

Historical Case Analysis of Uranium Plume Attenuation

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Abstract

Groundwater plumes containing dissolved uranium at levels above natural background exist adjacent to uranium ore bodies at uranium mines, milling locations, and at a number of explosive test facilities. Public health concerns require that some assessment of the potential for further plume movement in the future be made. Reaction-transport models, which might conceivably be used to predict plume movement, require extensive data inputs that are often uncertain. Many of the site-specific inputs are physical parameters that can vary spatially and with time. Limitations in data availability and accuracy mean that reaction-transport predictions can rarely provide more than order-of-magnitude bounding estimates of contaminant movement in the subsurface. A more direct means for establishing the limits of contaminant transport is to examine actual plumes to determine if, collectively, they spread and attenuate in a reasonably consistent and characteristic fashion. Here a number of U plumes from ore bodies and contaminated sites were critically examined to identify characteristics of U plume movement.

The magnitude of the original contaminant source, the geologic setting, and the hydrologic regime were rarely similar from site to site. Plumes also spanned a vast range of ages and no complete set of time-series plume analyses exist for a particular site. Despite the accumulated uncertainties and variabilities, the plume data set gave a clear and reasonably consistent picture of U plume behavior. Specifically, uranium plumes:

- Appear to reach steady-state, that is, they quit spreading rapidly (within a few years).
- Exceed roughly 2 km in length only in special cases e.g. where *in situ* leaching has been carried out. The majority is much smaller.
- Exhibit very similar U chemistry between sites. This implies analogous contaminant attenuation mechanisms despite their location.

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1.0 Introduction

Uranium plumes in groundwater have been produced in the course of mining, ore processing, and weapons testing. Because of concerns about off site movement of U in groundwater, many of the sites are being actively remediated while others are being considered for remediation; most will require some long-term monitoring. Natural processes, such as sorption, mineral growth and dispersion occur in the absence of, and often in parallel with, active remediation and collectively set limits on how far a particular plume can move. To assess performance and to guide long-term monitoring, it would be useful to know what these limits are.

Conceptually, a plume of dissolved U emanating from a point source can be expected to have the highest dissolved contaminant levels near the source and progressively lower level down-gradient. In theory, the direction of maximum advance of contaminants should be parallel to the hydrologic gradient. Once the source term is removed, or treated to stop further addition of contaminant to the groundwater, e.g. with a landfill cap, dilution by fresh recharge should lead to a decrease in dissolved phase concentrations. Indeed, dilution along the leading edge of a plume alone should ultimately arrest subsequent advance of a plume. Note though that, in

the absence of chemical removal mechanisms (natural or engineered), the contaminant mass in a plume is unaffected by dilution. Natural mechanisms that reduce the bioavailable mass of U in soil solutions include reversible and/or irreversible sorption and chemical transformation, e. g. reduction to less soluble forms. This natural reduction in mobile U mass would tend to hasten the cessation of plume advance. Recent work on leaking underground fuel tanks¹ suggests that if chemical processes were collectively the primary control over finite plume movement, we would expect to see similar behavior for U plume migration. For example, we might expect all U plumes to reach a similar length before halting. Knowledge of a general U plume cessation length would be useful in a public health sense as it would constitute a first step in assessing the potential impact of particular plumes on groundwater. It would also allow monitoring wells to be more effectively located for groundwater protection. We hypothesize that the natural history of U plumes is best developed by examining large numbers of individual plumes after the approach of Rice et al.¹ and resolving features of their collective advance and decay into specific chemical and physical processes.

2.0 Uranium in Soils

Feldspars in alkaline and granitic igneous rocks are the main source for much of the U present in near-surface natural environments (see table 1 for environmental concentrations of U). Transport of U dissolved from igneous source rocks typically occurs under

oxidizing conditions when U is present as the uranyl ion U (VI) (UO_2^{+2})^a. U is substantially less soluble under reducing

^a The oxidized uranyl(VI) species will be expressed hereafter as UO_2^{++} .

conditions, when it is typically present as aqueous $U(OH)_4$.

Table 1. Typical uranium contents of various rock types².

Rock Type	Range of Concentrations (ppm)
Igneous	
Diorite and Quartz Diorite	0.5-12
Alkaline Intrusive	0.04-20
Mafic	0.1-3.5
Sedimentary	
Sandstone	0.5-4
Orthoquartzite	0.2-0.6
Carbonate Shale	0.1-10
Black shale	1-15
Lignite	3-1250
Phosphorite	10-2500
Large Scale U Sources	
	Average Concentration (ppm)
High-grade U deposit	10^4 - 10^5
High-heat production granite	10
Continental Crust	1
Bulk Earth	2×10^{-2}

Figure 1 shows uranyl speciation under oxidizing conditions ($fO_2 = 0.2$ atm) and $pCO_2 = 10^{-3.5}$. Note the prevalence of UO_2^{2+} -carbonate complexes at $pH \geq 7$. In the absence of CO_2 , but at the same fO_2 , uranyl-hydroxy ions predominate at $pH > 5$. Under nominally reducing conditions ($\log fO_2 \approx -70$ atm), $25^\circ C$, and low total U concentrations (e.g., $[U]_T < 10^{-14}$ m), $U(IV)$ forms U -hydroxo complexes at $pH \geq 6$ (Fig. 2 and table 2). However, when $[U]_T > 10^{-14}$ m, uraninite (UO_2)_{cr} becomes stable.

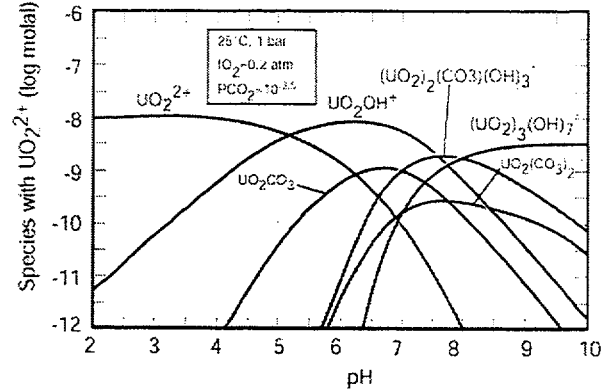


Figure 1 Uranyl speciation at $25^\circ C$ and 1 bar as a function of pH under oxidizing conditions; $fO_2 = 0.2$ atm and $pCO_2 = 10^{-3.5}$ atm. Uranium speciation data is from the Geochemical Workbench (GWB) software package³.

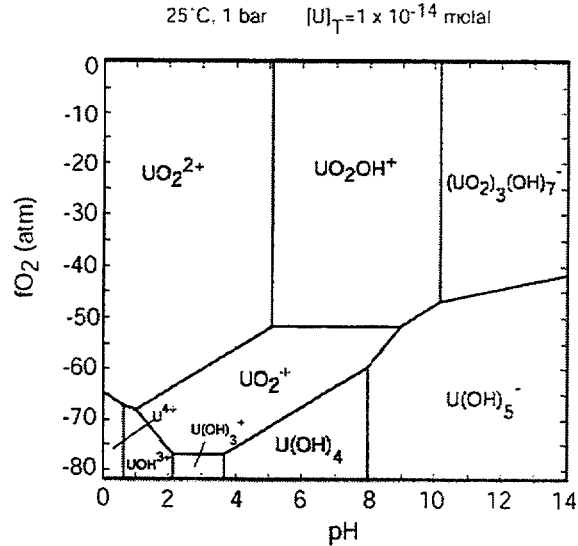


Figure 2. Uranium speciation at $25^\circ C$ and 1 bar as a function of fO_2 and pH ; $[U]_{total} = 10^{-14}$ molal. Uranium speciation data as in figure 1.

2.1 Ore Formation and Weathering

Subsurface accumulations of uranium (ore bodies) tend to occur at redox fronts where oxidizing U -rich solutions encounter electron-rich solids, typically organic, that are able to reduce soluble $U(VI)$ to insoluble $U(IV)$ -containing minerals. An exception is insoluble uranyl vanadates that form many of the ores found in the

Table 2. Equilibrium constants (K) for selected U(VI) aqueous complexes.

Chemical Reaction	Log K (25°C and 1 bar) ^{3,4}
$\text{UO}_2^{2+} + 2\text{H}^+ \Leftrightarrow \text{U}^{4+} + \text{H}_2\text{O} + 0.5\text{O}_2(\text{g})$	-33.78 ^a
$\text{UO}_2\text{OH}^+ + \text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + \text{H}_2\text{O}$	5.091 ^a
$\text{UO}_2(\text{OH})_2, \text{aq} + 2\text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + 2\text{H}_2\text{O}$	11.5 ^b
$\text{UO}_2(\text{OH})_3^- + 3\text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + 3\text{H}_2\text{O}$	20.00 ^b
$\text{UO}_2(\text{OH})_4^{2-} + 4\text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + 4\text{H}_2\text{O}$	33.00 ^b
$(\text{UO}_2)_2\text{OH}^{3+} + \text{H}^+ \Leftrightarrow 2\text{UO}_2^{2+} + \text{H}_2\text{O}$	2.70 ^b
$(\text{UO}_2)_2(\text{OH})_2^{2+} + 2\text{H}^+ \Leftrightarrow 2\text{UO}_2^{2+} + 2\text{H}_2\text{O}$	5.68 ^a
$(\text{UO}_2)_3(\text{OH})_4^{2+} + 4\text{H}^+ \Leftrightarrow 3\text{UO}_2^{2+} + 4\text{H}_2\text{O}$	11.90 ^b
$(\text{UO}_2)_3(\text{OH})_5^{5+} + 5\text{H}^+ \Leftrightarrow 3\text{UO}_2^{2+} + 5\text{H}_2\text{O}$	15.82 ^a
$(\text{UO}_2)_3(\text{OH})_7^- + 7\text{H}^+ \Leftrightarrow 3\text{UO}_2^{2+} + 7\text{H}_2\text{O}$	28.337 ^a
$(\text{UO}_2)_4(\text{OH})_7^+ + 7\text{H}^+ \Leftrightarrow 4\text{UO}_2^{2+} + 7\text{H}_2\text{O}$	21.9 ^b
$\text{UO}_2(\text{cr}) + 4\text{H}^+ \Leftrightarrow \text{U}^{4+} + 2\text{H}_2\text{O}$	-4.638 ^a
$\text{UO}_2^+ + 3\text{H}^+ \Leftrightarrow \text{U}^{4+} + 1.5\text{H}_2\text{O} + 0.25\text{O}_2(\text{g})$	-15.07 ^a
$\text{U}(\text{OH})_3^+ + 3\text{H}^+ \Leftrightarrow \text{U}^{4+} + 3\text{H}_2\text{O}$	4.88 ^a
$\text{U}(\text{OH})_4 + 4\text{H}^+ \Leftrightarrow \text{U}^{4+} + 4\text{H}_2\text{O}$	8.534 ^a
$\text{U}(\text{OH})_5^- + 5\text{H}^+ \Leftrightarrow \text{U}^{4+} + 5\text{H}_2\text{O}$	16.498 ^a
$\text{U}(\text{OH})_3^+ + \text{H}^+ \Leftrightarrow \text{U}^{4+} + \text{H}_2\text{O}$	0.6494 ^a
$\text{UO}_2\text{CO}_3, \text{aq} \Leftrightarrow \text{UO}_2^{2+} + \text{CO}_3^{2-}$	-9.67 ^b
$\text{UO}_2\text{CO}_3, \text{aq} + \text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + \text{HCO}_3^-$	0.694 ^a
$\text{UO}_2(\text{CO}_3)_2^{2-} \Leftrightarrow \text{UO}_2^{2+} + 2\text{CO}_3^{2-}$	-16.94 ^b
$\text{UO}_2(\text{CO}_3)_2^{2-} + 2\text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + 2\text{HCO}_3^-$	3.608 ^a
$\text{UO}_2(\text{CO}_3)_3^{4-} \Leftrightarrow \text{UO}_2^{2+} + 3\text{CO}_3^{2-}$	-21.60 ^b
$\text{UO}_2(\text{CO}_3)_3^{4-} + 3\text{H}^+ \Leftrightarrow \text{UO}_2^{2+} + 3\text{HCO}_3^-$	9.33 ^a
$(\text{UO}_2)_3(\text{CO}_3)_6^{6-} \Leftrightarrow 3\text{UO}_2^{2+} + 6\text{CO}_3^{2-}$	-54.0 ^b
$(\text{UO}_2)_2\text{CO}_3(\text{OH})_3^- + 4\text{H}^+ \Leftrightarrow 2\text{UO}_2^{2+} + \text{HCO}_3^- + 3\text{H}_2\text{O}$	11.524 ^a
$(\text{UO}_2)_3\text{CO}_3(\text{OH})_3^+ + 3\text{H}^+ \Leftrightarrow 3\text{UO}_2^{2+} + \text{CO}_3^{2-} + 3\text{H}_2\text{O}$	-0.66 ^b
$(\text{UO}_2)_{11}(\text{CO}_3)_6(\text{OH})_{12}^{2-} + 12\text{H}^+ \Leftrightarrow 11\text{UO}_2^{2+} + 6\text{CO}_3^{2-} + 12\text{H}_2\text{O}$	-36.43 ^b

