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Solubility Measurement of Uranium in Uranium-Contaminated Soils

S. Y. Lee Mark Elless Forrest Hoffman

Environmental Sciences Division Publication No 4138



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ABSTRACT

A short-term equilibration study involving two uranium-contaminated soils at the Department of Energy's Fernald Environmental Management Program (FEMP) site was conducted as part of the In Situ Remediation Integrated Program. The goal of this study is to predict the behavior of uranium during on-site remediation of these soils. Geochemical modeling was performed on the aqueous species dissolved from these soils following the equilibration study to predict the on-site uranium leaching and transport processes. Results showed that the soluble levels of the major components (total uranium, calcium, magnesium, and carbonate) increased continually for the first four weeks. After the first four weeks, these components either reached a steady-state equilibrium (in those components having solubilities approaching that of the controlling solid phase for that component) or continued linearity throughout the study (in those components having low solubilities). Other major components, such as aluminum, potassium, and iron, reached a steady-state concentration within three days. Silica levels approximated the predicted solubility of quartz throughout the study. A much higher level of dissolved uranium was observed in the soil contaminated from spillage of uranium-laden solvents and process effluents than in the soil contaminated from settling of airborne uranium particles ejected from the nearby incinerator. The high levels observed for soluble calcium, magnesium, and bicarbonate are probably the result of magnesium and/or calcium carbonate minerals dissolving in these soils. The increase in the total uranium levels with increasing carbonate levels is due to the complexation of uranium with carbonate species. Geochemical modeling confirms that the uranyl-carbonate complexes are the most stable and dominant in these solutions. The implication of this work is that the use of carbonate minerals on these soils for erosion control and road construction activities contributes to the leaching of uranium from contaminated soil particles. Dissolved carbonates promote uranium solubility, forming highly mobile anionic species. Mobile uranium species are contaminating the groundwater underlying these soils. Therefore, the development of a site-specific remediation technology is urgently needed for the FEMP site.

INTRODUCTION

Remediation of uranium-contaminated soils is currently considered a high priority within the U.S. Department of Energy (DOE) not only because these soils represent an environmental hazard, but also because these soils are a potential source of contamination of the underlying groundwater during natural leaching episodes. Therefore, the behavior of uranium under saturated conditions, particularly its complexation potential and mobility, must be understood for predicting the environmental impact these soils might have on the underlying groundwater quality.

The solubility of soil uranium depends upon the soil's physicochemical, mineralogical and micromorphological properties; the nature of the uranium association; and the mineralogical, morphological, and compositional characteristics of the uranium-bearing phases. In particular, uranium solubility is enhanced by the presence of dissolved carbonate species, especially if the uranium is in the hexavalent form (Bowie and Plant 1983; Francis et al. 1992). Uranyl-carbonate complexes are very strong and are often negatively charged. The negativity associated with these complexes allows them to be rather mobile in the soil environment and therefore represents a potential for groundwater contamination.

Uranium-contaminated soils from the Fernald Environmental Management Project (FEMP) site were selected for this study. Solubility of uranium-containing minerals and concentration of each uranium species in equilibrated soil solutions will be determined. The results of the FEMP study will be beneficial for the In Situ Remediation Integrated Program (ISRIP) and the Uranium Soil Integrated Demonstration (USID). This information is needed by ISRIP for its evaluation of FEMP as a potential in situ integrated demonstration site and by USID for chemical modeling and environmental assessment. As a part of the USID, two FEMP uranium-contaminated soils and several cores have been collected and partially characterized (Lee and Marsh 1992).

To ascertain the behavior of uranium solubility at the Fernald site, a short-term equilibration experiment using two uranium-contaminated FEMP soils was conducted. These two soils were selected because each represents a different mode of uranium contamination (airborne and spillage). This experimental approach has been commonly used to measure solubility of soil components and to identify soluble ionic and complex species of target components in soil solutions. The determination of solubility and chemical speciation will provide vital information for understanding contaminant behavior in these soils under natural conditions and during any in situ remediation demonstration.

Therefore, the objectives of this task are (1) to measure the solubility of the soil components in two uranium-contaminated soils, (2) to calculate the distribution of the dominant uranium species in the soil solution using a chemical speciation model, and (3) to compare the results of the soil equilibration approach with the known groundwater composition from this site. This study will submit results to (1) ISRIP, (2) the FEMP site remedial investigation team, and (3) the Characterization Task Group of the USID program.

