RECOMMENDED SITE-SPECIFIC SORPTION COEFFICIENTS FOR REVIEWING NON-HIGH-LEVEL WASTE DETERMINATIONS AT THE SAVANNAH RIVER SITE AND IDAHO NATIONAL LABORATORY

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EXECUTIVE SUMMARY

To satisfy non-high-level radioactive waste determinations for facilities at the Savannah River Site (SRS) and the Idaho National Laboratory (INL) under Section 3116 of the Ronald W. Reagan National Defense Authorization Act, the U.S. Department of Energy (DOE) commonly performs a deterministic performance assessment that includes evaluating potential radionuclide exposure via groundwater transport. In its consultative role in the waste determination process, the U.S. Nuclear Regulatory Commission reviews the performance assessments, including deterministic radionuclide geosphere transport simulations that use sorption coefficients, or K_ds. To aid in these reviews, this report documents a sorption coefficient database tailored to site-specific subsurface conditions for risk-significant radioelements at SRS (carbon, nickel, selenium, strontium, niobium, technetium, tin, iodine, cesium, lead, uranium, neptunium, plutonium, americium, and curium) and INL (carbon, strontium, technetium, iodine, cesium, lead, uranium, neptunium, plutonium, americium, and curium) and recommends K_d values appropriate for the respective hydrogeochemical environments. The recommended K_d values are based on the sorption behavior of each radioelement, available data on site hydrogeochemical and mineralogical conditions, and reported site-specific K_d values. When site-specific data were not available for a particular radioelement, K_d data on analogous systems and generic data from the open literature were evaluated to provide a technical basis for the K_d recommendation. The recommended values tended toward the lower, more conservative end of data ranges because of the uncertainties not only in the appropriate choice of sorption coefficients, but also in the simplified K_d-based groundwater transport model itself. The selection of low values is intended to ensure that groundwater pathway models will be unlikely to underestimate potential dose. Values were assigned a low or high level of uncertainty based on the availability of site-specific information. These recommendations should be used with care in light of new information that becomes available on site-specific sorption coefficients or hydrogeochemical conditions. This report is not to be construed as providing the technical basis for parameters DOE uses in its performance assessments. The results of this data review can be used, in a risk-informed manner, to identify potential areas of focused laboratory studies.

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QUALITY OF DATA, ANALYSES, AND CODE DEVELOPMENT

DATA: No original data were generated from the analyses presented in this report. Sources of data cited in this report should be consulted for determining the level of quality of those data.

ANALYSES AND CODE: No codes were used in the analyses contained in this report.

ABBREVIATIONS/ACRONYMS

CEC cation exchange capacity COC contaminant of concern

DEXOU D-Area expanded operable unit

DO dissolved oxygen

DOE U.S. Department of Energy

EPA U.S. Environmental Protection Agency

GSA General Separations Area

NRC U.S. Nuclear Regulatory Commission

INL Idaho National Laboratory

INTEC Idaho Nuclear Technology and Engineering Center

SDA Subsurface Disposal Area SRPA Snake River Plain Aquifer SRS Savannah River Site TDS total dissolved solid

1 INTRODUCTION

One of the requirements for satisfying non-high-level waste determinations for planned and existing facilities at the Savannah River Site (SRS) and Idaho National Laboratory (INL) under Section 3116 of the Ronald W. Reagan National Defense Authorization Act for Fiscal Year 2005 is that the U.S. Department of Energy (DOE) dispose of radioactive waste resulting from reprocessing in accordance with the performance objectives set out in 10 CFR Part 61, Subpart C. In satisfying one of these objectives—Protection of the General Population from Releases of Radioactivity (10 CFR 61.41)—DOE commonly performs a performance assessment that includes evaluating potential exposure via a groundwater pathway. Such calculations typically include groundwater transport calculations that involve the use of sorption coefficients (K_d) that assume equilibrium partitioning of a dissolved contaminant radionuclide between the groundwater solution and the host matrix. While the assumptions underlying the use of constant K_ds for given hydrostratigraphic units (e.g., invariable chemical conditions, relatively rapid sorption and desorption kinetics, linear concentration dependence) are not always satisfied, using K_ds to model the transport of dilute solutions of radionuclides in a homogeneous hydrochemical and geochemical environment can provide useful results. In performance assessments supporting waste determinations, DOE uses both site-specific and generic values in radionuclide transport models. This evaluation of site-specific data is not intended as an exhaustive treatise on sorption and contaminant transport. More detailed discussions of sorption mechanisms may be obtained from sources such as Hochella and White (1990). In addition, this review does not explicitly address in detail the effects of flow on sorption; hydraulic and physical properties of the transport medium are discussed only to the extent necessary for selecting appropriate K_d values.

1.1 Purpose

In its consultative role in the Section 3116 waste determination process, the U.S. Nuclear Regulatory Commission (NRC) reviews any relevant performance assessments and provides comments as appropriate. These reviews include evaluating whether sorption coefficients DOE uses for radionuclide transport in the geosphere reasonably reflect expected local conditions. In other words, NRC staff is interested in whether site-specific data were used, whether hydrochemical conditions along the groundwater pathway were considered in choosing $K_{\rm d}s$, and whether any uses of non-site-specific data were warranted and appropriate. For NRC staff reviewing DOE non-high-level waste determinations, the purposes of this report are (i) to document the sorption coefficient database tailored to site-specific subsurface conditions for risk-significant radionuclides and relevant decay chain members; (ii) to recommend sorption coefficient values, or ranges of values, for radionuclide species appropriate for those specific hydrochemical and geochemical environments found at SRS and INL; and (iii) to suggest appropriate ways to evaluate DOE values in light of the compiled information.

While this report does recommend appropriate sorption coefficients for transport based on careful analysis, these values should not be accepted without acknowledging any new data, considerations, or other information provided in DOE reports. For example, new understanding may emerge about the long-term evolution of SRS groundwater hydrochemistry. The inclusion of a recommended value in this report does not constitute a technical basis for DOE use of the value. Likewise, a recommendation in this report does not preclude the acceptability of different values DOE uses. DOE will always be expected to provide the technical basis, independent of

this report, for parameters employed in its models. Recommendations in this report tend to bound the lower range of measurements. If independent NRC staff analyses of K_ds were to be conducted, uncertainties reflected in the data reported here should be considered.

1.2 Use of Sorption Coefficients in Performance Assessment

The sorption coefficient commonly has units of mL/g1 or L/kg and represents the ratio of the amount of contaminant on the solid per unit mass to the amount of the contaminant in the fluid per unit volume. The effects of sorption are to slow transport relative to unretarded species and accentuate the attenuating effects of dispersion. Therefore, highly sorbing radionuclides (i.e., those with high K_d values) initially tend to be less important contributors to groundwater dose in the absence of colloidal effects. In relatively straightforward radionuclide transport models such as those DOE uses in non-high-level waste determinations, this ratio is typically assumed to be constant for a given stratum along the hydrologic transport pathway. Unit-specific values are intended to reflect the dependence of K_ds on mineralogical and surface area properties of the rocks in that unit. Sorption coefficients also depend on hydrogeochemical factors and, as such, should reflect the chemical characteristics of the transporting water. Dependence of K_as on such factors as pH and oxidation-reduction potential is well known and is related to the effects of these factors on aqueous speciation. For example, actinides such as neptunium and uranium show strong pH variation in K_d, with sharply defined peaks at moderate pH values. Aqueous complexation with organic and inorganic species (e.g., carbonate and sulfate) also affects surface-solution partitioning. The presence of colloids in groundwater can lead to higher mobility of strongly sorbing radioelements such as plutonium—an effect that is not captured in typical K_d approaches. The nature of the host rock may also affect transport in ways unaccounted for in simple porous-medium models (e.g., fractures may provide fast transport pathways).

Because these lithologic and hydrogeochemical factors are not always well constrained for a given transport model, it is often necessary to make conservative selections of K_d values that are at the low end of reasonable ranges. In this way, the model is less likely to overpredict the favorable effects of sorption. Such an approach is useful in the case of spatial and temporal variation. For example, the General Separations Area at SRS (see Chapter 2) was subject to extensive acid groundwater contamination during the 20^{th} century, imparting a range of groundwater pH from 3.4 to 7.1, with estimated "background" values of 4.8 to 6.8 (Kaplan, et al., 1994; Kaplan, 2004; Dai, et al., 2002; Serkiz and Johnson, 1994; Vyas, et al., 2001). Because actinide sorption coefficients can vary over one or two orders of magnitude over this pH range, it is clear that potential variation in pH must be considered in assigning model parameters.

1.3 Approach to Data Review

In this review, the correspondence between field conditions (i.e., subsurface mineralogy and aqueous chemistry) and the data used to estimate K_d values for each radionuclide contaminant of concern (COC) at SRS and INL were considered. In many cases, site-specific K_d data for the COCs at SRS and INL are available. When site-specific K_d data were not available, data from the open literature on comparable systems to SRS and INL were consulted and analyzed. The result of this data review is a recommended K_d value for each COC at SRS and INL. The

 1 The most widely accepted unit for K_{d} is mL/g and there are no comparable English units that are generally accepted.

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approach used here is to recommend a single, deterministic value; ranges of values presented are not intended explicitly to be applied directly as probability distributions for probabilistic models. The following general procedure was used in the K_d analysis:

- (1) The mineralogical and hydrogeochemical characteristics of subsurface units along transport pathways at SRS and INL are described (e.g., sandy aquifer sediments and clay-rich confining units at SRS and alluvium, interbed sediments, and basalt at INL).
- (2) The radionuclide contaminants of concern at SRS and INL are designated (see Sections 2.3 and 3.3).
- (3) For each COC at SRS and INL
 - The general chemistry and important sorptive aspects of the contaminant are briefly described.
 - Site-specific measurements of K_d values and/or measurements of K_d values under analogous conditions to SRS or INL are summarized.
 - Literature-based reference K_d estimates and recommendations prepared for use in previous transport assessments at SRS and INL, which take into account site hydrogeochemical conditions, are mentioned (e.g., Kaplan, 2006 for SRS and Cooper, et al., 2006 for INL).
 - K_d values from compendia of sediment and rock types comparable to subsurface geologic units at SRS and INL are mentioned (e.g., Sheppard and Thibault, 1990; Looney, et al., 1987).
 - Based on the above data review, a recommended K_d value is presented.

Recommended K_d values in this review are based on knowledge of the hydrogeochemistry of each COC, available data on site chemical conditions, and reported K_d values from site-specific experiments or experiments conducted under the most analogous conditions. The most analogous conditions are chosen on the basis of sediment lithology and mineralogy and site hydrogeochemistry, with greatest weight being given to results from experiments that closely emulate field conditions. The following sections provide some background information on how (i) measurement methods can affect the interpretation of reported K_d values and (ii) specific environmental factors affect contaminant subsurface transport.

1.4 K_d Measurements

Although detailed evaluation of all methods used for measuring K_d s relevant for SRS and INL is beyond the scope of this report, it is useful to understand these methods when evaluating radionuclide transport models. A reported K_d is partly a function of the method used to determine it—batch, column, or *in-situ* measurements—and of intramethod variations, such as solid–solution ratio in batch experiments, flow rates in column experiments, and particle-size fractions measured for *in-situ* measurements. Batch or column tests are the two most commonly used methods for measuring equilibrium K_d s. Each method may use either single-mineral samples, mineral composites, or actual sediment or rock samples. While

single-mineral analyses provide important data on processes, actual samples and mineral composites are preferred when site-specific empirical K_ds are needed for model application. These latter methods will more accurately capture all mineral effects, as well as any water–rock interactions affecting system chemistry. Some sorption experiments use synthetic, rather than sampled, groundwaters. The synthetic waters reflect major cation and anion chemistry and pH but may not account for characteristics such as the presence of organic substances. Nevertheless, the use of synthetic groundwaters may allow experimenters better chemical control by avoiding chemical sampling effects such as degassing.

The batch method, which involves placing a solid and solution containing the contaminant of interest into a vessel and mixing the sample for a prescribed time, has the advantage of being relatively fast and simple. Batch testing is generally the method of choice for strongly sorbing contaminants due to the extremely long time requirements of column tests for such contaminants. Batch tests have the disadvantage of being conducted under conditions far removed from those observed in the field. Specifically, they are carried out under static conditions (no advective flow), at solid—liquid ratios much lower than present in the field, and only under saturated conditions.

The column method consists of packing a glass or plastic column with a sediment or crushed rock sample. Solutions containing the contaminant are pumped through the column, and the effluents are collected and analyzed. Breakthrough occurs when the effluent concentration rises substantially. Column tests have the advantage of being conducted under conditions that more closely approximate those in the field. Specifically, columns may be packed to have bulk densities and porosities similar to those observed in the field, column tests are conducted under advective and dispersive flow conditions similar to those in the field, and column tests may be conducted under a variety of different moisture conditions. When available, reported K_d values from column transport experiments will generally be given greater credence in K_d recommendations than K_d values from batch equilibrium experiments. Nevertheless, some limitations of column experiments must be acknowledged when evaluating their results. For example, column experiments using crushed basalt may present excessive surface area compared to field conditions or may reflect sorption on fresh surfaces rather than the weathered or coated surfaces present in an aquifer. A column test may yield lower K_d values than an analogous batch test if, for example, sorption kinetics are overcome by high column flow velocities. Other considerations include column aspect ratios and column diameter relative to grain size (e.g., Relyea, 1982).

In-situ field measurements of K_d are sometimes determined by measuring the ratio of the concentration of the contaminant of interest in sampled sediment or rock to the concentration in sampled groundwater or porewater. These field measurements may more accurately indicate contaminant partitioning because the K_d is derived under actual field conditions. However, Payne, et al. (2001) reported that *in-situ* uranium K_d s varied over a wide range of about two orders of magnitude at the Koongarra uranium deposit in Northern Australia. Variations in K_d s were attributed to the heterogeneity of the field site, which reflects the complexity of most geologic systems.

Several experimental methods have been used to determine the concentration of metal contaminants in field samples. The total digestion method involves complete dissolution of all solid phase components in the sample. Measurements of this sort may often lead to overestimates of K_d values if the contaminant concentration on the solid includes mass that is

not exchangeable with the aqueous phase. A more commonly used procedure is the hot nitric acid procedure outlined in SW–846 Method 3050b (EPA, 1986). This method is intended to release exchangeable ions specifically adsorbed to clays, oxides, and organic matter, but would not release metals associated with insoluble minerals. More information may be gained from sequential extraction methods, which have been developed to selectively remove metals associated with the solid. These procedures are useful in categorizing the metals into several operationally defined geochemical fractions, such as exchangeable, specifically adsorbed, and metals associated with carbonates, organic matter, and/or iron and manganese oxides. Disadvantages of sequential extraction methods are that the extractant may remove metals associated with other fractions. Also, readsorption of extracted metals to the remaining solid phase may lead to artificially low concentrations of the metal being associated with that fraction.

1.5 Factors Affecting Sorption and Transport

In the subsurface, contaminants have the potential to be found (i) dissolved in the groundwater, (ii) occupying ion exchange sites on mineral constituents, (iii) adsorbed on mineral constituents, (iv) associated with insoluble organic matter, and (v) precipitated as pure or mixed solids. Sorption, defined as the accumulation of ions at the interface between a solid phase and an aqueous phase (and including both ion exchange and adsorption), is affected both by the nature of the solid substrate and the chemical form of the contaminant and solvent (Table 1-1). Some minerals, such as clays and iron oxyhydroxides, contribute to higher K_d s due to surface chemical characteristics and high effective surface areas. Quartz, the main constituent of sandstone, tends to be less sorptive. Cation sorption has been correlated with such properties as pH and oxidation-reduction potential and the content of clays, organic matter, iron and

Table 1-1. Example Effects of Environmental Factors on K _d						
Factor (Increasing) General Effect on K _d						
Clay content	Increase					
Iron and manganese oxide content	Increase					
Quartz content	Decrease					
Grain size	Decrease					
Specific surface area	Increase					
рН	Increase (cation) Decrease (anion)					
Aqueous carbonate	Decrease (actinides) at high pH					
Oxidation-reduction potential or Eh	Decrease (uranium) Decrease (plutonium) Decrease (technetium)					
Cation exchange capacity	Increase					
Total dissolved solids	Decrease					

manganese oxides, and calcium carbonate. Anion sorption has been correlated with iron and manganese oxide content, pH, and oxidation-reduction potential.

The pH is a very important parameter, directly influencing sorption/desorption, aqueous complex formation, and oxidation-reduction reactions. Many sorption sites of mineral constituents are pH dependent (i.e., iron and manganese oxides, organic matter, carbonates, and edge sites of clay minerals). As pH decreases, the number of negative surface sites for cation sorption diminishes while the number of sites for anion sorption increases. As the pH becomes more acidic, cations face competition from H⁺ for available charged sites. In general, although dependent on the aqueous speciation of a given contaminant, maximum retention of cationic metals occurs at pH>7, and maximum retention of anionic metals occurs at pH<7. However, cationic metal retention can decrease with increasing pH due to the formation of negatively charged metal ligand complexes. For example, the sorption behavior of actinides (e.g., uranium and plutonium), which is characterized by rapidly increasing sorption at pH values between 5 and 7, decreases at pH values greater than 7 to 9 in response to the formation of negatively charged mixed carbonate/hydroxyl and carbonate species.

Radionuclides exist in solution as either free (uncomplexed) metal ions or in various soluble complexes with inorganic or organic ligands. Common inorganic ligands are SO_4^{2-} , CI^- , OH^- , NO_3^- , and CO_3^{2-} . The formation of stable complexes in solution can significantly affect the transport of contaminants relative to the free metal ion. Due to complexation, the resulting species may be positively or negatively charged or be electrically neutral. Therefore, depending on the surface charge of available sites, the metal complex may be weakly sorbed or strongly sorbed relative to the free ion. Because ions compete for available surface sites, a high total dissolved solids content would tend to decrease sorption of any single contaminant.

Contaminant oxidation state strongly influences sorption behavior. For example, U(VI), or uranium in the 6+ oxidation state, is considerably more mobile than U(IV). The near-surface oxidizing conditions important to transport at INL and SRS, therefore, will favor uranium mobility in comparison with more reducing conditions found, for example, in deep, static aguifers.

The sorption effects of these hydrogeochemical and mineralogic effects can be simulated through the use of mechanistic surface complexation models. Kaplan (2006) discussed the use of such models for contaminant transport studies at SRS, but concluded that data support was insufficient for their general use in performance assessments at SRS. Surface complexation models have also not been widely applied at INL.

There is both laboratory and field evidence that fractions of some actinides (e.g., plutonium, americium, curium, and uranium) can migrate at rates that are greater than predicted by equilibrium K_d measurements. For example, Newman, et al. (1996) and Fjeld, et al. (2001) observed high mobility fractions of plutonium and americium in column tests conducted on basalt and sedimentary interbeds from INL. The enhanced-mobility fraction responsible for the early elution of plutonium and americium in the column tests raises questions about the mechanisms responsible for transport of plutonium and americium—specifically, the potential role of colloids. Dai, et al. (2002) found that a small fraction of total mobile plutonium in SRS sediments is associated with colloids, and Kaplan, et al. (1994) found that appreciable fractions of plutonium, thorium, and uranium in an acidic plume have been traveling through an aquifer underlying SRS in association with mobile colloids at rates appreciably greater than anticipated.

If these enhanced-mobility fractions are substantial and are being transported as colloids, their effects need to be considered in applying sorption-based transport models.

Aside from mineralogic constituents, hydrogeologic characteristics of the host rock also affect solute transport. For example, lower grain size and lower porosity may promote sorption by increasing the available specific surface area. Low permeability may lead to slower flow rates that could make sorption more effective because any kinetic barriers are more easily surmounted. Abundant fractures may provide fast water pathways that diminish water–rock interaction; in addition, fracture surfaces may provide contrasting mineralogic sorption environments versus the matrix. In the case of INL, for example, basalt fracture iron oxide coatings may be a significant factor for contaminant K_d s.

2 SAVANNAH RIVER SITE

2.1 Waste Management Background

Two types of radioactive waste disposal facilities governed by the Ronald W. Reagan National Defense Authorization Act for Fiscal Year 2005 consultative process are located at the Savannah River Site (SRS): grouted tanks and salt waste vaults. In the F and H-Areas of the General Separations Area (GSA), the U.S. Department of Energy (DOE) plans to close liquid radioactive waste tanks in place by removing waste to the maximum extent practical, filling the emptied tanks with grout (intended to immobilize any residual radionuclides), and constructing an engineered cap (e.g., DOE, 2005). In Z-Area of the GSA, DOE plans to dispose of a cementitious wasteform ("saltstone") containing the low-activity stream resulting from treatment of tank salt wastes (DOE, 2006a). For both disposal scenarios, waste determinations are supported by a performance assessment that includes modeled groundwater radionuclide transport. Because these facilities are all located on the same upland area in the GSA within a 5-km [3-mi] distance, they have the same environment with respect to hydrologic transport.

As mentioned in Section 1.2, the hydrochemistry of groundwater in the GSA bears the imprints of acid contamination, which is currently being actively remediated. These efforts are expected to lead to soil and groundwater cleanup and source isolation within the next few decades (e.g., Westinghouse Savannah River Company, 2002). By the time potential radionuclide releases would occur from closed tanks and the salt waste facility, it is expected that groundwater pH levels will have returned to natural background values.

2.2 Description of the General Separations Area

The GSA is an approximately 24-km² [9.3-mi²] area located in the central part of SRS. This area contains a number of radioactively contaminated facilities including a low-level radioactive waste disposal area, two large nuclear materials separation facilities, the Defense Waste Processing Facility, the Saltstone Disposal Facility, the Saltstone Processing Facility, the H-Tank Farm, and the F-Tank Farm. These facilities are located approximately 10 km [6.2 mi] from the nearest SRS site boundary on well-drained topographic highs bounded by three small black water streams called Upper Three Runs Creek, Fourmile Branch, and McQueen Branch.

SRS is underlain by Tertiary age unconsolidated and semiconsolidated sediments of the Floridan Aquifer System. The hydrostratigraphy of SRS consists of three vertically stacked aquifer zones and two confining zones that will be potentially impacted by release of contaminants from facilities within the GSA. The hydrostratigraphy of SRS has been the subject of several different classifications. This report incorporates the hydrostratigraphic nomenclature currently established for SRS vicinity by Aadland, et al. (1995). From top to bottom, the hydrostratigraphic units present beneath the GSA are: the Upper Three Runs aquifer, the Gordon confining unit, and the Gordon aquifer (Figure 2-1). The Upper Three Runs aquifer is locally divided into informal lower and upper aquifer zones separated by the tan clay confining zone.

The Upper Three Runs aquifer includes all sediments from the ground surface to the top of the Gordon confining unit. The upper aquifer zone (commonly referred to as the Water Table aquifer) consists of all sediments from the ground to the top of the tan clay confining zone and

	HYDROSTRATIGRAPIC UNIT	THICKNESS RANGE (FEET)		
Upper Three Runs Aquifer	Upper Zone (Water Table Aquifer)	0-110		
e Rı	Tan Clay	0-33		
Upper Thre	Lower Zone (Barnwell McBean Aquifer)	39-81		
G	ordon Confining Unit (Green Clay)	2-30		
	Gordon Aquifer Unit	52-107		
Crouch Branch Confining Unit >100				

Figure 2-1. Hydrostratigraphic Units at the GSA (Modified From Buice, et al., 2005)

includes the Upland unit, Tobacco Road Sand, and part of the Dry Branch Formation (Aadland, et al., 1995). This zone is characterized by sand and clayey sand with minor intercalated clay layers. The sediment within the Upland unit is commonly very dense and clayey and contains some gravelly sand. The term "Upland unit" is an informal name used to describe sediments at higher elevations located in the Upper Coastal Plain in southwestern South Carolina. The tan clay confining zone includes sediment of the Dry Branch Formation. The zone contains light-yellowish tan to orange clay and sandy clay interbedded with clayey sand and sand. The tan clay is dispersed vertically and horizontally and is not continuous over long distances (Aadland, et al., 1991). The lower aquifer zone (commonly referred to as the Barnwell McBean aquifer) consists dominantly of fine-grained, well-sorted sand and clayey sand of the Tinker/Santee Formation and parts of the Dry Branch Formation. The bulk of carbonate sediment present beneath the GSA is included in the lower aquifer zone. Descriptions of drill core indicate that the carbonate sediment in the SRS vicinity has a dominant siliciclastic component and consists of calcareous sand, micritic sand, shelly sand, and some sandy calcarenite and shelly limestone.

Strom and Kaback (1992) determined the mineralogical composition of 28 samples from the Upper Three Runs aquifer collected from wells within SRS by x-ray diffraction. The samples consisted of quartz with 2- to 40-wt% detrital clay. An examination of the mineralogical data reported by Strom and Kaback (1992) indicates that smectite and kaolinite are the dominant clay minerals in the upper zone of the Upper Three Runs aquifer and the tan clay confining unit and that smectite is the dominant clay mineral in the lower zone of the Upper Three Runs aquifer. Minor to trace amounts of illite, chlorite, pyrite, plagioclaise, and K-feldspar were present in both aquifer zones and the tan clay confining unit. Calcite was detected in minor to trace amounts in the lower zone of the Upper Three Runs aquifer.

Several researchers have described the lithological and mineralogical properties of sediments from the Upper Three Runs aquifer used in radionuclide partitioning studies at SRS. Serkiz and Johnson (1994) analyzed sediment samples taken at depths ranging from 4 to 28 m [13 to 93 ft] from the upper zone of the Upper Three Runs aquifer downgradient of the F- and H-Area seepage basins within the GSA. Sand, silt, and clay contents of the sediments were 57 to 99 wt%, 0 to 12 wt%, and 0.5 to 42 wt%, respectively. The cation exchange capacity (CEC) of the sediment varied from <0.1 to 2.9 meq/100 g. The CECs were noted to be relatively low due to the high percentage of sand in the sediments. No correlation between clay fraction of the soil and CEC was observed.

Kaplan (2003) described the characteristics of an upland sample collected at a depth of 2 m [6 ft] near the E-Area low-level waste facility and a wetland sample collected from the surface of a wetland area along the Savannah River. The particle size distribution of the two samples was similar (79- to 80-wt% sand, 14- to 15-wt% silt, and 5- to 6-wt% clay). Smectite and kaolinite were the dominant clays in the upland sample, and smectite was the dominant clay in the wetland sample. The CECs of the samples were 4.75 and 3.99 meq/100g for the wetland and upland sample, respectively. Although organic carbon contents were low in both samples, the wetland sample had an appreciably greater organic carbon concentration {1,395 mg/kg [1,395 ppm]} than the upland sample {<200 mg/kg [<200 ppm]}. The very low organic carbon concentration in the upland sample indicates that little organic matter is leached into the underlying sediments.

Hu, et al. (2005) described the characteristics of a surface soil, a subsoil {composite of sediments collected at a depth of 12 m [39 ft]}, and an aquifer sediment collected from the upper zone of the Upper Three Runs aquifer. Sand, silt, and clay weight percentages were 87.6, 2.9, and 9.5 for the surface soil; 76.7, 3.5, and 19.8 for the subsoil; and 97.2, 0.9, and 1.9 for the aquifer sediment. Smectite and kaolinite were the dominant clays in all the samples, and goethite (an iron oxyhydroxide) was noted to be present in the subsoil and aquifer sediment. Extractable iron contents (a measure of iron-oxide mineral coatings) were 2.82 g/kg [2,820 ppm] for the surface soil, 16.2 g/kg [16,200 ppm] for the subsoil, and 7.3 g/kg [7,300 ppm] for the aquifer soil. The CECs of the samples were 1.3, 1.8, and 0.064 meq/100 g for the surface soil, subsoil, and aquifer sediment, respectively. The surface soil contained 2.3-wt% organic matter, whereas the subsoil and aquifer soil contained 0.07-wt% and 0.01-wt% organic matter, respectively. Low organic matter contents in the subsoil and aquifer soil again indicates that little organic matter is leached into the underlying sediments.

Barnett, et al. (2000) described the chemical and mineralogical properties of a sediment sample acquired at a depth of 45 m [150 ft] from the lower zone of the Upper Three Runs aquifer (McBean Formation) near the old burial grounds within the GSA. The sample consisted of

69-wt% sand, 7-wt% silt, and 24-wt% clay and was dominated by sand-sized quartz coated with iron oxides. Smectite was the dominant clay mineral and also contained iron-oxide coatings. The sediment contained 0.12 g/kg [120 ppm] organic matter, 0.06 g/kg [60 ppm] manganese, and 25.7 g/kg [25,700 ppm] iron.

The Gordon confining unit includes sediments of the Warley Hill Formation and is commonly referred to as the "green clay." The unit comprises layers of interbedded silty and clayey sand, sandy clay, and clay that thicken and thin and pinch out abruptly. The clay is stiff and hard and is often fissile. Glauconite is a common constituent and imparts a distinctive greenish color to the sediment, hence the informal name "green clay." Zones of silica-cemented sand and clay are noted in some cores taken from SRS (Aadland, et al., 1995). In the vicinity of the GSA, the Gordon confining unit includes some calcareous sediment and limestone, primarily calcareous sand and clayey sand with subordinate calcareous and micritic clay, sandy micrite, and limestone.

X-ray diffraction analyses of four muddy-sand samples, collected from the Gordon confining unit from wells within SRS, were reported to consist mostly of quartz with 18- to 42-wt% clay minerals (Strom and Kaback, 1992). Smectite was the dominant clay mineral, with minor to trace amounts of illite and kaolinite. Textural analyses made on 10 sandy mud and muddy sand samples from the Gordon confining unit averaged 47.3-wt% mud, 52.2-wt% sand, and 0.5-wt% gravel (Bledsoe, et al., 1990).

The Gordon aquifer constitutes the basal unit of the Floridan Aquifer System. Beneath the GSA, the Gordon aquifer is made up of the loose sand and clayey sand of the Congaree and Fourmile Formations (Aadland, et al., 1991; 1995). The sand within the Gordon aquifer is yellowish to grayish orange in color and is sub to well rounded, moderately to poorly sorted, and medium to coarse grained. Pebbly layers and zones of iron and silica cement are common. Clay beds and stringers are present, but they are too thin and discontinuous to be more than local confining beds. Small amounts of calcareous sediment occur sporadically within the unit.

Petrographic analyses of 42 thin sections show that sand-rich sediments from the Gordon aquifer unit consist of 98.4-wt% framework quartz, 0.3-wt% feldspar, 0.5-wt% heavy minerals, 0.4-wt% opaque minerals, 0.4-wt% muscovite, and 0.1-wt% gluaconite (Robertson and Thayer, 1990). Strom and Kaback (1992) did semiquantitative x-ray diffraction analyses of 18 Gordon aquifer clay-bearing samples collected within SRS. Their results showed that the sand consists mostly of quartz with 5- to 18-wt% clay. Smectite and kaolinite were reported to be the dominant clay minerals, with minor or trace amounts of illite. Authigenic minerals included clinoptilolite, cristobalite, and chalcedony. X-ray diffraction analyses of 4 sandy mud samples from intra-aquifer confining layers were reported to consist of quartz and 51- to 64-wt% clay minerals, chiefly kaolinite with minor illite and smectite.

Groundwater beneath the GSA flows toward streams and swamps and into the Savannah River at rates ranging from inches to a few hundred feet per year. The depth at which nearby streams cut into sediments, the lithology of the sediments, and the orientation of the sediment formations control the horizontal and vertical movement of the groundwater. The valleys of smaller perennial streams, such as Fourmile Branch and McQueen Branch, allow discharge from the shallow saturated geologic formations (e.g., the Upper Three Runs aquifer). The valleys of major streams, such as Upper Three Runs Creek, drain formations of greater depth (e.g., the

Gordon aquifer). Because of these discharges, flow is not expected to penetrate into stratigraphic units beneath the Gordon aquifer within and in the vicinity of the GSA.

Strom and Kaback (1992) investigated the chemistry of water in aquifers at SRS. Along the northern boundary of SRS, groundwaters in the Upper Three Runs and Gordon aquifers are very low in total dissolved solids {< 20 mg/L [<20 ppm]} and contain high concentrations of dissolved oxygen {6 to 11 mg/L [6 to 11 ppm]}. Waters in this area are relatively acidic (pH 5 to 6), and there are no predominant cations or anions in the water. In this portion of SRS, the confining units that separate shallow aquifers are only of local extent, and the hydraulic gradient is downward from the Tertiary formations into the underlying Cretaceous formations.

South of Upper Three Runs Creek in the central and southern portions of SRS, which includes the GSA, shallow confining units are more laterally extensive, and waters within the Upper Three Runs and Gordon aquifers are geochemically distinctive. The chemical characteristics of waters in the shallow aquifers and confining units in this part of SRS are listed in Table 2-1. Groundwater in this area has total dissolved solids ranging from 5 to 150 mg/L [5 to 150 ppm]. Calcium and bicarbonate are the predominant ions, due largely to chemical reactions between groundwater and calcite-bearing units that are scattered sporadically through the Tertiary section. Waters in this area are slightly acidic to neutral (pH 5.2 to 7.7). Dissolved oxygen concentrations are variable {0.02 to 10 mg/L [0.02 to 10 ppm]} suggesting some interaction with organic matter.

Aqueous wastes from a nuclear materials processing facility on the GSA were released over a 33-year period to unlined seepage basins. Three basins covering approximately 2.8 ha [7 acres] were built in the F Area, and 4 basins covering 6.5 ha [16 acres] were constructed in the H Area. The seepage basins were designed to allow the effluent to percolate into the surrounding environment, where soil sorption processes were expected to retard the movement of the waste constituents. The wastes contained high concentrations of strong acids, sodium, tritium, and trace amounts of numerous radionuclides and metals (Kaplan, et al., 1994). Due to these releases, soils and groundwater beneath the GSA have widely varying groundwater chemistries and contaminant concentrations.

Several researchers have described the aqueous chemistry of acid-contaminated groundwater in the Upper Three Runs aquifer beneath the GSA (Serkiz and Johnson, 1994; Kaplan, et al., 1994; Dai, et al., 2002). These waters have acidic pHs (3.0 to 4.2); low inorganic and organic carbon contents; and high nitrate, tritium, and dissolved oxygen contents. Waters have elevated concentrations of cations such as Al³⁺, Fe³⁺, K⁺, Mg²⁺, Na⁺, and Si⁴⁺, consistent with acid leaching of aquifer matrix materials such as clays (e.g., kaolinite) and surface mineral coatings (e.g., iron oxyhydroxides). Contaminated waters are also enriched in radionuclides such as Pu-239, Am-241, Cm-243, Cm-244, and U-238 (Kaplan, et al., 1994) and in metals such as lead, zinc, cadmium, and copper (Dai, et al., 2002).

The chemistry of groundwater and mineralogic composition of sediments beneath the GSA will affect contaminant transport. Due to the release of acidic aqueous wastes, groundwater beneath the GSA currently has widely varying chemistry. However, in the future it is anticipated that acid-contaminated waters beneath the GSA will return to normal chemical conditions due to natural flushing of the aquifer and clay confining units by infiltrating rainwater. In addition, efforts are underway to remediate acid-contaminated areas beneath the GSA to speed the return to normal chemical conditions. These efforts are expected to lead to soil and

Table 2-1. Chemistry of Water in Shallow Aquifer and Confining Units in the Central and Southern Part of the Savannah River Site* **Upper Three** Runs Aquifer **Upper Three** (Upper Zone) **Runs Aquifer** Gordon and Tan Clay (Lower Zone) **Confining Unit Gordon Aquifer** 5.2 - 6.36.0 - 7.75.8-6.8 5.7-7.6 рΗ Specific 66–216 22-81 6.8-192 39-102 Conductance (µS) **Total Dissolved** 5-53 45-102 19-55 28-150 Solids (mg/L) 0.07-0.23 (-0.09)-0.12 Oxidation-0.09 - 0.30(-0.05)–0.13Reduction Potential (V) Eh (mV) 291-497 150-325 266-427 113-316 Dissolved Oxygen 7.33-9.6 0.41 - 9.825.35-9.98 0.02 - 4.33(mg/L) Ca²⁺ (mg/L) 1.2-12 7.2 - 384.4-17 9-45 Mg^{2+} (mg/L) 0.32 - 0.470.38 - 0.740.24 - 0.790.48 - 1.1Na⁺ (mg/L) 0.66 - 3.31.2 - 2.80.9 - 1.50.87 - 1.9K⁺ (mg/L) 0.3 - 2.20.43 - 2.90.52 - 1.60.35 - 4.31.2 - 3012-47 27-114 HCO_3^- (mg/L) 20-101 SO_4^{2-} (mg/L) 0.63-13.8 0.36 - 1.770.73 - 3.80.68 - 2.74Cl⁻ (mg/L) 1.95-2.88 1.50 - 7.582.16-3.31 2.40-2.74 F- (mg/L) <0.05-0.07 <0.05-0.34 <0.05-0.10 <0.05-0.11

groundwater cleanup, and source isolation within the next few decades (e.g., Westinghouse Savannah River Company, 2002). Due to natural flushing and remediation, groundwater in the hydrostratigraphic units beneath GSA is expected to have the following chemical characteristics that will affect contaminant transport:

- Slightly acidic to neutral pH (pH of 5.2 to 7.7; Table 2-1)
- Variable total dissolved solids (TDS)
- Cation concentrations dominated by Ca²⁺ and anion concentrations dominated by HCO₃⁻
- Variable dissolved O₂ concentrations due to interaction with organic matter

^{*}Strom and Kaback provide no chemistry data for the tan clay confining unit. Based on mineralogic similarities (e.g., the lack of calcite), the tan clay is expected to have chemical properties similar to the upper zone of the Upper Three Runs Aquifer. Strom, R.N. and D.S. Kaback. "SRP Baseline Hydrogeologic Investigation: Aquifer Characterization, Groundwater Geochemistry of the Savannah River Site and Vicinity." WSRC–RP–92–450. Aiken, South Carolina: Westinghouse Savannah River Company. 1992.

Natural organic matter is present in surface sediments and may be present in the less hydraulically conductive clay-rich confining units at GSA, potentially resulting in reducing conditions in some isolated, relatively immobile groundwater.

Mineralogic characteristics will affect the partitioning of contaminants between groundwater and solid materials in hydrostratigraphic units beneath GSA, including

- Compositions dominated by quartz grains and clays that are commonly coated with iron oxides and iron-oxyhydroxides
- Clay mineral contents accounting for 1 to 25 wt% of sediments composing aquifer units and up to 65 wt% of sediments composing confining units
- The presence of calcareous sediments in deeper hydrostratigraphic units (i.e., in the lower zone of the Upper Three Runs aquifer, Gordon confining unit, and Gordon aquifer)

Iron oxides and iron oxyhydroxides can strongly adsorb actinides, such as uranium, under certain chemical conditions. For example, uranium sorbs strongly to ferrihydrite over the pH range of 5 to 8 (Waite, et al., 1994). Clay minerals provide sites for sorption and ion exchange with contaminant-bearing water. For example, cesium adsorbs primarily via ion exchange with clay minerals. Smectite and kaolinite have been reported to be the most abundant clay minerals in sediments at SRS (Strom and Kaback, 1992). Smectite has the largest cation exchange capacities of the clay minerals present at SRS.

Reaction of groundwater and calcareous sediments can increase calcium and bicarbonate concentrations, thereby increasing groundwater pH and ionic strength. The sorption of many of the contaminants at SRS is sensitive to pH. For example, sorption of strontium and cesium, which exist as cationic species in natural waters, increases with increasing pH, while sorption of selenium, technetium, and iodine, which exist as anionic species in natural waters, decreases with increasing pH. On the other hand, the sorption behavior of actinides, such as uranium and plutonium, is characterized by rapidly increasing sorption at pH values between 5 and 7 and then decreasing sorption at pH values greater than 7 to 9 in response to the formation of mixed carbonate/hydroxyl and carbonate species. In addition, the increased concentration of calcium in groundwater resulting from dissolution of calcite-bearing sediments can reduce the sorption of contaminants such as strontium and lead due to competition for adsorption sites.

2.3 Designation of Risk-Significant Radionuclides

The COCs chosen for analysis in this report are based both on previous model results and on more general considerations of hydrochemical behavior and dose effects. For example, a radionuclide with a high-dose-conversion factor could be important if its sorption coefficient is not as high as supposed. In addition, inclusion on the DOE list of highly radioactive radionuclides was considered. DOE analyses (Rosenberger, et al., 2005; Buice, et al., 2005) have indicated that isotopes of selenium, technetium, iodine, and neptunium are potentially important in groundwater pathway dose calculations for salt waste and grouted tank facilities. Uranium is important in waste inventories and is relatively mobile in many environments. Other actinides—plutonium, americium, and curium—are included due to potentially important dose effects if sorption coefficients are overestimated. Isotopes of strontium and cesium, though

relatively short lived, have high inventories and could be important if groundwater travel pathways are short.

2.4 Sorption Coefficients

The hydrostratigraphic units that will be affected by release of contaminants from waste storage facilities at the GSA (e.g., the Saltstone Disposal Facility) are the upper and lower aquifer zones and the tan clay confining zone of the Upper Three Runs aquifer, the Gordon confining unit, and the Gordon aquifer. With the exception of surficial soils and sediments of the upper zone of the Upper Three Runs aquifer, little or no site-specific sorption data exists for the hydrostratigraphic units beneath the GSA. As a result, previous SRS contaminant transport models have classified aquifer units as sandy-textured sediment or soil and, for each COC, a single or uniform K_d has been assigned to all aquifer units (Cook, et al., 2005, 2000; Buice, et al., 2005; Kaplan, 2005; Rosenberger, et al., 2005). Similarly, confining units (i.e., the tan clay and Gordon confining unit) beneath the GSA have been classified as clay sediments and assigned a single or uniform K_d for each COC.

In this review, site-specific K_d measurements and literature-recommended K_d values for the COCs are analyzed and reviewed to derive a recommended K_d value for each hydrostratigraphic unit beneath SRS. In many instances where site-specific K_d measurements are lacking for hydrostratigraphic units, the mineralogic and hydrogeochemical characteristics of the units are evaluated to derive a K_d recommendation. The mineralogical and expected hydrogeochemical characteristics of each hydrostratigraphic unit are summarized in Table 2-2. Examination of the listed mineralogical and chemical properties in Table 2-2 indicates that the mineralogical and chemical properties of the aquifer units and the confining units are quite similar. The most notable difference is the presence of calcareous (calcite-bearing) sediments in the deeper hydrostratigraphic units (i.e., the lower zone of the Upper Three Runs aquifer, the Gordon confining unit, and the Gordon aquifer) and the related higher minimum pH of these units.

Sheppard and Thibault (1990) (see also Thibault, et al., 1990) provided a compilation of soil solid/liquid K_d values for use in environmental assessments for nuclear waste disposal. This compilation was originally prepared for assessing the concept of disposing Canadian spent fuel deep in Precambrian Shield plutonic rock. Due to the absence or lack of site-specific K_d measurements on soils and sediments for many of the COCs at SRS, K_d values from the compilation of Sheppard and Thibault (1990) have been treated as an independent source and used in performance assessments and special analyses at SRS (Cook, et al., 2005, 2000; Buice, et al., 2005; Kaplan, 2005; Rosenberger, et al., 2005).

The compilation of Sheppard and Thibault (1990) listed K_d values for 48 different elements in 4 different mineral soil types (sand, loam, clay, and organic). Values were compiled from measurements reported in the peer-reviewed literature. If no data existed in the literature for a given element, then the soil-to-plant concentration ratio was used to indicate the element's bioavailability and to predict a K_d value (Baes, et al., 1984). The soil-to-plant concentration ratio is a measure of the root uptake of a radionuclide incorporated into the surface horizon of soil and represents the ratio of radionuclide concentration in plant and soil at harvestable maturity. Based on a strong negative correlation between the soil-to-plant concentration ratio and K_d , Thibault, et al. (1990) used observed and predicted K_d values from regression analyses to

Table 2-2. Mineralogical and Hydrogeochemical Characteristics of Hydrostratigraphic Units at the Savannah River Site								
Unit	Lithology*	Mineralogy†	pH Range‡	Other Hydrogeochemical Aspects‡				
Upper Three Runs Aquifer: Upper Zone	Sand and clayey sand; minor intercalated clay layers	Major constituents: quartz, smectite, kaolinite Minor constituents: plagioclaise, K-feldspar, illite, pyrite, chlorite, iron oxyhydroxides	5.2–6.8	Variable dissolved O ₂ lons dominated by Ca ²⁺ , HCO ₃ ⁻ total dissolved solids: <50 mg/L [<50 ppm]				
Tan Clay Confining Unit	Clay and sandy clay interbedded with clayey sand and sand	Major constituents: quartz, smectite, kaolinite Minor constituents: plagioclaise, K-feldspar, illite, pyrite, chlorite, iron oxyhydroxides	5.2–6.8	Variable dissolved O ₂ lons dominated by Ca ²⁺ , HCO ₃ ⁻ total dissolved solids: <50 mg/L [<50 ppm]				
Upper Three Runs Aquifer: Lower Zone	Sand and clayey sand; interspersed calcareous sand, micritic sand, and shelly sand	Major constituents: quartz, smectite Minor constituents: kaolinite, plagioclaise, K-feldspar, illite, pyrite, chlorite, iron oxyhydroxides, calcite	6.0–7.7	Variable dissolved O ₂ lons dominated by Ca ²⁺ , HCO ₃ ⁻ total dissolved solids: 50 to 150 mg/L [50 to 150 ppm]				
Gordon Confining Unit	Interbedded silty and clayey sand, sandy clay and clay; some calcareous sediment and limestone	Major constituents: quartz, smectite, glauconite Minor constituents: kaolinite, illite, iron oxyhydroxides, calcite	5.8–6.8	Variable dissolved O ₂ lons dominated by Ca ²⁺ , HCO ₃ ⁻ total dissolved solids: <50 mg/L [<50 ppm]				

Table 2-2. Mineralogical and Hydrogeochemical Characteristics of Hydrostratigraphic Units at the Savannah River Site (continued)

Unit	Lithology*	Mineralogy†	pH Range‡	Other Hydrogeochemical Aspects‡
Gordon Aquifer	Loose sand and clayey sand; zones of iron and silica cement; small amounts of calcareous sediment	Major constituents: quartz, smectite, kaolinite Minor constituents: illite, feldspar, muscovite, pyrite, glauconite, clinoptilolite, cristobalite, chalcedony, iron oxyhydroxides, calcite	5.7–7.6	Variable dissolved O ₂ lons dominated by Ca ²⁺ , HCO ₃ ⁻ total dissolved solids: 50 to 150 mg/L [50 to 150 ppm]

^{*}Lithologic characterization based on data from Aadland, R.K., J.A. Gellici, and P.A. Thayer. "Hydrogeologic Framework of West-Central South Carolina." Report 5. PIT-MISC-0112. State of South Carolina Department of Natural Resources, Water Resources Division. 1995.