^aBethke³; ^bDavis⁴

Colorado Plateau in the USA⁵. Uranyl vanadates form during oxidative alteration of reduced U- and V-bearing minerals. Other reported U(VI) minerals that are not as common as the vanadates are the uranyl molybdates, tungstates, sulfates, selenites, and tellurites⁵. The most common U(IV)-bearing phases are uraninite (UO₂), pitchblende (highly impure uraninite),

coffinite (USiO₄·nH₂O), and Brannerite (U,Ca, Y, Ce)(Ti, Fe)₂O₆(OH)), all of which are typically found in U ore deposits (table 3). Other common uranyl phases, such as uranophane (Ca(UO₂)₂(SiO₃OH)₂(H₂O)₅), while limited in association to the presence of altered nuclear fuel rods and naturally weathered uraninite, are structurally and chemically complex (table 3).

Table 3. Common uranium-bearing minerals in rocks (after Finch and Murakami, 1999)⁵.

Phase	Chemical Formula	Comments
Reduced U(IV) Phases		
Uraninite	UO _{2+x}	Not purely stoichiometric in nature. It can incorporate minor REE and other cations.
Pitchblende	UO _{2+x}	Very finely grained highly impure uraninite; very common U ore phase
Coffinite	USiO ₄ ·nH ₂ O	Zircon structure, most important ore mineral after uraninite
Brannerite	(U,Ca, Y, Ce)(Ti, Fe) ₂ O ₆ (OH)	Metamict; most common ore phase after uraninite and coffinite
Orthobrannerite	(U ⁶⁺ , U ⁴⁺)(Ti, Fe) ₂ O ₆ (OH)	Metamict; mixed valence U phase
Ianthinite	U ⁴⁺ (U ⁶⁺ O ₂)O ₄ (OH) ₆ (H ₂ O) ₉	Mixed valence U phase; structure similar to β-U ₃ O ₈ ; it oxidizes to schoepite
Uranyl Oxyhydroxides		
Schoepite	(UO ₂) ₈ O ₂ (OH) ₁₂ (H ₂ O) ₁₂	Common uranyl phase
Metaschoepite	(UO ₂) ₈ O ₂ (OH) ₁₂ (H ₂ O) ₁₀	Partially dehydrated schoepite
Becquerilite	Ca(UO ₂) ₆ O ₄ (OH) ₆ (H ₂ O) ₈	It likely alters to schoepite at low pH's (Finch and Murakami, 1999) ⁵
Clarkeite	(Na,Ca)(UO ₂)(O,OH)(H ₂ O) _n (n=0-1)	-
Uranyl Carbonates		
Blatonite	UO ₂ CO ₃ ·H ₂ O	Monocarbonate; not a dehydration product of jolilite
Jolilite	UO ₂ CO ₃ ·nH ₂ O (n≈2)	Monocarbonate
Rutherfordine	UO ₂ CO ₃	Monocarbonate
Uranalcarite	Ca ₂ (UO ₂) ₃ (CO ₃)(OH) ₆ (H ₂ O) ₃	Monocarbonate
Wyartite	CaU ⁵⁺ (CO ₃)(UO ₂)O(OH) ₄ (H ₂ O) ₇	Monocarbonate; first known U phase to have pentavalent U

Table 3 (cont.). Common U-bearing minerals in rocks (after Finch and Murakami, 1999)⁵

Phase	Chemical Formula	Comments
Uranyl Carbonates (cont.)		
Fontanite	$\text{Ca}(\text{UO}_2)_3(\text{CO}_3)_4(\text{H}_2\text{O})_3$	Monocarbonate
Zellerite	$\text{Ca}(\text{UO}_2)_3(\text{CO}_3)_2(\text{H}_2\text{O})_5$	Dicarbonate
Metazellerite	$\text{Ca}(\text{UO}_2)_3(\text{CO}_3)_2(\text{H}_2\text{O})_n$ ($n < 5$)	Dicarbonate; dehydrated zellerite
Znucallite	$\text{CaZn}_{11-12}(\text{UO}_2)_3(\text{CO}_3)_3(\text{OH})_{20 \&-22}(\text{H}_2\text{O})_4$	Tricarbonate
Uranyl Silicates		
Uranophane	$\text{Ca}(\text{UO}_2)_2(\text{SiO}_3\text{OH})_2(\text{H}_2\text{O})_5$	Possibly the most common U mineral after uraninite
Soddyite	$(\text{UO}_2)_2\text{SiO}_4(\text{H}_2\text{O})_2$	-
Ursilite	$(\text{Mg,Ca})_4(\text{UO}_2)_4(\text{Si}_2\text{O}_5)_5(\text{OH})_6(\text{H}_2\text{O})_{15}$	-
Uranosilite	$(\text{UO}_2)\text{Si}_7\text{O}_{15}(\text{H}_2\text{O})_{0-1}$	Rare mineral
Sklodowskite	$\text{Mg}(\text{UO}_2)_2(\text{Si}_3\text{OH})_2(\text{H}_2\text{O})_6$	-
Weeksite	$\text{K}_{1-x}\text{Na}_x(\text{UO}_2)_2(\text{Si}_5\text{O}_{13})(\text{H}_2\text{O})_4$ ($x=0.4$)	-
Haiweeite	$\text{Ca}(\text{UO}_2)_2[\text{Si}_5\text{O}_{12}(\text{OH})_2](\text{H}_2\text{O})_{4.5}$	-
Uranyl Phosphates and Arsenates (Autunite group)		
Autunite	$\text{Ca}[(\text{UO}_2)(\text{PO}_4)]_2(\text{H}_2\text{O})_{10-12}$	-
Saleeite	$\text{Mg}[(\text{UO}_2)(\text{PO}_4)]_2(\text{H}_2\text{O})_{10}$	-
Novacekite	$\text{Mg}[(\text{UO}_2)(\text{AsO}_4)]_2(\text{H}_2\text{O})_8$	-
Vochtenite	$(\text{Fe}^{2+}, \text{Mg})\text{Fe}^{3+}[(\text{UO}_2)(\text{PO}_4)]_4(\text{OH})(\text{H}_2\text{O})_{12-13}$	-

Table 3 (cont.). Common U-bearing minerals in rocks (after Finch and Murakami, 1999)⁵.

Phase	Chemical Formula	Comments
Uranyl Vanadates		
Carnotite	$K_2(UO_2)_2(V_2O_8)(H_2O)_3$	Carnotite group mineral
Tyuyamunite	$Ca(UO_2)_2(V_2O_8)(H_2O)_8$	Carnotite group mineral
Metatyuyamunite	$Ca(UO_2)_2(V_2O_8)(H_2O)_8$	Carnotite group mineral
Uvanite	$(UO_2)_2V_6O_{17} \cdot 15H_2O$	-
Uranyl Molybdates		
Calcurmolite	$Ca(UO_2)_3(MoO_4)_3(OH)_2(H_2O)_{11}$	-
Cousinite	$Mg(UO_2)_2(MoO_4)_2(OH)_2(H_2O)_5$	-
Iriginite	$(UO_2)Mo_2O_7(H_2O)_3$	-
Tengchongite	$Ca(UO_2)_6(MoO_4)_2O_5(H_2O)_{12}$	-
Umohoite	$(UO_2)MoO_4(H_2O)_4$	-
Uranyl Sulfates, Selenites, and Tellurites		
Deliensite	$Fe(UO_2)_2(SO_4)_2(OH)_2(H_2O)_3$	-
Rabejacite	$Ca(UO_2)_4(SO_4)_2(OH)_6(H_2O)_6$	-
Uranopilite	$(UO_2)_6(SO_4)(OH)_{10}(H_2O)_{12}$	-
Zippeite	$K_4(UO_2)_6(SO_4)_3(OH)_{10}(H_2O)_4$	-
Haynesite	$(UO_2)_3(OH)_2(SeO_3)_3(H_2O)_5$	-
Cliffordite	UTe_3O_9	-
Schmitterite	$(UO_2)TeO_3$	-

Oxidative transformation of uraninite and oxidation/corrosion of synthetic UO_2 (spent nuclear fuel) are similar processes. Differences in solubilities between synthetic and natural UO_2 are observed though, and are believed to be caused by carbonate impurities in the natural samples⁶. Exposure to water versus dry air also affects UO_2 oxidation rates⁷. Typically the slow rate of oxygen diffusion at low temperatures makes the process rather sluggish⁵. Consequently, many studies are performed at relatively high temperatures ($T > 150^\circ\text{C}$)⁷. In general, uranyl mineral formation is preceded by the appearance of intermediate U oxides such as U_3O_7 and U_3O_8 irrespective of temperature, even though the latter minerals have yet to be observed in nature⁵.

2.2 Microbes and U

Microorganisms can affect the transport and deposition of uranium by, respectively, producing dissolved compounds, e.g., organic acids, capable of chelating uranium⁸, or by reducing uranyl to less soluble U(IV) minerals⁹. Also, U can sorb onto bacterial cell walls and migrate. Uranyl-chelates that are anionic, e.g. uranyl-citrate, tend to sorb to soil surfaces less than bare uranyl, and are therefore favored to be transported further. Microbial breakdown of uranyl chelates often determines maximum transport distance of microbially mobilized U and for this reason remains an important area of research^{8; 10}. Microbial reduction of U has been demonstrated using groundwater from the Tuba City, AZ (U.S.A) U mill tailing site¹¹. Column experiments with groundwater and sand sampled at this site showed a dramatic lowering of U(VI)

concentration from 250 to 14 $\mu\text{g/L}$ in a period of 21 days at 24°C and 0.5 $\mu\text{g/L}$ after 2 months in the presence of indigenous microorganisms¹¹⁻¹³. In these experiments, microorganisms were activated by addition of ethanol and metaphosphate to the groundwater/sand mixture.

