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Radionuclide Release from Spent Fuel Under Geologic Disposal Conditions: An Overview of Experimental and Theoretical Work Through 1985

P. W. Reimus S. A. Simonson

April 1988

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RADIONUCLIDE RELEASE FROM SPENT FUEL UNDER GEOLOGIC DISPOSAL CONDITIONS: AN OVERVIEW OF EXPERIMENTAL AND THEORETICAL WORK THROUGH 1985

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Pacific Northwest Laboratory Richland, Washington 99352 · •

FOREWORD

This overview of existing experimental and theoretical work on radionuclide release from spent fuel was completed in November 1985. Thus, any work on this topic that was published after that time is not reviewed in this document. As a result of this review, an informal library of literature on radionuclide release from spent fuel was created at Pacific Northwest Laboratory. This library will be maintained by the Waste Package Program of the Salt Repository Project.

SUMMARY

This report presents an overview of experimental and theoretical work on radionuclide release from spent fuel and uranium dioxide (UO_2) under geologic disposal conditions. The purpose of the report is to provide a source book of information that can be used to develop models that describe radionuclide release from spent fuel waste packages. Modeling activities of this nature will be conducted within the Waste Package Program (WPP) of the Department of Energy's Salt Repository Project (SRP).

The topics discussed in this report include:

- experimental methods for investigating radionuclide release
- how results have been reported from radionuclide release experiments
- theoretical studies of UO_2 and actinide solubility
- results of experimental studies of radionuclide release from spent fuel and UO₂ (i.e., the effects of different variables on radionuclide release)
- characteristics of spent fuel pertinent to radionuclide release
- status of modeling of radionuclide release from spent fuel.

Appendix A presents tables of data from spent fuel radionuclide release experiments. These data have been digitized from graphs that appear in the literature. An annotated bibliography of literature on spent fuel characterization is provided in Appendix B.

Based on what has been learned in spent fuel and UO₂ radionuclide release studies, a complete model describing the release of radionuclides from spent fuel should include the following submodels:

 an instant release submodel describing the rate of release of soluble radionuclides (e.g., cesium-134, cesium-137, iodine-129, and possibly technetium-99) that have accumulated in the fuel-cladding gap and at the grain boundaries of the spent fuel (the first surfaces to come in contact with leachant)

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- a kinetic submodel describing the rate of degradation (i.e., oxidation and dissolution) of the spent fuel matrix, which describes the rate at which many radionuclides become unbound from the spent fuel and available for transport
- a post-release submodel, which describes the accumulation of radionuclides in solution, the formation of precipitates, colloids, and alteration phases, and the constraints on these processes
- a mass transfer submodel, which describes the transport of the radionuclides away from the spent fuel according to the concentration of radionuclides in solution.

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1.0 INTRODUCTION

The Waste Package Program (WPP) at Pacific Northwest Laboratory (PNL) is being conducted for the Department of Energy's Salt Repository Project (SRP) to provide technical support in assessing the performance of nuclear waste packages in a conceptual geologic repository for high-level waste. The WPP includes a number of experimental activities to investigate the degradation of proposed waste package components and associated behavior that pertain to licensing a repository. These activities address the problem of demonstrating that waste packages can meet the criteria imposed by regulations such as the Nuclear Regulatory Commission (NRC) regulation 10 CFR 60 and the proposed Environmental Protection Agency (EPA) regulation 40 CFR 191.

The WPP includes a modeling task, which is intended to provide a close link between waste package experimentation and SRP performance assessment activities. The objective of WPP modeling is to develop and validate models that describe the behavior of waste package components under conditions expected in a geologic repository. These models can be easily incorporated by the SRP into a code such as the Waste Package Performance Assessment (WAPPA) code.

The predicted behavior of spent fuel in a geologic repository, after all other waste package barriers have failed (i.e., spent fuel in contact with ground water or brine), is of particular interest because commercial spent fuel is expected to be the major form of nuclear waste disposed in a repository. The release of radionuclides from spent fuel must be predicted to show compliance with the regulatory criteria cited above.

This document is intended to be a source book of information for future modeling efforts to be conducted within the WPP in the area of radionuclide release from spent fuel. It is intended to benefit both modelers and experimentalists who are involved in the assessment of spent fuel as a waste form.

