Solubility and Speciation Results from Over- and Undersaturation Experiments on Neptunium, Plutonium, and Americium in Water from Yucca Mountain Region Well UE-25p #1



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by

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ABSTRACT

Solubility and speciation are important in understanding aqueous radionuclide transport through the geosphere. They define the source term for transport retardation processes such as sorption and colloid formation. Solubility and speciation data are useful in verifying the validity of geochemical codes that are a part of predictive transport models. Ideally, solubility experiments should approach solution equilibrium from both oversaturation and undersaturation. Results are compared from solubility and speciation experiments from oversaturation and undersaturation of ²³⁷NpO₂+, ²³⁹Pu⁴⁺, and ²⁴¹Am³⁺/Nd³⁺ in a modified UE-25p #1 groundwater (from the Yucca Mountain region, Nevada, which is being investigated as a potential high-level nuclear waste disposal site) at 60°C and three pH values (6.0, 7.0, 8.5). In the oversaturation experiments, the solubilitycontrolling steady-state solids were identified and the speciation and/or oxidation states present in the supernatant solutions were determined. The characterized solids were then reintroduced into fresh solutions of the modified UE-25p #1 groundwater to approach the steady state from undersaturation. For the undersaturation experiments, the solubility-controlling steady-state solids were also identified and the speciation and/or oxidation states present in the supernatant solutions were determined. The neptunium solubility decreased with increasing pH in both the over- and undersaturation experiments. The steady-state concentrations from the two experiments agreed to within an order of magnitude. Plutonium concentrations from over- and undersaturation agree very well in the pH 6 and 7 experiments. The pH 8.5 oversaturation experiment resulted in a steadystate concentration one order of magnitude above its undersaturation counterpart. It appears that a new, more crystalline solid formed in the pH 8.5 undersaturation cell. All five of the other cells resulted in steady-state concentrations equivalent to each other within the error of the measurement. For the americium/neodymium solutions, the pH 6 and pH 7 experiments resulted in equivalent steady-state concentrations from both over and undersaturation. The pH 8.5 oversaturation experiment was 100 times more soluble than its undersaturation counterpart. This is probably due to the oversaturation cell not being at steady-state at the termination of the experiment. Otherwise, the solubility decreased with increasing pH.

PREFACE

This report is the third in a series documenting experimental solubility and speciation studies of radionuclides in groundwaters from the Yucca Mountain region. The objectives and experimental concepts were discussed in detail in the first report of this series (Milestone 3010), entitled "Measured Solubilities and Speciations of Neptunium, Plutonium, and Americium in a Typical Groundwater from the Yucca Mountain Region."¹ Sections 2, 3, and 4 of this report are, except for minor changes, identical to the respective sections of the first report. They are, however, included here to make this report a stand-alone document.

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1. EXECUTIVE SUMMARY

We studied the solubilities of neptunium, plutonium, and americium in a modified UE-25p #1 groundwater from the Yucca Mountain region (Nevada) from over- and undersaturation at 60°C and pH's 6, 7, and 8.5. Tables I, II, and III summarize the results for neptunium, plutonium, and americium, respectively. The nuclides were added to UE-25p #1 groundwater from oversaturation at the beginning of each experiment as ²³⁷NpO₂⁺, ²³⁹Pu⁴⁺, and Nd³⁺ with tracer ²⁴¹Am³⁺ added to facilitate nuclear counting. Because we maintained constant pH values of 6, 7, and 8.5 during the course of the experiments, the final solutions were closer to 0.1 M in total ionic strength with the primary constituents being sodium and perchlorate. The solids formed in the oversaturation experiment were then reintroduced into fresh UE-25p #1 groundwater to conduct the undersaturation experiments. At the conclusion of the undersaturation experiments the resulting solids were removed and characterized.

The neptunium solubility decreased with increasing pH in both the over- and undersaturation experiments. The steady-state concentrations from over- and undersaturation agreed within one order of magnitude. The soluble neptunium did not change oxidation state at steady-state. The pentavalent neptunium was increasingly complexed by carbonate with increasing pH. The steady-state solids were crystalline sodium neptunium carbonate hydrates with varying stoichiometry.

Plutonium concentrations showed no trend with pH. The oversaturation and undersaturation steady-state concentrations agreed within the error of the measurement for pH 6 and pH 7. The pH 8.5 oversaturation experiment resulted in a steady-state concentration one order of magnitude above the undersaturation experiment. This was due to formation of a more crystalline solid in the pH 8.5 undersaturation cell. The dominant oxidation state was Pu(VI) in all experiments, with a large contribution from Pu(V) in all of the undersaturation cells. The species responsible for this oxidation are unknown. The solubility-controlling solids from oversaturation at pH 6 and 7 were amorphous and contained Pu(IV) polymer. The solid produced in the pH 8.5 oversaturation cell is crystalline but unidentified. All of the undersaturation solids appear to be similar and yielded similar steady-state concentrations. They are crystalline but unidentified.

For the americium/neodymium solutions, the solubility decreased with increasing pH in all of the undersaturation experiments. It appears that the pH 8.5 oversaturation experiment did not reach steady-state until day 125 of the undersaturation experiment. The oxidation state in the supernatant of the americium/neodymium solutions remained in the trivalent state. The americium/neodymium solubility experiments in UE-25p #1 groundwater produced mixtures of orthorhombic NdOHCO₃ and Nd₂(CO₃)₃ • 2 H₂O.

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	Steady-State Concentration (M)		Oxidation State in Supernatant Solution (%)	
	Oversaturation	Undersaturation	Oversaturation	Undersaturation
рН б	$(2.5 \pm 0.2) \ge 10^{-3}$	$(3.4 \pm 0.1) \ge 10^{-3}$	V : 100 % uncomplexed	V : 100 % 24 % carbonate complexed
pH 7	$(3.4 \pm 1.0) \ge 10^{-5}$	$(1.0 \pm 0.1) \ge 10^{-4}$	V : 100 % 10 % carbonate complexed	V : 100 % 34 % carbonate complexed
pH 8.5	$(2.7 \pm 0.1) \ge 10^{-5}$	$(3.8 \pm 1.6) \ge 10^{-5}$	V : 100 % 100 % carbonate complexed	V : 100 % 100 % carbonate complexed

Table I.Summary of results for solubility experiments of neptunium in UE-25p #1
groundwater from over- and undersaturation at pH 6, 7, and 8.5 and 60°C.

	Eh (mV) vs. NHE		Solid Phase	
	Oversaturation	Undersaturation	Oversaturation	Undersaturation
рН б	(414 ± 15)	(406 ± 15)	Sodium Neptunyl(V) Carbonate Hydrates	Sodium Neptunyl(V) Carbonate Hydrates
рН 7	(367 ± 15)	(366 ± 15)	Sodium Neptunyl(V) Carbonate Hydrates	Sodium Neptunyl(V) Carbonate Hydrates
рН 8.5	(212 ± 15)	(239 ± 15)	Sodium Neptunyl(V) Carbonate Hydrates	Sodium Neptunyl(V) Carbonate Hydrates

	Steady-State Concentration (M)		Oxidation State in Supernatant Solution (%)			
	Oversaturation	Undersaturation	Oversa	turation	Undersa	aturation
pH 6	$(8.8 \pm 1.4) \ge 10^{-8}$	$(9.5 \pm 3.6) \ge 10^{-8}$	III + Poly. : IV : V : VI :	(1 ± 1) (1 ± 1) (4 ± 1) (94 ± 11)	III + Poly. : IV : V : VI :	(1 ± 1) (2 ± 1) (39 ± 4) (57 ± 6)
рН 7	$(9.1 \pm 1.2) \ge 10^{-8}$	$(6.8 \pm 3.1) \ge 10^{-8}$	III + Poly. : IV : V : VI :	(2 ± 1) (1 ± 1) (5 ± 1) (93 ± 11)	III + Poly. : IV : V : VI :	(1 ± 1) (1 ± 1) (9 ± 2) (89 ± 10)
рН 8.5	$(1.3 \pm 0.8) \ge 10^{-6}$	$(9.2 \pm 4.0) \ge 10^{-8}$	III + Poly. : IV : V : VI :	(5 ± 2) (10 ± 1) (0 ± 0) (86 ± 12)	III + Poly. : IV : V : VI :	(0 ± 0) (10 ± 2) (36 ± 4) (54 ± 7)

Table II.	Summary of results for solubility experiments of plutonium in UE-25p #1
	groundwater from over- and undersaturation at pH 6, 7, and 8.5 and 60°C.

	Eh (mV) vs. NHE		Solid Phase	
	Oversaturation	Undersaturation	Oversaturation	Undersaturation
рН б	(326 ± 15)	(312 ± 15)	Amorphous Pu(IV) polymer	Crystalline Unidentified
pH 7	(334 ± 15)	(313 ± 15)	Amorphous Pu(IV) polymer	Crystalline Unidentified
рН 8.5	(231 ± 15)	(218 ± 15)	Crystalline Unidentified	Crystalline Unidentified

Table III.Summary of results for solubility experiments of americium/neodymium in UE-
25p #1 groundwater from over- and undersaturation at pH 6, 7, and 8.5 and
60°C.

	Steady-State Concentration (M)		Oxidation State in Supernatant Solution (%)	
	Oversaturation	Undersaturation	Oversaturation	Undersaturation
рН б	$(2.7 \pm 0.4) \ge 10^{-9}$	$(7.0 \pm 7.3) \ge 10^{-10}$	III : 100 %	III : 100 %
pH 7	$(7.1 \pm 0.5) \ge 10^{-10}$	$(4.6 \pm 2.0) \ge 10^{-10}$	Not Available.	III : 100 %
рН 8.5	$(7.8 \pm 4.3) \ge 10^{-9}$	$(8.4 \pm 1.8) \ge 10^{-11}$	III : 100 %	III : 100 %

	Eh (mV) vs. NHE		Solid Phase	
	Oversaturation	Undersaturation	Oversaturation	Undersaturation
рН б	(370 ± 15)	(394 ± 15)	$(Am)_2(CO_3)_3 \bullet 2 H_2O$ and orthorhombic AmOHCO ₃	$(Am)_2(CO_3)_3 \cdot 2 H_2O$ and orthorhombic AmOHCO ₃
рН 7	Not Available.	(430 ± 15)	$(Am)_2(CO_3)_3 \cdot 2 H_2O$ and orthorhombic AmOHCO ₃	$(Am)_2(CO_3)_3 \cdot 2 H_2O$ and orthorhombic AmOHCO ₃
рН 8.5	(220 ± 15)	(215 ± 15)	$(Am)_2(CO_3)_3 \cdot 2 H_2O$ and orthorhombic AmOHCO ₃	$(Am)_2(CO_3)_3 \cdot 2 H_2O$ and orthorhombic AmOHCO ₃

2. PURPOSE AND OBJECTIVE

Yucca Mountain, Nevada, was identified for site characterization as the location for a potential repository of high-level nuclear waste. As a worst case scenario, intrusion of water into the potential repository must be considered for risk assessment. Water moving through the emplacement area towards the accessible environment can transport radionuclides in two ways: either as dissolved species in the water or as particulate material by the water. The Yucca Mountain Site Characterization Plan (SCP) requires "Studies to Provide the Information Required on Radionuclide Retardation by Precipitation Processes along Flow Paths to the Accessible Environment" before licensing and construction of the potential repository.² The purpose of this study is to supply data for calculating radionuclide transport along potential transport pathways from the potential repository to the accessible environment. Data derived from solubility studies are important for validating geochemical codes that are part of predictive radionuclide transport models. Such codes should be capable of predicting the results of solubility experiments, where the measured solubility is the sum of the equivalent concentrations of all of the species in equilibrium with a specified solid. Furthermore, agreement between geochemical calculations and experimental results can validate the thermodynamic data base used with the modeling calculation.

To predict behavior at higher temperatures, data bases used for modeling calculations must contain data on thermodynamic functions at elevated temperatures. To date, many of these data are unavailable and are therefore estimated by extrapolation from lower temperature data. Agreement between modeling calculations and experimental results would also validate such estimates, whereas significant discrepancies would identify the need for data base improvement. Improvements can be made by filling the gaps with basic experimental data.

In addition, experimental solubility data also provide the source terms or the starting concentrations for experimental sorption studies. To be valid, sorption studies should be conducted at or below the solubility limit because only soluble species can be transported and participate in the sorption process.

In selecting these experiments, we have considered the generic U.S. Nuclear Regulatory Commission (NRC) technical position entitled "Determination of Radionuclide Solubility in Groundwater for Assessment of High-Level Waste Isolation."³ This technical position served as guidance for our experiments to determine radionuclide solubility. It requires that if radionuclide solubility is used as a factor in limiting radionuclide release, experiments must be designed to determine solubility under site-specific conditions. Radionuclide concentrations in water passing through the emplacement area can be limited by two mechanisms: low dissolution rates of the solid waste form or solubilities of individual radionuclides. If solid waste dissolution rates are low enough, it may not be necessary to depend on solubilities to limit radionuclide concentrations. However, the solid waste forms have not yet been determined, and therefore the dissolution rates of the solid waste are unknown. Determination of radionuclide solubility limits provides an upper bound on radionuclide concentrations in solution and provides a basis for "extrapolation to long-term behavior." The rate of groundwater flow through the waste is expected to be sufficiently slow to permit saturation of water with radionuclides. Dissolution limited by saturation will provide maximum concentration limits. Therefore, an assessment of radionuclide release rates using a saturation-limited dissolution model represents the most conservative approach possible.

As radionuclides are transported along flow paths to the accessible environment, changing conditions of the water (pH, Eh, and concentrations of complexing species) can alter solubilities. Decreases in solubility can decrease radionuclide concentrations. A knowledge of radionuclide solubilities under the conditions along possible flow paths is necessary to assess this scenario. Solubility studies are very time-consuming because long times are often needed to reach equilibrium. Because we cannot investigate every possible solubility scenario, we selected pH and temperature values to bracket the expected range of conditions by choosing parameters that represent lower and upper limits.

Neptunium, plutonium, and americium are expected to be sparingly soluble with solubilitylimited dissolution. Water samples with compositions that bracket the range of waters expected in the vicinity of Yucca Mountain were chosen for solubility measurements.⁴ These samples come from two sources. Water from Well J-13 is a reference water for the unsaturated zone near the proposed emplacement area. Well UE-25p #1 taps the carbonate aquifer that underlies the emplacement horizon. This water has an ionic strength and total carbonate content higher by approximately an order of magnitude than Well J-13 water. UE-25p #1 water represents natural water with the highest concentrations of dissolved species expected in the vicinity of Yucca Mountain. The water from both wells is oxidizing. Generally, radionuclide solubility studies under oxidizing conditions lead to higher solubilities for a number of radionuclides than would occur under mildly or strongly reducing conditions. These experiments will therefore provide conservative results. In this study we are reporting on the results in UE-25p #1 water from overand undersaturation. The maximum temperature of the host rock in which liquid water is present is expected to be limited by the boiling point of water at Yucca Mountain (95°C). The solubility experiments that use Well J-13 water were conducted at temperatures between 25° and 90°C. This span covers the range from pre-emplacement temperatures to the maximum temperature at which solubility would be important. For Well UE-25p #1 water, solubility experiments were limited to a maximum temperature of 60°C. Maximum temperatures in the saturated zone under the emplacement area and those along the flow paths away from the emplacement area are expected to be less than $60^{\circ}C.^{5}$

3. CONCEPT OF SOLUBILITY STUDIES

Solubility establishes an upper limit for the dissolved components in the source term for radionuclide migration from a repository. Studies of the solubility of radionuclides in groundwaters from a repository horizon will provide limits on their potential concentrations in those waters. Such limits are important for (1) validating an essential part of the radionuclide transport calculations and (2) providing guidance in choosing the maximum starting concentrations for radionuclide sorption experiments. Compared with multi-parameter transport models, laboratory solubility experiments are controlled by fewer variables. If geochemical codes such as EQ3/6 are to be included in the transport model, the model should be capable of predicting the results of solubility experiments.

Complete solubility experiments should provide detailed knowledge of (1) the nature and chemical composition of the solubility-controlling solid, (2) the concentration of the components in solution, and (3) the identity and electrical charge of the species in the solution phase.

Meaningful thermodynamically defined solubility studies must satisfy four criteria: (1) attainment of equilibrium conditions; (2) determination of accurate solution concentrations; (3) attainment and identification of a well-defined solid phase; and (4) knowledge of the speciation/oxidation state of the soluble species at equilibrium.

3.1. Oversaturation and Undersaturation

Ideally, solubility experiments should approach solution equilibrium from both oversaturation and undersaturation. The approach from oversaturation consists of adding an excess amount of the element in soluble form to the aqueous solution and then monitoring the precipitation of insoluble material until equilibrium is reached. The solid formed must then be isolated and characterized. The approach from undersaturation consists of dissolving the same well-defined solid in an aqueous solution until equilibrium is reached. In both cases, the solution concentration is measured as a function of time until equilibrium is reached.