2.3 Adsorption

After precipitation, adsorption is the most important sink for U in natural systems^{4; 14-17}. A large number of studies have applied surface complexation models to explain U(VI) sorption on mineral surfaces. Tripathi¹⁸ and Hsi and Langmuir¹⁹ studied U uptake on iron oxides. Redden et al.²⁰ studied U(VI) adsorption on goethite, gibbsite, and kaolinite in the presence and absence of citric acid. Redden et al.²⁰, Payne et al.²¹, Payne and Waite¹⁶, and Payne and Waite¹⁷ studied U(VI) sorption onto kaolinite and ferrihydrite, and Pabalan et al.¹⁵ studied U(VI) sorption onto α -alumina (aluminum oxide), Na-clinoptilolite, and quartz. Davis⁴ combined surface complexation modeling and structural characterization of U(VI) on a variety of minerals. Jenne²², and more recently, Krupka et al.²³, examined U(VI) adsorption K_d 's obtained from sediment samples and those from various mineral adsorption studies reported in the literature. All these studies demonstrate that the peak for U(VI) uptake is at the near-neutral pH range of $\sim 6 - 7.5$ ²⁴. The sorption peak for phosphates is much lower - pH 2.5 to 4²⁵. Valsami-Jones et al.²⁶ and Arey et al.²⁷ showed that hydroxyapatite can sequester U inside its mineral structure.

In general, U(VI) sorption proceeds in the sequence: montmorillonite (clay) \approx kaolinite (clay) \approx gibbsite \approx goethite $>$ clinoptilolite $>$ α -alumina $>$ quartz $>$ phosphates^{15; 22; 25}. Relative sorption of U tends to depend on the type of aqueous complexes it forms in solution^{4; 15; 18; 19}. At high pH's, where anionic uranyl-carbonate complexes predominate, U is only weakly sorbed due to electrostatic repulsion by negatively charged mineral surfaces e.g.^{4; 12}. Uranyl-hydroxy complexes predominate in the pH range between 6-8 along with the mixed uranyl hydroxy carbonate complex^{4; 19}. When carbonate concentrations are low or absent, the predominant sorbing species are the uranyl-hydroxy complexes, e.g., $(\text{UO}_2)_3(\text{OH})_5^+$ ⁴. Krupka et al.²³ critically reviewed the environmental chemistry of uranium and showed that U sorption is primarily controlled by pH and carbonate levels. Table 4 gives Krupka et al.'s.²³ maximum and minimum soil Kd values. Despite the wide variation in Kd values, the results give a reasonably clear picture of U sorption trends on soil minerals. Irreversible sorption of U typically occurs when iron and manganese oxides are

present. Typically the irreversible fraction of U rarely exceeds 10% of the total²⁸.

2.4 Colloids

Colloidal transport of U has been investigated adjacent to both natural ore deposits and mill tailings^{12; 29-32}. Colloids are not an important transport mechanism for U. Colloidal transport depends on the amount and type of colloid present, its capacity to migrate in the media, and the predominant U aqueous species in the soil solution. The predominance of negatively charged uranyl-carbonate species often prevents adsorption onto negatively charged particulate material. Mielekey et al.²⁹ analyzed particulate material in waters at Poços de Caldas mine and observed that even when relatively large amounts of U and Th were associated with colloids, net transport was minor due to low colloidal concentrations in the waters flowing through the aquifer. A similar situation is observed at the Cigar Lake ore deposit in Canada where relatively low concentrations of colloids limit U migration even when the U concentrations in the particulate phase and waters are similar³⁰.

Table 4. Minimum and maximum soil uranium Kd's (after Krupka et al., 1999)²³

Kd (ml/g)	pH							
	3	4	5	6	7	8	9	10
Minimum	<1	0.4	25	100	63	0.4	<1	<1
Maximum	32	5000	160,000	1,000,000	630,000	250,000	7,900	5

3.0 Uranium Plumes

U-contaminated sites can be separated into three groups:

1. Uranium Mill Tailings Remedial Action (UMTRA) sites in the USA - sites where U ore was processed.
2. Refined U releases - sites where U was released during explosive testing or nuclear waste storage activities.
3. Natural analogues - natural uranium ore deposits.

3.1 Artificial Plumes

The UMTRA sites served as U ore processing and milling plants and typically witnessed the use of acid/alkaline leaching, sand-slime separation, and ion-exchange recovery processes (lime, sulfuric acid, nitric acid, sodium carbonate, kerosene, and ammonia gas were often used as part of the leaching processes). The UMTRA sites are classified in two types: Title I and Title II. Title I sites are those in which remedial action such as removal and relocation of the mill tailing piles has been performed. Conversely, Title II sites still have their mill tailings in place and remedial action or further mining and processing remain in consideration. There are 24 Title I UMTRA sites located mostly in the western United States (Fig. 3). Most were established in the 1950's and comprise the bulk of the U-plume data obtained as many of them have been the subject of recent (~15 years) monitoring and remediation. In general, these sites contained (Title I) or still containing (Title II) piles of mill tailings and other contaminated material forming alluvium-like surface deposits where the bulk of the material has been removed and relocated elsewhere in

specially engineered repositories. Title II UMTRA sites still have their mill tailings in place which typically contribute high levels of U to aquifers near the tailing source. This appears to result in longer plumes. The Title II sites will be discussed briefly because temporal monitoring of groundwater chemistry is scarce and evaluation of existing literature data is still in progress. Natural recharge through tailings piles is (or was) the primary source of dissolved U. Most of the aquifers beneath mill piles are contaminated, exceeding MCL levels of U (MCL = Maximum Concentration Limit defined by the U.S. Environmental Protection Agency = 44 ppb for U)³³. Additional contaminants in the effluent include lead, cadmium, molybdenum, nitrate, ammonium, and selenium. Some nearby rivers to Title I UMTRA sites are contaminated with U, but not at dangerous levels. For the majority of UMTRA sites, deep aquifers (even those close to the processing plants) do not show high levels of contamination.

Groundwater chemistry can be highly variable from site to site. Some sites, such as Riverton, Wyoming^{34; 35} and Monument Valley, Arizona³⁶ primarily contain uranyl carbonate (uranyl bi- and tri-carbonate) species whereas others, such as Falls City, Texas, have lower pH's (pH 2-5) in certain areas of the site, and have uranyl-hydroxyl complexes. Nevertheless, except for the anomalously low pH site of Falls City (Texas), the groundwater chemistry between sites is quite similar as will be exemplified later in the text. White et al.³⁴ envisioned three general transport mechanisms for U, which may well apply to all UMTRA sites:

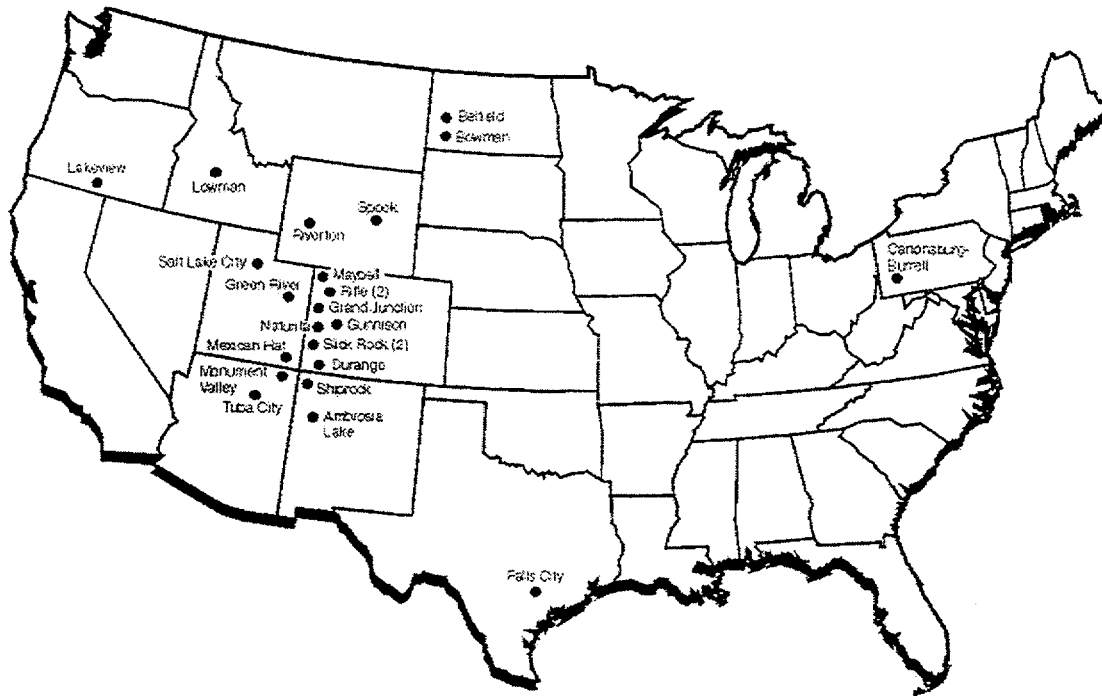


Figure 3. UMTRA Ground Water Project Title I site locations ³⁷.

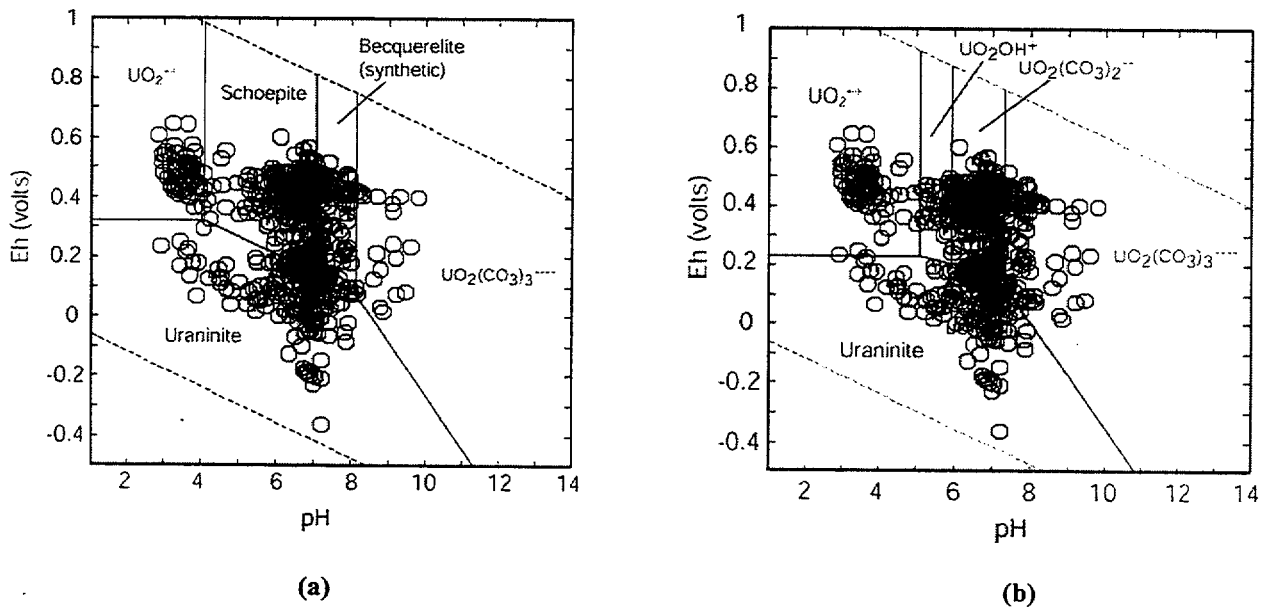


Figure 4a and b. Eh-pH diagram at 25°C and 1 bar showing U phase boundaries and data from nine title I UMTRA sites ($n = 778$ data points; see text). It is assumed that $[U]_{total} = [UO_2^{2+}]$ (molal units) in generating this diagram. (a) $[U]_{total} = 10^{-3}$ molal, $[Ca^{++}] = 10^{-1.8}$ molal, and $fCO_2 = 10^{-2.5}$ atm. Schoepite starts appearing at $[U] = 10^{-6.5}$ molal.; (b). $[U]_{total} = 10^{-7}$ molal, $[Ca^{++}] = 10^{-1.8}$ molal, and $fCO_2 = 10^{-2.5}$ atm. Uranium speciation data as in figure 1. Notice in this figure that only uranyl aqueous species are depicted in the diagram. Becquerelite (synthetic) solubility data is from Vochten and Van Haverbeke ³⁸

