of monovalent aqueous uranyl species, which were weakly bound to fixed-charge sites. At higher ionic strengths, competition with Na+ decreased the adsorption of UO22+ to fixed-charge sites. At higher pH, the most significant adsorption reactions were the binding of UO₂²⁺ to AlOH and of (UO₂)₃(OH)₅ to SiOH edge sites. Near-saturation of AlOH sites by UO₂²⁺ allowed significant contributions of SiOH sites to uranyl adsorption.

Key Words-Montmorillonite, Uranyl.

INTRODUCTION

Uranium is a widespread environmental contaminant resulting from mining and manufacturing activities related to nuclear power and weapons production. At U.S. Department of Energy facilities, for example, uranium is the most common radionuclide contaminant in groundwater/sediment systems (Riley *et al* 1992). The mobility of uranium is enhanced in oxidizing environments by the formation of uranyl (UO_2^{2+}) , which hydrolyzes to form a number of aqueous hydroxy species. The hydrolysis reactions considered here and their equilibrium constants (Grenthe 1992) are listed in Table 1; computed aqueous speciation of UO_2^{2+} at I=0.1, over pH of 4 to 9 is illustrated in Figure 1.

In near-surface groundwater systems, chemical interactions with solid surfaces act to retard uranyl mobility. The adsorption of uranyl ion to solid substances has been studied as a mobility-limiting process, particularly with respect to sorption by Fe oxides (Payne and Waite 1991, Hsi and Langmuir 1985, Ho and Miller 1985, Tripathi 1984, Maya 1982) and phyllosilicate minerals (Borovec 1981, Tsunashima *et al* 1981, Lieser *et al* 1992). In addition to constraining the mobility of contaminant uranium associated with mine tailings sites

(Bonotto 1989, Thomson et al 1986, Morin et al 1982, Morin and Cherry 1988), sorption reactions are thought to be an important component of the formation of economic uranium deposits (Fiala 1988, Giblin 1980). Quantitative modeling of uranyl adsorption to natural substrates has been limited to oxides and oxyhydroxides (e.g., Hsi and Langmuir 1985, Payne and Waite 1991); iron oxyhydroxides in particular have been studied because they have high adsorption capacity and are common accessory minerals or coatings in sediments and soils where uranyl adsorption reactions are significant. Iron oxyhydroxides are, however, only a small subset of the reactive constituents found in soils and sediments.

In this study we investigate the adsorption of uranyl to Wyoming montmorillonite SWy-1, a well-studied reference smectite. Smectites are ubiquitous 2:1 sheet silicates found as clay-sized components of soils and sediments, consisting of octahedrally coordinated aluminum atoms sandwiched between tetrahedrally coordinated silicon atoms. Cation substitutions within the crystal lattice can occur in either the octahedral (Mg²⁺ for Al³⁺) or tetrahedral (Al³⁺ for Si⁴⁺) sheets, resulting in a fixed negative charge that dominates the mineral's cation exchange capacity (CEC). Partially coordinated Al and Si are exposed at crystallite edges,

Table 1. Equilibrium constants.

Reaction	Log K	
Aqueous speciation		
$UO_2^{2+} + H_2O = UO_2(OH)^+ + H^+$	-5.20^{1}	
$2UO_2^{2+} + 2H_2O = (UO_2)_2(OH)_2^{2+} + 2H^+$	-5.62^{1}	
$3UO_2^{2+} + 5H_2O = (UO_2)_3(OH)_5^{+} + 5H^{+}$	-15.55^{1}	
$UO_2^{2+} + 3H_2O = UO_2(OH)_3^{-} + 3H^{+}$	-21.0 ¹	
Ion exchange reactions		
$UO_2^{2+} + 2X^- = UO_2X_2$	$30.9^{2,3}$	
$UO_2(OH)^+ + X^- = UO_2(OH)X$	8.63	
$(UO_2)_3(OH)_5^+ + X^- = (UO_2)_3(OH)_5X$	-1.75^{3}	
Edge-site reactions		
$SiOH + H^+ = SiOH_2^+$	-0.95^{4}	
$SiOH = SiO^- + H^+$	-6.95^{4}	
$SiOH + Na^+ = SiO^ Na^+ + H^+$	-6.6^{4}	
$SiOH + UO22+ = SiO \cdot UO2+ + H+$	0.1465	
$SiOH + (UO_2)_3(OH)_5^+ = SiO \cdot (UO_2)_3(OH)_5 + H^+$	-16.80^{5}	
$AlOH + H^+ = AlOH_2^+$	12.35	
$AlOH = AlO^- + H^+$	-13.6^{5}	
$AlOH + Na^+ = AlO^ Na^+ + H^+$	-10.37^{5}	
$AIOH + UO22+ = AIO \cdot UO2+ + H+$	7.10 ^s	
$AlOH + (UO_2)_3(OH)_5^+ = AlO \cdot (UO_2)_3(OH)_5 + H^+$	-15.8^{5}	

¹ Grenthe, 1992.

which hydrolyze to form aluminol (AlOH) and silanol (SiOH) groups (White and Zelazny 1988). These hydroxylated sites exhibit acid/base behavior and coordinative properties analogous to those of their component oxides [i.e., SiO₂, Al(OH)₃]; recent experimental and modeling studies have attempted to quantify the contributions of edge sites to metal ion retention by smectites and other layer silicates (Fletcher and Sposito 1989, Zachara et al 1993, Stadler and Schindler 1993, Zachara and Smith 1994).

The use of surface-site concentrations, surface-binding reactions, and adsorption constants as fitting parameters in adsorption modeling can lead to successful but underconstrained models of experimental systems. The studies of Hsi and Langmuir (1985) and Payne and Waite (1991), for example, each described uranyl adsorption to iron oxyhydroxides by trial and error fitting of the experimental data, but the surface-binding reactions and constants used did not agree between the two studies; the limitations of this approach were recognized in the latter study.