The review is divided into several chapters. The first chapter (Chapter 2.0) describes the experimental methods that have been used to measure radionuclide release rates from spent fuel and uranium dioxide (UO_2) . The second chapter (Chapter 3.0) describes the various ways that release and

release rates have been reported. A summary of all the radionuclide release studies that have been conducted on spent fuel, UO_2 , and simulated spent fuel under geologic disposal conditions is given in Chapter 4.0. This chapter contains tables that summarize the conditions and parameters studied in each of the experiments. Chapter 5.0 discusses theoretical studies of UO_2 dissolution and solubility, which are considered important because the reactions of spent fuel are dominated by the chemistry of UO_2 . Chapter 6.0 compares the experimental results of the studies discussed in Chapter 4.0 with the theoretical predictions from the studies in Chapter 5.0. Chapter 7.0 contains a brief discussion of the characteristics of spent fuel that are expected to be important to radionuclide release. Chapter 8.0 discusses the status of modeling of radionuclide release from spent fuel are made in Chapter 9.0. Two appendices are provided: an appendix of tabular spent fuel release data taken from the literature, and an annotated bibliography of spent fuel characterization.

2.0 EXPERIMENTAL METHODS FOR INVESTIGATING RADIONUCLIDE RELEASE

This chapter discusses the different types of test methods used in radionuclide release experiments. The test methods are divided into two major categories: non-accelerated and accelerated. Non-accelerated tests involve placing a waste form specimen in contact with a solution for a specified period of time without any interference from outside the system. As long as the waste form and solution remain in contact, they will approach a state of equilibrium with respect to each other. These tests are used to study dissolution kinetics, radionuclide release rates, solubility limits, and interactions of the waste form with other waste package barriers and/or the surrounding geology. In accelerated tests, an outside source of power (e.g., an electrical battery) is used to impose an unnatural driving force for reaction between the waste form and the solution. The source of power can generally be controlled to speed up, slow down, or reverse the reactions occurring in the system, which allows the experimentalist to collect data that are useful for deducing mechanisms of reactions. Because the imposed driving forces tend to keep the system from approaching equilibrium, however, it is difficult to study reaction rates and equilibrium conditions in accelerated tests.

2.1 NON-ACCELERATED RADIONUCLIDE RELEASE TESTS

Non-accelerated radionuclide release tests fall into two categories: dynamic tests and static tests. Discussions of variations of both types of tests and their objectives are given by Bradley, McVay and Coles (1980), McVay, Bradley and Kircher (1981), and Mendel (1982). Because data from different experiments are frequently difficult or impossible to compare, the Materials Characterization Center (MCC) at PNL has defined a series of standardized tests for nuclear waste forms (Mendel 1980, 1984a, 1984b). Most experimentalists in the United States now follow these standardized tests or modifications of them.

2.1.1 Dynamic Tests

Dynamic tests involve either 1) renewal of solution at specified time intervals or 2) continuous flow of solution past a waste form specimen. These tests generally simulate much higher flow rates than those expected in a

geologic repository. Dynamic tests have been used mainly to study the effects of different variables on forward reaction rates or degradation rates of spent fuel or UO_2 . Because the solution is constantly renewed, dynamic tests are not suited for observing saturation effects, formation of thermodynamically stable alteration phases, and long-term interactions with waste package constituents that are important at low flow rates. For this reason, most experimentalists have deemphasized or eliminated dynamic testing in their programs.

A brief review of the two types of dynamic tests is given below.

2.1.1.1 Renewal of Solution in Batches (IAEA and Modified IAEA Tests)

In 1971, Hespe proposed a standard test method for radionuclide release studies for the International Atomic Energy Agency (IAEA). In his procedure he specified a solution-volume-to-waste-form-surface-area ratio of 10 cm. He also suggested a sampling/solution renewal frequency of once a day at the start of the experiment, weekly after the first week, monthly after eight weekly changeouts, and finally semi-annually after six monthly changeouts (Hespe 1971). Many experimentalists have modified this procedure by changing the volume-to-surface area ratio and/or the sampling frequency (generally limiting the changeout to no more than once per month). Some experimentalists have also circulated the solution rather than allow it to sit in static contact with the solid sample. These tests are relatively easy to conduct and do not require a complicated apparatus.

A disadvantage of the IAEA or modified IAEA tests (in addition to those disadvantages already mentioned for dynamic tests in general) is that dissolution kinetics can cause misleading interpretations of the data. For example, if the rate of dissolution of a given component is such that the solubility limit in the solution is reached within one day of contact with the waste form, then the release rate will appear to decrease as contact time with solution is increased. In IAEA tests, the release rate may appear to decrease with time because the same amount of material will be released over successively longer periods of time. Although this is an oversimplified example, it serves to show that rates of dissolution that vary with time and concentration can give rise to misleading results from IAEA tests. A more detailed discussion of this topic is given by Ogard and Bryant (1982).

2.1.1.2 Continuous Flow of Solution (Continuous Flow Tests)

As the name implies, these tests are conducted with continuous flow of solution past the waste form. Since there is generally no recycling of solution, these tests are often called single-pass, continuous flow tests. Samples are taken periodically from the effluent stream.