Kinetic processes will control the equilibration speed in solubility experiments. Some solutions equilibrate rapidly, others more slowly. It must be demonstrated that equilibrium is reached. This can be accomplished by experimentally determining (for both oversaturation and undersaturation experiments) the solution concentration as a function of time. When the concentration stays constant for several weeks, it is assumed that equilibrium has been established. Because this assumption is based on judgment, the term "steady-state" instead of "equilibrium" is more precise. The U.S. Nuclear Regulatory Commission (U.S. NRC) defines "steady-state," as "the conditions where measurable changes in concentrations are not occurring over practical experimental times."³ At steady state, thermodynamic forces may still change the solution composition: solids may become less soluble as they change from a higher to a lower free energy. The change may be controlled by kinetics and may therefore be very slow and may not show in the experiment even after several years. These infinitesimal changes may require infinite experimental times. The steady-state solids formed in the experiments may therefore not represent the thermodynamically most stable solids with the lowest possible solubilities, but metastable solids having higher solubilities than the thermodynamically defined solids. The term "steady-state" implies this condition.

Despite this constraint, time-limited laboratory solubility experiments can supply valuable information. They provide good estimates on the upper limit of radionuclide concentrations in solution because the experimentally determined steady-state concentrations are higher than the equilibrium concentrations.

A reliable method of proving that an equilibrium has been reached is to approach steady-state from both oversaturation and undersaturation. When these two experimental approaches independently produce equal solution concentrations, the data are considered reliable. For unknown solubility systems, one should first perform experiments approaching steady-state concentration from oversaturation and then characterize the solids. This has the advantage of not specifying the solid that controls solubility but of allowing the system under investigation to determine the solid that will precipitate. These solids can then be used in confirmation experiments that approach steady-state from undersaturation. In this study we are reporting results for both the oversaturation and undersaturation experiments.

3.2. Phase Separation

The second criterion for meaningful solubility experiments is the derivation of accurate solution concentrations. This requires that phase separations must be as complete as possible. The separation of the solid from the solution often represents a significant practical problem in measuring solubility. Apparently higher or lower solubilities, compared with the steady-state values, can result from incomplete phase separation or from sorption of solute during and after the separation. Incomplete phase separations (leaving some of the solid with the solution phase) lead to higher radionuclide solubilities. Lower solubilities are measured if constituents of the steady-state solution have been sorbed on filters during a filtration and on container walls after the separation.

Experimentally, the solids and solutions are separated on the basis of differences in size (via filtration) or density (via sedimentation or centrifugation). Filtration is the more commonly applied technique because it physically partitions the solute and solids. Ultrafiltration (i.e., filtration using membranes $< 0.1 \,\mu\text{m}$) can effectively remove solids and colloidal particles from aqueous solution. A potential problem with ultrafiltration is adsorption of soluble species on filtration membranes. Effective filters for solubility studies must pass soluble species quantitatively; that is, either the filter should have no active sorption sites at all or any such sites should be irreversibly blocked. Filters are adequate if they have a small enough pore size to retain the solids and colloids and if they also show no sorption or only minimal sorption during multiple filtrations. Because adsorption of soluble radionuclide species on filters can be dependent on the solution's pH and on the solution species, it is mandatory to verify that possible sorption sites are indeed blocked. Usually the sorptive sites on a filter and filter housing are blocked by preconditioning of these materials. The filter is preconditioned by filtering a volume of the respective radionuclide solution through it and then discarding the filtrate. The volume required for preconditioning is determined experimentally. Details for this procedure are given in Section 4.5 Phase Separation.

3.3. Solid Phase

Solubility depends strongly on the state of the solid phase. Thermodynamically meaningful results require the existence of a well-defined solid phase, which ideally consists of crystalline material. The solids formed from the oversaturation in solubility tests must be clearly identified by physical or chemical characterization methods. Radionuclide solids formed in laboratory experiments and in nature are often thermodynamically ill-defined amorphous precipitates. Most amorphous solids, however, will become more crystalline with time. Freshly precipitated

microcrystalline solids can also convert in time to a macrocrystalline material. Improved bonding at the lattice surface results in decreasing surface area. Thus the crystalline solid of higher free energy changes to one of lower free energy (Ostwald ripening, Ostwald step rule) and becomes less soluble.^{6,7,8,9}

3.4. Determination of Oxidation States and Speciation

Information on oxidation states and speciation of the radionuclides in steady-state solubility solution is important for transport models simulating migration and sorption along the flow path to the accessible environment. The charge and speciation of radionuclides will control their sorption and transportation in the geologic host. Speciation measurements identify complexes that may form between radionuclides and complexing ions present in the groundwater near the potential repository. Radionuclides, like all nuclides, can have a single or several different oxidation states in solution. They can be present as simple ions or as complexes. When the ions react with one or several other solution components, they can form soluble complexes.

Oxidation states and speciation in solution are commonly determined by (1) absorption spectrophotometry, (2) ion exchange chromatography, (3) solvent extraction, (4) coprecipitation, (5) potentiometry, and (6) electrochemistry. Of these methods, only absorption spectrophotometry can provide information on speciation, while the others identify only the oxidation state in solution.

Absorption spectrophotometry of Neptunium, Plutonium and Americium in UE-25p #1 water has a detection limit of about 10⁻⁵ M. This relatively high concentration limits the application of spectrophotometry for speciation determination in solutions from radionuclide solubility studies because the solubilities can be several orders of magnitude below 10⁻⁶ M. Laser-Induced Photoacoustic Spectroscopy (LIPAS) provides much greater sensitivity, approaching 10⁻⁸ to 10⁻⁹ M. 10,11,12,13,14,15

The methods listed above as 2 through 6 determine only the oxidation state in solution because they cannot determine species. They detect the oxidation state of ions indirectly. This process is different from absorption spectrophotometry, which detects oxidation states and sometimes the solution species directly. The indirect methods, however, detect very small concentrations (10⁻¹⁰ M and below), which is useful for radionuclide solubility studies. Solvent extraction and coprecipitation are often used successfully to determine the oxidation states of ions in very dilute solutions.¹⁶

Ion exchange chromatography is less reliable for this purpose because the exchange resin often reduces the solution ions, which gives incorrect results for the oxidation state distribution. Electrochemical detection reduces or oxidizes the solution ions and measures the potentials of the reduction and oxidation reactions, respectively. The potential then identifies the individual ion. Electrochemistry needs fast kinetics and reversible thermodynamics for the reduction or oxidation step. These experiments greatly limit the method because many radionuclide ion redox reactions are irreversible and slow (e.g., the reactions of NpO_2^+/Np^{4+} , $PuO_2^+/Pu^{4+})^{17}$.

The neptunium solution species were determined by spectrophotometry because the solution concentration was greater than 10^{-5} M. The oxidation state of plutonium and americium species in solution were determined by a solvent extraction technique, which is described in principle by Nitsche, et al.¹⁶ and in detail in Section 5.2.2.

4. EXPERIMENTAL DETAILS

We studied the solubilities of neptunium, plutonium, and americium from over- and undersaturation at 60°C and pH values of 6, 7, and 8.5. Measurements were made in an inertatmosphere glovebox to avoid contamination of solutions by atmospheric CO₂. The solubilities were studied from oversaturation by injecting a small amount of actinide stock solution into 80 mL of groundwater obtained from Well UE-25p #1. The analysis of the water composition is listed in Table IV.⁴

Species	Concentration, mM
Na ⁺	7.43
K ⁺	0.34
Ca ²⁺	2.19
Mg ²⁺	1.31
SiO ₂	0.62
Cl-	1.04
SO_2^{2-}	1.34
F-	0.18
Total Alkalinity	11.44 mequiv./L
Total Carbonate	15.31
рН	6.7
Eh (mV)	360
Ionic Strength (mM)	~20

Table IV.UE-25p #1 Groundwater Composition4

The UE-25p #1 groundwater was obtained at the site by Los Alamos personnel. It was filtered at Los Alamos before it was shipped to LBL. The water's natural carbon dioxide partial pressure (pCO_2) could not be preserved during filtration and shipping. For the experiments, however, the natural state was induced by re-equilibrating the water with CO₂ gas. Details of this procedure are described in Sections 4.3. Pressure Control System, and 4.4. Solutions. Details of the filtration are described in Section 4.4. Solutions. The polyethylene shipping bottle was leached with acid and distilled water prior to its use for the groundwater. The leaching removes possible trace-level contaminants that may alter the composition of the UE-25p #1 water.

The solubilities were studied from undersaturation by placing the dry, washed solid obtained from the oversaturation experiment into a cell and introducing 80 mL of a fresh sample of groundwater obtained from Well UE-25p #1. The carbon dioxide partial pressure (pCO_2) was maintained in a similar manner to the oversaturation experiments.

4.1. Controlled-Atmosphere Glove Box

Due to the α -radiation hazard of the actinide elements under investigation, all experimental work was performed in glove boxes. External CO₂ control of the experimental solutions requires the exclusion of atmospheric CO₂. To satisfy both conditions, we used a controlled-atmosphere glove box.

4.2. Control System for pH and Temperature

Because the solubilities are highly sensitive to pH and temperature changes, close control of these parameters is necessary. For previous experiments, we had designed a computer-operated control system ("pH-stat") to maintain the aqueous actinide solutions at constant temperatures and pH values for the solubility experiments.¹⁸ The pH-stat records and adjusts the pH values of the experimental solutions (UE-25p #1 water) at the target values. However, due to frequent deviation from the target pH, the pH-stat began adding excess amounts of acid and base into our solutions. We therefore stopped the automatic addition of acid and base and used the pH-stat to monitor the pH only. Daily adjustments to the pH were performed manually, when necessary, and pH monitoring and data acquisition were continued using the pH-stat system. The temperature of 60°C was controlled within less than 1°C.

4.3. Pressure Control System

We designed and manufactured a pressure regulation system to maintain the well waters used in experiments at their nominal carbonate concentrations when their pH values are adjusted to conditions differing from their natural state. The system also ensured that no significant evaporative loss of the solutions occurred at elevated temperatures.

4.4. Solutions

The actinide stock solutions were prepared by using established methods.^{19 237}Np(V) stock solutions were prepared by dissolving its oxide in HCl. ²³⁹Pu(IV) stock was prepared from plutonium metal. Stable neodymium(III) was used as an analog for americium(III).¹ It was prepared by dissolving Nd₂O₃ in HClO₄. The solution was then spiked with purified ²⁴¹Am(III) tracer to enable the use of nuclear counting for the determination of the neodymium solution concentrations. Further details for these ²⁴¹Am/Nd mixtures are given in Section 5.3. The actinide solutions were purified from possible metal contaminants by ion exchange chromatography. For neptunium and plutonium, anion exchange was used, while cation exchange was employed for americium²⁰. The solutions were converted to a non-complexing perchlorate system. The neptunium and plutonium stock solutions were in the +6 oxidation state after their conversion to perchloric acid (2 to 3 M) and were reduced electrolytically to NpO_2^+ and Pu^{3+} , respectively. Pu^{4+} was prepared by electrolytic oxidation of pure Pu^{3+} immediately before the start of the plutonium solubility experiments in order to minimize the disproportionation of Pu⁴⁺.^{19,21,22} Valence purity of the stock solutions was established by absorption spectrophotometry.^{23,24} Oxidation states other than Pu(IV) were not detected. With our absorption spectrophotometer (Guided Wave Model 260, El Dorado Hills, CA), the limits of detection for Pu(III), Pu(V), and Pu(VI) are approximately 10⁻⁴, 3 x 10⁻⁴, and 10⁻⁵ M, respectively.

The groundwater was filtered through 0.05 μ m polycarbonate membrane filters (Nuclepore Corp., Pleasanton, CA). This filtration was carried out by Los Alamos personnel prior to shipping the UE-25p #1 water sample to LBL. The actinide stock solutions, and all other solutions utilized in this experiment were filtered through 0.22 μ m polyvinylidene difluoride syringe filter units (Millipore Corp., Bedford, MA). Filtration was used to remove suspended particulate material, e.g., dust or silica, that could absorb the actinide ions to form pseudocolloids. For the oversaturation experiments, before adding between 1 and 4 mL of the actinide stock solutions to approximately 80 mL of UE-25p #1 water, a small amount of CO₂-free sodium hydroxide solution was added in order to keep the pH values at or above the desired solution pH. Letting the

pH drop below the target value would necessitate addition of concentrated base to the system while the actinide ion is already present in the solution. Addition of strong base can result in unpredictable microprecipitation and formation of microcolloids.

When we started the neptunium oversaturation experiments, we added small volumes of 5 M sodium hydroxide to the UE-25p #1 groundwater to neutralize the perchloric acid that would be introduced with the stock solution. We added the acidic stock followed by 1 M perchloric acid, 1 M sodium hydroxide, or both if necessary, to attain the desired pH for the experiment. At pH 6, the added acid and base led to additional concentrations of Na⁺ and ClO₄⁻ of 53 and 8.9 mM, respectively, resulting in a higher initial concentration of Na⁺ by a factor of about 8.2 and an initial ionic strength about 4.9 times higher than that of UE-25p #1 water. At pH 7, addition of only sodium hydroxide increased the Na⁺ concentration by 9.6 mM. This increased the ionic strength by only 25 percent. At pH 8.5, additions of NaOH and HClO₄ increased the respective Na⁺ and ClO₄⁻ concentrations by 23 and 7.5 mM. This increased the Na⁺ content and the ionic strength by factors of about 4.1 and 1.8, respectively.

For the plutonium experiments, we also added 5 M NaOH to the UE-25p #1 water before we introduced the acidic plutonium stock solution. The additions of acid and base were small and the resulting changes in the water composition were not large. The plutonium stock solution was 7.9 x 10^{-3} M in total plutonium and 3.0 M in HClO₄. At pH 6 and 7, adjustments led to additional concentrations of Na⁺ and ClO₄⁻ of 21 and 80 mM, respectively. These additions increased the sodium content and the ionic strength in both experiments by factors of about 3.8 and 3.6, respectively. For the pH 8.5 experiment, acid and base adjustments led to additional concentrations of Na⁺ and ClO₄⁻ of 22 and 95 mM, respectively. This increased the Na⁺ content and the ionic strength by factors of about 4.0 and 3.9, respectively.

For the neodymium/americium experiments, we used even smaller amounts of 5 M NaOH because the Nd/Am stock solution was 1 M in HClO₄. At pH 6, addition of acid and base, including the stock solution that was again 1 M in HClO₄, increased the Na⁺ and ClO₄⁻ concentrations by 14 and 20 mM, respectively. This increased the initial amount of sodium and the ionic strength by factors of about 2.9 and 1.9, respectively. At pH 7, additions increased the Na⁺ and ClO₄⁻ concentrations by 15 and 17 mM, respectively, increasing the sodium content and the ionic strength by factors of about 3.0 and 1.8, respectively. At pH 8.5, additions increased the Na⁺ and ClO₄⁻ concentrations by 19 and 17 mM, respectively. This increased the initial amount of sodium and the ionic strength by factors of about 3.0 and 1.8, respectively. At pH 8.5, additions increased the Na⁺ and ClO₄⁻ concentrations by 19 and 17 mM, respectively. This increased the initial amount of sodium and the ionic strength by factors of about 3.5 and 1.9, respectively.

In summary, initial additions of sodium hydroxide and perchloric acid to the neptunium experiments increased the total Na⁺ content by factors that ranged from 2.3 to 8.2 times the original amount found in UE-25p #1 water and increased the ionic strength of the solutions by factors ranging from 1.3 to 4.9. Additions of acid and base at the start of the plutonium experiments increased the total Na⁺ content by factors that ranged from 3.8 to 4.0, resulting in ionic strengths that were higher than that of UE-25p #1 well water by factors that ranged from 3.6 to 3.9. And for the neodymium/americium experiments, additions of acid and base at the start of the start of the start of the experiments increased the initial sodium content by factors that ranged from 2.9 to 3.5, resulting in initial ionic strengths higher by factors that ranged from 1.8 to 1.9.

The well water's total dissolved carbonate $(1.531 \times 10^{-2} \text{ M})$ was preserved at each individual pH by equilibrating the solution with mixtures of CO₂ in argon.⁴ The amount of CO₂ at a given pH and temperature was calculated from Henry's constant and the dissociation constants of carbonic acid from literature data.²⁵ If the value at the given temperature was not available, the number was derived by interpolation of adjacent values. Activity coefficients were adjusted for ionic strength using the Davies equation. The concentrations of the equilibration gas mixtures are given in Table V together with the dissociation constants used to determine them.

The test solutions were kept in 90 mL cells that were made of Polyether etherketone (PEEK).²⁶ All cells had sealed ports at the top that accommodate the permanent emplacement of a pH electrode, an opening to draw samples, and three 1/16" diameter Teflon lines for addition of acid, base, and the CO₂-argon mixture. The temperature was controlled by placing the test cells in a heated aluminum block of LBL design. The electric heater was mounted on an orbital shaker (Lab-Line Inc., Melrose Park, IL), and all solutions were shaken continuously at approximately 100 rpm. The solutions' pH values were monitored by a computer-operated pH control system (pH-stat, see Section 4.2), and were controlled by hand. Combination pH electrodes from Broadley-James Corp., Model E-1393EC1-A03BC were used to monitor the solutions pH values. Table V. Concentrations (in percent) of carbon dioxide gas in argon to maintain a total dissolved carbonate concentration of 1.531×10^{-2} M in UE-25p #1 water at different pH's together with the values for Henry's constant and the dissociation constants for carbonic acid used to determine the pCO₂ values.