†Mineralogic characterization based on data from Strom, R.N. and D.S. Kaback. "SRP Baseline Hydrogeologic Investigation: Aquifer Characterization, Groundwater Geochemistry of the Savannah River Site and Vicinity." WSRC–RP–92–450. Aiken, South Carolina: Westinghouse Savannah River Company. 1992; Kaplan, D.I. "Influence of Surface Charge of an Fe-Oxide and an Organic Matter Dominated Soil on Iodide and Pertechnetate Sorption." *Radiochimica Acta.* Vol. 91, No. 3. pp. 173–178. 2003; Hu, Q., Z. Pihong, J.E. Moran, and J.C. Seaman. "Sorption and Transport of Iodine Species in Sediments from Savannah River and Hanford Sites." *Journal of Contaminant Hydrology.* Vol. 78. pp. 185–205. 2005; and Barnett, M.O., P.M. Jardine, S.C. Brooks, and H.M. Selim. "Adsorption and Transport of Uranium(VI) in Subsurface Media." *Soil Science Society American Journal.* Vol. 64. pp. 908–917. 2000.

‡Chemical characterization based on data from Strom, R.N. and D.S. Kaback. "SRP Baseline Hydrogeologic Investigation: Aquifer Characterization, Groundwater Geochemistry of the Savannah River Site and Vicinity." WSRC–RP–92–450. Aiken, South Carolina: Westinghouse Savannah River Company. 1992. The pH range does not include lower values resulting from acid contamination.

develop a model to estimate the K_d of a given radionuclide for each soil type from a measured soil-to-plant concentration ratio.

In the compilation of Sheppard and Thibault (1990), the soils that contained at least 70-wt% sand-sized particles were classified as sand soils, and those containing at least 35-wt% clay-sized particles were classified as clay soils. Loam soils had an even distribution of sand-, clay-, and silt-sized particles or consisted of up to 80-wt% silt-sized particles. Organic soils contained >30-wt% organic matter. Based on particle size analyses, aquifer sediments beneath the GSA are most comparable to sand soils, and clay confining units are most comparable to clay soils defined by Sheppard and Thibault (1990). The range and "default" K_d values for sand and clay reported by Sheppard and Thibault (1990) for each COC in this review are graphically illustrated in Figures 2-2 through 2-11 and Figures 2-13 through 2-17. "Default" K_d s for a given COC were the geometric mean of all reported K_d observations or were predicted using soil-to-plant concentration ratios for COCs with no observations (Sheppard and Thibault, 1990).

To aid in environmental assessments of waste sites at the Savannah River Plant, Looney, et al. (1987) estimated K_d ranges and "reference" soil K_d values for 31 elements. The estimated K_d ranges presented by Looney, et al. (1987) were based on the literature and represent, as available, onsite measurements, values for silty soils, and values for all soils. Literature sources used to estimate the K_d ranges included Sheppard, et al. (1984); Baes and Sharp (1983); Hoeffner (1985); Ames and Rai (1978); Callahan, et al. (1979); and U.S. Environmental Protection Agency (EPA) (1985). The "reference" K_d values specified by Looney, et al. (1987) for SRS soils were predicted based on the general solution and solid surface properties and reactions expected in the subsurface at SRS.

A compilation of estimated K_d values for contaminants at the SRS tabulated by Kaplan (2006) has been useful in this data review. Kaplan (2006) estimated "best" and "reasonably conservative" K_d values for 40 elements in sandy subsurface sediment and clayey subsurface sediment for use in SRS-related performance assessment calculations related to SRS. The "best" estimates provide guidance on the most likely K_d value for a given contaminant and are based on site-specific measurements and literature experimental data. When site-specific and literature data were not available, chemical analogs were used in the "best" estimates of K_a values. The "reasonably conservative" values represent lower bounding values that consider the range of physical, chemical, and mineralogical conditions that often lead to enhanced radionuclide migration. Ideally, the "reasonably conservative" value was based on the lower limit of multiple K_d measurements. In the absence of sufficient data, the "reasonably conservative" value was based on an assumed range of values, using the "best" value as a central point in the range. The sandy subsurface sediment was conceptualized as a sediment containing a clay and silt content less than 25 wt%, and the clayey subsurface sediment was conceptualized as a sediment containing a clay and silt content of 25 to 45 wt%. Based on these particle size characteristics, aguifer units beneath the GSA are most comparable to the sandy subsurface sediments, and clay confining units are most comparable to the clayey subsurface sediments.

When little or no site-specific or analogous K_d data could be found for a COC at SRS, the default soil and clay K_d values specified by Sheppard and Thibault (1990), the soil reference K_d values estimated for SRS by Looney, et al. (1987), and the "reasonably conservative" and "best" K_d values estimated for sandy and clayey subsurface sediment at SRS by Kaplan (2006) were in

some cases directly incorporated into the K_d recommendations in this review. In other cases, these values are presented to help determine whether the recommended K_d values for SRS are reasonable.

2.4.1 **Carbon**

The C-14 present in SRS waste streams originates from nuclear fuel processing and is predominantly inorganic in nature. In the GSA environment, inorganic carbon transport occurs via both slow aqueous transport as HCO_3^- ion and rapid transport of CO_2 gas. Carbon-14 transport is likely to be affected by reactions other than sorption, but an empirical K_d approach may be sufficient for understanding potential transport.

McIntyre (1988) performed batch experiments using carbon, Savannah River Plant soil, and Savannah River Plant burial ground groundwater. The Savannah River Plant soil composition was 72-wt% sand, 5-wt% silt, and 23-wt% clay with a moisture content of near 16 percent. The groundwater used in the experiments had a pH of 6.0. After 7 hours of equilibration, carbon K_d values ranged from 1.6 to 2.3 mL/g (average of 2.0 mL/g). After 72 hours of equilibration, carbon K_d values ranged from 46.9 to 63.2 mL/g (average of 55 mL/g).

Sheppard and Thibault (1990) estimated a carbon default K_d for sand of 5.0 mL/g (three observations ranging from 1.7 to 7.1 mL/g) and a carbon default K_d for clay of 1.0 mL/g (based on a soil-to-plant concentration ratio). Looney, et al. (1987) estimated a reference SRS soil K_d of 0.01 mL/g for carbon. Kaplan (2006) estimated "best" and "reasonably conservative" K_d values of 0 mL/g for carbon in both sandy subsurface sediment and clayey subsurface sediment at SRS. The estimation of Kaplan (2006) was based on the judgment that carbon is expected to exist as HCO_3^- in the aqueous phase, and therefore, little sorption is expected.

Measured, estimated, and recommended carbon K_d values are summarized in Table 2-3. Site-specific carbon K_d values from the batch experiments of McIntyre (1988) using Savannah River Plant soil (upper zone of Upper Three Runs aquifer) ranged from 1.6 to 2.3 mL/g after 7 hours of equilibration. This K_d range is bracketed by the carbon K_d value estimated by Kaplan (2006) (0 mL/g) for sandy subsurface sediment at SRS, the SRS reference soil carbon K_d value estimated by Looney, et al. (1987) (0.01 mL/g), and the carbon K_d value estimated for sand by Sheppard and Thibault (1990) (5 mL/g). A carbon K_d value of 2 mL/g, which is the average of the measured carbon K_d values in the 7-hour batch test of McIntyre (1988), is recommended for the upper zone of the Upper Three Runs aquifer. The mineral composition of the aquifer units is not significantly different and should not greatly affect carbon sorption. Due to the presence of calcareous sediments, groundwater in the lower zone of the Upper Three Runs aquifer and the Gordon aquifer have higher bicarbonate contents than the upper zone of the Upper Three Runs aquifer. These higher bicarbonate contents could potentially enhance the mobility of carbon as HCO_3^- . Therefore, a carbon K_d value of 1 mL/g is recommended for the lower zone of the Upper Three Runs Aquifer and the Gordon Aquifer.

No site-specific data are available for carbon partitioning to the clay confining units at SRS. Based on the carbon K_d value of 1 mL/g estimated by Sheppard and Thibault (1990) for clay and the carbon K_d value of 0 mL/g estimated by Kaplan (2006) for SRS clayey subsurface

Table 2-3. Measured, Estimated, and Recommended K _d Values for Carbon in Aquifer Sediments and Clay Confining Units at the Savannah River Site							
	Measured Carbon K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Carbon K _d (mL/g)*	Generic Estimated Carbon K _d (mL/g)†	Recommended Carbon K _d (mL/g)		
Upper Three Runs Aquifer: Upper	1.6–63.2‡	Batch	0/0	5	2		
Tan Clay Confining Unit	_		0/0	1	1		
Upper Three Runs Aquifer: Lower	_		0/0	5	1		
Gordon Confining Unit	_		0/0	1	1		
Gordon Aquifer	_		0/0	5	1		

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

sediment and considering the carbon K_d values of 1 and 2 mL/g recommended for aquifer units at SRS, a carbon K_d value of 1 mL/g is recommended for confining units at SRS.

The position of the carbon K_d values recommended in this review with respect to measured and estimated site-specific and generic carbon K_d ranges and values is graphically displayed in Figure 2-2. This figure illustrates the uncertainty associated with the carbon K_d values recommended for SRS aquifer units and clay confining units. The recommended carbon K_d values are associated with high uncertainty due to limited site-specific sorption data. The recommended carbon K_d values of 1 and 2 mL/g for SRS aquifer units are based solely on site-specific carbon sorption data reported by McIntyre (1988). The recommended carbon K_d value of 1 mL/g for SRS confining units is based on a carbon soil-to-plant concentration ratio reported by Sheppard and Thibault (1990) for clay and expert judgment (Kaplan, 2006).

2.4.2 Nickel

The most common oxidation state of nickel is +2, although 0, +1, +3, and +4 nickel complexes are observed. Available nickel sorption studies indicate that sorption of nickel is strongly pH dependent and increases with increasing pH (Christensen, et al., 1996). Several studies on

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds , for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

[‡]McIntyre, P.F. "Sorption Properties of Carbon-14 on Savannah River Plant Soil." DPST–88–900. Aiken, South Carolina: E.I. du Pont de Nemours and Company, Savannah River Plant. 1998.

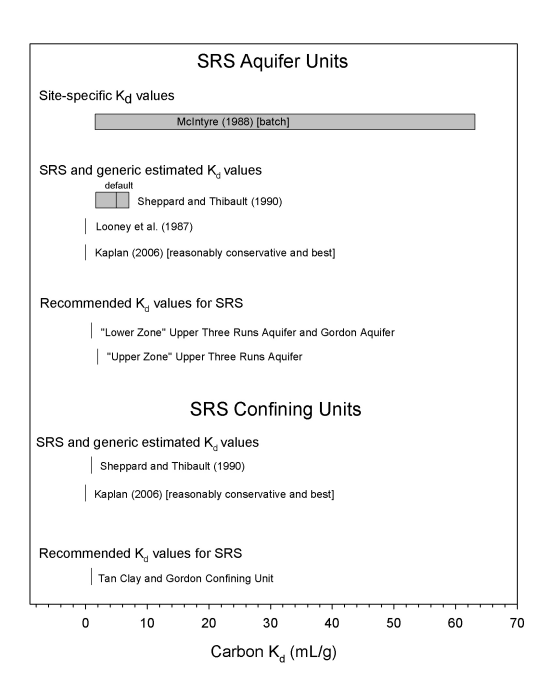


Figure 2-2. Measured and Estimated Site-Specific and Generic Carbon $K_{\rm d}$ Ranges and Values and Recommended Carbon $K_{\rm d}$ Values for Aquifer Units and Confining Units at SRS (Table 2-3). Recommended Carbon $K_{\rm d}$ Values Are Associated With High Uncertainty Due to Limited Site-Specific Sorption Data.

sorption of heavy metals onto soils indicate that nickel is among the most mobile of heavy metals (Kabata-Pendias and Pendias, 1984; Alloway, 1995).

Crapse, et al. (2004) performed sequential extractions to estimate nickel K_d values on two major soil types at the D-Area of SRS: upland subsurface soils associated with a low pH/high sulfate/metals plume downgradient of the D-Area Coal Pile Runoff Basin at the D-Area expanded operable unit (DEXOU) and surface ash material discharged to the wetland from the D-Area Ash Basin. The upland subsurface soils were collected from the lower zone of the Upper Three Runs aguifer, and the wetland samples were collected from the upper zone of the Upper Three Runs aguifer. The upland subsurface soils consisted of ash-impacted and nonash-impacted samples. Ash impacted upland soils had higher pHs and higher K₁s than nonash-impacted upland soils. Estimated nickel K_d values for ash-impacted upland subsurface soils ranged from 3,100 to 28,000 mL/g over the pH range of 7.5 to 7.9. Estimated nickel K_d values for nonash-impacted soils ranged from 0.19 to >347 mL/g over the pH range of 3.2 to 6.3. Estimated K_d values for the wetland soil ranged from 284 to 6,500 mL/g over the pH range of 4.5 to 6.1. In both soils, nickel K_d values increased with increasing pH. The increase in nickel K_d was attributed in part to the increased cation exchange capacity and increased iron-oxyhydroxide concentrations in the sediments expected with increased pH. Over the pH range expected in the subsurface at the SRS (5.2 to 7.7), nickel K_d values in the nonash-impacted upland soil ranged from 81 to >347 mL/g, and nickel K_d values in the wetland soil ranged from 284 to 6,500 mL/g.

Kaplan and Serkiz (2004) conducted batch experiments to measure nickel K_d values on a clayey sediment and a sandy sediment collected from the E-Area at SRS as a function of pH and dissolved oxygen (DO) content. The sediments were collected from the upper zone of the Upper Three Runs aquifer and represented end members with respect to contaminant sorption properties expected in the E-Area subsurface. Average nickel $K_{\rm d}$ values were reported at DO concentrations ranging from 0 to 300 mg/L [0 to 300 ppm] for pHs of 3.9, 5.3, and 6.7. Table 2-1 indicates that DO concentrations ranging from 0 to 10 mg/L [0 to 10 ppm] are characteristic of sandy aquifer and clay confining units beneath the GSA. At pH 3.9 and DO concentrations of 0 and 10 mg/L [0 to 10 ppm], Kaplan and Serkiz (2004) reported average nickel K_d values of 2.3 and 1.8 mL/g for clayey sediment and average nickel K_d values of 0.6 and 1.3 mL/g for sandy sediment. At pH 5.3 and DO concentrations of 0 and 10 mg/L [0 to 10 ppm], Kaplan and Serkiz (2004) reported average nickel K_d values of 29.5 and 65.3 mL/g for clayey sediment and average nickel K_d values of 6.8 and 7.7 mL/g for sandy sediment. At pH 6.7 and DO concentrations of 0 and 10 mg/L, Kaplan and Serkiz (2004) reported average nickel K_d values of 287 and 521 mL/g for clayey sediment and average nickel K_d values of 150 and 279 mL/g for sandy sediment.

Powell, et al. (2001) reported *in-situ* nickel K_d values for seven "matched" pore water and soil pairs collected at the D-Area expanded operable unit (DEXOU) located in the southwestern part of the SRS. The soils were collected from the upper zone of the Upper Three Runs aquifer at depths of 0.3 to 10 m [1 to 34 ft]. Field measurements of pH were in the range of 4.59 to 6.57 for all the soil collection locations. The nickel K_d values were estimated using EPA Method 3050b (EPA, 1986) and ranged from 0.63 to >310 mL/g. Over the pH range expected in the subsurface at SRS (5.2 to 7.7), Powell, et al. (2001) reported nickel K_d values ranging from 27 to >310 mL/g. EPA Method 3050b is a hot nitric extraction that likely overestimates the metals actually available for transport. Therefore, the nickel K_d values measured by Powell,

et al. (2001) (0.63 to >310 mL/g) likely underestimated actual nickel $K_{\rm d}$ values on soils at DEXOU.

Kaplan and Serkiz (2000) performed desorption measurements to measure nickel K_d values on shallow contaminated sediments from the TNX Outfall Operable Unit at the SRS. The desorption K_d values were based on extractable portions that comprised the exchangeable, organic-bound, amorphous Fe-oxide bound, and water extractable fractions. Basing the K_d estimates on these extractable fractions instead of the more easily determined total concentrations in the sediment, more conservative and technically defensible K_d values were measured. The contaminated sediments were collected at depths of 0 to 0.6 m [0 to 2 ft] by hand auger and were composed dominantly of sand and silt (i.e., sand and silt contents ranging from 80 to 94 wt%). The pH of the sediments ranged from 4.00 to 4.53. Measured nickel K_d values ranged from 257 to 404 mL/g.

Examination of K_d values listed in the compendium of default soil solid/liquid K_d values of Thibault, et al. (1990) indicates three clayey sand K_d values listed for soils from South Carolina. These values are from a study by Neiheisal (1983) and are 115, 116, and 120 mL/g. The composition and pH of solutions used in the Neiheisal (1983) study were not listed in the compendium of Thibault, et al. (1990).

Several researchers have measured nickel K_d values in sediments and soils from localities other than SRS. These studies are described next.

Christensen, et al. (1996) performed batch equilibrium experiments to measure nickel K_d values as a function of pH for sandy aquifer materials obtained at 12 locations in Denmark. Sorption experiments consisted of 10 g [0.02 lb] of aquifer material and 50 mL [1.7 oz] of 10^{-3} M CaCl₂ solution equilibrated for 48 hours. Aquifer materials consisted of >95-wt% sand-sized particles. Measured nickel K_d values ranged from 3 to 7,250 mL/g over the pH range of 4.9 to 8.87 and increased strongly with increasing pH.

Reddy and Dunn (1986) conducted batch equilibrium experiments to measure nickel K_d values on three soils as a function of nickel concentration in solution. Measured K_d values for nickel ranged from 152 to 388 mL/g. The Mecklenburg clay, with the highest CEC, gave the highest K_d values. Measured nickel K_d values increased as the concentration of nickel increased in test solutions.

Veeresh, et al. (2003) performed batch equilibrium experiments to measure nickel K_d values on three soils from different climatic regions of India. A 0.005 M $Ca(NO_3)_2 \cdot 2H_2O$ solution was used in the batch experiments. Soils had varying particle size characteristics and pHs: a silty sand soil (85-wt% sand and silt, 15-wt% clay) with a pH of 5.2, a clay-rich silty sand soil (72-wt% sand and silt, 28-wt% clay) with a pH of 7.1, and a clay-rich soil (63-wt% clay, 27-wt% sand and silt) with a pH of 8.6. K_d values determined from sorption isotherms were 9 mL/g for the silty sand soil, 24 mL/g for the clay-rich silty sand soil, and 329 mL/g for the clay-rich soil.

Sheppard and Thibault (1990) estimated a nickel default K_d for sand of 400 mL/g (11 observations ranging from 60 to 3,600 mL/g) and a nickel default K_d for clay of 650 mL/g (10 observations ranging from 305 to 2,467). Looney, et al. (1987) estimated a reference SRS soil K_d of 100 mL/g for nickel from an estimated range of 10 to 1,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" nickel K_d values of 5 and 7 mL/g for sandy

subsurface sediment and "reasonably conservative" and "best" nickel $\rm K_{\rm d}$ values of 21 and 30 mL/g for clayey subsurface sediment at SRS. The estimated nickel K_d values estimated by Kaplan (2006) were based on the nickel sorption data of Kaplan and Serkiz (2004) measured at a pH of 5.3.

Measured, estimated, and recommended nickel K_d values are summarized in Table 2-4. At pHs expected in the subsurface for SRS (5.2 to 7.7), site-specific K_d values ranging from 81 to 6,500 mL/g (Crapse, et al., 2004), 6.8 to 279 mL/g (Kaplan and Serkiz, 2004), and 27 to >310 mL/g (Powell, et al., 2001) were reported for sandy soils and sediments collected from the Upper Three Runs aguifer.

Table 2-4. Measured, Estimated, and Recommended K_d Values for Nickel in Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Nickel K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Nickel K _d (mL/g)*	Generic Estimated Nickel K _d (mL/g)†	Recommended Nickel K _d (mL/g)
Upper Three Runs Aquifer: Upper	284-6,500‡ 0.6-521§ 0.63->310¶ 257-404#	In situ Batch In situ In situ	5/7	400	10
Tan Clay Confining Unit			21/30	650	21
Upper Three Runs Aquifer: Lower	0.19–28,000‡	In situ	5/7	400	80
Gordon Confining Unit			21/30	650	150
Gordon Aquifer			5/7	400	80

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC-TR-2006-00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, Kas, for Four Major Soil

Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990. ‡Crapse, K.P., S.M. Serkiz, A. Pishko, P.C. McKinsey, R.L. Brigmon, E.P. Shine, C. Fliermans, and A.S. Knox. "Monitored Natural Attenuation of Inorganic Contaminants Treatability Study Final Report."

WSRC-TR-2004-00124. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company, Savannah River Site. 2004.

[§]Kaplan, D.I. and S.M. Serkiz. "Influence of Dissolved Organic Carbon and pH on Contaminant Sorption to Sediment." WSRC-RP-2004-00593. Aiken, South Carolina: Westinghouse Savannah River Company. 2004. ¶Powell, K.R., S. Serkiz, and D.I. Kaplan. "Evaluation of the D-Area Expanded Operable Unit for Monitored Natural Attenuation of Inorganic Constituents of Concern: Interim Report." WSRC-TR-2001-00574. Aiken, South Carolina: Westinghouse Savannah River Company. 2001.

[#]Kaplan, D.I. and S.M. Šerkiz. "In-Situ K_d Values and Geochemical Behavior for Inorganic and Organic Constituents of Concern at the TNX Outfall Delta." WSRC-TR-99-00488. Aiken, South Carolina: Westinghouse Savannah River Company. 2000.

The nickel K_d values reported by Neiheisal (1983) for soils from South Carolina (115 to 120 mL/g) and the reference SRS soil nickel K_d value of 100 mL/g estimated by Looney, et al. (1987) are within the ranges of SRS for these site-specific nickel K_d values. With the exception of nickel K_d values for wetland soil at the DEXOU reported by Crapse, et al. (2004) (284 to 6,500 mL/g), the default nickel K_d value of 400 mL/g recommended by Sheppard and Thibault (1991) lies outside or at the high end of SRS site-specific nickel K_d values reported by Kaplan and Serkiz (2000, 2004) and Powell, et al. (2001). The "reasonably conservative" and "best" nickel K_d values of 5 and 7 mL/g estimated by Kaplan (2006) for sandy subsurface sediment at SRS were based on the K_d range of 6.8 to 7.7 mL/g measured on sandy sediment at the E-Area at pH 5.3 by Kaplan and Serkiz (2004). Based on the previous observations and considering that pH in the upper zone of the Upper Three Runs aquifer may be as low as 5.2 (Table 2-2), a nickel K_d value of 10 mL/g is recommended for the upper zone of the Upper Three Runs aquifer. This value lies between the nickel K_d values reported by Kaplan and Serkiz (2004) at a pH of 5.3 (6.8 to 7.7 mL/g) and those reported by Powell, et al. (2001) at pHs expected at the SRS (27 to >310) for sandy sediments from the upper zone of the Upper Three Runs aguifer. Because nickel K_d values are highly sensitive to pH (i.e., nickel K_d increases significantly with increasing pH) and pH is higher in deeper aquifer units at SRS (Table 2-2), a nickel K_d value of 80 mL/g is recommended for the lower zone of the Upper Three Runs aquifer and the Gordon aquifer. This value is at the low end of the nickel K_d range of 81 to >347 mL/g reported for sediments from the lower zone of the Upper Three Runs aquifer by Crapse, et al. (2004) at pHs expected in the subsurface at SRS.

Kaplan and Serkiz (2004) reported nickel K_d values ranging from 29.5 to 65.3 at a pH of 5.3 and from 287 to 521 mL/g at a pH of 6.7 on clayey sediment from the E-Area of SRS. Veeresh, et al. (2003) reported a nickel K_d value of 329 mL/g for a clay-rich soil from India. Reddy and Dunn (1986) reported nickel K_d values on soils ranging from 152 to 388 mL/g and observed that soils with the highest cation exchange capacities gave the highest K_d values. Sheppard and Thibault (1990) recommend a nickel default K_d value of 650 mL/g for clay. The "reasonably conservative" and "best" nickel K_d values of 21 and 30 mL/g estimated by Kaplan (2006) for clayey sediments at SRS are based on the nickel K_d range of 29.5 to 65.3 at a pH of 5.3 reported by Kaplan and Serkiz (2004) for clayey sediment from the E-Area. Based on the previous observations and considering that pH in the tan clay confining unit may be as low as 5.2 (Table 2-2), the "reasonably conservative" nickel K_d value of 21 mL/g estimated by Kaplan (2006) for SRS sandy subsurface sediments is recommended for the tan clay. Because nickel K_d values are highly sensitive to pH (i.e., nickel K_d increases significantly with increasing pH) and pH is higher in the Gordon confining unit (Table 2.2), a nickel K_d value of 150 mL/g is recommended for the Gordon confining unit. This value lies at the approximate midpoint between the nickel K_d ranges reported by Kaplan and Serkiz (2004) on clayey subsurface sediments at SRS at pHs of 5.3 (29.5 to 65.3 mL/g) and 6.7 (287 to 521 mL/g). The position of the nickel K_d values recommended in this review with respect to measured and estimated site-specific and generic nickel K_d ranges and values is graphically displayed in Figure 2-3. This figure illustrates the uncertainty associated with the nickel K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific nickel sorption data (Crapse, et al., 2004; Kaplan and Serkiz, 2000, 2004; Powell, et al., 2001), the recommended nickel K_d values are associated with low uncertainty.

2.4.3 Selenium

Se-79 can exist in the -2, 0, +2, +4, and +6 valance states. Selenium in the +6 or selenate state (SeO₄²⁻) is stable under alkaline and oxidizing conditions. Selenium in the +4 state occurs naturally as selenite (SeO₃²⁻). Selenite tends to oxidize to the +6 state in alkaline oxidizing solutions, but not in an acid medium. Selenite binds tightly to iron and aluminum oxides. Thus, selenite is quite insoluble in soils and generally not present in waters in any appreciable amount.

Kaplan and Serkiz (2006) conducted batch experiments to measure selenium K_d values on a clayey sediment and a sandy sediment collected from the E-Area at SRS as a function of pH and DO content. The sediments were collected from the upper zone of the Upper Three Runs aquifer and represented end members with respect to contaminant sorption properties expected in the E-Area subsurface. Average selenium K_d values were reported at DO concentrations ranging from 0 to 300 mg/L [0 to 300 ppm] for pHs of 3.9, 5.3, and 6.7. Table 2-1 indicates that DO concentrations ranging from 0 to 10 mg/L [0 to 10 ppm] are characteristic of sandy aquifer and clay confining units beneath the GSA. At pH 3.9 and a DO concentration of 0 mg/L, Kaplan and Serkiz (2006) reported an average selenium K_d value of 1,041 mL/g for clayey sediment and an average selenium K_d value of 1,041 mL/g for sandy sediment. At pH 5.3 and a DO concentration of 0 mg/L [0 ppm], Kaplan and Serkiz (2006) reported an average selenium K_d value of 1,041 mL/g for clayey sediment and an average selenium K_d value of 1,311 mL/g for sandy sediment. At pH 6.7 and a DO concentration of 0 mg/L [0 ppm], Kaplan and Serkiz (2006) reported an average selenium K_d value of 1,041 mL/g for clayey sediment and an average selenium K_d value of 601 mL/g for sandy sediment. Kaplan and Serkiz (2006) noted that the sandy sediment, but not the clayey sediment, showed the characteristic decrease in K_d values as pH increased.

Powell, et al. (2001) reported *in-situ* selenium K_d values for seven "matched" pore water and soil pairs collected at DEXOU located in the southwestern part of SRS. The soils were collected from the upper zone of the Upper Three Runs aquifer at depths of 0.3 to 10 m [1 to 34 ft]. Field measurements of pH were found to be in the range of 4.59 to 6.57 for all the soil collection locations. The selenium K_d values were estimated using EPA Method 3050b and ranged from 0.7 to 270 mL/g. Over the pH range expected in the subsurface at the SRS (5.2 to 7.7), Powell, et al. (2001) reported selenium K_d values that also ranged from 0.7 to 270 mL/g. EPA Method 3050b (EPA, 1986) is a hot nitric extraction that likely overestimates the metals actually available for transport. Therefore, the selenium K_d values measured by Powell, et al. (2001) (0.7 to 270 mL/g) likely underestimated actual selenium K_d values on soils at the DEXOU.

Kaplan and Serkiz (2000) performed desorption measurements to measure selenium K_d values on shallow contaminated sediments from the TNX Outfall Operable Unit at SRS. The desorption K_d values were based on extractable portions that comprised the exchangeable, organic-bound, amorphous Fe-oxide bound, and water extractable fractions. Basing the K_d estimates on these extractable fractions instead of the more easily determined total concentrations in the sediment, more conservative and technically defensible K_d values were measured. The contaminated sediments were collected at depths of 0 to 0.6 m [0 to 2 ft] by hand auger and were composed dominantly of sand and silt (i.e., sand and silt contents ranging from 80 to 94 wt%). The pH of

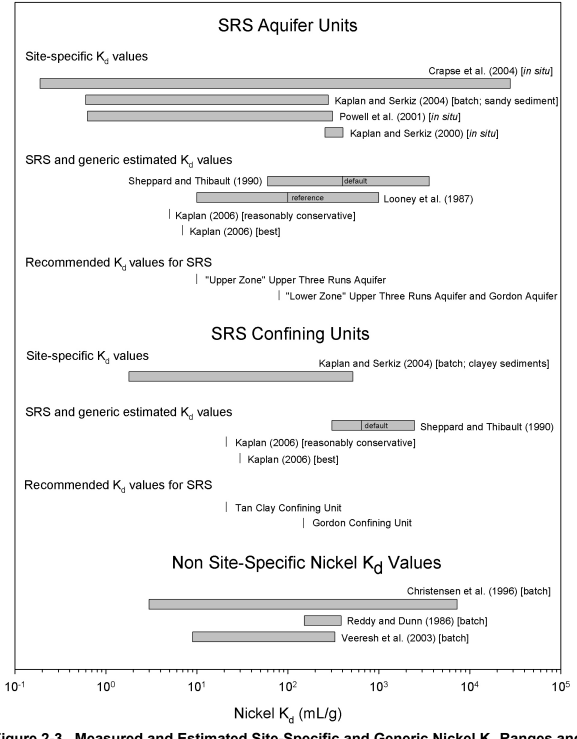


Figure 2-3. Measured and Estimated Site-Specific and Generic Nickel $K_{\rm d}$ Ranges and Values and Recommended Nickel $K_{\rm d}$ Values for Aquifer Units and Confining Units at SRS (Table 2-4). Recommended Nickel $K_{\rm d}$ Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

the sediments ranged from 4.00 to 5.78. Measured selenium K_d values ranged from 219 to 267 mL/g. Over the pH range expected in the subsurface at SRS (5.2 to 7.7), Kaplan and Serkiz (2000) reported selenium K_d values of 254 and 248 mL/g for sediments with pHs of 5.26 and 5.78, respectively.

Kaplan (2004) estimated K_d values for selenium in sandy soil and clay for use in a special analysis of the Slit and Engineered Trenches Special Analysis and the Intermediate Level Vault Special Analysis at SRS. The values estimated by Kaplan (2004) were extracted from the compendium of soil solid/liquid partition coefficients for four major soil types (Thibault, et al., 1990). Selenium K_d values for sand soil types between pH 5.3 and 6.3 were reported as 70, 70, and 36 mL/g, and generally increased as pH decreased, which is expected for anionic radionuclides such as $SeO_4^{\ 2^-}$ and $SeO_3^{\ 2^-}$. According to Kaplan (2004), selenium K_d values for clay sediments between pH 5.0 and 6.0 were reported as 76, 140, 80, 246, and 170 mL/g. Based on this tabulation, Kaplan (2004) estimated a selenium K_d value of 36 mL/g for soils and a selenium K_d value of 76 mL/g for clay for use in transport modeling.

Sheppard and Thibault (1990) specified a selenium default K_d for sand of 55 mL/g (three observations ranging from 36 to 70 mL/g) and a selenium default K_d for clay of 115 mL/g (14 observations ranging from 36 to 246 mL/g). Looney, et al. (1987) recommended a reference SRS soil K_d of 2.5 mL/g for selenium from an estimated range of 1 to 100 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" selenium K_d values of 800 and 1,000 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" selenium K_d values of 800 and 1,000 mL/g for clayey subsurface sediment at SRS. The estimations of Kaplan (2006) were based on the selenium sorption data of Kaplan and Serkiz (2006) described previously.

Measured, estimated, and recommended selenium K_d values are summarized in Table 2-5. Kaplan and Serkiz (2006) reported selenium K_d values ranging from 601 to 1,311 mL/g on sandy sediment from the E-Area of SRS over the pH range of 3.9 to 6.7. This same selenium K_{d} range was observed at pHs expected in the subsurface at SRS (pH 5.2 to 7.7). Powell, et al. (2001) reported selenium K_d values ranging from 0.7 to 270 mL/g for sediments collected from the upper zone of the Upper Three Runs aguifer at the DEXOU over the pH range of 4.59 to 6.57. Over the pH range expected in the subsurface at the SRS (5.2 to 7.7), Powell, et al. (2001) reported selenium K_d values that also ranged from 0.7 to 270 mL/g. The selenium K_d values reported by Powell, et al. (2001) were determined using EPA Method 3050b, which is an extraction method that likely underestimates actual K_d values. Kaplan and Serkiz (2000) reported selenium K_d values ranging from 219 to 267 mL/g on shallow contaminated soils from the TNX Outfall Operable Unit at the SRS over the pH range of 4.0 to 5.78. Over the pH range expected in the subsurface at SRS (5.2 to 7.7), Kaplan and Serkiz (2000) reported selenium K_d values of 254 and 248 mL/g for sediments with pHs of 5.26 and 5.78, respectively. Over the pH range of 5.3 to 6.3, Kaplan (2004) estimated a selenium K_d value of 36 mL/g for aquifer units (i.e., soils) at SRS based on an analysis of K_d data for selenium in the compilation of soil/liquid partition coefficients of Thibault, et al. (1990). The compilation of Sheppard and Thibault (1990) specified a selenium K_d of 55 mL/g for sand soils. Looney, et al. (1987) estimated a much lower

Table 2-5. Measured, Estimated, and Recommended K_d Values for Selenium in Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Selenium K _d (mL/g)	Test Method	Savannah River Site- Specific "Reasonably Conservative"/ "Best" Estimated Selenium K _d (mL/g)*	Generic Estimated Selenium K _d (mL/g)†	Recommended Selenium K _d (mL/g)
Upper Three Runs Aquifer: Upper	601–1,311‡ 0.7–>270§ 219–267¶	Batch In situ In situ	800/1,000	55	425
Tan Clay Confining Unit	_	_	800/1,000	115	800
Upper Three Runs Aquifer: Lower	_	_	800/1,000	55	425
Gordon Confining Unit	_	_	800/1,000	115	800
Gordon Aquifer	_	_	800/1,000	55	425

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC-TR-2006-00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

selenium Kd value of 2.5 mL/g for SRS soil. Based on the selenium sorption data of Kaplan and Serkiz (2006), Kaplan (2006) estimated a "reasonably conservative" selenium K_d of 800 mL/g for sandy sediments at SRS. Based on the previous observations, a selenium K_d value of 425 mL/g is recommended for aquifer units at SRS. This value lies between the measured selenium K_d value of 248 mL/g reported by Kaplan and Serkiz (2000) at a pH of 5.78 and the

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_d s, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

[‡]Kaplan, D.I. and S.M. Serkiz. "Influence of Dissolved Organic Carbon and pH on Anion Sorption to Sediment." WSRC–RP–2006–00005. Aiken, South Carolina: Westinghouse Savannah River Company. 2006. §Powell, K.R., S. Serkiz, and D.I. Kaplan. "Evaluation of the D-Area Expanded Operable Unit for Monitored Natural Attenuation of Inorganic Constituents of Concern: Interim Report." WSRC–TR–2001–00574. Aiken, South Carolina: Westinghouse Savannah River Company. 2001.

[¶]Kaplan, D.I. and S.M. Šerkiz. *"In-Situ* K_d Values and Geochemical Behavior for Inorganic and Organic Constituents of Concern at the TNX Outfall Delta." WSRC–TR–99–00488. Aiken, South Carolina: Westinghouse Savannah River Company. 2000.

measured selenium K_d value of 601 mL/g reported by Kaplan and Serkiz (2006) for sandy subsurface sediment at SRS at a pH of 6.7.

Kaplan and Serkiz (2006) reported a selenium K_d value of 1,041 mL/g on clayey sediment from the E Area of SRS over the pH range of 3.9 to 6.7. Over the pH range of 5.0 to 6.0, Kaplan (2004) estimated a selenium K_d value of 76 mL/g for clay units at SRS. This recommendation was based on an analysis of K_d data for selenium in the compilation of soil/liquid partition coefficients of Thibault, et al. (1990). The compilation of Sheppard and Thibault (1990) (also Thibault, et al., 1990) recommended a selenium K_d of 115 mL/g for clay soils. Based on the site-specific selenium sorption data of Kaplan and Serkiz (2006), Kaplan (2006) estimated a "reasonably conservative" selenium K_d of 800 mL/g for clayey sediments at SRS, and this value is recommended for clay confining units at SRS.

The position of the selenium K_d values recommended in this review with respect to measured and estimated site-specific and generic selenium K_d ranges and values is graphically displayed in Figure 2-4. This figure illustrates the uncertainty associated with the selenium K_d values recommended for SRS aquifer units and clay confining units. Due to the wide range in measured site-specific selenium K_d values, the recommended selenium K_d values are associated with high uncertainty. For example, Kaplan and Serkiz (2006) reported selenium K_d values ranging from 601 to 1,311 mL/g over the pH range of 3.9 to 7.6 for sandy sediment from the E-Area, whereas Powell, et al. (2001) reported selenium K_d values ranging from 219 to 267 mL/g over the pH range of 4.0 to 5.78 for soil samples from the D-Area.

2.4.4 Strontium

Sr-90 exists primarily as the uncomplexed Sr^{2+} ion over the pH range of natural waters (pH 3 to 10). Sr^{2+} adsorbs to both clay and carbonate minerals. For clays, the extent to which strontium partitions from the aqueous phase to the solid phase is expected to be controlled primarily by the CEC of the solid phase. Other factors affecting strontium K_d values are pH and ionic strength. Strontium K_d values have been reported to increase with increasing pH (Prout, 1958; Rhodes, 1957). This trend is likely the result of hydrogen ions competing with Sr^{2+} for exchange sites. Elevated concentrations of major cations (e.g., Na^+ , K^+ , Ca^{2+} , Mg^{2+}) can greatly reduce strontium K_d values (Bunde, et al., 1998, 1997). Strontium K_d values may decrease from 100 to 200 mL/g in low ionic strength solutions to less than 5 mL/g in high ionic strength solutions (Routson, et al., 1980). There is little tendency for strontium to form complexes with inorganic or organic ligands (Faure and Powell, 1972; Stevenson and Fitch, 1986).

Hoeffner (1985) summarized strontium sorption studies conducted to determine the performance of the low-level waste burial ground at SRS. The majority of studies used a Savannah River Plant soil taken from depths 2.5–3.0 m [8–10 ft] below grade and a low ionic strength groundwater representative of the burial ground (pH 5.0). Hoeffner (1985) reported an expected range of strontium K_d values from 1.5 to 3,000 mL/g over a pH range of 3.4 to 7.3, and for a groundwater of typical composition at the SRS burial ground (pH 4.7), a K_d value of 7.5 mL/g is expected.

Ryan (1982) performed batch strontium sorption experiments using SRS burial ground core samples and groundwater. The measured strontium K_d values from the experiments ranged from 1.1 to 10.9 mL/g.

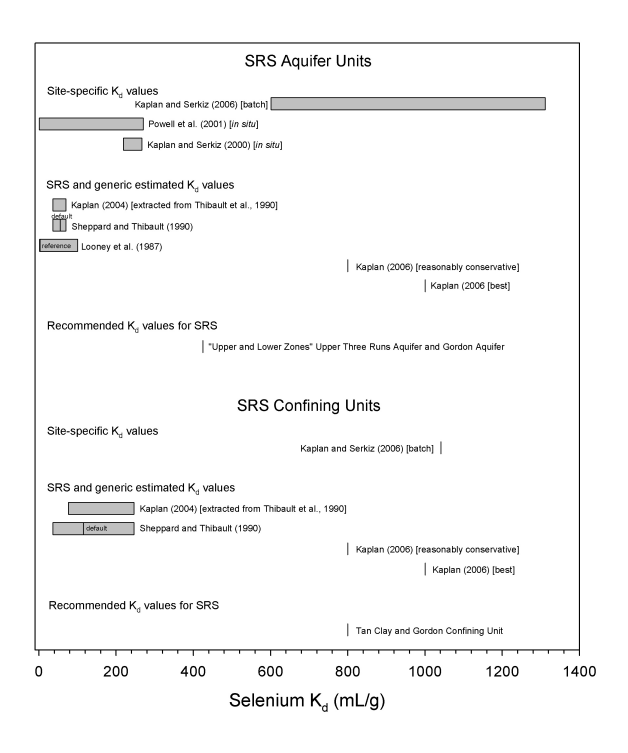


Figure 2-4. Measured and Estimated Site-Specific and Generic Selenium $K_{\rm d}$ Ranges and Values and Recommended Selenium $K_{\rm d}$ Values for Aquifer Units and Confining Units at SRS (Table 2-5). Recommended Values Are Associated With High Uncertainty Due to the Wide Range in Measured Site-Specific Selenium $K_{\rm d}$ Values.

McConnell, et al. (1995) reported strontium K_d values ranging from 20 to 40 ml/g for SRS soil used in lysimeter tests to obtain information on the performance of radioactive wasteforms and DO. The sediments were collected from the upper zone of the Upper Three Runs aquifer These values were obtained using a batch method. The pH and composition of the solution used in the batch test was not reported.

Kaplan and Serkiz (2004) conducted batch experiments to measure strontium $\rm K_d$ values on a clayey sediment and a sandy sediment collected from the E-Area at SRS as a function of pH and DO content. The sediments were collected from the upper zone of the Upper Three Runs aquifer and represented end members with respect to contaminant sorption properties expected in the E-Area subsurface. Average strontium $\rm K_d$ values were reported at DO concentrations ranging from 0 to 300 mg/L [0 to 300 ppm] for pHs of 5.3 and 6.7. Table 2-1 indicates that DO concentrations ranging from 0 to 10 mg/L [0 to 10 ppm] are characteristic of sandy aquifer and clay confining units beneath the GSA. At pH 5.3 and DO concentrations of 0 to 10 mg/L [0 to 10 ppm], Kaplan and Serkiz (2004) reported average strontium $\rm K_d$ values of 16.6 and 42.6 mL/g for clayey sediment and average strontium $\rm K_d$ values of 4.5 and 5.5 mL/g for sandy sediment. At pH 6.7 and DO concentrations of 0 to 10 mg/L [0 to 10 ppm], Kaplan and Serkiz (2004) reported average strontium $\rm K_d$ values of 97.4 and 67.7 mL/g for clayey sediment and average strontium $\rm K_d$ values of 46.9 and 60.4 mL/g for sandy sediment.

Kaplan and Serkiz (2000) performed desorption measurements to measure strontium K_d values on shallow contaminated sediments from the TNX Outfall Operable Unit at SRS. The desorption K_d values were based on extractable portions that comprised the exchangeable, organic-bound, amorphous Fe-oxide bound, and water extractable fractions. Basing the K_d estimates on these extractable fractions instead of the more easily determined total concentrations in the sediment, more conservative and technically defensible K_d values were measured. The contaminated sediments were collected at depths of 0 to 0.6 m [0 to 2 ft] by hand auger and were composed dominantly of sand and silt (i.e., sand and silt contents ranging from 80 to 94 wt%). The pH of the sediments ranged from 4.00 to 5.78. Measured strontium K_d values ranged from 1,676 to 5,518 mL/g. Over the pH range expected in the subsurface at the SRS (5.2 to 7.7), Kaplan and Serkiz (2000) reported strontium K_d values of 1,676 and 1,886 mL/g for sediments with pHs of 5.26 and 5.78, respectively.

Sheppard and Thibault (1990) recommended a strontium default K_d for sand of 15 mL/g (81 observations ranging from 0.05 to 190 mL/g) and a strontium default K_d for clay of 110 mL/g (24 observations ranging from 3.6 to 32,000 mL/g). Looney, et al. (1987) specified a reference SRS soil K_d of 8.0 mL/g for strontium from an estimated range of 1 to 1,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" strontium K_d values of 4 and 5 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" strontium K_d values of 12 and 17 mL/g for clayey subsurface sediment at SRS. The estimations of Kaplan (2006) were based on the strontium sorption data of Kaplan and Serkiz (2004) measured at a pH of 5.3.