(1) initial dewatering of the slurry material in the tailings pile and concomitant downward flow soon following deposition, (2) mixing and dilution of the tailing pore waters with local ground and seepage waters, and (3) periodic meteoric water recharge into the tailing pile by precipitation or snowmelt. Subsequent U transport through the alluvium will be highly dependent upon solution chemistry (i.e., pH, ionic strength, and composition) and redox state of the system. Bulk groundwater Eh (redox potential), U, and pH analyses from nine Title I UMTRA sites taken over a 3-8 year time span are shown in figures 4 through 7. The database source is from the DOE-GJO Site Environmental Evaluation Uranium Mill Tailings Remedial Action (SEE UMTRA database). Note first of all that the Eh-pH data extends into the uranyl (UO_2^{++}), uranyl hydroxide (UO_2OH^+), and uranyl-carbonate $\text{UO}_2(\text{CO}_3)_2^{2-}$ species fields when $[\text{U}] = 10^{-7}$ m and $[\text{Ca}^{++}] \sim 10^{-2}$ m (Fig. 4a). If $[\text{U}] = 10^{-3}$ m and with the same Ca concentration (Fig.

4b), all of the data falls between the stability fields of uraninite (UO_2), schoepite, becquerelite, and uranyl fields. If the most scattered data represented by the Falls City (Texas) site is removed, the other 8 sites show a cluster of data points forming a characteristic vertical trend plotting at a pH ~ 7 (Fig. 5). The Falls City, Texas data set differs from the others by possessing a bimodal pH distribution³⁹ (Fig. 6) and a pH-independent $[\text{U}]$ level opposite to the other UMTRA sites. Bimodal pH distributions are not uncommon in contaminated groundwater aquifers associated with mill tailings or mining sites that have been widely exposed to pervasive *in-situ* leaching (ISL). Other examples of these are Tuba City (Arizona), Konigstein (Saxony, Germany), and Split Rock (Wyoming - Title II). At all these sites, the low pH groundwater is spatially associated with the mill tailing repository area and/or chemically associated with

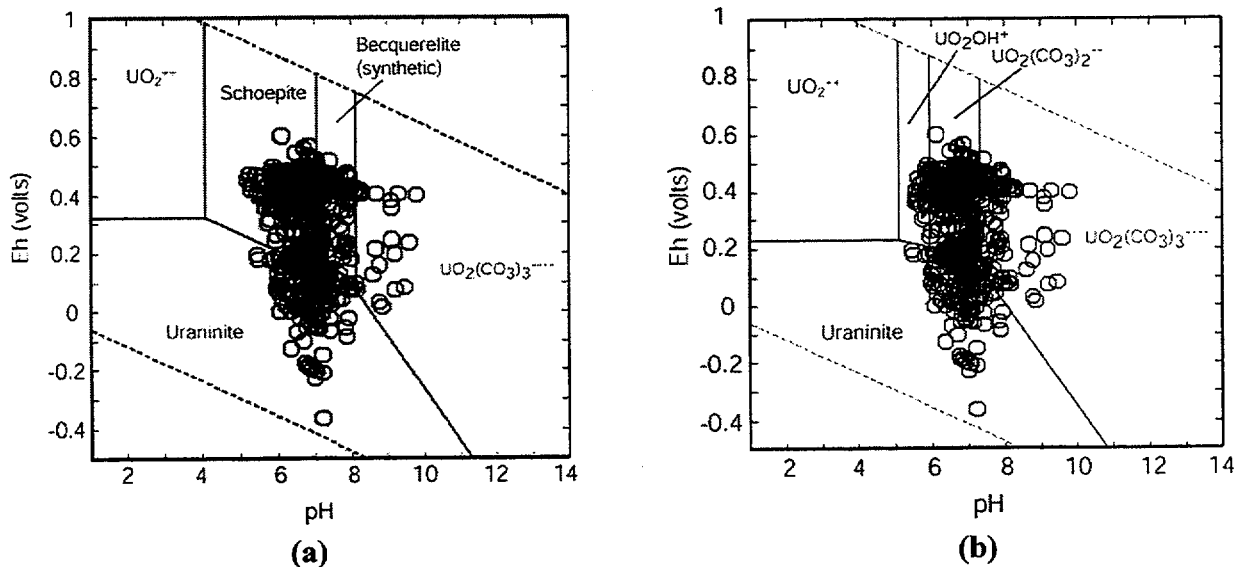


Figure 5a and b. Eh-pH diagram at 25°C and 1 bar showing U phase boundaries and data from eight title I UMTRA sites (Falls City (Texas) data removed; n = 634 data points). It is assumed that $[\text{U}]_{\text{total}} = [\text{UO}_2^{++}]$ (molal units) in generating this diagram. (a) $[\text{U}]_{\text{total}} = 10^{-3}$ molal, $[\text{Ca}^{++}] = 10^{-1.8}$ molal, and $f\text{CO}_2 = 10^{-2.5}$ atm. Notice that solid uranyl phases appear in the diagram for this $[\text{U}]_{\text{total}}$; (b) $[\text{U}]_{\text{total}} = 10^{-7}$ molal, $[\text{Ca}^{++}] = 10^{-1.8}$ molal, and $f\text{CO}_2 = 10^{-2.5}$ atm. Schoepite starts appearing at $[\text{U}]_{\text{total}} = 10^{-6.5}$ molal. The uranyl species field covered by the data points is $\text{UO}_2(\text{CO}_3)_2^{2-}$. Uranium speciation data as in figure 1. Becquerelite (synthetic) solubility data as in figure 4.

residual mine leaching waters. Even though the low pH waters in Falls City (Texas) and the above mentioned sites usually contain significant amounts of SO_4^- and PO_4^- , U does not exceed the maximum range of concentrations of $\sim 10^{-4} - 10^{-5}$ m common to all the studied mill tailing sites. Therefore, the low pH seen in these anomalous groundwater may reflect the presence of residual sulfate remaining in the aquifer pores as precipitates or as fluid pockets which were dispersed by present water recharges or remediation activities.

The vertical trend in Fig. 5 originates from the uraninite stability field overlapping in a parallel fashion to the schoepite-becquerelite equilibrium boundary (Fig. 5a) or plotting inside the field of the U-carbonate complex (Fig. 5b) using a total

CO_2 fugacity ($f\text{CO}_2$) of $10^{-2.5}$ atm typical of soils. If a lower CO_2 fugacity of $10^{-3.5}$ atm typical of equilibrium with the atmosphere is considered, the becquerelite- $\text{UO}_2(\text{CO}_3)_3^{4-}$ equilibrium boundary in Fig. 5a shifts to a lower pH diminishing the becquerelite stability field. High levels of dissolved organic carbon in UMTRA groundwater (~ 200 ppm) suggest that a relatively high $f\text{CO}_2$ value of $10^{-2.5}$ atm is a reasonable input to use in the chemographic analyses below. The becquerelite solubility value at 25°C and 1 bar chosen for this analysis is taken from Vochten and Van Haverbeke ³⁸ for a synthetic material. Casas et al. ⁴⁰ and Sandino and Grambow ⁴¹ reported becquerelite solubility values for a natural sample from the Shinkolobwe U deposit in

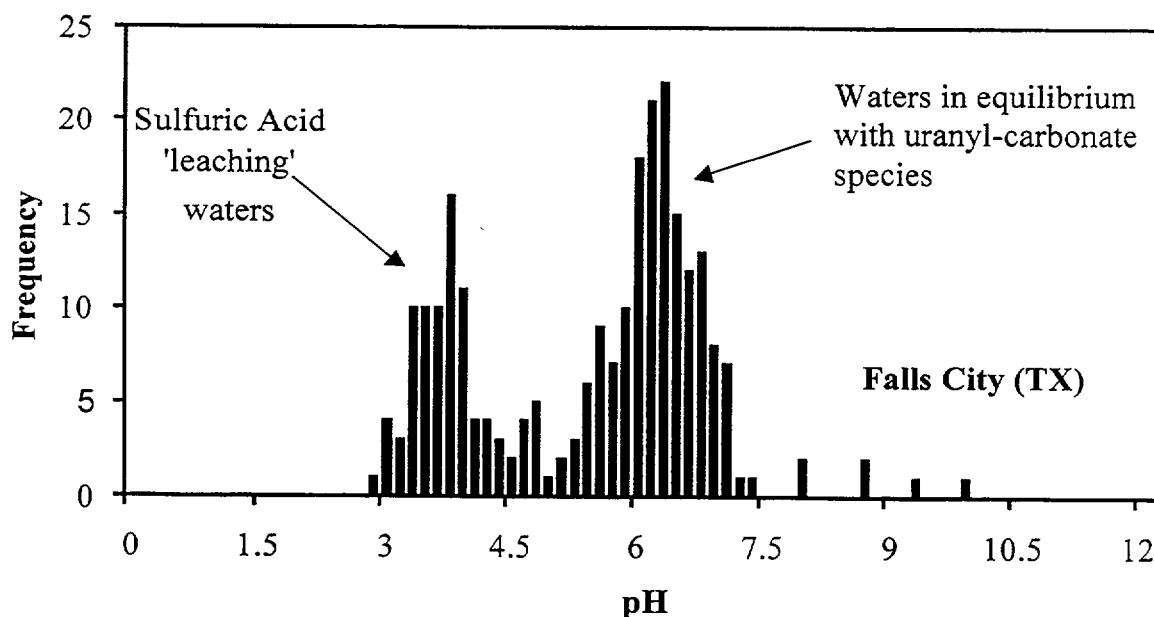


Figure 6. Bimodal distribution of pH for groundwater measured at Falls City (Texas) Title I UMTRA site (n = 249 data points; see text).