The system studied here included a heterogeneous suite of surface sites and multiple aqueous species that could potentially bind at the solid surface; the abundance of adjustable parameters in this system could lead to a poorly constrained chemical model. Our objective was to construct a chemical model of uranyl/smectite interactions with a minimum number of parameters, using measurements of site concentrations

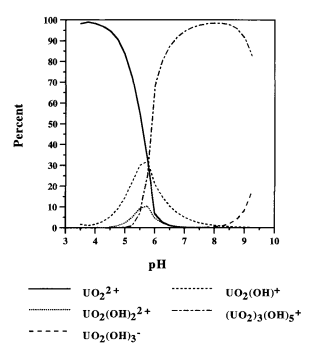


Figure 1. Uranyl speciation at varying pH. Calculations are for 8×10^{-6} mol L^{-1} uranyl in solution of ionic strength 0.1. Speciation constants from Grenthe (1992).

² Fixed relative to Na+ constant.

³ Calculated as explained in the text.

⁴ Zachara and Smith 1994.

⁵ This study, Model 1.

and adsorption constants that were independent of the experimental smectite adsorption data. A multiple-site adsorption model containing fixed-charge (X-) and hydroxylated edge sites (AlOH, SiOH), similar to those developed by Schindler et al (1987) for kaolinite, and by Stadler and Schindler (1993) for SWy-1, was used to compute aqueous speciation and uranyl adsorption to surface sites. We have applied assumptions of likely surface reactions on the analogue solids, silica and gibbsite, to derive adsorption constants describing binding reactions to comparable sites on the edge of the clay. Although the identity of responsible surface chemical reactions cannot be determined definitively through macroscopic modeling, our approach constrains the model to a greater degree than would direct parameter fitting.

MATERIALS AND METHODS

Sorbent preparation

The less than 2- μ m size fraction of SWy-1 was separated from the bulk material and pretreated with 0.001 N HClO₄ to remove surface precipitates, with 3% hydrogen peroxide (H₂O₂) to remove organic matter, and washed repeatedly with 0.1 mol L⁻¹ NaClO₄. The resultant stock suspension was stored under ultraviolet light to suppress microbial activity. The proportions of clay, salt, and H₂O in the suspensions were determined according to the procedure described by Sposito et al (1981).

Gibbsite [Al(OH)₃, ALCAN SF-4, Alcoa] was pretreated by first shaking a suspension of 5 g of gibbsite in 20 ml of 0.01 N NaOH for 30 min, followed immediately by seven successive washes with 0.01 mol L⁻¹ NaClO₄. Silica (SiO₂, Aerosil 200, Degussa, Inc.) was placed in suspension at the desired density and equilibrated by shaking overnight.

Chemical and physical measurements

The external surface area of the solid reactants was measured by multipoint N₂ adsorption. Particle sizes, used to calculate smectite surface areas and edge-site concentrations for modeling purposes, were determined using a Phillips 300 transmission electron microscope (TEM). Smectite particles from dilute suspensions were affixed to carbon-coated copper grids. Grids were examined at high magnification to select the optimum resultant particle spacing (i.e., a high particle density with a minimal number of overlapping particles). Several such observation fields were photographed. The images were digitized, particle overlaps were removed by editing the digital image (less than 5% of the total number of grains overlapped), and the mean particle size and equivalent particle diameter (the diameter of a circle with equivalent area) were calculated using image acquisition and processing software. Over 200 particles were analyzed by this process.

Acid-base titration of gibbsite was done by a batch method, in which replicate suspensions of gibbsite were equilibrated with known masses of added acid and base. Gibbsite suspensions were prepared at a massto-liquid ratio of 90 g L⁻¹ (300 m² L⁻¹) and equilibrated over a pH range of 4.5 to 9.5 using carbonatefree NaOH and HClO4. After measurement of equilibrium pH values, suspension aliquots were filtered through 1.8-nm cone filters, and the filtrate was acidified with ultrapure HCl and analyzed for dissolved aluminum using an inductively coupled plasma-mass spectrometry system (with a detection limit of 0.1 ng ml⁻¹) to determine the mass of gibbsite dissolved during the titration. Blank titration data were collected and modelled using FITEQL (Westall 1982a,b, 1993), and experimental data were corrected for aqueous protolysis effects using the results of that modeling.

Isotopic exchange (²²Na; Babcock and Schultz 1970) was used to determine the CEC of the sorbent over the pH range from 4.0 to 8.0 in the presence of 0.01 and 0.1 mol L⁻¹ NaClO₄. Suspensions of the Na⁺-saturated sorbent materials were equilibrated for 24 h after pH adjustment with NaOH or HClO₄. Following an additional 14-h equilibration period in the presence of ²²Na⁺, the suspensions were centrifuged at 30,000 rcf for 30 min; then supernatant ²²Na⁺ activity (scintillation counting), Na⁺ concentration (ion chromatography), and pH were determined.

Bulk chemical composition of SWy-1 was determined on a sodium-saturated sample that was dialyzed before freeze-drying to remove excess salt. Clay powder (500 mg) was pressed into a wafer for X-ray fluorescence determination of Al, Si, K, Ti, Mn, and Fe. The clay (50 mg) was mixed with 600 mg LiBO₂, fused at 900 °C, and dissolved in dilute HNO₃ for determination of Na and Mg by atomic absorption spectroscopy. Gravimetric moisture content was determined after heating the clay to 105 °C and again after heating it to 900 °C.

Adsorption experiments

Adsorption experiments were conducted under N_{2(g)} to preclude the complexation of uranyl by dissolved carbonate. Experimental conditions are summarized in Table 2. Suspensions were prepared using degassed electrolyte solutions diluted with deionized water to compositions of 0.001 mol L^{-1} to 0.100 mol L^{-1} NaClO₄. A mass of stock clay suspension containing the desired mass of SWy-1 was transferred to a centrifuge tube and washed three times in the electrolyte solution. The total mass of SWy-1 suspension, placed in a jacketed 500-ml reaction vessel, was adjusted to yield a final suspension charge concentration of 0.5 meq kg⁻¹, based on the measured CEC at pH 6. An autotitrator was used to monitor and adjust pH using 0.01 mol L⁻¹ NaOH or HClO₄. A glass pH electrode and calomel reference electrode (isolated from the sus-

Table 2. Experimental conditions for uranyl adsorption to SWy-1, gibbsite, and SiO₂.