As in the case of IAEA and modified IAEA tests (Section 2.1.1.1), misleading results can be obtained from continuous flow tests when the dissolution rate of the waste form varies with time and/or concentration. Generally, the release rate of a given component will be greater when the flow rate is higher because the steady-state concentration of a component near the surface of the waste form will decrease as the flow rate increases, which leads to a greater driving force for dissolution.

The continuous flow test was first used at Lawrence Livermore National Laboratory (Weed and Jackson 1979; Coles 1981), and it has been adopted by the MCC as leach test MCC-4 (Mendel 1980, 1984a, 1984b). This review of the literature has revealed no published studies in which a continuous flow test was conducted on spent fuel.

2.1.2 Static Tests

The distinguishing feature of static tests is that the solution is never renewed. These tests simulate the no-flow or very low flow conditions that are expected in a geologic repository. These conditions allow experimentalists to observe saturation effects, the formation of alteration phases (i.e., precipitates and colloids), and interactions of the waste form constituents with the surrounding geology and failed engineered barriers if such materials are included in the test environment along with the waste form.

Some of the conditions used in static tests are summarized below.

• Low temperature (less than 100°C) with an intact waste form. This test is described by MCC leach test MCC-1 (Mendel 1980, 1984b).

- High temperature and frequently at elevated pressures (hydrothermal conditions) with an intact waste form. These tests are generally carried out in autoclaves, and are described by MCC leach test MCC-2 (Mendel 1980, 1984b, 1985).
- Either of the above conditions with a crushed waste form. This test is described by MCC leach test MCC-3 (Mendel 1980, 1984a, 1984b).
- Any of the above conditions in the presence of geologic materials. This type of test is currently being incorporated as one of the standard MCC leach tests.
- Any of the above conditions in the presence of engineered barrier materials.
- Any of the above conditions in the presence of both geologic materials and engineered barrier materials. This type of test is currently being incorporated as one of the standard MCC leach tests.
- Any of the above conditions with the system being rocked rather than kept static.

The reader is referred to McVay, Bradley and Kircher (1981) for a discussion of some of the advantages and disadvantages of static tests.

2.2 ACCELERATED DISSOLUTION (ELECTROCHEMICAL DISSOLUTION)

Accelerated test methods are generally used for investigating mechanisms of dissolution rather than rates of dissolution or solubility limits of various waste form constituents (e.g., Wang 1981a; Sunder et al. 1982). These methods are frequently used in conjunction with sophisticated surface analysis techniques (e.g., scanning electron microscopy, X-ray photoelectron spectroscopy), which provide information on the composition, thickness, and uniformity of films that form on the waste form surface under various conditions. Accelerated test methods involve an electrochemical cell in which a piece of UO_2 or spent fuel is used as one of the electrodes. An outside source of electric power is generally used to step or ramp the potential of this electrode (which is measured against a standard electrode such as the saturated calomel electrode). This induces accelerated oxidation and/or dissolution of the electrode surface. The current observed as a function of time or potential is a direct indication of 1) the oxidation or reduction of a layer on the surface of the electrode, or 2) the dissolution of species from the surface of the electrode. An anodic step or scan (increasing potential) will cause oxidation and consequent dissolution of the electrode surface, while a cathodic step or scan (decreasing potential) will cause reduction of oxidized species on the surface. By abruptly removing the applied potential at various points in time or at various potentials and removing the electrodes from the cell to examine their surfaces, the investigator can determine which oxidized species are formed at the surface and in what order they are formed.

During a scan, sluggish increases in current with increasing potential generally indicate the presence or growth of an oxidized surface layer which serves to passivate the surface, while rapid increases in current with increasing potential indicate that dissolution is occurring. After an anodic step or scan, a reverse cathodic sweep can provide additional information on the thickness and nature of oxidized layers that formed on the surface of the electrode. These reverse scans also allow the experimentalist to determine the fraction of anodic current that went towards oxidation of the electrode surface and the fraction that went towards dissolution (since cathodic scans cause only the reduction of oxidized layers on the electrode surface and not the precipitation of dissolved species).

Electrochemical dissolution experiments have been carried out in a number of different solutions at various temperatures and pH's. Mechanisms of oxidation and dissolution of UO_2 derived from electrochemical experiments have been proposed by Nicol and Needes (1975, 1977), and by a team of Canadian investigators (Sunder et al. 1981, 1982, 1983; Shoesmith et al. 1983, 1984). Wang and Katayama used electrochemical methods to propose mechanisms for dissolution of

both UO_2 and spent fuel, but they did not propose mechanisms for oxidation (Wang and Katayama 1980; Wang 1981; Wang and Katayama 1981a, b, and c, 1982). Their results indicated, however, that dissolution of uranium occurred primarily after oxidation of the surface of the UO_2 or spent fuel. A more detailed discussion of the mechanisms of oxidation and dissolution is given in Section 6.8.