2. MATERIAL AND METHODS

2.1 SOIL SAMPLING AND PRELIMINARY CHARACTERIZATION

FEMP personnel sampled and homogenized two FEMP soils for treatability tests by the USID Decontamination Task Group. One sample (B-16, Drum No. 6) was collected near the Plant 1 Storage Pad Area within the production area. The other sample (A-14, Drum No. 12) was collected near the Incinerator area located a few hundred yards east of the main plant area. The Characterization Task Group of the USID program took soil core samples from the same area and characterized their physicochemical and mineralogical properties (Lee and Marsh 1992). Each excavated area was ~ 25 x 20 ft, with an excavation depth of 6 to 8 inches (Kneff et al. 1992). Preliminary characterization of the samples was completed (Lee and Marsh 1992), but detailed characterization for treatability studies will be continued by the Characterization Task Group of the USID. Particle-size separations were performed on both FEMP soils to determine the uranium partitioning among the various size fractions. Particle size separations were performed by field moist sieving using 4- and 2-mm stainless steel sieves. Size fractions larger than 2 mm were designated as gravel. The <2-mm fractions were further separated into sand particles measuring 2 to 0.053 mm, silt particles ranging from 0.053 to 0.002 mm, and clay particles measuring < 0.002 mm by wet sieving and centrifugation (Jackson 1975). These size-fractionated samples and the whole soils were then analyzed by neutron activation analysis for uranium quantification (Wade et al. 1992). Heavy liquid separations were also performed on the sand and silt fractions of both soils in an attempt to isolate a uranium-bearing fraction for later mineralogical analysis. Lithium metatungstate (LMT, $\rho = 3.0 \text{ g/cm}^3$) was the medium used in all density separations. Uranium quantification on all density fractions was determined by gamma spectroscopy (Larsen et al. 1992).

2.2 MINERALOGICAL CHARACTERIZATION

Scanning electron microscopy (SEM), using both secondary and backscattered electron imaging coupled with energy-dispersive X-ray analysis (EDX), was used for morphological analysis and particle-size/elemental distributions. A small amount of dry whole soil and size-fractionated samples of both soils was embedded in epoxy resin under vacuum. Vacuum removal of the soil air allows complete resin migration into the soil micropores. After resin polymerization, microscopic specimens that have cross-sectional areas of ~ 1 to 4 cm² were prepared for SEM examination by coarse sanding and fine polishing.

Mineralogical analyses by X-ray diffraction (XRD) of uranium-enriched size and density fractions of both soils were conducted to determine the nature of the uranium phases occurring in these soils. Phase identification was determined using the Joint Committee of Powder Diffraction Standards data base. Additionally, quantitative analysis on the carbonate minerals in these soils was performed by the reference intensity ratio (RIR) method (Chung 1974a, 1974b, 1975). The authors determined the RIR constants for dolomite, calcite, and quartz as well as the relative intensities of each pure mineral rather than using published values, because these constants strongly depend on the slide preparation for XRD. All XRD analyses were performed on a Scintag 2000XDS equipped with CuK α -radiation. Power settings for all XRD analyses were 45 kV and 40 ma. All diffractograms were collected from 2 to 70° 2 Θ at 1° 2 Θ min⁻¹.

2.3 EQUILIBRATION STUDY AND GEOCHEMICAL MODELING

For the equilibration study, 200 g of air-dried A-14 and B-16 soil were equilibrated with 2500 mL of deionized distilled water in a 1-gal polyethylene container. Duplicate samples of each soil and one experimental blank were also analyzed as an internal check on the precision of the analytical results and on the base-line quality of the water used in this study. For the first 10 weeks of the study, this mixture was shaken manually three times daily for 30 s each time and allowed to rest between these shakings. Periodically, these samples were also allowed to aerate to atmospheric conditions. After ten weeks, the samples were allowed to rest until the final sampling episode.

For each sampling episode, the samples were allowed to rest for 1 h following the first shaking before a 50-mL aliquot was taken from each sample. Each aliquot was then vacuum-filtered through $0.45-\mu m$ millipore membrane paper to remove all coarser particulates from the aliquot. Following filtration, the samples were brought to the analytical laboratory for immediate analysis of pH, dissolved cation and anions, total uranium, and alkalinity. Cation (Na⁺, Li⁺, K⁺, Ca²⁺, Mg²⁺, Fe³⁺, Al³⁺, Si⁴⁺, and many others) and anion (Cl⁻, F⁻, NO₃⁻, SO₄²⁻, PO₄³⁻) concentrations were determined by inductively coupled plasma (ICP) spectroscopy and ion chromatography, respectively. Charge balance, based upon the analytical results, was used to check the performance of the analyses. A small average net positive charge of 1.52 x 10⁻⁴ for soil A-14 and a small average net negative charge of 4.45 x 10⁻⁴ for soil B-16 were calculated. Radionuclide concentrations (²³⁵U and ²³⁸U) and alkalinity were determined with the use of mass spectrometer and acid titration, respectively. Samples were taken according to the following schedule (in days after initial mixing): 1, 3, 7, 14, 21, 28, 70, and 300.