Sorbent	Susp. (g kg ⁻¹)	S.A. (m ² g ⁻¹)	SiOH (µmol L ⁻¹)	AlOH (µmol L-1)	U (µmol L-1)	Na (mmol L ⁻¹)
SWy-1	0.59	31	11	9.9	8.7	96
	0.59		11	9.9	8.5	9.7
	0.59		11	9.9	8.6	4.0
SiO ₂	0.13	180	18		9.0	91
	0.15		21		9.4	9.1
	0.013		1.8		8.9	97
Gibbsite	5.6	3.4		19	9.1	1.2
	5.7			19	9.3	95
	0.55			1.9	9.1	98

pension by an agar bridge) were used to minimize contamination of the suspension by KCl. The stirred suspension was allowed to equilibrate at pH 4.0 for 16 h before adding uranyl.

A uranyl nitrate [UO₂(NO₃)₂ 6H₂O; Fluka Chemical Co., greater than 99.0% purity] stock solution containing 195 mg U kg-1 was added by mass to the equilibrated SWy-1 suspensions to yield an initial U concentration of 2 mg L^{-1} (8.4 × 10⁻⁶ mol L^{-1}). The pH of the suspension was increased incrementally by addition of NaOH, and duplicate 15-ml samples were removed at each 0.5-pH-unit step. The samples were equilibrated for 20 h in a controlled-environment shaker (150 rpm, 25°C). The equilibrated samples were centrifuged at 5,000 rcf for 1 h, and separate 4-ml samples of the supernatant were removed for determination of pH, solution-phase uranyl concentration, and Na concentration. The final pH of the supernatant was measured under a N₂ atmosphere using a combination electrode. The quantity of uranyl adsorbed was calculated from the difference between initial and final solutionphase uranyl concentrations. Uranium concentration was determined using a kinetic phosphorescence analyzer (Model KPA-11, CHEMCHEK Instruments, Inc.); the detection limit for these analyses was $0.3 \mu g$ U L⁻¹, with an analytical precision of 0.09 μ g U L⁻¹. These limits compare to a range in experimental samples of 3 to 2,000 μ g U L⁻¹.

The adsorption behavior of uranyl on gibbsite and silica was determined using an experimental approach identical to that described above for uranyl adsorption to SWy-1. Two suspension concentrations, differing by approximately one order of magnitude for each adsorbent (described below), were used to determine the extent of adsorption under conditions of different aqueous uranyl speciation.

RESULTS

Chemical and physical properties

The elemental analysis (Table 3), reduced using the method of Newman and Brown (1987), yielded a 2:1 clay-mineral structural formula of (Ca_{0.001}Na_{0.707}-

Table 3. Physical and chemical properties of Na-saturated SWy-1 montmorillonite.

Composition:	-
SiO ₂ , wt. %	61.6
$Al_2\tilde{O}_3$	21.2
Fe ₂ O ₃	4.29
TiO_2	0.08
MnŌ	0.03
MgO	2.80
CaO	0.01
Na ₂ O	2.86
K_2O	0.02
H_2^- O	<u>_7.1_</u>
Total	99.96

Mean equivalent particle diameter: 0.48 μm

Surface area: 31 m² g⁻¹

 $K_{0.003}$)[Al^{3.04}Fe(III)_{0.41}Mg_{0.532}](Si_{7.85}Al_{0.147})O₂₀OH₄ (montmorillonite).

The CEC of Swy-1 shows a relatively small dependence on pH and ionic strength (Figure 2); CEC at pH 4 is $0.63 \text{ mmol}_{c} \text{ g}^{-1}$; CEC at pH 10 is $0.75 \text{ mmol}_{c} \text{ g}^{-1}$. Changes in CEC with pH result from the additive contribution of fixed-charge and ionized edge sites; i.e., SiOH and AlOH (White and Zelazny 1988; Sposito 1989). The dissolution and readsorption of Al³⁺ at lower pH may also contribute to this trend in CEC. A calculation based on composition and unit cell parameters (Newman 1987) gives a fixed charge (pH-independent CEC) of 0.95 mmol_c g⁻¹ arising from interlayer cation charge. Our measurement gives 0.74 mmol_c g⁻¹ in 0.01 mol L^{-1} to 0.1 mol L^{-1} NaClO₄ at pH 6.0; this value was used as the unit-mass concentration of fixedcharge sites in model calculations. The discrepancy between theoretical and experimental determinations of CEC is unexplained.

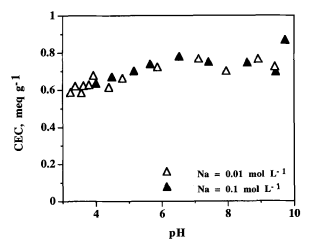


Figure 2. Cation exchange capacity of treated SWy-1 at varying pH and two ionic strengths.

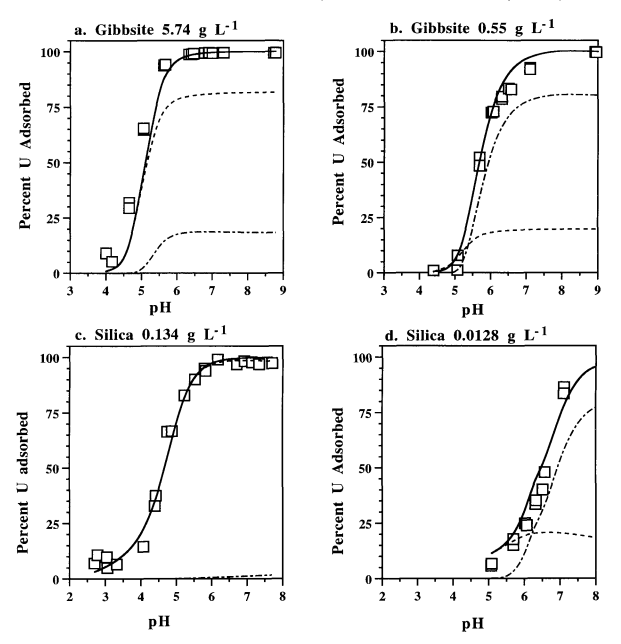


Figure 3. Experimental results of uranyl interaction with gibbsite (a,b) and silica (c,d). Total computed contributions to adsorption=solid line; monomeric (UO_2^{2+}) surface complex=dashed line; trimeric $[(UO_2)_3(OH)_5^+]$ surface complex=dot-dashed line.