3.0. REPORTING OF RESULTS

Results from radionuclide release tests have been reported in numerous ways in the literature. An excellent review of these is given by McVay, Bradley and Kircher (1981). The most common methods are summarized below:

Release Rates

- Normalized release rate, g/m²-day
- Release rate, g/day
- Activity release rate, Bq/day or Ci/day
- Fractional release rate, fraction/day (day⁻¹)
- Penetration rate, m/day

Integrated Release

- Normalized release, g/m^2
- Fractional release, fraction (no units)
- Weight loss, g
- Concentration, g/mL

Each of these methods of reporting release can be based on different elements in the waste form. Usually the fractional release with respect to a given element is determined by measuring the activity level of that element in the solution and dividing this by the total activity of that element in the waste form specimen. The latter quantity is generally calculated from a computer program for radionuclide generation and depletion such as the ORIGEN-2 code developed at Oak Ridge National Laboratory (Croff 1980).^(a) The fractional release can be used to calculate the release or release rate by any of the units given above provided the following parameters are known:

⁽a) ORIGEN-2 calculations for inventory determinations in small test samples may be subject to large errors. The calculations are frequently relied on for relative inventory determinations while the absolute amounts are determined by "calibrating" the calculations with the measurement of an easily measurable fission product that is soluble in the UO_2 matrix (e.g., 144Ce).

- weight of the waste form specimen, g
- surface area of the waste form specimen, m^2
- density of the waste form specimen, g/m^3
- volume of the solution sample, mL
- number of days solution has been in contact with waste form, days.

In this report (Chapter 6.0), all results from dynamic tests have been presented as fractional release rates, and all results from static tests have been presented as fractional release or concentration. Whenever release or release rates were reported in the literature by other methods, the units of the reported values were converted accordingly. These conversions were done primarily to offer consistency in presentation of the data.

It must be acknowledged that fractional release rates are expected to be directly proportional to the surface area of spent fuel samples. For this reason, release rate data are often reported as normalized release rates (g/m^2-day) . This method of presentation removes the surface area dependence from the reported values and provides a common basis for comparison of the data. The data in this report are not presented on a normalized basis for two reasons: 1) surface areas are often not reported in the literature, and 2) there is a great deal of uncertainty associated with the measurement of surface area. Most of the comparisons of data in this report are based on results from tests in which the spent fuel samples were obtained from the same source and prepared in the same way. Therefore, it is assumed that the surface areas of the samples are approximately the same, and any observed differences in the release behavior between samples are caused by factors other than surface area. In cases in which there is a reason to expect a difference in surface area between samples, it is pointed out that some of the observed difference in release behavior may be caused by a difference in surface area.

4.0 STUDIES OF RADIONUCLIDE RELEASE FROM URANIUM DIOXIDE AND SPENT FUEL

Most studies of radionuclide release from uranium dioxide (UO_2) and spent fuel that are relevant to disposal in a geologic repository have been done within the past 10 years. Before that, virtually no studies were conducted on spent fuel. Those that were conducted on UO_2 were frequently directed at intentional dissolution processes such as leach mining (Pearson and Wadsworth 1958; Parkhaeva and Gromov 1975; Hiskey 1979, 1980), or at the collection of basic thermodynamic data (Gayer and Leider 1957; Tremaine et al. 1981; Ryan and Rai 1983). This section presents all the repository-relevant studies of radionuclide release from UO_2 and spent fuel that have been found in the literature. Part of this section is devoted to studies on simulated spent fuel, which is essentially UO_2 fabricated to contain quantities of fission product compounds that are typically found in spent fuel.

It should be noted that there are three major differences between unirradiated UO_2 and spent fuel that can affect radionuclide release behavior. (The effects that each of these characteristics have on release will be discussed in later sections.) First, spent fuel typically has an intense radiation field associated with it while UO_2 has essentially no radiation field. Second, spent fuel pellets are generally fractured due to thermal cycling and large radial and axial temperature gradients that develop during irradiation. In contrast, UO_2 pellets are almost always structurally intact. This difference is important when intact waste forms are studied. Third, spent fuel frequently contains a number of secondary phases and inclusions consisting of fission product compounds, actinide compounds, and fission gases that are not incorporated in the original fuel matrix whereas UO_2 is generally a homogeneous material.