	60°C		60°C
рН б	52.03	pK _H	1.77
pH 7	11.22	pK ₁	6.17
pH 8.5	0.3909	pK ₂	9.91

The combination pH electrodes from Broadley-James Corp. were used in the experiments because of their expectedly better long-term pH stability. These electrodes, however, went out of calibration quite often and the computer controlled monitoring of the pH showed deviations of up to 0.2 units. These deviations were the result of the electrode's going out of calibration, and not a result of the instability of the experimental equilibria. Therefore, we did not use the pH-stat for pH adjustment. We calibrated the electrodes often and only adjusted the solution's pH by hand after each calibration. The deterioration of the electrode is mainly due to the dissolution of the Ag/AgCl layer of the reference electrode wire and also of the wire used in the pH sensing compartment itself; the solubility of AgCl increases approximately 240 times when the temperature changes from 10° to 100°C. Although the manufacturer claims the working range of these electrodes is up to 100°C, we were unable to use the electrodes continuously with pH-stat computer controlled pH adjustment. Therefore, we allowed pH-stat to continuously monitor the pH, but we performed the pH adjustment by hand.

The effect on the composition of the UE-25p #1 groundwater due to the addition of acid and/or base during the experiment is a rather complicated one. At the start of the oversaturation experiments, the ionic strengths of the solutions were increased by factors that ranged from 1.3 to 4.9 simply with the introduction of the actinide stock solutions and the sodium hydroxide needed to neutralize the perchloric acid in the stock solutions. Therefore, the initial ionic strengths ranged from ~0.03 to 0.1 M. During the course of the solubility experiments, 0.1 M perchloric acid and sodium hydroxide were used to maintain the desired pH of the solubility experiments. Also during that time, we assayed the solutions for concentration and speciation determinations. The volumes of acid and base added to the experiments ranged from 2 to over 20 mL; whereas, to perform all of the concentration and speciation determinations throughout the experiment, we removed ~30 to 50 mL of solution. The net effect on 70 mL of an initially 0.04 M solution in

removing 50 mL for assays and then adding 20 mL of 0.1 M acid and/or base solutions to adjust the pH would result in increasing the ionic strength to approximately 0.07 M. For solutions that were initially 0.1 M, they would remain at roughly the same ionic strength. Therefore, in these cases we would have final ionic strengths that range from ~0.07 to 0.1 M. In all cases, however, the constituents in UE-25p #1 groundwater other than sodium (and perchlorate) would be diluted by roughly a factor of 7. Of course the solutions to which we added very little acid and/or base, the final ionic strengths will be closer to their initial ionic strengths. The addition of acid and/base, however, ran concurrently with the removal of solution volumes for assays, so it is rather difficult to establish exactly the true ionic strength of these solutions at the conclusion of the experiments.

The addition of acid and base to the solutions during the course of the experiments appears to have had an averaging effect on the ionic strength of the solutions when compared with the effect of introducing the actinide stock solutions. Upon starting the experiments, the ionic strengths ranged from ~0.03 to 0.1 M, but by adding 0.1 M solutions of acid and base during the course of the experiments, we narrowed that range. With an approximate analysis based on final volumes, we can only say that the final ionic strengths in all of the solubility experiments were probably closer to 0.1 M, \pm 50%, and that the primary constituents are sodium and perchlorate. This five fold increase in the ionic strength of UE-25p #1 groundwater definitely shows the difficulties in trying to maintain constant solution conditions in long term solubility experiments such as these.

4.5. Phase Separation

Achievement of steady-state conditions for the solubility measurements was monitored by sampling aliquots of the solution phases and analyzing for the respective radioisotope as a function of time. We used Centricon-30 centrifugal filters (Amicon Corp., Danvers, MA) for separating the phases of the neptunium, plutonium, and americium solutions. For the separations, the centrifuge (High-speed centrifuge, Model HSC-1000, Savant Instruments Inc.) was heated with a circulating water bath to 60°C. The filters contain a YM-type membrane with a calculated pore size of 4.1 nm²⁷. To ascertain that we achieved complete phase separation and minimal adsorption on the filters during the preparation of the solution assays, we conducted a series of filtration tests.

For each solution, these tests were done at different times during the equilibration period. We used one filter per solution and filtered consecutive 500 μ L portions of solution through it. Each filtrate was acidified to minimize sorption in the filtrate-collection container and an assay was taken. The concentration was plotted as a function of total volume passed through the filter. This was repeated until the assays showed a constant concentration. The volume necessary to saturate

the filter was the cumulative amount of volumes used until the assay concentration remained constant. The presaturation volume was radionuclide-dependent.

We determined and used the following preconditioning volumes: 500 μ L, 500 μ L, and 2500 μ L for the neptunium, plutonium, and neodymium/americium solutions, respectively.

4.6. Analysis

After separation of the solution and the solid phases, the two components were analyzed separately. Concentration measurements of the supernatants were made by counting liquid aliquots with a germanium low-energy counting system (LBL design). For ²³⁷Np and ²⁴¹Am, the 29.38 keV and 59.54 keV- γ -ray lines were used, respectively. ²³⁹Pu was analyzed by utilizing the U *L* x-rays coming from the α -decay of the plutonium. Possible contributions to the *L* x-rays from the decays of other radionuclides, also present in small amounts, were corrected by subtraction.²⁸ In selected cases, liquid scintillation counting was also used for plutonium concentration determinations (LKB Instruments, Inc., Wallac Oy, Model 1219 RackBeta). We used the "Pulse Shape Analysis" feature of the 1219 RackBeta to discriminate all β -emitting solution contaminants from the plutonium α -radiation. Repeated sample counting and the observation of a constant count rate in the α -window ensured no β -contribution to the α -count.

4.7. Criteria for Steady-State Concentrations

Constant concentrations over time with minimal deviation during that time span are the criteria for determining the average steady-state concentration from the individual concentration measurements. For experiments in which the aqueous concentration continually increases (or decreases), the final steady-state concentration will be equal to the final concentration measurement taken from the experiment. This concentration may not be the actual steady-state concentration, and all that can be stated is that the steady-state concentration is probably greater than (or less than) or equal to the value reported.

4.8. Eh Measurements

We are aware of the problems associated with Eh measurements in groundwater systems. Lindberg and Runnells²⁹ point out that in the apparent absence of an internal redox equilibrium, as is the case for many groundwater systems, Eh measurements may not accurately predict the equilibrium chemistry of the system. Because our experimental systems contained a very limited number of components and may be well poised, we measured the Eh at the end of each solubility

experiment. In several cases, we obtained stable readings only after measuring as long as 24 hours. This drifting does not lend much confidence in the obtained Eh value. Despite these limitations, we conducted these measurements in order to supply future chemical modeling efforts (neptunium, plutonium, and americium solubilities in UE-25p #1 water solutions) with an approximate value indicating the general oxic nature of the system. Without modeling, however, the Eh measurements are only of limited value, because they may represent a combination of many different redox reactions for each individual solubility solution despite the limited number of components.

We measured the Eh with a platinum electrode versus a Ag/AgCl/sat. NaCl reference. We cleaned the platinum electrode with 6 M HNO₃ before and after each measurement. The electrode setup was checked with "Zobell Solution" before and after each measurement.^{30,31}

4.9. Identification of Solids

The solid compounds were analyzed by x-ray powder diffraction measurements. A few micrograms of each actinide precipitate were placed in a 0.33 mm diameter quartz capillary tube, and the tube was sealed with an oxy-butane microtorch. The tube was mounted in an 11.46 cm diameter Debye-Scherrer camera and then irradiated with x-rays from a Norelco III x-ray generator (Phillips Electronics, Inc.). Copper K_{α} radiation filtered through nickel was used.

5. RESULTS AND DISCUSSION

5.1. Neptunium. *5.1.1. Solubility.* The comparison of the steady-state concentrations of both the over- and undersaturation neptunium experiments are shown in Figure 1. For the oversaturation experiments the neptunium was initially introduced as NpO_2^+ into the UE-25p #1 groundwater. For the undersaturation experiments, the solids formed from oversaturation were added to fresh solutions of UE-25p #1 groundwater. The steady-state concentrations and the solutions' Eh values are given in Table VI. Both the neptunium over- and undersaturation experiments provided the same steady-state solution potentials within the uncertainties of the measurement. For both experiments, the pH 6 solubility experiment had the highest redox potential and the pH 8.5 solubility experiment had the lowest potential. Concentration profiles as a function of equilibration time and pH for both over- and undersaturation experiments are shown in Figure 2. The individual measurements are listed in Appendix A. We experienced some difficulties with the neptunium undersaturation cell at pH 6. Near day 189 of this experiment we measured a pH of 7.00. This rise in pH precipitated some of the aqueous neptunium and lowered the soluble neptunium concentration. At this point, we restarted the undersaturation experiment by removing

the solid phase and placing it in fresh UE-25p #1 water at pH 6. We kept it at this pH for the remainder of the experiment. The aqueous concentration of neptunium did not increase back to the concentration before the pH increase. There was a slight increase in the aqueous concentration, but with time the concentration leveled off to a steady-state value midway between the pH 6 and pH 7 steady-state concentrations. At the point that the pH increase was first discovered, we removed a small portion of the solid phase for x-ray analysis. The powder pattern obtained was different from both those obtained in the pH 6 and pH 7 oversaturation experiments, and contained nearly twice as many diffraction lines. The fact that the solid phase removed from the second undersaturation experiment remained in contact with the UE-25p #1 water at pH 6 for nearly 400 days, combined with the lower solubility and the different powder pattern, suggests that a different, more crystalline solid phase formed with a more thermodynamically stable crystal lattice which led to lower solubilities in the groundwater.

Results from neptunium filtration experiments have been given elsewhere.³² The filtration experiments are described in Section 4.5. We found no volume effect for the neptunium in either the over- or undersaturation experiments. We therefore used 500 μ L of solution as presaturation volume for routine separations.

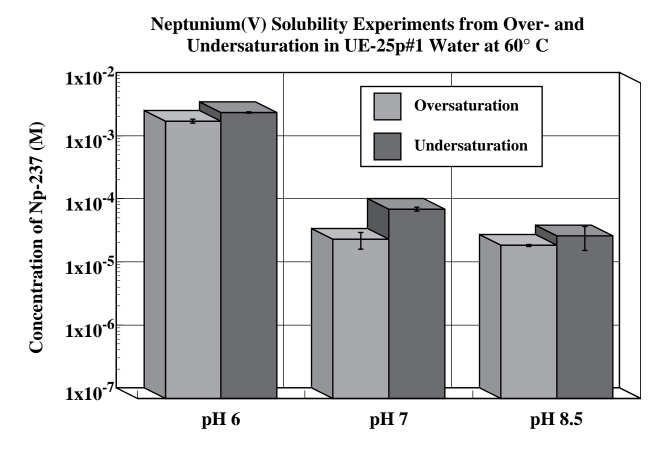
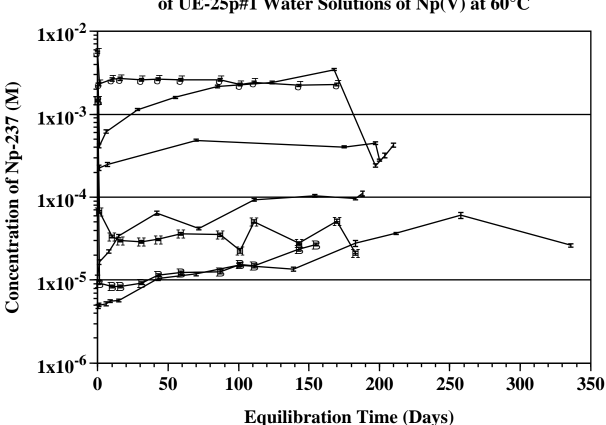


Figure 1. Results of the neptunium solubility experiments from over- and undersaturation in UE-25p #1 groundwater as a function of pH.

Table VI.Comparison of steady-state solution concentrations and Eh for neptunium over-
and undersaturation experiments in UE-25p #1 groundwater at 60°C.

	Neptunium(V)				
pН	Concentration (M)		Eh (mV vs. NHE)		
	Oversaturation	Undersaturation	Oversaturation	Undersaturation	
6	$(2.5 \pm 0.2) \ge 10^{-3}$	$(3.4 \pm 0.1) \ge 10^{-3}$	(414 ± 15)	(406 ± 15)	
7	$(3.4 \pm 1.0) \times 10^{-5}$	$(1.0 \pm 0.1) \ge 10^{-4}$	(367 ± 15)	(366 ± 15)	
8.5	$(2.7 \pm 0.1) \ge 10^{-5}$	$(3.8 \pm 1.6) \ge 10^{-5}$	(212 ± 15)	(239 ± 15)	



Approach to Equilibrium from Over- and Undersaturation of UE-25p#1 Water Solutions of Np(V) at 60°C

-J pH 6, Oversaturation	— pH 6, Undersaturation
——— pH 7, Oversaturation	— pH 7, Undersaturation
B pH 8.5, Oversaturation	— pH 8.5, Undersaturation
— pH 6, Undersaturation	

Solution concentrations of 237 Np in contact with precipitate obtained from supersaturation of UE-25p #1 groundwater at 60°C as a function of time. pH Figure 2. 6.0 ± 0.1 (closed circles), pH 7.0 ± 0.1 (closed triangles), and pH 8.5 ± 0.1 (closed squares). The neptunium was added initially (day 0) as NpO_{2^+} ; initial concentrations were 5.6 x 10^{-3} M (pH 6), 1.5 x 10^{-3} M (pH 7), and 1.5 x 10^{-3} M (pH 8.5). Undersaturation experiments at pH 6.0 (open circles, open diamonds), pH 7.0 (open triangles), and pH 8.5 (open squares) were started with precipitates obtained in the supersaturation experiments at their respective pH values.

In summary, the neptunium solubility decreased with increasing pH in both the over- and undersaturation experiments. All of the undersaturation cells had steady-state aqueous neptunium concentrations slightly higher than the steady-state concentrations from oversaturation. However the steady-state concentrations from over- and undersaturation agreed to within an order of magnitude.

5.1.2. Speciation. The supernatant solutions from the neptunium solubility experiments were analyzed by absorption spectrophotometry to determine the oxidation state and speciation. The spectra for the 60°C oversaturation experiments at pH 6, 7, and 8.5 are shown in Figures 3, 4, and 5, respectively. The spectra for the 60°C undersaturation experiments at pH 6, 7, and 8.5 are shown in Figures 6, 7, and 8, respectively. The solutions underwent phase separation in a centrifuge heated to 60°C; however, the quartz cuvettes used for the spectral measurements were not thermostated. Even though spectral measurements were made immediately after the separation, the supernatant solutions may have cooled slightly before finishing the measurements. Depending on the pH, the spectra show either the NpO₂⁺ main absorption band at 980.6 nm and/or an

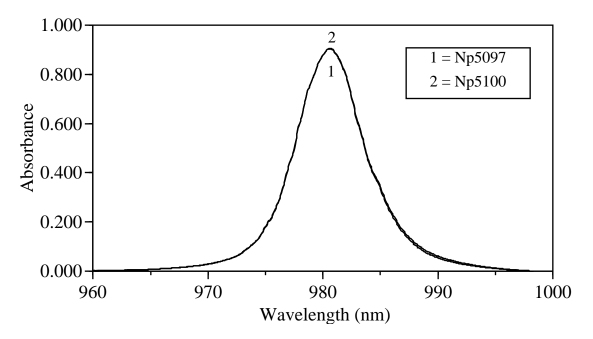


Figure 3. Near-IR absorption spectra of Np supernatant solutions at steady-state formed in UE-25p #1 groundwater from oversaturation at 60°C in pH 6: (1) at the experimental pH and (2) after acidification with HClO₄ to pH 2.

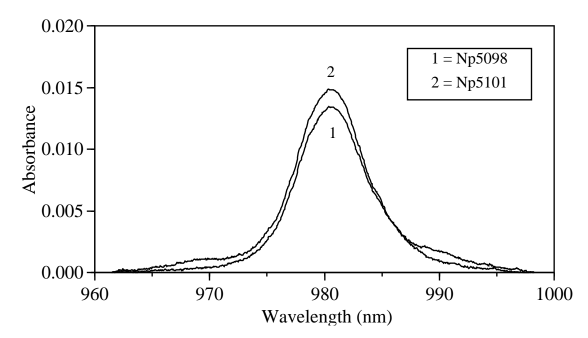


Figure 4. Near-IR absorption spectra of Np supernatant solutions at steady-state formed in UE-25p #1 groundwater from oversaturation at 60°C in pH 7: (1) at the experimental pH and (2) after acidification with HClO₄ to pH 1.

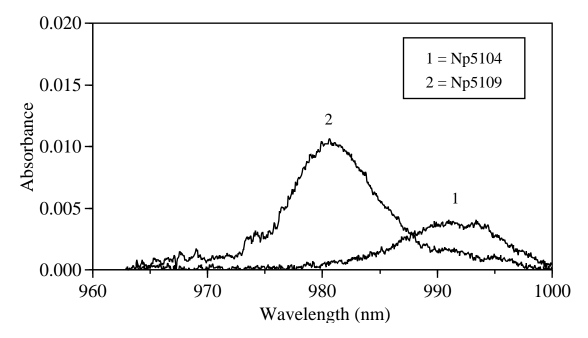


Figure 5. Near-IR absorption spectra of Np supernatant solutions at steady-state formed in UE-25p #1 groundwater from oversaturation at 60°C in pH 8.5: (1) at the experimental pH; (2) after acidification with HClO₄ to pH 1.