Adsorption experiments

Suspension concentrations of gibbsite and silica at higher solid-to-solution ratios (Table 2) were sufficient to produce greater than 50% adsorption of uranyl below a pH of 5.2 (Figure 3), where UO₂²⁺ predominated in solution (Figure 1). Experiments at a solids concentration of one-tenth the higher value had lower adsorption at equivalent pH; i.e., the adsorption edge was

shifted to higher pH where multiple, hydrolyzed uranyl species existed. In an investigation of uranyl adsorption to silica gel (SiO₂:xH₂O), Lieser *et al* (1992) found an adsorption maximum at approximately pH 6.7. Data from Allard *et al* (1983) were calculated by Lieser *et al* (1992) to yield similar results for Al₂O₃. Lieser *et al* (1992) concluded that decreased adsorption at elevated pH, similar for differing adsorbents, showed the effects of changes in aqueous uranyl speciation toward pre-

dominance of an anionic species [(UO₂)₃(OH)₇⁻)]. In contrast, our experiments showed a continued increase in adsorption at pH above 6.7, suggesting that the formation of an anionic species was not a significant determinant of overall adsorption results.

The adsorption of uranyl by SWy-1 varied with pH and ionic strength (Figure 4). Adsorption was most dependent on ionic strength at low pH, with adsorption edges coalescing at all ionic strengths above pH 6.5. This behavior may result from adsorption to two classes of sites on the solid surface: fixed-charge sites sensitive to competitive exchange between sodium and uranyl ions, and edge sites to which uranyl is specifically adsorbed without significant competition from sodium.

Hydrolysis of UO₂²⁺ may also affect adsorption. (In the remainder of the text, we use "uranyl" to refer to UO₂²⁺ and its hydrolysis products without differentiation, and restrict the use of "UO22+" to mean unhydrolyzed uranyl ion.). At low ionic strength, for example (Figure 4), the fractional adsorption of uranyl decreased over the pH range of 4 to 6.5. This decrease was paralleled by a reduction in the aqueous concentration of UO22+ in favor of UO2(OH)+ and (UO₂)₃(OH)₅+ (Figure 1). One may speculate that ion exchange was the predominant adsorption mechanism for uranyl over this pH range, and that the divalent UO22+ ion competed with Na+ more effectively than did the monovalent uranyl hydroxide ion $[UO_2(OH)^+]$. This hypothesis parallels the exchange reaction modelled by Stadler and Schindler (1993) with respect to copper adsorption to SWy-1. At slightly alkaline pH, copper forms a single hydrolyzed species that is analogous to $UO_2(OH)^+$ (CuOH+; Log K = -8.29); in nonpreference exchange neither monovalent species is significantly removed from solution in the presence of excess Na+. In contrast, the ion exchange of weakly hydrolyzing divalent metals, such as Cd2+, which does not form a monovalent complex over the pH range of interest, does not decrease with increasing pH unless other competitive sites, sorbents, or ligands exist (Zachara et al 1993, Zachara and Smith 1994).

ADSORPTION MODELING

The smectite surface was considered to consist of three discrete sites: fixed-charge ion-exchange sites (X⁻) on the smectite basal plane, and hydroxylated SiOH and AlOH edge sites. The chemical modeling program FITEQL (Westall 1982a, b) was used to compute the contribution of each site to adsorption using the Triple-Layer Model (TLM, Davis and Leckie 1978, Sposito 1984, Hayes and Leckie 1987) for the amphoteric edge sites, and a half-reaction approach was used (Fletcher and Sposito 1989) for the fixed-charge sites. The integral capacitance densities, C₁ and C₂, were fixed at 1.2 and 0.20 F m⁻², respectively, for all TLM calculations. Consistent with the results shown in Figure 4,

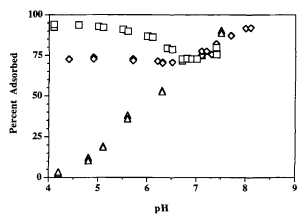


Figure 4. Combined adsorption results for uranyl binding to SWy-1. Suspension concentration was approximately 0.5 g L⁻¹; total uranium was 8.4 x 10^{-6} mol L⁻¹; the sodium perchlorate electrolyte concentration was varied: 0.001 mol L⁻¹ = squares, 0.01 mol L⁻¹ = diamonds, 0.1 mol L⁻¹ = triangles.

ion exchange to fixed-charge sites was hypothesized to dominate adsorption at low pH, while adsorption to edge sites was speculated to be more important at higher pH. Ion-exchange contributions to adsorption were evaluated first, then combined with edge-site complexation to construct a unified adsorption model.

Edge-site concentrations on SWy-1 (Table 2) were determined from TEM data and bulk composition using the method described in detail by Zachara and Smith (1994). In this method, the mean equivalent diameter of 0.48 μm , the bulk composition, and the edge-site density for montmorillonite calculated by White and Zelazny (1988) [N_{S,T} = 6.39 $\mu mol~m^{-2}$ and N_{S,O} = 5.32 $\mu mol~m^{-2}$, where N_{S,T} is the number of tetrahedral (SiOH) sites and N_{S,O} the number of octahedral (AlOH) sites] were used to estimate the total number of edge sites on SWy-1 at 3.52 \times 10⁻⁵ mol g⁻¹.

Derivation of constants

Reactions with fixed-charge sites. Exchange constants for UO_2^{2+} and its monovalent hydrolysis species $[UO_2OH^+$ and $(UO_2)_3(OH)_5^+]$ were calculated using the equation of Fletcher and Sposito (1989):

$$K_1 = [K_a K_2^{Za} (a_a^{Zb}/a_b^{Za})(m_{aX} + m_{bX})^{(Zb-Za)}]^{1/Zb}$$
 (1)

which was modified to yield

$$K_1 = [K_a K_2^{Za} (mX^-)^{(Zb-Za)}]^{1/Zb}$$
 (2)

where K_1 describes the half-reaction equilibrium constant (mol L^{-1}) for interaction of a uranyl species of charge Z_a with fixed-charge X^- sites (molal concentration), in competition with an ion of charge Z_b whose binding is described by K_2 . The value of log K_2 was arbitrarily fixed at 13.8 for computation of K_1 values.