4.1 URANIUM DIOXIDE

Studies of uranium release from unirradiated UO_2 provide useful insights into the behavior of spent fuel under geologic disposal conditions because spent fuel is generally over 95% UO_2 and therefore its chemistry can be expected to be dominated by UO_2 . Also, UO_2 studies are much less expensive to conduct than spent fuel studies because they do not require the extensive

radioactive shielding that spent fuel studies require. Experiments have been conducted under a variety of conditions using both natural and enriched UO_2 pellets and crushed UO_2 .

The literature contains several references to studies that provide useful insights into the dissolution kinetics and solubility of uranium species under various conditions, but in many cases these conditions are not relevant to geologic disposal. Many studies have pointed out the importance of acids and carbonate species in enhancing the rate of dissolution of UO_2 (Pearson and Wadsworth 1958; Schortmann and DeSesa 1958; Parkhaeva and Gromov 1975; Grandstaff 1976; Hiskey 1979, 1980; Gromov 1981). However, these studies were generally carried out at higher concentrations of acids and carbonates than are expected in deep geologic ground waters. Also, in many studies the uranium sample being dissolved contained significant quantities of uranium in higher oxidation states than UO_2 , as is typical in natural uranium ores (Gayer and Leider 1955; Grandstaff 1976; Rich, Holland and Peterson 1977; Scott, Glasser and Nicol 1977; Holland and Brush 1978). The higher oxides of uranium have been observed to dissolve faster than UO_2 under most conditions.

Many of the solubility studies of uranium have been conducted under much more alkaline conditions than are considered to be possible in a geologic repository (Gayer and Leider 1957; Tremaine et al. 1981; Ryan and Rai 1983). The purpose of these studies has been to obtain basic thermodynamic data on species and reactions that are predominant under alkaline conditions (pH>12). In this section, only studies of uranium release from UO₂ under conditions that are considered to be possible in a geologic repository are presented.

Table 1 summarizes some of the UO_2 studies relevant to geologic disposal that have appeared in the literature. Included among these are two studies of uranium solubility in ground waters and sodium chloride (NaCl) solutions (Dosch 1981; Silva and Yee 1981). The source of uranium in these experiments was the uranyl ion (UO_2^{+2}) , which is expected to be the primary species dissolving from UO_2 under disposal conditions. The results from these studies are discussed in Section 6.0.

TABLE 1. Studies of Uranium Release from UO2

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Laboratory	Description of Solid Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
Pacific Northwest Laboratory (PNL), USA	Samples: Single crystal UO2 Mass: 3-6 g Surface Area: ~4 cm ²	1) Deionized water pH = 5.5-6.8 200 ppm 0 ₂ 2) 0.03 <u>M</u> NaHCO ₃ pH = 9.1-9.4 200 ppm 0 ₂ 3) WIPP "B" Brine pH = 6.9-7.5(a) 200 ppm 0 ₂	Purpose: Determine solubility and dissolution rate in 3 different solutions at different temperatures Method: Static test Container Material: Hastelloy C [®] (Auto- clave) (U0 ₂ contained in Ti capsules) Solution Volume: 40 mL (40:1 volume/SA ratio, cm) Runs: Total of six; once with each solu- tion at two different temps Temps: 78°C 150°C Duration: 81 days Plateout: Yes	Wang 1981 Wang and Katayama 1981b Wang and Katayama 1981c Wang and Katayama 1982
Los Alamos National Laboratory, USA	Samples: Finely divided UO2 Mass: 8-100 mg Surface Area: ~1.30 cm ² /mg (calculated)	1) Deionized water $pH = 4.2 \pm 0.2$ $Eh = -0.15 \pm 0.03 V$ H_2 bubbled through 2) Deionized water $pH = 4.1 \pm 0.2$ $Eh = +0.48 \pm 0.04 V$ $20% 0_2$ in N ₂ bubbled through 3) Deionized water $pH = 3.5 \pm 0.2$ $Eh = -0.18 \pm 0.3 V$ H_2 bubbled through	 Purpose: Determine solubility under oxidizing and reducing conditions at different temperatures Method: Static test with H₂ or 20% O₂ in N₂ bubbled through solution Container Material: Polyethylene (sealed from atmosphere) Solution Volume: 2.0-2.1 liters Runs: Total of 4; 1. Solution 1 at high temp for 55 days 2. Solution 2 at high temp then low temp for last 5 days of 12-day run 4. Solution 3 at low temp for 200 days with conditions altered after 84 days Temps: 25°C 70°C Plateout: No mention 	Norris 1978 Bryant 1979 Norris 1979b Norris 1979d Norris 1980a Norris 1980b Norris 1980c Ogard and Duffy 1981

(a) pH values for hrine solutions are inexact because it is difficult to accurately measure pH in high ionic strength solutions.
 ? = Not reported.