Zaire (Africa) and from the equilibria during the transformation of synthetic schoepite to becquerelite, respectively. The lower solubility value obtained by Casas et al.⁴⁰ for natural becquerelite predicts supersaturation of the mineral with the groundwater chemistry at the considered UMTRA sites. However, the much larger solubility values obtained for synthetic becquerelite^{38; 41} are somewhat similar and show a near equilibrium relationship between schoepite-becquerelite for the average Ca concentrations observed in these groundwaters. This is a more realistic association considering the commonly observed presence of schoepite as a hydration product of U oxide in natural and synthetic systems. The reason for the gross oversaturation, with respect to natural becquerelite, is that Casas et al.⁴⁰ obtained a solubility value for natural becquerelite

that is considerably lower than those obtained for synthetic analogues, possibly a result of lower surface areas present in macroscopic crystals yielding much lower solubilities.

Figures 7a and 7b show U mineral solubilities as a function of pH and was constructed using a mean value of 0.0158 m for the maximum Ca concentrations obtained from several groundwater analyses at various UMTRA sites. Using a median value of 0.0065 m for all Ca concentrations at the UMTRA sites that were considered, causes a negligible enlargement of the schoepite field in Fig. 7. These figures show that for the assumed $f\text{CO}_2$ value of $10^{-2.5}$ atm the majority of data points fall within the stability field of the $\text{UO}_2(\text{CO}_3)_2^{2-}$ species. Using a lower

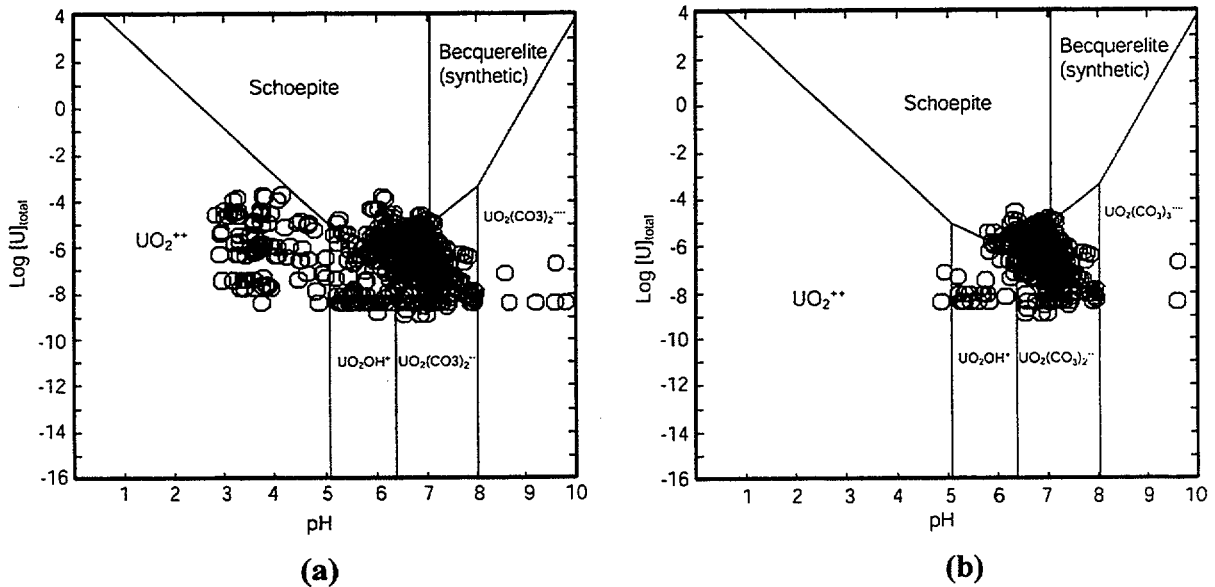
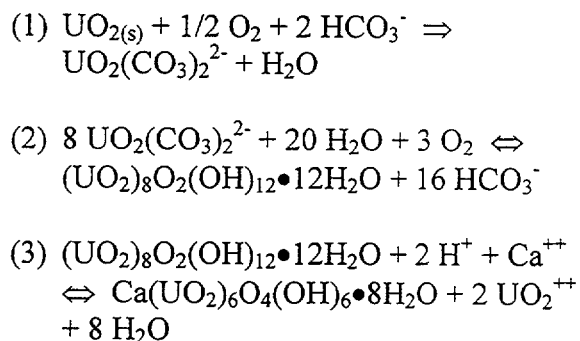


Figure 7a and b. Uranyl phase solubilities as a function of pH at 25°C and 1 bar. It is assumed that $[\text{U}]_{\text{total}} = [\text{UO}_2^{++}]$ (molal units) in generating this diagram. (a) $[\text{Ca}^{++}] = 10^{-1.8}$ molal and $f\text{CO}_2 = 10^{-2.5}$ atm (all data from the Title I UMTRA sites; $n = 778$ data points); (b) $[\text{Ca}^{++}] = 10^{-1.8}$ molal and $f\text{CO}_2 = 10^{-2.5}$ atm (no Falls City (Texas) data; $n = 634$ data points). Uranium speciation data as in figure 1. Becquerelite (synthetic) solubility as in figure 4. Notice the absence of data below $[\text{U}]_{\text{total}} \approx 10^{-8}$ molal which denotes the lower level of analytical detection.

$f\text{CO}_2$ value of $10^{-3.5}$ atm would extend the schoepite field to lower uranyl concentration values causing some of the data points to fall well inside this field. In general, the maximum uranyl concentrations correspond to the schoepite- $\text{UO}_2(\text{CO}_3)_2^{2-}$ equilibrium suggesting a solubility and, therefore, transport control on dissolved uranyl in the most contaminated aquifers that are adjacent to the former mill tailing impoundments. The close association of schoepite and becquerelite during the alteration of uranium oxide has been previously reported in the literature ^{38; 41-45}. Becquerelite is considered to be a thermodynamically stable product of schoepite (or meta-schoepite) alteration in the presence of alkaline elements such as Ca ^{45; 46}. Both phases have been observed to coexist in shallow weathered zones of natural uraninite deposits, and in short- and long-term UO_2 corrosion experiments simulating alteration of anthropogenic uranium solids like nuclear spent fuel ^{5; 41; 44-46}. However, their existence in groundwater reservoirs directly associated with mill tailing sites still needs to be confirmed. In the lack of unambiguous evidence for noticeable weathering mineralization in the studied mill tailing sites (Title I and II UMTRA sites in this case), we propose that the uranyl-hydroxy and uranyl-carbonate complexes such as $\text{UO}_2(\text{CO}_3)_2^{2-}$ and $\text{UO}_2(\text{OH})^+$, respectively, along with their intrinsic local carbonate inputs near and far from the tailings impoundment, would be the most likely aqueous species to buffer the groundwater chemistry to neutral pH's. Maximum U concentrations observed at various contaminated aquifers in UMTRA sites might be controlled by the schoepite-uranyl-carbonate (possibly in combination with becquerelite) equilibrium at near neutral pH's. This observation is also

applicable to Title II UMTRA sites where the mill tailings are still in place and some milling operations along with possible mining activities also remain active. Dilution and sorption are the most likely mechanisms responsible for lowering U concentrations away from the plume source.

The reaction mechanisms described before, based on the large DOE Grand Junction Project Office groundwater chemical database (SEE UMTRA - Title I UMTRA sites), plus recent DOE and the Nuclear Regulatory Commission (NRC) public reports (Title II UMTRA sites), clearly suggest a reaction path controlled by uraninite dissolution and movement of groundwater compositions towards the schoepite (or meta-schoepite)/fluid equilibrium at a pH between ~ 6.5 and ~ 7.5 . As shown in Fig. 5a, the proposed path begins with oxidative dissolution of uraninite to uranyl-carbonate (reaction 1) and ends with the equilibrium between schoepite and $\text{UO}_2(\text{CO}_3)_2^{2-}$ carbonate species (reaction 2). Equilibrium between schoepite and becquerelite may occur in the presence of Ca (reaction 3).



The second type of U plumes arise from releases of refined U, such as those involving explosive activity or nuclear waste storage. One located at Lawrence Livermore National Laboratory (LLNL) involved explosive activity in pits and is

still under investigation. Preliminary U well data for this site obtained near the explosive pits show plumes lengths of ~0.5 kilometer measured to the 11 pCi/L (≈ 16 ppb) contour. Some important nuclear waste sites that are under current investigation include the Hanford Tank Farm ⁴⁷ in Washington and the Savannah River Site (SRS) in Georgia, but were not considered in the present study for several reasons: (1) reports on groundwater chemistry needed to assess plume length are not yet available (SRS) or are in the process of being updated or published (Hanford), and (2) the monitoring well network distribution in some cases (e.g., tank farm at Hanford) is not adequate for the purpose of this study. At this moment, the SRS study has not reported U plumes beyond the tank farm and most of the currently available data is for Tritium and TCE plumes ⁴⁸. Even when there are insufficient monitoring wells at a distance from the Hanford tanks to accurately estimate further plume extensions beyond the source, we report only one relatively large U plume in one process trench (see table 5).

3.2 Natural Plumes

Natural uranium ore deposits are often used as analogues for long-term high level nuclear waste repositories but are also useful analogues for existing contaminant plumes. The principal U ore source mineral in these deposits is reduced U oxide (UO_2) which is subsequently altered during fluid-mineral interaction under oxidizing conditions to uranyl- oxides, phosphates, and silicates. Most of the U migration, away from the concentrated uranium ore, occurred after oxidation through groundwater mass transport. Subsequently, U became incorporated into surrounding rock through sorption and precipitation of

weathering phases. This sequence of oxidation, transport, and deposition is expected to be a reasonably good analogue for transport of U present in man-made radioactive waste repositories, mine mill tailings, and ore processing plants. Since U-contaminated soils are chemically analogous to U ore bodies, the extent of the contaminated "halo" around the latter might also provide useful information about plume migration.

Roughly 40% of U ore bodies occur as epigenetic sedimentary deposits such as those found in the Wyoming Tertiary basins and in coastal plain systems, e.g. the South Texas Uranium province ⁴⁹. Other U occurrences are associated with alkaline volcanic and granitic igneous provinces, e.g. the Sanerliu granite-hosted U deposit in southern China ⁵⁰, the Peña Blanca district rhyolite tuff in the state of Chihuahua, Mexico ⁵¹⁻⁵⁷, the Palmottu U deposit in Finland ^{58; 59}, the Maqarin U hydrothermal deposit in northern Jordan ^{60; 61}, the Valles natural analogue at Valles Caldera in New Mexico ⁶², the Konigstein U mine in Saxony, Germany ^{63; 64}, and the Poços de Caldas alkaline complex in the state of Minas des Gerais, Brazil e.g. ^{65; 66}. We focus here on those ore bodies that have been well studied and monitored for long periods of time and whose groundwater chemistry provide an objective plume length analysis. These include the Alligator Rivers region (the Koongarra ore deposit) in Australia, the natural uranium reactors/mines of Oklo in the Republic of Gabon in equatorial Africa, the Morro do Ferro and Osamu Utsumi U mines at Poços de Caldas alkaline complex in the state of Minas Gerais, Brazil, and Cigar Lake in Saskatchewan, Canada.