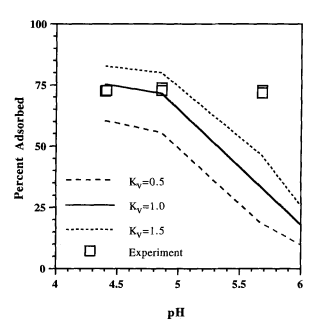


Figure 5. The effect of different values for the cation exchange selectivity constant (K_v) on calculated uptake of uranyl by fixed-charge exchange sites. A non-selective constant (K_v) = 1) best reproduced the experimental results at pH < 5.

The constant Ka is equivalent to the Vanselow selectivity coefficient, K_v, when the ratio of surface activity coefficients (a_a and a_b) is unity. Because the total X concentration is approximately 500 times that of total uranyl, mX is an appropriate approximation of $(m_{ax}+m_{bx})$, the summed molal concentrations of X⁻ site complexes derived from ions of different charge; mX_{Total} was not significantly changed by proportionation of X⁻ sites between complexes with univalent and divalent ions. In our computations, nonpreference exchange $(K_v = K_a = 1.0)$ was assumed between Na and all of the uranyl species. The validity of this assumption was evaluated by comparing calculated and experimental results (NaClO₄ = $0.01 \text{ mol } L^{-1}$) using K, values of 0.5, 1.0, and 1.5 (Figure 5). Nonpreference exchange was found to most closely approximate lowpH results for experiments at intermediate ionic strength. Although nonpreference exchange was used as an approximation of uranyl interaction with fixedcharge sites at all ionic strengths, exchange selectivity showed a small dependence on ionic strength. Experimental results were matched more closely using a slight preference for Na+ at lower ionic strength (0.001 M Na) and a slight preference for UO22+ at higher ionic strength (0.1 M Na).

Reactions to edge sites. Gibbsite and silica surface sites were considered as analogues of the smectite edge. For gibbsite, acidity constants were fit to gibbsite acid-base titration data, then uranyl adsorption constants were

fit to uranyl adsorption edges. Published acidity constants were used for the silica surface in fitting uranyl adsorption constants to silica/uranyl adsorption edges. The use of gibbsite as an analogue to smectite edge sites (AlOH) formed by octahedrally coordinated aluminum atoms is complicated by differences in surface chemistry and morphology. Gibbsite consists of octahedrally coordinated aluminum, but its surface is populated by doubly coordinated hydroxyl groups on the basal plane as well as singly coordinated groups on the crystal edge. The doubly coordinated hydroxyl groups are less reactive than edge groups, however, and have been treated as nonreactive sites with respect to proton uptake (Hiemstra et al 1987). In addition, the edge structure of gibbsite is not identical to that of the octahedral layer in smectites. Gibbsite edges are composed of singly coordinated hydroxyl groups, whereas the octahedral layer in smectites includes singly coordinated hydroxyls as well as bridging oxygens between aluminum and silicon (White and Zelazny 1988). We have treated gibbsite as an analogue of the edge sites exposed at the surface of the smectite octahedral layer by discounting the reactivity of doubly coordinated hydroxyl ions and by independently estimating the concentration of sites at smectite and gibbsite edges.

Reactivity of AIOH sites. Initial modeling calculations indicated that the ability to simulate uranyl adsorption to smectite at high ionic strength and pH was dependent on the ionization and electrolyte complexation constants used for the AIOH surface. The surface chemistry of gibbsite is difficult to define uniquely because of uncertainty regarding edge and basal site concentrations and reactivity and because of complications arising from Al dissolution. Accordingly, different sets of adsorption constants, site densities, and surface areas can provide equivalent fits to titration and adsorption data on gibbsite. We evaluated several self-consistent sets of gibbsite surface parameters for their ability to describe uranyl adsorption to SWy-1.

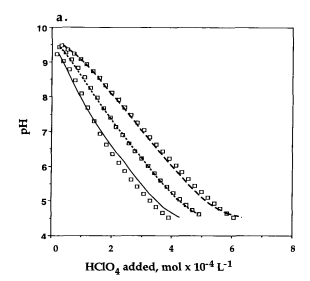
Three sets of surface parameters were obtained for the gibbsite used in the uranyl adsorption studies; one from Wieland et al (1993), and two from the fitting of acid-base titration data (Figure 6a). In evaluating titration data, the consumption of titrant from the dissolution of gibbsite was accounted for by fitting an operational K_{so} for Al(OH)₃ to pH-dependent Al concentrations in gibbsite suspensions (Figure 6b). The K_{sp} was used as a component within FITEQL to drive H+ consumption by dissolution and aqueous Al speciation reactions. The correction of titration data for dissolution of the solid had a small, but discernible, effect on fitted values for acidity constants. In the modeling, no distinction was made between basal plane Al₂OH and AlOH edge sites. The site concentration, [AlOH]_T, acidity constants, and electrolyte complexation constants were derived by modeling in two ways, described below.

The first set of parameters was derived from the titration data by simultaneously optimizing [AlOH]_T, K_{XOH}^{2+} , and K_{XOH}^{2+} — C_{IO4-} , then iteratively fitting K_{XOH}^{-} and K_{XOH}^{-} — K_{IO4-} and a log K_{IO4} (AlOH₂+ = AlOH + H⁺) of 12.3 (Model 1 calculations, Table 4). Our log K_{IO4} value was higher and our site concentration was lower than the single-p K_{IO4} result of Hiemstra et al (1987; pzc = 10, no correction for Al dissolution, with 54.2 μ mol sites g⁻¹).

The number of fitting parameters was reduced to three to arrive at a second set of parameters, by assuming that the intrinsic ion-pair formation constants for the electrolyte species $AlOH_2^+$ – ClO_4^- and AlO^- – Na^+ were equal and that and the ΔpK_a was 5. The fitting parameters were then $[AlOH]_T$, K_{a1} , and the combined electrolyte binding constant. The fitted site concentration was 23.9 μ mol sites g^{-1} , with a log K_{a1} of 7.87 (Model 2 calculations, Table 4). The site concentration was lower than that for Model 1 and the first acidity constant (K_{a1}) was close to the value (7.6) from Wieland *et al* (1993; Model 3 calculations, Table 4); Wieland *et al* (1993) assumed a higher site concentration for their gibbsite (74.5 μ mol g^{-1}).

Reactivity of SiOH sites. The ionization and electrolyte complexation constants used here (Table 1) were taken from Zachara and Smith (1994), refitted from the data of Riese (1982) for experiments conducted on α –SiO₂ in NaNO₃ electrolyte. Zachara and Smith (1994) described the adsorption of Cd²⁺ to SiOH edge sites on smectites, whereas Riese (1982) derived silica surface constants to model the adsorption of radium and thorium on quartz.