One caveat for this procedure is warranted. The withdrawal of 50-mL aliquots per sampling episode causes a reduction in the solution:solid ratio for each sampling episode of 2% of the initial solution volume. Initially, this ratio is 12.5:1, whereas at the end of the study the ratio is 10:1. Although extraction efficiencies are expected to become lower with lower solution:solid ratios, the authors believe that this ratio varies rather narrowly and is not expected to significantly retard the dissolution of these soils.

Geochemical modeling was performed using the Geochemical Expert System (GES) prototype (Hoffman and Tripathi 1993), a software system designed to analyze interactions between solution and mineral phases in nature. GES uses the MINEQL (Westall et al. 1976) equilibrium model to assess the rate and extent of geochemical interactions. GES then uses the results of the model to create qualitative geochemical interpretations similar to those written by expert geochemists. This program attempts to describe important characteristics and salient features of the prescribed geochemical composition using as much quantitative information obtained from the equilibrium model, as possible. GES actually generates English text to describe the state of oxidation/reduction, complexation, and precipitation/dissolution of the geochemical system.

Because of some assumptions made by GES, additional modeling was performed using MINEQL directly. These simulations were used to predict the complexation potential, activity, and degree of complexation of each dissolved species and the saturation indices of each solid phase. Kinetic processes were not modeled because kinetic data are uncertain or nonexistent.

3. RESULTS AND DISCUSSION

3.1 GENERAL SOIL PROPERTIES

The A-14 soil was covered by fescue grass. Soils near the concrete curb of the driveway were highly disturbed, but the soils away from the curb were less disturbed. This soil had a well-developed Ap horizon with a dark grayish-brown (7.5YR 4/2) color, 5% gravel content, high organic matter content, pH 7.2, and a silt loam texture. Undisturbed soils of this area are classified as the Fincastle soil series (USDA 1979).

The B-16 soil was also covered by rescue grass. In general, the soils near the Storage Pad were highly disturbed, but the soils away from the Storage Pad were less disturbed. This soil had a dark brown (7.5YR 3/3) color, 5% gravel content, pH 7.5, and a silt loam texture. These soil properties suggest that the B-16 soil may be similar to the Ap horizon of the Henshaw soil series (USDA 1979).

Even though soils A-14 and B-16 share the same textural class, differences in the actual percentages for each fraction are evident. The A-14 soil has a greater silt percentage and lesser sand and clay percentages than the spatially adjacent B-16 soil (Table 1).

	Size range	Size distribution	Uranium concentration	Uranium contribution		
Soil	Class mm	(%)	mg/kg	mg/kg	(%)	
A-14	Gravel >2	5	<1	<1	<1	
	Sand 2-0.05	12	1043	125	27	
	Silt 0.05-0.002	70	288	203	44	
	Clay < 0.002	13	1026	133	29	
B -16	Gravel >2	5	<1	<1	<1	
	Sand 2-0.05	21	117	25	7	
	Silt 0.05-0.002	54	240	129	37	
	Clay < 0.002	20	989	195	56	

Table 1. Results of particle size and uranium distribution of the soils equilibrated

3.2 NATURE OF URANIUM CONTAMINATION

The uranium partitioning among the particle-size fractions of both soils is shown in Table 1. The distribution of uranium among the particle-size fractions indicates that the nonclay fractions contain most of the uranium in soil A-14 and 44% of the uranium in soil B-16. The results are conclusive evidence that the uranium in these soils mainly occurs as a particulate form rather than

an adsorbed form on the external and internal surfaces of layer silicate minerals (clay minerals). SEM analyses of the samples also confirm this interpretation (Plates 1 and 2).

Results of the heavy liquid separations for both soils show a predominance of light minerals over heavy minerals (Table 2). For both A-14 silt and sand, >96% of the total sample occurred in the light fraction. For B-16 silt, >80% of the sample occurred in the light fraction. For B-16 solt, >80% of the sample occurred in the light fraction. For B-16 solt, >80% of the sample occurred in the light fraction.