Measured, estimated, and recommended strontium K_d values are summarized in Table 2-6. Over the pH range of 3.4 to 7.3, Hoeffner (1985) reported an expected range of strontium K_d values from 1.5 to 3,000 mL/g. These values are based on a summary of strontium sorption studies of shallow sediments at the SRS burial ground. For a groundwater of typical composition at the SRS burial ground (pH 4.7), Hoeffner (1985) reports an expected strontium K_d value of 7.5 mL/g. This expected value is in agreement with strontium K_d values measured by Ryan (1982) using shallow soil samples and groundwater from the SRS burial ground (1.1 to

Table 2-6. Measured, Estimated, and Recommended K_d Values for Strontium in Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Strontium K _d (mL/g)	Test Method	Savannah River Site- Specific "Reasonably Conservative"/ "Best" Estimated Strontium K _d (mL/g)*	Generic Estimated Strontium K _d (mL/g)†	Recommended Strontium K _d (mL/g)
Upper Three Runs Aquifer: Upper	1.5–3,000‡ 1.1–10.9§ 20–40¶ 4.5–97.4# 1,676–5,518**	Batch Batch Batch Batch In situ	4/5	15	5
Tan Clay Confining Unit			12/17	110	12
Upper Three Runs Aquifer: Lower			4/5	15	25
Gordon Confining Unit			12/17	110	50
Gordon Aquifer			4/5	15	25

*Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

†Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

‡Hoeffner, S.L. "Radionuclide Sorption on Savannah River Plant Burial Ground Soil—A Summary and Interpretation of Laboratory Data." DP–1702. Aiken, South Carolina: E.I. du Pont de Nemours and Company, Savannah River Laboratory. 1985.

§Ryan, J.P. "Batch and Column Strontium Distribution Coefficients with Water-Saturated Soil Strata from the Savannah River Plant Burial Ground." Environmental Migration of Long-Lived Radionuclides. International Symposium Proceedings, Knoxville, Tennessee. Vienna, Austria: International Atomic Energy Agency. 1982. ¶McConnell, Jr., J.W., R.D. Rogers, J.D. Jastrow, W.E. Sanford, S.R. Cline, T.M. Sullivan, and M. Fuhrmann. "Results after Ten Years of Field Testing Low-level Radioactive Waste Forms Using Lysimeters." ANL/ER/CP 95275. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1995.

#Kaplan, D.I. and S.M. Serkiz. "Influence of Dissolved Organic Carbon and pH on Contaminant Sorption to Sediment." WSRC–RP–2004–00593. Aiken, South Carolina: Westinghouse Savannah River Company. 2004. **Kaplan, D.I. and S.M. Serkiz. "In-Situ K_d Values and Geochemical Behavior for Inorganic and Organic Constituents of Concern at the TNX Outfall Delta." WSRC–TR–99–00488. Aiken, South Carolina: Westinghouse

Savannah River Company. 2000.

10.9 mL/g). The strontium $\rm K_d$ values measured by McConnell, et al. (1995) on SRS soil (20 to 40 mL/g) are slightly higher than those measured by Ryan (1982). The pH of the solution used by McConnell, et al. (1995) was not reported but may have had a higher pH than the burial ground groundwater used in the Ryan (1982) study. Kaplan and Serkiz (2004) reported strontium $\rm K_d$ values on sandy sediment from the E-Area of SRS that ranged from 4.5 to

5.5 mL/g at a pH of 5.3 and from 46.9 to 60.4 mL/g at a pH of 6.7. Kaplan and Serkiz (2000) reported strontium K_d values of 1,676 and 1,886 mL/g on contaminated sediments from the SRS with pHs of 5.26 and 5.78, respectively. With the exception of the strontium K_d values reported by Kaplan and Serkiz (2000), the reported site-specific strontium K_d values correspond well to the recommended strontium K_d value for sand of Sheppard and Thibault (1990) (15 mL/g) and to the strontium K_d value of 8 mL/g estimated by Looney, et al. (1987) for SRS soil. The "reasonably conservative" and "best" strontium K_d values of 4 and 5 mL/g estimated by Kaplan (2006) for sandy subsurface sediment at SRS were based on the K_d range of 4.5 to 5.5 mL/g measured on sandy sediment at the E-Area at pH 5.3 by Kaplan and Serkiz (2004). Based on the above observations, a strontium K_d value of 5 mL/g is recommended for the upper zone of the Upper Three Runs aguifer at SRS. This value is the median of the strontium K_d values measured by Kaplan and Serkiz (2004) on sandy subsurface sediments at SRS at a pH of 5.3 (4.5 to 5.5 mL/g), which corresponds to the low end of the pH range expected at SRS (pH of 5.2 to 6.0). The lower zone of the Upper Three Runs aquifer and the Gordon aquifer have reported pH values above 6.0 and 5.7, respectively. Because strontium K_d values are sensitive to pH (i.e., strontium K_d increases with increasing pH), a strontium K_d value of 25 mL/g is recommended for the lower zone of the Upper Three Runs aquifer and the Gordon aquifer. This value lies at the approximate midpoint between the strontium K_d ranges reported by Kaplan and Serkiz (2004) on sandy subsurface sediments at SRS at pHs of 5.3 (4.5 to 5.5 mL/g) and 6.7 (46.9 to 60.4 mL/g).

Kaplan and Serkiz (2004) reported strontium K_d values on clayey sediment from the E-Area of SRS that ranged from 16.6 to 42.6 mL/g at a pH of 5.3 and from 67.7 to 97.6 mL/g at a pH of 6.7. These measured values are lower than the default strontium K_d value of 110 mL/g estimated by Sheppard and Thibault (1990) for clay. Based on the strontium K_d values reported by Kaplan and Serkiz (2004) at a pH of 5.3, Kaplan (2006) estimated a "reasonably conservative" strontium K_d value of 12 for clayey sediments at SRS. Based on the above observations and considering that pH in the tan clay confining unit may be as low as 5.2 (Table 2-2), the "reasonably conservative" strontium K_d value of 12 mL/g estimated by Kaplan (2006) for SRS sandy subsurface sediments is recommended for the tan clay. Because strontium K_d values are sensitive to pH (i.e., strontium K_d increases with increasing pH) and pH is higher in the Gordon confining unit (Table 2-2) (pH 5.8 to 6.8), a strontium K_d value of 50 mL/g is recommended for the Gordon confining unit. This value lies at the approximate midpoint between the strontium K_d ranges reported by Kaplan and Serkiz (2004) on clayey subsurface sediments at SRS at pHs of 5.3 (16.6 to 42.6 mL/g) and 6.7 (67.7 to 97.6 mL/g).

The position of the strontium K_d values recommended in this review with respect to measured and estimated site-specific and generic strontium K_d ranges and values is graphically displayed in Figure 2-5. This figure illustrates the uncertainty associated with the strontium K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific strontium sorption data (Hoeffner, 1985; Kaplan and Serkiz, 2004, 2000), the recommended strontium K_d values are associated with low uncertainty.

2.4.5 Niobium

Niobium is present in nuclear waste as a result of the activation of zirconium pellets associated with nuclear fuel. Radioactive niobium is a metal that is present in radioactive wastes as Nb-95 (half-life of 35 days), Nb-93 (half-life of 13.6 years), and Nb-94 (half-life of 20,300 years). Knowledge of the behavior of niobium in the environment is limited. Niobium in soils is reported

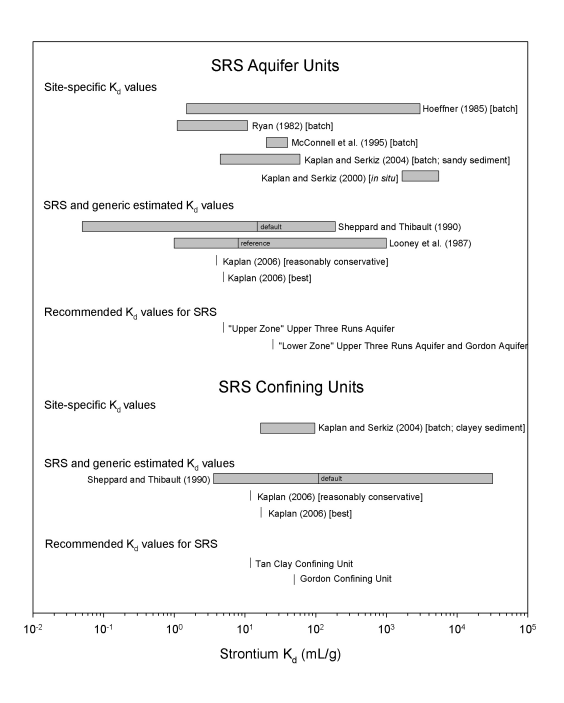


Figure 2-5. Measured and Estimated Site-Specific and Generic Strontium K_d Ranges and Values and Recommended Strontium K_d Values for Aquifer Units and Confining Units at SRS (Table 2-6). Recommended Strontium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

to be rather immobile (Rhodes, 1957), being present as insoluble Nb(V) oxides and hydroxides. Nb(OH) $_{\rm 5}$ is expected to be the predominant form according to its hydrolysis properties in the pH and Eh ranges found in most soils (Baes and Mesmer, 1976). No site-specific sorption data for niobium on subsurface units at SRS were found in the literature. Rhodes (1957) performed a batch equilibrium experiment to measure niobium $K_{\rm d}$ values on a calcareous sandy soil. Rhodes (1957) reported a $K_{\rm d}$ value of >1,980 mL/g for combined zirconium and niobium. The extent to which this value is valid for niobium only was not specified due to analytical difficulties.

Echevarria, et al. (2005) performed batch experiments to study niobium sorption on three soils from northeastern France. Soils consisted of one Luvisol (a sand-rich soil) and two Cambisols (clay-rich soils). Batch experiments consisted of soil–liquid ratios of 1 to 10 using deionized water. Measured niobium K_d values after 3 days were 1,978 mL/g for the Luvisol and 3,228 and 2,512 mL/g for the Cambisols. These values continued to increase after 3 days. Extrapolation of isotherm data suggested that after 30 days, niobium K_d values would have reached 4,524 mL/g for the Luvisol and 8,370 and 4,729 mL/g for the Cambisols.

Based on soil-to-plant concentration ratios, Sheppard and Thibault (1990) estimated a niobium default K_d for sand of 160 mL/g and a niobium default K_d for clay of 900 mL/g. A reference soil K_d for niobium at SRS was not included in the compilation of Looney, et al. (1987). Kaplan (2006) estimated "best" and "reasonably conservative" K_d values of 0 mL/g for niobium in both sandy subsurface sediment and clayey subsurface sediment at SRS. The niobium K_d values estimated by Kaplan (2006) were not based on site-specific data for niobium sorption on SRS sediments.

Recommended and estimated niobium K_d values are summarized in Table 2-7. Sheppard and Thibault (1990) recommended a niobium default K_d for sand of 160 mL/g. This recommended value is much lower than the measured niobium K_d value of a sand-rich soil from northeastern France (1,978 mL/g after 3 days of equilibrium) (Echevarria, et al., 2005). A niobium K_d value of 160 mL/g is, therefore, considered reasonably conservative and is recommended for aquifer units at SRS.

Sheppard and Thibault (1990) recommended a niobium default K_d for clay of 900 mL/g; this value is much lower than the measured niobium K_d values of two clay-rich soils from northeastern France (3,228 and 2,512 mL/g after 3 days of equilibrium) (Echevarria, et al., 2005). The niobium K_d value of 900 mL/g recommended by Sheppard and Thibault (1990) is, therefore, considered reasonably conservative and is recommended for clay confining units at SRS.

The position of the niobium K_d values recommended in this review with respect to estimated site-specific and generic niobium K_d ranges and values is graphically displayed in Figure 2-6. This figure illustrates the uncertainty associated with the niobium K_d values recommended for SRS aquifer units and clay confining units. The recommended niobium K_d values are associated with high uncertainty due to the lack of site-specific sorption data.

2.4.6 Technetium

Tc-99 occurs in the oxidized (+7) state, predominantly as pertechnetate (TcO₄⁻), over the complete pH range of natural waters, generally sorbs poorly to sediments, and is typically highly mobile (e.g., Denys, et al. 2003; Kaplan, 2003; Mashkin and Shikov 2000). The sorption

Table 2-7. Recommended K_d Values for Niobium in Aquifer Sediments and Clay
Confining Units at the Savannah River Site

	Measured Niobium K _d (mL/g)	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Niobium K _d (mL/g)*	Generic Estimated Niobium K _d (mL/g)†	Recommended Niobium K _d (mL/g)
Upper Three Runs Aquifer: Upper		0/0	160	160
Tan Clay Confining Unit		0/0	900	900
Upper Three Runs Aquifer: Lower	_	0/0	160	160
Gordon Confining Unit	_	0/0	900	900
Gordon Aquifer	_	0/0	160	160

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

behavior of technetium is highly redox sensitive. Experimental data indicate that technetium mobility increases with increasing pH (Denys, et al., 2003) and decreases under organic-rich, reducing conditions (Mashkin and Shidov, 2000). The pH effect may arise from pH-dependent changes in surface charge, or may reflect the role of organic matter, which both provide anion sorption sites for TcO_4^- and reduce pH (Kaplan, 2003).

Hoeffner (1985) summarized radionuclide sorption studies conducted to determine the performance of the low-level waste burial ground at the Savannah River Plant. A technetium K_d value of 0.16 mL/g is reported for a batch experiment using a Savannah River Plant soil and a low ionic strength groundwater representative of the SRS burial ground (pH 5.0). Hoeffner (1985) also reported the results of a technetium sorption study on soils with varying clay contents. Measured technetium K_d values for sand-rich soils (clay contents of 10 to 11 wt%) ranged from 0.1 to 0.23 mL/g, while technetium K_d values for clay-rich soils (clay contents of 30 to 45 wt%) ranged from 0.33 to 1.32 mL/g.

Kaplan (2003) measured K_d values for technetium as a function of pH on two sediments from SRS. Over the pH range of 2 to 9, technetium K_d values ranged from -0.1 to 0.1 mL/g for a silty-sand upland SRS sediment (0.3 wt% Fe₂O₃, <0.02 wt% organic carbon). Over the pH

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

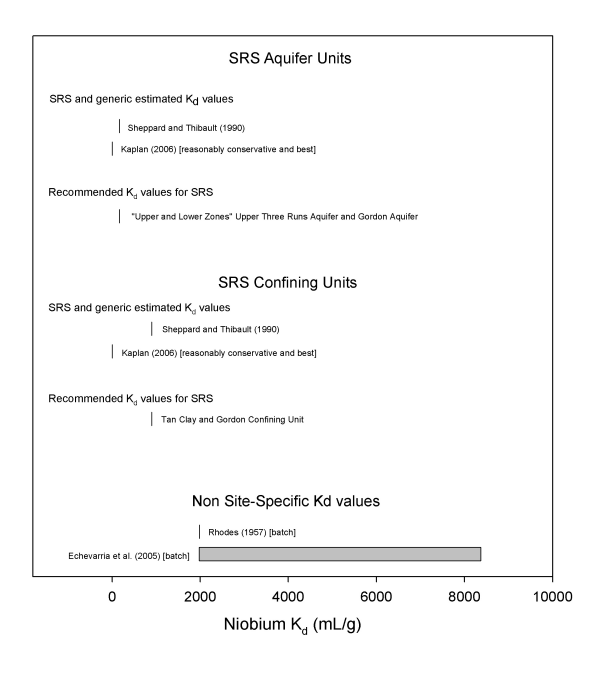


Figure 2-6. Measured and Estimated Site-Specific and Generic Niobium $K_{\rm d}$ Ranges and Values and Recommended Niobium $K_{\rm d}$ Values for Aquifer Units and Confining Units at SRS (Table 2-7). Recommended Niobium $K_{\rm d}$ Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

range of 3 to 7, technetium K_d values ranged from 0 to 0.3 mL/g for a silty-sand wetland SRS sediment (0.01 wt% Fe_2O_3 , 0.14 wt% organic carbon). For both sediments, K_d tended to decrease with increasing pH, as expected for an anionic species. Kaplan suggested that the negative K_d values measured for technetium were due to anion exclusion effects.

Routson, et al. (1977) used batch equilibrium experiments to measure K_d values for technetium on soil from South Carolina as a function of the concentration of dissolved sodium bicarbonate. The location where the soil sample was taken was not revealed, but it could have been collected from SRS. The soil contained <0.2 mg/g [200 ppm] $CaCO_3$, 3.6-wt% silt, 37.2-wt% clay, and had a CEC of 2.5 meq/100 g and a pH of 5.1. The measurements indicated that technetium did not sorb to the soil sample. The measured K_d values for technetium were 0.02, -0.05, -0.033, and 0.010 mL/g for 0.002, 0.008, 0.020, and 0.200 M NaHCO₃ solutions, respectively.

Sheppard and Thibault (1990) estimated a technetium default K_d for sand of 0.1 mL/g (19 observations ranging from 0.01 to 16 mL/g) and a technetium default K_d for clay of 1.0 mL/g (4 observations ranging from 1.16 to 1.32 mL/g). Looney, et al. (1987) estimated a reference SRS soil K_d of 0.001 mL/g for technetium from a specified range of 0.001 to 100 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" technetium K_d values of 0.05 and 0.1 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" technetium K_d values of 0.1 and 0.2 mL/g for clayey subsurface sediment at SRS. The technetium K_d values estimated by Kaplan (2006) were based on the technetium sorption data of Hoeffner (1985) and Kaplan (2003) described previously.

Measured, estimated, and recommended technetium K_d values are summarized in Table 2-8. Hoeffner (1985) reports a technetium K_d value of 0.16 mL/g for a shallow soil collected from the SRS burial ground and technetium K_d values ranging from 0.1 to 0.23 mL/g for shallow sand-rich soils from SRS. Kaplan (2003) measured technetium K_d values for an upland soil and wetland soil at SRS over pH ranges of 2 to 9 and 3 to 7, respectively. For the upland soil, technetium K_d values ranged from −0.1 to 0.1, and for the wetland soil, technetium K_d values ranged from 0 to 0.3. These values are in agreement with the technetium K_d values reported by Hoeffner (1985). Routson, et al. (1977) reported measured technetium K_d values ranging from −0.05 to 0.02 for soils from South Carolina. The technetium K_d value for sand estimated by Sheppard and Thibault (1990) (0.1 mL/g) and the reference technetium K_d value for SRS soil estimated by Looney, et al. (1987) (0.001 mL/g) are within the technetium K_d values measured at SRS (Hoeffner, 1985; Kaplan, 2003). Based on the technetium K_d values reported by Hoeffner (1985) and Kaplan (2003), Kaplan (2006) estimated a "reasonably conservative" technetium K_d value of 0.05 mL/g for sandy subsurface sediments at SRS. An examination of the technetium sorption data of Kaplan (2003) indicates that over the pH range expected at SRS (5.2 to 7.7), technetium K_d values in the upland and wetland sediments were at or below 0. The mineral composition of the aguifer units at SRS is not significantly different and, thus, these units would be expected to exhibit similar technetium sorption. Based on the above analysis, a technetium K_d value of 0 is recommended for aquifer units at SRS.

Hoeffner (1985) reported technetium K_d values ranging from 0.33 to 1.32 mL/g for clay-rich soils from SRS. The hydrostratigraphic unit from which these soils were collected was not specified. However, the reported clay contents of these soils (30 to 45 wt%) are comparable to clay contents found in the tan clay and Gordon confining units. Therefore, it is assumed that these technetium K_d values are, at least, analogous to values for the tan clay and the Gordon

Table 2-8. Measured, Estimated, and Recommended Technetium K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Technetium K _d (mL/g)	Test Method	Savannah River Site- Specific "Reasonably Conservative"/ "Best" Estimated Technetium K _d (mL/g)*	Generic Estimated Technetium K _d (mL/g)†	Recommended Technetium K _d (mL/g)
Upper Three Runs Aquifer: Upper	0.1-0.23‡ -0.1-0.3§ -0.05-0.02¶	Batch Batch Batch	0.05/0.1	0.1	0
Tan Clay Confining Unit	0.33–1.32‡	Batch	0.1/0.2	1	0.3
Upper Three Runs Aquifer: Lower	_		0.05/0.1	0.1	0
Gordon Confining Unit	_		0.1/0.2	1	0.3
Gordon Aquifer	_		0.05/0.1	0.1	0

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

§Kaplan, D.I. "Influence of Surface Charge of an Fe-Oxide and an Organic Matter Dominated Soil on Iodide and Pertechnetate Sorption." *Radiochimica Acta.* Vol. 91, No. 3. pp. 173–178. 2003.

¶Routson, R.C., G. Jansen, and A.V. Robinson. "²⁴¹Am, ²³⁷Np, and ⁹⁹Tc Sorption on Two United States Subsoils From Differing Weathering Intensity Areas." *Health Physics*. Vol. 33. pp. 311–317. 1977.

confining units. The technetium K_d value for clay estimated by Sheppard and Thibault (1990) (1.0 mL/g) is within the range of values for the clay-rich soils reported by Hoeffner (1985). Based on the technetium K_d values reported by Hoeffner (1985) and Kaplan (2003), Kaplan (2006) estimated "reasonably conservative" and "best" technetium K_d values of 0.1 and 0.2 mL/g, respectively, for clayey subsurface sediments at SRS. However, the technetium K_d values of -0.1 to 0.3 mL/g reported by Kaplan (2003) were measured on silty-sand sediments. Therefore, a technetium K_d value of 0.3 mL/g is recommended for clay confining units at SRS.

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_d s, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

[‡]Hoeffner, S.L. "Radionuclide Sorption on Savannah River Plant Burial Ground Soil—A Summary and Interpretation of Laboratory Data." DP–1702. Aiken, South Carolina: E.I. du Pont de Nemours and Company, Savannah River Laboratory. 1985.

This value is at the low end of the range of technetium K_d values reported by Hoeffner (1985) (0.33 to 1.32 mL/g) for clay-rich soils and at the high end of the range of technetium K_d values reported by Kaplan (2003) (-0.1 to 0.3 mL/g) for silty-sand sediments.

The position of the technetium K_d values recommended in this review with respect to measured and estimated site-specific and generic technetium K_d ranges and values is graphically displayed in Figure 2-7. This figure illustrates the uncertainty associated with the technetium K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific technetium sorption data (Hoeffner, 1985; Kaplan, 2003), the recommended technetium K_d values are associated with low uncertainty.

2.4.7 Tin

The sorptive properties of Sn-121 are poorly known. Sn(IV) is expected to be the predominant tin species in natural waters. Limited evidence seems to indicate that $Sn(OH)_4^0$ is a major dissolved species controlling the solubility of $SnO_2(s)$ in the pH range 2 to 10 (Kedziorek, et al., 2007).

No site-specific tin sorption data are available for SRS. Kedziorek, et al. (2007) performed batch experiments in the pH range of 2 to 12 to study the solubility of Sn(IV) and its adsorption by solids under consideration for radioactive waste storage (MX-80 bentonite and Callovo-Oxfordian argillite from the Bure site in Eastern France). Measured tin K_d values ranged from 725 to 12,000 mL/g. Kedziorek, et al. (2007) indicated that tin sorption was independent of pH and was most likely controlled by clay minerals.

Sheppard and Thibault (1990) specified a tin default K_d for sand of 130 mL/g and for clay of 670 mL/g; both values were based on a soil-to-plant concentration ratio. Looney, et al. (1987) estimated a reference SRS soil K_d of 100 mL/g for tin. Kaplan (2006) estimated "reasonably conservative" and "best" tin K_d values of 500 and 2,000 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" tin K_d values of 2,500 and 5,000 mL/g for clayey subsurface sediment at SRS. The tin K_d values estimated by Kaplan (2006) were based partly on reported lead K_d values measured on SRS burial ground sediment by Bibler and Marson (1992). Although lead is a divalent metal and tin exists dominantly in the tetravalent state [(Sn(IV)], they were considered to be chemically analogous and to have a similar sorptive behavior by Kaplan (2006). Other than being in Group IVA of the periodic chart, Kaplan (2006) gave no other technical basis for the similar chemical behavior of tin and lead. Bibler and Marson (1992) reported lead K_d values ranging from 63 to >2 million mL/g. However, it is likely that these experiments were conducted at concentrations above the solubility limit of lead {e.g., initial Pb²+ of 5 mg/L [5 ppm]} and therefore reflect precipitation more than adsorption.

Recommended and estimated tin K_d values are summarized in Table 2-9. The tin K_d value for sand estimated by Sheppard and Thibault (1990) (130 mL/g) is in good agreement with the tin K_d value for SRS soil specified by Looney, et al. (1987) (100 mL/g). The tin K_d value for sand estimated by Sheppard and Thibault (1990) was based on a soil-to-plant concentration ratio, and the tin K_d value for soil estimated by Looney, et al. (1987) was based on nonsite-specific literature data. Based on reported lead K_d values measured on SRS burial ground sediment by Bibler and Marson (1992), Kaplan (2006) estimated a "reasonably conservative" tin K_d value of 500 mL/g for sandy subsurface sediments at SRS. Lead and tin were considered chemically analogous and to have a similar sorptive behavior. Lacking site-specific tin sorption data, the

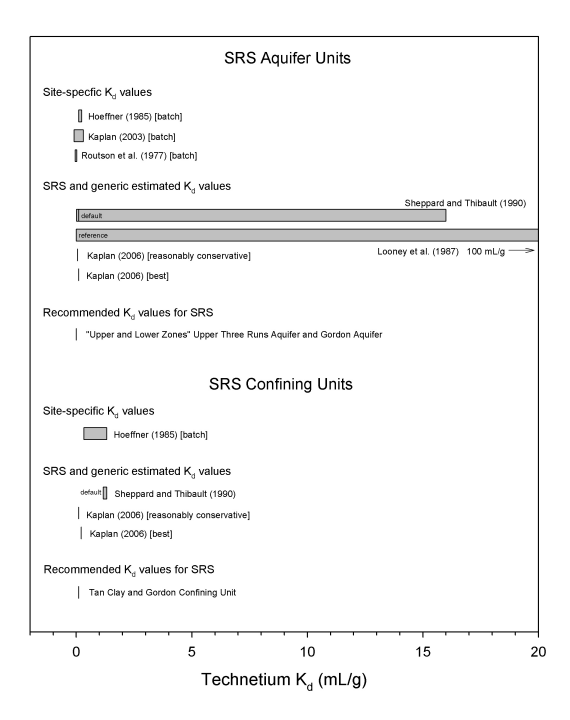


Figure 2-7. Measured and Estimated Site-Specific and Generic Technetium K_d Ranges and Values and Recommended Technetium K_d Values for Aquifer Units and Confining Units at SRS (Table 2-8). Recommended Technetium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

Table 2-9. Estimated and Recommended Tin K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Tin K _d (mL/g)	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Tin K _d (mL/g)*	Generic Estimated Tin K _d (mL/g)†	Recommended Tin K _d (mL/g)
Upper Three Runs Aquifer: Upper	ı	500/2,000	130	500
Tan Clay Confining Unit	_	2,500/5,000	670	2,500
Upper Three Runs Aquifer: Lower	_	500/2,000	130	500
Gordon Confining Unit	1	2,500/5,000	670	2,500
Gordon Aquifer	_	500/2,000	130	500

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

†Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_d s, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

"reasonably conservative" tin K_d value of 500 mL/g estimated by Kaplan (2006) is recommended for aquifer units at SRS. Sheppard and Thibault (1990) estimated a tin K_d value for clay of 670 mL/g based on a soil-to-plant concentration ratio. Based on reported lead K_d values measured on SRS burial ground sediment by Bibler and Marson (1992), Kaplan (2006) estimated a "reasonably conservative" tin K_d value of 2,500 mL/g for clayey subsurface sediments at SRS. The conservative tin K_d value of 2,500 mL/g estimated by Kaplan (2006) is near the low end of the tin K_d range measured on clay-rich materials by Kedziorek, et al. (2007) (725 to 12,000 mL/g). Therefore, the "reasonably conservative" tin K_d value of 2,500 mL/g estimated by Kaplan (2006) is recommended for clay confining units at SRS.

The position of the tin K_d values recommended in this review with respect to estimated site-specific and generic tin K_d ranges and values is graphically displayed in Figure 2-8. This figure illustrates the uncertainty associated with the tin K_d values recommended for SRS aquifer units and clay confining units. The recommended tin K_d values are associated with high uncertainty due to the lack of site-specific sorption data.

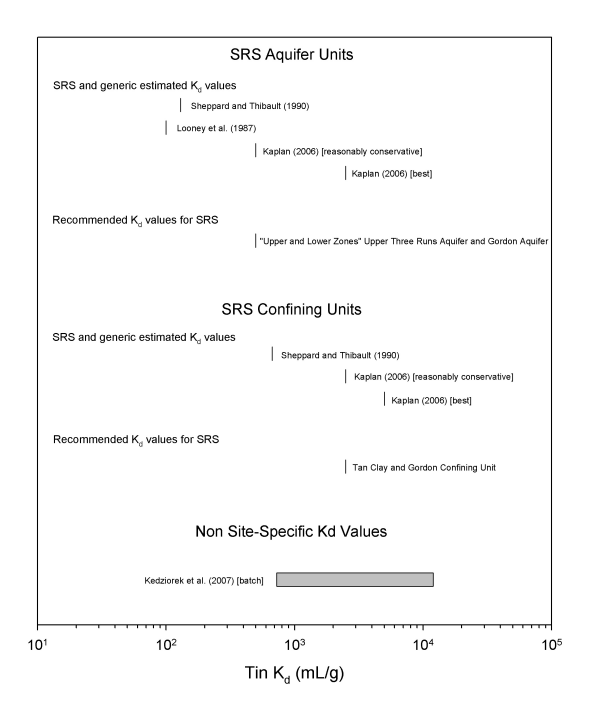


Figure 2-8. Measured and Estimated Site-Specific and Generic Tin K_d Ranges and Values and Recommended Tin K_d Values for Aquifer Units and Confining Units at SRS (Table 2-9). Recommended Tin K_d Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

2.4.8 **lodine**

I-129 generally occurs as iodate (IO_3^-), iodide (I^-), or organic iodine in natural environments. Sorption of iodine species appears to be controlled in part by soil organic matter and in part by iron and aluminum oxides, with adsorption of iodine becoming increasingly important under more acid conditions. Although the extent of sorption is typically low, especially in systems containing little or no organic matter, I^- and IO_3^- are sorbed to a measurable extent by soils and some oxide minerals at near-neutral and alkaline pH conditions. The adsorption behavior of IO_3^- also appears to be appreciably different from that of I^- , because IO_3^- sorbs much more strongly than I^- to soil and mineral surfaces (Ticknor and Cho, 1990). Mechanisms causing this sorption behavior of iodine at near-neutral and alkaline pH conditions are not completely understood. Some have proposed that this observed adsorption behavior in soils may be the result of the oxidation of I^- and/or reduction of IO_3^- to the more reactive molecular I_2^0 (aq) and/or its hydrolysis products (Ticknor and Cho, 1990).

Hoeffner (1985) summarized iodine sorption studies conducted to determine the performance of the low-level waste burial ground at SRS. Hoeffner (1985) describes a batch equilibrium study to measure $I^ K_d$ values as a function of I^- concentration using a Savannah River Plant soil taken from depths 2.5–3.0 m [8–10 ft] below grade and a low ionic strength groundwater representative of the burial ground (pH 5.0). K_d values ranged from 10 mL/g at 5 ppb I^- to 3.6 mL/g at 500 ppb I^- . Differences in pH were also noted to influence I^- sorption. Hoeffner (1985) reported an expected range of I-129 K_d values from 3 to 10 mL/g over a pH range of 3.4 to 7.3, and an expected K_d of 5 mL/g with a groundwater of typical composition at the SRS burial ground (pH 4.7).

Kaplan (2003) measured K_d values for I^- as a function of pH on two sediments from SRS using batch techniques (14-day equilibrium). Over the pH range of 3.5 to 7, $I^ K_d$ values ranged from 0.6 to 0.7 mL/g for a silty-sand upland SRS sediment (0.3 wt% Fe_2O_3 , <0.02 wt% organic carbon) and from 1 to 2 mL/g for a silty-sand wetland SRS sediment (0.01 wt% Fe_2O_3 , 0.14 wt% organic carbon). The K_d values were consistently greater for the wetland sediment and were attributed to its greater organic matter content. The measured $I^ K_d$ values also generally decreased with increasing pH, as would be expected for dissolved anionic contaminants.

Hu, et al. (2005) conducted column studies of I⁻ and iodate (IO₃⁻) sorption and transport on three uncontaminated sediments from SRS. The sediments were collected at different depths (surface soil, subsoil, and aquifer soil), providing an opportunity to examine the influence of organic matter, clay mineralogy, soil pH, and texture on the transport behavior of iodine species. For SRS surface soil {pH 4.9, 2.3-wt% organic matter, 9.5-wt% clay, 2.8 g/kg iron [2,800 ppm], 1.3 g/kg Al [1,300 ppm]}, retardation factors ranged from 0.93 to 0.99 for I⁻ and 2.29 to 2.92 for IO₃⁻. For the SRS subsoil {pH 5.36; 0.072-percent organic matter, 19.8-wt% clay, 16.2 g/kg [16,200 ppm] iron, 2.9 g/kg [2,900 ppm] Al}, retardation factors for I⁻ ranged from 5.5 to 6.3, and no breakthrough for IO₃⁻ was observed. For aquifer soil {pH 5.42, 0.013-percent organic matter, 1.9-wt% clay, 7.3 g/kg [7,300 ppm] iron, 0.42 g/kg [420 ppm] Al}, retardation factors ranged from 1.00 to 1.35 for I⁻ and 7.92 to 16.2 for IO₃⁻. Results indicate very limited sorption of I⁻ during transport in surface soil and aquifer soil. Retardation factors ranged from

0.93 to 1.35, which equate to K_d values of approximately (-0.015) to 0.09 mL/g.² However, there was significant retardation of I^- in the subsoil sample; the retardation factors of 5.5 to 6.3 equate to K_d values of approximately 1 to 1.4 mL/g. The subsoil sample contained appreciable amounts of iron and aluminum oxide minerals, which possess positively charged surfaces and contribute to anion adsorption. Iodate is expected to exhibit more retarded transport than I^- because it interacts more strongly with clays and organic matter. This behavior was confirmed in this study; IO_3^- consistently showed greater sorption than I^- . Measured retardation factors for IO_3^- in surface and aquifer soils ranged from 2.29 to 16.2, which equate to K_d values of approximately 0.3 to 4.0 mL/g.

Kaplan and Serkiz (2006) conducted batch experiments to measure iodine K_d values, as Γ , on a clayey sediment and a sandy sediment collected from the E-Area at SRS as a function of pH and DO. The sediments were collected from the upper zone of the Upper Three Runs aquifer and represented end members with respect to contaminant sorption properties expected in the E-Area subsurface. Iodine K_d values were reported at DO concentrations ranging from 0 to 300 mg/L [0 to 300 ppm] for pHs of 3.9, 5.3, and 6.7. Table 2-1 indicates that DO concentrations ranging from 0 to 10 mg/L are characteristic of sandy aquifer and clay confining units beneath GSA. At pH 3.9 and a DO concentration of 0 mg/L [0 ppm], Kaplan and Serkiz (2006) reported an iodine K_d value of -0.2 ± 0.2 mL/g (\pm 95-percent confidence interval) for clayey sediment and an iodine K_d value of 0 mL/g for sandy sediment. At pH 5.3 and a DO concentration of 0 mg/L [0 ppm], Kaplan and Serkiz (2006) reported an iodine K_d value of 0.1 \pm 0.4 mL/g for clayey sediment and an iodine K_d value of 0 mL/g for sandy sediment. At pH 6.7 and a DO concentration of 0 mg/L [0 ppm], Kaplan and Serkiz (2006) reported an iodine K_d value of 0.2 \pm 0.1 mL/g for clayey sediment and an iodine K_d value of 0 mL/g for sandy sediment.

Sheppard and Thibault (1990) specified an iodine default K_d for sand of 1.0 mL/g (22 observations ranging from 0.04 to 81 mL/g) and an iodine default K_d for clay of 1.0 mL/g (8 observations ranging from 0.2 to 29). Looney, et al. (1987) estimated a reference SRS soil K_d of 0.2 mL/g for iodine from an estimated range of 0.001 to 100 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" iodine K_d values of 0 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" iodine K_d values of 0.3 and 0.6 mL/g for clayey subsurface sediment at SRS. The iodine K_d values estimated by Kaplan (2006) were based on the iodine sorption data of Hoeffner (1985), Kaplan (2003), and Kaplan and Serkiz (2006) described previously.

Measured, estimated, and recommended iodine K_d values are summarized in Table 2-10. Hoeffner (1985) reported a measured range of iodine K_d values for SRS soils of 3 to 10 mL/g over a pH range of 3.4 to 7.3 and an expected iodine K_d of 5 mL/g with a groundwater of typical composition at the SRS burial ground (pH 4.7). These values are higher than iodine K_d values measured by Kaplan (2003) on silty-sand upland and wetland sediments from SRS. Over the pH range of 3.5 to 7, Kaplan (2003) reported an iodine K_d range of 0.6 to 0.7 mL/g for the upland soil and an iodine K_d range of 1 to 2 mL/g for the wetland soil. Hu, et al. (2005) performed column experiments to measure I^- and IO_3^- retardation factors in a surface soil, a

²Retardation factor (R_f converted to K_d based on following equation: $K_d = \frac{\left(R_f - 1\right)}{\rho_b}\theta$, where θ is porosity and ρ_b is bulk density.

Table 2-10. Measured, Estimated, and Recommended Iodine K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Iodine K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Iodine K _d (mL/g)*	Generic Estimated Iodine K _d (mL/g)†	Recommended lodine K _d (mL/g)
Upper Three Runs Aquifer: Upper	3–10‡ 0.6–2§ -0.015–4¶ -0.4–0.5#	Batch Batch Column Batch	0/0	1.0	0
Tan Clay Confining Unit			0.3/0.6	1.0	0.2
Upper Three Runs Aquifer: Lower			0/0	1.0	0
Gordon Confining Unit			0.3/0.6	1.0	0.2
Gordon Aquifer			0/0	1.0	0

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

†Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

‡Hoeffner, S.L. "Radionuclide Sorption on Savannah River Plant Burial Ground Soil—A Summary and Interpretation of Laboratory Data." DP–1702. Aiken, South Carolina: E.I. du Pont de Nemours and Company, Savannah River Laboratory. 1985.

§Kaplan, D.I. "Influence of Surface Charge of an Fe-Oxide and an Organic Matter Dominated Soil on Iodide and Pertechnetate Sorption." *Radiochimica Acta.* Vol. 91, No. 3. pp. 173–178. 2003.

¶Hu, Q., Z. Pihong, J.E. Moran, and J.C. Seaman. "Sorption and Transport of Iodine Species in Sediments from Savannah River and Hanford Sites." *Journal of Contaminant Hydrology*. Vol. 78. pp. 185–205. 2005. #Kaplan, D.I. and S.M. Serkiz. "Influence of Dissolved Organic Carbon and pH on Anion Sorption to Sediment." WSRC–RP–2006–00005. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

subsoil, and an aquifer soil from SRS. For I^- , retardation factors measured for the 3 soils ranged from 0.93 to 6.3, which equate to iodine K_d values ranging from -0.015 to 1.4 mL/g. For IO_3^- , retardation factors measured on the three soils ranged from 2.29 to 16.2, which equate to K_d values of 0.3 to 4.0 mL/g. The range of iodine K_d values measured by Hu, et al. (2005) (-0.015 to 4.0 mL/g) are consistent with the measured iodine K_d values of Kaplan (2003). The estimated iodine K_d value for sand of Sheppard and Thibault (1990) (1.0 mL/g) and the reference iodine K_d value of 0.2 mL/g specified by Looney, et al. (1987) for SRS soil are within the range of iodine K_d values measured on SRS sediments by Kaplan (2003) and Hu, et al. (2005). Kaplan and Serkiz (2006) reported iodine K_d values of 0 mL/g on sandy subsurface sediments from SRS over the pH range of 3.9 to 6.7. Based on the iodine K_d values reported by

Hoeffner (1985), Kaplan (2003), and Kaplan and Serkiz (2006), Kaplan (2006) estimated a "reasonably conservative" iodine K_d value of 0 mL/g for sandy sediments at SRS. The mineral compositions of the aquifer units at SRS are not significantly different and, thus, these units should exhibit similar iodine sorption. Based on the range of reported site-specific iodine K_d values summarized previously and considering the "reasonably conservative" iodine K_d estimated by Kaplan (2006) for SRS sandy sediments, an iodine K_d value of 0 mL/g is recommended for aquifer units at SRS.

Kaplan and Serkiz (2006) reported iodine K_d values of 0.1 ± 0.4 mL/g and 0.2 ± 0.1 mL/g on clayey subsurface sediments from SRS at pHs of 5.3 and 6.7, respectively. Based on the iodine K_d values reported by Hoeffner (1985), Kaplan (2003), and Kaplan and Serkiz (2006), Kaplan (2006) estimated a "reasonably conservative" iodine K_d value of 0.3 mL/g for clayey subsurface sediments at SRS. Sheppard and Thibault (1990) recommended a default iodine K_d value of 1.0 for clay. Based on the iodine K_d values reported by Kaplan and Serkiz (2006) at pHs of 5.3 and 6.7 and considering the "reasonably conservative" iodine K_d estimated by Kaplan (2006) for SRS clayey sediments, an iodine K_d value of 0.2 mL/g is recommended for clay confining units at SRS.

The position of the iodine K_d values recommended in this review with respect to measured and estimated site-specific and generic iodine K_d ranges and values is graphically displayed in Figure 2-9. This figure illustrates the uncertainty associated with the iodine K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific iodine sorption data (Kaplan, 2003; Hu et al., 2005; Kaplan and Serkiz, 2006), the recommended iodine K_d values are associated with low uncertainty.

2.4.9 **Cesium**

Cs-137 exists in the environment in the +1 oxidation state. Cesium forms few stable complexes and is likely to exist in groundwater as the uncomplexed Cs⁺ ion. Cs⁺ primarily adsorbs via ion exchange reactions with sedimentary clay minerals. In general, most soils sorb cesium rather strongly (Ames and Rai, 1978). The extent to which adsorption will occur will depend on the concentration of clays in the soil and the concentration of major cations, such as K⁺, that can effectively compete with Cs⁺ for adsorption sites. Cesium may also adsorb to iron oxides (Schwertmann and Taylor, 1989). Sorption of Cs⁺ is pH dependent and increases with increasing pH. Prout (1958) reported that cesium adsorption to iron-oxide-dominated soils from South Carolina decreased dramatically when the pH was less than 6.

Hoeffner (1985) summarized cesium sorption studies conducted to determine the performance of the low-level waste burial ground at SRS. The majority of studies used a Savannah River Plant soil taken from depths 2.5–3.0 m [8–10 ft] below grade and a low ionic strength groundwater representative of the burial ground (pH 5.0). Hoeffner (1985) found that cesium sorption was very dependent on pH and solution concentration. Hoeffner (1985) reported an expected range of cesium K_d values from 330 to 1800 mL/g over a pH range of 3.4 to 7.3 and an expected K_d value of 500 for a groundwater of typical composition at SRS burial ground (pH 4.7).

McConnell, et al. (1995) reported a cesium K_d value of 37 mL/g for SRS soil used in lysimeter tests to obtain information on the performance of radioactive wasteforms. This value was

obtained using a batch method. The composition and pH of the solution used in the batch test was not reported.

Elprince, et al. (1977) performed batch sorption experiments on surficial sediments collected from five streams (Upper Three Runs, Four Mile Creek, Pen Branch, Steel Creek, and Lower Three Runs), Par Pond, and the Savannah River at SRS. The Par Pond, Steel Creek, and Upper Three Runs sediments had a high percentage of quartz (54–66 wt%) and a lower CEC than the Pen Branch, Lower Three Runs, Four Mile Creek, and Savannah River sediments (<29- through 40-wt% quartz). The solution used in the batch experiments was a low ionic strength sodium solution (0.005 M sodium); the pH of the solution was not reported. Cesium K_d values for samples with the low CEC (Par Pond, Steel Creek, and Upper Three Runs) ranged from 18.3 to 40.1 mL/g. Cesium K_d values for sediments with a high CEC (Pen Branch, Lower Three Runs, Four Mile Creek, and Savannah River) ranged from 83.6 to 130 mL/g. The low CEC samples with high percentages of quartz were assumed to be sand-rich soils, and the samples with high CEC and low percentages of quartz were assumed to be clay-rich or clay soils.

Sheppard and Thibault (1990) estimated a cesium default K_d for sand of 280 mL/g (81 observations ranging from 0.2 to 10,000 mL/g) and a cesium default K_d for clay of 1,900 mL/g (28 observations ranging from 37 to 31,500). Looney, et al. (1987) estimated a reference SRS soil K_d of 500 mL/g for cesium from an expected range of 10 to 100,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" cesium K_d values of 15 and 50 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" cesium K_d values of 180 and 250 mL/g for clayey subsurface sediment at the SRS. The cesium K_d values estimated by Kaplan (2006) were based on cesium sorption data of Hoeffner (1985), Bibler and Marson (1992), Johnson (1995), and Goto (2001). As reported in Kaplan (2006), Bibler and Marson (1992) performed batch sorption experiments to measure cesium K_d values on four soils collected from the E-Area at SRS. Reported cesium K_d values ranged from 7 to 37 mL/g on these soils. Kaplan (2006) reported that Johnson (1995) performed field tests (squeezed pore water out of aquifer sediment) using subsurface F-Area sediments and measured cesium K_d values ranging from 5 to 95 mL/g over the pH range of 4.2 to 6.4. The measured cesium K_d values were very pH dependent and increased with increasing pH. At pH 5.5, Johnson (1995) measured cesium K_d values ranging from 20 to 35 mL/g. Kaplan (2006) reported that Goto (2001) conducted cesium sorption experiments with 5 SRS sediments of varying clay content. Measured cesium K_d values ranged from 21 ± 7 to 225 ± 51 mL/g (± 95-percent confidence interval) and varied directly with clay content. Sediments with clay content >10 wt% had cesium K_d values >139 mL/g, and those with clay content <10 wt% had cesium K_d values ranging from 15 to 73 mL/g.