Uranyl adsorption to gibbsite and silica. The solution species UO₂²⁺, UO₂OH⁺, and (UO₂)₃(OH)₅⁺, which are predominant over the experimental pH range (Figure 1), were considered when fitting adsorption constants to silica and gibbsite. Surface reactions between these species and surface sites leading to inner-sphere surface complexes were considered singly, in pairs, and altogether for their ability to describe the adsorption data. First, adsorption constants were fit individually for the three solution species on gibbsite and silica using the high-solids adsorption data. Simulative modeling of the low-sorbent suspensions was then attempted, but each of the calculations with only a single adsorbing species either overpredicted or underpredicted actual uranyl adsorption. Iterative fitting of constants between the two suspension concentrations was then attempted for pairs of aqueous species; constants for UO₂OH⁺ with either of the other species could not simultaneously predict uranyl adsorption in both suspensions. However, for both adsorbents, uptake of ura-



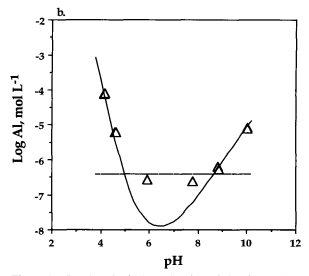


Figure 6. Results of acid/base titration of gibbsite at three ionic strengths (0.001, 0.01, and 0.1 mol L^{-1} NaClO₄. (a) Lines indicate data simulation where surface acidity constants, surface area, and electrolyte complexation constants were fit independently. (b) Aluminum dissolution observed at each experimental pH, with the conditional K_{sp} fit to experimental data (solid line); and the effective detection limit for dissolved aluminum (dashed line).

nyl could be well-described using UO_2^{2+} and $(UO_2)_3(OH)_5^+$ as adsorbing species (Figure 3). Addition of a UO_2OH^+ surface complex, to form a three-sorbate model, did not markedly change constants for the other species or significantly improve fits to or simulations of the experimental data.

The shift in adsorption to higher pH as the sorbent concentration decreased was accompanied by a reversal in the relative importance of each adsorbing species. Comparison of contributions to total adsorption (Model 1) at suspensions of 5.74 g L^{-1} gibbsite (Figure 3a)

Table 4. Gibbsite constants used in fitting uranyl adsorption constants $K_{UO_2^{2+}}$ and $K_{(UO_2)_{MOH_3^{+}}}$ Model 3 constants (other than uranyl adsorption constants) from Wieland *et al* (1993).

	Sites -	_		Log	••••		
	μmol g ⁻¹	K _{a1}	K _{a2}	K _a -	K _c +	K _{UO22+}	K _{(UO2)3(OH)5} +
Model 1	33.8	12.3	-13.16	10.15	-10.37	7.10	-15.80
Model 2	23.9	7.87	-12.87	10.38	10.36	6.25	-15.80
Model 3	74.5	7.6	-10.6	10.7	-7.3	4.50	-15.70

 $K_{a1} (AlOH + H^+ = AlOH_2^+).$

 K_{a2} (AlOH = AlO⁻ + H⁺).

 $K_{a^{-}}^{a2}(AlOH_{2}^{+} + ClO_{4}^{-} = AlOH_{2}^{+} - ClO_{-}).$

 K_c + (AlOH + Na⁺ = AlO⁻ - Na⁺ + H⁺).

 $K_{UO_2}^{2+}$ (AlOH + UO_2^{2+} = AlO· UO_2^{+} + H⁺).

 $K_{(UO_2)_3(OH)_5}^+$ [AlOH + $(UO_2)_3(OH)_5^+$ = AlO· $(UO_2)_3(OH)_5$ + H⁺].

and $0.55 \,\mathrm{g} \,\mathrm{L}^{-1}$ gibbsite (Figure 3b), for example, showed a computed predominance of the $\mathrm{UO_2}^{2+}$ surface complex in the higher-mass gibbsite suspension, and predominance of the $(\mathrm{UO_2})_3(\mathrm{OH})_5^+$ surface complex in the lower-mass gibbsite suspension. The relative importance of the two adsorbing species was dependent, in part, on aqueous uranyl speciation. At constant pH, as the suspension concentration of gibbsite decreased, the fractional abundance of the polymeric aqueous species $(\mathrm{UO_2})_3(\mathrm{OH})_5^+$ increased. At pH 5.7, for example, aqueous $(\mathrm{UO_2})_3(\mathrm{OH})_5^+$ was 1% of dissolved species at 5.74 g gibbsite L^{-1} , and 17% at 0.55 g gibbsite L^{-1} . The relationship of aqueous speciation to surface speciation for gibbsite was also observed for silica (Figure 3c, d).

Uranyl adsorption constants (Table 4) were determined for each set of surface parameters on gibbsite (Models 1, 2, and 3). Equivalent descriptions of experimental data were obtained using each set of surface parameters (Figure 7; 5.74 g L⁻¹ gibbsite). The effect of ionic strength on model results was evaluated by simulative calculations using fitted adsorption constants for Models 1, 2, and 3, in all of which the concentration of NaClO₄ was reduced from 0.1 to 0.001 mol L⁻¹. The results of these calculations are indicated as dashed lines in Figure 7. For Models 1 and 2, the

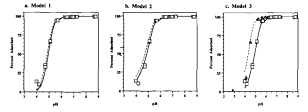


Figure 7. Experimental and calculated results of uranyl adsorption to gibbsite, 5.74 g L⁻¹. Results for each model are shown with calculations at NaClO₄ concentrations of 0.1 mol L⁻¹ (lines) and 0.001 mol L⁻¹ (dashes). Also shown for comparison with Model 3 calculations are experimental results of adsorption at two ionic strengths (squares = 0.1 mol L⁻¹, triangles = 0.001 mol L⁻¹).

effect of ionic strength was minimal (Figure 7a, b), as shown previously for the inner-sphere complex formalism using the TLM (Hayes and Leckie 1987). Calculations using Model 3, however, showed a significant increase in adsorption at lower ionic strength (Figure 7c). This increased adsorption resulted directly from the strength of the electrolyte anion surface complex in this model, which led to competition between the electrolyte anion (ClO₄⁻) and uranyl for surface sites at higher pH. Regardless of the chemical reality of this strong electrolyte ion surface complex, comparison with experimental data collected at a NaClO₄ concentration of 0.001 mol L⁻¹ (Figure 7c) suggested that Model 3 most closely approximated the gibbsite surface properties.