Results of the uranium partitioning among all particle density fractions show much higher levels of uranium phases (on an equivalent weight basis) in A-14 than B-16. Furthermore, higher uranium levels were associated with the sand fractions than with the silt fractions of each soil. A predominance of uranium phases was observed in the heavy sand and silt fractions of A-14, and the highest levels of uranium of any density fraction were observed in the heavy sand fraction of this soil. Nearly equal uranium levels were observed between the light and heavy fractions of B-16, thereby indicating an ineffective density separation for this sample. All uranium-rich heavy fractions were later examined by XRD for determination of their uranium mineralogy.

		Wei	ght distribution	Density fraction		
Soil	Size fraction	Heavy	Light %	Heavy	Light pCi/g	
A-14	Silt	2	98	516	34	
	Sand	4	96	1740	414	
B-16	Silt	21	80	15	21	
	Sand	4	96	40	37	

 Table 2. Results of the uranium partitioning among the particle density fractions of the equilibrated soils

3.3 MINERALOGICAL PROPERTIES

SEM micrographs of the A-14 sand fraction show a wide variety of sizes and shapes of minerals (Plate 1). Some minerals occur as a stable, large-sized aggregate. The unusually high stability of the aggregates is expected to develop during the incinerating process. The aggregates contained uranium particles as well as other heavy minerals. Uranium occurred also as a microfracture-filling mineral in the aggregate. Uranium-containing particles were typically composed of calcium, silicon, or phosphorus. A cerium phosphate mineral was also found in this sample. Quartz was the dominant mineral in the sand fraction.



Plate 1. Four scanning electron micrographs were taken from the sand fraction of the A-14 sample. Particle A has uranium as a major component and a small amount of oxygen. Particles B and D have uranium and a lesser amount of calcium, phosphorus, and oxygen. Particle C has phosphorus, cerium, neodymium, and a lesser amount of thorium and oxygen. The microfracture-filling mineral (marked E) has uranium, silicon, and a smaller amount of calcium and phosphorus. The matrix of the aggregate is aluminosilicate clays (marked F) and silt-size quartz (marked G).



Plate 2. Two scanning electron micrographs were taken from the silt fraction of the B-16 sample. The uranium particle A also contains calcium, silicon, and aluminum. The fine silty aggregate is an aluminosilicate mineral (marked B). Particle C is an iron oxide mineral and particle D is a phosphate mineral that contains cerium, neodymium, and lanthanum (possibly monazite mineral). Particles E and F are an iron/titanium oxide (ilmenite) mineral. Other silt particles are quartz, dolomite (marked G), and feldspars.

In the B-16 silt fraction (Plate 2), uranium particles either were composed entirely of uranium or coexisted with calcium and/or silicon. In future analytical determinations, detailed microscopic analyses will be performed after heavy liquid separation in order to examine concentrated uranium-bearing fractions.

Mineralogically, these two soils are dominated by quartz and carbonate minerals in their bulk soil. Calcite $[CaCO_3]$ and dolomite $[CaMg(CO_3)_2]$ have been identified as the carbonate minerals in these soils. These two carbonate minerals are anthropogenic artifacts because these two minerals are not present in the nearby off-site soils at this same depth. They occur in these soils because FEMP personnel placed these minerals on the uranium-contaminated soils for erosion control and road construction activities. Quantitative estimates of the carbonate minerals and quartz on a whole soil basis are 20% calcite, 2% dolomite, 65% quartz, and 13% clay minerals in soil A-14 and 15% calcite, 19% dolomite, 46% quartz, and 20% clay minerals in soil B-16. Feldspars were minor phases in both soils. Kaolinite and illite are the two most dominant clay-sized minerals, and lesser amounts of quartz and traces of smectite and/or vermiculite occur in this size fraction of both soils.

In terms of uranium mineralogy, uraninite $[UO_2]$ is the only uranium mineral identified in these soils. This mineral was observed only in the heavy-sand nonmagnetic fraction of core SP-9, which was sampled in the same area as soil A-14. This tetravalent form of uranium is much less soluble than the hexavalent form and may be partly responsible for the lower extraction efficiencies of the A-14 soil than the B-16 soil as reported by the leaching task group (Francis et al. 1992). Other minerals observed in the heavy-sand and silt fractions of both soils include amphiboles, anatase, and iron oxides.

3.4 EQUILIBRATION STUDY AND GEOCHEMICAL MODELING

Results showed that the soluble levels of the major components (total uranium, calcium, magnesium, nitrate, and carbonate) continually increased for the first 28 d and then either approached a steady-state condition in those components having solubilities nearing that of the controlling solid phase for that component or continued linearity throughout the study in those components having low solubilities. Other major components, such as aluminum, potassium, and iron, reached a steady-state concentration within 3 d. Silica levels approximated the predicted solubility of quartz throughout the study.