Measured, estimated, and recommended cesium K_d values are summarized in Table 2-11. Hoeffner (1985) reported a range of cesium K_d values for SRS soils ranging from 330 to 1,800 mL/g over a pH range of 3.4 to 7.3 and an expected cesium K_d of 500 mL/g for a groundwater of typical composition at the SRS burial ground (pH 4.7). McConnell, et al. (1995) reported a measured cesium K_d value of 37 mL/g for an SRS soil; however, the pH and composition of the water used in the batch sorption experiments were not reported. Elprince, et al. (1977) report measured cesium K_d values ranging from 18.3 to 40.1 for sand-rich, low CEC surficial sediments from SRS and cesium K_d values ranging from 83.6 to 130 mL/g for

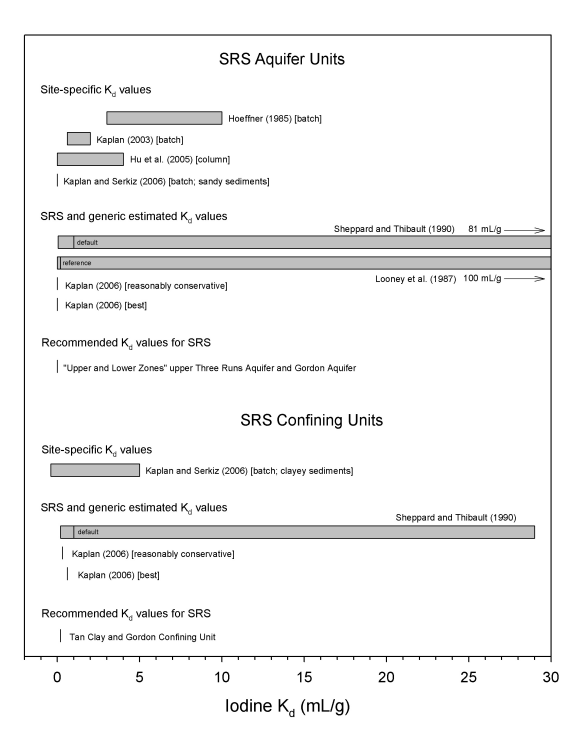


Figure 2-9. Measured and Estimated Site-Specific and Generic Iodine K_d Ranges and Values and Recommended Iodine K_d Values for Aquifer Units and Confining Units at SRS (Table 2-10). Recommended Iodine K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

clay-rich, high CEC surficial sediments from SRS. Kaplan (2006) estimated a "reasonably conservative" cesium K_d of 15 mL/g for sandy subsurface sediments at SRS. This "reasonably conservative" K_d value was based on site-specific cesium K_d measurements reported by Bibler and Marson (1992) (7 to 37 mL/g), Johnson (1995) (5 to 95 mL/g), and Goto (2001) (15 to 73 mL/g; sediments with clay content <10 wt%). These values are consistent with the measured site-specific cesium K_d values reported by Elprince, et al. (1977) for sand-rich surficial sediments (18.3 to 40.1 mL/g), but lower than the range of cesium K_d values for SRS soils reported by Hoeffner (1985) (330 to 1,800 mL/g). The default cesium K_d value for sand of 280 mL/g from the compilation of Sheppard and Thibault (1990) and the reference soil cesium K_d of 500 mL/g estimated by Looney, et al. (1987) are greater than the site specific cesium K_d values measured by Elprince, et al. (1977), Bibler and Marson (1992), Johnson (1995), and Goto (2001). Based on the measured cesium K_d values of Elprince, et al. (1997), Bibler and Marson (1992), Johnson (1995), and Goto (2001) and considering the "reasonably conservative" cesium K_d of 15 mL/g estimated by Kaplan (2006) for sandy subsurface sediments at SRS, a cesium K_d value of 15 mL/g is recommended for the upper zone of the Upper Three Runs aguifer. The lower zone of the Upper Three Runs aguifer and the Gordon aquifer have reported pH values mainly above 6.0 and 5.7, respectively. Because cesium sorption increases with increasing pH, a cesium K_d value of 50 mL/g is recommended for these aquifer units. This recommended value lies between cesium K_d values of 20 to 35 mL/g measured by Johnson (1995) at a pH of 5.5 and the maximum cesium K_d value of 95 mL/g measured by Johnson (1995) over the pH range of 4.2 to 6.4.

Elprince, et al. (1977) reported measured cesium K_d values ranging from 83.6 to 130 mL/g for clay-rich, high CEC surficial sediments from SRS. Kaplan (2006) estimated a "reasonably conservative" cesium K_d of 180 mL/g for clayey subsurface sediments at SRS. This "reasonably conservative" K_d value was based on site-specific cesium K_d measurements reported by Goto (2001) (cesium K_d >139 mL/g for sediments with clay content >10 wt%). The site-specific cesium K_d values reported by Elprince, et al. (1977) and Goto (2001) are much lower than the cesium K_d value of 1,900 mL/g recommended by Sheppard and Thibault (1990) for clay. Based on the measured cesium K_d values of Elprince, et al. (1997) and Goto (2001) and considering the "reasonably conservative" cesium K_d of 180 mL/g estimated by Kaplan (2006) for clayey subsurface sediments at SRS, a cesium K_d value of 150 mL/g is recommended for the clay confining units at SRS.

The position of the cesium K_d values recommended in this review with respect to measured and estimated site-specific and generic cesium K_d ranges and values is graphically displayed in Figure 2-10. This figure illustrates the uncertainty associated with the cesium K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific cesium sorption data (Elprince, et al., 1977; Bibler and Marson, 1992; Johnson, 1995; Goto, 2001), the recommended cesium K_d values are associated with low uncertainty.

2.4.10 Lead

Pb-210 has 3 known oxidation states, 0, +2, and +4, and the most common redox state found in the environment is the divalent form. Dissolved lead in natural systems may exist in free ionic form (Pb²⁺) and also as hydrolytic and complex species. At pH values exceeding 7, aqueous lead exists mainly as carbonate complexes [Pb(CO)₃⁰(aq) and Pb(CO₂)₂²⁻]. Studies have shown

Table 2-11. Measured, Estimated, and Recommended Cesium $K_{\rm d}$ Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Cesium K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Cesium K _d (mL/g)*	Generic Estimated Cesium K _d (mL/g)†	Recommended Cesium K _d (mL/g)
Upper Three Runs Aquifer: Upper	330–1,800‡ 37§ 18.3–130¶ 7–37# 5–95** 15–377††	Batch Batch Batch Batch In situ Batch	15/50	280	15
Tan Clay Confining Unit	1	1	180/250	1,900	150
Upper Three Runs Aquifer: Lower	_	_	15/50	280	50
Gordon Confining Unit	_	_	180/250	1,900	150
Gordon Aquifer	_	_	15/50	280	50

*Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

†Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

‡Hoeffner, S.L. "Radionuclide Sorption on Savannah River Plant Burial Ground Soil—A Summary and Interpretation of Laboratory Data." DP–1702. Aiken, South Carolina: E.I. du Pont de Nemours and Company, Savannah River Laboratory. 1985.

§McConnell, Jr., J.W., R.D. Rogers, J.D. Jastrow, W.E. Sanford, S.R. Cline, T.M. Sulllivan, and M. Fuhrmann. "Results After Ten Years of Field Testing Low-level Radioactive Waste Forms Using Lysimeters." ANL/ER/CP 95275. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1995.

¶Elprince, A.M., C.I. Rich, and D.C. Martens. "Effect of Temperature and Hydroxy Aluminum Interlayers on the Adsorption of Trace Radioactive Cesium by Sediments near Water-Cooled Nuclear Reactors." *Water Resources Research.* Vol. 13, No. 2. pp. 375–380. 1977.

#Bibler, J.P. and D.B. Marson. "Behavior of Mercury, Lead, Cesium, and Uranyl Ions on Four SRS Soils." WSRC-RP-92-326. Aiken, South Carolina: Westinghouse Savannah River Company. 1992.

**Johnson, W.H. "Sorption Models for U, Cs, and Cd on Upper Atlantic Coastal Plain Soils." A Thesis Presented to the Academic Faculty at Georgia Institute of Technology, Atlanta, Georgia. 1995.

††Goto, M. "Development of a Quantitative Model for Binding Cesium to SRS soils." Masters' Degree Thesis. Georgia Institute of Technology, Atlanta, Georgia. 2001.

that, in neutral to high pH conditions, Pb²+ can preferentially exchange for Ca²+ and K⁺ (Zimdahl and Hassett, 1977; Bittel and Miller, 1974). Under acidic pH conditions, hydrogen ions and aluminum ions will displace lead from mineral exchange sites. Lead is known to adsorb onto soil constituent surfaces such as clays, oxides, hydroxides, oxyhydroxides, and organic matter. Lead adsorption studies on bulk soils indicate that the adsorption is strongly correlated with pH and the CEC values of soils (Bittel and Miller, 1974; Griffin and Shimp, 1976; Haji-Djafari, et al., 1981; Scrudato and Estes, 1975; Zimdahl and Hassett, 1977). Properties that affect CEC of soils, such as organic matter content, clay content, and surface area, have a greater effect on lead adsorption than soil pH.

Bibler and Marson (1992) performed batch sorption experiments to measure lead K_d values on four Savannah River soils collected from the E-Area. Bibler and Marson (1992) reported lead K_d values ranging from 63 to >2,000,000 mL/g. However, it is likely that these experiments were conducted at concentrations above the solubility limit of lead (e.g., initial Pb²⁺ of 5 mg/L), and therefore reflect precipitation more than adsorption.

Kaplan and Serkiz (2000) performed desorption measurements to measure lead K_d values on shallow contaminated sediments from the TNX Outfall Operable Unit at SRS. The desorption K_d values were based on extractable portions that were comprised of the exchangeable, organic-bound, amorphous Fe–oxide bound, and water extractable fractions. Basing the K_d estimates on these extractable fractions instead of the more easily determined total concentrations in the sediment, more conservative and technically defensible K_d values were measured. The contaminated sediments were collected at depths of 0 to 0.6 m [0 to 2 ft] by hand auger and were composed dominantly of sand and silt (i.e., sand and silt contents ranging from 80 to 94 wt%). The sediment pH of the sediments ranged from 4.00 to 5.78. Measured lead K_d values ranged from >11,460 to >38,751 mL/g. Over the pH range expected in the subsurface at SRS (5.2 to 7.7), Kaplan and Serkiz (2000) reported lead K_d values of >34,511 and >11,460 mL/g for sediments with pHs of 5.26 and 5.78, respectively.

Haji-Djafari, et al. (1981) measured lead K_d values on sandy sediments from the Split Rock Formation in central Wyoming as a function of pH. The K_d values reported in this study are listed in EPA (1999). Lead K_d values for pHs of 2.0, 4.5, 5.75, and 7.0 were 20, 100, 1,500, and 4,000 mL/g, respectively. As expected, lead adsorption increased with increasing pH. Gerriste, et al. (1982) examined lead adsorption onto generic soils as a function of pH. Results indicated that lead adsorption increased with increasing soil pH (from 4 to 8). A summary of K_d values reported by Gerriste, et al. (1982) are also listed in EPA (1999). Lead K_d values for pHs of 4.5, 5.0, 7.5, and 8.0 were 280; 1,295; 3,000; and 4,000 mL/g, respectively.

Sheppard and Thibault (1990) estimated a lead default K_d for sand of 270 mL/g (3 observations ranging from 19 to 405 mL/g) and a lead default K_d for clay of 550 mL/g (based on a soil-to-plant concentration ratio). Looney, et al. (1987) estimated a reference SRS soil K_d of 100 mL/g for lead from an expected range of 1 to 10,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" lead K_d values of 500 and 2,000 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" lead K_d values of 2,500 and 5,000 mL/g for clayey subsurface sediment at the SRS. The Kaplan (2006) estimations were based in part on the measured lead K_d values reported by Bibler and Marson (1992).

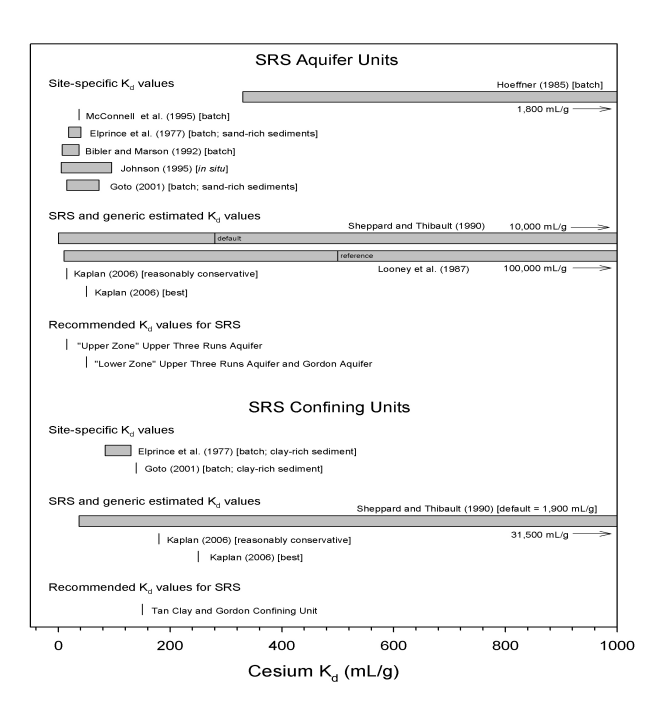


Figure 2-10. Measured and Estimated Site-Specific and Generic Cesium $K_{\rm d}$ Ranges and Values and Recommended Cesium $K_{\rm d}$ Values for Aquifer Units and Confining Units at SRS (Table 2-11). Recommended Cesium $K_{\rm d}$ Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

Measured, estimated, and recommended lead $\rm K_d$ values are summarized in Table 2-12. Kaplan and Serkiz (2000) reported lead $\rm K_d$ values of >34,511 and >11,460 mL/g for contaminated sediments collected at the SRS with pHs of 5.26 and 5.78, respectively. Haji-Djafari, et al. (1981) reported measured lead $\rm K_d$ values on sandy sediments ranging from 20 to 4,000 mL/g over the pH range of 2.0 to 7.0. At pHs of 5.75 and 7.0, lead $\rm K_d$ values were 1,500 and 4,000 mL/g, respectively. Gerriste, et al. (1982) reported measured lead $\rm K_d$ values on generic soils ranging from 280 to 4,000 mL/g over the pH range of 4.5 to 8.0. At a pH of 5.0 and 7.5, lead $\rm K_d$ values of 1,295 and 3,000 mL/g were reported. The default lead $\rm K_d$ value for sand of 270 mL/g from the compilation of Sheppard and Thibault (1990) and the reference SRS soil lead $\rm K_d$ of 100 mL/g estimated by Looney, et al. (1987) are much lower than the lead $\rm K_d$ values reported by Haji-Djafari, et al. (1981) at pHs of 5.75 and 7.0 (1,500 and 4,000 mL/g, respectively) and by Gerriste, et al. (1982) at pHs of 5.0 and 7.5 (1,295 and 3,000 mL/g, respectively). The "reasonably conservative" lead $\rm K_d$ value of 500 mL/g estimated by Kaplan (2006) for sandy subsurface sediment at SRS is also lower than the lead $\rm K_d$ values reported by Haji-Djafari, et al. (1981) at pHs of 5.75 and 7.0 and by Gerriste, et al. (1982) at pHs of 5.0 and

Table 2-12. Measured, Estimated, and Recommended Lead K _d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site						
	Measured Lead K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Lead K _d (mL/g)*	Generic Estimated Lead K _d (mL/g)†	Recommended Lead K _d (mL/g)	
Upper Three Runs Aquifer: Upper	63–2.0 × 10 ⁶ ‡ >11,460– >38,751§	Batch In situ	500/2,000	270	500	
Tan Clay Confining Unit	1		2,500/5,000	550	2,500	
Upper Three Runs Aquifer: Lower	1		500/2,500	270	1,500	
Gordon Confining Unit	I		2,500/5,000	550	2,500	
Gordon Aquifer	_		500/2,500	270	1,500	

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

 $[\]dagger$ Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_d s, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

[‡]Bibler, J.P. and D.B. Marson. "Behavior of Mercury, Lead, Cesium, and Uranyl Ions on Four SRS Soils." WSRC–RP–92–326. Aiken, South Carolina: Westinghouse Savannah River Company. 1992.

 $[\]S$ Kaplan, D.I. and S.M. Serkiz. "In Situ K_d Values and Geochemical Behavior for Inorganic and Organic Constituents of Concern at the TNX Outfall Delta." WSRC-TR-99-00488. Aiken, South Carolina: Westinghouse Savannah River Company. 2000.

7.5 and is significantly lower than site-specific lead $\rm K_d$ values reported by Kaplan and Serkiz (2000) at pHs of 5.26 and 5.78 (>34,511 and >11,460, respectively). Based on these observations, the "reasonably conservative" lead $\rm K_d$ value of 500 mL/g recommended for SRS sandy subsurface sediments by Kaplan (2006) is recommended for the upper zone of the Upper Three Runs aquifer. The lower zone of the Upper Three Runs aquifer and the Gordon aquifer have reported pH values above 6.0 and 5.7, respectively. Because lead $\rm K_d$ values are highly sensitive to pH (i.e., lead $\rm K_d$ increases significantly with increasing pH), a lead $\rm K_d$ value of 1,500 mL/g is recommended for the lower zone of the Upper Three Runs aquifer and the Gordon aquifer. This value was approximated for a pH of 6.0 based on the lead $\rm K_d$ values of 1,500 and 4,000 mL/g reported by Haji-Djafari, et al. (1981) at pHs of 5.75 and 7.0 and the lead $\rm K_d$ values of 1,295 and 3,000 mL/g reported by Gerriste, et al. (1982) at pHs of 5.0 and 7.5.

Sheppard and Thibault (1990) recommend a default lead K_d value of 550 mL/g for clay based on a soil-to-plant concentration ratio. Kaplan (2006) estimated a "reasonably conservative" lead K_d value of 2,500 mL/g for clayey subsurface sediment at the SRS based in part on measured lead K_d values reported by Bibler and Marson (1992). Lacking additional site specific or generic lead sorption data on clay sediments, the "reasonably conservative" lead K_d value of 2,500 mL/g estimated by Kaplan (2006) is recommended for clay confining units at SRS.

The position of the lead K_d values recommended in this review with respect to measured and estimated site-specific and generic lead K_d ranges and values is graphically displayed in Figure 2-11. This figure illustrates the uncertainty associated with the lead K_d values recommended for SRS aquifer units and clay confining units. The recommended lead K_d values are associated with high uncertainty due to wide variation in measured site-specific sorption data. For example, Bibler and Marson (1992) reported lead K_d values ranging from 2 to 2 million mL/g on soils from the E-Area and Kaplan and Serkiz (2000) measured lead K_d values ranging from >11,460 to >38,751 on contaminated SRS sediments over the pH range of 4.0 to 5.78.

2.4.11 **Uranium**

In natural environments, the most important oxidation states of uranium are +4 and +6. U(VI) species dominate in oxidizing environments; the dominant species are UO₂²⁺ at pH values less than 5 and mixed carbonate/hydroxyl and carbonate species at pH values between 5 and 9 (Langmuir, 1997). In low ionic strength solutions with low U(VI) concentrations, UO₂²⁺ concentrations will likely be controlled by cation exchange and adsorption processes. The uranyl ion and its complexes adsorb onto clays (Ames, et al. 1982), organics (Read, et al. 1993; Shanbhag and Choppin, 1981), and oxides (Hsi and Langmuir, 1985; Waite, et al. 1994). As ionic strength increases, other ions, notably Ca²⁺, Mg²⁺, and K⁺, displace the uranyl ion from soil exchange sites, forcing it into solution. U(VI) retention by soils and rocks in alkaline conditions is poor because of the predominance of neutral and negatively charged species. An increase in pCO₂ in soil solutions reduces U(VI) adsorption by promoting the formation of poorly sorbing carbonate complexes (Pabalan, et al., 1998). U(IV) species dominate in reducing environments. U(IV) tends to hydrolyze and form strong hydrolytic complexes. U(IV) also forms strong complexes with naturally occurring organic materials.

Serkiz and Johnson (1994) investigated the partitioning of uranium on soil in contaminated groundwater downgradient of the F and H Areas Seepage Basins at SRS. Their study tabulated an extensive set of field-derived K_d values for U-238 and U-235 for 48 soil/porewater samples.

The soil/porewater samples were collected over a range of geochemical conditions (e.g., pH and contaminant concentration). The K_d values determined by Serkiz and Johnson (1994) are plotted as a function of pH in Figure 2-12. The uranium K_d values vary from 1.2 to 34,000 mL/g over a pH range of 3 to 6.8. The K_d values indicate that uranium sorption increases with increasing pH over the range 3 to 5.2. The adsorption of uranium is at a maximum at approximately pH 5.2 and then decreases with increasing pH over the pH range of 5.2 to 6.8. Serkiz and Johnson found that the field-derived K_d values for U-238 and U-235 were not well correlated with the weight percentage of clay-sized particles or CEC of the soil samples, and they proposed that the uranium was not binding to clays by a cation exchange reaction, but rather to a mineral surface coating with a variable surface charge due to varying porewater pH.

Johnson (1995) conducted field and laboratory studies to determine the processes controlling uranium sorption to SRS sediments. A portion of this work involved determining K_d values on paired sediment-pore water samples collected within a contaminated plume from the F Area at SRS. Sequential extractions were used to determine uranium concentrations in sediment and pore water uranium concentrations in a saturated past extract. Johnson (1995) reported uranium K_d values of 87 mL/g at pH 4.7, 100 mL/g at pH 4.75, 37 mL/g at pH 5.05, and 93 mL/g at pH 5.12. Johnson (1995) attributed the lack of a trend in the uranium K_d values with pH to mineralogical variations in the sampled sediments. Johnson (1995) also performed batch sorption tests using a sandy subsurface F-Area sediment to measure uranium K_d values as a function of pH. At pHs of 4.82, 4.87, 5.01, and 5.04, Johnson (1995) reported uranium K_d values of 116, 153, 198, and 238 mL/g, respectively.

Kaplan and Serkiz (2000) performed desorption measurements to measure uranium K_d values on shallow contaminated sediments from the TNX Outfall Operable Unit at SRS. The desorption K_d values were based on extractable portions that comprise the exchangeable, organic-bound, amorphous Fe-oxide bound, and water extractable fractions. Basing the K_d estimates on these extractable fractions instead of the more easily determined total concentrations in the sediment, more conservative and technically defensible K_d values were measured. The contaminated sediments were collected at depths of 0 to 0.6 m [0 to 2 ft] by hand auger and were composed dominantly of sand and silt (i.e., sand and silt contents ranging from 80 to 94 wt%). The sediment pH of the sediments ranged from 4.00 to 5.78. Measured uranium K_d values ranged from 170 to 6,493 mL/g. Over the pH range expected in the subsurface at the SRS (5.2 to 7.7), Kaplan and Serkiz (2000) reported uranium K_d values of 170 and 2,110 mL/g for sediments with pHs of 5.26 and 5.78, respectively.

Barnett, et al. (2000) conducted batch tests to study uranium sorption onto a sediment sample from SRS. The sediment sample was acquired from a depth of 45 m [148 ft] in the McBean Formation (lower zone of the Upper Three Runs aquifer) near the burial ground site. The sediment consisted of 69-wt% sand, 7-wt% silt, and 24-wt% clay. Sand-sized particles were noted to be coated with iron oxides. The sediment had 25.7 g/kg [25,700 ppm] iron, 0.12 g/kg [120 ppm] organic matter, and a pH of 4.1. A 0.01 M NaNO $_3$ solution that was adjusted to the pH of the solid phase (4.1) was used in adsorption isotherm measurements. Barnett, et al. (2000) reported a K $_d$ value of 230 \pm 7 mL/g (\pm 95-percent confidence interval) for the SRS sediment.

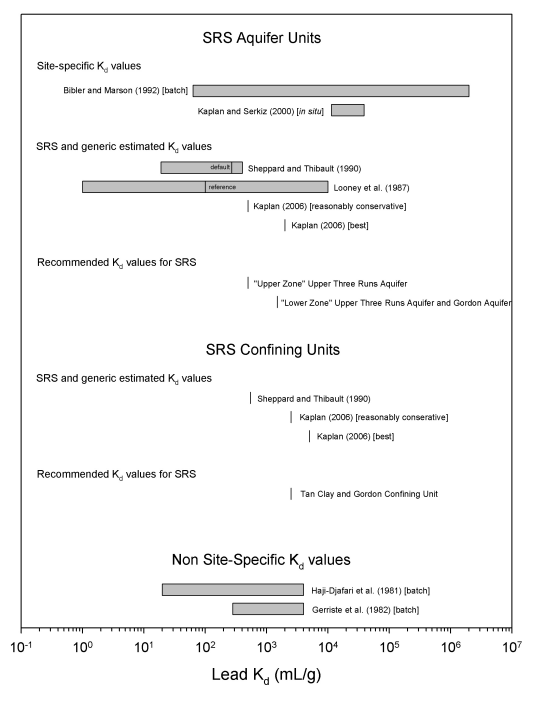


Figure 2-11. Measured and Estimated Site-Specific and Generic Lead $K_{\rm d}$ Ranges and Values and Recommended Lead $K_{\rm d}$ Values for Aquifer Units and Confining Units at SRS (Table 2-112). Recommended Lead $K_{\rm d}$ Values Are Associated With High Uncertainty Due to Wide Variation in Measured Site-Specific Sorption Data.

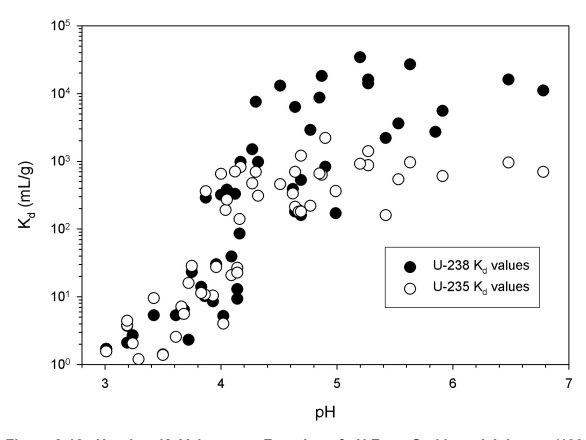


Figure 2-12. Uranium K_d Values as a Function of pH From Serkiz and Johnson (1994)

Sheppard and Thibault (1990) estimated a uranium default K_d for sand of 35 mL/g (24 observations ranging from 0.03 to 2,200 mL/g) and a uranium default K_d for clay of 1,600 (7 observations ranging from 46 to 395,100). Looney, et al. (1987) specified a reference SRS soil K_d of 40 mL/g for uranium from an estimated range of 0.1 to 1,000,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" uranium K_d values of 100 and 200 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" uranium K_d values of 150 and 300 mL/g for clayey subsurface sediment at SRS. The uranium K_d values estimated by Kaplan (2006) were based on the uranium sorption data of Serkiz and Johnson (1994) and Johnson (1995) described previously.

Measured, estimated, and recommended uranium K_d values are summarized in Table 2-13. Serkiz and Johnson (1994) reported K_d values for U-238 and U-235 in soil/porewater samples collected from the upper zone of the Upper Three Runs aquifer that ranged from 1.2 to 34,000 mL/g over the pH range of 3 to 6.8 (Figure 2-4). Over the pH range expected in aquifer units beneath SRS (5.2 to 7.7), uranium K_d values range from about 100 to 34,000 mL/g. Johnson (1995) reported uranium K_d values in sediment/porewater samples collected from the upper zone of the Upper Three Runs aquifer that ranged from 37 to 100 mL/g over the pH range of 4.7 to 5.12. Johnson (1995) also reported uranium K_d values measured by batch methods using sandy subsurface F-Area sediment that ranged from 116 to 238 mL/g over the pH range of 4.82 to 5.04. Kaplan and Serkiz (2000) reported uranium K_d values of 170 and 2,110 mL/g

for contaminated sediments at the SRS with pHs of 5.26 and 5.78, respectively. Barnett, et al. (2000) reported a uranium K_d value of 230 ± 7 mL/g for a sediment sample acquired from the lower zone of the Upper Three Runs aquifer near the burial ground site. This uranium K_d value was determined in a solution at low pH (i.e., pH 4.1). Clays in the lower zone of the Upper Three Runs aguifer are dominated by smectite, which has a high CEC capacity, and this may account for the higher minimum uranium K_d value observed for this single sediment sample from this unit when compared to measured uranium K_d values reported by Serkiz and Johnson (1994) for the upper zone of the Upper Three Runs aquifer at a similar pH of 4.1. Based on the measured uranium K_d values reported by Serkiz and Johnson (1994) and Johnson (1995). Kaplan (2006) estimated a "reasonably conservative" uranium K_d of 100 mL/g for sandy subsurface sediments at SRS. When compared to the uranium sorption data of Serkiz and Johnson (1994) (Figure 2-12), the estimated default uranium K_d value of 35 mL/g of Sheppard and Thibault (1990) for sand and the estimated uranium K_d value of 40 mL/g of Looney, et al. (1987) for soil at SRS correspond to measured uranium K_d values in low pH groundwater (e.g., pH < 4.3). Groundwater in the upper zone of the Upper Three Runs aguifer has an expected minimum pH of 5.2. Based on the uranium sorption data of Serkiz and Johnson (1994) and Johnson (1995), a uranium K_d value of 100 mL/g is recommended for the upper zone of the Upper Three Runs aquifer. Groundwater in the lower zone of the Upper Three Runs aguifer and the Gordon aguifer have expected pHs of 5.7 to 7.7. Uranium K_d values tend to reach a maximum at a pH of about 6.5 and then decrease with increasing pH (Pabalan, et al., 1998). Due to the absence of measured site-specific uranium K_d values at pHs above 6.8, the same value of 100 mL/g is recommended for the lower zone of the Upper Three Runs aguifer and the Gordon aguifer at SRS.

No site-specific uranium sorption data were found for clay confining units at SRS. Sheppard and Thibault (1990) recommended a default uranium K_d of 1,600 mL/g for clay. Based on the measured uranium K_d values reported by Serkiz and Johnson (1994), Johnson (1995), and Kaplan (2006) estimated a "reasonably conservative" uranium K_d of 150 mL/g for clayey subsurface sediments at SRS and this value is recommended for the tan clay unit at SRS. Groundwater in the tan clay confining unit has an expected minimum pH of 5.2, whereas groundwater in the Gordon confining unit has an expected pH range of 5.8 to 6.8 (Table 2-2). Because uranium K_d values tend to reach a maximum at a pH of about 6.5, a higher uranium K_d value of 300 mL/g is recommended for the Gordon confining unit. This value corresponds to the "best" uranium K_d value estimated by Kaplan (2006) for clayey subsurface sediments at SRS.

The position of the uranium K_d values recommended in this review with respect to measured and estimated site-specific and generic uranium K_d ranges and values is graphically displayed in Figure 2-13. This figure illustrates the uncertainty associated with the uranium K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific uranium sorption data (Serkiz and Johnson, 1994; Johnson (1995); Kaplan and Serkiz, 2000), the recommended uranium K_d values are associated with low uncertainty.

2.4.12 Neptunium

Np-237 may exist in several valance states; Np(VI), Np(V), and Np(IV) being the most important in natural waters. Over the pH range of most natural waters, Np(V) is present primarily as the cation NpO₂⁺. Np(V) aqueous species sorb to some extent to iron oxides and clays, but do not sorb to a major degree on most common minerals. Therefore, dissolved Np(V) is considered to

Table 2-13. Measured, Estimated, and Recommended Uranium K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Uranium K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Uranium K _d (mL/g)*	Generic Estimated Uranium K _d (mL/g)†	Recommended Uranium K _d (mL/g)
Upper Three Runs Aquifer: Upper	1.2-34,000‡ 1706,493§ 37-100¶ 116-238¶	In situ In situ In situ Batch	100/200	35	100
Tan Clay Confining Unit	_	_	150/300	1,600	150
Upper Three Runs Aquifer: Lower	223–237#	Batch	100/200	35	100
Gordon Confining Unit	_	_	150/300	1,600	300
Gordon Aquifer	_	_	100/200	35	100

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

†Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

‡Serkiz, S.M. and W.H. Johnson. "Uranium Geochemistry in Soil and Groundwater at the F and H Seepage Basins (U)." EPD–SGS–94–307. Aiken, South Carolina: Westinghouse Savannah River Company. 1994. §Kaplan, D.I. and S.M. Serkiz. "In-Situ K_d Values and Geochemical Behavior for Inorganic and Organic Constituents of Concern at the TNX Outfall Delta." WSRC–TR–99–00488. Aiken, South Carolina: Westinghouse Savannah River Company. 2000.

¶Johnson, W.H. "Sorption Models for U, Cs, and Cd on Upper Atlantic Coastal Plain Soils." A Thesis Presented to the Academic Faculty at Georgia Institute of Technology, Atlanta, Georgia. 1995.

#Barnett, M.O., P.M. Jardine, S.C. Brooks, and H.M. Selim. "Adsorption and Transport of Uranium(VI) in Subsurface Media." *Soil Science Society American Journal*. Vol. 64. pp. 908–917. 2000.

be relatively mobile in soil systems. Because NpO_2^+ does not compete favorably with dissolved Ca^{2+} and other divalent ions for adsorption sites on soils, the K_d values for Np(V) are relatively low (Kaplan and Serne, 2000). The adsorption of Np(V) is strongly dependent on pH. Typically, the sorption of Np(V) on iron oxides is negligible at pH values less than 5 and increases rapidly at pH values between 5 and 7. On silicates, the sorption of Np(V) is negligible at pH values less than 6 and increases rapidly at pH values between 6 and 8. This pH dependency is expected for ions present in solution primarily as cations. In carbonate-containing solutions, the adsorption of Np(V) on iron oxides and silicates has been observed to decrease at pH values

greater than 7 to 9 and 8 to 9, respectively. in response to formation of aqueous Np(V) carbonate complexes. Aqueous organic complexes can enhance Np(V) solubility, but sedimentary organic matter is not known to notably affect Np(V) sorption.

Routson, et al. (1977) used batch equilibrium experiments to measure K_d values for Np-237 on a South Carolina soil as a function of the concentrations of dissolved calcium and sodium. The location of the soil sample was not revealed, but it could have been collected from SRS. The soil consisted of 3.6-wt% silt and 37.2-wt% clay and had a moderate exchange capacity (2.5 meq/100 g) and a pH of 5.1. The K_d values were measured in 0.002 and 0.2 M $Ca(NO_3)_2$ solutions and 0.015, 0.03, 0.30, 0.75, and 3.0 M $NaNO_3$ solutions. Measured K_d values for the 0.002 and 0.2 M $Ca(NO_3)_2$ solutions were 0.25 and 0.16 mL/g, respectively. Measured K_d values for the $NaNO_3$ solutions ranged from 0.43 to 0.66 mL/g and decreased with increasing concentration of dissolved sodium.

Nishita, et al. (1981) studied the extractability of Np(V) for several types of soils as a function of pH. For sandy soils with organic matter contents ranging from 2.4 to 5.7 percent, moderate exchange capacities (15 to 20 meq/100 g) and final solution pHs in the range of groundwater pHs expected at SRS (pH 4.1 to 6.8), Nishita, et al. (1981) reported neptunium K_d values ranging from 3 to 95 mL/g.

Sheppard and Thibault (1990) estimated a neptunium default K_d for sand of 5 mL/g (16 observations ranging from 0.5 to 390 mL/g) and a neptunium default K_d for clay of 55 mL/g (4 observations ranging from 0.4 to 2,575). Looney, et al., (1987) specified a reference SRS soil K_d value of 10 mL/g for neptunium from an estimated range of 0.1 to 1,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" neptunium K_d values of 0.2 and 0.6 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" neptunium K_d values of 25 and 35 mL/g for clayey subsurface sediment at the SRS. The neptunium K_d values estimated by Kaplan (2006) were based on a neptunium sorption study using SRS sediments conducted by Sheppard, et al. (1979). As reported by Kaplan (2006), Sheppard, et al. (1979) measured neptunium K_d values of 0.25 and 0.16 mL/g for a sediment with a pH of 5.1 and a clay content of 0.5 wt%. Kaplan (2006) indicated that the pH and clay content of the sediment was lower than assumed for typical sandy and clayey sediments used in the K_d estimations at SRS and, therefore, higher neptunium K_d values were reported.

Measured, estimated, and recommended neptunium K_d values are summarized in Table 2-14. Routson, et al. (1977) reported measured neptunium K_d values ranging from 0.16 to 0.66 mL/g on a South Carolina soil having a pH of 5.1. Nishita, et al. (1981) reported neptunium K_d values ranging from 3 to 95 mL/g for generic sandy soils over the pH range of 4.1 to 6.8. The estimated neptunium K_d value of 5 mL/g of Sheppard and Thibault (1990) for sand and the specified reference neptunium K_d values of 10 mL/g of Looney, et al. (1987) for SRS soil are at the low end of the neptunium K_d range reported by Nishita, et al. (1981). Based on limited site-specific neptunium K_d measurements of Sheppard, et al. (1979), Kaplan (2006) estimated a "reasonably conservative" neptunium K_d value of 0.2 mL/g for sandy subsurface sediments at SRS. This value is in agreement with the measured neptunium K_d values reported by Routson, et al. (1977) on a South Carolina soil with a pH of 5.1. Like uranium, sorption of neptunium is strongly dependent on pH. Groundwater in the upper zone of the Upper Three Runs aquifer has an expected minimum pH of 5.2. Based on these observations, the "reasonably conservative" neptunium K_d value of 0.2 mL/g estimated by Kaplan (2006) is recommended for the upper zone of the Upper Three Runs aquifer. Groundwater in the lower zone of the Upper

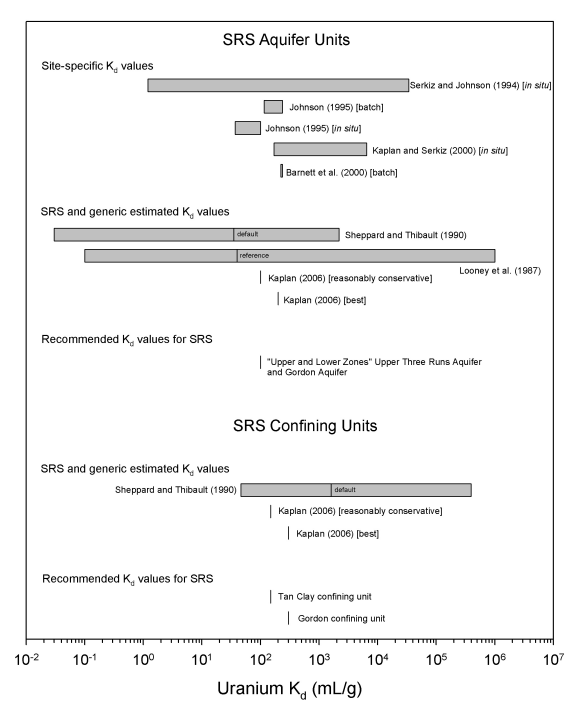


Figure 2-13. Measured and Estimated Site-Specific and Generic Uranium K_d Ranges and Values and Recommended Uranium K_d Values for Aquifer Units and Confining Units at SRS (Table 2-13). Recommended Uranium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

Three Runs aquifer and the Gordon aquifer have expected pH ranges of 6.0 to 7.7 and 5.7 to 7.6, respectively. Neptunium sorption on silicates increases with increasing pH with maximum sorption occurring at a pH of about 8.0 (Bertetti, et al., 1998). Therefore, the "best" neptunium K_d value of 0.6 mL/g estimated by Kaplan (2006) is recommended for the lower zone of the Upper Three Runs aguifer and the Gordon aguifer at SRS.

No site-specific neptunium sorption data for clay confining units at SRS were found in the literature. Sheppard and Thibault (1990) estimated a default neptunium $K_{\rm d}$ value of 55 mL/g for clay. Kaplan (2006) estimated a "reasonably conservative" neptunium $K_{\rm d}$ value of 25 mL/g for clayey subsurface sediments at SRS. This "reasonably conservative" value is recommended for clay confining units at SRS.

The position of the neptunium K_d values recommended in this review with respect to measured and estimated site-specific and generic neptunium K_d ranges and values is graphically displayed in Figure 2-14. This figure illustrates the uncertainty associated with the neptunium K_d values recommended for SRS aquifer units and clay confining units. The recommended neptunium K_d values are associated with high uncertainty due to limited site-specific sorption data. For example, the recommended neptunium K_d value of 0.2 mL/g for the upper zone of the Upper Three Runs Aquifer based on reported neptunium K_d values measured by Routson, et al. (1977) on South Carolina soils and two neptunium K_d values measured by Sheppard, et al. (1979) on SRS sediments. No site-specific neptunium sorption data exists for SRS confining units.

2.4.13 Plutonium

In the range of pH and conditions typically encountered in natural waters, Pu-239 can exist in all 4 oxidation states, namely +3, +4, +5, and +6. Under oxidizing conditions, Pu(IV), Pu(V), and Pu(VI) are common, whereas under reducing conditions, Pu(III) and Pu(IV) exist. Plutonium is known to adsorb onto soil components such as clays, oxides, hydroxides, oxyhydroxides, aluminosilicates, and organic matter. Depending on the properties of the substrates, pH, and the composition of solutions, plutonium would adsorb with affinities varying from low ($K_d = 10 \text{ mL/g}$) to extremely high ($K_d = 300,000 \text{ mL/g}$) (Baes and Sharp, 1983; Thibault, et al.,1990). Under conditions of low pH and high concentrations of dissolved organic carbon, it appears that plutonium-organic complexes may control adsorption and mobility of plutonium. At neutral to alkaline pH (>7), dissolved plutonium forms very strong hydroxy–carbonate mixed ligand complexes, resulting in desorption and increased mobility in the environment. Studies have shown the potential importance of colloid-facilitated transport of plutonium in soil systems (Penrose, et al., 1990). Dai, et al. (2002) and Kaplan, et al. (1994) have found that small fractions of total mobile plutonium in SRS sediments are associated with colloids.

A number of studies indicate that iron hydroxides adsorb and reduce Pu(V) and Pu(VI) to a tetravalent state at the solid surface. Fjeld, et al. (2003) have described a conceptual model where plutonium transport through sediment columns is controlled by the net rate of reduction of adsorbed Pu(V) to adsorbed Pu(IV). Field studies have supported this model in SRS vadose zone sediments (Kaplan, et al., 2004). In this model, oxidized plutonium can mobilize as Pu(V), and the Pu(V) can readsorb onto iron-bearing minerals and be slowly reduced to Pu(IV). In this way, kinetic control of plutonium transport can allow a small fraction to move while the bulk of

Table 2-14. Measured, Estimated, and Recommended Neptunium K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Neptunium K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Neptunium K _d (mL/g)*	Generic Estimated Neptunium K _d (mL/g)†	Recommended Neptunium K _d (mL/g)
Upper Three Runs Aquifer: Upper	0.16-0.66‡ 0.16-0.25§	Batch Batch	0.2/0.6	5	0.2
Tan Clay Confining Unit	_		25/35	55	25
Upper Three Runs Aquifer: Lower	_		0.2/0.6	5	0.6
Gordon Confining Unit	_		25/35	55	25
Gordon Aquifer	_		0.2/0.6	5	0.6

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990. ‡Routson, R.C., G. Jansen, and A.V. Robinson. "²⁴¹Am, ²³⁷Np, and ⁹⁹Tc Sorption on Two United States Subsoils

[‡]Routson, R.C., G. Jansen, and A.V. Robinson. "²⁴¹Am, ²³⁷Np, and ⁹⁹Tc Sorption on Two United States Subsoils from Differing Weathering Intensity Areas." *Health Physics*. Vol. 33. pp. 311–317. 1977.

[§]Sheppard, J.C., M.J. Campbell, J.A. Kittrick, and T.L. Hardt. "Retention of Neptunium, Americium, and Curium by Diffusible Soil Particles." *Environmental Science and Technology*. Vol. 13. pp. 680–684. 1979.

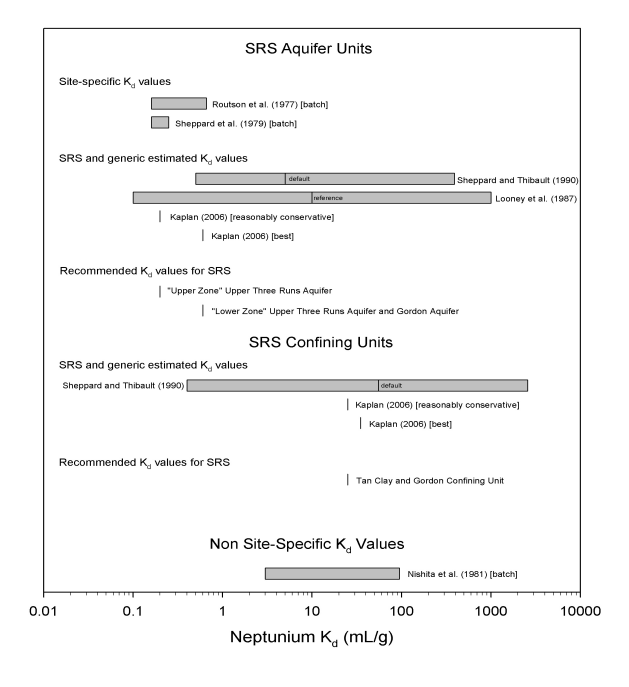


Figure 2-14. Measured and Estimated Site-Specific and Generic Neptunium $\rm K_d$ Ranges and Values and Recommended Neptunium $\rm K_d$ Values for Aquifer Units and Confining Units at SRS (Table 2-14). Recommended Neptunium $\rm K_d$ Values Are Associated With High Uncertainty Due to Limited Site-Specific Sorption Data.

plutonium is largely immobile. This type of nonequilibrium behavior is difficult to capture with a K_d approach, but K_d can be used if the system is assumed to have reached steady state with respect to plutonium redox kinetics.

Prout (1958) studied adsorption of plutonium +3, +4, and +6 redox states on a Savannah River Plant soil as a function of pH. The calculated K_d ranges were from <10 to >10,000 mL/g for Pu(III); ~100 to ~10,000 mL/g for Pu(IV); and <10 to ~3,000 mL/g for Pu(VI). The large range in values is due to the broad pH range studied. Maximum K_d values were observed between pH values of about 6.5 to 8.5. Because the initial concentrations of plutonium used in these experiments were about 10^{-6} M, precipitation reactions may have accounted for some of the observed removal of plutonium from solution.