3.2.1 Koongarra

The Koongarra ore deposit is located in northern Australia and is part of the Alligator Rivers Uranium Field. The host rock for the Koongarra ore deposit is a quartz chlorite schist and the zone of primary mineralization is about 100 meters deep⁶⁷. The primary mineralized region is composed of uraninite and is situated between a graphite rich region bounding the Kombolgie Sandstone and the Koongarra reverse fault⁶⁷. According to Yanase et al.³², the groundwater flowpath of the groundwater begins at the surface and percolates through the porous Kombolgie Sandstone and the Koongarra reverse fault that separates the sandstone from the ore body. Fluids move deeper into the primary mineralized zone through the fault, but shift away from it to a horizontal flow regime creating a weathered zone of secondary U mineralization. The close proximity of the primary mineralized region and the graphitic schist layer suggest that reducing reactions involving this strata were responsible for the precipitation of uraninite from fluids percolating through and near the fault. A very similar scenario caused the formation of uraninite in Oklo natural reactors where the mineralized region is also bounded by a fault^{68; 69}. The primary ore body at Koongarra has been affected by weathering forming a secondary ore alteration zone extending the limits of the U migration^{67; 70}. The presence of this weathered zone is also a good indication of groundwater flow through the ore body. Weathering processes such as precipitation and concomitant transformation of secondary minerals, e.g., chlorite → vermiculite + Fe-minerals → kaolinite + Fe-minerals, coupled with sorption by these phases, retards U transport and migration downstream from the main ore deposit. The

precipitation of saléeite ($\text{Mg}(\text{UO}_2)_2(\text{PO}_4) \cdot 10\text{H}_2\text{O}$) and the irreversible sorption of U onto Fe-oxides produced by chlorite weathering likely controlled U migration and/or dispersion at Koongarra⁷¹. According to Murakami et al.⁷¹, saléeite, along with metatorbernite, precipitated between apatite and sklodowskite grains at the expense of these two latter phases. These authors suggested on the basis of micro-textural observations that localized interfacial precipitation is the most plausible way of forming these phases in undersaturated fluid conditions for the Koongarra groundwater. Moreover, Sato et al.⁷² reported significant U scavenging associated with Fe oxides (nodules) having concentrations many times larger than in the groundwater.

The Koongarra ore is exposed to monsoonal climate, but seasonal sampling suggests that a sharp change in water recharge has little effect on groundwater bulk chemistry. In general, the groundwater of this deposit is dilute, with total dissolved solids in the range of 200 mg/L³². The deeper groundwater is enriched in CO_2 indicating that U in solution, within the weathered zone, is mobilized as uranyl-carbonate complexes^{32; 73}. The primary mineralized zone is traversed by the Koongarra fault which may well serve as a conduit for surface water to permeate into the deeper formations and dilute the groundwater^{32; 73}. U concentrations in the groundwater wane away from the primary mineralized zone for about 200 meters until background concentrations are observed. Groundwater flow modeling⁷⁴ suggests that water moving away from the porous sandstone and the fault are primarily transmitted through a very heterogeneous fractured

media in the weathered zone, making the characterization of flow velocities at the scale of the ore very difficult. Therefore, simplistic 1-D solute transport simulations using single velocities cannot be satisfactorily applied for the description of U migration in this area⁷⁴.

3.2.2 Oklo

The Oklo natural reactors in The Republic of Gabon, equatorial Africa, consist of two natural reactors, Bangombe and Okelobondo, which represent an extreme case where large amounts of transuranium series elements were produced over long periods of time, after the reactor went critical ~2 billion years ago⁶⁸. Furthermore, the tropical climate assures abundant recharge. The Oklo reactors have been extensively studied due to potential similarities to proposed long-term high-level nuclear waste repositories^{68; 75}. The reactor cores are composed of uraninite surrounded by Precambrian pelitic sandstones and unmetamorphosed volcanic rocks⁶⁸. U Mineralization of the Oklo cores occurs within a sedimentary layer of fluvial sandstones and conglomerates associated with a carbonate-rich/argillaceous matrix⁶⁸. An interesting similarity between Oklo and Koongarra ore deposits is that both are bounded by a major fault which serves as a natural barrier, and as a conduit, limiting U migration and/or dispersion through groundwater percolation near the fault and along the weathered zone. In both cases, the fault served as a conduit for fluid recharge and as the initial point for the oxidation zone to originate. The estimated maximum axial plume length for Bangombe and Okelobondo are 0.25 and ~1.3 km, respectively. Recent reactive transport models on Bagombe and Okelobondo by Ayora et al.⁶⁹ and Salas et

al.⁷⁶, respectively, suggest that local geology and mineralization in the host strata controls fluid chemistry. At Okelobondo, $\text{Fe}^{2+}/\text{Fe}(\text{OH})_3$ and Mn-minerals control the reaction paths for two types of site groundwater⁷⁶. At Bangombe the pore water chemistry appears to be controlled by the $\text{Fe}^{2+}/\text{Fe}(\text{OH})_3$ equilibria⁶⁹.

3.2.3 Poços de Caldas

The Poços de Caldas ore deposit has been the subject of a number of studies that have characterized its geology, geochemistry, hydrology, and hydrochemistry in order to understand ore formation and U transport⁶⁵. Poços de Caldas is a round-shaped Mesozoic igneous alkaline complex of ~33 km in diameter comprising suites of alkaline volcanic and intrusive rocks mainly phonolites and syenites⁷⁷. Most of the radioactive and REE mineralization is associated with heavy hydrothermal activity and the formation of volcanic breccias that host these deposits^{77; 78}. Subsequent episodes of ultramafic magmatism and lamprophyric dyke formation, followed by intense weathering, resulted in the formation of supergene zones producing redox fronts that mobilized and enriched U to lateritic levels⁷⁷. There are two sites for U mineralization in this area: Osamu Utsumi and Morro do Ferro mines. U mineralization at Osamu Utsumi is the result of supergene weathering beneath a lateritic soil extending to an approximate depth between 80 and 140 meters⁷⁹. Oxidized and reduced zones along the redox front are distinguished by sharp changes in color. Most of the minerals hosting the bulk of U in this mine are uraninite, pitchblende, and brannerite. Precipitation of these U-bearing minerals is associated with oxidation of pyrite and secondary precipitation of Fe oxy-hydroxides

throughout the reaction front ⁷⁹. A reactive transport model by Lichtner and Waber ⁸⁰ for the Osamu Utsumi mine successfully predicts the migration of the redox front and the resulting phases precipitated along the reaction path. In this case, pyrite oxidation causes the fluid to be reduced, resulting in precipitation of uraninite in the redox front ^{80; 81}. Morro do Ferro is richer in Th and depleted in U relative to Osamu Utsumi ²⁹. Groundwater sampled close to the surface of this deposit is oxidized as expected, but more reduced in deeper samples in the boreholes. The high concentration of Th in the groundwater is thought to be associated to colloidal matter because of its low solubility and its high partition with colloidal matter at the Osamu Utsumi site ²⁹. The Morro do Ferro plume was measured using a vertical profile along a transect comprising a limited set of sampling boreholes. The plume length is ~0.15 km.

3.2.4 Cigar Lake

The Cigar Lake unconformity-type U ore deposit in northern Saskatchewan, Canada ⁸²⁻⁸⁴ is hydrothermal in origin, was formed ~1.3 Ma ago, and is confined to an altered

sandstone ~430 meters below the surface ^{84; 85}. The primary U minerals are uraninite and pitchblende. Waters in contact with the ore originate from an overlying permeable sandstone aquifer. Because the Cigar Lake ore deposit is not exposed at the surface, U release is extremely slow. Weathering, formation of a surrounding clay-rich matrix, and capping by an impermeable quartz-cemented zone cause the groundwater in contact with the ore body to be highly reduced. The system has consequently been assumed to be closed with respect to U. U transport in this ore body has been modeled by Liu et al. ⁸⁶ using a near-field release model assuming molecular diffusion perpendicular to the clay zone and advective groundwater flow parallel to the clay zone. Bruno et al. ⁸⁷ modeled the fluid chemical evolution along different flowpaths, using a simple kinetic mass transfer calculation entailing oxidative uraninite dissolution, assuming long residence times. Indeed, Liu et al. ⁸⁶ model predicts very low U concentrations as observed in the field. The confinement of U and other radionuclides in the clay zone arrest their migration, therefore producing plumes in the porous overlying sandstone that are too narrow to be detected ⁸⁶.

4.0 Plume Analysis

The maximum surface extension of both artificial and natural plumes or **maximum plume axial length** is used as the index criteria to assess plume behavior. Note though that the concept of maximum plume axial length, as applied here, is by necessity operational because of the random, limited, and in most cases, subjectively biased well sampling or monitoring used by different workers at different sites. Moreover, the highly distorted morphology of groundwater plumes, the presence of daughter plumes,

and the presence of background levels of U causes more uncertainty as to the real extent of the 2-D surface coverage of these plumes. To establish an objective basis of comparison, visual inspection of plume contour maps and U concentration data in sample wells were used jointly to establish the maximum plume axial lengths. Specifically, the maximum plume axial length is defined here as *the maximum distance between two points encompassing the farthest boundaries of the plume as*

constrained and/or permitted by the sampling well network in a particular site where measurable U concentrations in the range of 10-20 ppb have been obtained.

Previous workers⁸⁸ conversely, have considered the farthest distance between the source (or highest contaminant concentration) and the plume boundary. This approach is useful in intuitively assessing the limits for potential spreading of a plume within a given area if the data set is sufficiently large and reliable. For most U plumes, temporal and spatial limitations in well sampling and the generation of daughter plumes through ongoing remediation activities, or natural recharge, makes identification of the source within a waste site a very difficult task for a given well monitoring network. This could lead to underestimates of plume length that can only be overcome by a large and fairly dependable data set. Given the limited amount of useful data, the irregular spatial distribution of monitoring wells, and the scarce number of the latter at each site, we found the maximum axial length provides (in the extreme case of U mobilization) a reasonably good estimate of the 2-D contaminant surface coverage. Note there is a general lack of temporal data for periods longer than 5 years for most sites. Many of the sites possessed a very large (hundreds of meters across) and disperse source term. The width of the source term is implicitly counted in the maximum plume axial length measurement. In other words, if the actual plume advance were modeled as emanating from a point source, the calculated plume lengths would be a great deal less. In some cases, particularly for the large scale natural analogues (e.g., Poços de Caldas and Oklo), the plume lengths were estimated based on a vertical profile using a linear monitoring well transect. To illustrate the manner in which plumes were measured, figures 8

through 11 show a number of the Title I UMTRA plumes (Wyoming, Colorado, and Arizona) and the labeled plume lengths.

Many of the UMTRA sites are located within 2 or 3 km of rivers. There are a few cases where groundwater plumes were truncated by discharge into rivers, e.g. Figure 8 - Riverton, Wyoming, as might be expected where rivers are fed by groundwater. In arid regions though, rivers often lose water to adjacent aquifers and many of the plumes we observed spread parallel to, or away from, nearby rivers, suggesting that measured plume lengths reflect groundwater transport. Table 5 shows all the U plumes considered in this study along with estimated maximum axial plume lengths. The frequency distribution of maximum axial plume lengths for all sites listed in table 5 is shown in figure 12, and suggest that the maximum observed distance of migration is a little more than 2 kilometers. Note again that this distance is the maximum observed spread of the 10-20 ppb U plume contour, and that it includes both upgradient and downgradient limbs of the plume. This means, the downgradient (maximum) reach of plumes from the source is substantially less than 2 km. If we calculate plume length using contours of 44 ppb U, the MCL, most of the plumes (if not all of them) would have an axial length of approximately 0.5 km or less. An anomalous long outlier is the plume associated with the Konigstein mine^{63; 64}, located 25 km southeast of the city of Dresden, Germany and the UMTRA site Falls City, Texas. *In situ* leaching (ISL) was conducted in the Konigstein mine using periodic inputs of sulfuric acid (H₂SO₄) that mixed and diluted with local groundwater needing further additions of the acid to continue the leaching process⁶³.