On average, total uranium levels increased linearly throughout the study from <0.1 to 3.8 mg U/L (<1 to 36 μ g U/g soil) in soil A-14 <Fig. 1a>. The maximum concentration represents only 7% of the total uranium in the whole soil of A-14. A similar increase along a logarithmic function from 2.5 to 10.0 mg U/L (30 to 95 μ g U/g soil) was observed for the B-16 soil <Fig. 1b>. This maximum concentration represents 21% of the total uranium in the whole soil of B-16.



Fig. 1. Total soluble uranium concentrations versus time for (a) soil A-14 and (b) soil B-16. Data for both duplicates are shown.

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Quite evidently, much higher absolute and relative levels of soluble uranium were observed throughout the study in the B-16 than the A-14 soil. In fact, the uranium concentration in B-16 soil solution after 300 d is nearly three times the corresponding uranium concentration in A-14 soil solution. The low-soluble uranium levels observed for soil A-14 cause this soil to remain undersaturated with respect to the uranium minerals considered by MINEQL in this study. The linearity in uranium solubility for soil A-14 indicates that saturation with respect to uranium minerals has not been approached in this soil. In soil B-16, the much higher levels appear to approach a steady-state condition, suggesting that saturation or near saturation with respect to uranium minerals has been reached. The overall greater solubility of uranium in the B-16 soil may be due to the higher carbonate content of this soil compared with that of A-14 and to the presence of less soluble tetravalent uranium-bearing refractive phases (i.e., uraninite) formed in the incinerating process in soil A-14.

Calcium and magnesium were the two dominant soluble cationic species observed throughout this study. For the A-14 soil, calcium levels increased along a logarithmic function from 8.7 to 31.8 mg/L (107 to 325 μ g Ca/g soil) before decreasing to 30.9 mg/L (294 μ g Ca/g soil) at 300 d, whereas magnesium levels increased along a logarithmic function from 2.0 to 6.2 mg/L (25 to 63 μ g Mg/g soil) before decreasing to 4.1 mg/L (39 μ g Mg/g soil) at 300 d < Fig. 2a>. For the B-16 soil, calcium levels increased rapidly along a logarithmic function from 19.0 to 36.0 mg/L (232 to 397 μ g Ca/g soil) through the first 4 weeks and afterward have been nearly constant through 300 d. Magnesium behaved similarly; levels also increased along a logarithmic function from 2.9 to 5.1 mg/L (36 to 56 μ g/g soil) through the first 4 weeks and afterward have been nearly constant through 300 d < Fig. 2b>. The nearly constant calcium and magnesium levels after 28 d in both soils is interpreted as the attainment of saturation or near-saturated conditions with respect to the controlling calcium and magnesium phases in this soil. Other soluble cationic species (i.e., sodium, iron, and aluminum) remained at low concentrations throughout the study in both soils (see Appendix). The slightly higher concentrations of soluble calcium and magnesium in soil B-16 probably reflect the higher calcite and dolomite mineral contents in this soil than those in soil A-14.

Total carbonate species or alkalinity was the dominant anionic species throughout the study for both soils. Bicarbonate was the dominant carbonate species present throughout the study. For soil A-14, total soluble bicarbonate levels increased along a logarithmic function from 13.0 to 81.5 mg/L (159 to 774 μ g/g soil) < Fig. 3a>. For soil B-16, total soluble bicarbonate levels increased along a logarithmic function from 39.5 to 110.0 mg/L (484 to 1045 μ g/g soil) < Fig. 3b>. The logarithmic response of alkalinity for both soils signifies an approach to saturated conditions with respect to the controlling carbonate species. The observed increase in the total uranium levels with increasing carbonate levels is very likely due to the enhanced solubility and complexation of uranium by carbonate species.

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(a)



Fig. 2. Total soluble calcium and magnesium concentrations versus time for (a) soil A-14 and (b) soil B-16. Data for both duplicates are shown.



Fig. 3. Total soluble alkalinity concentration versus time for (a) soil A-14 and (b) soil B-16. Data for both duplicates are shown.