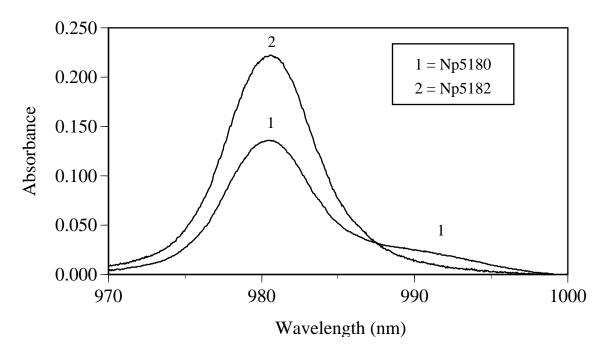


Figure 6. Near-IR absorption spectra of Np supernatant solutions at steady-state formed in UE-25p #1 groundwater from undersaturation at 60°C in pH 6: (1) at the experimental pH; (2) after acidification with HClO₄ to pH 0.

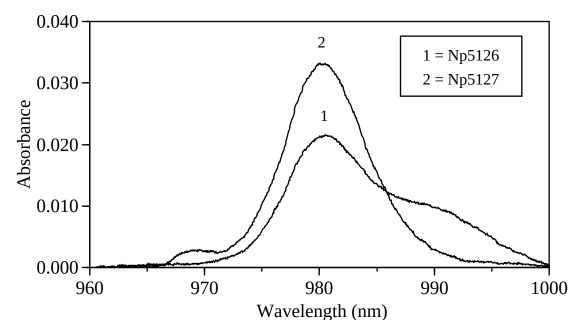


Figure 7. Near-IR absorption spectra of Np supernatant solutions at steady state formed in UE-25p #1 groundwater from undersaturation at 60°C in pH 7: (1) at the experimental pH; (2) after acidification with HClO₄ to pH 0.

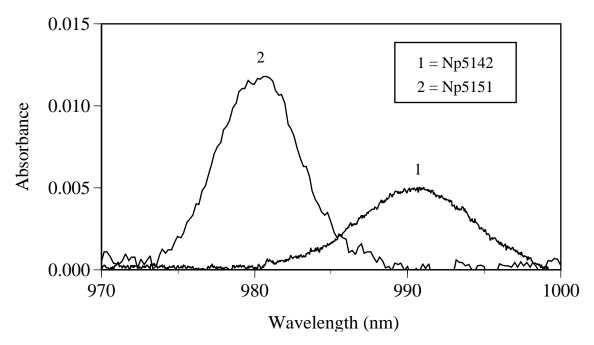


Figure 8. Near-IR absorption spectra of Np supernatant solutions at steady-state formed in UE-25p #1 groundwater from undersaturation at 60°C in pH 8.5: (1) at the experimental pH; (2) after acidification with HClO₄ to pH 0.

additional band at 991 nm that increases with pH due to the increasing carbonate complexation. The band at 980.6 nm is characteristic of uncomplexed NpO_2^+ . The band at 991 nm was established by Nitsche, et al.³³ It is typical for neptunyl(V) carbonate complexation.

To show that the band at 991 nm was indeed due to neptunyl(V) complexation by carbonate, we acidified the solutions to liberate the carbonate as carbon dioxide. Each of the solutions was acidified to pH ~ 0 with HClO₄ and observed for gas formation. All solutions, with the exception of the pH 6 oversaturation solution, gave off detectable amounts of CO_2 gas. The spectra of the acidified cell was then immediately taken. The absorption band at 991 nm disappeared, and only the band at 980.6 nm from uncomplexed neptunium(V) remained. Through this procedure, we established that all solutions contained neptunyl(V) carbonate complex(es) with the exception of the pH 6 sample from oversaturation. The spectrum of this sample, before acidification, showed only the absorption band at 980.6 nm, and upon addition of HClO₄ showed no gas formation or change in the absorption spectrum. We determined the relative amounts of free neptunium and complexed neptunium (present as carbonate complex) from the difference between the total amount of neptunium in solution (determined by γ -spectroscopy) and the free NpO₂⁺ determined from the 980.6 nm absorption band. The results of our neptunium speciation study are summarized in Table VII. It should be noted that we analyzed the pH 6 undersaturation solution after the drop in pH occurred (as discussed in the solubility section). The spectra for this solution was taken several hundred days after we adjusted the pH to 6. However, the relative

amounts of complexed/uncomplexed neptunium was slightly higher than in the pH 6 oversaturation solution where no complexation was evident.

	Nej	ptunium Carbonate Co	omplexation (%)	
pН	Oversa	turation	Undersa	aturation
	NpO ₂ +	NpO ₂ (CO ₃) ⁻	NpO ₂ +	NpO ₂ (CO ₃) ⁻
6	100	0	76	24
7	90	10	66	34
8.5	0	100	0	100

Table VII.	Comparison of extent of carbonate complexation for steady-state solutions of
	neptunium in UE-25p #1 groundwater from over- and undersaturation at 60°C.

We used a non-linear, least-squares peak fitting routine (Spectra Calc, Galactic Industries Corporation, Salem, New Hampshire) to deconvolute the five absorption spectra that showed carbonate complexation in order to determine the complex stoichiometry. The results for the spectrum from the pH 7 undersaturation solution, shown in Figure 9, indicate that the spectrum is composed of only two peaks: one at 980.6 nm for the uncomplexed NpO₂⁺ and a second one at 991 nm for the NpO₂CO₃⁻ complex. Higher complexes such as the NpO₂(CO₃)₂³⁻ complex, did not fit the spectrum. This was the case for all of the complexed spectra. The pH 8.5 solutions, from both over- and undersaturation cells, fit only one species; that of NpO₂CO₃⁻ because all of the neptunium in solution was complexed.

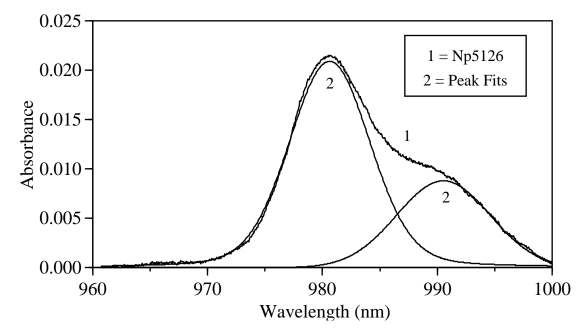


Figure 9. Deconvolution of the spectrum from the neptunium undersaturation experiment at pH 7 and 60°C showing the two deconvoluted peaks at 980.6 and 991 nm.

5.1.3. Identification of Solids. The precipitates formed in the neptunium solutions were collected by centrifugation, washed with a small amount of CO_2 -free water, and dried with a jet of argon. When wet, all precipitates, from both the over- and undersaturation experiments, were a very faint green color; however, when completely dry, they all became white crystalline solids. X-ray powder diffraction patterns taken from precipitates produced distinct lines; crystal lattice dspacings and relative intensities are listed in Tables VIII, IX, and X.

All of the solids obtained from the three neptunium oversaturation experiments matched well with each other. Twelve of the 26 lines from the pH 6 solid matched with the other two oversaturation solids to within ± 0.02 Å. Twelve of the 14 lines from the pH 7 solid matched, and twelve of the 21 lines matched from the pH 8.5 solid. The solids from the pH 6 and pH 8.5 oversaturation cells matched even better. Twenty of the 26 lines in the pH 6 solid matched with the pH 8.5 solid to within ± 0.02 Å, and 18 of 21 lines in the pH 8.5 solid matched with the pH 6 solid. The pH 7 solid had relatively fewer "weak" and "trace" lines, probably as a result of its shorter exposure time. However, the strong and medium intensity lines matched well with the pH 6 and pH 8.5 solids. The comparison of the three oversaturation solids is given in Table VIII.

As was indicated in a previous report on the UE-25p #1 solids³², the comparison of the 60°C oversaturation solid patterns with patterns in the literature agreed to varying degrees. All of the solids obtained were similar to ternary sodium neptunium carbonate hydrates of different

stoichiometry and/or water content. It should be noted that many of the corresponding solids described in the literature are metastable, difficult to reproduce, and show the extent to which line spacings vary.^{34,35,36,37}

Four solids were obtained from the neptunium experiments from undersaturation: one from each of the pH 7 and pH 8.5 cells at the conclusion of these experiments, and two from the pH 6 cell. The solids from the pH 6 cell were removed at two separate times during the experiment: 1) at day 210, when the solubility dropped an order of magnitude, and 2) at the end of the experiment after almost 400 days of equilibration time. Table X lists the d-spacings and intensities for all of the final solids obtained in these experiments. Table IX lists the d-spacings and intensities for the solid removed at day 210 from the pH 6 undersaturation cell.

The solid removed from the pH 6 undersaturation cell, after the drop in solubility, gave a powder pattern with d-spacings that were very different from those of the starting solid from the oversaturation experiment (from either the pH 6 or the pH 7 cell). Table IX shows a comparison of the first solid removed from the pH 6 cell with the solids obtained from the oversaturation experiments at pH 6 and pH 7. The first undersaturation solid from the pH 6 cell contains nearly twice as many diffraction lines, very few of which match within ± 0.03 Å of the oversaturation solids. The fact that the solubility dropped one order of magnitude, combined with the different powder pattern obtained, suggests that a different, more crystalline solid phase formed. The solid, most likely, has a more thermodynamically stable crystal lattice which lead to a lower solubility in the groundwater. Comparison of the powder pattern from this solid with patterns found in the literature are given in Tables XI and XII. The solid fit poorly with all of the references it was compared to. The best fit was obtained with the published patterns NaNpO₂CO₃•3.5H₂O and NaNpO₂CO₃.³⁶ In the pH 6 solid, 10 of the 48 lines matched the NaNpO₂CO₃•3.5H₂O solid to within ± 0.03 Å and only 5 matched to within ± 0.01 Å. The match with the NaNpO₂CO₃ solid was slightly better; with 11 of the 48 lines matching to within ± 0.03 Å and 7 of the 48 lines matching within ± 0.01 Å. At this time the solid remains unidentified, but based on the predominance of sodium in the groundwater accompanied by the high carbonate content, leads us to conclude that this solid is of the sodium neptunium(V) carbonate type.

The solid removed from the pH 6 undersaturation cell after almost 400 days of equilibration time gave a different powder pattern than the solid removed at 200 days. The pattern of the solid removed from the cell at the end of the experiment is shown in Table XIII along with a comparison to $KNpO_2CO_3$ and $NaNpO_2CO_3$ reference patterns from the literature. The powder pattern of the pH 6 solid matched 11 out of 34 lines from the $KNpO_2CO_3$ to within ± 0.01 Å and 14 out of 34 lines to within ± 0.03 Å. It matched 11 out of 34 lines from the $NaNpO_2CO_3$ literature reference to within \pm 0.01 Å and 18 out of 34 lines to within \pm 0.03 Å. Again, due to the predominance of sodium in the UE-25p #1 groundwater, it would appear that the solid formed was very similar to NaNpO₂CO₃.

The powder pattern from the pH 7 undersaturation solid matched that of the literature reference pattern of Na_{0.6}NpO₂(CO₃)_{0.8} · 2.5 H₂O³⁴ relatively well. The comparison of the two solids is given in Table XIV. Twenty two out of 48 lines fit to within \pm 0.01 Å, and 31 out of 48 lines fit to within \pm 0.03 Å. It should be noted that the x-ray diffraction pattern for the Na_{0.6}NpO₂(CO₃)_{0.8} · 2.5 H₂O solid has much in common with hydrate forms of sodium monocarbonateneptunylate³⁴ NaNpO₂CO₃, and it is possible that we obtained a mixture of the Na_{0.6}NpO₂(CO₃)_{0.8} · 2.5 H₂O and the NaNpO₂CO₃ solids. In light of the difficulties in justifying the stoichiometry of the Na_{0.6}NpO₂(CO₃)_{0.8} · 2.5 H₂O solid, it is also possible that the solid obtained was simply NaNpO₂CO₃.

The powder pattern from the pH 8.5 undersaturation solid matched that of the literature reference pattern of $Na_3NpO_2(CO_3)_2 \cdot nH_2O^{35}$ fairly well. A comparison of the two patterns is given in Table XV. Nine out of 30 lines fit to within \pm 0.01 Å and 19 out of 30 lines fit to within \pm 0.03 Å. The fit with the solid of higher carbonate content is consistent with the increase in pH and carbonate available for complexation and solid formation in the cell. This is also consistent with the solubility decrease, as the higher pH increased the free carbonate in solution and precipitated more of the aqueous neptunium resulting in a lower neptunium solubility. This is simply the pH effect on the solubility of the carbonate containing solid in solution. Since there are such discrepancies in the literature, we suggest that a detailed study of these solids be performed, so that differences can be clearly explained. We also suggest measuring the solubility of defined ternary neptunium carbonate solids as a function of solution pH in order to obtain the solubility products required for modeling calculations.

pH 6	5	pH 7	7	pH 8	.5
(oversat.)		(oversat.)		(oversat.)	
<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī
9.41	vs	9.41	S	9.43	S
4.87	m	4.88	m-	4.86	w
4.70	s	4.72	s-	4.70	s
4.34	s	4.33	s-	4.33	s
3.93	s-	3.95	m	3.93	s-
3.40	t	3.38	W	3.38	w
3.26	m	3.24	W	3.25	w+
3.20	t				
3.15	w	3.14	t	3.15	w
3.07	s	3.06	m-	3.06	s
2.67	m	2.67	W-	2.68	w
2.55	w			2.54	t
2.43	w			2.44	t
2.36	m	2.36	t	2.36	w-
2.25	t				
2.17	t	2.17	t	2.16	t
2.12	m	2.12	t	2.12	w
2.07	w	2.07	t	2.08	w
1.97	t			1.97	t
1.93	w			1.94	t
1.90	t			1.90	t
1.89	t			1.89	t
1.76	w+			1.76	w
1.62	t				
1.60	t				
1.57	t				

Table VIII.X-ray powder diffraction patterns of neptunium solid phases from
oversaturation in UE-25p #1 groundwater at 60°C and pH 6, 7,
and 8.5.

 (a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

Table IX.X-ray powder diffraction pattern of neptunium solid phase from
undersaturation in UE-25p #1 groundwater at 60°C and pH 6
removed after day 210, compared with oversaturation solids from pH
6 and 7.

pH 6 (1st ppt.)		рН б		pH 7	
(unders		(oversat.)		(oversat.)	
	,				
<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī
9.85	w				
		9.41	V S	9.41	s
7.48	w				
6.57	S				
5.11	t	4.97		4.00	
4.80	t	4.87	m	4.88	m-
4.80	ι	4.70	s	4.72	s-
4.55	m	4.70	3	4.72	3-
		4.34	s	4.33	s-
4.24	w+				
4.06	m-				
		3.93	S-	3.95	m
3.72	t				
3.51	s	3.40	t	3.38	w
3.34	t	5.40	ι	5.30	w
		3.26	m	3.24	w
3.21	m+	3.20	t		
		3.15	W	3.14	t
3.00	t	3.07	s	3.06	m-
2.96	t				
2.84	t				
2.77	t				
2.68	t	2.67	m	2.67	W-
2.61 2.55	t m-	2.55	w		
2.33	t	2.33	w		
2.40	t				
		2.36	m	2.36	t
2.31 2.26	t t	2.25	t		
2.20	w	2.25	L		
2.17	w	2.17	t	2.17	t
2.10	w	2.12	m	2.12	t
1.00		2.07	w	2.07	t
1.99 1.93	s t	1.97 1.93	t w		
1.90	w	1.90	t		
		1.89	t		
1.85	t				
1.81 1.80	t w-				
1.75	w- w	1.76	w+		
1.72	w				
1.66	t				
		1.62 1.60	t t		
1.56	t	1.60	t t		
1.50	w-	1.07	-		
1.49	t				
1.47	t				
1.41	t t				
1.39	t				
1.31	t				
1.27	t				
1.24	w-				
1.21	t t				
1.15	t				
	-				

⁽a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

pH 6	6	"""	7		5
(unders		pH (unders		pH 8 (unders	
(unders	ai.)	(unders	al.)	(unders	a)
<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī
0.21		9.43	s		
9.31	s			8.63	w
				6.92	t
6.55	W	6.50 6.13	t t	6.58	t
5.08	m	0.15	t	5.09	m+
4.87	w	4.89	w		
4.70	m	4.70	S-		
4.39	W	1.24		4.43	S-
		4.34 4.21	S-		
4.07	s	4.21	t t	4.06	s
3.99	t	1.05	ı	1.00	5
		3.94	m+		
		3.57	t		
		3.40	w+		
3.33	m			3.34	m
3.23	m-	3.25	m	3.25	t
3.14	W-	3.19 3.14	m-		
3.05	t v-	3.07	m- m+	3.10	t
5.05	·	2.88	W-	2.93	w
				2.82	m-
2.68	m-	2.68	m-	2.68	w
2.55	m+	2.55	W	2.55	S-
2.44	t	2.43	W-		
2.36	t	2.37	W		
2.28	w+	2.26	W-	2.27 2.21	W
2.21	t m-	2.18	m	2.21	w m
2.10	III	2.13	m+	2.10	
		2.08	m		
2.03	t			2.03	t
1.99	s	2.00	m+	1.99	s
		1.94	m-		
		1.91	W-		
1.05		1.88	t	1.05	
1.85 1.80	W W	1.86 1.83	t t	1.85	w w
1.00	vv	1.85	w-	1.00	vv
1.73	w+	1.73	w-	1.73	w
1.65	w+	1.67	t	1.65	w
1.62	t	1.63	W-		
1.59	t	1.60	W-	1.59	t
		1.58	w-		
1.50		1.52	t t	1.50	
1.50 1.48	w t	1.50 1.48	t t	1.50	t t
1.40	ι	1.48	t	1.4/	ι
1.42	t	1.41	w-	1.41	t
	-	1.39	ť		-
		1.37	t		
1.30	t	1.30	W-	1.29	t
1.28	t			1.28	w
1.25	w+	1.25	W-		
1.22		1.24	W-	1.24	W-
1.22	t	1.22 1.14	t t	1.22	t
		1.14	ı		
		Į		Į	

Table X.X-ray powder diffraction patterns of neptunium solid phases from
undersaturation in UE-25p #1 groundwater at 60°C and pH 6, 7,
and 8.5 (solids removed at the conclusion of all experiments).