Hoeffner (1985) summarized plutonium sorption studies conducted to determine the performance of the low-level waste burial ground at SRS. The majority of studies used a Savannah River Plant soil taken from depths 2.5–3.0 m [8–10 ft] below grade and a low ionic strength groundwater representative of the burial ground (pH 5.0). Hoeffner (1985) reported expected ranges of Pu(III), Pu(IV), and Pu(VI) K_d values. Over a pH range of 3.4 to 7.3 and for a groundwater of typical composition at SRS (pH 4.7) (i) Pu(III) K_d values are expected to range from 800 to >10,000 mL/g and a K_d value of 8,000 mL/g is expected; (ii) Pu(IV) K_d values are expected to range from 120 to 7,100 mL/g; and a K_d of 150 mL/g is expected; and (iii) Pu(VI) K_d values are expected to range from 7 to 250 mL/g, and a K_d value of 9 mL/g is expected.

Powell, et al. (2002) performed Pu(V) and Pu(IV) batch tests on sediments collected from SRS as a function of pH and contact time. Sediments were selected to represent end members with respect to sediment properties existing on SRS that were likely to influence plutonium sorption. Characteristics of the sediments are listed in Table 2-15a. Powell, et al. (2002) did not specify the location, depth, or hydrostratigraphic unit from which the sediments were collected. However, based on particle size characteristics, the sandy subsurface and surface sediments are most comparable to aquifer units beneath the GSA, and the clayey subsurface and surface sediments are most comparable to confining units beneath the GSA. All experiments were performed using 0.02 M NaClO₄ solutions. For each sediment, K_d values for Pu(IV) were determined after 24 hours of reaction, and K_d values for Pu(V) were determined after 24 hours and 33 days of reaction. The range of plutonium K_d values measured for all the sediments are listed in Table 2-15b.

Results of the plutonium sorption experiments of Powell, et al. (2002) indicated a general trend of increasing Pu(IV) sorption with increasing pH for all the sediments. Maximum sorption for each sediment was observed at higher pHs (pH >6.0). The K_d values for Pu(V) shifted with time (i.e., from 24 hours to 33 days) to resemble those of Pu(IV) for all the sediments studied, suggesting reduction of Pu(V) to Pu(IV). Stronger sorption to the surface clayey sediment was attributed to the greater surface area and organic matter concentration of this sediment. The plutonium sorption data of Powell, et al. (2002) were analyzed with respect to the pH range of groundwater expected at SRS (pH 5.2 to 7.7). The range of plutonium K_d values measured for all the sediments at this pH range is listed in Table 2-15c. Examination of the data indicates that the Pu(IV) K_d values at 24 hours resemble the Pu(V) K_d values at 33 days except for the surface clayey sediment. For the surface clayey sediment, Pu(V) K_d values are much greater at 33 days than Pu(IV) K_d values at 24 hours. Stronger sorption to the surface clayey sediment can again be attributed to the greater surface area and organic matter concentration of this sediment.

Table 2-15a. Ch	Table 2-15a. Characteristics of Sediments in Plutonium Batch Sorption Experiments*					
Sediment	Sand/Silt/Clay (wt%)	Surface Area (m²/g)	рН	Organic matter (wt%)		
Subsurface Sandy	97/2/1	1.27	5.10	<0.01		
Subsurface Clayey	58/30/12	15.31	4.55	Not applicable		
Surface Sandy	93/6/1	1.37	4.70	<0.01		
Surface Clayey	67/25/8	9.24	5.83	3.2		

^{*}Used by Powell, B.A., R.A. Fjeld, J.T. Coates, D.I. Kaplan, and S.M. Serkiz. "Plutonium Oxidation State Geochemistry in the SRS Subsurface Environment (U)." WSRC–TR–2003–00035. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company, Savannah River Site. 2002.

Table 2-15b. Range of Plutonium K_d Values Measured With Respect to Sediment Type, Plutonium Oxidation State, and Reaction Time. The pH Range Over Which Measurements Were Collected for Each Sediment Type and Oxidation State Is Also Shown.*

Sediment	Plutonium Oxidation State	pH Range	Plutonium K _d Range (mL/g)
Subsurface Sandy	Pu(IV) at 24 hours	1.91–9.28	8–9,300
Subsurface Sandy	Pu(V) at 24 hours	1.81–6.83	6–1,100
Subsurface Sandy	Pu(V) at 33 days	1.91–5.62	3–4,900
Subsurface Clayey	Pu(IV) at 24 hours	1.71–7.91	90–50,000
Subsurface Clayey	Pu(V) at 24 hours	1.67–8.95	4–660
Subsurface Clayey	Pu(V) at 33 days	1.63-8.89	60–10,000
Surface Sandy	Pu(IV) at 24 hours	2.05–10.47	100–1,100
Surface Sandy	Pu(V) at 24 hours	1.90-8.26	7–110
Surface Sandy	Pu(V) at 33 days	1.91–7.86	150–20,000
Surface Clayey	Pu(IV) at 24 hours	1.78–9.95	500–4,600
Surface Clayey	Pu(V) at 24 hours	1.68–9.69	300–2,600
Surface Clayey	Pu(V) at 33 days	1.68–9.15	500–33,000

^{*}Measured in Powell, B.A., R.A. Fjeld, J.T. Coates, D.I. Kaplan, and S.M. Serkiz. "Plutonium Oxidation State Geochemistry in the SRS Subsurface Environment (U)." WSRC–TR–2003–00035. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company, Savannah River Site. 2002.

Sheppard and Thibault (1990) estimated a plutonium default $K_{\rm d}$ for sand of 550 mL/g (39 observations ranging from 27 to 3,600 mL/g) and a plutonium default $K_{\rm d}$ for clay of 5,100 mL/g (18 observations ranging from 316 to 190,000). Looney, et al. (1987) estimated a reference SRS soil $K_{\rm d}$ value of 100 mL/g for plutonium from an estimated range of 10 to 100,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" plutonium $K_{\rm d}$ values of 180 and 270 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" plutonium $K_{\rm d}$ values of 2,950 and 5,900 mL/g for clayey subsurface sediment at the SRS. The plutonium $K_{\rm d}$ values estimated by Kaplan (2006) were based on the plutonium sorption data of Powell, et al. (2002).

Measured, estimated, and recommended plutonium K_d values are summarized in Table 2-15d. Hoeffner (1985) reported expected plutonium K_d ranges for Pu(III) (800 to >10,000 mL/g),Pu(IV) (120 to 7,100 mL/g), and Pu(VI) (7 to 250 mL/g) for SRS soil over a pH range of 3.4 to 7.3. For a groundwater of typical composition at the SRS burial ground (4.7), Hoeffner (1985) reports expected plutonium K_d value of 8,000 mL/g for Pu(III), 150 mL/g for Pu(IV), and 9 mL/g for

Table 2-15c. Range of Plutonium K_d Values Measured in Sediments Over the pH Range of Groundwater Expected at the Savannah River Site (pH 5.2 to 7.7)*

Sediment	Plutonium Oxidation State	pH Range	Plutonium K _d Range (mL/g)
Subsurface Sandy	Pu(IV) at 24 hrs	5.85–7.70	390–2,800
Subsurface Sandy	Pu(V) at 24 hrs	5.30-6.83	9–1,100
Subsurface Sandy	Pu(V) at 33 days	5.10–5.62	600–4,900
Subsurface Clayey	Pu(IV) at 24 hrs	4.81–6.68	2,100–18,000
Subsurface Clayey	Pu(V) at 24 hrs	5.30-6.02	50–440
Subsurface Clayey	Pu(V) at 33 days	4.95–5.68	4,700–10,000
Surface Sandy	Pu(IV) at 24 hrs	6.73	130
Surface Sandy	Pu(V) at 24 hrs	5.26–6.98	20–110
Surface Sandy	Pu(V) at 33 days	5.26–6.98	270–2,900
Surface Clayey	Pu(IV) at 24 hrs	6.09–6.83	1,000–4,600
Surface Clayey	Pu(V) at 24 hrs	5.10–6.89	1,300–2,600
Surface Clayey	Pu(V) at 33 days	4.67–6.81	8,200–33,000

*Measured in Powell, B.A., R.A. Fjeld, J.T. Coates, D.I. Kaplan, and S.M. Serkiz. "Plutonium Oxidation State Geochemistry in the SRS Subsurface Environment (U)." WSRC-TR-2003-00035. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company, Savannah River Site. 2002.

Table 2-15d. Measured, Estimated, and Recommended Plutonium K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Plutonium K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Plutonium K _d (mL/g)*	Generic Estimated Plutonium K _d (mL/g)†	Recommended Plutonium K _d (mL/g)
Upper Three Runs Aquifer: Upper					
Pu(III)	10–10,000‡ 800–10,000§	Batch Batch			
Pu(IV)	100-10,000‡ 120-7,100§ 8-50,000¶	Batch Batch Batch			
Pu(V)	3-33,000¶	Batch			
Pu(VI) Pu(total)	10–3,000‡ 7–250§	Batch Batch	180/270	550	250
Tan Clay Confining Unit	_		2,950/5,900	5,100	2,500
Upper Three Runs Aquifer: Lower	_		180/270	550	250
Gordon Confining Unit	_		2,950/5,900	5,100	2,500
Gordon Aquifer	_		180/270	550	250

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

¶Powell, B.A., R.A. Fjeld, J.T. Coates, D.I. Kaplan, and S.M. Serkiz. "Plutonium Oxidation State Geochemistry in the SRS Subsurface Environment (U)." WSRC–TR–2003–00035. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company, Savannah River Site. 2002.

 $[\]dagger$ Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_d s, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

[‡]Prout, W.E. "Adsorption of Radioactive Wastes by Savannah River Plant Soil." *Soil Science*. Vol. 84. pp. 13–17. 1958.

[§]Hoeffner, S.L. "Radionuclide Sorption on Savannah River Plant Burial Ground Soil—A Summary and Interpretation of Laboratory Data." DP–1702. Aiken, South Carolina: E.I. du Pont de Nemours and Company, Savannah River Laboratory. 1985.

Pu(VI). Powell, et al. (2002) reported measured plutonium K_d values for four different soil types from SRS as a function of pH and contact time. Over the pH range expected at SRS (5.2 to 7.7), the plutonium K_d range for Pu(IV) at 24 hours ranged from 130 to 18,000 mL/g, the plutonium K_d range for Pu(V) at 24 hours ranged from 9 to 2,600 mL/g, and the plutonium K_d range for Pu(V) at 33 days ranged from 270 to 33,000 mL/g. Except for the surface clayey sediment, the K_d range for Pu(IV) at 24 hours resembled the K_d range for Pu(V) at 33 days. The expected plutonium K_d value reported by Hoeffner (1985) for Pu(IV) (150 mL/g) is at the low end of the range of plutonium K_d values measured by Powell, et al. (2002) for $Pu(\tilde{IV})$ at 24 hours (130 to 18,000 mL/g) and for Pu(V) at 33 days (270 to 33,000 mL/g). The default plutonium K_d value of 550 mL/g estimated by Sheppard and Thibault (1990) for sand and the reference plutonium K_d value of 100 mL/g specified by Looney, et al. (1987) for soil at SRS lie at the low end of the Pu(IV) K_d range reported by Hoeffner (1985) (120 to 7,100 mL/g) and at the low end of the range of plutonium K_d values measured by Powell, et al. (2002) for Pu(IV) at 24 hours (130 to 18,000 mL/g) and for Pu(V) at 33 days (270 to 33,000 mL/g). Based on these observations and considering the apparently rapid rate of surface reduction of sorbed Pu(V), a plutonium K_d value of 250 mL/g is recommended for all the aquifer units at SRS. This value is below the lowest observed Pu(IV) and longer term Pu(V) K_ds observed by Powell, et al. (2002) for subsurface sandy sediments in the pH range of interest (Table 2-15c). This value is also in agreement with the "reasonably conservative" (180 mL/g) and "best" (270 mL/g) plutonium K_d values estimated by Kaplan (2006), which are based on the plutonium sorption data of Powell, et al. (2002).

Sheppard and Thibault (1990) estimated a default plutonium K_d of 5,100 mL/g for clay. Powell, et al. (2002) reports measured Pu(IV) and Pu(V) K_d ranges on a subsurface clayey and a surface clayey sediment collected at SRS. Although these sediments were collected in the upper zone of the Upper Three Runs aquifer, their mineralogic compositions and high surface areas may be comparable to clay confining units at SRS (see Table 2-15a). Over the pH range expected at SRS (5.2 to 7.7), the plutonium K_d range for Pu(IV) in these clayey sediments was 1,000 to 18,000 mL/g at 24 hours. The plutonium K_d range for Pu(V) in the clayey sediments was 4,700 to 33,000 mL/g at 33 days. Based on the Pu(IV) K_d range at 24 hours and the Pu(V) K_d range at 33 days and considering the apparently rapid rate of surface reduction of sorbed Pu(V), a plutonium K_d of 2,500 mL/g is recommended for clay confining units at SRS. This value is in agreement with the "reasonably conservative" plutonium K_d value of 2,950 mL/g estimated by Kaplan (2006), which is also based on the plutonium sorption data of Powell, et al. (2002).

In the performance assessment for the salt waste facility (Cook, et al., 2005), DOE employed a plutonium sorption model that considered two composite oxidation state species with different sorption properties—Pu(III/IV) and Pu(V/VI)—and calculated species abundance based on kinetic oxidation-reduction rates (Kaplan, 2004). If such an approach were employed, it would be appropriate to use the lowest Powell, et al. (2002) 24-hour sorption coefficients for Pu(IV) and Pu(V) for the pH range of interest for the two composite species (Table 2-15c). Kaplan (2004) summarized the Powell, et al. (2002) data; if Kaplan's values for appropriately bounding pH (5.25) were lower than those suggested by Table 2-15c, the lower value was selected. Thus, for the plutonium oxidation state model, the following K_d values are recommended:

Pu(III/IV) in aquifers (subsurface sandy) = 350 mL/g (Kaplan, 2004, at pH 5.25)
 Pu(V/VI) in aquifers (subsurface sandy) = 9 mL/g (Table 2-15c)

Pu(III/IV) in confining units (subsurface clayey) = 2,100 mL/g (Table 2-15c)
 Pu(V/VI) in confining units (subsurface clayey) = 50 mL/g (Table 2-15c)

The position of the plutonium K_d values recommended in this review with respect to measured and estimated site-specific and generic plutonium K_d ranges and values is graphically displayed in Figure 2-15. This figure illustrates the uncertainty associated with the plutonium K_d values recommended for SRS aquifer units and clay confining units. Due to available site-specific plutonium sorption data (Hoeffner, 1985; Powell, et al., 2002), the recommended plutonium K_d values are associated with low uncertainty.

2.4.14 Americium

Am-241 can exist in the +3, +4, +5, and +6 valance states, but americium predominantly occurs as Am(III) in natural environments. Sorption studies indicate that Am(III) readily sorbs to minerals, crushed rock, and soil materials and, along with plutonium, is considered one of the most immobile actinide elements in the environment (Lu, et al., 1998; Ticknor, et al., 1996). Based on studies of Am(III) on single mineral phases and crushed rock, the adsorption of Am(III) is strongly pH dependent and increases with increasing pH with peak adsorption occurring between pH values of 5 and 6. This observed pH dependence is expected, because the dominant aqueous species of americium in the pH range of natural waters are primarily Am³⁺ and cationic carbonate complexes at acidic and basic pH values, respectively. High levels of soil organic matter can enhance americium mobility (Artinger, et al., 1998; Schuessler, et al., 2000; Ticknor, et al., 1996). In soil environments, clay minerals are expected to control americium sorption, and K_d values are expected to correlate with CEC. Studies have shown the potential importance of colloid-facilitated transport of americium in soil systems (Penrose, et al., 1990). Kaplan, et al. (1994) found that small fractions of total mobile americium in SRS sediments are associated with colloids.

Routson, et al., (1977) used batch equilibrium experiments to determine K_d values for Am-241 on a subsoil sample from South Carolina as a function of solution concentration of calcium (Ca^{2+}) and of sodium (Na^+). The exact location of the South Carolina soil used in this study was not revealed, but it could have been collected at SRS. The soil had a moderate exchange capacity (2.5 meq/100 g), a pH of 5.1, and consisted of 3.6-wt% silt and 37.2-wt% clay. The americium K_d values were determined in 0.002, 0.02, 0.05, 0.1, and 0.2 M $Ca(NO_3)_2$ solutions and 0.015, 0.03, 0.3, 0.75, and 3.0 M $NaNO_3$ solutions. The pH values for selected samples of the americium solutions in the calcium and sodium systems were 7.1 and 6.1. For the solution concentrations used in the experiments, the K_d values for americium ranged from 1.0 to 67 mL/g as a function of dissolved calcium and 1.6 to 280 mL/g as a function of dissolved sodium. K_d values decreased in all cases with increasing solution concentrations of calcium and sodium.

Sheppard and Thibault (1990) suggested an americium default K_d for sand of 1,900 mL/g (29 observations ranging from 8.2 to 300,000 mL/g) and an americium default K_d for clay of 8,400 mL/g (11 observations ranging from 25 to 400,000). Looney, et al. (1987) estimated a reference SRS soil K_d value of 100 mL/g for americium from an estimated range of 1 to 100,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" americium K_d values of 550 and 1,100 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" americium K_d values of 4,250 and 8,500 mL/g for clayey subsurface sediment at the SRS. The Kaplan (2006) estimations are based on reported cerium K_d values measured on

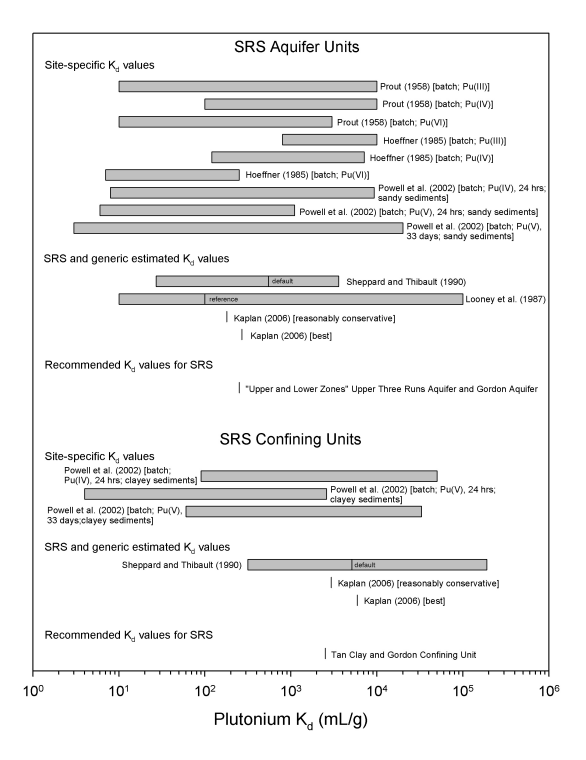


Figure 2-15. Measured and Estimated Site-Specific and Generic Plutonium K_d Ranges and Values and Recommended Plutonium K_d Values for Aquifer Units and Confining Units at SRS (Table 2-15d). Recommended Plutonium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

SRS sandy and clayey sediments by Kaplan and Serkiz (2004). Kaplan (2006) used cerium as a chemical analog for americium; both cerium and americium are trivalent cations. At a pH of 5.3, Kaplan and Serkiz (2004) reported a cerium K_d of 1,220 mL/g for SRS sandy sediment and a cerium K_d of 8,697 mL/g for SRS clayey sediment. Kaplan and Serkiz (2004) also measured europium K_d values on the SRS sandy and clayey sediments. Both europium and cerium are trivalent cations and had similar K_d values for the given SRS sediments. At a pH of 5.3, Kaplan and Serkiz (2004) reported a europium K_d of 1,168 mL/g for SRS sandy sediment and a europium K_d of 9,020 mL/g for the SRS clayey sediment.

Measured, estimated, and recommended americium K_d values are summarized in Table 2-16. Routson, et al. (1977) reported measured americium K_d values ranging from 1.0 to 280 for a soil from South Carolina as a function of Ca2+ and Na+ solution concentration. The pH values of selected samples of solutions used in the americium sorption experiments were 6.1 and 7.1. Routson, et al. (1977) reported that measured americium K_d values increased with decreasing calcium and sodium concentration. At the low concentrations of Ca2+ and Na+ expected in groundwater at SRS, americium K_d values in sediments at SRS would most likely correspond to the high end of the americium K_d range reported by Routson, et al. (1977). The default americium K_d of 1,900 mL/g for sand estimated by Sheppard and Thibault (1990) is much greater than the reported americium K_d values of Routson, et al. (1977). The reference americium K_d value of 100 mL/g estimated by Looney, et al. (1987) for soil at SRS is within the range of measured americium K_d values reported by Routson, et al. (1977). Kaplan (2006) estimated a "reasonably conservative" americium K_d value of 550 mL/g for sandy subsurface sediments at SRS based on measured cerium K_d values reported by Kaplan and Serkiz (2004). The "reasonably conservative" americium K_d value of Kaplan (2006) is slightly higher than the americium K_d values reported by Routson, et al. (1997). The mineral compositions of the aquifer units at SRS are not significantly different and, thus, should not affect americium sorption. Sorption of americium is strongly pH dependent with maximum sorption occurring at pHs between 5 and 6. These pHs correspond to the minimum pHs of all the aguifer units at SRS. As indicated above, due to the low Ca²⁺ and Na⁺ content of the groundwater at SRS, americium K_d values for sediments at SRS would most likely correspond to the high end of the americium K_d range reported by Routson, et al. (1977). Therefore, an americium K_d value of 250 is recommended for the upper zone of the Upper Three Runs aguifer at SRS. Groundwater in the lower zone of the Upper Three Runs aguifer and the Gordon aguifer have minimum expected pHs of 6.0 and 5.7, respectively, but in the absence of a more quantitative pH dependence, the same 250 mL/g value is recommended for the lower zone of the Upper Three Runs aguifer and the Gordon aguifer at SRS. The use of the high Sheppard and Thibault (1990) value of 1,900 mL/g would be qualitatively inconsistent with observations of some degree of dissolved americium mobility in the F Area (Kaplan, et al., 1994).

Sheppard and Thibault (1990) estimated a default americium K_d of 8,400 mL/g for clay. Kaplan (2006) estimated a "reasonably conservative" americium K_d value of 4,250 mL/g for clayey subsurface sediments at SRS based on measured cerium K_d values reported by Kaplan and Serkiz (2004). Both cerium and americium are trivalent cations and Kaplan (2006) used cerium as a chemical analog for americium. Lacking site-specific americium sorption data for clay sediments, the "reasonably conservative" americium K_d value of 4,250 mL/g estimated by Kaplan (2006) is recommended for clay confining units at SRS.

Table 2-16. Measured, Estimated, and Recommended Americium K_d Values for Aquifer Sediments and Clay Confining Units at the Savannah River Site

	Measured Americium K _d (mL/g)	Test Method	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Americium K _d (mL/g)*	Generic Estimated Americium K _d (mL/g)†	Recommended Americium K _d (mL/g)
Upper Three Runs Aquifer: Upper	1–280‡	Batch	550/1,100	1,900	250
Tan Clay Confining Unit	_		4,250/8,500	8,400	4,250
Upper Three Runs Aquifer: Lower	_		550/1,100	1,900	250
Gordon Confining Unit	_		4,250/8,500	8,400	4,250
Gordon Aquifer	_		550/1,100	1,900	250

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC-TR-2006-00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

The position of the americium K_d values recommended in this review with respect to measured and estimated site-specific and generic americium K_d ranges and values is graphically displayed in Figure 2-16. This figure illustrates the uncertainty associated with the americium K_d values recommended for SRS aguifer units and clay confining units. The recommended americium K_d values are associated with high uncertainty due to the lack of site-specific sorption data.

2.4.15 Curium

Cm-243 is a transuranic (actinide) element, and exists in the +3 oxidation state in natural waters. Curium is quite similar geochemically to americium (Guillaumont, et al., 2003). The dominant aqueous species of curium in the pH range of natural waters are primarily cations such as Cm³⁺ and Cm(III) carbonate complexes at acidic and basic pH values, respectively. Available curium sorption studies indicate that sorption of curium is strongly pH dependent and increases with increasing pH with maximum adsorption occurring between pH values of 5 and 6. Compared to other actinides, curium is considered to be immobile in soil environments, and exhibits high K_d values. However, the tendency of curium, like americium, to strongly adsorb to

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, Kds, for Four Major Soil

Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990. ‡Routson, R.C., G. Jansen, and A.V. Robinson. "²⁴¹Am, ²³⁷Np, and ⁹⁹Tc Sorption on Two United States Subsoils from Differing Weathering Intensity Areas." Health Physics. Vol. 33. pp. 311-317. 1977.

soil particles indicates that there is potential for colloid-facilitated transport of curium. Kaplan, et al. (1994) found that small fractions of total mobile curium in SRS sediments are associated with colloids.

Using cerium (a trivalent cation) as an analog to curium, Kaplan and Serkiz (2000) estimated curium K_d values ranging from 255 to 4,074 mL/g on contaminated sediments at SRS. Kaplan and Serkiz (2000) performed desorption measurements to measure cerium K_d values on shallow contaminated sediments from the TNX Outfall Operable Unit at SRS. The desorption K_d values were based on extractable portions that comprise the exchangeable, organic-bound, amorphous Fe–oxide bound, and water extractable fractions. Basing the K_d estimates on these extractable fractions instead of the more easily determined total concentrations in the sediment, more conservative and technically defensible K_d values were measured. The contaminated sediments were collected at depths of 0 to 0.6 m [0 to 2 ft] by hand auger and were composed dominantly of sand and silt (i.e., sand and silt contents ranging from 80 to 94 wt%). The sediment pH of the sediments ranged from 4.00 to 5.78. Measured cerium K_d values ranged from 255 to 4,074 mL/g.

Nishita, et al. (1981) studied the extractability of Cm-244 for several types of soils as a function of pH. For sandy soils with organic matter contents ranging from 2.4 to 5.7 wt%, moderate CEC (15 to 20 meq/100 g) and final solution pHs ranging from 4.39 to 6.17, Nishita, et al. (1981) reported curium K_d values ranging from 186 to 23,350 mL/g.

Sheppard and Thibault (1990) suggested a curium default K_d for sand of 4,000 mL/g (2 observations ranging from 780 to 22,970 mL/g) and a curium default K_d for clay of 6,000 (based on a soil-to-plant concentration ratio). Looney, et al., (1987) estimated a reference SRS soil K_d value of 3,160 mL/g for curium from an estimated range of 100 to 100,000 mL/g. Kaplan (2006) estimated "reasonably conservative" and "best" curium K_d values of 550 and 1,100 mL/g for sandy subsurface sediment and "reasonably conservative" and "best" curium K_d values of 4,250 and 8,500 mL/g for clayey subsurface sediment at the SRS. The Kaplan (2006) estimations were based on reported cerium K_d values measured on SRS sandy and clayey sediments by Kaplan and Serkiz (2004). Due to similarities in aqueous chemistry, the sorptive characteristics of curium on sandy and clayey sediments at SRS are expected to be very similar to those of cerium. At a pH of 5.3, Kaplan and Serkiz (2004) reported a cerium K_d of 1,220 mL/g for SRS sandy sediment and a cerium K_d of 8,697 mL/g for SRS clayey sediment. Kaplan and Serkiz also measured europium K_d values on the SRS sandy and clayey sediments. Both europium and cerium are trivalent cations and had similar K_d values for the given SRS sediments. At a pH of 5.3, Kaplan and Serkiz (2004) reported a europium K_d of 1,168 mL/g for SRS sandy sediment and a europium K_d of 9,020 mL/g for the SRS clayey sediment.

Recommended and estimated curium K_d values are summarized in Table 2-17. Using cerium (a trivalent cation) as an analog to curium, Kaplan and Serkiz (2000) estimated curium K_d values ranging from 255 to 4,074 mL/g on contaminated sediments at SRS. Likewise, Kaplan (2006) estimated a "reasonably conservative" curium K_d value of 550 mL/g for sandy subsurface sediments at SRS based on measured cerium K_d values reported by Kaplan and Serkiz (2004). Nishita, et al. (1981) reported curium K_d values ranging from 186 to 23,350 mL/g for generic soils over the pH range of 4.39 to 6.17. The default curium K_d value of 4,000 mL/g estimated by Sheppard and Thibault (1990) for sand and the reference curium K_d of 3,160 mL/g specified

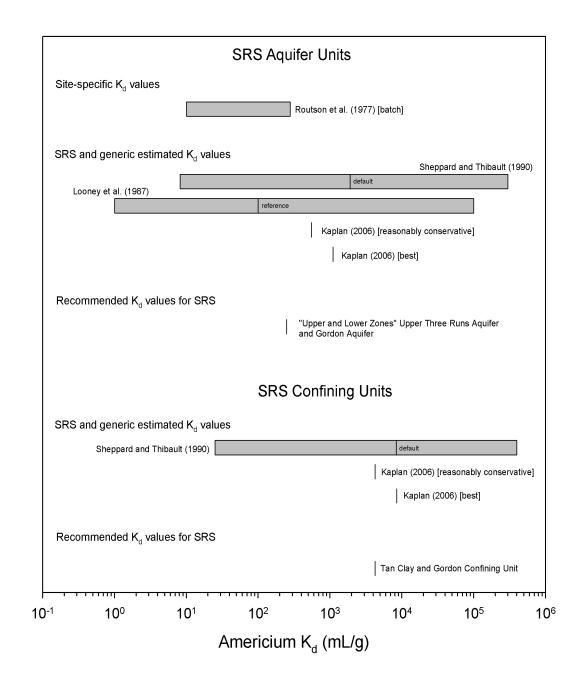


Figure 2-16. Measured and Estimated Site-Specific and Generic Americium K_d Ranges and Values and Recommended Americium K_d Values for Aquifer Units and Confining Units at SRS (Table 2-16). Recommended Americium K_d Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

by Looney, et al. (1987) for soil at SRS are within the curium K_d range reported by Nishita, et al. (1981). However, these values are at the high end of the curium K_d ranges estimated by Kaplan and Serkiz (2000) for SRS contaminated soils (255 to 4,074 mL/g) and are greater than the "reasonably conservative" curium K_d value of 550 mL/g estimated by Kaplan (2006). Based on the estimated curium K_d values of Kaplan and Serkiz (255 to 4,074 mL/g) and considering the "reasonably conservative" curium K_d value of 550 mL/g estimated by Kaplan (2006), a curium K_d value of 300 mL/g is recommended for aquifer units at SRS. The use of the high Sheppard and Thibault (1990) value of 4,000 mL/g for the aquifer units would be qualitatively inconsistent with observations of some degree of dissolved curium mobility in the F Area (Kaplan, et al., 1994).

Sheppard and Thibault (1990) estimated a default curium K_d of 6,000 mL/g for clay. Kaplan (2006) estimated a "reasonably conservative" curium K_d value of 4,250 mL/g for clayey subsurface sediments at SRS based on measured cerium K_d values reported by Kaplan and Serkiz (2004). Both cerium and curium are trivalent cations and Kaplan (2006) used cerium as a chemical analog for curium. Lacking site-specific curium sorption data for clay sediments, the "reasonably conservative" curium K_d value of 4,250 mL/g estimated by Kaplan (2006) is recommended for clay confining units at SRS.

Table 2-17. Estir	ommended Curium hing Units at the Sava	Հ _ս Values for Aquifer Տ annah River Site	Sediments and
	Savannah Biyar		

	Measured Curium K _d (mL/g)	Savannah River Site-Specific "Reasonably Conservative"/ "Best" Estimated Curium K _d (mL/g)	Generic Estimated Curium K _d (mL/g)†	Recommen ded Curium K _d (mL/g)
Upper Three Runs Aquifer: Upper	_	550/1,100	4,000	300
Tan Clay Confining Unit	_	4,250/8,500	6,000	4,250
Upper Three Runs Aquifer: Lower	_	550/1,100	4,000	300
Gordon Confining Unit	_	4,250/8,500	6,000	4,250
Gordon Aquifer	_	550/1,100	4,000	300

^{*}Kaplan, D.I. "Geochemical Data Package for Performance Assessment Calculations Related to the Savannah River Site (U)." WSRC–TR–2006–00004. Rev. 0. Aiken, South Carolina: Westinghouse Savannah River Company. 2006.

[†]Sheppard, M.I. and D.H. Thibault. "Default Soil Solid/Liquid Partition Coefficients, K_ds, for Four Major Soil Types: A Compendium." *Health Physics*. Vol. 59, No. 4. pp. 471–482. 1990.

The position of the curium K_d values recommended in this review with respect to measured and estimated site-specific and generic curium K_d ranges and values is graphically displayed in Figure 2-17. This figure illustrates the uncertainty associated with the curium K_d values recommended for SRS aquifer units and clay confining units. The recommended curium K_d values are associated with high uncertainty due to the lack of site-specific sorption data.

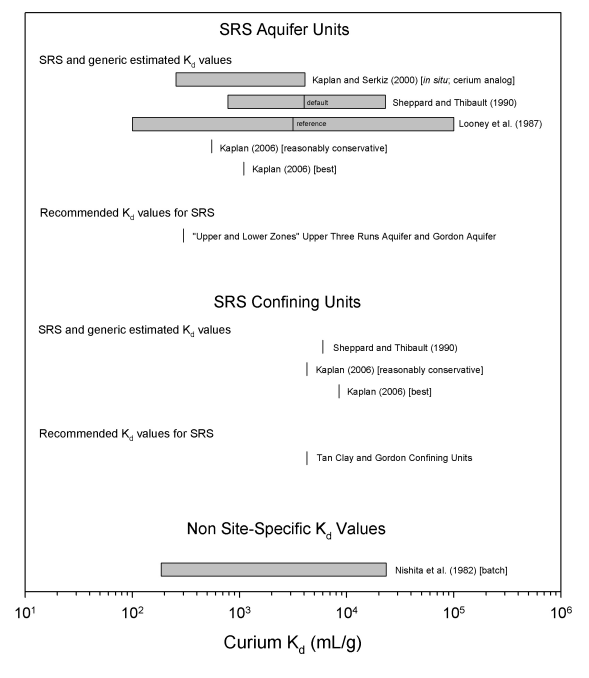


Figure 2-17. Measured and Estimated Site-Specific and Generic Curium K_d Ranges and Values and Recommended Curium K_d Values for Aquifer Units and Confining Units at SRS (Table 2-17). Recommended Curium K_d Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

3 IDAHO NATIONAL LABORATORY

3.1 Waste Management Background

At Idaho National Laboratory (INL), the U.S. Department of Energy (DOE) has used tanks at the Idaho Nuclear Technology and Engineering Center (INTEC) Tank Farm Facility to store a variety of radioactive liquid wastes. DOE has cleaned the tanks to remove the bulk of the waste and now plans to close the facility by stabilizing any residual radioactive materials in the tanks by filling them, in place, with grout (DOE, 2006b). In addition to the residual wastes at the bottom of the tanks, in two of the tanks, radionuclides have contaminated the sand pad underlying the tank as a result of inadvertent back-siphoning of first-cycle extraction fluid from reprocessing from the large tanks into the concrete vaults. As part of the demonstration of compliance with performance objectives, the DOE has modeled groundwater transport from the tank heels and sandpads into the Tank Farm Facility subsurface.

As at Savannah River, the INTEC subsurface has been locally contaminated by previous disposal activities and spills. There are two sources of contamination in the subsurface at INTEC: (i) contamination from a former injection well that released low activity waste water directly into the aquifer south of the Tank Farm Facility and (ii) contamination from piping leaks. The subsurface at the Tank Farm Facility was significantly contaminated as a result of a 1972 leak of 70,400 L [18,600 gal] of sodium-bearing waste, which entered the vadose zone during an unsuccessful attempt at transferring the waste between tanks. Approximately 5.9 × 10⁸ MBq [1.59 × 10⁴ Ci] of strontium-90 and 1.1 × 10⁵ MBq [3 Ci] of technetium-99 were released during the event (DOE–Idaho, 2006a). Maximum concentrations measured in shallow perched water underneath the Tank Farm Facility are 200,000 pCi/L for strontium-90, while maximum measured concentrations in the saturated zone are 3,000 pCi/L for technetium-99 (DOE–Idaho, 2006a,b). Other COCs detected in the subsurface as a result of the release of radioactivity into the environment from the 1972 event and from the injection well located south of the Tank Farm Facility include tritium, neptunium-237, plutonium-239, plutonium-240, and uranium-234.

3.2 Description of Idaho Nuclear Technology and Engineering Center Tank Farm Facility

The INTEC is a large industrial complex located in the south-central portion of INL. INL is an approximately 2,305-km² [890-mi²] reservation in southeastern Idaho owned by the U.S. government. The INTEC facility is located approximately 29 km [18 mi] from the closest eastern boundary, approximately 23 km [14 mi] from the closest western boundary, approximately 16 km [10 mi] from the closest southern boundary, and approximately 29 km [18 mi] from the closest northern boundary. INL and INTEC are located in a broad, relatively flat plain in the Pioneer Basin—a closed drainage basin. INTEC is approximately 460 m [1,500 ft] from the Big Lost River channel. The Big Lost River is an intermittent stream that sinks into the permeable vadose zone and the Snake River Plain Aquifer (SRPA) below. The Big Lost River enters INL in the southern portion of its western boundary, and in the wettest years, flows east and north where it ends in the Big Lost River playas and sinks located approximately 15 km [9 mi] north of INTEC.

The historical mission of INTEC, formerly known as the Idaho Chemical Processing Plant, was to recover fissile uranium by reprocessing spent nuclear fuel. The resulting liquid waste

generated from this process was acidic and radioactive. The liquid contained fission products, activation products, transuranic radionuclides, and various metals. The liquid waste was temporarily stored in the underground Tank Farm Facility at INTEC until the liquid radioactive waste was converted to a solid granular form. The Tank Farm Facility, located within the northern portion of INTEC, comprises eleven 1,000-m³ [300,000-gal] stainless steel tanks in unlined concrete vaults of various construction, four inactive 100-m³ [30,000-gal] stainless steel tanks, interconnecting waste transfer lines, and associated support instrumentation and valves.

INTEC is underlain by an alluvial veneer of Pleistocene-to-Holocene Big Lost River flood plain deposits and a sequence of Quaternary volcanic rocks and sedimentary interbeds (Whitehead, 1992). The Tank Farm Facility is embedded in 13 m [43 ft] of alluvial silt, sand, and gravel that lie above an alternating sequence of basalt lava flows and interbedded sediments (Figure 3-1). Stratigraphic sections derived from well characterization activities at or near INTEC indicate that basalt flows and sedimentary interbeds comprise horizontal to slightly inclined layers. More than 30 geologic units compose the vadose zone and the upper portion of the SRPA, including 19 basalt flow groups, 11 sedimentary interbeds, and surficial alluvium.

Alluvium and interbed sediments at INL include particles from clay to gravel sized. The alluvium typically is a few meters thick and has undergone little artificial disturbance over much of INL. Sedimentary interbeds may be as thick as 20 m [60 ft]. Sedimentary interbeds accumulated above basalt flows for hundreds to hundreds of thousands of years during periods of volcanic quiescence. Much of the alluvium and interbed sediment at INL was deposited by meltwater discharge and periodic floods along the ancestral channel and flood plain of the Big Lost River during past glacial declines (Rathburn, 1993). Another major part of the alluvium and sedimentary interbeds is loess and aeolian material derived from fine alluvial deposits having grain sizes ranging from fine silt to very fine sand. Clay- and sand-sized sediment was also transported by water and wind into the fractures of the underlying basalt flows. Besides silt, sand, and gravel, sedimentary interbeds may contain small clay lenses, scoria, and basalt rubble (Nimmo, et al., 2004).

During periods of volcanic quiescence, sediment accumulated in the topographic depressions of underlying basalt flows to form the sedimentary interbeds. Mineralogically, the interbed sediments closely resemble the alluvium. However, the interbed sediments have a greater density, more uniform structure, and are less aggregated than the alluvium (Nimmo, et al., 2004). Some interbed sediments are highly stratified and have internal layers and lenses that differ substantially in texture, structure, and geochemical composition due to varying depositional conditions or baking by a flow of fresh lava.

As determined by sieve analysis, optical scattering, and other methods, grain sizes of alluvium and interbed sediments range from clay to pebble sized (Bartholomay, et al., 1989; Perkins and Nimmo, 2000; Perkins, 2003; Winfield, 2003). Most alluvium and interbed sediment falls within the silt loam textural class, containing about 0 to 27-wt% clay, 55- to 80-wt% silt, and 10- to 35-wt% sand. Significant lithologic variations occur within surficial sediment and sedimentary interbeds as a result of changes in the depositional environment through time (Rightmire and Lewis, 1987). Weakly developed to well-developed paleosols containing carbonized zones and abundant clay cutans occur in both the alluvium and sedimentary interbeds (Forman, et al., 1993). Carbonized zones represent vegetation that was inundated and baked by lava flows during the geologic past. Besides paleosols, sedimentary structures in the alluvium and interbed sediments that could influence the movement of contaminants include cracks formed

by hydrocompaction and dessication, freeze—thaw features, burrow and rootlet holes, caliche development, horizontal laminations, ripple cross-stratification, planar cross-stratification, lenticular bedding, flaser bedding, rip-up clasts, load casts, and varves (Rightmire and Lewis, 1987).

Analyses of bulk mineralogy of INL alluvium and interbed sediments show the presence of quartz, plagioclaise feldspar, K-feldspar, pyroxene, olivine, calcite, dolomite, and clay minerals (Bartholomay, 1990; Bartholomay, et al., 1989). In general, quartz and plagioclaise feldspar are the most abundant minerals in the sediment in the south and western portions of INL, which includes INTEC, while calcite and quartz are the most abundant minerals to the north (Bartholomay, 1990). Clay minerals are also present in most of the sediment samples from INL. Analyses of the clay minerals sampled throughout INL indicate that illite predominates (ranging from 10 to 100 percent) and that lesser amounts of smectite, mixed-layer illite/smectite, kaolinite, and possible chlorite are present (Bartholomay, et al., 1989; Reed and Bartholomay, 1994). Abundance of clay minerals in interbed samples from the Subsurface Disposal Area (SDA), located approximately 12 km [7.5 mi] southwest of INTEC, ranges from 0 to 60 percent and averages about 20 percent (Bartholomay, 1990). Mineralogical analyses indicated that most of the carbonate in INL sediments is in the form of calcite (Bartholomay, et al., 1989). Calcite contents in samples from the SDA range from zero, for most of the samples, to 54 percent. Rightmire and Lewis (1987) reported trace amounts of iron oxyhydroxides, hematite, siderite, and dolomite in some INL sediment samples.

Basalt flows are mostly thin {3 to 9 m [10 to 30 ft]}, tube-fed pahoehoe flows (Anderson, et al., 1999). Outcrop exposures indicate that individual basalt flows generally form long, sinuous lobes having median length/width ratios of about 3:1 (Welhan, et al., 1997). Some flows are as long as several kilometers (Kuntz, et al., 1994). Many of the flows have closely spaced fractures up to a few meters wide and tens of meters long. These may be opened or filled with rubble, sediment, or younger lava. In some places, basalt flows are cut by vertical fissures, dikes, and tension cracks (Anderson, et al., 1999).

Basalt flows and other volcanic deposits combine into basalt flow groups that range from 3 to 20 m [10 to 60 ft] thick and are internally interbedded with discontinuous scoria and thin layers of sediment (Nimmo, et al., 2004). Each basalt flow group is a complex assemblage of overlapping flows and deposits related to a single eruption. A basalt flow group comprises several, perhaps hundreds, of distinct basalt flows that occurred in a time interval brief enough that little or no sediment accumulated between them.

Most basalt flows at INL have the chemical characteristics of both tholeittic and alkali olivine basalts. Typical basalt samples consist mainly of plagioclaise feldspar, pyroxene, and olivine and contain lesser amounts of ilmenite, magnetite, hematite, and accessory apatite (Kuntz, et al., 1980; Rightmire and Lewis, 1987; Knobel, et al., 1997). The typical chemical composition of basalt samples at INL is shown in Table 3-1. The basalts are rich in iron and calcium and poor in sodium and potassium.

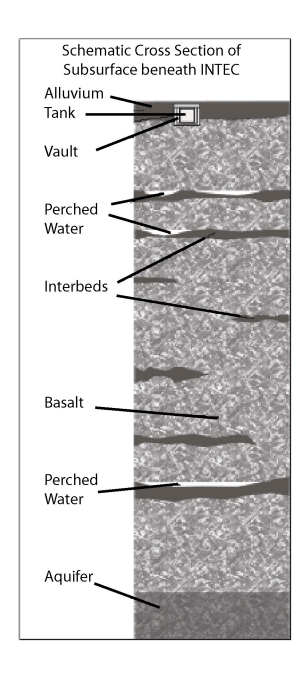


Figure 3-1. Schematic of the Subsurface at INTEC (Not to Scale, Taken From DOE–Idaho, 2006c)

Table 3-1. Typical Chemical Composition of Idaho National Laboratory Basalt.* Oxide Contents in wt% and Strontium and Barium Contents in mg/L.										
SiO ₂	TiO ₂	Al ₂ O ₃	Fe ₂ O ₃	MnO	MgO	CaO	Na₂O	K ₂ O	Strontium	Barium
45–48	1.3–3.8	12–15	12–17	0.17-0.23	5–10	9–11	2.1–2.7	0.3-0.9	230–352	193–507

*Colello, J.J., J.J. Rosentreter, R.C. Bartholomay, and M.J. Liszewski. "Strontium Distribution Coefficients of Basalt Core Samples from the Idaho National Engineering and Environmental Laboratory, Idaho." U.S. Geological Survey Water Resource Investigations Report 98-4256. 1998.