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Nitrate was the next dominant anionic species. Overall, nitrate levels generally increased logarithmically through the first 10 weeks in both soils, and slightly higher levels were observed in the A-14 soil < Figs. 4a, 4b>. After 10 weeks until the final sampling at 300 d, nitrate levels dropped precipitously to <25% of the maximum levels for both soils. The authors believe that the microorganisms depleted the dissolved oxygen in the closed containers from a 70- to 300-d period. Anaerobic bacteria (either chemoautotrophs or chemoheterotrophs) would then use nitrate as their electron sink and dramatically lower nitrate levels The presumed lowering of the pe may be responsible for the increase in pH (>8.0 for both soils) during this final sampling episode because of the inverse relationship of pe and pH in the Nernst equation (Ponnamperuma 1972). This pH increase may in turn induce calcite precipitation in these soils which may account for the observed evening to lowering of both the calcium and magnesium levels with time. Excluding the levels associated with the 300-d sampling episode, sulfate concentrations similar to those of nitrate were observed in the B-16 soil; however, sulfate levels in the A-14 soil remained quite low (see Appendix). This observed anaerobicity is unlikely to occur in the field because these soils are seldom saturated for long periods of time to induce anaerobic conditions. The extension of the best-fit line in these two graphs beyond the first 10 weeks is suspect; however, the authors believe that this logarithmic extension, which assumes little to no anaerobic conditions occurring in these soils, more accurately models the natural situation of these soils. Other soluble anionic species (i.e., chloride, fluoride, and phosphate) remained at low concentrations throughout the study in both soils (see Appendix).

In terms of pH, both soils showed a rapid increase in pH within the first 3 d and then achieved a steady-state condition through the first 10 weeks of the study. Both soils, however, showed a strong increase in pH (>0.5 pH unit increase) at the 300-d sampling episode (see Appendix). As stated earlier, this pH increase may be related to the observed nitrate reduction process, which occurred within these closed containers from 70 to 300 d. Reduction processes are known to consume excess protons, causing a concomitant increase in pH. One potential ramification of increasing pH may be calcite precipitation, which lowers soluble calcium, alkalinity, and even magnesium levels. The more alkaline pH at the start of the study in soil B-16 than in soil A-14 probably reflects the higher carbonate mineral content of this soil.



Fig. 4. Total soluble nitrate concentrations versus time for (a) soil A-14 and (b) soil B-16. Data for both duplicates are shown. Best-fit line excludes 300 d data points.

Geochemical modeling was employed to determine the complexation potential of soluble uranium. Given that the pH for both soils is near neutral, the redox potential of the soil solutions was assumed to be zero voltage. This is also a conservative estimate based upon the typical Eh-pH pairs found in natural aqueous environments (Garrels and Christ 1965). This estimate is not a critical factor for the uranium speciation results because the most important uranium redox couples occur at even more oxidizing conditions than the assumed condition (Garrels and Christ 1965). Results indicate that the three most dominant soluble uranium species for both soils uranyl dicarbonate $[UO_2(CO_3)_2]^{2-},$ uranyl tricarbonate $[UO_2(CO_3)_3]^4$, were and trihydroxocarbonatodiuranyl $[(UO_2)_2CO_3(OH)_3]^{1-}$. Uranyl dicarbonate accounted for 55% of the total dissolved uranium, whereas uranyl tricarbonate and trihydroxocarbonatodiuranyl accounted for 27% and 9% of the total dissolved uranium, respectively, for soil A-14. Similarly for soil B-16, uranyl dicarbonate accounted for 40% of the total dissolved uranium, whereas uranyl tricarbonate and trihydroxocarbonatodiuranyl accounted for 30% and 13%, respectively. It is important to note that among these three most dominant soluble uranium species, anionic uranium species account for 91 and 83% of the total soluble uranium species in soils A-14 and B-16. respectively. Therefore a large portion of the uranium solubilized in this experiment is calculated to be in an anionic form that would be quite mobile in the soil environment and would possibly contaminate the underlying groundwater.

Table 3. Comparison of the average major chemical components between the two Fernald soils after equilibrium and the groundwater wells at the Fernald site. Concentrations of all dissolved species, excluding pH, are in milligrams per liter.

Dissolved species	Equilibra A-14	ation data B-16	Grou High	ndwater Low	
pН	8.0	8.2	7.5	6.6	
Calcium	31	40	261	92	
Magnesium	4	6	75	36	
Alkalinity	82	110	323	214	
Uranium	4	10	12	1	
Major uranium species	[UO ₂ ([UO ₂ ([(UO ₂) ₂ CO	CO ₃) ₂] ²⁻ CO ₃) ₃] ⁴⁻ D ₃ (OH) ₃] ¹⁻	[UO ₂ (CO ₃) ₂] ² [UO ₂ (CO ₃) ₃] ⁴ [(UO ₂) ₂ CO ₃ (OH) ₃] ⁻¹		

Similar ranges in pH and total uranium were observed between the two Fernald soils and the groundwater (Tidwell et al. 1992). Additionally, similar aqueous uranium species predicted to occur in this study were also predicted to occur in the groundwater wells at the Fernald site <Tab 3>. However, much higher levels of soluble calcium, magnesium, and total carbonate were observed in the groundwater wells at the site. This direct comparison of the soil solution results with the groundwater results is tentative because (1) the solution:soil ratio associated with the groundwater is unknown, (2) the recharge/discharge behavior of the groundwater is also unknown, and (3) the transient character or "seasonality" of the groundwater was not modeled in this study.