SWy-1 adsorption simulation

No adjustable parameters were used in simulating experimental results for uranyl adsorption to SWy-1 (Figure 8). The estimation of exchange site and edge site concentrations on SWy-1, from CEC measurements and clay-platelet diameters, resulted in good model approximations of experimental data using Model 1. Uranyl adsorption could be explained by the formation of surface complexes with fixed-charge (UO₂X₂), aluminol (AlO UO_2^+), and silanol [SiO $(UO_2)_3(OH)^+$] sites. These complexes were consistent with, but did not correspond closely to, the changes in aqueous uranyl speciation; e.g., the surface complex AlO·UO₂+ persisted above the pH where UO₂²⁺ predominated. Although a hydrolyzed surface species might be expected under conditions favoring hydrolysis in the aqueous phase, adsorbed species distinct from coexistent aqueous species may exist. Uranyl adsorbed to montmorillonite surfaces in particular has yielded spectroscopic evidence for lesshydrolyzed species at the mineral surface (Chisholm-Brause et al 1994). The relatively small number of significant surface complexes modelled here is also supported by spectroscopic evidence; X-ray absorption spectroscopy (Chisholm-Brause et al 1994) and optical spectroscopy (Morris et al 1994) have both shown the existence of three predominant uranyl species on the fixed-charge and amphoteric aluminol and silanol edge sites of montmorillonite. Dent *et al* (1992) detected uranyl bound to fixed-charge sites on the SWy-1 surface at pH less than 5.

TEM measurements, which obviated the need to use edge site concentrations as fitting parameters, yielded edge site concentrations that were different from fitted values obtained by other researchers. In a recent model of Cu2+ adsorption to SWy-1, Stadler and Schindler (1993) fitted edge-site concentrations to acid-base titration data because the binding of protons by individual surface sites could not be resolved experimentally. The AlOH concentrations of Stadler and Schindler (1993) were approximately three and one-half times the value obtained here, and the ratio of concentrations (AlOH: SiOH) was 1.68, twice the theoretical ratio (0.83; White and Zelazny 1988); an independent determination of site concentrations may have produced different results for acidity constants and Cu2+ binding constants.

Adsorption to exchange sites, treated identically in all three models, was affected by aqueous uranyl speciation (Figure 1) and the Na concentration in the electrolyte, as illustrated by Model 1 results (Figure 8). As pH increased at low Na concentration, fixed-charge uranyl complexes declined rapidly in concentration as UO_2^{2+} was replaced in solution by monovalent species, and more abundant Na⁺ successfully competed for fixed-charge sites. As Na concentration increased, the effect of aqueous uranyl speciation and competition was increased as Na⁺ displaced uranyl through non-preference exchange, even where UO_2^{2+} was the dominant aqueous species.

Three issues warrant discussion in relation to the edge-site model calculations presented in Figure 8: 1) the application of different model assumptions to simulations of experimental data; 2) the potential effects of edge site saturation and site coverage on model results; and 3) the effects of ionic strength on model and experimental results.

Model 1 and 2 calculations of adsorption to SWy-1 had differences that resulted from the way in which gibbsite titration data were modeled (Table 4). Gibbsite surface constants used to derive the uranyl adsorption constants for Model 1 included a relatively low [AlOH]_T (18.76 μ mol sites L⁻¹ at gibbsite = 0.555 g L⁻¹), and a high K_{a1} (log $K_{a1} = 12.3$), resulting in a high value of the binding constant for AlO UO₂+. In applying Model 1 to SWy-1 data, the gibbsite surface acidity, electrolyte adsorption, and uranyl adsorption constants were used, but the AlOH site concentration (9.9 µmol sites L^{-1}) was determined by calculations on TEM measurements of SWy-1 particle size. Although the value for the AlO UO₂+ adsorption constant was the highest of the three models (log K = 7.10), the high acidity constants specified a smectite edge that was

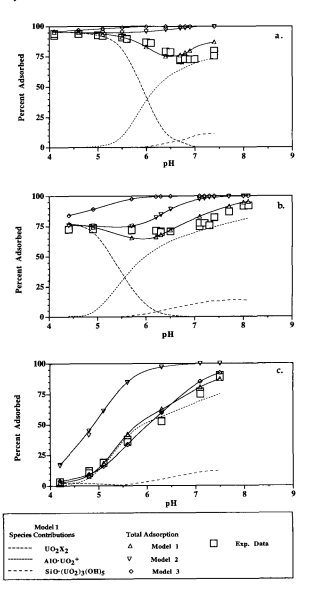


Figure 8. Model results for uranyl/SWy-1 interaction. Total predicted uptakes for constants derived by three methods are indicated (see text). Contributions to adsorption from UO₂X₂, AlO·UO₂+, and SiO·(UO₂)₃(OH)₅ are shown for the calculations that best predicted adsorption at all three ionic strengths (Model 1, see text).

positively charged throughout the experimental range in pH. Model 1 results simulated the SWy-1 experimental data best at all three ionic strengths. Gibbsite surface constants used to derive uranyl binding constants for Model 2 included still lower [AlOH]_T (13.27 μ mol sites L⁻¹ at gibbsite = 0.555 g L⁻¹), and a relatively low value for K_{a1} (log K_{a1} = 7.87); the resultant AlO·UO₂+ binding constant was approximately 0.9 log units lower in value than the Model 1 constant. When Model 2 was applied to SWy-1 data, the combination of a K_{a1} that was more than 5 log units lower than that

for Model 1, and an AlO·UO₂+ binding constant that was still relatively high, resulted in overprediction of uranyl binding at all ionic strengths. The published gibbsite surface constants (Wieland et al 1993) resulted in a high [AlOH]_T (41.34 μ mol sites L⁻¹ at gibbsite = 0.555 g L⁻¹) with surface acidity constants that were similar to those for Model 2; the resultant AlO·UO₂+ binding constant was lower than the Model 2 value ($\log K = 4.50$). When applied to SWy-1 experimental data, Model 3 simulated uranyl adsorption well at a Na concentration of 0.1 mol L⁻¹ (the gibbsite uranyl binding constants were fit at this ionic strength). At lower ionic strengths, the shift toward increased adsorption caused by the relatively high value of the anion-binding constant resulted in an overprediction of adsorption.