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Laboratory	Description of Solid Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
Technical Research Center, Finland	Samples: UO2 pellets Mass: 13-14 g Surface Area: 7.9 cm ²	 Deionized water pH = ? Eh = ? Saturated with air Olkiluoto groundwater pH = 8.2 initially deaerated prior to use Stripa groundwater pH = 8.3 initially deaerated prior to use 	Purpose: Determine rate of dissolution in 3 different solutions at different • temperatures Method: Modified IAEA test Container Material: Polyethylene Solution Volume: 77 mL Runs: Total of 10 2 with solution 1 at low temp 2 with solution 2 at low temp 2 with solution 2 at high temp 2 with solution 3 at low temp 2 with solution 3 at high temp Temps: $25^{\circ}C$ $60^{\circ}C$ Duration = 612 to 732 days Plateout: No mention	01111a 1985
Technical Research Center, Finland	Samples: UO ₂ pellets Mass: 13-14 [°] g Surface Area: 7.9 cm ²	 Olkiluoto groundwater pH = 8.2 initially Eh = -0.40 to -0.20 V Stripa groundwater pH = 8.3 initially Eh = -0.3 to 0.0 V Olkiluoto groundwater pH = 8.2 initially "oxidizing conditions" Stripa groundwater pH = 8.3 initially "oxidizing conditions" 	 Purpose: Determine solubility and rate of approach to solubility under oxidizing and reducing conditions in different solutions Method: Static test for oxidizing conditions Container Material: Polyethylene Solution Volume: 77 mL for reducing 38 mL for oxidizing Runs: Several at low temp under reducing conditions for both groundwaters 3 each at high temp under oxidizing conditions for both groundwaters (6, 12, and 22 months) Temps: 25°C 60°C Duration: 6, 12, and 22 months Plateout: No mention, but precipitates suggested saturation 	011i1a 1985

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? = Not reported.

Laboratory	Description of Solid Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
Ontario Hydro Research, Canada	Samples: Polished UO ₂ pellets Mass: ~18.5 g Surface Area: 7.9 cm ²	1) Distilled, deionized water pH = see below Eh = see below 2) Granite groundwater pH = see below 3) WN-1 simulated saline groundwater pH = ? Eh = ? The following range of parameters were applied to solutions 1 and 2: a) $PO_2 = 0.25$ to 0.75 atm b) 0.1 to 1.0 M CO_3^{-2} and HCO_3 c) pH from 1 to 12	Purpose: Study the effect of several varjables (pH, Eh, temp., solution, CO3 ² concentration) on rate of dissolution Method: Static test Container Material: Glass Lechant Volume: 100-400 mL Runs: Several under numerous conditions Temps: Varied from 30 to 90°C Duration: 7 to 8 days Plateout: Yes	Thomas and Till 1984
Lawrence Livermore National Laboratory, USA	Samples: UO ₂ pellets Mass: ~16 g Surface Area: ~10.68 cm ²	 WIPP "B" Brine pH = ?(a) Eh = ? Simulated basalt groundwater: pH = ? Eh = ? 0.03 <u>M</u> NaHCO₃ pH = ? Eh = ? 	Purpose: Determine rate of dissolution in different solutions and at different temperatures Method: Single-pass, continuous flow test Container Material: Plastic Solution Flow rates: 1, 10, 300 mL/day Runs: Several using each solution. In the case of solution 1, sandstone was sometimes included in the container with the UO ₂ . In the case of solution 2, Umtanum Basalt was sometimes included. Temps: 25°C 75°C Duration: 435 days Plateout: Not looked for	Bazan et al. 1984

(a) pH values for brine solutions are inexact because it is difficult to accurately measure pH in high ionic strength solutions. ? = Not reported.