The water chemistries from this study were also used to predict undersaturation, equilibrium, and supersaturation conditions with respect to chemically similar minerals. The water chemistries at steady state of both soils were either near equilibrium or supersaturated with respect to uraninite $[UO_2]$, calcium autunite $[Ca(UO_2)_2(PO_4)_2]$, fluorapatite $[Ca_5(PO_4)_3F]$, hydroxyapatite $[Ca_5(PO_4)_3OH]$, calcite $[CaCO_3]$, dolomite $[CaMg(CO_3)_2]$, and quartz. The water chemistries at steady state of both soils were undersaturated with respect to sodium autunite $[Na_2(UO_2)_2(PO_4)_2]$ and ningyoite $[UCa(PO_4)_2*2H_2O]$. It is interesting to note that the majority of the minerals predicted to be near equilibrium or supersaturated are phosphate minerals. Because the uranium incorporated into calcium phosphate minerals is insoluble, the phosphate concentration controls the rate of uranium release from these minerals. Therefore, the phosphate concentration may be considered the master variable in influencing the dissolution behavior of the Fernald soils.

Thermodynamically, $UO_2(HPO_4)_2^{2-}$ is the most soluble uranyl species between pH 4 and 7.6, whereas $UO_2(CO_3)_2^{2-}$ and $UO_2(CO_3)_3^{4-}$ become the most soluble uranyl species above pH 7.6 in a purely aqueous system (Langmuir 1978). However, in soil-solution systems, phosphate anions readily adsorb onto the surfaces of minerals that are pH dependently charged (i.e., kaolinite and sesquioxides) (Lindsay et al. 1989). Adsorption of phosphate reduces its availability in the soil solution and causes the dominance of uranyl-carbonate complexes in both soils.

Tidwell et al. (1992) found that the water chemistries from all six groundwater wells investigated at the Fernald site showed supersaturation with respect to soddyite $[(UO_2)_2SiO_42H_2O]$. Additionally, four of these wells showed saturation with respect to haiweeite $[Ca(UO_2)_2Si_6O_{15}5H_2O]$, whereas the other two were slightly undersaturated with respect to haiweeite. In one well, that contained the highest phosphate levels, supersaturation with respect to $(UO_2)_3(PO_4)_24H_2O)$ and saleeite $[(UO_2)_2Mg(PO_4)_2]$ was observed. Rutherfordine $[UO_2CO_3]$ and schoepite were also observed near saturation in all wells. Supersaturation with respect to uranium silicate minerals at equilibrium in their groundwater wells was predicted because they assume silicon saturation with respect to quartz in the well waters.

The observed dissolution behavior of both Fernald soils was used to estimate the leaching of uranium from these soils under natural conditions (i.e., a severe or prolonged storm event). Linear regression analysis using soluble uranium levels during the first 3 d of this study was performed to determine the expected level of uranium solubilized under these conditions. This estimate is conservative because the uranium values during the first 3 d for both soils are undersaturated with respect to any uranium-bearing mineral. Additionally, some uranium-bearing pore water is still held in the soil even after a storm. Even upon uranium precipitation during the next dry cycle, this freshly precipitated form would be much more easily redissolved in the next wet cycle as a uranyl carbonate form rather than initially when it is not bound with carbonate. Results of this analysis predict that 0.4 μ g U/g soil (0.09%) and 15.6 μ g U/g soil (4.47%) would become solubilized for A-14 and B-16, respectively. Because the mode of uranium contamination associated with soil B-16 represents the typical mode of uranium contamination at the Fernald site, the higher predicted soluble uranium levels associated with soil B-16 impart severe ramifications for groundwater contamination at the Fernald site. If this solubilized uranium does not become readsorbed onto soil particles, significant contamination of the groundwater underlying this soil will result.