Table 5. Summary of estimated maximum axial plume lengths and their site characteristics. The listed UMTRA sites are the only ones for which plume length data can be extracted.

Site	Type	Max. Axial Plume Length (km)	Min. Axial Plume Length* (km)	Sampled Depth (m)	Sources	Comments
Canonsburg, PA	UMTRA (Title I)	0.3-0.37	-	2-8	89	Groundwater table can be found at shallow depths in the fill. Humid continental climate.
Crow Butte Uranium Mine Unit 1, NB	In situ leaching	0.63	0.07	-	90	Pre-operational/baseline maximum plume length measured to ~ 20 ppb. Post-operational ISL mining caused [U] to be orders of magnitude larger in monitoring groundwater wells.
Falls City, TX	UMTRA (Title I)	4.95	3.94	-	39	Plume analysis comprises tailings pile areas 1, 2, 3, 4, 5, 6, and 7. Largest UMTRA plume.
Fernald Processing Site (OH)	UMTRA (Title I)	1.3	0.61-0.78	-	91	Private well monitoring locations (1992-1996)
Grand Junction, CO	UMTRA (Title I)	2.5	0.47 - 0.6	-	92	Bulk groundwater composition is SO ₄ ⁻ rich and relatively HCO ₃ ⁻ poor. Close to saturation with respect to calcite.
Gunnison, CO	UMTRA (Title I)	2	0.4	50-150	93 ^a	Lindgren ^a reported a plume length value of 1.5 km interpolated distance to [U]=40 ppb.
Hanford (WA) 300 Area process trench	UMTRA (Title I)	0.79	0.52	50-150	94	Plume bounded by the Columbia River
Kennecott Uranium Facility (WY)	UMTRA - Mine (Title II)	0.69	0.26	~15	95	Highly irregular plume shape. Maximum plume length measured to ~8 pCi/L
Konigstein Mine, Germany	In situ leaching	3.6-4.0	~1.5	~15-350	63; 64	In situ leaching (ISL) of U with sulfuric acid (H ₂ SO ₄). Among longest plume measured.

^a E. R. Lindgren (unpublished report)

Table 5 (cont.). Summary of estimated maximum axial plume lengths and their site characteristics.

Site	Type	Max. Axial Plume Length (km)	Min. Axial Plume Length* (km)	Sampled Depth (m)	Sources	Comments
LLNL-plume 1 pit 4-5, CA	Explosive Activity	0.43	0.08	-	96	Sampled 2 nd quarter 1994; plume length measured to the [²³⁴ U + ²³⁸ U]=10 pCi/L (~30 ppb) contour.
Maybell, CO	UMTRA (Title I)	0.4	0.15	40-50	97	U/TDS* ratio indicates that soluble salts move further than U beyond the mill tailing limits.
Monticello Millsite, CO	UMTRA (Title I)	2.2-2.4	0.42	-	98	Plume length distance measured to a ²³⁴ U + ²³⁸ U concentration level of ~18 pCi/L (~54 ppb).
Monument Valley, AZ	UMTRA (Title I)	1.4	1.1	17-47	36	Plume length for the [U] _T >44ppb region (deep De Chelly aquifer) is ~0.7 km. Max. plume length determined for the alluvial aquifer.
Naturita, CO	UMTRA (Title I)	0.7	0.2	3-76	99	Plume length may be larger than estimated value. Groundwater sampling restricted to the shallow river alluvium.
New Rifle, CO	UMTRA (Title I)	1.6	0.6	30-95	100	U/TDS ratio is similar in all sampling wells suggesting that U salts and U migrate at the same rate.
Rio Algom, Moab - Lisbon Facility, UT	UMTRA - Mine (Title II)	2.52	1.71	13-45	101	Maximum plume length measured to 10 -20 pCi/L natural U sampling well - among the largest Title II plumes
Riverton, WY	UMTRA (Title I)	1.7	1.2	7-8	34; 35; 102	Lindgren ^a reported a plume length value of 0.9 km interpolated distance to the 44 ppb [U] point
Slick Rock (NC), CO	UMTRA (Title I)	0.24	0.12	20-50	103	Sampling restricted to tailings pile. Plume may be bigger than estimated. Monitoring wells at plume boundary show [U] ≈ 900-1000 ppb.

Table 5 (cont.). Summary of estimated maximum axial plume lengths and their site characteristics.

Site	Type	Max. Axial Plume Length (km)	Min. Axial Plume Length (km)	Sampled Depth (m)	Sources	Comments
Slick Rock (UC), CO	UMTRA (Title I)	0.5	0.2	20-50	103	Site is bounded by a topographic high and a river.
Sohio Western L-Bar, NM	UMTRA - Mine (Title II)	1.34	0.96	-	104	Maximum and minimum plume lengths are approximate - few wells available for measuring natural U sampling
Split Rock (WY) Northwest Valley	UMTRA (Title II)	2.63	0.75	0-30	105	Mill tailings still remain in place. Long plume length for an UMTRA site.
Split Rock, (WY) Southwest Valley	UMTRA (Title II)	2.51	0.86	0-30	105	Mill tailings still remain in place. Long plume length for an UMTRA site.
Split Rock (WY) Between Northwest and Southwest Valley	UMTRA (Title II)	2	-	0-30	105	Mill tailings still remain in place. Plume length measured between two valleys containing the mill processing plants and tailings.
Tuba City, AZ	UMTRA (Title I)	1.12	0.5	15-18	12; 37; 106; 107	Maximum plume length measured to [U] \approx 40 ppb ¹⁰⁷ .
Weldon Springs Site, Missouri (WSOW)	UMTRA (Title I)	0.6	-	-	108	Plume length value is very approximate. [U] well data is very heterogeneous. Multiple plumes observed. Very localized plume lengths with [U] > 15 pCi/L (~45 ppb) are only reported.
Weldon Springs Site, Missouri (WSCP)	UMTRA (Title I)	1.1	-	-	108	Multiple plumes observed. Same explanation as above.

Table 5 (cont.). Summary of estimated maximum axial plume lengths and their site characteristics.

Site	Type	Max. Axial Plume Length (km)	Min. Axial Plume Length (km)	Sampled Depth (m)	Sources	Comments
Koongarra ore deposit, Alligator River Uranium Field, Australia	Natural Analogue	0.48-0.5	0.38	13-25	32; 67; 73	Presence of a weathered zone. Uranyl-carbonate complexes predominant due to high HCO_3^- concentration in deeper groundwater.
Bangombe, Oklo natural reactors, Gabon	Natural Analogue	0.25	-	25-500	68; 69; 75; 109; 110	Presence of a weathered zone. Groundwater chemistry controlled by the $\text{Fe}^{2+}/\text{Fe}(\text{OH})_3$ equilibria. Fluids are not enriched in CO_2 .
Okelobondo, Oklo natural reactors, Gabon	Natural Analogue	0.9-1.0	-	6-100	68; 69; 75; 76; 109; 110	Presence of a weathered zone. Groundwater chemistry controlled by the $\text{Fe}^{2+}/\text{Fe}(\text{OH})_3$ (reduced) and $\text{Mn}^{2+}/\text{MnOOH}$ (oxidized) equilibria. The latter is richer in CO_2 .
Osama Utsumi, Poços de Caldas, Brazil	Natural Analogue	0.5-0.6	-	0-125	78; 111-113	Presence of a weathered zone. Pyrite oxidation induces reduction of fluids and subsequent UO_2 precipitation in the redox front.
Morro do Ferro, Poços de Caldas, Brazil	Natural Analogue	0.15	-	0-85	78; 111; 112; 114	Presence of a weathered zone. Th rich deposit. The presence in groundwater is probably associated to colloids. Ore zone is very close to the surface.
Cigar Lake ore deposit, Canada	Natural Analogue	0.4	-	0-500	84; 87	Deep (~430 m) and concealed unconformity type U deposit. Capped by an impermeable quartz barrier. Considered a closed system.

* Minimum axial lengths are measured perpendicular to maximum axial length.

Leached contaminants have therefore been spread further than they would have otherwise. ISL has been used in many U mines in the United States, e.g., Falls City, Texas, and worldwide ¹¹⁵ and is being currently considered as a cheaper option for future U mining by various countries ¹¹⁶; ¹¹⁷. Falls City (Texas) mill site show the largest observed plume for a Title I UMTRA site. It also has a fairly recent history of secondary solution mining operations between 1978 to 1982 which may be attributed to its spatial extent of contamination¹¹⁸. Some examples of previous and presently planned use of ISL solution mining are Germany (Konigstein), Czech Republic (Stráz mine in north Bohemia), Bulgaria, Ukraine, Russia, Kazakhstan, Uzbekistan, China, United States, and Australia ¹¹⁶; ¹¹⁷; ¹¹⁹. The

Konigstein mine is probably the best studied example of intensive use of ISL in U mining and its consequences on aquifer and groundwater contamination ¹¹⁵. A recent example of ISL solution mining by injection of an oxidant and a carbonate-rich solution in the USA is the Crow Butte U mine unit 1 in Nebraska ⁹⁰. The groundwater chemical patterns of post-operational ISL activities show a plausible maximum plume length increase that may exceed ~3-4 times that of pre-operational/baseline standards (baseline max. plume length ≈ 0.62 km)⁹⁰. Even the subsequent restoration/stabilization activity of groundwater quality at this site shows U concentrations that exceed MCL limits further beyond the monitoring well network.