The relatively large SWy-1 clay-platelet size resulted in a low abundance of edge sites in proportion to fixedcharge sites, so the computed surface speciation of uranyl on edge sites was affected by site saturation. In Model 1 calculations, 70% of AlOH sites were occupied by UO22+ at maximum adsorption. The reduced concentration of available AlOH sites in combination with the relative magnitude of SiOH and AlOH binding constants was computed to reduce the potential for formation of AlO·(UO₂)₃(OH)₅. Strong binding of uranyl to AlOH sites (log $K_{AlO \cdot UO2}^+ = 7.1$) suppressed the formation of SiO·UO₂⁺ (log $K_{SiO·UO2}$ ⁺ = 0.15). The complex SiO·(UO₂)₃(OH)₅+ was computed to form because the values for $K_{AlO\cdot (UO2)_3(OH)_5}{}^+$ and $K_{SiO\cdot (UO2)_3(OH)_5}{}^+$ were similar (10^{-16.8} and 10^{-15.8}, respectively), and the concentration of AlOH sites available to form AlO. (UO₂)₃(OH)₅ was reduced by formation of AlO·UO₂⁺. SiOH sites, on the other hand, were ionized to SiOat alkaline pH, and SiOH were not occupied by uranyl complexes.

Because the sorbing ions are relatively large, multiple site coverage was considered for its potential effect on adsorption. The diameter of the UO22+ ion is approximately 0.58 nm (Aberg et al 1983). Based on considerations of crystal chemistry, the (UO₂)₃(OH)₅⁺ ion consists of three uranium atoms at the apices of an equilateral triangle with an average edge length of 0.381 nm; each atom is surrounded by hydroxyls and waters of hydration with an interatomic U-O distance of 0.244 nm (Aberg 1978). The longest dimension presented by the $(UO_2)_3(OH)_5^+$ ion is, therefore, approximately 0.869 nm. An estimation of surface-site dimensions obtained by inverting the overall site density for smectites (approximately 6 μ mol sites m⁻²; White and Zelazny 1988) yields an area per site of 2.77 x 10⁻¹⁹ m², equivalent to a square with an edge length of 0.53 nm. The (UO₂)₃(OH)₅⁺ ion, therefore, covers an area at the smectite surface that is potentially larger than that occupied by the complexing site. The effect of interference with adsorption to adjacent sites was evaluated by calculations in which three sites were removed from further interaction for each $(UO_2)_3(OH)_5^+$ ion adsorbed to either AlOH or SiOH sites. The computed maximum reduction to overall adsorption was less than 2% at the highest pH modelled, indicating that the relative size of adsorbing ions did not significantly affect modelled results.

An unexpected computational result was the response of uranyl edge complexation at low pH and varying ionic strength for Model 1. As ionic strength increased, the percentage adsorption of UO₂²⁺ to AlOH sites was computed to increase; at pH 5.2, the edge complex contribution increased from approximately 4% at I = 0.004 to 17% at I = 0.1. This increase resulted from aqueous uranyl speciation and changes in competitive adsorption by edge and exchange sites. As uranyl binding to exchange sites was inhibited with increasing Na, the aqueous concentration of uranyl increased. Concentration and ionic-strength effects on speciation caused the fraction of UO₂²⁺ in the aqueous phase to increase as well. At pH 5.2, for example, the percent of dissolved uranyl as UO22+ increased from 54% to 62% as ionic strength changed from 0.004 to 0.1. This increase led, by mass action, to a concomitant increase in AlO·UO₂+.

CONCLUSIONS

The pH and ionic strength-variant system of uranyl on SWy-1 could be described as a composite of reactions between the sorbate and fixed-charge cation-exchange sites and variable-charge SiOH and AlOH edge sites. The relative concentrations of edge and basalplane surface (exchange) sites were fixed by solid-phase composition and morphology. The contribution to adsorption of the modelled sites could be modelled discretely if their concentrations were estimated independently. The use of the analogue components, silica and gibbsite, to predict uranyl edge complexation reactions was highly dependent on the reaction parameters selected for the aluminol site. For our application, the best suite of surface constants for the AlOH site involved a log K_{a1} of 12.3, specifying negligible development of neutral or negatively charged aluminol sites at the smectite edge. Whether the success of this particular suite of constants was fortuitous or indeed had mechanistic implications was not established. The high value for K_{a1}, although derived from gibbsite titration data, may reflect conditions at the smectite edge that are distinct from the gibbsite surface, because uranyl adsorption to gibbsite was better modelled using surface acidity and electrolyte complexation constants with more conventional values.

Model calculations made without fitting parameters indicate the presence of three predominant species on the smectite surface: UO₂²⁺ bound to exchange sites, UO₂²⁺ bound to AlOH sites, and (UO₂)₃(OH)₅⁺ bound to SiOH sites. The least significant surface complex, SiO·(UO₂)₃(OH)₅, contributed significantly to uranyl

adsorption above a pH of 6. The aqueous speciation of uranyl was a significant determinant of uranyl adsorption. The formation of monovalent aqueous uranyl species as pH increased caused desorption of uranyl from basal-plane sites in the presence of a more abundant monovalent electrolyte cation. Changes in uranyl speciation, e.g., the increased abundance of UO₂²⁺ at low pH as ionic strength increased, caused uranyl adsorption to AlOH sites to increase slightly with increasing ionic strength, and increased abundance of (UO₂)₃(OH)₅⁺ at higher pH accounted in part for the formation of SiO (UO₂)₃(OH)₅. The concentration ratio of UO₂²⁺: (UO₂)₃(OH)₅⁺ at elevated pH, combined with surface acidity and surface binding constants that caused a large fraction of AlOH sites to be occupied by UO₂²⁺, gave rise to SiO (UO₂)₃(OH)₅ to the exclusion of AlO (UO₂)₃(OH)₅. These results differed from those for metals that did not have significant pH effects on speciation, such as Cd, where only a single aqueous metal species was proportionated between surface sites (Zachara and Smith 1994).

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