Fractures and vesicles within basalts commonly are coated with fine-grained sediment infill and sometimes with secondary minerals consisting of calcite, clays, zeolites, silica, iron oxides, and iron oxyhydroxides (Rightmire and Lewis, 1987; Morse and McCurry, 1997). Fractures within and contacts between individual basalt flows provide a complex network of potential vertical and horizontal pathways for the movement of water and wastes within the vadose zone and the aquifer.

The vadose zone is about 150 m [450 ft] thick near the INTEC and overlies the SRPA (Martian, et al., 2006). The SRPA is one of the largest and most productive aquifers in the United States, receiving natural recharge at and near the INTEC from precipitation and Big Lost River underflow (Bennett, 1990). Several perched water zones underlie the INTEC facility. Occurrences of perched water at the INTEC correspond to flow barriers created by low permeability portions of sedimentary interbeds. The perching of water in the vadose zone is brought about primarily by operational processes and landscape watering and secondarily by ephemeral Big Lost River seepage that releases water into the subsurface.

Analyses of water samples from eight SRPA wells at INL indicate that the waters are a calcium magnesium bicarbonate type (Knobel, et al., 1997, 1992), which is typical of most water in the SRPA. The composition of major cations and anions in the SRPA based on Wood and Low's (1986) characterization is listed in Table 3-2. These data indicate that groundwater from the SRPA is typically alkaline with significant bicarbonate concentration.

Water chemistry data for perched groundwater at INL can vary depending on the location of the wells and the timing of sample collection. Some samples collected from the SDA contain waste constituents leached from radiochemical and organic wastes (Bartholomay and Tucker, 2000).

Water chemistry data from perched groundwater samples from INL generally show chemical characteristics similar to SRPA water (Barraclough, et al., 1976; Rightmire and Lewis, 1987;

Table 3-2. Composition of Eastern Snake River Plain Aquifer Groundwater*				
Component	t Mean (mg/L) Maximum (mg/L) Minimum (
Ca ²⁺	51	120	5	
K⁺	4	12	0	
Mg ²⁺	18	54	0.2	
Na⁺	26	140	4	

Table 3-2. Composition of Eastern Snake River Plain Aquifer Groundwater* (continued)				
Component	Mean (mg/L)	Maximum (mg/L)	Minimum (mg/L)	
CI ⁻	28	230	1.8	
SO ₄ ²⁻	41	218	1.8	
HCO ₃ -	220	510	52	
F ⁻	0.6	11	0.03	
SiO ₂	50	138	5.8	
рН	7.8	9.1	7.0	
Ionic strength (mol/L)	0.008	0.02	0.004	

^{*}Wood, W.W. and W.H. Low. "Aqueous Geochemistry and Diagenesis in the Eastern Snake River Plain Aquifer System, Idaho." *Geological Society of America Bulletin 97*. 1986.

Knobel, et al., 1992; Bartholomay, 1998; Del Debbio and Thomas, 1989). The chemical composition of a perched water sample from the vadose zone taken at a depth of 38 m [125 ft] south of the Test Reactor Area near the Big Lost River and aquifer water taken at a depth of 140 m [460 ft] west of the Test Reactor Area are listed in Table 3-3 (Del Debbio and Thomas, 1989). The data indicate that both groundwaters are buffered by a carbonate/bicarbonate system and have an average pH of 8. Oxygen analyses indicate that the groundwaters are saturated with oxygen thus creating oxidizing conditions.

Several chemical properties of water will affect the mobility of contaminants at INL including pH, alkalinity, dissolved organic carbon, system oxidation state, and the concentration of aqueous complexing agents. Water from perched zones at INL and from the SRPA have the following characteristics that can significantly affect contaminant transport:

- pH that ranges from 7.2 to 8.4 (Nimmo, et al., 2004)
- Cation concentrations dominated by Ca²⁺ and anion concentrations dominated by HCO₃⁻
- Dissolved O₂ concentrations that are nearly saturated to slightly supersaturated with respect to air

Table 3-3. Composition of Perched Water and Aquifer Water Collected At the Idaho National Laboratory*				
Component	Perched Water (mg/L)	Aquifer Water (mg/L)		
Ca ²⁺	44	48		
Mg ²⁺	14	18		
SiO ₂	18	26		

Table 3-3. Composition of Perched Water and Aquifer Water Collected At the Idaho National Laboratory* (continued)				
Component	Perched Water (mg/L)	Aquifer Water (mg/L)		
Iron (total)	<0.02	<0.02		
Na⁺	8.2	8.5		
K⁺	1.6	2.1		
Cl ⁻	7.4	12		
SO ₄ ²⁻	22	24		
HCO ₃ -	207	206		
O ₂	7.4	7.6		
рН	8.1	8.0		
Eh	750 mV	750 mV		

*Del Debbio, J.A. and T.R. Thomas. "Transport Properties of Radionuclides and Hazardous Chemical Species in Soils at the Idaho Chemical Processing Plant." WINCO-1068. Idaho Falls, Idaho: Westinghouse Idaho Nuclear Company, Inc. 1989.

- Dissolved organic carbon concentrations that are generally smaller than 1 mg/L [1 ppm] (Knobel, et al., 1992) or about 1 mg/L [1 ppm] (Busenberg, et al., 2000)
- Saturation indices that indicate the water is typically nearly saturated with respect to calcite (Nimmo, et al., 2004)

Most groundwaters at the INL site have dissolved O_2 , which restricts the extent to which redox conditions can vary at least in mobile contaminated water at the site. Natural organic matter can be present in some of the less hydraulically conductive materials at INL, thereby potentially resulting in reducing conditions in some isolated, relatively immobile waters at the site. Lower redox potentials have been observed at a few locations beneath the INTEC, where the perched water appears to have been affected by past releases of fuel hydrocarbons (Golder Associates, Inc., 1991).

Mineralogical characteristics will also affect the mobility of contaminants at INL. The mineralogic properties that are important to the partitioning of contaminants between groundwater and the sediment and basalt units beneath INL include

- The presence of calcite in both basalt fractures and in sedimentary interbeds
- Iron oxyhydroxides, which can strongly adsorb actinides under certain conditions, are commonly observed on fracture surfaces in basalt and in the interbed sediments
- Clay minerals, most commonly identified as illite, which account for an average of 20 wt% of the interbed sediments (Bartholomay, 1990)

The clay minerals that are present in the system provide sites for sorption or ion exchange with contaminant-bearing water. Both smectite and mixed-layer smectite clays were observed in some interbed samples; these clays have the largest cation exchange capacities (CEC) of the clay minerals present. For example, samples from the SDA with low CECs contained illite, while high CEC samples contained from 9- to 33-wt% expandable layer clays, mixed layer clays, and smectite (Rightmire and Lewis, 1987).

The previously noted chemical and mineralogic characteristics define the environment that will control contaminant mobility at INL. For example, the presence of O_2 and dissolved organic carbon affects the oxidation-reduction characteristics of the system and hence the affinity of the chemical species for the aqueous or solid phase. Oxygen in the vadose zone may be depleted by reaction with organic materials released from waste tanks, creating localized reducing conditions that may affect contaminant mobility. However, due to the mineralogic characteristics of the basalt and interbed sediments at INL, the buffering capacity of the vadose zone and aquifer system beneath INL is great, and reducing conditions potentially caused by waste releases are not expected to persist as water moves vertically through the system.

3.3 Designation of Risk-Significant Radionuclides

The COCs chosen for analysis in this report are based both on previous model results and on more general considerations of hydrochemical behavior and dose effects. In addition, inclusion on the DOE list of highly radioactive radionuclides was considered (DOE, 2006b). DOE performance assessment results suggested that isotopes of strontium, technetium, and iodine may be most important to groundwater dose (DOE–Idaho, 2003). In the models, the transported strontium arose chiefly from the sandpad inventories, because tank-derived strontium would have mostly decayed by the time tanks failed. Uranium and neptunium are significant in waste inventories and are relatively mobile in the subsurface environment at the site. Other actinides—plutonium, americium, and curium—are included due to potentially important dose effects if sorption coefficients are overestimated. DOE included potentially mobile carbon and cesium radioisotopes in their list of highly radioactive radionuclides; lead is also significant in inventories and potentially mobile in the subsurface.

3.4 Sorption Coefficients

The hydrostratigraphic units that will be affected by release of contaminants from storage facilities at INTEC are alluvium, sedimentary interbeds, and basalt. For many of the COCs in this review, site-specific K_d data for subsurface media at the INTEC or for subsurface media at INL facilities in the vicinity of INTEC (e.g., the Surface Disposal Area and Test Reactor Area) are available. In this review, site-specific K_d measurements and literature-recommended K_d values for the COCs at INL are analyzed and reviewed to derive a recommended K_d value for alluvium, sedimentary interbeds, and basalt beneath INL. In some instances where site-specific K_d measurements are lacking for hydrostratigraphic units, the mineralogic and hydrogeochemical characteristics of the units (summarized in Table 3-4) are used in the K_d recommendation. Table 3-4 indicates that the lithologic, mineralogic, and chemical composition of the alluvium and sedimentary interbed units are not significantly different. Therefore, when site-specific K_d data are available for a COC for one of these units but not the other, the K_d recommendation based on the site-specific K_d data is applied to both units.

Many researchers have measured K_d values in sediments and basalt from the Hanford site in southeastern Washington for the COCs in this review. The hydrogeochemical environment at the Hanford site is analogous to INL in many aspects (i.e., alkaline pH, high carbonate concentrations, oxidizing conditions, and similar though not identical sediment and basalt mineral compositions). When available, reported K_d values from Hanford for the COCs in this review are presented to help determine whether the recommended K_d values for INL are reasonable.

To aid in transport model simulations at INTEC, Cooper, et al. (2006) estimated K_d ranges and recommended K_d values for a number of radionuclides in alluvium, interbed sediments, and basalt. For many of the designated radionuclides in this review, these estimated K_d ranges and recommended K_d values were useful in determining whether the K_d values recommended for the INL in this review are reasonable. In the K_d estimates of Cooper, et al. (2006), site-specific K_d data from INTEC was given the highest priority in the K_d recommendations. When site-specific INTEC data was not available, INL-SDA data were judged to be most representative. Sediments from the SDA have been studied more throughly than INTEC sediments, originate from the same geologic source, and possess similar geochemical properties. The main difference between these two subsurface regions is in the grain size distribution, CEC, and reactive surface area of the respective sediments. Based on reported data from Bartholomay, et al. (1989), K_d estimates based on SDA data were scaled according to typical CEC of the respective sediments (INTEC/SDA = 1:5). When SDA data were not available, data from the open literature on comparable sediments were employed, using INTEC and SDA geochemistry as a guide. The results of the Cooper, et al. (2006) analyses were three K_d values for each contaminant in each subsurface unit (alluvium, interbed sediments, and basalt): a minimum estimated K_d value, a maximum estimated K_d value, and a recommended K_d value. For each of the COCs in this review analyzed by Cooper, et al. (2006), the estimated range (i.e., the minimum and maximum estimated K_d value) and recommended K_d are graphically illustrated in Figures 3-2 through 3-12 for alluvium, interbed sediments, and basalt.

3.4.1 Carbon

The C-14 present in INL waste streams originates from nuclear fuel processing and is predominantly inorganic in nature. Inorganic carbon transport occurs via both slow aqueous transport as HCO_3^- ion and rapid transport of CO_2 gas. Carbon transport is likely to be affected by reactions other than sorption, but an empirical K_d approach may be sufficient for understanding potential transport.

Plummer, et al. (2004) conducted an unsaturated column study to determine carbon transport in INL surficial sediments collected from playas adjacent to the SDA. The water used in the column experiments was a synthetic vadose zone water (pH 7.8) formulated to approximate the composition of natural infiltrating water at the SDA. Results indicated that approximately 70 percent of total carbon was released from the surface of the column as $\rm CO_2$ gas, 4 percent exited as the aqueous phase, 2 percent was lost to sampling, and 24 percent remained in the column (1 percent in soil gas, 3 percent in soil water, and 20 percent adsorbed). After fitting the carbon data to a multiphase transport equation that accounted for minor variations in pH, carbon $\rm K_d$ values of $\rm 0.5 \pm 0.1~mL/g$ were calculated for the first 3 months following carbon injection and ranged from 0.8 to 2.4 mL/g after 1 year of transport.

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Table 3-4. Mineralogic and Hydrogeochemical Characteristics of Units at the Idaho National Laboratory				
Unit	Lithology*	Mineralogy†	pH Range‡	Other Hydrochemical Aspects‡
Alluvium	Silt loam (0–27-percent clay, 55–80-percent silt, 10–35-percent sand)	Major constituents: quartz, plagioclaise, illite. Minor constituents: K-feldspar, pyroxene, olivine, calcite, dolomite, iron oxyhydroxides, smectite, mixed layer illite/smectite, kaolinite, chlorite, hematite, siderite	7.2–8.4	High dissolved O ₂ Cations dominated by Ca ²⁺ Anions dominated by HCO ₃ Low dissolved organic carbon Water near calcite saturation
Sedimentary Interbeds	Silt loam (0–27-percent clay, 55–80-percent silt, 10–35-percent sand); some small clay lenses, scoria, and basalt rubble	Major constituents: quartz, plagioclaise, illite. Minor constituents: K-feldspar, pyroxene, olivine, calcite, dolomite, iron-oxyhydroxides, smectite, mixed layer illite/smectite, kaolinite, chlorite, hematite, siderite	7.2–8.4	High dissolved O ₂ Cations dominated by Ca ²⁺ Anions dominated by HCO ₃ Low dissolved organic carbon Water near calcite saturation

Table 3-4. Mineralogic and Hydrogeochemical Characteristics of Units At the Idaho National Laboratory (continued)				
Unit	Lithology*	Mineralogy†	pH Range‡	Other Hydrochemical Aspects‡
Basalt	Tube fed pahoehoe flows; closely spaced fractures often filled with rubble, sediments, or younger lava	Major constituents: plagioclaise, pyroxene, olivine. Minor constituents: ilmenite, magnetite, hematite, apatite. Fractures and vesicles commonly coated with calcite, clays, zeolites, silica, iron oxides, and iron oxyhydroxides	7.2– 8.4	High dissolved O ₂ Cations dominated by Ca ²⁺ Anions dominated by HCO ₃ ⁻ Low dissolved organic carbon Water near calcite saturation

*Lithologic characterization based on data from Bartholomay, R.C., L.L. Knobel, and L.C. Davis. "Mineralogy and Grain Size of Surficial Sediment from the Big Lost River Drainage and Vicinity, with Chemical and Physical Characteristics of Geologic Materials from Selected Sites at the Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Open-File Report 89-384. 1989; Nimmo, J.R., J.P. Rousseau, K.S. Perkins, K.G. Stollenwerk, P.D. Glynn, R.C. Bartholomay, and L.L. Knobel. "Hydraulic and Geochemical Framework of the Idaho National Engineering and Environmental Laboratory Vadose Zone." *Vadose Zone Journal.* Vol. 3. pp. 6–34. 2004; and Anderson, S.R., M.A. Kuntz, and L.C. Davis. "Geologic Controls of Hydraulic Conductivity in the Snake River Plain Aquifer at and Near the Idaho National Engineering and Environmental Laboratory, Idaho." U.S. Geological Survey Water Resources Investigations Report 99-4033. 1999.

†Mineralogic characterization based on data from Bartholomay, R.C., L.L. Knobel, and L.C. Davis. "Mineralogy and Grain Size of Surfical Sediment from the Big Lost River Drainage and Vicinity, with Chemical and Physical Characteristics of Geologic Materials from Selected Sites at the Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Open-File Report 89-384. 1989; Bartholomay, R.C. "Mineralogical Correlation of Surficial Sediment from Area Drainages With Selected Sedimentary Interbeds at the Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Water Resources Investigations Report 90-4147. 1990; Rightmire, C.T. and B.D. Lewis. "Hydrogeology and Geochemistry of the Unsaturated Zone, Radioactive Waste Management Complex, Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Water Resources Investigations Report 87-4198. 1987; Kuntz, M.A., G.B. Dalrymple, D.E. Champion, and D.J. Doherty. "Petrography, Age, and Paleomagnetism of Volcanic Rocks at the Radioactive Waste Management Complex, Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Open-File Report 80-388. 1980; and Knobel, L.L., R.C. Bartholomay, and B.R. Orr. "Preliminary Delineation of Natural Geochemical Reactions, Snake River Plain Aquifer System, Idaho National Engineering Laboratory and Vicinity, Idaho." U.S. Geological Survey Water Resources Investigations Report 97-4093. DOE/ID-22139. 1997.

‡Chemical characterization based on data from Nimmo, J.R., J.P. Rousseau, K.S. Perkins, K.G. Stollenwerk, P.D. Glynn, R.C. Bartholomay, and L.L. Knobel. "Hydraulic and Geochemical Framework of the Idaho National Engineering and Environmental Laboratory Vadose Zone." *Vadose Zone Journal.* Vol. 3. pp. 6–34. 2004; Wood, W.W. and W.H. Low. "Aqueous Geochemistry and Diagenesis in the Eastern Snake River Plain Aquifer System, Idaho." *Geological Society of America Bulletin* 97. 1986; Del Debbio, J.A. and T.R. Thomas. "Transport Properties of Radionuclides and Hazardous Chemical Species in Soils at the Idaho Chemical Processing Plant." WINCO–1068. Idaho Falls, Idaho: Westinghouse Idaho Nuclear Company, Inc. 1989; Knobel, L.L., B.R. Orr, and L.D. Cecil. "Summary of Background Concentrations of Selected Radiochemical and Chemical Constituents in Groundwater from the Snake River Plain Aquifer, Idaho—Estimated from an Analysis of Previously Published Data." *Journal of Idaho Academy of Sciences.* Vol. 28. pp. 48–61. 1992; and Busenberg, E., L.N. Plummer, M.W. Doughten, P.K. Widman, and R.C. Bartholomay. "Chemical and Isotopic Composition and Gas Concentrations of Ground Water and Surface Water from Selected Sites at and near the Idaho National Engineering and Environmental Laboratory, Idaho, 1994–97." U.S. Geological Survey Open-File Report 00-81. 2000.

Based on the Plummer, et al. (2004) study, Cooper, et al. (2006) estimated that the K_d for carbon in alluvium and interbed sediments could range from 0.5 to 2.4 mL/g and recommended a carbon K_d value of 1.6 mL/g for alluvium and interbed sediments at INTEC. Cooper, et al. (2006) found no data for carbon partitioning in basalt at INL and recommended a K_d of 0 mL/g.

Measured and recommended carbon K_d values are summarized in Table 3-5. Site-specific carbon K_d values from the column experiment of Plummer, et al. (2004) using INL alluvium ranged from 0.5 to 2.4 mL/g. Cooper, et al. (2006) recommended a carbon K_d value of 1.6 mL/g for alluvium and interbed sediments at INTEC, which is the median of the range of carbon K_d values reported for the 1-year transport experiment of Plummer, et al. (2004). In this review, a carbon K_d value of 1 mL/g is recommended for INL alluvium and interbed sediments. This recommendation is estimated from the minimum of the range of values reported by Plummer, et al. (2004) for the 1-year transport experiment (0.8 mL/g). No site-specific or literature-derived data are available for carbon partitioning to basalt, and a value of 0 mL/g is recommended. For basalt, there is no clear basis for using carbon K_d values reported for sediments or soils (e.g., Thibault, et al., 1990).

The position of the carbon K_d values recommended in this review with respect to measured and estimated site-specific carbon K_d ranges and values is graphically displayed in Figure 3-2. This figure illustrates the uncertainty associated with the carbon K_d values recommended for alluvium, interbed sediments, and basalt at INL. The recommended carbon K_d values are associated with high uncertainty due to limited site-specific sorption data. The recommended carbon K_d value of 1 mL/g for INL alluvium and interbed sediments is based solely on site-specific carbon sorption data reported by Plummer, et al. (2004). No site-specific carbon sorption data exists for INL basalt.

Interbed Sediments, and Basalt at the Idaho National Laboratory				
	Measured Carbon K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Carbon K _d (mL/g)*	Recommended Carbon K _d (mL/g)
Alluvium	0.5–2.4†	Column	1.6	1
Sedimentary Interbeds	_	_	1.6	1

arred and Decommanded K. Values for Carbon in Alluvium

0

0

Basalt

^{*}Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

[†] Plummer, M.A., L.C., Hull, and D.T. Fox. "Transport of Carbon-14 in a Large Unsaturated Soil Column." *Vadose Zone Journal.* Vol. 3. pp.109–121. 2004.

3.4.2 Strontium

Sr-90 exists primarily as the uncomplexed Sr^{2+} ion over the pH range of natural waters (pH 3 to 10). Sr^{2+} adsorbs to both clay minerals and carbonate minerals. The extent to which strontium partitions from the aqueous phase to the solid phase is expected to be controlled primarily by the CEC of the solid phase. Other factors affecting strontium K_d values are pH and ionic strength. Strontium K_d values have been reported to increase with increasing pH (Prout, 1958; Rhodes, 1957). This trend is likely the result of hydrogen ions competing with Sr^{2+} for exchange sites. Elevated concentrations of major cations (e.g., Na^+ , K^+ , Ca^{2+} , Mg^{2+}) can greatly reduce strontium K_d values (Bunde, et al., 1998, 1997). Strontium K_d values may decrease from 100 to 200 mL/g in low ionic strength solutions to less than 5 mL/g in high ionic strength solutions (Routson, et al., 1980). There is little tendency for strontium to form complexes with inorganic or organic ligands (Faure and Powell, 1972; Stevenson and Fitch, 1986).

A number of researchers have investigated strontium sorption onto INL alluvium, interbed sediments, and basalt. In general, these studies indicate that strontium sorption is strongly affected by variability in both solution composition and mineral content.

Liszewski, et al. (1997) measured strontium K_d s for 20 surficial sediment samples collected at INL. Batch experimental techniques were used to determine K_d s using a synthesized aqueous solution representative of wastewater in INL waste disposal ponds (pH 8.0 ± 0.1). Strontium K_d values of the 20 surficial sediments ranged from 36 to 275 mL/g, indicating significant variability in the strontium sorptive capacities. Liszewski, et al. (1997) attributed the strontium K_d variability to physical and chemical properties of the sediments and compositional changes in the equilibrated solutions after mixing with the sediments. In similar batch experiments conducted on interbed sediments from INL, Liszewski, et al. (1998) report strontium K_d values that ranged from 66 to 266 mL/g using low ionic strength solutions.

Hemming, et al. (1997) conducted batch experiments using INL surficial sediments to examine the effect of different experimental techniques on experimentally derived strontium K_d values at a fixed pH of 8.0. Strontium K_d s values ranged from 11 to 23 mL/g among the three experimental variables examined: method of sample agitation (rotating mixer and shaker table), solid/liquid ratio (1:2 and 1:20), and method of sediment preparation (crushed and noncrushed).

Del Debbio and Thomas (1989) measured strontium K_d values on alluvium, interbed sediments, and basalt collected at INL. Batch experiments were performed on the materials at a 1:20 solid/liquid ratio using a low ionic strength groundwater (pH 8.0) collected from an onsite well. Measured strontium K_d values ranged from 35 to 52 mL/g for the alluvium, 110 to 186 mL/g for the interbed sediments, and 1.1 to 2.7 mL/g for the basalt.

Schmalz (1972) performed batch experiments to measure strontium K_d values using alluvium from INL and a low ionic strength pond water with a pH of 9.2 collected from the Test Reactor Area. The pond water was spiked with varying concentrations of strontium. Strontium K_d values measured on four soil samples varied over a narrow range (23 to 26 mL/g).

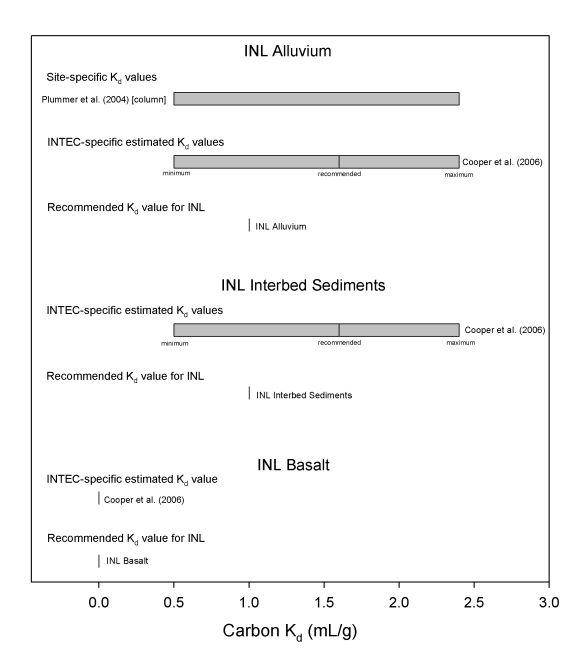


Figure 3-2. Measured and Estimated Site-Specific Carbon K_d Ranges and Values and Recommended Carbon K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-5). Recommended Carbon K_d Values Are Associated With High Uncertainty Due to Limited Site-Specific Sorption Data.

Hawkins and Short (1965) performed batch tests to measure strontium K_d values on alluviual soil from INL as a function of solution chemistry. A solution for use in these tests was synthesized to simulate INL groundwater. The chemistry of this solution was then varied with respect to Ca^{2+} , Mg^{2+} , Sr^{2+} , K^+ , Na^+ , and NH_4^+ . K_d values measured for strontium ranged from 8.3 to 16.6 mL/g. Based on these tests, equations were developed to predict the sorption of strontium at varying solution chemistries.

Newman, et al. (1996) summarized results from short-term (400-hour) batch experiments and column tests measuring strontium K_d for basalt and interbed sediments. The batch and column tests were conducted using a synthetic groundwater formulated to approximate the composition observed in SRPA field samples (ionic strength = 0.0048 M; pH 8.0) (Wood and Low, 1986). For batch tests, strontium K_d values ranged from 9 to 13 mL/g for basalt and from 42 to 63 mL/g for interbed sediments. For column tests, strontium K_d values ranged from 6.3 to 6.5 mL/g for basalt and >47 mL/g for interbed sediments.

Fjeld, et al. (2001) performed laboratory column tests to characterize the mobility of strontium in a basalt sample and a composite of sedimentary interbed from INL. The solution used in the tests was a low ionic strength (0.004 M) synthetic groundwater (pH 8.0) formulated to approximate the major ion concentration of SRPA groundwater samples (Wood and Low, 1986). Strontium retardation factors ranged from 25 to 33 for basalt and >200 for interbed sediments. Based on the reported porosities and bulk densities of the basalt and interbed columns, these retardation factors equate to K_d values of 6.8 to 9.7 mL/g¹ for basalt and >47 mL/g for interbed sediments.

Colello, et al. (1998) measured strontium K_d values for 24 crushed basalt core samples collected at INL. Batch experiments were used to measure the K_d values using an aqueous solution representative of wastewater in waste disposal ponds (pH 8.0). Calculated strontium K_d values of the crushed basalt core samples ranged from 3.6 to 29.4 mL/g. The results indicated a narrow range of variability in the strontium sorptive capacities of basalt relative to those of sedimentary materials at INL. No correlation was observed between strontium K_d and any physical properties of the basalt.

Porro, et al. (2000) performed column and batch adsorption tests to determine strontium K_d values in crushed basalt from INL. All column and batch tests were conducted using a synthetic groundwater formulated to approximate the composition observed in SRPA samples. The synthetic groundwater had a low ionic strength (0.0048 M) and a pH of 8.0. Column tests were conducted at saturated and unsaturated moisture contents. Measured strontium K_d values for the column tests ranged from 2.37 to 4.26 mL/g and were not significantly affected by moisture content. Batch tests were conducted at several solid/liquid ratios (0.25 to 2.33). Measured strontium K_d values for the batch tests ranged from 0.58 mL/g at the highest solid/liquid ratio to 2.56 mL/g at the lowest solid/liquid ratio. Column tests conducted at various moisture contents from saturation to 20 percent consistently resulted in essentially the same K_d value and, therefore, were considered to be more robust than the batch tests.

¹Retardation factor (R_f converted to K_d based on following equation: $K_d = \frac{\left(R_f - 1\right)}{\rho_b}\theta$, where θ is porosity and ρ_b is bulk density.

Cooper, et al. (2006) estimated a strontium K_d range of 8 to 20 mL/g for INTEC alluvium and recommended a strontium K_d value of 12 mL/g for INTEC alluvium. For sedimentary interbeds, Cooper, et al. (2006) estimated a range of strontium K_d values from 25 to 84 mL/g with a recommended strontium K_d value of 50 mL/g. The estimated K_d values for alluvium and sedimentary interbeds were determined using equations developed by Hawkins and Short (1965) to predict the sorption of strontium at varying solution chemistries. The K_d values were calculated using the PHREEQC geochemical code and water analyses for perched zone wells at the INTEC compiled by Roddy (2005). For basalt, Cooper, et al. (2006) estimated a strontium K_d range of 0 to 15 mL/g and recommended a strontium K_d value of 0.5 mL/g. Cooper, et al. (2006) indicate that the strontium K_d estimates and recommendations for basalt were taken from Colello, et al. (1998), with the recommended strontium K_d value equal to the minimum strontium K_d value observed for basalt. However, Colello, et al. (1998) reported measured strontium K_d values ranging from 3.6 to 29.4 mL/g for basalt, which is not in agreement with the minimum and maximum estimated strontium K_d values for basalt reported by Cooper, et al. (2006).

Measured and recommended strontium K_d values are summarized in Table 3-6. For alluvium, site-specific strontium K_d values for the 5 listed batch experiments range from a minimum of 8.3 mL/g (Hawkins and Short, 1965) to a maximum of 275 mL/g (Liszewski, et al., 1997). Review of the laboratory procedures used by Liszewski, et al. (1997) to measure strontium K_d values on alluvium indicates that they may be biased toward higher values because of the water used in the measurements (i.e., synthesized aqueous solution representative of wastewater from disposal ponds). Therefore, the measured values of Liszewski, et al. (1997) may not be representative of natural geochemical conditions in the alluvium. Removing the Liszewski, et al. (1997) strontium K_d values from the analysis results in a maximum measured K_d value of 52 mL/g for the alluvium (Del Debbio and Thomas, 1989). A reasonably conservative strontium K_d value of 12 mL/g is recommended for INL alluvium based on the minimum measured K_d value of 8.3 mL/g (Hawkins and Short, 1965) and the maximum measured K_d value of 52 mL/g (Del Debbio and Thomas, 1989). This recommended strontium K_d value is in agreement with the strontium K_d value of 12 mL/g recommended by Cooper, et al. (2006) for INTEC alluvium.

For interbed sediments, site-specific strontium K_d values for the three listed batch experiments range from a minimum of 42 mL/g (Newman, et al., 1996) to a maximum of 266 mL/g (Liszewski, et al., 1997). The K_d value for 2 listed column experiments is >47 mL/g (Newman, et al., 1996; Fjeld, et al., 2001). Again the measured strontium K_d values of Liszewski, et al. (1997) may be biased high and not representative of natural chemical conditions in the interbed sediments because of the water used in the experiments (i.e., synthesized aqueous solution representative of wastewater from disposal ponds). Removing the Liszewski, et al. (1997) strontium K_d values from the analysis results in a maximum K_d value of 186 mL/g for the interbed sediments (Del Debbio and Thomas, 1989). Based on the strontium K_d values reported for the two column experiments (>47 mL/g), which are within the range of strontium K_d values reported for the batch experiments (42 to 186 mL/g), a K_d value of 50 mL/g appears to be appropriate for interbed sediments at INL. A strontium K_d value of 50 mL/g is in agreement with the strontium K_d value of 50 recommended by Cooper, et al. (2006) for INTEC interbed sediments. However, sensitivity studies conducted on models simulating past contaminant transport at the site (Rodriguez, et al., 1997, Appendix F) concluded that a strontium K_d of 12–24 mL/g matched observations well for both the unsaturated and saturated zones. On this basis, a conservative K_d of 12 mL/g is recommended for strontium in the interbed sediments.

Table 3-6. Measured and Recommended $K_{\rm d}$ Values for Strontium in Alluvium, Interbed Sediments, and Basalt at the Idaho National Laboratory

	Measured Strontium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Strontium K _d (mL/g)*	Recommended Strontium K _d (mL/g)
Alluvium	36-275† 11-23‡ 35-52§ 23-26¶ 8.3-16.6#	Batch Batch Batch Batch Batch	12	12
Sedimentary Interbeds	66-266** 110-186§ 42-63†† >47†† >47‡‡	Batch Batch Batch Column Column	50	12
Basalt	1.1-2.7§ 9.0- 3†† 6.3-6.5†† 3.6-29.4§§ 0.58-2.56¶¶ 2.37-4.26¶¶	Batch Batch Column Batch Batch Column	0.5	2

*Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

†Liszewski, M.J., J.J. Rosentreter, and K.E. Miller. "Strontium Distribution Coefficients of Surficial Sediment Samples from the Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Water Resource Investigations Report 97-4044. 1997.

‡Hemming, C.H., R.L. Bunde, M.J. Liszewski, J.J. Rosentreter, and J. Welhan. "Effect of Experimental Technique on the Determination of Strontium Distribution Coefficients of a Surficial Sediment from the Idaho National Engineering Laboratory, Idaho." *Water Research.* Vol. 31, Issue 7. pp. 1,629–1,636. 1997.

Engineering Laboratory, Idaho." *Water Research*. Vol. 31, Issue 7. pp. 1,629–1,636. 1997. §Del Debbio, J.A. and T.R. Thomas. "Transport Properties of Radionuclides and Hazardous Chemical Species in Soils at the Idaho Chemical Processing Plant." WINCO–1068. Idaho Falls, Idaho: Westinghouse Idaho Nuclear Company, Inc. 1989.

Company, Inc. 1989.

¶Schmalz, B.L. "Radionuclide Distribution in Soil Mantle of the Lithosphere as a Consequence of Waste Disposal at the National Reactor Testing Station." IDO–10049. Idaho Falls, Idaho: United States Atomic Energy Commission, Idaho Operations Office. 1972.

#Hawkins, D.B. and H.L. Short. "Equations for the Sorption of Cesium and Strontium on Soil and Clinoptilolite." IDO–12046. Idaho Falls, Idaho: Atomic Energy Commission, Idaho Operations Office. 1965.

**Liszewski, M.J., J.J. Rosentreter, K.E. Miller, and R.C. Bartholomay. "Strontium Distribution Coefficients of Surficial and Sedimentary Interbed Sediment Samples From the Idaho National Engineering Laboratory, Idaho." U.S. Geological Survey Water Resource Investigations Report 98-4037. 1989.

††Newman, M.E., I. Porro, R. Scott, R.M. Dunnivant, R.W. Goff, M.D. Blevins, S.M. Ince, J.D. Leyba, T.A. DeVol, A.W. Elzerman, and R.A. Fjeld. "Evaluation of the Mobility of Am, Cs, Co, Pu, Sr, and U Through INEL Basalt and Interbed Materials: Summary Report of the INEL/Clemson University Laboratory Studies." ER–WAG7–82, INEL –95/282 Idaho Falls, Idaho National Engineering Laboratory, 1996

INEL-95/282. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1996.
‡ ‡Fjeld, R.A., T.A. DeVol, R.W. Goff, M.D. Blevins, D.D. Brown, S.M. Ince, A.W. Elzerman, and M.E. Newman.
"Characterization of the Mobilities of Selected Actinides and Fission/activation Products in Laboratory Columns
Containing Subsurface Material from the Snake River Plain." *Nuclear Technology*. Vol. 135. pp. 92–108. 2001.

§\$Colello, J.J., J.J. Rosentreter, R.C. Bartholomay, and M.J. Liszewski. "Strontium Distribution Coefficients of Basalt Core Samples from the Idaho National Engineering and Environmental Laboratory, Idaho." U.S. Geological Survey Water Resource Investigations Report 98-4256. 1998.

Survey Water Resource Investigations Report 98-4256. 1998.
¶¶Porro, I., M.E. Newman, and F.M. Dunnivant. "Comparison of Batch and Column Methods for Determining Strontium Distribution Coefficients for Unsaturated Transport in Basalt." *Environmental Science and Technology*. Vol. 34. pp. 1,679–1,686. 2000.

For basalt, site-specific strontium K_d values for 4 listed batch experiments range from a minimum of 0.58 mL/g (Porro, et al., 2000) to a maximum of 29.4 mL/g (Colello, et al., 1998). Site-specific strontium K_d values for three listed column experiments range from a minimum of 2.37 mL/g (Porro, et al., 2000) to a maximum of 9.7 mL/g (Fjeld, et al., 2001). Review of the laboratory procedures used by Colello, et al. (1998) to measure strontium K_d values on basalt indicates that they may be biased toward higher values because of the water used in the measurements (i.e., synthesized aqueous solution representative of wastewater from disposal ponds). Therefore, the measured values of Colello, et al. (1998) may not be representative of natural chemical conditions in the basalt. Removing the Colello, et al. (1998) strontium K_d values from consideration results in a maximum measured K_d value of 13 mL/g for the basalt (Newman, et al., 1996). Based on the range of strontium K_d values reported for the column experiments (2.37 to 9.7 mL/g), which are within the range of strontium K_d values reported for the batch experiments (0.58 to 13 mL/g), a K_d value of 2 mL/g is recommended for basalt at INL.

The position of the strontium K_d values recommended in this review with respect to measured and estimated site-specific strontium K_d ranges and values is graphically displayed in Figure 3-3. This figure illustrates the uncertainty associated with the strontium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific strontium sorption data (Figure 3-3), the recommended strontium K_d values are associated with low uncertainty.

3.4.3 Technetium

Tc-99 predominantly occurs in oxidized (+7) state, as pertechnetate (TcO_4^-) over the complete pH range of natural waters, generally sorbs poorly to sediments, and is typically highly mobile (e.g., Denys, et al., 2003; Kaplan, 2003; Mashkin and Shikov, 2000). The sorption behavior of technetium is highly redox sensitive. Experimental data indicate that technetium mobility increases with increasing pH (Denys, et al., 2003) and decreases under organic-rich, reducing conditions (Mashkin and Shikov, 2000). The pH effect may arise from pH-dependent changes in surface charge or may reflect the role of organic matter; both provide anion sorption sites for TcO_4^- and reduce pH (Kaplan, 2003).

Del Debbio and Thomas (1989) performed column experiments to determine technetium sorption onto alluvium, interbed sediments, and basalt at INL. The experiments used a low ionic strength groundwater (pH 8.0) collected from an onsite well. Measured retardation factors were <1.0 on alluvium and interbed sediments, indicating no retention of technetium. The measured retardation factor for one basalt column experiment was 1.1, which equates to a technetium K_d of 0.016 mL/g using the reported bulk density and porosity of the basalt column.

Cooper, et al. (2006) used the technetium sorption data reported by Del Debbio and Thomas (1989) to recommend a technetium K_d value of 0 mL/g for alluvium and interbed sediments at INTEC. Cooper, et al. (2006) examined technetium behavior in other systems to establish a range of minimum and maximum technetium K_d estimates at INTEC. For example, Kaplan (2003) reported technetium K_d values that ranged from -0.1 to 0.1 mL/g for a silty-sand upland Savannah River Site (SRS) sediment (0.3 wt% Fe₂O₃, <0.02 wt% organic carbon) and from 0 to 0.3 mL/g for a silty-sand wetland SRS sediment (0.01 wt% Fe₂O₃, 0.14 wt% organic carbon). Sheppard and Thibault (1991) conducted a 4-year lysimeter study of technetium transport in

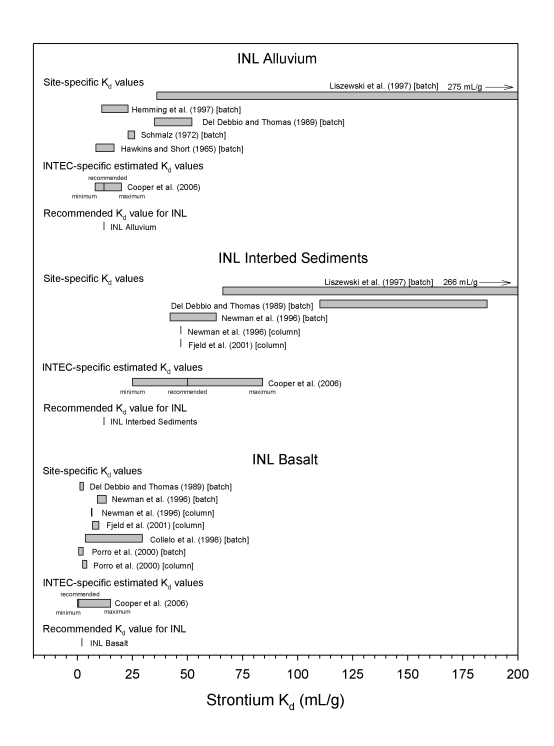


Figure 3-3. Measured and Estimated Site-Specific Strontium K_d Ranges and Values and Recommended Strontium K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-6). Recommended Strontium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

surficial soils and reported technetium K_d values ranging from 0.1 to 1.4 mL/g for sediments with 0.2 to 0.3 wt% organic carbon. Cooper, et al. (2006) indicated that the SRS upland sediment was most similar to INTEC sediment interbeds, while the SRS wetland sediment and lysimeter studies were most comparable to INTEC alluvium. Therefore, Cooper, et al. (2006) estimated a technetium K_d range of -0.1 to 1.4 mL/g for alluvium and a technetium K_d range of -0.1 to 0.1 mL/g for interbed sediments. Noting the lack of direct measurements, Cooper, et al. (2006) estimated a minimum and recommended technetium K_d for INTEC basalt of 0 mL/g.

Several studies have measured technetium K_d values in sediments from the Hanford site. The sediments and hydrogeochemical environment at the Hanford site are analogous to INL (i.e., alkaline pH, high carbonate concentrations, oxidizing conditions, and similar though not identical sediment mineral compositions). These studies are summarized next.

Kaplan, et al. (1998a,b) used batch equilibrium techniques to measure K_d values for technetium under oxic conditions on sediment samples from the Hanford site. The groundwater selected for the sorption studies was an uncontaminated groundwater for the Hanford site with a low ionic strength and pH of 8.4. The mean K_d values of three replicates measured for technetium on each Hanford sediment sample ranged from -0.04 to 0.01 mL/g. The measurements indicated essentially no sorption of technetium on Hanford sediments. Kaplan, et al. (1998b) suggested that the negative K_d values measured for technetium were due to anion exclusion effects.

Kaplan and Serne (1998) and Kaplan, et al. (1996) used batch techniques to measure the technetium K_d values, as TcO_4^- , on three sediments for the Hanford site. The measurements were conducted with a groundwater sample (pH 8.3) taken from a well located in an uncontaminated area of the Hanford site. The K_d values at 266 days of contact time were very low for all sediments and ranged from -0.18 to 0.11 mL/g.

Serne, et al. (1993) measured K_d values for TcO_4^- on three samples of sediment (two loamy sands and one sand) from the Hanford Formation. Batch equilibrium tests under oxidizing conditions were conducted using an uncontaminated groundwater (pH 8.14) sample from the Hanford site. Under the conditions studied, no significant sorption of technetium was measured; technetium K_d values ranged from 0.1 to 0.2 mL/g.

Measured and recommended technetium K_d values are summarized in Table 3-7. Site-specific technetium K_d values for alluvium and sedimentary interbeds from the column experiments of Del Debbio and Thomas (1989) indicate essentially no sorption of technetium. These results are in agreement with batch experiments of technetium sorption on sediments at the Hanford site, which showed technetium K_d values ranging from -0.04 to 0.2 mL/g (Kaplan, et al., 1998a,b, 1996; Kaplan and Serne, 1998; Serne, et al., 1993). Based on the column experiments of Del Debbio and Thomas (1989), a technetium K_d value of 0 mL/g is recommended for INL alluvium and interbed sediments. This value is in agreement with the technetium K_d value recommended by Cooper, et al. (2006) for INTEC alluvium and interbed sediments, which was also based on the column experiments of Del Debbio and Thomas (1989). A site-specific technetium K_d value of 0.016 mL/g was reported for basalt from the column experiments of Del Debbio and Thomas (1989). Lacking no other site-specific measurements, a technetium K_d value of 0 mL/g is recommneded for basalt at INL.

Table 3-7. Measured and Recommended Technetium K_d Values for Alluvium, Interbed Sediments, and Basalt at the Idaho National Laboratory

	Measured Technetium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Technetium K _d (mL/g)*	Recommended Technetium K _d (mL/g)
Alluvium	0†	Column	0	0
Sedimentary Interbeds	0†	Column	0	0
Basalt	0.016†	Column	0	0

^{*}Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

The position of the technetium K_d values recommended in this review with respect to measured and estimated site-specific and analogous technetium K_d ranges and values is graphically displayed in Figure 3-4. This figure illustrates the uncertainty associated with the technetium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific sorption data (Del Debbio and Thomas, 1989) and analogous technetium sorption data from the Hanford site (Kaplan, et al., 1998a,b; Kaplan and Serne, 1998; Serne, et al., 1993), the recommended technetium K_d values are associated with low uncertainty.

3.4.4 **lodine**

I-129 generally occurs as iodate (IO_3^-), iodide (I^-), or organic iodine in natural environments. Sorption of iodine species appears to be controlled in part by soil organic matter and in part by iron and aluminum oxides, with adsorption of iodine becoming increasingly important under more acid conditions. Although the extent of sorption is typically low, especially in systems containing little or no organic matter, I^- and IO_3^- are sorbed to a measurable extent by soils and some oxide minerals at near-neutral and alkaline pH conditions. The adsorption behavior of IO_3^- also appears to be appreciably different from that of I^- , in that IO_3^- sorbs much more strongly than I^- to soil and mineral surfaces (Ticknor and Cho, 1990). Mechanisms causing this sorption behavior at near-neutral and alkaline pH conditions are not completely understood. Some have proposed that this observed adsorption behavior in soils may be the result of the oxidation of I^- and/or reduction of IO_3^- to the more reactive molecular I_2^0 (aq) and/or its hydrolysis products (Ticknor and Cho, 1990).

[†]Del Debbio, J.A. and T.R. Thomas. "Transport Properties of Radionuclides and Hazardous Chemical Species in Soils at the Idaho Chemical Processing Plant." WINCO-1068. Idaho Falls, Idaho: Westinghouse Idaho Nuclear Company, Inc. 1989.