4. SUMMARY AND RECOMMENDATIONS

Results from a 300-d equilibration study involving two uranium-contaminated soils from the DOE Fernald site have predicted that the solubilized uranium exists mainly as anionic uranyl-carbonate species. Because of their anionic character, these species are considered quite mobile in the soil environment and may contaminate the underlying groundwater under prolonged and/or severe leaching conditions. High levels of soluble calcium, magnesium, and bicarbonate are probably the result of the dissolution of magnesium and/or calcium carbonate minerals in these soils. The increase of carbonnet levels in the solute by progressive dissolution of the minerals would enhance the solubility of uranium-bearing minerals by carbonate complexation on the surface of the particulates.

The implication of this work is very important in many ways. For example, uranium transport assessment to off-site areas as a part of risk assessment should recognize that the solubility of uranium-bearing minerals is the critical factor rather than uranium distribution coefficient (Kd) in soils. Anionic uranyl-carbonate complexes are very stable in this carbonate-dominated system (i.e. carbonate-rich soil, parent material, and groundwater). Therefore, as long as the pH of these soils is maintained near neutrality, sorption of uranium onto soil particles will be unlikely in both Fernald soils. FEMP site management should realize that the use of carbonate minerals on these soils for the control of erosion and for road construction activities actually aids in the leaching of uranium from contaminated soil particles. The dissolved carbonates promote uranium solubility, forming highly mobile anionic species. Unfortunately, such contamination has been documented in groundwater wells at the Fernald site. Therefore, the development of a site-specific remediation technology is urgently needed for the FEMP site.

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APPENDIX

Soil Equilibration Results of the Major Cations Soil Equilibration Results of the Major Anions

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Soil Equilibration Results of the Major Cations

Sample	Sampling Period	Sampling Volume	pH	Total Uranium	Al	Ca	Fe	Si	Mg	Na
	days	mL		<			-mg/L			>
A-14	1 3 7 14 21 28 70 300	2450 2400 2350 2300 2250 2200 2050 1900	6.6 6.9 7.7 6.9 7.3 7.1 8.0	0.035 0.050 0.098 0.205 0.260 0.443 0.743 3.780	1.10 1.40 0.35 0.25 0.20 BDL 1.05 BDL	8.70 12.25 14.15 19.70 23.10 27.05 31.75 30.90	0.90 1.10 0.25 0.30 0.15 BDL 0.85 BDL	ND ND 3.10 4.25 ND 6.00 4.50	2.00 2.80 3.10 4.10 4.70 5.45 6.15 4.10	1.10 1.85 0.75 2.10 0.65 0.90 0.95 1.00
B-16	1 3 7 14 21 28 70 300	2450 2400 2350 2300 2250 2200 2050 1900	7.6 7.4 7.9 7.4 7.7 7.7 8.2	2.480 3.290 4.295 5.325 5.470 7.020 8.370 10.040	1.85 BDL BDL BDL BDL BDL BDL BDL	18.95 22.15 23.75 29.25 31.90 36.05 40.70 39.75	1.75 BDL BDL BDL BDL BDL BDL BDL	ND ND 2.45 3.00 ND 3.00 3.90	2.90 3.05 3.30 3.95 4.35 5.05 5.65 6.25	2.30 2.25 2.00 2.20 1.85 2.15 2.10 2.45

Note: BDL = Below detection limit (For Al, BDL < 0.2 mg/L. For Fe, BDL < 0.3 mg/L).

ND = Not determined.

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Sample	Sampling Period	Sampling Volume	Cl	F	NO3	P04-P	SO4	TOTAL ALK
	days	mL	>]	mg/L		>
A-14	1 3 7 14 21 28 70 300	2450 2400 2350 2300 2250 2200 2050 1900	1.02 1.93 0.57 2.19 0.82 0.78 1.38 2.92	0.58 1.83 1.97 1.99 1.90 1.10 0.97 1.52	11.45 5.25 13.65 15.70 17.75 10.45 20.60 4.39	1.17 0.29 BDL BDL 4.01 2.95 4.20 2.52	1.89 2.18 2.75 3.88 3.77 2.80 3.87 4.70	13.00 22.50 25.00 38.50 50.50 66.50 74.00 81.50
B-16	1 3 7 14 21 28 70 300	2450 2400 2350 2300 2250 2200 2050 1900	1.36 0.90 0.50 0.52 0.80 0.54 0.93 4.07	2.59 3.39 6.43 8.09 8.24 4.10 3.62 3.95	8.86 7.72 11.00 9.91 12.15 10.25 12.45 0.92	BDL BDL BDL BDL BDL BDL 0.29	8.71 7.51 9.33 9.74 10.95 9.35 12.65 18.40	39.50 47.00 46.50 66.50 70.00 86.00 99.50 110.00

Soil Equilibration Results of the Major Anions

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Note: BDL = Below detection limit. For PO₄, BDL < 0.125 mg/L.

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