⁽a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

pH 6 (1st ppt.) (undersat.)	NaNaOaCOac	2.511.0
		NaNpO ₂ CO ₃ •	3.5H2U
1 (²)	-		-
<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī
9.85	w	9.87	V S
7.48	w	,,	
6.57	S		
5.11	t		
	-	4.92	s
4.80	t		
4.55	m		
		4.41	m-
		4.35	m-
4.24	w+	4.23	m+
4.06	m-		
3.72	t		
		3.61	m-
3.51	S	3.50	W-
		3.43	W-
3.34	t		
		3.28	W-
3.21	m+	3.20	S
3.00	t		
2.96	t		
		2.91	m-
		2.89	m-
2.84	t		
2.77	t		
2.68	t		
2.61	t		
2.55	m-	2.52	W-
2.42	t	2.44	m-
2.40	t	2.37	t
		2.35	W-
2.31	t		
2.26	t		
2.20	w	2.20	t
2.17	W	2.17	W-
2.10	W	2.12	m+
1.99	S		

Table XI.X-ray powder diffraction pattern of neptunium solid phase from
undersaturation in UE-25p #1 groundwater at 60°C and pH 6 removed
after day 210, compared with the pattern of NaNpO₂CO₃•3.5 H₂O.³⁶

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

(b) The pH 6 pattern has 21 additional lines (w and t), not listed here, ranging from 1.93 Å to 1.22 Å.

pH 6 (1st	nnt)		
(unders		NaNpO ₂ 0	0
(unders	at.)	NaryO ₂	.03
<u>d(Å)</u>	Ι	<u>d(Å)</u>	Ι
	1	<u>u(11)</u>	1
9.85	w		
7.48	W		
6.57	S		
		5.35	V S
5.11	t		
4.80	t		
4.55	m		
		4.44	V S
4.24	w+		
4.06	m-	4.05	V S
3.72	t		
3.51	S		
3.34	t		
		3.25	V S
3.21	m+		
3.00	t		
2.96	t		
2.84	t	2.87	W
2.77	t	2.77	V S
2.68	t	2.67	m+
2.61	t		
2.55	m-		
2.42	t	2.43	m+
2.40	t		
2.31	t		
2.26	t		
2.20	W	2.21	W-
2.17	W	2.18	m+
2.10	W	2.08	V S
		2.06	W-
		2.01	W
1.99	S	1.96	V S
1.93	t	1.91	W

Table XII.X-ray powder diffraction pattern of neptunium solid phase from
undersaturation in UE-25p #1 groundwater at 60°C and pH 6 removed
after day 210, compared with the pattern of NaNpO2CO3.36

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

(b) The pH 6 pattern has 20 additional lines (w and t), not listed here, ranging from 1.90 Å to 1.22 Å.

pH 6 (2nd ppt.) (undersat.)		NaNpO ₂ CO ₃		KNpO ₂ CO ₃	
1/ 8	T		Ŧ	1(Å)	Ţ
<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī
9.31	S				
6.55	W				
		5.35	V S		
5.08	m			5.01	S
4.87	W				
4.70	m				
4.39	W	4.44	V S	4.44	m
4.07	S	4.05	VS	4.06	V S
3.99	t				
3.33	m			3.31	S
3.23	m-	3.25	V S		
3.14	W-				
3.05	t				
		2.87	w		
		2.77	V S		
2.68	m-	2.67	m+	2.66	m
2.55	m+			2.56	S
				2.49	m
2.44	t	2.43	m+		
2.36	t				
2.28	w+			2.28	m
2.21	t	2.21	W-	2.22	W
2.16	m-	2.18	m+	2.16	S
		2.08	V S		
		2.06	W-		
2.03	t	2.01	W	2.02	m
1.99	S	1.96	V S		
		1.91	w		
1.85	W			1.85	w+
				1.82	w+
1.80	W			1.79	m
1.73	w+				
1.65	w+			1.65	m+
1.62	t				
1.59	t			1.59	W
				1.55	t
1.50	W			1.50	W
1.48	t			1.48	W

Table XIII. X-ray powder diffraction pattern of neptunium solid phase from undersaturation in UE-25p #1 groundwater at 60°C and pH 6 removed at day 410 compared with the patterns of NaNpO₂CO₃ and KNpO₂CO₃.

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

(b) The pH 6 pattern has 5 additional lines (w and t), not listed here, ranging from 1.42 Å to 1.22 Å.

pH (under		Na _{0.6} NpO ₂ •2.5H		pH (under		Na _{0.6} NpO ₂ •2.5H	
<u>d(Å)</u>	Īa	<u>d(Å)</u>	Īa	<u>d(Å)</u>	Īa	<u>d(Å)</u>	Īa
		13.64	t			1.96	W-
		9.96	S-	1.94	m-	1.94	t
9.43	S			1.91	W-		
6.50	t			1.88	t		
		6.22	W	1.86	t		
6.13	t			1.83	t	1.83	w-
		4.96	S -			1.80	t
4.89	W	4.82	W	1.76	W-	1.77	w
4.70	S -			1.73	W-	1.72	t
4.34	S -	4.33	V S	1.67	t	1.70	w
4.21	t			1.63	W-	1.63	t
4.05	t			1.60	W-	1.61	w-
3.94	m+	3.97	S	1.58	W-	1.58	t
		3.73	t	1.52	t	1.53	W-
3.57	t			1.50	t	1.50	t
		3.44	t	1.48	t	1.49	t
		3.30	t			1.47	t
3.40	w+			1.45	t	1.44	w
3.25	m	3.22	S-			1.43	t
3.19	m-			1.41	W-	1.40	t
3.14	m-			1.39	t	1.38	t
3.07	m+	3.06	m-	1.37	t	1.37	t
2.88	W-	2.89	t			1.35	t
		2.80	W-			1.33	t
		2.77	W-	1.30	W-	1.29	t
2.68	m-	2.70	m+			1.28	t
		2.62	w+	1.25	W-	1.25	t
2.55	w			1.24	W-	1.24	t
2.43	W-	2.47	w	1.22	t	1.23	t
2.37	W	2.40	w+			1.21	t
		2.34	t			1.20	t
		2.30	t			1.16	t
2.26	W-			1.14	t	1.15	t
2.18	m	2.16	m-		-	1.13	t
2.13	m+	2.10	w			1.11	t
2.08	m	2.06	m				2
2.00	m+	1.98	w-				

Table XIV. X-ray powder diffraction pattern of neptunium solid phase from undersaturation in UE-25p #1 groundwater at 60°C and pH 7 compared with the pattern of $Na_{0.6}NpO_2(CO_3)_{0.8}$ •2.5H₂O.

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

Table XV. X-ray powder diffraction pattern of neptunium solid phase from undersaturation in UE-25p #1 groundwater at 60°C and pH 8.5 compared with the pattern of $Na_3NpO_2(CO_3)_2$ •n H₂O.

pH 8.5 (undersat.)		Na ₃ NpO ₂ (CC	O ₃)₂•n H₂O
<u>d(Å)</u>	<u>I</u> a	<u>d(Å)</u>	<u>I</u> a
8.63	w		
6.92	t		
6.58	t		
		6.07	W
		5.75	w+
5.09	m+		
4.43	S-	4.39	VS
		4.29	W-
4.06	s		
		4.00	w+
3.34	m		
3.25	t		
3.10	t	3.11	W-
		2.99	W-
2.93	w	2.91	W-
2.82	m-	2.80	W-
2.68	w	2.71	t
2.55	S-	2.53	m
		2.34	t
2.27	w		
2.21	w	2.19	m-
2.16	m	2.14	t
		2.10	t
2.03	t		
1.99	s	2.00	W-
		1.97	W-
		1.94	W-
		1.90	W-
1.85	w	1.84	t
1.80	W	1.80	t
1.73	W	1.74	t
		1.70	t
1.65	W	1.67	t
1.59	t	1.60	t
		1.57	t
		1.54	t
		1.51	t
1.50	t	1.50	t
1.47	t	1.46	W-
1.41	t	1.43	t
		1.38	t
1.29	t	1.28	t
1.28	W	1.27	t
1.24	w-		
1.22	t		

⁽a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

5.2. Plutonium. *5.2.1. Solubility*. Results of the plutonium solubility studies are shown in Figure 10. The plutonium was initially introduced as Pu^{4+} into the UE-25p #1 groundwater. The steady-state concentrations and the solutions' Eh values are given in Table XVI. Concentration profiles as a function of time and pH for over- and undersaturation experiments at 60°C are shown in Figure 11. The individual measurements are listed in Appendix B. Results from plutonium filtration experiments at 60°C have been given elsewhere.³² Methods used for the filtration experiments are described in Section 4.5. We could not find any volume effect for the plutonium, so we used 500 μ L of solution as presaturation volume for routine separations.

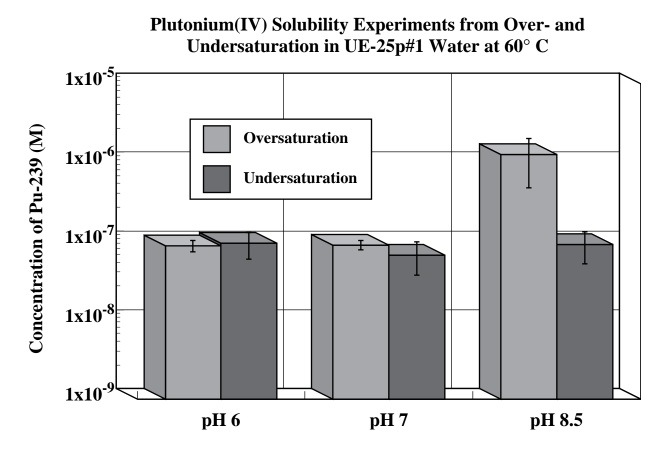
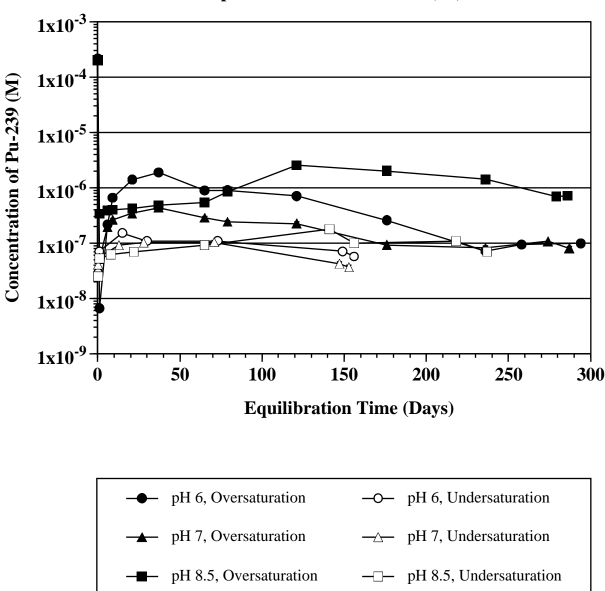


Figure 10. Results of the plutonium solubility experiments from over- and undersaturation in UE-25p #1 groundwater at 60°C as a function of pH.

	Plutonium(IV)				
pН	Concentr	ation (M)	Eh (mV	vs. NHE)	
	Oversaturation	Undersaturation	Oversaturation	Undersaturation	
6	$(8.8 \pm 1.4) \ge 10^{-8}$	$(9.5 \pm 3.6) \ge 10^{-8}$	(326 ± 15)	(312 ± 15)	
7	$(9.1 \pm 1.2) \ge 10^{-8}$	$(6.8 \pm 3.1) \ge 10^{-8}$	(334 ± 15)	(313 ± 15)	
8.5	$(1.3 \pm 0.8) \ge 10^{-6}$	$(9.2 \pm 4.0) \ge 10^{-8}$	(231 ± 15)	(218 ± 15)	

Table XVI.	Comparison of steady-state solution concentrations and Eh for plutonium from
	over- and undersaturation in UE-25p #1 water at 60°C.

In summary, plutonium steady-state concentrations from the over- and undersaturation experiments at pH 6 and 7 agree very well. There was no change in the steady-state concentrations. The concentration at pH 8.5 decreased by one order of magnitude. It appears that a new, more crystalline solid has formed, different from that from the oversaturation experiment. All of the undersaturation steady-state concentrations at the three pH's agree within the error of the measurements.



Approach to Equilibrium from Over- and Undersaturation of UE-25p#1 Water Solutions of Pu(IV) at 60°C

Figure 11 Solution concentrations of 239 Pu in contact with precipitate obtained from supersaturation of UE-25p #1 groundwater at 60°C as a function of time. pH 6.0 ± 0.1 (closed circles), pH 7.0 ± 0.1 (closed triangles), and pH 8.5 ± 0.1 (closed squares). The plutonium was added initially (day 0) as Pu⁴⁺; initial concentrations were 2.2 x 10⁻⁴ M (pH 6), 2.2 x 10⁻⁴ M (pH 7), and 2.0 x 10⁻⁴ M (pH 8.5). Undersaturation experiments at pH 6.0 (open circles), pH 7.0 (open triangles), and pH 8.5 (open squares) were started with precipitates obtained in the supersaturation experiments at their respective pH values.

5.2.2. Speciation

The plutonium supernatant solutions at steady-state were analyzed for their oxidation state distributions. Speciation studies are made difficult by the low solubility of plutonium in UE-25p #1 groundwater. The solutions' concentration levels lie below the sensitivity range of methods such as absorption spectrophotometry, which would allow the direct measurement of the species present. Therefore, we developed a method to determine the plutonium oxidation states indirectly. The method involves a combination of solvent extractions that allows us to determine the relative amounts of plutonium oxidation states in solution. Table XVII lists the methods used to determine the distribution of plutonium oxidation states in both the over- and undersaturation experiments at 60°C.

Table XVII. Methods used for determining Plutonium oxidation states in solution.

	Method	Oxidation State Distribution				
		Organic Phase	Aqueous Phase			
1.	0.025 M PMBP Extraction at pH = 0	(+4)	(+3,+5,+6, p) ^a			
2.	0.025 M PMBP Extraction at $pH = 0$ with $Cr_2O_7^{2-}$	(+3,+4)	(+5,+6, p) ^a			
3.	0.5 M HDEHP Extraction at $pH = 0$	(+4,+6)	(+3,+5, p) ^a			
4.	0.5 M HDEHP Extraction at $pH = 0$ with $Cr_2O_7^{2-}$	(+3,+4,+5,+6)	(p) ^a			

(a) (p) = Pu(IV) polymer

For the experiments, we used 4-benzoyl-3-methyl-1-phenyl-2-pyrazolin-5-one (PMBP) as the extractant for Pu(IV). At pH 0, PMBP extracts Pu(IV) from aqueous solutions without significant decomposition by oxidants, such as dichromate. We used relatively small concentrations of extractant, therefore aiding in the nuclear counting of the fractions. Using PMBP and dichromate, we extract Pu(III) and Pu(IV) together by oxidation of Pu(III) to Pu(IV). At pH 0, di-(2-ethylhexyl)-orthophosphoric acid (HDEHP) extracts Pu(IV) and Pu(VI). Upon introducing dichromate to the aqueous phase before extraction, Pu(III) is oxidized to Pu(IV) and Pu(V) is oxidized to Pu(VI). HDEHP with dichromate extracts Pu(III), Pu(IV), Pu(V), and Pu(VI) by oxidation of Pu(III) to Pu(IV) and of Pu(V) to Pu(VI). The oxidation of Pu(III) to Pu(IV) is instantaneous when dichromate is introduced to the aqueous solution. After several hours, disproportionation and/or the oxidation of Pu(IV) to Pu(V) by dichromate may occur, however,

experiments we conducted show the kinetics of these undesirable reactions are very slow and do not effect the extractions because they are completed in less than thirty minutes. This method was tested on plutonium solutions containing millimolar carbonate concentrations. The oxidation state distribution was measured at the same time by Laser Induced Photoacoustic Spectroscopy (LIPAS) and by our extraction method.³⁸ In testing this method, we also performed extractions with 2-thenoyltrifluoroacetone (TTA) at pH 0 in addition to the PMBP separations to verify that the two extractants produce the same results. At pH 0, TTA also extracts Pu(IV) from aqueous solutions. In all cases the results were the same.

For the experiments, we tried to maintain the 60°C temperature as well as it was experimentally possible. Although we mixed the aqueous and organic phase for one minute at ambient temperature using a Vortex mixer, we separated the phases in a centrifuge heated to 60°C. There is, however, the possibility that the oxidation state distributions at 60°C may have been affected by the handling-induced temperature change. Therefore, the derived results may be more semiquantitative.