No site-specific measurement of iodine K_d values for alluvium, interbed sediment, or basalt at INL were found in the literature. Based on transport model calibration (i.e., visual fitting of a model predicted plume and an I-129 plume interpreted from measured data), Arnett, et al. (1990) estimated a retardation factor of 2 for I-129 at INL.

Cooper, et al. (2006) analyzed comparable systems and estimated iodine K_d values ranging from 0.04 to 8.7 mL/g with a recommended iodine K_d value of 1.5 mL/g for INTEC alluvium. For sedimentary interbeds, Cooper, et al. (2006) estimated iodine K_d values ranging from 0.04 to 3 mL/g with a recommended iodine K_d value of 0.7 mL/g. The estimated iodine K_d ranges for alluvium and interbed sediments are based on measured iodine K_d values from a field lysimeter experiment conducted by Sheppard and Thibault (1991). In soil horizons most comparable to INTEC alluvium (0.2 to 0.3 wt% organic carbon), iodine K_d values ranged from 0.1 to 8.7 mL/g at 1 year of transport and ranged from 0.04 to 0.06 mL/g at 4 years of transport. The recommended iodine K_d values for alluvium and interbed sediments are based on batch experiments conducted by Kaplan (2003) on two SRS sediments that are comparable to INTEC sediments. An upland sediment roughly corresponding to INTEC interbed sediments displayed an iodine K_d of 0.7 ± 0.1 mL/g. A wetland sediment roughly corresponding to INTEC alluvium (higher organic carbon) displayed an iodine K_d of 1.5 ± 0.5 mL/g. Noting the lack of direct measurements, Cooper, et al. (2006) estimated a minimum and recommended iodine K_d for INTEC basalt of 0 mL/g.

Several studies have measured iodine K_d values in sediments from the Hanford site. Kaplan, et al. (2000, 1996) performed batch experiments to measure the K_d values for iodine, as I^- , on three surficial sediment samples from the Hanford site. The measurements were conducted with a groundwater sample (pH 8.46) taken from a well located in an uncontaminated area of the Hanford site. The average measured K_d values at 330 days were 9.83, 6.83, and 4.72 mL/g for a very coarse sand, sand, and silty loam sediments, respectively. Kaplan, et al. (2000, 1996) noted that K_d values increased as a function of time; for example, K_d values increased from 0.19 to 9.83 over 330 days of contact time for the very coarse sand.

Kaplan, et al. (1998a) used batch equilibrium tests to measure the K_d values for I^- under oxic conditions on 20 sediment samples taken from a borehole in the Hanford formation at the Hanford site. The solution used in the tests was an uncontaminated groundwater with a low ionic strength and a pH of 8.4. Measured K_d values ranged from -0.03 to 0.23 mL/g and indicated essentially no sorption of I^- on the Hanford sediments under these conditions.

Serne, et al. (1993) conducted batch equilibrium tests to measure the K_d values for I^- on three surficial sediments: two loamy sands and one sand. The measurements were conducted using an uncontaminated groundwater (pH 8.14) sample from the Hanford site. After 35 days of contact time, the K_d values measured for I^- were 1.5 mL/g and 2.6 mL/g for the two loamy sand samples and 1.2 mL/g for the sand sample.

Hu, et al. (2005) conducted column studies of I^- and iodate (IO_3^-) sorption and transport on a sediment sample from the Hanford site. The sediment consisted of a composite of core samples taken at 9 to 12 m [30 to 80 ft] below surface. For I^- , retardation factors ranged from 1.05 to 1.33, which equate to K_d values of approximately 0.01 to 0.1 mL/g. For IO_3^- , retardation factors ranged from 1.41 to 1.93, which equate to K_d values of approximately 0.1 to 0.25 mL/g.

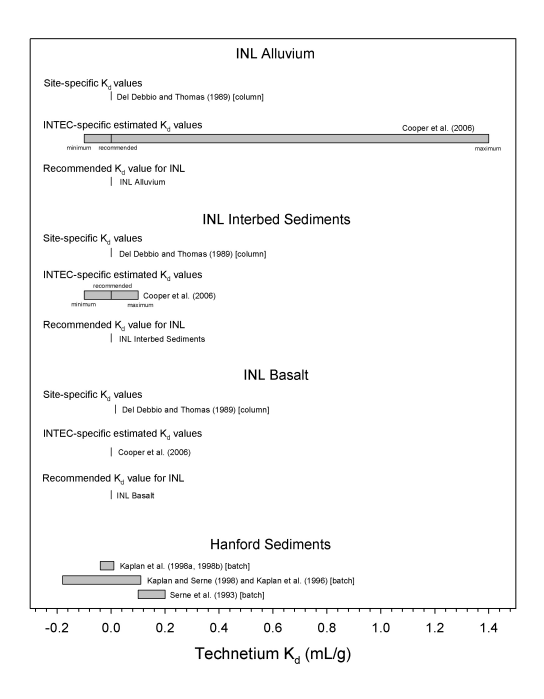


Figure 3-4. Measured and Estimated Site-Specific and Analogous Technetium $K_{\rm d}$ Ranges and Values and Recommended Technetium $K_{\rm d}$ Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-7). Recommended Technetium $K_{\rm d}$ Values Are Associated With Low Uncertainty Due to Available Site-Specific and Analogous Sorption Data.

Table 3-8. Red		dine $\mathbf{K}_{\scriptscriptstyle \mathrm{d}}$ Values for the Idaho Nation	or Alluvium, Interbed S onal Laboratory	ediments, and
	Measured lodine K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Iodine K _d (mL/g)*	Recommended lodine K _d (mL/g)
Alluvium	_	_	1.5	0.1
Sedimentary Interbeds		_	0.7	0.1
Basalt	_	_	0	0

*Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

Recommended iodine K_d values are summarized in Table 3-8. Based on an analysis of comparable systems, Cooper, et al. (2006) estimated that iodine K_d values for alluvium at INL should range from 0.04 to 8.7 mL/g and recommended an iodine K_d value of 1.5 mL/g. For interbed sediments, Cooper, et al. (2006) estimated iodine K_d values ranging from 0.04 to 3 mL/g and recommended an iodine K_d value of 0.7 mL/g. The range of iodine K_d values estimated by Cooper, et al. (2006) is within the range of iodine K_d values measured in batch experiments of sediments from the Hanford site (-0.03 to 9.84 mL/g) (Kaplan, et al., 2000, 1998a, 1996; Serne, et al., 1993), but is greater than the range of iodine K_d values measured in a column experiment using Hanford sediments (0.01 to 0.25 mL/g) (Hu, et al., 2005).

There is clear evidence that iodine is unlikely to travel unretarded in the INL subsurface. However, lacking any site-specific data, it is reasonable to employ a conservative K_d that represents the low end of values from subsurface samples from the analogous Hanford site

(Kaplan, et al., 1998a; Hu, et al., 2005). Therefore, an iodine K_d value of 0.1 mL/g is recommended for alluvium and sedimentary interbeds at INL. Lacking any direct or analogous measurements, the iodine K_d of 0 mL/g recommended by Cooper, et al. (2006) for INTEC basalt is recommended for basalt at INL.

The position of the iodine K_d values recommended in this review with respect to measured and estimated site-specific and analogous iodine K_d ranges and values is graphically displayed in Figure 3-5. This figure illustrates the uncertainty associated with the iodine K_d values recommended for alluvium, interbed sediments, and basalt at INL. Although analogous iodine sorption data is available for sediments from the Hanford site (Kaplan, et al., 2000, 1998a, 1996; Serne, et al., 1993; Hu, et al., 2005), the recommended iodine K_d values are associated with high uncertainty due to the lack of site-specific iodine sorption data.

3.4.5 Cesium

Cs-137 exists in the environment in the +1 oxidation state. Cesium forms few stable complexes and is likely to exist in groundwater as the uncomplexed Cs⁺ ion. Cs⁺ primarily adsorbs via ion exchange reactions with sedimentary clay minerals. In general, most soils sorb cesium rather strongly (Ames and Rai, 1978). The extent to which adsorption will occur will depend on the concentration of clays in the soil and the concentration of major cations, such as K⁺, that can effectively compete with Cs⁺ for adsorption sites. Cesium may also adsorb to iron oxides (Schwertmann and Taylor, 1989).

Schmalz (1972) performed batch experiments to measure cesium K_d values using alluvium from INL and a low ionic strength pond water with a pH of 9.2 collected from the Test Reactor Area. The pond water was spiked with varying concentrations of cesium. Cesium K_d values measured on 4 soil samples ranged from 589 to 1,253 mL/g.

Hawkins and Short (1965) performed batch tests to measure cesium K_d values on alluvial soil from INL as a function of solution chemistry. A solution for use in these tests was synthesized to simulate INL groundwater. The chemistry of this solution was then varied with respect to Ca^{2+} , Mg^{2+} , Cs^+ , K^+ , Na^+ , and NH_4^+ . K_d values measured for cesium ranged from 75 to 225 mL/g. Based on these tests, equations were developed to predict the sorption of cesium at varying solution chemistries.

Newman, et al. (1996) summarized results from short-term (400-hour) batch experiments and column tests measuring cesium K_d values for basalt and interbed sediments. The batch and column tests were conducted using a synthetic groundwater formulated to approximate the composition observed in SRPA field samples (ionic strength = 0.0048 M; pH 8.0) (Wood and Low, 1986). For batch tests, cesium K_d values ranged from 39 to 43 mL/g for basalt and from 2,230 to 3,260 mL/g for interbed sediments. For column tests, cesium K_d values ranged from 39 to 44 mL/g for basalt and >49 mL/g for interbed sediments. The cesium K_d value of >49 mL/g for interbed sediments represents a retardation factor of >200, which corresponds to the maximum number of pore volumes eluted during the column tests.

Fjeld, et al. (2001) performed laboratory column tests to characterize the mobility of cesium in a basalt sample and a composite of sedimentary interbed from INL. The solution used in the tests was a low ionic strength (0.004 M) synthetic groundwater (pH 8.0) formulated to approximate the major ion concentration of SRPA groundwater samples (Wood and Low, 1986). Cesium retardation factors ranged from 70 to 88 for basalt and >200 and >800 for interbed sediments. Based on the reported porosities and bulk densities of the basalt and interbed columns, these retardation factors equate to K_d values of 21 to 24.7 mL/g for basalt and >49 and >194 mL/g for interbed sediments.

Cooper, et al. (2006) estimated a cesium K_d range of 10 to 160 mL/g for alluvium and interbed sediments at INTEC and recommended a cesium K_d value of 50 mL/g for alluvium and interbed sediments. The estimated cesium K_d range for the alluvium and interbed sediments was calculated using the equation of Hawkins and Short (1965) and minimum and maximum values for measured geochemical parameters in INTEC alluvium and interbed sediments (see Cooper, et al., 2006, Table D–2–2). The recommended cesium K_d values of 50 mL/g for alluvium and

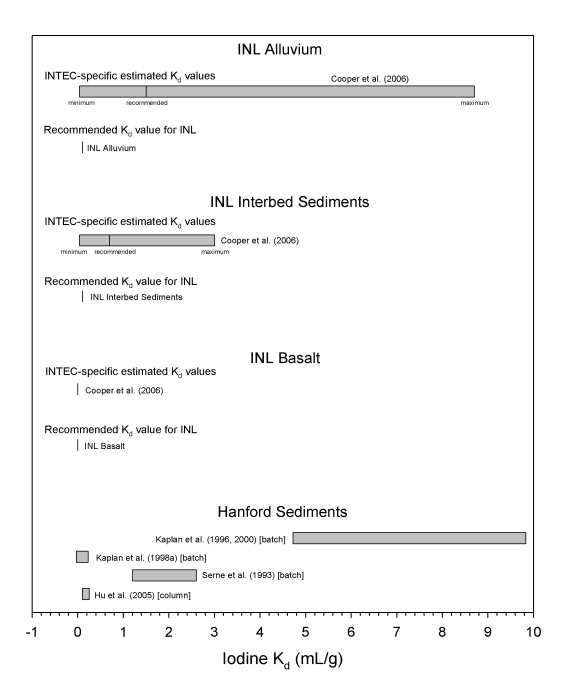


Figure 3-5. Measured and Estimated Site-Specific and Analogous Iodine K_d Ranges and Values and Recommended Iodine K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-8). Recommended Iodine K_d Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

interbed sediments were based on the study of Fjeld, et al. (2001). For basalt, Cooper, et al. (2006) estimated a cesium K_d range of 0 to 44 mL/g with a recommended cesium K_d value of 25 mL/g. The recommended cesium K_d value for basalt was based on the study of Fjeld, et al. (2001).

Researchers have measured cesium K_d values for sediments and basalt from the Hanford site. Serne, et al. (1993) conducted batch equilibrium tests to measure the K_d values for cesium on three surficial sediments: two loamy sands and one sand. The measurements were conducted using an uncontaminated groundwater (pH 8.14) sample from the Hanford site. The K_d values measured for cesium were 620 mL/g and 7,610 mL/g for the two loamy sand samples and 2,190 mL/g for the sand sample.

Ames, et al. (1982) conducted batch equilibrium tests to measure cesium sorption onto Umtanum, Flow E, and Pomona basalts from the Hanford site as a function of groundwater and aqueous cesium concentration. The pH of solutions used in the experiments ranged from 8.2 to 8.7; aqueous cesium contents ranged from approximately $2.5\times10^2~\mu\text{M}$ to approximately $2\times10^{-6}~\mu\text{M}$. For Umtanum basalt, cesium K_d values ranged from 27 to 1,084 mL/g using a 0.002 M solution and from 28 to 1,025 mL/g using a 0.013 M solution. For Flow E basalt, cesium K_d values ranged from 18 to 487 mL/g using a 0.002 M solution and from 20 to 617 mL/g using a 0.013 M solution. For Pomona Basalt, cesium K_d values ranged from 48 to 1,281 mL/g using a 0.002 M solution and from 56 to 902 mL/g using a 0.013 M solution. In all the tests cesium K_d values decreased with increasing cesium content; however, solution concentration had little effect on K_d values.

Measured and recommended cesium K_d values are summarized in Table 3-9. Site-specific cesium K_d values for alluvium from batch experiments range from a minimum of 75 mL/g (Hawkins and Short, 1965) to a maximum of 1,253 mg/L (Schmalz, 1972). These values are near the lower end of the range of cesium K_d values measured on surficial sediments from the Hanford site by batch methods (620 to 7,610 mL/g) (Serne, et al., 1993). Review of the laboratory procedures used by Schmalz (1972) to measure cesium K_d values on alluvium indicates that they may be biased high because of the water used in the measurements (i.e., low ionic strength pond water with a pH of 9.2). Therefore, the measured values of Schmalz (1972) may not be representative of natural chemical conditions in the alluvium. Removing the Schmalz (1972) strontium K_d values from the analysis results in a maximum measured K_d value of 225 mL/g for the alluvium (Hawkins and Short, 1965). Based on the range of cesium K_d values measured by Hawkins and Short (1965) on alluvial soil from INL (75 to 225 mL/g), a cesium K_d value of 75 mL/g is recommended for alluvium. This value is slightly higher than the cesium K_d value of 50 mL/g recommended by Cooper, et al. (2006). However, the cesium K_d value recommended by Cooper, et al. (2006) is based on column experiments conducted by Fjeld, et al. (2001) using interbed sediments rather than alluvium. In addition, cesium breakthrough was not achieved in the column experiments of Fjeld, et al. (2001) and the cesium K_d value of 50 mL/g cited by Cooper, et al. (2006) is considered to provide only an approximate lower boundary for cesium retention.

Table 3-9. Measured and Recommended Cesium \mathbf{K}_{d} Values for Alluvium, Interbed Sediments, and Basalt at the Idaho National Laboratory				
	Measured Cesium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Cesium K _d (mL/g)*	Recommended Cesium K _d (mL/g)
Alluvium	589–1,253† 75–225‡	Batch Batch	50	75
Sedimentary Interbeds	2,230-3,260§ >49§ >194¶	Batch Column Column	50	200
Basalt	39–43§ 39–44§ 21–24.7¶	Batch Column Column	25	25

^{*}Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

‡Hawkins, D.B. and H.L. Short. "Equations for the Sorption of Cesium and Strontium on Soil and Clinoptilolite." IDO–12046. Idaho Falls, Idaho: Atomic Energy Commission, Idaho Operations Office. 1965.

§Newman, M.E., I. Porro, R. Scott, R.M. Dunnivant, R.W. Goff, M.D. Blevins, S.M. Ince, J.D. Leyba, T.A. DeVol, A.W. Elzerman, and R.A. Fjeld. "Evaluation of the Mobility of Am, Cs, Co, Pu, Sr, and U Through INEL Basalt and Interbed Materials: Summary Report of the INEL/Clemson University Laboratory Studies." ER–WAG7–82, INEL–95/282. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1996.

¶Fjeld, R.A., T.A. DeVol, R.W. Goff, M.D. Blevins, D.D. Brown, S.M. Ince, A.W. Elzerman, and M.E. Newman. "Characterization of the Mobilities of Selected Actinides and Fission/activation Products in Laboratory Columns Containing Subsurface Material from the Snake River Plain." *Nuclear Technology*. Vol. 135. pp. 92–108. 2001.

For sedimentary interbed materials, site-specific cesium K_d values from a batch sorption experiment ranged from 2,230 to 3,260 mL/g (Newman, et al., 1996). Site-specific cesium K_d values from 2 column experiments using sedimentary interbed materials were >49 and >194 mL/g (Newman, et al., 1996; Fjeld, et al., 2001). In the column experiments, cesium breakthrough was not observed in the interbed sediments. The reported cesium K_d estimate of Fjeld, et al. (2001) (>194 mL/g) is based on gamma-spectroscopic analyses of intact cores and is considered to provide an approximate lower boundary for cesium retention. Based on the column experiment of Fjeld, et al. (2001), a cesium K_d value of 200 mL/g is recommended for interbed sediments at INL. This value is higher than the cesium K_d value of 50 mL/g recommended by Cooper, et al. (2006), which was also based on the column experiments of Fjeld, et al. (2001). However, Cooper, et al. (2006) did not cite the reported cesium K_d estimate of >194 mL/g based on gamma-spectroscopic analyses of intact cores from the Fjeld, et al. (2001) study, which is considered to provide a more accurate approximate lower boundary for cesium retention.

Site-specific cesium K_d values from a batch sorption experiment using crushed basalt from INL ranged from 39 to 43 mL/g (Newman, et al., 1996). These values are in agreement with

[†]Schmalz, B.L. "Radionuclide Distribution in Soil Mantle of the Lithosphere as a Consequence of Waste Disposal at the National Reactor Testing Station." IDO–10049. Idaho Falls, Idaho: United States Atomic Energy Commission, Idaho Operations Office. 1972.

site-specific cesium K_d values from two column experiments using crushed basalt from INL. One basalt column experiment reported cesium K_d values ranging from 39 to 44 mL/g (Newman, et al., 1996), and the other column experiment reported cesium K_d values ranging from 21 to 24.7 mL/g (Fjeld, et al., 2001). These values correspond to the low end of the range of cesium K_d values reported from batch experiments on basalts from the Hanford site (18 to 1,281 mL/g) (Ames, et al., 1982). Based on the range of cesium K_d values measured in the column experiments of Newman, et al. (1996) and Fjeld, et al. (2001) (21 to 44 mL/g), a cesium K_d value of 25 mL/g is recommended for basalt at INL. This value is in agreement with the cesium K_d value of 25 mL/g recommended by Cooper, et al. (2006), which was based on the column experiments of Fjeld, et al. (2001).

The position of the cesium K_d values recommended in this review with respect to measured and estimated site-specific and analogous cesium K_d ranges and values is graphically displayed in Figure 3-6. This figure illustrates the uncertainty associated with the cesium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific sorption data (Hawkins and Short, 1965; Newman, et al., 1996; Fjeld, et al., 2001), the recommended cesium K_d values are associated with low uncertainty.

3.4.6 Lead

Pb-210 has 3 known oxidation states—0, +2, and +4—and the most common redox state found in the environment is the divalent form. Dissolved lead in natural systems may exist in free ionic form (Pb²⁺) and also as hydrolytic and complex species. At pH values exceeding 7, aqueous lead exists mainly as carbonate complexes {Pb[(CO₃)⁰(aq)] and Pb(CO₂)₂²⁻}. Studies have shown that, in neutral to high pH conditions, Pb²⁺ can preferentially exchange for Ca²⁺ and K⁺ (Zimdahl and Hassett, 1977; Bittel and Miller, 1974). Under acidic pH conditions, hydrogen ions and aluminum ions will displace lead from mineral exchange sites. Lead is known to adsorb onto soil constituent surfaces such as clays, oxides, hydroxides, oxyhydroxides, and organic matter. Lead adsorption studies on bulk soils indicate that the adsorption is strongly correlated with pH and the CEC values of soils (Bittel and Miller, 1974; Griffin and Shimp, 1976; Haji-Djafari, et al., 1981; Scrudato and Estes, 1975; Zimdahl and Hassett, 1977). Properties that affect CEC of soils, such as organic matter content, clay content, and surface area, have a greater effect on lead adsorption than soil pH.

No site-specific data on lead sorption on geologic media at INL were found in the literature. Rodriguez, et al. (1997) analyzed systems comparable to the INL and estimated that lead K_d values for sediments and basalt at INL should range from 30 to 1,000 mL/g and 5 to 50 mL/g, respectively. Based on these ranges, Rodriguez, et al. (1997) recommended a lead K_d value of 270 mL/g for INL sediments and a lead K_d value of 10 mL/g for INL basalt.

Gerriste, et al. (1982) examined lead adsorption onto generic soils as a function of pH. Results indicated that lead adsorption increased with increasing soil pH (from 4 to 8). A summary of K_d values reported by Gerriste, et al. (1982) is listed in EPA (1999). Lead K_d values for pHs of 4.5, 5.0, 7.5, and 8.0 were 280; 1,295; 3,000; and 4,000 mL/g; respectively.

Rhoads, et al. (1992) investigated lead adsorption by a subsurface soil from the Hanford site. Adsorption data from these experiments showed that K_d values increased with decreasing lead

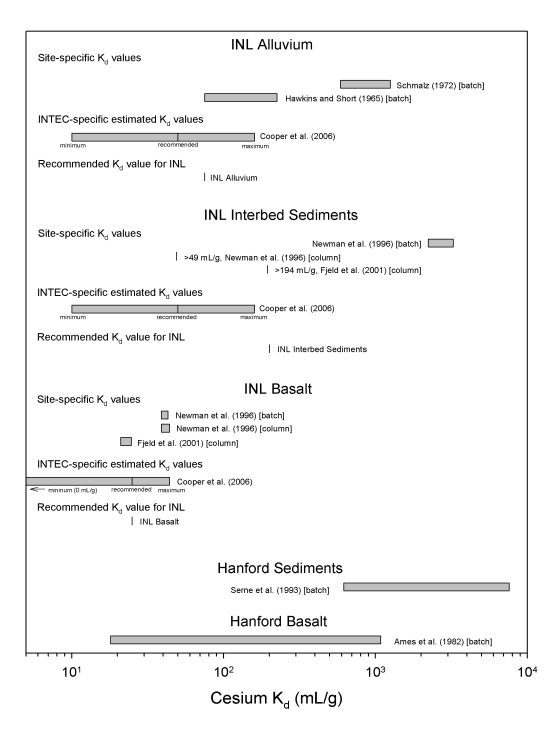


Figure 3-6. Measured and Estimated Site-Specific and Analogous Cesium $K_{\rm d}$ Ranges and Values and Recommended Cesium $K_{\rm d}$ Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-9). Recommended Cesium $K_{\rm d}$ Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

concentrations in solution {from 0.2 to 0.0001 mg/L [0.2 to 0.0001 ppm]}. At a fixed pH of 8.35, the authors found that K_d values, which ranged from 13,000 to 79,000 mL/g, were log-linearly correlated with equilibrium concentrations of lead in solution.

Based on the data sets reported by Gerriste, et al. (1982) and Rhoads, et al. (1992), a lookup table was developed to estimate the range of lead K_d values as a function of soil pH and equilibrium lead concentration (EPA, 1999). Gerriste, et al. (1982) measured adsorption data for 2 well-characterized soils using a range of lead concentrations {0.001 to 0.1 mg/L [0.001 to 0.1 ppm]}, which precluded the possibility of precipitation reactions. Similarly, the adsorption data reported by Rhoads, et al. (1992) encompassed a range of lead concentrations from 0.0001 to 0.2 mg/L [0.0001 to 0.2 ppm] at a fixed pH value. Both of these data sets were used for estimating the range of K_d values for the range of pH and aqueous lead concentration values associated with soils. For the range of pH values expected at INL (pH 6.4–8.7), equilibrium lead concentrations from 0.1 to 0.9 μ g/L [0.1 to 0.9 ppb], 1.0 to 9.9 μ g/L [1.0 to 9.9 ppb], 10 to 99.9 μ g/L [10 to 99.9 ppb], and 100 to 200 μ g/L [100 to 200 ppb] were estimated to have lead K_d values ranging from 4,360 to 23,270 mL/g; 1,950 to 10,760 mL/g; 900 to 4,970 mL/g; and 710 to 2,300 mL/g; respectively.

Veeresh, et al. (2003) performed batch equilibrium experiments to measure lead K_d values on three soils. A 0.005 M $Ca(NO_3)_2 \cdot 2H_2O$ solution was used in the batch experiments. Soils had varying particle size characteristics: a silty sand soil (85-wt% sand and silt, 15-wt% clay) at pH 5.1, a clay-rich silty sand soil (72-wt% sand and silt, 28-wt% clay) at pH 7.1, and a clay-rich soil (63-wt% clay, 27-wt% sand and silt) at pH 8.6. K_d values determined from sorption isotherms were 87 mL/g for the silty sand soil, 726 mL/g for the clay-rich silty sand soil, and 7,083 mL/g for the clay-rich soil.

Recommended lead K_d values are summarized in Table 3-10. No site-specific K_d data are available for lead adsorption on INL subsurface media. Based on an analysis of comparable systems, Rodriguez, et al. (1997) determined that lead K_d values in sediments at INL could range from 30 to 1,000 mL/g and recommended a lead K_d value of 270 mL/g. Measured lead K_d values from batch sorption studies at alkaline pH on generic soils (3,000 to 4,000 mL/g) (Gerriste, et al., 1982) and sediments from the Hanford site (13,000 to 79,000 mL/g) (Rhoads, et al., 1992) have much higher values than estimated by Rodriguez, et al. (1997). From the batch sorption experiments of Veeresh, et al. (2003), the silty sand soil and clay-rich silty sand soil noted previously are most comparable to sediments at INL, and the low ionic strength, calcium-rich solution used in the experiments is comparable to groundwater at INL. Measured K_d values for the silty sand soil and the clay-rich silty sand soil were 87 mL/g and 726 mL/g, respectively; however, the 87 mL/g value was from a pH 5.1 experiment. Considering the strong pH dependence of the lead K_d (e.g., Gerriste, et al., 1982), this especially low result can be discounted for the INL subsurface environment. The lowest value from the EPA (1999) lookup table for INL conditions was 710 mL/g. Based on these observations, the lead K_d value of 270 mL/g recommended by Rodriguez, et al. (1997) for sediments at INL is conservative and is recommended in this review for alluvium and sedimentary interbeds. Based on an analysis of comparable systems, Rodriguez, et al. (1997) estimated that lead K_d values in basalt at INL could range from 5 to 50 mL/g and recommended a lead K_d value of 10 mL/g. Lacking direct sorption data for lead, a lead K_d value of 10 mL/g is recommended for basalt.

Table 3-10.	Recommended Lead K _d Values for Alluvium, Interbed Sediments, and
	Basalt at the Idaho National Laboratory

	Measured Lead K _d (mL/g)	Test Method	Idaho National Laboratory-Specific Recommended Lead K _d (mL/g)*	Recommended Lead K _d (mL/g)
Alluvium	1	1	270	270
Sedimentary Interbeds	_	_	270	270
Basalt	_	_	10	10

^{*}Rodriguez, R.R., A.L. Schafer, J. McCarthy, P. Martian, D.E. Burns, D.E. Raunig, N.A. Burch, and R.L. VanHorn. DOE/ID–10534, "Comprehensive RI/FS for the Idaho Chemical Processing Plant OU 3-13 at the INEEL—Part A, RI/BRA Report (Final)." Idaho Falls, Idaho: Idaho National Engineering Laboratory, DOE–Idaho. 1997.

The position of the lead K_d values recommended in this review with respect to measured and estimated site-specific and analogous lead K_d ranges and values is graphically displayed in Figure 3-7. This figure illustrates the uncertainty associated with the lead K_d values recommended for alluvium, interbed sediments, and basalt at INL. The recommended lead K_d values are associated with high uncertainty due to the lack of site-specific lead sorption data.

3.4.7 Uranium

In natural environments, the most important oxidation states of uranium are +4 and +6. U(VI) species dominate in oxidizing environments; the dominant species are UO₂²⁺ at pH values less than 5 and mixed carbonate/hydroxyl and carbonate species at pH values between 5 and 9 (Langmuir, 1997). In low ionic strength solutions with low U(VI) concentrations, UO₂²⁺ concentrations will likely be controlled by cation exchange and adsorption processes. The uranyl ion and its complexes adsorb onto clays (Ames, et al. 1982), organics (Read, et al. 1993; Shanbhag and Choppin, 1981), and oxides (Hsi and Langmuir, 1985; Waite, et al., 1994). As ionic strength increases, other ions, notably Ca²⁺, Mg²⁺, and K⁺, displace the uranyl ion from soil exchange sites, forcing it into solution. U(VI) retention by soils and rocks in alkaline conditions is poor because of the predominance of neutral and negatively charged species. An increase in pCO₂ in soil solutions reduces U(VI) adsorption by promoting the formation of poorly sorbing carbonate complexes (Pabalan, et al., 1998). U(IV) species dominate in reducing environments. U(IV) tends to hydrolyze and form strong hydrolytic complexes. U(IV) also forms strong complexes with naturally occurring organic materials.

Several researchers have conducted sorption studies of uranium onto INL sediments and basalts. These studies are described next.

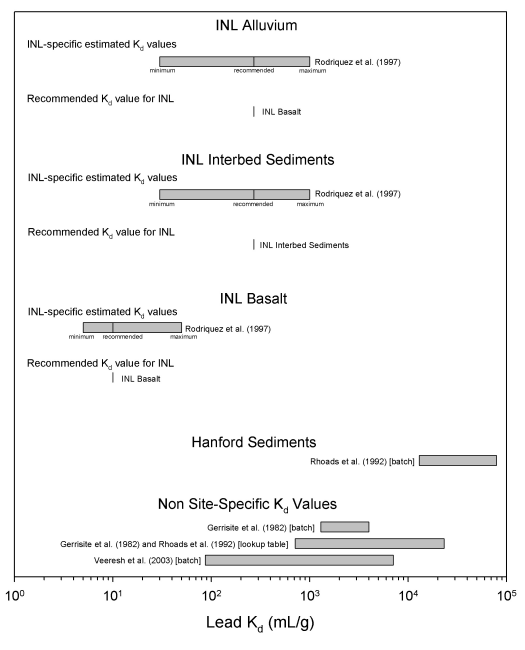


Figure 3-7. Measured and Estimated Site-Specific and Analogous Lead K_d Ranges and Values and Recommended Lead K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-10). Recommended Lead K_d Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

Leecaster and Hull (2004) conducted batch experiments to measure uranium K_d values on 36 SDA interbed sediment samples in a synthetic groundwater. The simulated groundwater had a low ionic strength, a pH of 8.0 ± 0.2 , and contained important cations and anions present in vadose zone waters at the SDA. Batch tests were carried out for time periods of 48 to 56 days. Measured uranium K_d values ranged from 0.6 to 48 mL/g with a median of 17.9 mL/g. When compared to sediment properties, uranium K_d values correlated most strongly to CEC, suggesting that surface reactions with clay minerals control uranium sorption chemistry in the INL subsurface.

Grossman, et al. (2001) conducted batch sorption experiments to characterize the sorption behavior of uranium at INL. Batch experiments were conducted with depth-discrete core samples of sedimentary interbed from beneath the Radioactive Waste Management Complex. The solution used for the batch experiments was a low ionic strength synthetic groundwater (pH 8.0) based on Wood and Low's (1986) characterization of the SRPA. Partitioning data were collected for between 48 and 56 days. Measured uranium K_d values ranged from 23 to 74 mL/g.

Newman, et al. (1996) summarized results from short-term (400-hour) batch experiments and column tests measuring uranium K_d values for basalt and interbed sediments. The batch and column tests were conducted using a synthetic groundwater formulated to approximate the composition observed in SRPA field samples (ionic strength = 0.0048 M; pH 8.0) (Wood and Low, 1986). For batch tests, uranium K_d values ranged from 4 to 6 mL/g for basalt and from 3 to 6 mL/g for interbed sediments. For column tests, uranium K_d values ranged from 0.2 to 0.3 mL/g for basalt and 7.34 to 11.43 mL/g interbed sediments.

Fjeld, et al. (2001) performed laboratory column tests to characterize the mobility of uranium in a basalt sample and a composite of sedimentary interbed from INL. The solution used in the tests was a low ionic strength (0.004 M) synthetic groundwater (pH 8.0) formulated to approximate the major ion concentration of SRPA groundwater samples (Wood and Low, 1986). Uranium retardation factors ranged from 5.1 to 6.1 for basalt and 30 to 31 for interbed sediments. Based on the reported porosities and bulk densities of the basalt and interbed columns, these retention factors equate to K_d values of 1.30 to 1.42 mL/g for basalt and 7.38 to 8.18 mL/g for interbed sediments.

An unpublished study of uranium transport through unsaturated sediments in a near field-scale column reactor packed with SDA surficial sediments (Cooper, et al., 2006) indicated that uranium $K_{\rm d}$ values can range from 4 to 50 mL/g.

Cooper, et al. (2006) estimated that uranium K_d values for alluvium and interbed sediments at INTEC should range from 0.12 to 12 mL/g and recommended a uranium K_d value of 1.6 mL/g for the alluvium and interbed sediments. The minimum and maximum uranium K_d values estimated by Cooper, et al. (2006) represented the range of values from batch and column experiments conducted on SDA sediments by Leecaster and Hull (2004), Grossman, et al. (2001), and Fjeld, et al. (2001), as scaled to CEC (INTEC/SDA = 1:5). The recommended uranium K_d value for alluvium and interbed sediments was based on the column experiments of Fjeld, et al. (2001), as scaled to CEC. Based on the column experiments of Fjeld, et al. (2001), Cooper, et al. (2006) estimated a uranium K_d range of 0 to 1.4 mL/g for INTEC basalt and recommended a uranium K_d value of 0 mL/g.

Researchers have measured uranium K_d s in basalt and surficial sediments from the Hanford site. These measurements are described next.

Kaplan and Serne (1995) and Lindenmeier, et al. (1995) performed unsaturated column tests to measure uranium sorption onto a loamy sand sediment taken from the Hanford site. The solution used in the experiments was a low ionic strength, low organic content, uncontaminated groundwater (pH 8.3) from the site. Kaplan and Serne (1995) reported measured K_d values ranging from 0.08 to 2.81 mL/g, and Lindenmeier, et al. (1995) reported K_d values ranging from 0.2 to 2.7 mL/g. In both studies, uranium K_d values typically increased with increasing degree of column saturation.

Kaplan, et al. (1998c) conducted batch tests to measure uranium sorption onto a silty loam and coarse sand taken from trenches at the Hanford site. The solution used in the tests was an uncontaminated groundwater from the Hanford site (pH 8.3). Measured K_d values ranged from 1.07 to 2.44 mL/g for the silty loam and from 0.14 to 0.2 mL/g for the coarse sand.

Kaplan, et al. (1996) performed batch and unsaturated column tests to measure uranium sorption onto surficial sediments from the Hanford site using an uncontaminated groundwater (pH 8.46). Measured K_d values ranged from 0.9 to 3.5 mL/g for the batch tests and 0.1 to 0.64 mL/g for the unsaturated column tests.

Ames, et al. (1982) conducted batch equilibrium tests to measure uranium sorption onto Umtanum, Flow E, and Pomona basalts from the Hanford site under oxidizing conditions using two synthetic groundwater solutions. The compositions of the solutions were based on those of groundwater samples taken at depth from the Columbia River basalt formations. Four initial concentrations of uranium were used for the measurements (1.0×10^{-4} , 1.0×10^{-5} , 1.0×10^{-6} , and 1.0×10^{-7} M uranium). The pH values of the final solutions ranged from 7.65 to 8.48. Measured uranium K_d values for the basalts ranged from 0.19 to 20 mL/g and decreased with increasing uranium content.

In a similar study, Salter, et al. (1981) conducted batch experiments to measure uranium sorption onto Columbia River basalts (Umtanum, Flow E, and Pomona basalts). Measurements were conducted with two synthetic groundwater solutions having compositions representative of the groundwater present in basalt formations at the Hanford site. K_d values were determined for initial concentrations of 1.0×10^{-4} , 1.0×10^{-5} , 1.0×10^{-6} , 1.0×10^{-7} , and 2.15×10^{-8} M uranium. Measured uranium K_d values for the basalts ranged from 0.2 to 56 mL/g, and as expected, decreased with increasing uranium content.

Measured and recommended uranium K_d values are summarized in Table 3-11. Measured uranium K_d values from column experiments conducted on INL alluvium range from 4 to 50 mL/g (Cooper, et al., 2006). Cooper, et al. (2006) also estimated a uranium K_d range of 0.12 to 12 mL/g for INTEC alluvium and recommended a uranium K_d value of 1.6 mL/g. The estimated uranium K_d range and recommended uranium K_d for alluvium reported by Cooper, et al. (2006) represent the range of uranium K_d values from batch and column experiments conducted on SDA sediments by Leecaster and Hull (2004), Grossman, et al. (2001), and Fjeld, et al. (2001), scaled to CEC (INTEC/SDA = 1:5). Measured uranium K_d values from batch and column experiments conducted on surficial sediments from Hanford have K_d values at the low end of the uranium K_d range estimated by Cooper, et al. (2006). Measured uranium K_d values from batch experiments range from 0.9 to 3.5 mL/g (Kaplan, et al., 1998c, 1996). Measured

Table 3-11. Measured and Recommended Uranium K_d Values for Alluvium, Interbed Sediments, and Basalt at the Idaho National Laboratory

	Measured Uranium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Uranium K _d (mL/g)*	Recommended Uranium K _d (mL/g)
Alluvium	4–50†	Column	1.6	1.6
Sedimentary Interbeds	0.6-48‡ 23-74§ 3-6¶ 7.43-11.43¶ 7.38-8.18#	Batch Batch Batch Column Column	1.6	1.6
Basalt	4–6¶ 0.2–0.3¶ 1.3–1.42#	Batch Column Column	0	0.2

*Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

 \dagger Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

 \ddagger Leecaster, M.K. and L.C. Hull. "Spatial Distribution of Neptunium and Uranium Partition Coefficients (K_d) for Interbed Sediments at a Radioactive Waste Subsurface Disposal Area." ICP/EXT-03-00088. Rev. 0. Idaho Falls, Idaho: Idaho National Engineering and Environmental Laboratory. 2004.

§Grossman, C.J., R.A. Fjeld, J.T. Coates, and A.W. Elzerman. "The Sorption of Selected Radionuclides in Sedimentary Interbed Soils from the Snake River Plain." INEEL/EXT–01–01106. Rev. 0. Idaho Falls, Idaho: Idaho National Engineering and Environmental Laboratory. 2001.

¶Newman, M.E., I. Porro, R. Scott, R.M. Dunnivant, R.W. Goff, M.D. Blevins, S.M. Ince, J.D. Leyba, T.A. DeVol, A.W. Elzerman, and R.A. Fjeld. "Evaluation of the Mobility of Am, Cs, Co, Pu, Sr, and U Through INEL Basalt and Interbed Materials: Summary Report of the INEL/Clemson University Laboratory Studies." ER–WAG7–82, INEL–95/282. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1996.

#Fjeld, R.A., T.A. DeVol, R.W. Goff, M.D. Blevins, D.D. Brown, S.M. Ince, A.W. Elzerman, and M.E. Newman. "Characterization of the Mobilities of Selected Actinides and Fission/activation Products in Laboratory Columns Containing Subsurface Material from the Snake River Plain." *Nuclear Technology.* Vol. 135. pp. 92–108. 2001.

uranium K_d values from column experiments range from 0.08 to 2.81 mL/g (Kaplan and Serne, 1995; Lindenmeier, et al., 1995). The uranium K_d value of 1.6 mL/g recommended by Cooper, et al. (2006) is within the range of reported uranium K_d values measured on surficial sediments from Hanford (0.08 to 3.5 mL/g). Based on the recommendation of Cooper, et al. (2006), a uranium K_d value of 1.6 mL/g is recommended for alluvium at INL.

Measured uranium K_d values from batch experiments conducted on INL sedimentary interbed materials range from 0.6 to 74 mL/g (Leecaster and Hull, 2004; Grossman, et al., 2001; Newman, et al., 1996). Measured uranium K_d values from column experiments conducted on INL interbed sediments vary over a much narrower range (7.38 to 11.43 mL/g) (Newman, et al., 1996; Fjeld, et al., 2001). Cooper, et al. (2006) scaled the uranium K_d values reported by

Leecaster and Hull (2004), Grossman, et al. (2001), and Fjeld, et al. (2001) with respect to observed differences in the CEC of INTEC and SDA interbed sediments (INTEC/SDA = 1:5). Based on this scaling, Cooper, et al. (2006), estimated a uranium K_d range of 0.12 to 12 mL/g for interbed sediments at INTEC and recommended a uranium K_d value of 1.6 mL/g. Based on the recommendation of Cooper, et al. (2006), a uranium K_d value of 1.6 mL/g is recommended for interbed sediments at INL.

Measured uranium K_d values from a batch experiment conducted on INL basalt range from 4 to 6 mL/g (Newman, et al., 1996). Measured uranium K_d values from column experiments conducted on INL basalt range from 0.2 to 1.42 mL/g (Newman, et al., 1996; Fjeld, et al., 2001). These values are comparable to the low end of uranium K_d values measured from batch experiments using basalt from the Hanford site (0.19 to 56 mL/g) (Ames, et al., 1982; Salter, et al., 1981). Based on the column experiments of Newman, et al. (1996) and Fjeld, et al. (2001), a uranium K_d value of 0.2 mL/g is recommended for basalt. This value is greater than the uranium K_d value of 0 mL/g recommended by Cooper, et al. (2006), which was based solely on the column experiments of Fjeld, et al. (2001).

The position of the uranium K_d values recommended in this review with respect to measured and estimated site-specific and analogous uranium K_d ranges and values is graphically displayed in Figure 3-8. This figure illustrates the uncertainty associated with the uranium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific uranium sorption data (e.g., Leecaster and Hull, 2004; Newman, et al., 1996; Fjeld, et al., 2001) and analogous uranium sorption data from the Hanford site (e.g., Kaplan and Serne, 1995; Kaplan, et al., 1998c, 1996), the recommended uranium K_d values are associated with low uncertainty.

3.4.8 Neptunium

Np-237 may exist in several valance states. Np(VI), Np(V), and Np(IV) are the most important valance states in natural waters. Over the pH range of most natural waters, Np(V) is dominant and present primarily as the cation NpO_2^+ . Np(V) aqueous species sorb to some extent to iron oxides and clays, but do not sorb to a major degree on most common minerals. Therefore, dissolved Np(V) is considered to be relatively mobile in soil systems. Because NpO₂⁺ does not compete favorably with dissolved Ca2+ and other divalent ions for adsorption sites on soils, the K_d values for Np(V) are relatively low (Kaplan and Serne, 2000). The adsorption of Np(V) is strongly dependent on pH. Typically, the sorption of Np(V) on iron oxides is negligible at pH values less than 5 and increases rapidly at pH values between 5 and 7. On silicates, the sorption of Np(V) is negligible at pH values less than 6 and increases rapidly at pH values between 6 and 8. This pH dependency is expected for ions present in solution primarily as cations. In carbonate-containing solutions, the adsorption of Np(V) on iron oxides and silicates has been observed to decrease at pH values greater than 7 to 9 and 8 to 9, respectively, in response to formation of aqueous Np(VI) carbonate complexes. Aqueous organic complexes can enhance Np(V) solubility, but sedimentary organic matter is not known to notably affect Np(V) sorption.

Mincher, et al. (2003) conducted batch experiments to investigate neptunium sorption onto SDA soils collected at depths of 0.5 to 6 m [1.6 to 20 ft] at INL. A simulated low ionic strength simulated groundwater (pH 7.55) was used in the batch experiments. At 3-hour contact times, neptunium K_d values of 49 \pm 12 mL/g were reported.

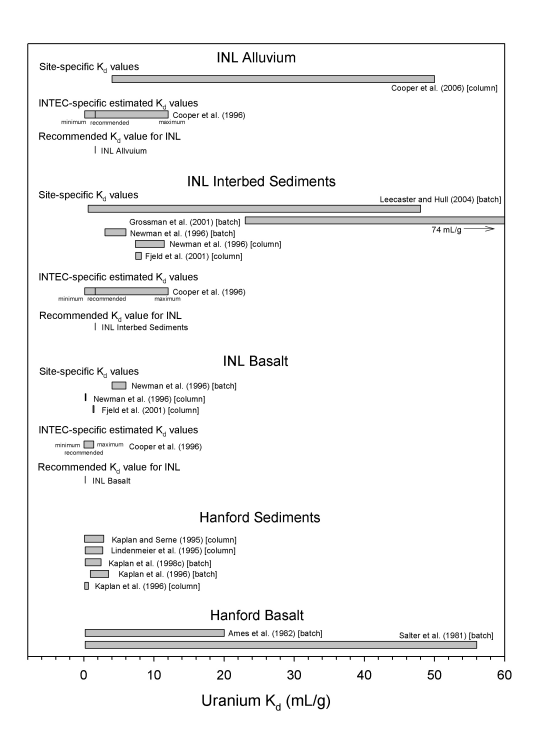


Figure 3-8. Measured and Estimated Site-Specific and Analogous Uranium K_d Ranges and Values and Recommended Uranium K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-11). Recommended Uranium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

Leecaster and Hull (2004) conducted batch experiments to measure neptunium K_d values on 36 SDA interbed sediment samples in a synthetic groundwater. The simulated groundwater had a low ionic strength, a pH of 8.0 \pm 0.2, and contained important cations and anions present in vadose zone waters at the SDA. Batch tests were carried out for time periods of 48 to 56 days. Measured neptunium K_d values ranged from 0.1 to 251 mL/g with a median of 35 mL/g. When compared to sediment properties, neptunium K_d values correlated most strongly to CEC, suggesting that surface reactions with clay minerals control neptunium sorption chemistry in the INL subsurface.