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Laboratory	<pre>Vescription of Solid Specimen(s)</pre>	Description of Solution(s)	Experimental Procedure and Conditions	References
Pacific Northwest Laboratory (PNL), USA	Samples: UU ₂ pellets Mass: ? Surface Area: 3.60 cm ² 5.85 cm ²	 Synthetic Permian Basin Brine Final pH = 4.4 to 7.5^(a) atmosphere Deionized water Final pH = 3.9 to 8.1 air atmosphere 	<pre>Purpose: Study the effects of the presence of various constituents on dissolution in various solutions at different temperatures Method: Static test (MCC-1) Container Material: Teflon lined Solution Volume: 36 mL (10:1 Volume/ SA ratio, cm) Runs: Several with each solution at dif- ferent temps and for different durations Systems studied- 1. U02-solution 2. U02-iron-solution 3. U02-zirconium-solution 4. U02-iron-zirconium-solution Temps: 25°C 150°C Duration: 2, 5, 7, 14, 28, and 60 days Plateout: On Iron</pre>	Gray and McVay 1984 Barner et al. 1986 Gray and McVay 1986
Sandia National Laboratory, USA	Sample: dissolved U(VI) nitrate salt added to solutions	 Simulant representing WIPP brine in contact with potash materials pH = 7, 8, 9 (adjusted) Simulant representing WIPP brine in contact with halites pH = 7, 8, 9 (adjusted) Simulant representing "typical" potable WIPP groundwater pH = 7, 8, 9 (adjusted) 	Purpose: Determine solubility and sorption of uranium in different solutions Method: Static test Container Material: Polyethylene Solution Volume: 10-15 mL Runs: 1 for each solution at each pH. Starting values of U(VI) concentration were 1, 10, and 50 mg/L Temp: Room temp Duration: 5 days Plateout: No mention	Dosch 1981

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⁽a) pH values for brine solutions are inexact because it is difficult to accurately measure pH in high ionic strength solutions. ? = Not reported.

Laboratory	Description of Solid Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
Lawrence Berkeley Laboratory, USA	Sample: dissolved U(VI) salt added to solutions	1) U.001 <u>M</u> NaC1 pH = 5 to 10 Eh = ? 2) U.1 <u>M</u> NaC1 pH = 5 to 10 Eh = ?	Purpose: Determine solubility and sorption of uranium in different solutions at various pH Method: Static test Container Material: No mention Solution Volume: No mention Runs: 2 at each pH with initial concen- tration of $U(VI) = 10^{-4}$ M Temp: 23°C Duration: 8.5 and 10 weeks Plateout: No mention	Silva and Yee 1981
South Africa	<pre>Samples: (electrodes): 1) Sintered UO₂ pellets 2) Single crystal UO₂ 3) Fused polycrystal- line UO₂ with U₄O₉ inclusions Mass: ?</pre>	1) 0.1 to 1.1 <u>M</u> HClO ₄ pH = varied from 2 to 13 2) 0.1 <u>M</u> to 1.0 <u>M</u> NaClO ₄ pH = ? 3) 1 <u>M</u> NaClO ₄ plus 10 ⁻³ to 10 ⁻¹ <u>M</u> Na ₂ CO ₃ pH = 9.8 4) 0.5 <u>M</u> and 1.0 <u>M</u> Na ₂ CO ₃ pH = 9 to 11	Purpose: Study mechanisms of UO ₂ dissolution in carbonate and perchlorate solutions Method: Electrochemical Container Material: Titanium Solution Volume: ? Runs: Several with each sample and each solution Temp: 25°C	Nicol and Needes 1975 Nicol and Needes 1977

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? = Not reported.

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Laboratory	Description of Solid Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
Whiteshell Nuclear Research Establishment, Canada	Samples: (electrodes): Polycrystalline UO2 from unused CANDU fuel bundle	1) $0.5 \text{ mol}/\text{dm}^3 \text{ Na}_2\text{SO}_4$ pH = 6 to 12 N ₂ atmosphere 2) $0.5 \text{ mol}/\text{dm}^3 \text{ Na}_2\text{SO}_4 \text{ plus}$ 0 to 0.5 mol/dm ³ CO ₃ pH = ? N ₂ atmosphere 3) $0.1 \text{ mol}/\text{dm}^3 \text{ CO}_3^=$ pH = 9.5 N ₂ atmosphere	Purpose: Study mechanisms of oxidation and dissolution of UO ₂ Method: Electrochemical Container Material: Glass Solution Volume: ? Runs: Several in each solution Temps: 25° to 30°C	Sunder et al. 1981 Sunder et al. 1982 Sunder et al. 1983 Shoesmith et al. 1983 Shoesmith et al. 1984
Pacific Northwest Laboratory (PNL), USA	Samples: Single crystal ¹³⁰ 2	1) Deionized water pH = $5.3-5.7$ 2) $0.03 \text{ M} \text{ NaHCO}_3$ pH = $9.0-9.3$ air atmosphere 3) WIPP "B" brine pH = $7.0-7.3$ ^(a) air atmosphere	Purpose: Study mechanisms of dissolution of UO2 Method: Electrochemical Container Material: Pyrex® glass Solution Volume: <250 mL Runs: At least one in each solution at two different temperatures Temps: 25°C 75°C	Wang 1981 Wang and Katayama 1981a Wang and Katayama 1981b Wang and Katayama 1981c Wang and Katayama 1982

(a) pH values for brine solutions are inexact because it is difficult to accurately measure pH in high ionic strength solutions. ? = Not reported.