In order to minimize the changing of plutonium oxidation states in solution, each of the extractions was carried out independently on a new solution sample. By taking differences in the relative amounts obtained in the four separations, the percent of each oxidation state can then be determined. Results of these studies are given in Table XVIII, and displayed in Figures 12, 13, and 14, for pH 6, 7, and 8.5, respectively.

The 60°C oversaturation solutions contained Pu(VI) almost completely. A small amount of Pu(IV) was present in the pH 8.5 solution. Pu(III) and Pu(IV) polymer and Pu(V) were present in insignificant amounts. The 60°C undersaturation solutions were a little less consistent. In the pH 6 and pH 8.5 solutions, the relative amounts of Pu(VI) dropped significantly and the Pu(V) percentage increased. Only the pH 7 solution was completely consistent with its oversaturation counterpart within the error of the measurement.

The observed valence distributions cannot be explained by disproportionation equilibria and complex stabilization.³⁹ It is possible that oxidation products formed by α -radiolysis of the water may cause the predominance of higher oxidation states. It is noteworthy that the solutions were filtered through a 4.1 nm filter prior to the oxidation state determination because we wanted to determine only the true soluble plutonium fraction without any colloidal/polymeric plutonium being present. Therefore, we refer to the Pu(IV) polymer, in the context of this determination, as to the fraction that is smaller than 4.1 nm. This colloidal or polymeric plutonium does not

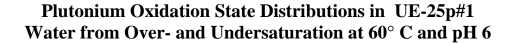
participate in the ionic solution equilibrium, and its removal will not immediately affect the oxidation state distribution in the solution⁴⁰.

Another possible explanation for the predominance of high oxidation states at steady-state may lie in the plutonium stock solution itself. Because of the limits of detection associated with our spectrophotometer, the solubility experiments may have contained initial concentrations of Pu(III), Pu(V), and Pu(VI) as high as 2×10^{-6} , 6×10^{-6} , and 2×10^{-7} M, respectively. These concentrations could result based on the maximum concentrations that would have been undetected in the 10⁻² M Pu(IV) stock solution. Assuming complete precipitation of the lower oxidation states, one could conclude that the potential initial concentrations of Pu(V) and Pu(VI)may represent the steady-state oxidation state distributions and the total aqueous plutonium concentrations. However, after completing oxidation state determinations in the corresponding plutonium undersaturation experiments in UE-25p #1 groundwater at 60° C, the above speculation cannot be confirmed, that trace level oxidation state impurities in the stock solution used to start the oversaturation experiments may be causing the plutonium solubility limits and the oxidation state distribution in solution. The undersaturation experiments were started with only the solid phases obtained in the oversaturation experiments. The results from oxidation state determinations on the undersaturation experiments showed combined relative amounts for Pu(V)and Pu(VI) of 96%, 98%, and 90% for solutions at pH 6, 7, and 8.5, respectively. The undersaturation experiment at pH 8.5 also showed a small amount of Pu(IV), approximately 10%. This finding is in excellent agreement with the result from the corresponding oversaturation experiment at pH 8.5. We also note that the slightly lower, yet still oxic, Eh value measured for the pH 6 and 8.5 experiments also support the increased presence of Pu(IV) and Pu(V). Here again, all three measurements were reproduced in the corresponding undersaturation experiments to within 21 mV.

In light of this, the predominance of higher oxidation states appears to be a real effect, yet more investigation into the species responsible for the oxidation of the plutonium in solution is desirable.

	Oxidation States (%)											
pН	Pu(IV)-polymer + Pu(III) Over Under		Pu(IV)		Pu(V)		Pu(VI)					
			Over	Under	Over	Under	Over	Under				
6	(1 ± 1)	(1 ± 1)	(1 ± 1)	(2 ± 1)	(4 ± 1)	(39 ± 4)	(94 ± 11)	(57 ± 6)				
7	(2 ± 1)	(1 ± 1)	(1 ± 1)	(1 ± 1)	(5 ± 1)	(9 ± 2)	(93 ± 11)	(89 ± 10)				
8.5	(5 ± 2)	(0)	(10 ± 1)	(10 ± 2)	(0)	(36 ± 4)	(86 ± 12)	(54 ± 7)				

Table XVIII.Plutonium oxidation state determinations in UE-25p #1 at pH values 6, 7, and 8.5and 60°C.



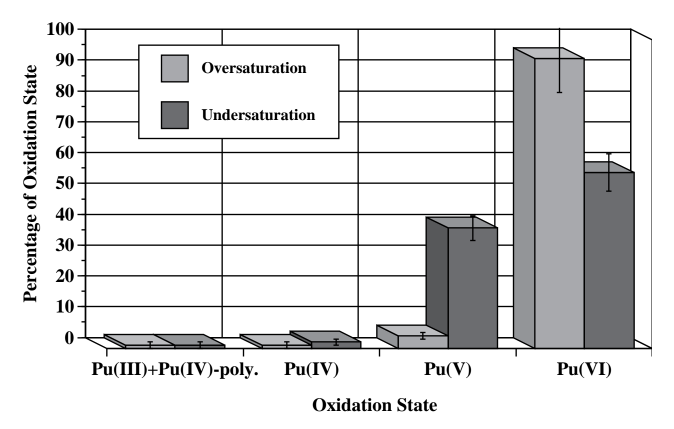
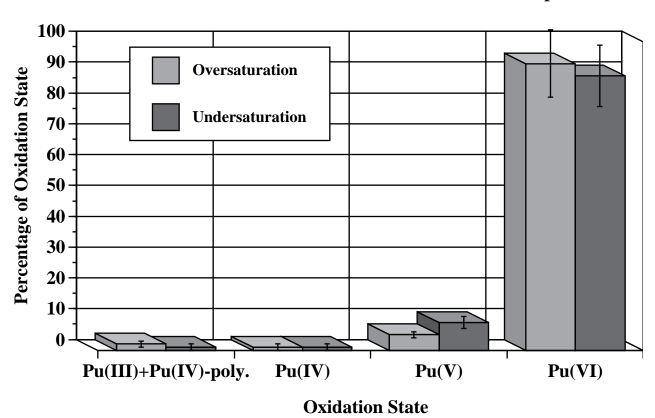
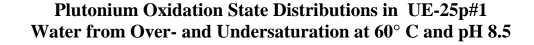


Figure 12. Plutonium oxidation state distributions of the supernatant at steady-state for Pu⁴⁺ solubility experiments from over- and undersaturation in UE-25p #1 groundwater at pH 6 and 60°C. The solutions were filtered through 4.1 nm filters.



Plutonium Oxidation State Distributions in UE-25p#1 Water from Over- and Undersaturation at 60° C and pH 7

Figure 13. Plutonium oxidation state distributions of the supernatant at steady-state for Pu^{4+} solubility experiments from over- and undersaturation in UE-25p #1 groundwater at pH 7 and 60°C. The solutions were filtered through 4.1 nm filters.



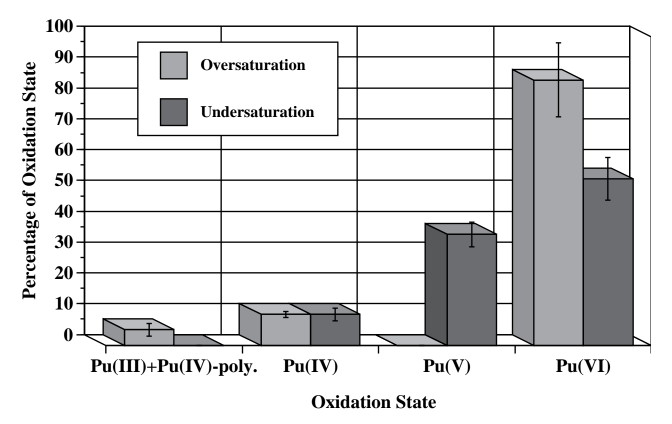


Figure 14. Plutonium oxidation state distributions of the supernatant at steady-state for Pu⁴⁺ solubility experiments from over- and undersaturation in UE-25p #1 groundwater at pH 8.5 and 60°C. The solutions were filtered through 4.1 nm filters.

5.2.3. Identification of Solids

The plutonium precipitates from both over- and undersaturation solutions were collected by centrifugation, washed with a small amount of CO_2 -free water, and dried with a jet of argon. All precipitates from the oversaturation experiments were a light brown-slightly greenish color. All precipitates from the undersaturation experiments were a very light brown color. Crystal lattice d-spacings and relative intensities of the x-ray powder diffraction patterns from the precipitates are listed in Table XIX.

The plutonium solid from the oversaturation experiment at pH 6 showed only a diffuse band indicating a low degree of crystallinity, which may be indicative of Pu(IV)-polymer. The powder pattern from the solid obtained in the pH 7 experiment showed the same diffuse band plus one weak line at 3.03 Å. Again, this shows a very low degree of crystallinity and is consistent with results for polymeric Pu(IV). The solid from the pH 8.5 experiment at 60°C produced a powder pattern that contained 7 diffraction lines. All of the diffraction lines were weak and trace lines with the exception of one, at a d-spacing of 3.26 Å. This powder pattern was compared to patterns of crystalline PuO₂⁴¹, PuO₃•0.8H₂O⁴², KPuO₂CO₃⁴³, NH₄PuO₂CO₃⁴³, and PuO₂CO₃⁴⁴. No match was found among these patterns, so all we can conclude is that the solid obtained at pH 8.5 and 60°C is crystalline but cannot be identified.

The plutonium solids removed from the solubility cells after completion of the undersaturation experiment are somewhat different. The pH 6 solid gave five distinct diffraction lines, all at least weak in intensity, and the pH 7 solid gave six distinct lines. Both of these results indicate that a more crystalline solid formed. The pH 8.5 solid gave six lines in its diffraction pattern. These lines were completely different from those obtained from the pH 8.5 oversaturation solid. All three solids where compared to patterns of crystalline PuO₂⁴¹, PuO₃•0.8H₂O⁴², KPuO₂CO₃⁴³, NH₄PuO₂CO₃⁴³, and PuO₂CO₃⁴⁴. No match was found among these patterns, so all we can conclude is that the solids are crystalline but not yet identified. The solids obtained from the undersaturation experiments gave diffraction patterns that matched each other very well and it appears that a similar solid was formed in at all three pH's.

In summary, the solids formed from oversaturation at pH 6 and 7 appear to be amorphous Pu(IV)-polymer; whereas, the solids produced from oversaturation at pH 8.5 and from undersaturation at all pH's are crystalline but unidentified.

pH 7 (ov	ersat.)	pH 8		pH		pН	7	pH 8	.5
		(overs	sat.)	(undersat.)		(undersat.)		(undersat.)	
<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī	<u>d(Å)</u>	Ī
		3.40	w+						
		3.26	S-						
3.03	W	3.04	t			3.06	w		
				2.74	w	2.77	t	2.81	t
		2.71	t						
								2.65	w-
		2.52	t						
								2.31	t
		2.03	w						
		1.98	w	1.99	s	1.99	s	1.99	s+
				1.95	w	1.95	w		
				1.74	m	1.73	m	1.73	m-
				1.25	w	1.25	m-	1.24	w

Table XIX.X-ray powder diffraction patterns of plutonium solid phases from
oversaturation in UE-25p #1 groundwater at 60°C and pH 6, 7, and 8.5.

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

5.3. Americium

We used non-radioactive neodymium in place of americium to minimize radiation-induced degradation of the solubility cell. Neodymium is chemically similar to americium.⁴⁵ It has an ionic radius of 0.983 Å that is very close to that of 0.975 Å for trivalent americium.⁴⁶ The neodymium was spiked with a small amount of ²⁴¹Am to facilitate sample counting using the 59.54-keV photo-peak. The use of the neodymium spiked with ²⁴¹Am tracer reduced the alpharadiation to a fraction of the radiation that would have been present if we had used pure ²⁴³Am. The mole ratio (([Am] + [Nd])/[Am]) used in these solubility experiments was 113.7:1.

We confirmed in an earlier study that neodymium is a good stand-in element for americium.¹ The differences between the solubilities of Nd/²⁴¹Am and pure ²⁴³Am were insignificant at each of the studied pH values. This validates our solubility tests in UE-25p #1 groundwater at 60°C where we used neodymium mixed with trace amounts of ²⁴¹Am as a substitute for ²⁴³Am.

5.3.1. Solubility. Results of the solubility studies are shown in Figure 15. The steady-state concentrations and the solutions' Eh values are given in Table XX. Concentration profiles as a function of equilibration time and pH for both the over- and undersaturation experiments are shown in Figure 16. Individual measurements are listed in Appendix C. Results for the americium/neodymium solubility experiment from oversaturation at pH 7 are incomplete because the experiment was stopped early due to equipment failure. The test cell, equipped with a faulty O-ring, allowed the experimental solution to evaporate. Steady-state may not have been reached, and the solution concentration reported in Table XX was determined from only the last two samplings. However, the steady-state concentration values from the undersaturation experiment are nearly identical to the steady-state concentration values from the oversaturation experiment, confirming that the solution was very near steady-state.

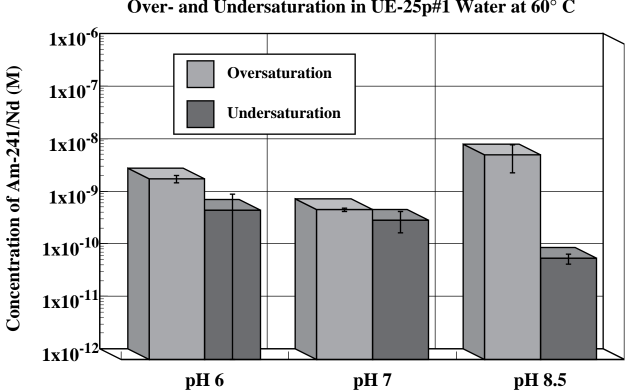
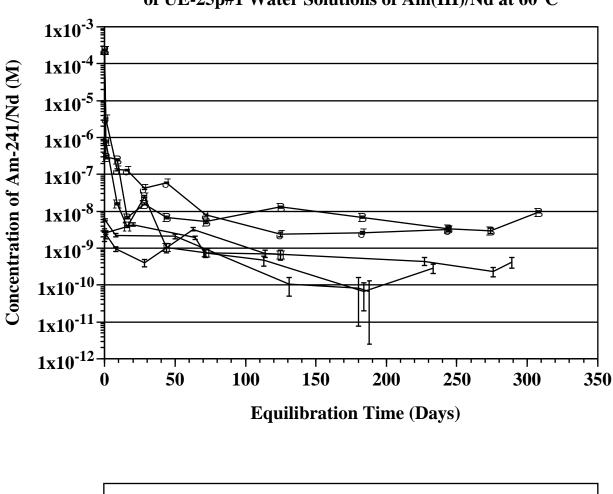


Figure 15. Results for ${}^{241}Am^{3+}/Nd^{3+}$ solubility experiments from over- and undersaturation in UE-25p #1 groundwater at 60°C as a function of pH.

Table XX.	Comparison	of	steady-state	solution	concentrations	and	Eh	for	americium/
	neodymium fr	rom	over- and u	ndersatura	ation in UE-25)#1 v	vater	at 60	°C.

	Americium(III)/Neodymium										
pН	Concentr	ation (M)	Eh (mV vs. NHE)								
	Oversaturation	Undersaturation	Oversaturation	Undersaturation							
6	$(2.7 \pm 0.4) \ge 10^{-9}$	$(7.0 \pm 7.3) \ge 10^{-10}$	(370 ± 15)	(394 ± 15)							
7	$(7.1 \pm 0.5) \ge 10^{-10}$	$(4.6 \pm 2.0) \ge 10^{-10}$	(NA)	(430 ± 15)							
8.5	$(7.8 \pm 4.3) \ge 10^{-9}$	$(8.4 \pm 1.8) \ge 10^{-11}$	(220 ± 15)	(215 ± 15)							

Americium(III)/Neodymium Solubility Experiments from Over- and Undersaturation in UE-25p#1 Water at 60° C



Approach to Equilibrium from Over- and Undersaturation of UE-25p#1 Water Solutions of Am(III)/Nd at 60°C

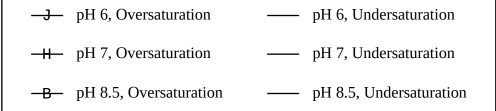


Figure 16. Solution concentrations of 241 Am/Nd in contact with precipitate obtained from supersaturation of UE-25p #1 groundwater at 60°C as a function of time. pH 6.0 \pm 0.1 (closed circles), pH 7.0 \pm 0.1 (closed triangles), and pH 8.5 \pm 0.1 (closed squares). The americium/neodymium was added initially (day 0) as 241 Am³⁺/Nd³⁺; initial (241 Am³⁺ + Nd³⁺) concentrations were 2.3 x 10⁻⁴ M (pH 6), 2.3 x 10⁻⁴ M (pH 7), and 2.3 x 10⁻⁴ M (pH 8.5). Undersaturation experiments at pH 6.0 (open circles), pH 7.0 (open triangles), and pH 8.5 (open squares) were started with precipitates obtained in the supersaturation experiments at their respective pH values.

Results for americium/neodymium filtration experiments have been given elsewhere.³² The filtration experiments are described in Section 4.5. Filtration experiments on day one of the 60°C oversaturation experiments at pH 6, 7, and 8.5 required 500, 1500, and 2500 μ L, respectively, as preconditioning volumes. Toward the end of the 60°C oversaturation experiments, as concentrations dropped from 10⁻⁸ to 10⁻¹⁰ M, we routinely used 2500 μ L as a presaturation volume for all experiments to ensure adequate saturation of the filters before sampling. We repeated the filtration experiments for the undersaturation cells and confirmed that indeed 2500 μ L was a sufficient preconditioning volume to use over the time of the entire solubility experiment.