Grossman, et al. (2001) conducted batch sorption experiments to characterize the sorption behavior of neptunium at INL. Batch experiments were conducted with depth-discrete core samples of sedimentary interbed from beneath the Radioactive Waste Management Complex. The solution used for the batch experiments was a low ionic strength synthetic groundwater (pH 8.0) based on the Wood and Low (1986) SRPA characterization. Partitioning data were collected for 56 days. Measured neptunium K_d values ranged from 38 to 280 mL/g.

Dicke (1997) analyzed comparable systems and estimated that neptunium K_d values for sediments and basalt at INL should range from 1 to 80 mL/g and 1.3 to 51 mL/g, respectively. Based on these ranges, Dicke (1997) recommended a neptunium K_d value of 8 mL/g for both INL sediments and INL basalt.

Cooper, et al. (2006) estimated that neptunium K_d values for alluvium and interbed sediments at INTEC should range from 0.1 to 60 mL/g and recommended a neptunium K_d value of 2 mL/g for the alluvium and interbed sediments. The minimum and maximum neptunium K_d values estimated by Cooper, et al. (2006) represented the range of values from batch experiments conducted on SDA sediments by Mincher, et al. (2003), Leecaster and Hull (2004), and Grossman, et al. (2001), as scaled to CEC (INTEC/SDA = 1:5). The recommended neptunium K_d value of 2 mL/g was based on field lysimeter experiments conducted on a sandy sediment roughly comparable to INTEC alluvium and interbed sediments (Sheppard and Thibault, 1991). In these experiments, Sheppard and Thibault (1991) reported neptunium K_d values ranging from 0.5 to 1.6 mL/g at 1-year aging and 0.6 to 5 mL/g at 4-year aging with a median value of 2 mL/g. Lacking site-specific K_d data for basalt, Cooper, et al. (2006) assumed that the neptunium K_d value of 8 mL/g recommended by Dicke (1997) for basalt provided a reasonable maximum value and recommended a neptunium K_d value of 0 mL/g for basalt.

Researchers have measured neptunium K_d values in surficial sediments from the Hanford site. Serne, et al. (1993) measured K_d values for Np(V) in groundwater contacting surficial sediments from the Hanford site. The pH of the Hanford groundwater was 8.3. The sediment used in the experiment contained 87-wt% sand, 7-wt% silt, and 6-wt% clay. The K_d values measured at 5 and 44 days were 13.5 \pm 3 and 29.1 \pm 3.6 mL/g, respectively.

Kaplan, et al. (1996) conducted batch experiments to measure the K_d values for Np(V) on three surficial sediment samples from Hanford. The measurements were conducted with a groundwater (pH 8.46) taken from an uncontaminated well on the Hanford site. Sediments included a sand, silty loam, and very coarse sand. Kaplan, et al. (1996) observed that the K_d values increased as a function of time. K_d values measured at 7 days were 2.17 mL/g for the sand, 2.67 mL/g for the silty loam, and 14.17 mL/g for the coarse sand. K_d values measured at 77 days were 3.62 mL/g for the sand, 13.48 mL/g for the silty loam, and 19.86 mL/g for the coarse sand.

Measured and recommended neptunium K_d values are summarized in Table 3-12. Mincher, et al. (2003) measured site-specific neptunium K_d values ranging from 37 to 61 mL/g for alluvium at INL. Measured neptunium K_d values for surficial sediments from Hanford determined using batch methods ranged from 2.17 to 29.1 mL/g (Kaplan, et al., 1996; Serne, et al., 1993). Based on field lysimeter experiments conducted by Sheppard and Thibault (1991) on a sandy sediment roughly comparable to INTEC alluvium, Cooper, et al. (2006) recommended a neptunium K_d of 2 mL/g for INTEC alluvium. The value is comparable to the low end of the range of neptunium K_d values measured on surficial sediments at Hanford (2.17 to 29.1 mL/g). Based on the neptunium K_d recommendation of Cooper, et al. (2006), a neptunium K_d value of 2 mL/g is recommended for alluvium at INL.

Measured neptunium K_d values for sedimentary interbed materials from batch experiments range from a minimum of 0.1 mL/g (Leecaster and Hull, 2004) to a maximum of 280 mg/L (Grossman, et al., 2001). Based on an analysis of comparable systems, Dicke (1997) estimated a neptunium K_d range of 1 to 80 mL/g for INL sediments and recommended a neptunium K_d of 8 mL/g. Based on field lysimeter experiments conducted by Sheppard and Thibault (1991) on a sandy sediment roughly comparable to INTEC interbed sediments, Cooper, et al. (2006) recommended a neptunium K_d of 2 mL/g for INTEC interbeds. Based on these observations, a neptunium K_d value of 5 mL/g is recommended for sedimentary interbed materials at INL. This value is the midpoint between the neptunium K_d of 8 mL/g recommended by Dicke (1997) for

Table 3-12. Measured and Recommended Neptunium K _d Values for Alluvium, Interbed
Sediments, and Basalt at the Idaho National Laboratory

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	Measured Neptunium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Neptunium K _d (mL/g)*	Recommended Neptunium K _d (mL/g)
Alluvium	37–61†	Batch	2	2
Sedimentary Interbeds	0.1–251‡ 38–280§	Batch Batch	2	5
Basalt	_	_	0	0.2

^{*}Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

[†]Mincher, B.J., R.V. Fox, D.C. Cooper, and G.S. Groenewold. "Neptunium and Plutonium Sorption to Snake River Plain, Idaho Soil." *Radiochimica Acta*. Vol. 91. pp. 397–401. 2003.

[‡]Leecaster, M.K. and L.C. Hull. "Spatial Distribution of Neptunium and Uranium Partition Coefficients (K_d) for Interbed Sediments at a Radioactive Waste Subsurface Disposal Area." ICP/EXT–03–00088. Rev. 0. Idaho Falls, Idaho: Idaho National Engineering and Environmental Laboratory. 2004.

[§]Grossman, C.J., R.A. Fjeld, J.T. Coates, and A.W. Elzerman. "The Sorption of Selected Radionuclides in Sedimentary Interbed Soils from the Snake River Plain." INEEL/EXT–01–01106. Rev. 0. Idaho Falls, Idaho: Idaho National Engineering and Environmental Laboratory. 2001.

INL sediments and the neptunium K_d of 2 mL/g recommended by Cooper, et al. (2006) for INTEC interbed sediments.

No site-specific neptunium sorption data are available for basalt from INL. Dicke (1997) estimated a neptunium K_d range of 1.3 to 51 for INL basalt based on analyses of comparable systems and recommended a neptunium K_d of 8 mL/g. Cooper, et al. (2006) recommended a neptunium K_d value of 0 mL/g for INTEC basalt. Beasley, et al. (1998) observed migration of contaminant radionuclides in the SRPA fractured basalt aquifer and calculated relative concentrations of radionuclides along the transport pathway. Based on limited U-236 and Np-237 results, and consistent with an earlier basalt aquifer modeling study, Beasley, et al. (1998) concluded that neptunium and uranium were retarded similarly. Therefore, we recommend the basalt K_d for neptunium be the same as for uranium—0.2 mL/g.

The position of the neptunium K_d values recommended in this review with respect to measured and estimated site-specific and analogous neptunium K_d ranges and values is graphically displayed in Figure 3-9. This figure illustrates the uncertainty associated with the neptunium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific neptunium sorption data (e.g., Leecaster and Hull, 2004; Grossman, et al., 2001) and analogous neptunium sorption data from the Hanford site (Serne, et al., 1993; Kaplan, et al., 1996), the recommended neptunium K_d values are associated with low uncertainty.

3.4.9 Plutonium

In the range of pH and conditions typically encountered in natural waters, Pu-239 can exist in all 4 oxidation states, namely +3, +4, +5, and +6. Under oxidizing conditions, Pu(IV), Pu(V), and Pu(VI) are common, whereas under reducing conditions, Pu(III) and Pu(IV) exist. Plutonium is known to adsorb onto soil components such as clays, oxides, hydroxides, oxyhydroxides, aluminosilicates, and organic matter. Depending on the properties of the substrates, pH, and the composition of solutions, plutonium would adsorb with affinities varying from low $(K_d = 10 \text{ mL/g})$ to extremely high $(K_d = 300,000 \text{ mL/g})$ (Baes and Sharp, 1983; Thibault, et al., 1990). Under conditions of low pH and high concentrations of dissolved organic carbon, it appears that plutonium-organic complexes may control adsorption and mobility of plutonium. At neutral to alkaline pH (>7), dissolved plutonium forms very strong hydroxy–carbonate mixed ligand complexes, resulting in desorption and increased mobility in the environment. Studies have shown the potential importance of colloid-facilitated transport of plutonium in soil systems. Laboratory studies by Penrose, et al. (1990) predicted that the movement of plutonium would be limited to less than a few meters through a shallow aquifer within the site of the Los Alamos National Laboratory. However, plutonium was detected in monitoring wells as far as 3,390 m [11,120 ft] downgradient from the point source. Almost all of the plutonium in the groundwater at the 3,390-m [11,120-ft] well was associated with colloids.

A number of studies indicate that iron hydroxides adsorb and reduce Pu(V) and Pu(VI) to its tetravalent state at the solid surface. Fjeld, et al. (2003) have described a conceptual model for plutonium transport through sediment columns where plutonium transport is controlled by the net rate of reduction of adsorbed Pu(V) to adsorbed Pu(IV). In this model, oxidized plutonium can mobilize as Pu(V), and the Pu(V) can readsorb onto iron bearing minerals and be slowly reduced to Pu(IV). In this way, kinetic control of plutonium transport can allow a small fraction to move while the bulk of plutonium is largely immobile. This type of nonequilibrium behavior is

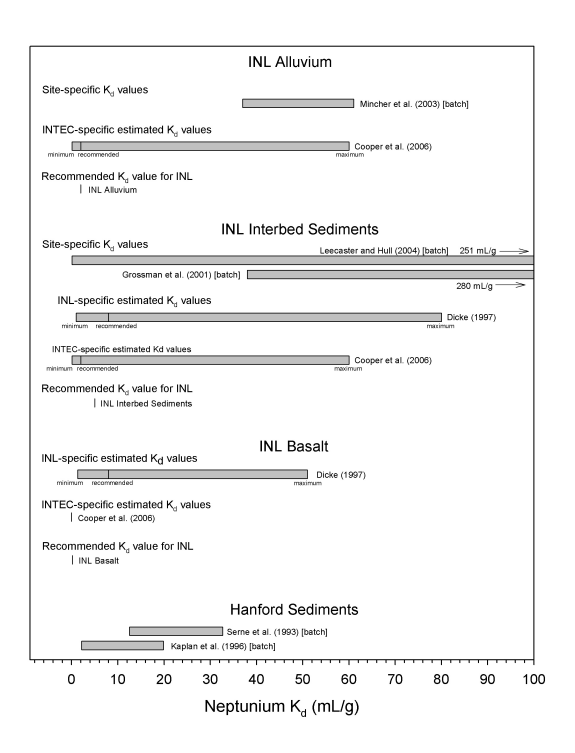


Figure 3-9. Measured and Estimated Site-Specific and Analogous Neptunium $K_{\rm d}$ Ranges and Values and Recommended Neptunium $K_{\rm d}$ Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-12). Recommended Neptunium $K_{\rm d}$ Values Are Associated With Low Uncertainty Due to Available Site-Specific and Analogous Sorption Data.

difficult to capture with a K_d approach, but K_d can be used if the system is assumed to have reached steady state with respect to plutonium redox kinetics.

Mincher, et al. (2003) conducted batch experiments to investigate plutonium sorption onto SDA soils collected at depths of 0.5 to 6 m [1.6 to 20 ft] at INL. A low ionic strength simulated groundwater (pH 7.55) was used in the batch experiments. At 3-hour contact times, plutonium K_d values of 54 \pm 26 mL/g were reported.

Newman, et al. (1996) summarized results from short-term (400-hour) batch experiments measuring plutonium K_d values for basalt, interbed sediments, and surficial soils and column tests measuring plutonium K_d values for basalt and interbed sediments. The batch and column tests were conducted using a synthetic groundwater formulated to approximate the composition observed in SRPA field samples (ionic strength = 0.0048 M; pH 8.0) (Wood and Low, 1986). A series of batch tests was also conducted on basalt, interbed sediments, and surficial soils in the presence of EDTA, a strong chelating agent that forms negative or neutral complexes with metals. For batch tests, $Pu(V) K_d$ values ranged from 70 to 130 mL/g for basalt; 5,100 to 7,900 mL/g for interbed sediments; and 7,800 to 22,000 mL/g for surficial soils. Pu(VI) K_d values for the batch tests ranged from 12 to 24 mL/g for basalt; 110 to 690 mL/g for interbed sediments; and 1,800 to 4,900 mL/g for surficial soils. Plutonium K_d values in the presence of EDTA for the batch tests ranged from 0.04 to 0.05 mL/g for basalt, 4 to 5 mL/g for interbed sediments, and 40 to 490 mL/g for surficial soils. However, organic complexants such as EDTA are not expected to be important for subsurface plutonium transport at INL. For column tests. total plutonium mobility was characterized by high and low mobility fractions. For basalt, the high mobility fraction (10 to 55 percent) had an estimated K_d value of 0.17 mL/g and a remaining low mobility fraction with an estimated K_d of >60 mL/g. For interbed sediments, the high mobility fraction (~1 percent) was too small to determine a K_d value, and the remaining low mobility fraction had an estimated K_d of >47 mL/g.

Fjeld, et al. (2001) performed laboratory column tests to characterize the mobility of plutonium in a basalt sample and a composite of sedimentary interbed from INL. The solution used in the tests was a low ionic strength (0.004 M) synthetic groundwater (pH 8.0) formulated to approximate the major ion concentration of SRPA groundwater samples (Wood and Low, 1986). Plutonium mobility was characterized by two fractions in both basalt and interbed sediments. For basalt, a high mobility fraction (17 to 29 percent) had retardation factors ranging from 1.2 to 5.0, and a remaining low mobility fraction had a retardation factor >200. For interbed sediments, a high mobility fraction (1.1 to 2.4 percent) had retardation factors of ~3, and a remaining low mobility fraction had a retardation factor >200. Based on reported porosities and bulk densities of packed columns, retardation factors for basalt equate to K_d values ranging from 0.05 to 1.14 mL/g for the high mobility fraction and >60 mL/g for the low mobility fraction and of rinterbed sediments a K_d value of ~0.46 mL/g for the high mobility fraction and >47 mL/g for the low mobility fraction.

Dicke (1997) analyzed comparable systems and estimated a plutonium K_d range of 5,100 to 22,000 mL/g for sediments at INL and a plutonium K_d range of 70 to 130 mL/g for basalt. Dicke (1997) recommended a plutonium K_d value of 5,100 mL/g for sediments, and a value of 100 mL/g for basalt. J. Navratil, cited in Cooper, et al. (2006), reviewed historical studies of plutonium sorption on SDA interbed sediments and recommended a plutonium K_d between 300 and 3,500 mL/g for interbed material. Miner, et al. (1982) reported plutonium K_d values that range from 120 to 5,000 mL/g for batch experiments with Snake River Plain soils.

Cooper, et al. (2006) estimated a plutonium K_d range of 96 to 12,700 mL/g for alluvium and interbed sediments at INTEC and recommended a plutonium K_d value of 1,000 mL/g. The minimum and maximum plutonium K_d values estimated by Cooper, et al. (2006) represented the range of values from batch experiments conducted by Miner, et al. (1982) and Mincher, et al. (2003) and estimated plutonium K_d values for INL sediments reported by Dicke (1997). The recommended plutonium K_d value of 1,000 mL/g for alluvium and interbed sediments was based on a lysimeter experiment using a sandy clay sediment roughly comparable to INTEC alluvium and interbed sediments (Kaplan, et al., 2004). Results of the lysimeter experiment indicated a retention factor of 3,000 for Pu(IV), which equated to a K_d of 1,000 mL/g. Cooper, et al. (2006) estimated a plutonium K_d range of 0 to 130 mL/g for INTEC basalt and recommended a plutonium K_d value of 70 mL/g. The plutonium K_d value of 70 mL/g recommended by Cooper, et al. (2006) for basalt is equal to the minimum value estimated by Dicke (1997).

Researchers have measured plutonium $\rm K_d$ values in surficial sediments and basalt from the Hanford site. Barney (1992) studied sorption of plutonium in +4, +5, and +6 redox states on a shallow sediment from the Hanford site to elucidate any differences in rate and amount of plutonium sorption in different redox states. A synthetic groundwater was used in the experiments, and initial plutonium concentrations were varied between about 10^{-11} and 10^{-9} M. The data indicated that the $\rm K_d$ values ranged from 2,100 to 11,600 mL/g; 2,700 to 4,600 mL/g; and 1,000 to 4,600 mL/g for plutonium in +4, +5, and +6 redox states, respectively. The data also indicated that Pu(V) and Pu(VI) upon adsorption were reduced to the tetravalent state.

Ames, et al. (1981) and Salter, et al. (1981) conducted batch equilibrium tests to measure plutonium sorption onto Umtanum Basalt from the Hanford site. Measurements were conducted with synthetic groundwater solutions having compositions representative of the groundwater present in basalt formations at the Hanford site. Ames, et al. (1981) reported a plutonium K_d value of 20 mL/g, and Salter, et al. (1981) reported plutonium K_d values ranging from 20 to 25 mL/g.

Measured and recommended plutonium K_d values are summarized in Table 3-13. Site-specific total plutonium K_d values for alluvium from INL determined by batch methods range from 28 to 80 mL/g (Mincher, et al., 2003) at 3-hour contact times. These experiments may have been too short to provide reliable K_d values. Newman, et al. (1996) conducted batch experiments to measure plutonium sorption on INL alluvium at 400-hour contact times and reported Pu(V) K_d values ranging from 7,800 to 22,000 mL/g; Pu(VI) K_d values ranging from 1,800 to 4,900 mL/g; and total plutonium K_d values ranging from 40 to 490 mL/g in the presence of EDTA. Miner, et al. (1982) reported total plutonium K_d values ranging from 120 to 5,000 mL/g for Snake River Plain soils. Based on the 400-hour batch experiments of Newman, et al. (1996) and considering the range of plutonium K_d values reported by Miner, et al. (1982), a total plutonium K_d value of 1,000 mL/g is recommended for alluvium at INL. This value is in agreement with the plutonium K_d value of 1,000 mL/g recommended by Cooper, et al. (2006) for INTEC alluvium.

For sedimentary interbeds at INL, Newman, et al. (1996) conducted batch experiments and reported Pu(V) K_d values ranging from 5,100 to 7,900 mL/g; Pu(VI) K_d values ranging from 110 to 690 mL/g; and plutonium K_d values ranging from 4 to 5 mL/g in the presence of EDTA. J. Navratil, cited in Cooper, et al. (2006), recommended a total plutonium K_d between 300 and

Table 3-13. Measured and Recommended Plutonium K_d Values for Alluvium, Interbed Sediments, and Basalt at the Idaho National Laboratory

	Measured Plutonium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Plutonium K _d (mL/g)*	Recommended Plutonium K _d (mL/g)
Alluvium Pu(V) Pu(VI) Pu(EDTA) Pu(total)	7,800–22,000† 1,800–4,900† 40–490† 28–80‡ 120–5,000§	Batch Batch Batch Batch Batch	1,000	1,000
Sedimentary Interbeds Pu(V) Pu(VI) Pu(EDTA) Pu(total)	5,100-7,900† 110-690† 4-5† >47† >47¶ 0.46¶ (hmf)#	Batch Batch Batch Column Column Column	1,000	500 0.5 (hmf)#
Basalt Pu(V) Pu(VI) Pu(EDTA) Pu(total)	7-130† 12-24† 0.04-0.05† >60† 0.17† (hmf)# >60¶ 0.05-1.14¶ (hmf)#	Batch Batch Batch Column Column Column Column	70	70 0.2 (hmf)#

*Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

†Newman, M.E., I. Porro, R. Scott, R.M. Dunnivant, R.W. Goff, M.D. Blevins, S.M. Ince, J.D. Leyba, T.A. DeVol, A.W. Elzerman, and R.A. Fjeld. "Evaluation of the Mobility of Am, Cs, Co, Pu, Sr, and U Through INEL Basalt and Interbed Materials: Summary Report of the INEL/Clemson University Laboratory Studies." ER–WAG7–82, INEL–95/282. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1996.

‡Mincher, B.J., R.V. Fox, D.C. Cooper, and G.S. Groenewold. "Neptunium and Plutonium Sorption to Snake River Plain, Idaho Soil." *Radiochimica Acta*. Vol. 91. pp. 397–401. 2003.

§Miner, F.J., P.A. Evans, and W.L. Polzer. "Plutonium Behavior in the Soil/water Environment." RFP–2480. Golden, Colorado: Rockwell International. 1982.

¶Fjeld, R.A., T.A. DeVol, R.W. Goff, M.D. Blevins, D.D. Brown, S.M. Ince, A.W. Elzerman, and M.E. Newman. "Characterization of the Mobilities of Selected Actinides and Fission/activation Products in Laboratory Columns Containing Subsurface Material from the Snake River Plain." *Nuclear Technology*. Vol. 135. pp. 92–108. 2001. #(hmf) denotes a high mobility fraction.

3,500 mL/g for interbed sediments based on a review of historical studies. Total plutonium K_d values for interbed sediments determined from column experiments were >47 ml/g (Newman, et al., 1996; Fjeld, et al., 2001). High mobility fractions of plutonium were detected in the column experiments. In the column experiments of Newman, et al. (1996), the high mobility fraction was too small (~1 percent) to determine a K_d . In the column experiments of Fjeld, et al. (2001), the high mobility plutonium fraction was 1.1 to 2.4 percent and had a K_d value of 0.46 mL/g. Based on the recommended plutonium K_d values of J. Navratil, cited in Cooper, et al. (2006), and considering the results of column experiments (plutonium K_d >47 mL/g), a plutonium K_d of 500 mL/g is recommended for interbed sediments. This value was chosen to be lower than the alluvium K_d based on the overall lower values for interbed sediments when compared to alluvium (Table 3-13). Use of the lower Pu(VI) values is not recommended in light of evidence for surface reduction of sorbed plutonium. A K_d value of 0.5 mL/g is recommended for the high mobility fraction of plutonium in interbed sediments. This high plutonium mobility fraction likely results from colloid-facilitated transport, and its effects should be considered in groundwater dose models.

For INL basalt, Newman, et al. (1996) conducted batch sorption experiments and reported K_d values for Pu(V) ranged from 7 to 130 mL/g, Pu(VI) K_d values ranged from 12 to 24 mL/g, and plutonium K_d values in the presence of EDTA ranged from 0.04 to 0.05 mL/g. The Pu(V) and Pu(VI) K_d values measured by Newman, et al. (1996) are in good agreement with the range of plutonium K_d values measured for basalt from Hanford (20 to 25 mL/g) using batch techniques (Ames, et al., 1981; Salter, et al., 1981). Total plutonium K_d values for basalt determined from column experiments were >60 mL/g (Newman, et al., 1996; Fjeld, et al., 2001). In the column experiments of Newman, et al. (1996), the high mobility plutonium fraction was 10 to 55 percent and had an estimated K_d value of 0.17 mL/g. In the column experiments of Fjeld, et al. (2001), the high mobility plutonium fraction was 17 to 29 percent and had estimated K_d values ranging from 0.05 to 1.14 mL/g. Based on the basalt column experiments of Newman, et al. (1996) and Fjeld, et al. (2001), a plutonium K_d of 70 mL/g is recommended for basalt. This value is in agreement with the plutonium K_d value of 70 mL/g recommended by Cooper, et al. (2006) for INTEC basalt. Use of the lower Pu(VI) values is not recommended in light of evidence for surface reduction of sorbed plutonium. A K_d value of 0.2 mL/g is recommended for the high mobility fraction of plutonium in basalt. This high plutonium mobility fraction likely results from colloid-facilitated transport, and its effects should be considered in groundwater dose models.

The position of the plutonium K_d values recommended in this review with respect to measured and estimated site-specific and analogous plutonium K_d ranges and values is graphically displayed in Figure 3-10. This figure illustrates the uncertainty associated with the plutonium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific plutonium sorption data (e.g., Newman, et al., 1996; Fjeld, et al., 2001) and analogous plutonium sorption data from the Hanford site (e.g., Barney, 1992; Salter, et al., 1981), the recommended plutonium K_d values are associated with low uncertainty.

3.4.10 Americium

Am-241 can exist in the +3, +4, +5, and +6 valance states. But, americium predominantly occurs as Am(III) in natural environments. Sorption studies indicate that Am(III) readily sorbs to minerals, crushed rock, and soil materials and, along with plutonium, is considered one of the most immobile actinide elements in the environment (Lu, et al., 1998; Ticknor, et al., 1996).

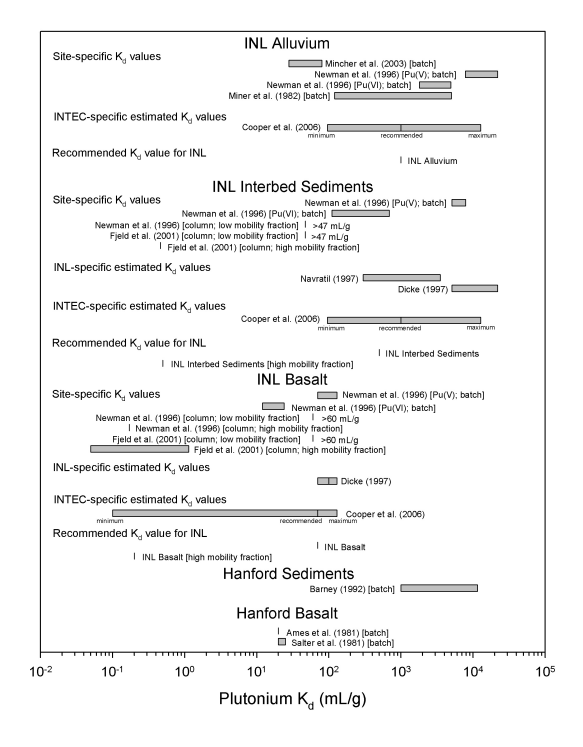


Figure 3-10. Measured and Estimated Site-Specific and Analogous Plutonium K_d Ranges and Values and Recommended Plutonium K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-13). Recommended Plutonium K_d Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

Based on studies of Am(III) on single mineral phases and crushed rock, the adsorption of Am(III) is strongly pH dependent, and increases with increasing pH, with peak adsorption occurring between pH values of 5 and 6. This observed pH dependence is expected, because the dominant aqueous species of americium in the pH range of natural waters are primarily Am³+ and cationic carbonate complexes at acidic and basic pH values, respectively. High levels of soil organic matter can enhance americium mobility (Artinger, et al., 1998; Schuessler, et al., 2000; Ticknor, et al., 1996). In soil environments, clay minerals are expected to control americium sorption, and K_d values are expected to correlate with CEC. Studies have shown the potential importance of colloid-facilitated transport of americium in soil systems (Penrose, et al., 1990).

Newman, et al. (1996) summarized results from short-term (400-hour) batch experiments and column tests measuring americium K_c values for basalt and interbed sediments. The batch and column tests were conducted using a synthetic groundwater formulated to approximate the composition observed in SRPA field samples (ionic strength = 0.0048 M; pH 8.0) (Wood and Low, 1986). For batch tests, americium K_d values ranged from 70 to 280 mL/g for basalt and from 450 to 1,100 mL/g for interbed sediments. For column tests, americium mobility was characterized by high and low mobility fractions. For basalt, the high mobility fraction (9 to 13 percent) had an estimated K_d value of 0.18 mL/g and a remaining low mobility fraction with an estimated K_d of >60 mL/g. For interbed sediments, the high mobility fraction (~1 percent) was too small to determine a K_d , and the remaining low mobility fraction had an estimated K_d >49 mL/g.

Fjeld, et al. (2001) performed laboratory column tests to characterize the mobility of americium in a basalt sample and a composite of sedimentary interbed from INL. The solution used in the tests was a low ionic strength (0.004 M) synthetic groundwater (pH 8.0) formulated to approximate the major ion concentration of SRPA groundwater samples (Wood and Low, 1986). For basalt, americium mobility was characterized by two fractions: a high mobility fraction (7 to 12 percent) with retardation factors ranging from 2.6 to 5.8 and a remaining low mobility fraction with a retardation factor >200. For interbed sediments, retardation factors were >800. Based on reported porosities and bulk densities of packed columns, retardation factors for basalt equate to K_d values of 0.5 to 1.36 mL/g for the high mobility fraction and >57 mL/g for the low mobility fraction, and a K_d value of >194 mL/g for interbed sediment.

Cooper, et al. (2006) estimated an americium K_d range of 100 to 10,000 mL/g for alluvium and interbed sediments at INTEC and recommended an americium K_d value of 400 mL/g for the alluvium and interbed sediments. The minimum and maximum americium K_d values estimated by Cooper, et al. (2006) represented the range of values from the batch experiments of Newman, et al. (1996) and long-term contact data (see Cooper, et al., 2006, Figure D–3–2), which indicated that americium K_d values should range from 100 to 10,000 mL/g in INTEC sediments. The recommended americium K_d value of 400 mL/g for alluvium and interbed sediments was based on column experiments conducted by Artinger, et al. (2002). Artinger, et al. (2002) reported americium K_d values ranging from 1,950 to 2,150 mL/g for SDA interbed sediments. Adjusting for CEC (INTEC/SDA = 1:5) lead to the recommended americium K_d value of 400 mL/g for alluvium and interbed sediments. Cooper, et al. (2006) estimated an americium K_d range of 0 to 140 mL/g for basalt at INTEC and recommended an americium K_d value of 0.85 mL/g for basalt. The maximum estimate of 140 mL/g was based on the highest

value measured in the batch experiments of Newman, et al. (1996) for basalt and corrected for the sediment/water ratio. The recommended americium K_d value of 0.85 mL/g was estimated to be the average K_d reported in the column experiments of Fjeld, et al. (2001).

Americium K_d values have been determined for Columbia River basalts collected from the Hanford site under experimental conditions similar though not identical to the hydrogeologic system at INL (i.e., alkaline pH, high carbonate concentrations, oxidizing conditions, and similar though not identical basalt mineral compositions). Barney (1981) reported americium K_d values for Umtanum ranging from 50–315 mL/g. Salter, et al. (1981) reported americium K_d values for Umtanum ranging from 50–600 mL/g.

Measured and recommended americium K_d values are summarized in Table 3-14. Measured americium K_d values for interbed sediments using batch methods range from 450 to 1,100 mL/g (Newman, et al., 1996). Measured americium K_d values for interbed sediments from column experiments were >49 mL/g and >194 mL/g (Newman, et al., 1996; Fjeld, et al., 2001). In the column experiments of Newman, et al. (1996), a high americium mobility fraction was also detected; however, the high americium mobility fraction was too small (~1 percent) to determine a K_d . Based on the column experiments of Fjeld, et al. (2001), which indicated an americium K_d value >194 mL/g, and considering the minimum americium K_d value reported from the batch experiments of Newman, et al. (1996) (450 mL/g), an americium K_d value of 400 mL/g is recommended for alluvium and interbed sediments. This value is in agreement with the americium K_d value of 400 mL/g recommended by Cooper, et al. (2006) for INTEC alluvium and interbed sediments.

For basalt at INL, measured americium K_d values using batch methods ranged from 70 to 280 mL/g (Newman, et al., 1996). Measured americium K_d values for basalt from the Hanford site determined by batch methods have similar ranges of 50 to 315 mL/g (Barney, 1981) and 50 to 600 mL/g (Salter, et al., 1981). In column experiments using INL basalts, high and low americium mobility fractions were detected (Newman, et al., 1996; Fjeld, et al., 2001). The high mobility americium fraction in the column experiments had a K_d range of 0.18 to 1.36 mL/g, and the low mobility fraction had K_d values of >57 and >60 mL/g. Based on the americium K_d value from the column experiments of Fjeld, et al. (2001) (>60 mL/g) and considering the minimum americium K_d value reported from the batch experiments of Newman, et al. (1996) (70 mL/g), an americium K_d value of 70 mL/g is recommended for basalt. This value is much higher than the americium K_d value of 0.85 ml/g recommended by Cooper, et al. (2006) for INTEC basalt.

However, the recommendation of Cooper, et al. (2006) considered only K_d data measured on the high mobility fraction from the column experiments of Fjeld, et al. (2001). The high americium mobility fraction in basalt ranged from 9 to 13 percent in the column experiments of Newman, et al. (1996) and from 7 to 12 percent in the column experiments of Fjeld, et al. (2001). Based on the range of high mobility americium K_d values measured in the column experiments (0.18 to 1.36 mL/g), a K_d value of 0.2 mL/g is recommended for the high mobility fraction of americium in basalt. This high americium mobility fraction likely results from colloid-facilitated transport, and its effects should be considered in groundwater dose models.

Table 3-14. Measured and Recommended Americium K_d Values for Alluvium, Interbed Sediments, and Basalt at the Idaho National Laboratory

	Measured Americium K _d (mL/g)	Test Method	Idaho Nuclear Technology and Engineering Center-Specific Recommended Americium K _d (mL/g)*	Recommended Americium K _d (mL/g)
Alluvium	_	l	400	400
Sedimentary Interbeds	450–1,100† >49† >194‡	Batch Column Column	400	400
Basalt	70–280† 0.18† (hmf§) >60† 0.5–1.36‡ (hmf) >57‡	Batch Column Column Column Column	0.85	70 0.2 (hmf)

^{*}Cooper, D.C., G.D. Redden, and L.C. Hull. DOE/NE–ID–11227, "Operable Unit 3-14 Tank Farm Soil and Groundwater Remedial Investigation/Baseline Risk Assessment—Appendix D: Estimation of K_d Values for INTEC Groundwater Model." Idaho Falls, Idaho: DOE–Idaho. 2006.

The position of the americium K_d values recommended in this review with respect to measured and estimated site-specific and analogous americium K_d ranges and values is graphically displayed in Figure 3-11. This figure illustrates the uncertainty associated with the americium K_d values recommended for alluvium, interbed sediments, and basalt at INL. Due to available site-specific americium sorption data (e.g., Newman, et al., 1996; Fjeld, et al., 2001) and analogous americium sorption data from the Hanford site (Barney, 1981; Salter, et al., 1981), the recommended americium K_d values are associated with low uncertainty.

3.4.11 Curium

The actinide curium exists in the +3 oxidation state in natural waters. The dominant aqueous species of curium in the pH range of natural waters are primarily cations such as Cm³⁺ and Cm(III) carbonate complexes at acidic and basic pH values, respectively. Available curium sorption studies indicate that sorption of curium is strongly pH dependent and increases with increasing pH with maximum adsorption occurring between pH values of 5 and 6. Curium geochemistry is very similar to that of americium (Guillaumont, et al., 2003). Both are

[†]Newman, M.E., I. Porro, R. Scott, R.M. Dunnivant, R.W. Goff, M.D. Blevins, S.M. Ince, J.D. Leyba, T.A. DeVol, A.W. Elzerman, and R.A. Fjeld. "Evaluation of the Mobility of Am, Cs, Co, Pu, Sr, and U Through INEL Basalt and Interbed Materials: Summary Report of the INEL/Clemson University Laboratory Studies." ER–WAG7–82, INEL–95/282. Idaho Falls, Idaho: Idaho National Engineering Laboratory. 1996.

[‡]Fjeld, R.A., T.A. DeVol, R.W. Goff, M.D. Blevins, D.D. Brown, S.M. Ince, A.W. Elzerman, and M.E. Newman. "Characterization of the Mobilities of Selected Actinides and Fission/activation Products in Laboratory Columns Containing Subsurface Material from the Snake River Plain." *Nuclear Technology*. Vol. 135. pp. 92–108. 2001. §(hmf) denotes a high-mobility fraction.

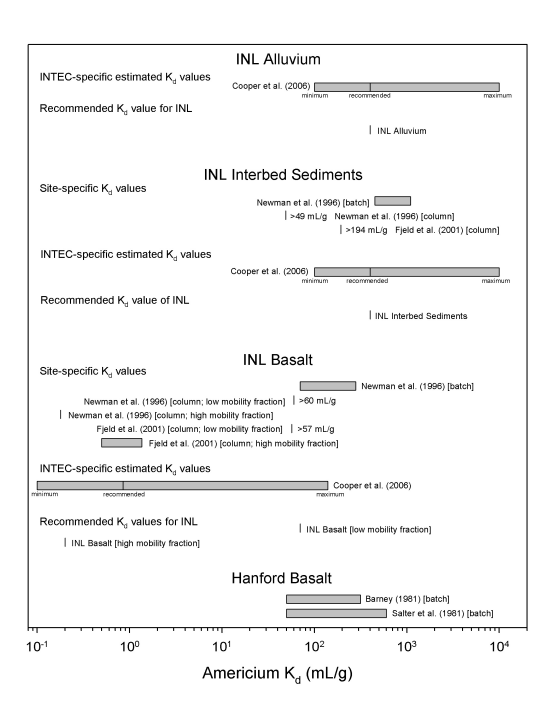


Figure 3-11. Measured and Estimated Site-Specific and Analogous Americium $K_{\rm d}$ Ranges and Values and Recommended Americium $K_{\rm d}$ Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-14). Recommended Americium $K_{\rm d}$ Values Are Associated With Low Uncertainty Due to Available Site-Specific Sorption Data.

transuranic elements that occur in the +3 oxidation state in natural waters. Both curium and americium form cationic carbonate complexes at acidic and alkaline pH. Both are considered to be immobile in soil environments, and both exhibit high $K_{\rm d}$ values. However, the tendency of curium, like americium, to strongly adsorb to soil particles indicates that there is potential for colloid-facilitated transport of curium.

No site-specific sorption data for curium at INL were found in the literature. Using site-specific sorption data for americium from Newman, et al. (1996) as an analog for curium, Rodriguez, et al. (1997) recommended a curium K_d range of 400 to 1,000 mL/g for sediments at INL and a curium K_d range of 70 to 280 mL/g for basalt. Rodriguez, et al. (1997) recommended values of 400 mL/g for sediments and 70 mL/g for basalt.

Recommended curium K_d values are summarized in Table 3-15. No site-specific curium sorption data are available for subsurface materials at INL. Due to similarities in aqueous chemistry, the sorptive characteristics of curium onto alluvium, interbed sediments, and basalt at INL are expected to be very similar to those of americium. Using americium as an analog for curium, a curium K_d value of 400 mL/g is recommended for interbed sediments and alluvium at INL, and a curium K_d value of 70 mL/g is recommended for basalt at INL. In addition, the possibility of a high mobility fraction with characteristics similar to that assigned to americium (Table 3-14) should be considered for curium.

The position of the curium K_d values recommended in this review with respect to estimated site-specific curium K_d ranges and values is graphically displayed in Figure 3-12. This figure illustrates the uncertainty associated with the curium K_d values recommended for alluvium, interbed sediments, and basalt at INL. The recommended curium K_d values are associated with high uncertainty due to the lack of site-specific curium sorption data.

Basalt at the Idaho National Laboratory*					
	Measured Curium K _d (mL/g)	Test Method	Idaho National Laboratory-Specific Recommended Curium K _d (mL/g)	Recommended Curium K _d (mL/g)	
Alluvium	_	_	400	400	
Sedimentary Interbeds	_	_	400	400	

*Rodriguez, R.R., A.L. Schafer, J. McCarthy, P. Martian, D.E. Burns, D.E. Raunig, N.A. Burch, and R.L. VanHorn. DOE/ID–10534, "Comprehensive RI/FS for the Idaho Chemical Processing Plant OU 3-13 at the INEEL—Part A, RI/BRA Report (Final)." Idaho Falls, Idaho: Idaho National Engineering Laboratory, DOE–Idaho. 1997.

Basalt

70

70

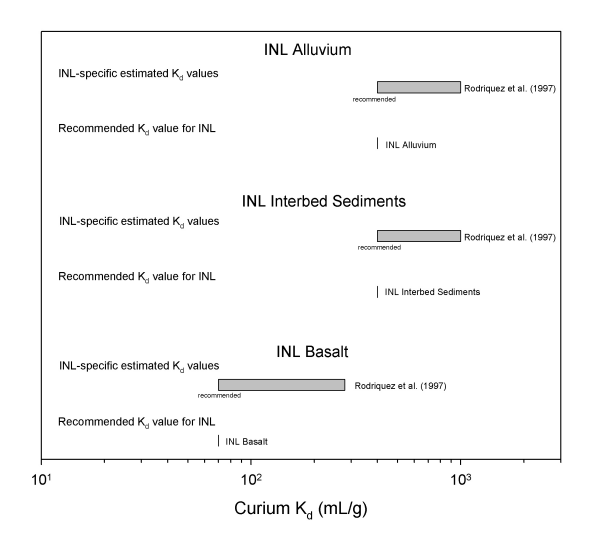


Figure 3-12. Estimated Site-Specific Curium K_d Ranges and Values and Recommended Curium K_d Values for Alluvium, Interbed Sediments, and Basalt at INL (Table 3-15). Recommended Curium K_d Values Are Associated With High Uncertainty Due to the Lack of Site-Specific Sorption Data.

4 SUMMARY AND CONCLUSIONS

To satisfy non-high-level radioactive waste determinations for facilities at the Savannah River Site (SRS) and the Idaho National Laboratory (INL) under Section 3116 of the Ronald W. Reagan National Defense Authorization Act, the U.S. Department of Energy (DOE) commonly performs a performance assessment that includes evaluating potential exposure to radionuclides via groundwater transport. Deterministic performance assessment models DOE currently uses to estimate the rate of migration of contaminants at SRS and INL require K_d s to describe solute retardation along subsurface flow paths. In its consultative role in the Section 3116 waste determination process, the U.S. Nuclear Regulatory Commission (NRC) reviews relevant performance assessments and, therefore, is interested in whether the K_d s DOE uses for radionuclide transport in the geosphere reasonably reflect expected local conditions.

This report documents a sorption coefficient database tailored to site-specific subsurface conditions for risk-significant radionuclides at SRS and INL and recommends K_d values for the risk-significant radionuclides that are deemed appropriate for the hydrogeochemical environments found at SRS and INL. A summary of the K_d recommendations for the risk-significant radionuclides in the hydrostratigraphic units that will be affected by waste releases at SRS and INL is provided in Tables 4-1 and 4-2. The recommended K_d values are based on knowledge of the sorption behavior of each radionuclide of concern, available data on site hydrogeochemical and mineralogical conditions, and reported K_d values from site-specific experiments. When site-specific K_d values were not available for a particular radionuclide, K_d data on analogous systems to SRS or INL and generic data from the open literature were evaluated to provide a technical basis for the K_d recommendation. The recommended values tended toward the lower, more conservative end of data ranges. This is because of the uncertainties not only in the appropriate choice of sorption coefficients, but also in the simplified K_a-based groundwater transport model itself. For example, the fractured basalt strata at INL may contain fast pathways that will make surface sorption processes less effective. The selection of low values is intended to ensure that groundwater pathway models will be unlikely to underestimate potential dose.

These recommendations should be used with care in light of new information that becomes available on site-specific sorption coefficients or on temporal or spatial variability in hydrochemical conditions. This report is not to be construed as providing the technical basis for parameters DOE uses in its performance assessments.

For plutonium, americium, and curium at INL, a high mobility fraction with much lower K_d values than those in Table 4-2 should be considered in transport models (see Tables 3-13 and 3-14).

The results of this data review can be used in a risk-informed manner to identify potential areas of focused laboratory studies. For each site, starting with the list of radionuclides included in this analysis, staff could narrow the list based on results of performance assessments or other modeling studies. Among this smaller list, those with an insufficient quantity of high-quality, site-specific sorption data (considering both laboratory and field-based studies) may be considered for laboratory analyses that would complement any past and ongoing DOE efforts.

Table 4-1. Summary of Recommended ${\rm K_d}$ Values (mL/g) for Risk-Significant Radionuclides in Hydrostratigraphic Units at the Savannah River Site

Radionuclide	Upper Three Runs Aquifer: Upper	Tan Clay Confining Unit	Upper Three Runs Aquifer: Lower	Gordon Confining Unit	Gordon Aquifer
Carbon	2	1	1	1	1
Nickel	10	21	80	150	80
Selenium	425	800	425	800	425
Strontium	5	12	25	50	25
Niobium	160	900	160	900	160
Technetium	0	0.3	0	0.3	0
Tin	500	2,500	500	2,500	500
lodine	0	0.2	0	0.2	0
Cesium	15	150	50	150	50
Lead	500	2,500	1,500	2,500	1,500
Uranium	100	150	100	300	100
Neptunium	0.2	25	0.6	25	0.6
Plutonium	250	2,500	250	2,500	250
Americium	250	4,250	250	4,250	250
Curium	300	4,250	300	4,250	300

Table 4-2. Summary of Recommended $K_{\rm d}$ Values (mL/g) for Risk-Significant Radionuclides in Hydrostratigraphic Units at the Idaho National Laboratory

Radionuclide	Alluvium	Sedimentary Interbeds	Basalt
Carbon	1	1	0
Strontium	12	12	2
Technetium	0	0	0
lodine	0.1	0.1	0
Cesium	75	200	25
Lead	270	270	10
Uranium	1.6	1.6	0.2
Neptunium	2	5	0.2
Plutonium	1,000	500	70
Americium	400	400	70
Curium	400	400	70

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