5.0 Discussion and Conclusions

The hydrologic conductivities, Kds, and original contaminant source masses for the various sites probably vary by orders of magnitude (see e.g. table 4). Nevertheless, actual plume trajectories seem to cluster, and suggest that the combined effects of dispersion and chemical reaction are sufficient to arrest most uranium plumes before they move more than roughly a kilometer from their source. The natural life cycle of a uranium plume appears to involve an initial movement away from a source region that takes place within a few years and does not exceed 2 kilometers, followed by a geologically long period of immobile quiescence. Natural plumes from ores that have been weathered and subjected to periodic meteoric inputs for long periods of time do not migrate appreciably beyond their known natural barriers, even during mining. Similarly, the UMTRA sites do not show a

significant dispersion of contaminants beyond the limits of the contaminated area, even though these are not as deeply buried and are in more porous strata than those found in the natural analogues and ore U mining sites. The plume length and the U concentration in monitoring wells remain relatively constant, or change insignificantly, for periods of time approaching 15 years in many cases^a. It appears that sorption, dilution, and precipitation are sufficiently effective sinks to limit short-term (years to decades) the advance of artificial U plumes. In long-term situations (thousands to millions of years), weathering processes and secondary precipitation of oxidized uranyl phases appears to limit advance of natural plumes. This picture of U plume behavior has a number of implications for activities

^a E. R. Lindgren (unpublished report)

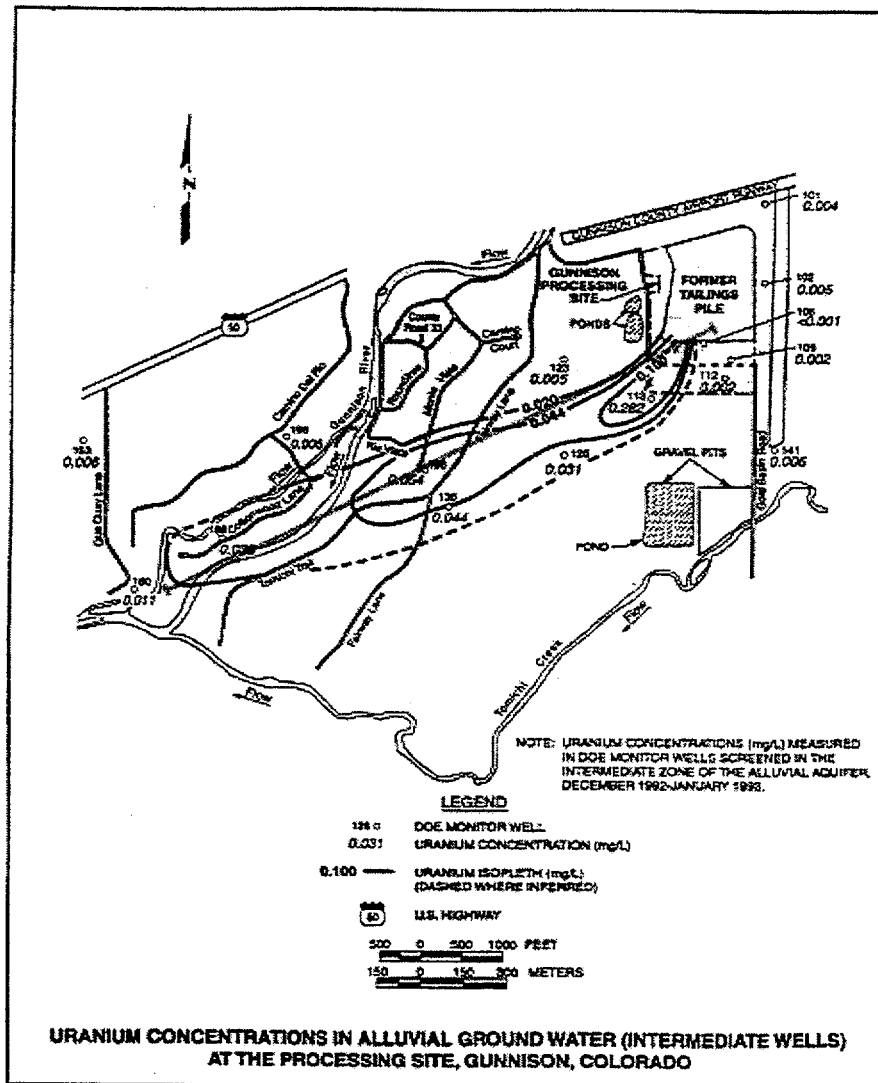


Figure 9. Gunnison (Colorado) plume ⁹³. The thick light-gray line is the estimated maximum axial plume length as in figure (see figure 8).

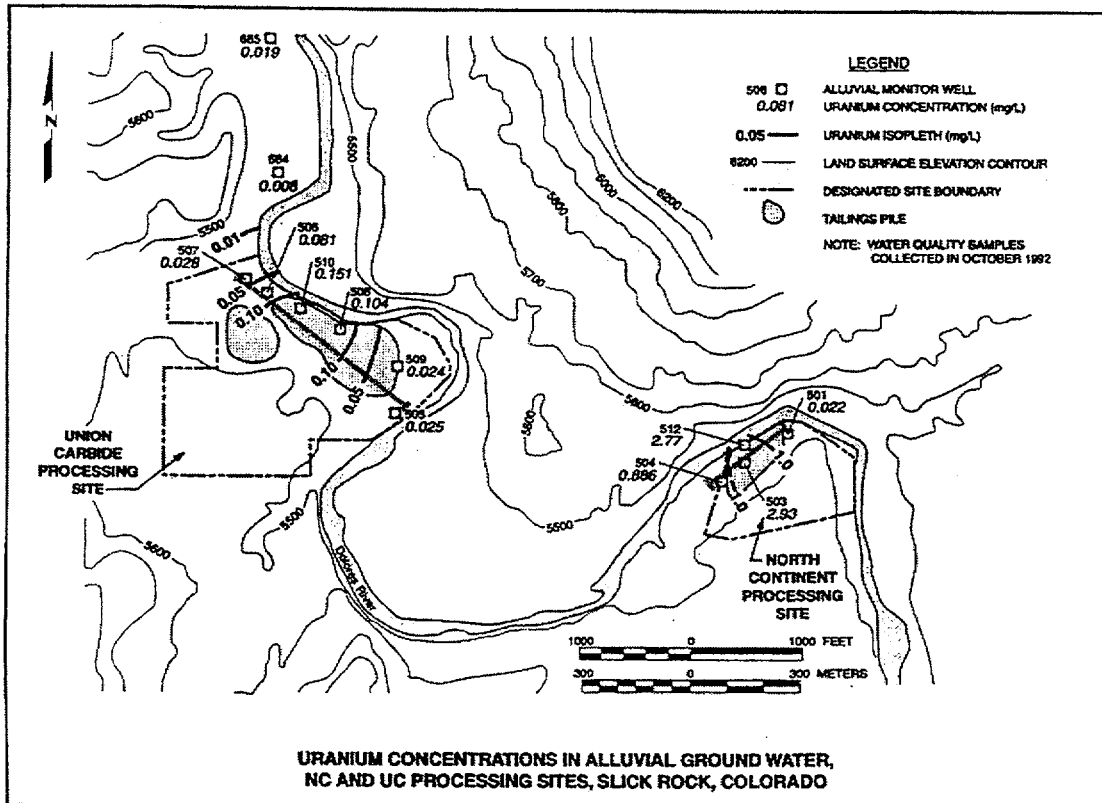


Figure 11. Slick Rock (Colorado) plume¹⁰³. The thick light-gray line is the estimated maximum axial plume length as in figure (see figure 8).

associated with remediation and monitoring. To begin with, these results set very clear limits on what reaction-transport model outputs should look like. Although input functions and sub-models can vary widely because "every site is different", output predictions of U movement in subsurface environments do not exceed roughly 2 kilometers and represent natural plume behavior, unless special chemistries, and possibly hydrologies are involved, e.g., active sulfuric acid leaching or perhaps, demonstrable fast path fractures. Long-term monitoring wells placed ahead of plumes assuming steady long-term plume advance may never detect their targets. *In situ* remediation that relies on mobilization by chelating ligands may ultimately result in anomalous long plume movement. Source term removal alone seems to limit plume advance. This picture assumes no change in the geochemical state of the attenuated uranium near the source, such as the introduction of chelating agents and/or a deleterious shift in redox conditions.

Only two of the few plumes examined for the Title II sites Rio Algom (Utah) and Split Rock (Wyoming) exceed plume lengths >2.5 km, which represent the longest estimated distances in this study. It is expected that Title II sites produce longer plumes because the source term is still in place; however the rest of these fall well within the plume length distribution range (i.e., <2km) obtained for both UMTRA Title I and Title II sites.

Lastly, it should be noted that geochemical factors favor uranium transport to be greater than the transport of many other cationic metals and radionuclides such as Pb, Cd, ⁹⁰Sr, ¹³⁷Cs because U is a relatively weak sorber and/or because the soil minerals it forms are relatively soluble. Moreover, the fraction of U that sorbs irreversibly in soils is relatively small. For these reasons, we should expect a plume advance for most other cationic metals and radionuclides to be substantially less than the 2 kilometers observed for U.

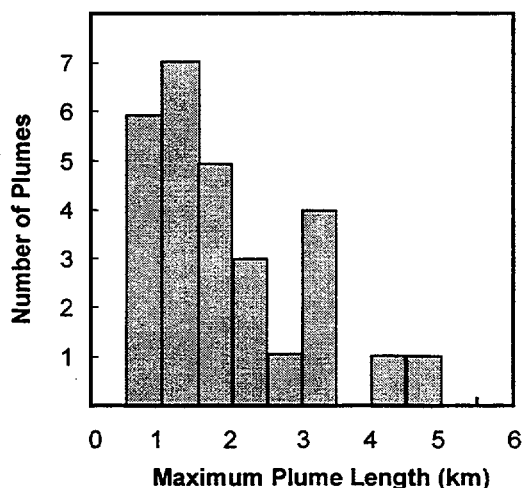


Figure 12. Histogram of maximum plume lengths for all considered uranium plumes in this study (n=28; see text).

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11. ABSTRACT *(200 words or less)*

Groundwater plumes containing dissolved uranium at levels above natural background exist adjacent to uranium ore bodies at uranium mines, milling locations and at a number of explosive test facilities. Public health concerns require that some assessment of the potential for further plume movement in the future be made. Reaction-transport models, which might conceivably be used to predict plume movement, require extensive data inputs that are often uncertain. Many of the site-specific inputs are physical parameters that can vary spatially and with time. Limitations in data availability and accuracy mean that reaction-transport predictions can rarely provide more than order-of-magnitude bounding estimates of contaminant movement in the subsurface. A more direct means for establishing the limits of contaminant transport is to examine actual plumes to determine if collectively, they spread and attenuate in a reasonably consistent and characteristic fashion. Therefore, a number of U plumes from ore bodies and contaminated sites were critically examined to identify characteristics of U plume movement. The magnitude of the original contaminant source, the geologic setting, and the hydrologic regime were rarely similar from site to site. Plumes also spanned a vast range of ages and no complete set of time-series plume analyses exist for a particular site. Despite the accumulated uncertainties and variabilities, the plume data set gave a clear and reasonably consistent picture of U plume behavior. Specifically, uranium plumes: (1) Appear to reach steady-state, that is, they quit spreading rapidly (within a few years). (2) Exceed roughly 2 km in length only in special cases e.g. where in situ leaching has been carried out. The majority is much smaller. (3) Exhibit very similar U chemistry between sites. This implies analogous contaminant attenuation mechanisms despite their location.

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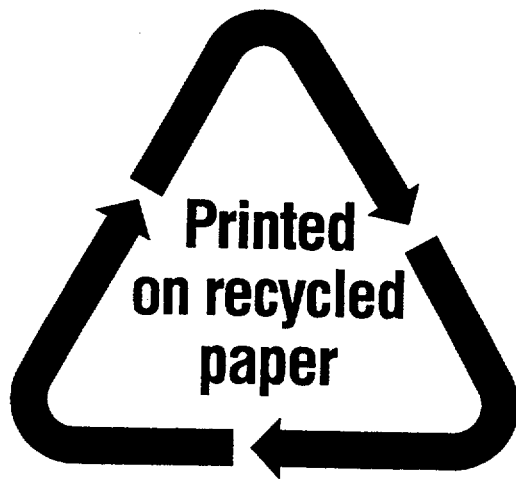
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