In summary, the solubility of americium/neodymium from over- and undersaturation at pH 6 and 7 agreed within the error of the measurements. The pH 8.5 experiment from undersaturation was nearly two orders of magnitude less soluble than its oversaturation counterpart. This result shows that the steady-state condition may not have been reached in the oversaturation cell at pH 8.5. It appears that the americium/neodymium concentration from the first few days of the undersaturation experiment continued at the level of the final assays from the oversaturation experiment. In time, the concentration dropped by two orders of magnitude and a new steady-state condition was reached. The solubility of americium/neodymium showed no trend with pH in the oversaturation experiment, but showed decreasing concentration with increasing pH in the undersaturation experiment. If we assume that steady-state was not reached for the pH 8.5 cell until after day 125 of the undersaturation experiment, then the dependence of the concentration on the pH is confirmed.

5.3.2. Speciation. Speciation measurements could not be carried out because of the low solution concentrations. The trivalent neodymium is expected, under these experimental conditions, not to change its oxidation state. In determining whether the ²⁴¹Am tracer undergoes a change in oxidation state, we used extractions with 0.5 M thenoyltrifluoroacetone (TTA) or 0.025 M 1-phenyl-3-methyl-4-benzoyl-pyrazolin-5-one (PMBP) at pH 0 and coprecipitations with LaF₃. TTA or PMBP extracts the oxidation state IV and leaves the oxidation states III, V, and VI in the aqueous solution. The lanthanum fluoride (with holding oxidant) coprecipitates the oxidation states III and IV, and leaves the V and VI in the supernatant. As could be expected for the experimental conditions, we found no higher americium oxidation states. The trivalent americium did not change its oxidation state.

5.3.3. *Identification of Solids.* The neodymium/americium-241 precipitates found in both experiments were collected by centrifugation, washed with a small amount of CO₂-free water,

dried with an argon jet, and analyzed by x-ray powder diffraction. The crystal lattice d-spacings and relative intensities of all of the solids obtained are listed in Table XXI.

The solids from oversaturation were analyzed in an earlier report.³² They were found to be mixtures of Nd₂(CO₃)₃ • 2 H₂O and orthorhombic NdOHCO₃ in the case of the pH 7 and 8.5 solids. The pH 6 solid yielded very few lines and was considered to be "possibly a mixture of Nd₂(CO₃)₃ • 2 H₂O and hexagonal NdOHCO₃". It was shown in the literature that americium hydroxycarbonates are isostructural to the analogous neodymium hydroxycarbonates.45,47,48 The pH 6 undersaturation solid produced many more diffraction lines than the solid from oversaturation. With these additional lines it appears that the solid formed is a mixture of Nd₂(CO₃)₃ • 2 H₂O and orthorhombic NdOHCO₃, as was found in the pH 7 and pH 8.5 oversaturation solids. The pH 7 undersaturation solid yielded fewer lines in its diffraction pattern than the solid obtained from oversaturation. However, the lines that are present fit well with both the pH 7 and pH 8.5 oversaturation solids. Again, it appears to be a mixture of $Nd_2(CO_3)_3 \cdot 2$ H₂O and orthorhombic NdOHCO₃. The pH 8.5 undersaturation solid fits well with the pH 8.5 oversaturation solid. The solid from undersaturation has fewer trace lines probably due to shorter exposure times. This solid fits well with the $Nd_2(CO_3)_3 \cdot 2 H_2O^{45}$ solid, but still has some lines in common with the orthorhombic form of NdOHCO₃.⁴⁸ A comparison of all solids obtained in the undersaturation experiments with the reference patterns of: 1.) $Nd_2(CO_3)_3 \cdot 2H_2O^{45}$; and 2.) orthorhombic NdOHCO₃⁴⁸ are shown in Table XXII.

With regard to literature reference patterns containing diffraction lines with d-spacings larger than 6 Å, we have found that with the published powder patterns for hydrated neptunium solids, strong lines with large d-spacings were influenced by varying hydrate content in the solid. If the solids from the Am/Nd experiments contained Nd₂(CO₃)₃ with a varying hydrate content and the diffraction characteristic of hydrate water behaves as it does in neptunium solids, then we may in fact see some diffraction lines at d-spacings greater than 10-12 Å. The only conclusion we can draw at this point regarding the solids produced is that they appear to be mixtures of orthorhombic NdOHCO₃ and Nd₂(CO₃)₃ • 2 H₂O with possibly varying water content.

In summary, the UE-25p #1 solubility experiments at all pH's appear to have produced a mixture of orthorhombic NdOHCO₃ and Nd₂(CO₃)₃•2H₂O with possibly varying water content.

		H 0 7		11.7	pII 9 5	
pH 6	pH 7	pH 8.5	pH 6 Undersat.	pH 7	pH 8.5 Undersat.	
Oversat.	Oversat.	Översat.	Undersat.	Undersat.	Undersat.	
$\underline{d}(\underline{\mathring{A}}) = \underline{I}^{a}$	$\underline{d}(\underline{\mathring{A}}) = \underline{I}^{a}$	<u>d(Å) I</u> ^a	$\underline{d}(\underline{A}) = \underline{I}^{a}$	$\underline{d}(\underline{A}) = \underline{I}^{a}$	$\underline{d}(\underline{A}) = \underline{I}^{a}$	
	9.38 s					
	4.88 m-					
	4.71 m+					
		4.59 m				
	4.32 m	4.28 s				
			4.15 m-			
	3.93 m					
			3.51 w			
	3.37 t		3.32 t			
	3.24 m	3.25 w	3.25 w	3.24 w-	3.24 m	
	3.19 t					
	3.13 m-					
3.06 w	3.07 m-		3.06 s-		3.09 t	
		3.04 m-			3.04 t	
					2.94 t	
2.86 s		2.83 m	2.84 s	2.83 w+	2.83 t	
					2.73 t	
	2.67 w					
		2.61 t	2.59 t			
	2.55 w					
		2.48 t				
	2.43 w-		2.42 t			
	2.36 w					
		2.32 t				
	2.26 t		2.27 t			
	2.17 w-	2.15 t	2.16 t			
	2.12 t		2.13 w+			
	2.07 w-	2.05 t				
1.99 m	1.97 w-	1.99 w-	1.99 w+	1.98 s	1.99 s	
	1.94 t				1.93 t	
	1.91 t					
	1.88 t		1.87 t		1.87 t	
			1.86 t			
					1.79 t	
		1.72 t	1.72 w-	1.72 w+	1.72 t	
					1.65 t	
	1.62 t				1.63 t	
	1.60 t				· · · · ·	
	1.57 t					
	1.52 t	1.53 m-			1.53 w	
	1.47 t				1.00 17	
	1.44 t					
	1.1.7 L	1.31 t	1.32 t		1.32 t	
			1.02 (1.24 t	
(a) Relativ		lly estimated: vs = very stroi		1 1' 1		

Table XXI.X-ray powder diffraction patterns of Nd/241Am solid phases in UE-25p #1
groundwater from over- and undersaturation at 60°C and pH 6, 7, and 8.5.

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.

Table XXII.	X-ray powder diffraction patterns of $Nd/^{241}Am$ solid phases in UE-25p #1 groundwater from undersaturation at 60°C and pH 6, 7, and 8.5 compared with
	reference patterns of: 1.) Nd ₂ (CO ₃) ₃ •2H ₂ O; 2.) orthorhombic NdOHCO ₃ . ^{45,48}

pH 6, 6	50°C	pH 7, 0	50°C	pH 8.5,	60°C		Ref. a	#1	Ref. #2	2 ^(b)
<u>d(Å)</u>	<u>I</u> ^a	<u>d(Å)</u>	<u>I</u> ^a	<u>d(Å)</u>	<u>I</u> ^a		<u>d(Å)</u>	<u>I</u> ^a	<u>d(Å)</u>	<u>I</u> ^a
							7.56	m		
							5.68	s	5 50	
							4.67	s	5.50	m-
							4.07	8	4.28	m
									4.24	vs
4.15	m-									
							3.93	s		
							3.83	w		
									3.68	t
2.51							3.62	m	3.65	m
3.51	w								2.22	
3.32 3.25	t w	3.24	w-	3.24	m				3.32	m
3.06	w s-	5.24	vv-	3.24	t		3.09	s		
5.00	5			3.04	t		3.02	s		
				2.94	t		2.98	w	2.94	t
									2.91	m
2.84	s	2.83	w+	2.83	t		2.87	w		
				2.73	t		2.75	w	2.75	m
2.59	t								2.63	m
							2.58	s	2.49	
2.42	t								2.48	s
2.42	ι								2.40 2.32	t w
									2.32	m-
2.27	t						2.25	w	2.51	
2.16	t						2.17	w	2.14	w
2.13	w+								2.12	s
									2.10	t
							2.08	w	2.05	w
1.00		1.00		1.00			2.02	m	2.03	w
1.99	w+	1.98	s	1.99	s		1.98	w	1.96	w
1.87	t			1.93 1.87	t t		1.87	m	1.93 1.88	w t
1.86	t			1.07	ť		1.83	m	1.83	s
				1.79	t		1.78	m	1.81	w
1.72	w-	1.72	w+	1.72	t		1.74	w	1.72	w
									1.69	w
									1.68	t
				1.65	t				1.66	t
				1.63	t				1.62	w
									1.60 1.58	w t
							1.56	m	1.58	w
				1.53	w				1.54	m
									1.52	w
									1.50	t
									1.47	w
									1.46	w
									1.44	t
									1.42 1.37	t t
1.32	t			1.32	t				1.37	t w
1.52	i			1.24	t				1.51	
L		L				I j				

(a) Relative intensities visually estimated: vs = very strong, s = strong, m = medium, w = weak, t = trace.
(b) Ref. #3 contains 7 more "weak" and "trace" lines with d-spacings smaller than 1.31 Å that are not listed.

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APPENDIX A

RESULTS OF NEPTUNIUM SOLUBILITY EXPERIMENTS IN UE-25p #1 WATER

Sample I.D.	Days	рН	Concentration (M)
Initial amount added	0	_	$(5.6 \pm 0.2) \ge 10^{-3}$
1-1A6U6	1	6.04	$(2.32 \pm 0.11) \ge 10^{-3}$
1-2A2U6	10	5.98	$(2.62 \pm 0.11) \ge 10^{-3}$
1-3A2U6	16	6.07	$(2.67 \pm 0.12) \ge 10^{-3}$
1-4A1U6	31	6.04	$(2.60 \pm 0.11) \ge 10^{-3}$
1-5A1U6	43	6.02	$(2.65 \pm 0.09) \ge 10^{-3}$
1-6A1U6	59	5.99	$(2.59 \pm 0.10) \ge 10^{-3}$
1-7A2U6	87	6.02	$(2.61 \pm 0.10) \ge 10^{-3}$
1-8A1U6	101	6.02	$(2.27 \pm 0.09) \ge 10^{-3}$
1-9A1U6	111	5.97	$(2.43 \pm 0.09) \ge 10^{-3}$
1-10A1U6	143	6.04	$(2.22 \pm 0.07) \ge 10^{-3}$
1-11A1U6	170	6.01	$(2.28 \pm 0.08) \times 10^{-3}$

Appendix IA. Results of Neptunium solubility experiments from oversaturation in UE-25p #1 water at 60°C and pH 6.0

average 1-1A6U6 through 1-11A1U6

 $(2.48 \pm 0.18) \ge 10^{-3}$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(1.5 \pm 0.1) \ge 10^{-3}$
2-1A6U6	1	7.02	$(6.66 \pm 0.22) \ge 10^{-5}$
2-2A2U6	10	6.96	$(3.35 \pm 0.12) \ge 10^{-5}$
2-3A2U6	16	7.09	$(2.98 \pm 0.11) \ge 10^{-5}$
2-4A1U6	31	7.10	$(2.90 \pm 0.12) \ge 10^{-5}$
2-5A1U6	43	7.10	$(3.09 \pm 0.11) \ge 10^{-5}$
2-6A1U6	59	6.88	$(3.62 \pm 0.14) \ge 10^{-5}$
2-7A2U6	87	7.09	$(3.54 \pm 0.16) \ge 10^{-5}$
2-8A1U6	101	7.03	$(2.26 \pm 0.14) \ge 10^{-5}$
2-9A1U6	111	6.97	$(5.07 \pm 0.22) \ge 10^{-5}$
2-10A1U6	143	7.07	$(2.79 \pm 0.09) \ge 10^{-5}$
2-11A1U6	170	7.07	$(5.14 \pm 0.28) \ge 10^{-5}$
2-12A1U6	183	7.06	$(2.11 \pm 0.11) \ge 10^{-5}$

Appendix IIA. Results of Neptunium solubility experiments from oversaturation in UE-25p #1 water at 60° C and pH 7.0

average 2-2A2U6 through 2-12A1U6

 $(3.35 \pm 0.99) \ge 10^{-5}$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(1.5 \pm 0.1) \ge 10^{-3}$
3-1A6U6	1	8.45	$(9.22 \pm 0.44) \ge 10^{-6}$
3-2A2U6	10	8.34	$(8.43 \pm 0.31) \ge 10^{-6}$
3-3A2U6	16	8.31	$(8.43 \pm 0.34) \ge 10^{-6}$
3-4A1U6	31	8.32	$(9.17 \pm 0.39) \ge 10^{-6}$
3-5A1U6	43	8.54	$(1.15 \pm 0.05) \ge 10^{-5}$
3-6A1U6	59	8.40	$(1.23 \pm 0.06) \ge 10^{-5}$
3-7A6U6	87	8.50	$(1.26 \pm 0.06) \ge 10^{-5}$
3-8A1U6	101	8.50	$(1.55 \pm 0.09) \ge 10^{-5}$
3-9A1U6	111	8.37	$(1.50 \pm 0.09) \ge 10^{-5}$
3-10A1U6	143	8.45	$(2.35 \pm 0.13) \ge 10^{-5}$
3-11A1U6	155	8.50	$(2.68 \pm 0.09) \ge 10^{-5}$

Appendix IIIA. Results of Neptunium solubility experiments from oversaturation in UE-25p #1 water at 60° C and pH 8.5

assay 3-11A1U6

 $(2.68 \pm 0.09) \ge 10^{-5}$

Sample I.D.	Days	pH	Concentration (M)
Initial solid added	0	6.00	-
1-12A1U6	0.05	6.35	$(2.28 \pm 0.08) \ge 10^{-3}$
1-13A1U6	1	6.03	$(4.15 \pm 0.22) \ge 10^{-4}$
1-14A1U6	6	5.92	$(6.18 \pm 0.29) \ge 10^{-4}$
1-15A1U6	28	6.00	$(1.14 \pm 0.03) \ge 10^{-3}$
1-16A1U6	55	6.09	$(1.59 \pm 0.06) \ge 10^{-3}$
1-17A1U6	85	5.97	$(2.16 \pm 0.07) \ge 10^{-3}$
1-18A1U6	124		$(2.42 \pm 0.08) \ge 10^{-3}$
1-19A1U6	168	5.64	$(3.44 \pm 0.10) \ge 10^{-3}$
1-20A1U6	197	5.93	$(2.40 \pm 0.10) \ge 10^{-4}$
1-21A1U6	204	6.98	$(3.18 \pm 0.20) \ge 10^{-4}$
1-22A1U6	210	5.96	$(4.23 \pm 0.24) \ge 10^{-4}$
1-23A1U6*	1	6.12	$(2.27 \pm 0.16) \ge 10^{-4}$
1-24A1U6 [*]	7	6.01	$(2.49 \pm 0.13) \ge 10^{-4}$
1-25A1U6*	70	5.92	$(4.85 \pm 0.14) \ge 10^{-4}$
1-26A1U6 [*]	175	5.98	$(4.02 \pm 0.13) \ge 10^{-4}$
1-27A1U6*	197	6.02	$(4.48 \pm 0.19) \ge 10^{-4}$
1-28A1U6*	200	6.31	$(2.81 \pm 0.08) \ge 10^{-4}$

Appendix IVA. Results of Neptunium solubility experiments from undersaturation in UE-25p#1 water at 60° C and pH 6.0

assay 1-19A1U6

 $(3.44 \pm 0.10) \ge 10^{-3}$

(*) = Assays taken after cell was restarted

Sample I.D.	Days	pH	Concentration (M)
Initial solid added	0	_	-
2-13A1U6	0.02	7.28	$(1.08 \pm 0.07) \ge 10^{-5}$
2-14A1U6	1	7.12	$(1.65 \pm 0.10) \ge 10^{-5}$
2-15A1U6	8	7.08	$(2.21 \pm 0.10) \ge 10^{-5}$
2-16A1U6	15	7.05	$(3.39 \pm 0.16) \ge 10^{-5}$
2-17A1U6	42	6.93	$(6.39 \pm 0.34) \ge 10^{-5}$
2-18A1U6	72	7.15	$(4.21 \pm 0.18) \ge 10^{-5}$
2-19A1U6	111		$(9.32 \pm 0.32) \ge 10^{-5}$
2-20A1U6	154	7.08	$(1.04 \pm 0.03) \ge 10^{-4}$
2-21A1U6	183	7.36	$(9.71 \pm 0.38) \ge 10^{-5}$
2-22A1U6	188	7.00	$(1.11 \pm 0.07) \ge 10^{-4}$

Appendix VA. Results of Neptunium solubility experiments from undersaturation in UE-25p#1 water at 60° C and pH 7.0

average 2-19A1U6 through 2-22A1U6

 $(1.01 \pm 0.07) \times 10^{-4}$

Sample I.D.	Days	pН	Concentration (M)
Initial solid added	0	8.50	-
3-12A1U6	0.16	8.35	$(4.79 \pm 0.26) \ge 10^{-6}$
3-13A1U6	1	8.45	$(5.03 \pm 0.21) \ge 10^{-6}$
3-14A1U6	6	8.49	$(5.20 \pm 0.28) \ge 10^{-6}$
3-15A1U6	9	8.50	$(5.60 \pm 0.24) \ge 10^{-6}$
3-16A1U6	15	8.46	$(5.68 \pm 0.21) \ge 10^{-5}$
3-17A1U6	43	8.49	$(1.05 \pm 0.06) \ge 10^{-5}$
3-18A1U6	70	8.48	$(1.19 \pm 0.06) \ge 10^{-5}$
3-19A1U6	100	8.48	$(1.52 \pm 0.09) \ge 10^{-5}$
3-20A1U6	139		$(1.36 \pm 0.08) \ge 10^{-5}$
3-21A1U6	183	8.65	$(2.80 \pm 0.23) \ge 10^{-5}$
3-22A1U6	212	8.39	$(3.66 \pm 0.11) \ge 10^{-5}$
3-23A1U6	258	8.53	$(6.03 \pm 0.48) \ge 10^{-5}$
3-25A1U6	336	8.50	$(2.63 \pm 0.11) \ge 10^{-5}$

Appendix VIA. Results of Neptunium solubility experiments from undersaturation in UE-25p#1 water at 60° C and pH 8.5

average 3-21A1U6 through 3-25A1U6

 $(3.78 \pm 1.57) \ge 10^{-5}$

APPENDIX B

RESULTS OF PLUTONIUM SOLUBILITY EXPERIMENTS IN UE-25p #1 WATER

Sample I.D.	Days	рН	Concentration (M)
Initial amount added	0	_	$(2.2 \pm 0.2) \ge 10^{-4}$
4-1A6U6	1	5.87	$(6.65 \pm 0.76) \ge 10^{-9}$
4-2A1U6	6	5.85	$(2.18 \pm 0.14) \ge 10^{-7}$
4-3A1U6	9	5.67	$(6.64 \pm 0.42) \ge 10^{-7}$
4-4A1U6	21	6.04	$(1.41 \pm 0.09) \ge 10^{-6}$
4-5A1U6	37	5.96	$(1.90 \pm 0.12) \ge 10^{-6}$
4-6A2U6	65	5.90	$(8.97 \pm 0.56) \ge 10^{-7}$
4-7A1U6	79	6.90	$(9.04 \pm 0.57) \ge 10^{-7}$
4-8A1U6	121	6.04	$(7.11 \pm 0.45) \ge 10^{-7}$
4-9A1U6	176	6.16	$(2.59 \pm 0.22) \ge 10^{-7}$
4-10A1U6	236	6.08	$(7.22 \pm 0.63) \ge 10^{-8}$
4-11A1U6	258	5.90	$(9.42 \pm 0.84) \ge 10^{-8}$
4-12A1U6	286	5.93	$(9.91 \pm 0.89) \ge 10^{-8}$

Appendix IB. Results of Plutonium solubility experiments from oversaturation in UE-25p #1 water at 60° C and pH 6.0

average 4-10A1U6 through 4-12A1U6

 $(8.85 \pm 1.43) \ge 10^{-8}$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(2.2 \pm 0.2) \ge 10^{-4}$
5-1A6U6	1	7.33	$(5.13 \pm 0.37) \ge 10^{-8}$
5-2A1U6	6	7.29	$(1.96 \pm 0.12) \ge 10^{-7}$
5-3A1U6	9	7.35	$(2.62 \pm 0.14) \ge 10^{-7}$
5-4A1U6	21	7.29	$(3.45 \pm 0.22) \ge 10^{-7}$
5-5A1U6	37	7.31	$(4.39 \pm 0.28) \ge 10^{-7}$
5-6A2U6	65	7.41	$(2.88 \pm 0.18) \ge 10^{-7}$
5-7A1U6	79	7.64	$(2.43 \pm 0.15) \ge 10^{-7}$
5-8A1U6	121	7.01	$(2.26 \pm 0.14) \ge 10^{-7}$
5-9A1U6	176	7.25	$(9.32 \pm 0.80) \ge 10^{-8}$
5-10A1U6	236	7.36	$(8.13 \pm 0.71) \ge 10^{-8}$
5-11A1U6	274	6.88	$(1.07 \pm 0.10) \ge 10^{-7}$
5-12A1U6	287	7.09	$(8.07 \pm 0.73) \ge 10^{-8}$

Appendix IIB. Results of Plutonium solubility experiments from oversaturation in UE-25p #1 water at 60° C and pH 7.0

average 5-9A1U6 through 5-12A1U6

 $(9.06 \pm 1.24) \ge 10^{-8}$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(2.0 \pm 0.2) \ge 10^{-4}$
6-1A6U6	1	7.90	$(3.40 \pm 0.22) \ge 10^{-7}$
6-2A1U6	6	7.98	$(3.89 \pm 0.24) \ge 10^{-7}$
6-3A1U6	9	8.23	$(4.01 \pm 0.25) \ge 10^{-7}$
6-4A1U6	21	8.54	$(4.21 \pm 0.26) \ge 10^{-7}$
6-5A1U6	37	8.04	$(4.84 \pm 0.30) \ge 10^{-7}$
6-6A2U6	65	8.49	$(5.45 \pm 0.34) \ge 10^{-7}$
6-7A1U6	79	8.42	$(8.59 \pm 0.54) \ge 10^{-7}$
6-8A1U6	121	8.44	$(2.57 \pm 0.16) \ge 10^{-6}$
6-9A1U6	176	8.58	$(2.02 \pm 0.17) \ge 10^{-6}$
6-10A1U6	236	8.51	$(1.42 \pm 0.12) \ge 10^{-6}$
6-11A1U6	279	8.50	$(7.00 \pm 0.63) \ge 10^{-7}$
6-12A1U6	294	8.43	$(7.15 \pm 0.64) \ge 10^{-7}$

Appendix IIIB. Results of Plutonium solubility experiments from oversaturation in UE-25p #1 water at 60° C and pH 8.5

average 6-6A2U6 through 6-12A1U6

 $(1.26 \pm 0.78) \ge 10^{-6}$

Sample I.D.	Days	pH	Concentration (M)
Initial solid added	0	6.00	-
4-13A1U6	0	6.27	$(2.91 \pm 0.27) \ge 10^{-8}$
4-14A1U6	1	6.38	$(7.09 \pm 0.64) \ge 10^{-8}$
4-15A1U6	15	6.14	$(1.53 \pm 0.13) \ge 10^{-7}$
4-16A1U6	30	6.24	$(1.10 \pm 0.09) \ge 10^{-7}$
4-17A1U6	73	5.98	$(1.10 \pm 0.09) \ge 10^{-7}$
4-18A1U6	149	6.03	$(7.07 \pm 0.61) \ge 10^{-8}$
4-19A1U6	156	6.08	$(5.74 \pm 0.50) \ge 10^{-8}$

Appendix IVB. Results of Plutonium solubility experiments from undersaturation in UE-25p#1 water at 60° C and pH 6.0

average 4-14A1U6 through 4-19A1U6

 $(9.53 \pm 3.58) \ge 10^{-8}$

Sample I.D.	Days	pH	Concentration (M)
Initial solid added	0	7.00	-
5-13A1U6	0	7.31	$(4.06 \pm 0.40) \ge 10^{-8}$
5-14A1U6	1	7.40	$(5.51 \pm 0.50) \ge 10^{-8}$
5-15A1U6	13	7.24	$(9.22 \pm 0.81) \ge 10^{-8}$
5-16A1U6	28	7.20	$(1.03 \pm 0.09) \ge 10^{-7}$
5-17A1U6	71	6.95	$(1.04 \pm 0.09) \text{ x } 10^{-7}$
5-18A1U6	147	7.40	$(4.26 \pm 0.37) \ge 10^{-8}$
5-19A1U6	153	6.86	$(3.71 \pm 0.33) \ge 10^{-8}$

Appendix VB. Results of Plutonium solubility experiments from undersaturation in UE-25p#1 water at 60° C and pH 7.0

average 5-13A1U6 through 5-19A1U6

 $(6.78 \pm 3.06) \ge 10^{-8}$

Sample I.D.	Days	pН	Concentration (M)
Initial solid added	0	8.50	-
6-13A1U6	0	8.43	$(2.50 \pm 0.26) \ge 10^{-8}$
6-14A1U6	1	8.37	$(5.30 \pm 0.48) \ge 10^{-8}$
6-15A1U6	8	8.34	$(6.24 \pm 0.55) \ge 10^{-8}$
6-16A1U6	22	8.47	$(6.94 \pm 0.60) \ge 10^{-8}$
6-17A1U6	65	8.44	$(9.28 \pm 0.80) \ge 10^{-8}$
6-18A1U6	141	8.73	$(1.78 \pm 0.15) \ge 10^{-7}$
6-20A1U6	156		$(1.00 \pm 0.08) \ge 10^{-7}$
6-22A1U6	218	8.54	$(1.08 \pm 0.10) \ge 10^{-7}$
6-23A1U6	237	8.52	$(6.96 \pm 0.62) \ge 10^{-8}$

Appendix VIB. Results of Plutonium solubility experiments from undersaturation in UE-25p#1 water at 60° C and pH 8.5

average 6-14A2U6 through 6-23A1U6

 $(9.17 \pm 3.99) \ge 10^{-8}$

APPENDIX C

RESULTS OF AMERICIUM/NEODYMIUM SOLUBILITY EXPERIMENTS IN UE- $25p\ \#1\ WATER$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(2.3 \pm 0.1) \times 10^{-4}$
7-1A6U6	1	6.05	$(3.18 \pm 0.19) \ge 10^{-6}$
7-2A1U6	9	5.99	$(1.33 \pm 0.07) \ge 10^{-7}$
7-3A1U6	16	5.89	$(1.30 \pm 0.07) \ge 10^{-7}$
7-4A1U6	28	6.01	$(4.20 \pm 0.23) \ge 10^{-8}$
7-5A1U6	44	6.00	$(5.94 \pm 0.33) \ge 10^{-8}$
7-6A6U6	72	6.05	$(7.88 \pm 0.64) \ge 10^{-9}$
7-7A1U6	125	6.06	$(2.37 \pm 0.36) \ge 10^{-9}$
7-8A1U6	183	5.90	$(2.59 \pm 0.20) \ge 10^{-9}$
7-9A1U6	244	6.57	$(3.18 \pm 0.40) \ge 10^{-9}$

Appendix IC. Results of Americium/Neodymium solubility experiments from oversaturation in UE-25p #1 water at 60°C and pH 6.0

average 7-7A1U6 through 7-9A1U6

 $(2.71 \pm 0.42) \times 10^{-9}$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(2.3 \pm 0.1) \ge 10^{-4}$
8-1A6U6	1	7.07	$(7.92 \pm 0.65) \ge 10^{-7}$
8-2A1U6	9	7.06	$(1.62 \pm 0.09) \ge 10^{-8}$
8-3A1U6	16	7.00	$(3.80 \pm 0.24) \ge 10^{-9}$
8-4A1U6	28	7.08	$(2.49 \pm 0.15) \ge 10^{-8}$
8-5A1U6	44	7.02	$(1.06 \pm 0.30) \ge 10^{-9}$
8-6A6U6	72	7.06	$(7.46 \pm 1.95) \text{ x10}^{-10}$
8-7A1U6	125	7.11	$(6.73 \pm 1.92) \text{ x10}^{-10}$

Appendix IIC. Results of Americium/Neodymium solubility experiments from oversaturation in UE-25p #1 water at 60°C and pH 7.0

average 8-6A6U6 through 8-7A1U6

 $(7.09 \pm 0.52) \ge 10^{-10}$

Sample I.D.	Days	pН	Concentration (M)
Initial amount added	0	_	$(2.3 \pm 0.1) \ge 10^{-4}$
9-1A6U6	1	7.78	$(2.96 \pm 0.36) \ge 10^{-7}$
9-2A1U6	9	8.38	$(2.49 \pm 0.13) \ge 10^{-7}$
9-3A1U6	16	8.64	$(6.66 \pm 0.42) \ge 10^{-9}$
9-4A1U6	28	8.39	$(1.58 \pm 0.09) \ge 10^{-8}$
9-5A1U6	44	8.42	$(6.89 \pm 0.59) \ge 10^{-9}$
9-6A6U6	72	8.51	$(5.19 \pm 0.44) \ge 10^{-9}$
9-7A1U6	125	8.46	$(1.30 \pm 0.09) \ge 10^{-8}$
9-8A1U6	183	8.53	$(6.89 \pm 0.52) \ge 10^{-9}$
9-9A1U6	244	8.61	$(3.41 \pm 0.40) \ge 10^{-9}$
9-10A1U6	274	8.56	$(2.99 \pm 0.47) \ge 10^{-9}$
9-11A1U6	308	8.54	$(9.60 \pm 0.71) \ge 10^{-9}$

Appendix IIIC. Results of Americium/Neodymium solubility experiments from oversaturation in UE-25p #1 water at 60°C and pH 8.5

average 9-3A1U6 through 9-11A1U6

 $(7.83 \pm 4.28) \times 10^{-9}$

Sample I.D.	Days	pH	Concentration (M)
Initial solid added	0	6.00	-
7-13A1U6	0	6.09	$(3.04 \pm 0.37) \ge 10^{-9}$
7-14A1U6	1	6.12	$(2.63 \pm 0.26) \ge 10^{-9}$
7-15A1U6	20	5.89	$(4.36 \pm 0.42) \ge 10^{-9}$
7-16A1U6	64	6.12	$(1.93 \pm 0.02) \ge 10^{-9}$
7-17A1U6	71	6.15	$(7.56 \pm 1.82) \ge 10^{-10}$
7-18A1U6	113	6.09	$(4.71 \pm 1.48) \ge 10^{-10}$
7-19A1U6	184	5.97	$(6.74 \pm 4.73) \ge 10^{-11}$
7-20A1U6	233	6.03	$(2.91 \pm 0.84) \ge 10^{-10}$

Appendix IVC. Results of Americium/Neodymium solubility experiments from undersaturation in UE-25p #1 water at 60°C and pH 6.0

average 7-16A1U6 through 7-20A1U6

 $(7.03 \pm 7.31) \ge 10^{-10}$

Sample I.D.	Days	pН	Concentration (M)
Initial solid added	0	7.00	-
8-9A1U6	0	7.38	$(2.10 \pm 0.58) \ge 10^{-9}$
8-10A1U6	1	7.32	$(2.36 \pm 0.59) \ge 10^{-9}$
8-11A1U6	8		$(9.49 \pm 1.21) \ge 10^{-10}$
8-12A1U6	28	7.19	$(4.07 \pm 0.90) \ge 10^{-10}$
8-13A1U6	63	6.79	$(3.26 \pm 0.32) \ge 10^{-9}$
8-14A1U6	114	7.18	$(7.20 \pm 1.59) \text{ x10}^{-10}$
8-17A1U6	227	7.14	$(4.43 \pm 1.06) \text{ x10}^{-10}$
8-18A1U6	276	7.04	$(2.37 \pm 0.68) \text{ x10}^{-10}$
8-19A1U6	289	6.93	$(4.19 \pm 1.28) \text{ x10}^{-10}$

Appendix VC. Results of Americium/Neodymium solubility experiments from undersaturation in UE-25p #1 water at 60°C and pH 7.0

average 8-14A1U6 through 8-19A1U6

 $(4.55 \pm 1.99) \ge 10^{-10}$

Sample I.D.	Days	pH	Concentration (M)
Initial solid added	0	8.50	-
9-13A1U6	0	8.43	$(5.81\pm 0.38) \ge 10^{-9}$
9-15A1U6	8	8.47	$(2.24 \pm 0.23) \ge 10^{-9}$
9-16A1U6	50	8.42	$(2.09 \pm 0.33) \ge 10^{-9}$
9-17A1U6	131	8.54	$(1.03 \pm 0.54) \ge 10^{-10}$
9-18A1U6	180	8.65	$(8.23 \pm 7.47) \ge 10^{-11}$
9-19A1U6	188	8.51	$(6.63 \pm 6.38) \ge 10^{-11}$

Appendix VIC. Results of Americium/Neodymium solubility experiments from undersaturation in UE-25p #1 water at 60°C and pH 8.5

average 9-17A1U6 through 9-19A1U6

 $(8.39 \pm 1.84) \ge 10^{-11}$

The data used to write this report are recorded in the following YMP Laboratory Record Books:

TWS-LBL-02-90-04, pp. 231 - 299

TWS-LBL-10-91-04, pp. 1 - 211

TWS-LBL-02-91-01, pp. 39-70

TWS-LBL-01-92-02, pp. 1 - 209

TWS-LBL-01-93-02, pp. 1- 77 (open)

TWS-LBL-01-93-03, pp. 1- 37 (open)

TWS-LBL-01-93-04, pp. 1- 79 (open)

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