4.2 SPENT FUEL

Several studies of radionuclide release from spent fuel relevant to geologic disposal have been conducted within the last 10 years. These have been carried out under a variety of conditions and for numerous purposes. Most studies have been conducted on light-water reactor fuels enriched in 235 U, but the Canadians have done several studies on natural uranium fuel irradiated in deuterium water reactors. The fuels that have been studied have had several different irradiation histories. Fuel specimens have included pellets with cladding, pellets without cladding, and crushed spent fuel; solutions have included deionized water and all types of ground waters. In some experiments, host rock and/or waste package barrier materials have been included in the testing systems. Table 2 summarizes the spent fuel radionuclide release studies that have appeared in the literature. The results from these studies are discussed in Chapter 6.0.

4.3 SIMULATED SPENT FUEL

Studies of release from simulated spent fuel (SSF) have been conducted by Rockwell Hanford Operation's Basalt Waste Isolation Project (Apted and Myers 1982; Grandstaff et al. 1983; Myers, Apted and Mazer 1984; McKeon et al. 1984). The simulated spent fuel contains the same stoichiometric quantities of fission product compounds (albeit non-radioactive isotopes) as 10-year-old pressurized water reactor (PWR) fuel with a burnup of 33 MWd/kg U. It is fabricated at the Hanford Engineering Development Laboratory (operated by Westinghouse Hanford Company). The SSF provides a waste form of the same bulk composition as spent fuel without the radiation field associated with spent fuel. Details of the composition and the fabrication procedure are given by Woodley, Wilson and Hervig (1981).

Although the simulated spent fuel has the same bulk composition as spent fuel, investigators should be careful when comparing SSF release test results to spent fuel release test results for two reasons: 1) unlike the spent fuel, SSF does not have a radiation field associated with it and 2) the various components of SSF are probably not distributed in the grains and along the grain boundaries in the same way that they are in spent fuel. Simulated spent fuel

TABLE 2. Studies of Radionuclide Release from Spent Fuel

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Surface

4.9 9.4

Surface Area, cm²

> 5.6 9.5

Mass.g Area, cm²

 Quad Cities-11 fuel fragments (no cladding)
 Burnup: 9.0 MMd/kg U
 Enrichment: ?

Mass,g

4.853 4.846

<u>Id</u> <u>Mass.g</u> HBR-1 4.944 HBR-2 5.109

<u>ld</u>

QC-1 QC-2

Laboratory	Description of Spent Fuel Specimen(s)	Description of Solutions	Experimental Procedure and Conditions	References
Pacific Northwest Laboratory (PNL), USA	Light water reactor spent fuels: 1) Zorita Reactor fuel fragments (no cladding) Burnup: 54.5 MWd/kg U Enrichment: 5.81 wtX 235 U Surface 1d Mass.q Area. cm 2-2 5.183 11.7 7-3 5.048 11.4 2-6 5.516 12.6 2-7 4.431 10.0 Z-8 2.722 37.6 2) H. B. Robinson-11 fuel fragments (no cladding) Burnup: 28.0 MWd/kg U Enrichment: 2.55 wtX 235 U	 Deionized water pH = ? air saturated Synthetic sea water solution pH = ? high in NaCl air saturated Hanford groundwater pH = 7.9 ~168 ppm HCO² air saturated 	 Purpose: Study the effects of burnup and different solutions on release rates Method: Modified IAEA in Paige apparatus with solution recirculated by air bubbling Container Material: Glass Solution Volume: 500 mL Runs: Z-6, Z-7, HBR-1, HBR-2, QC-1, QC-2 in deionized water Z-2, Z-3, in Hanford groundwater Z-8 in synthetic sea water Temp: 25°C Duration: Approx 350 days for groundwater tests, 822 to 1399 days for others Plateout: Yes, solution samples acidified 	Katayama 1976 Katayama and Mendel 1977 Katayama 1979 Katayama et al. 1980b Katayama and Bradley 1980

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? * Not reported.

Laboratory	Description of Spent Fuel Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
Pacific Northwest Laboratory (PNL), USA	Samples: H. B. Robinson spent fuel fragments (no cladding) Burnup: 28.0 MWd/kg U Enrichment: 2.55 wt% ²³⁵ U Mass: ~15 g Surface Area: ~30 cm ²	1) Deionized water pH = 6.6 decreasing to 4.3 at end of run 2) 0.03 M NaC1 $pH = \frac{7}{2}$ 3) WIPP "B".Brine pH = 6.5(a) 4) 0.03 M NaHCO ₃ pH = 7 5) 0.015 M CaC1 ₂ pH = 7	Purpose: Study the effects of different solutions on release rates Method: Modified IAEA Container Material: High density polypropylene (sealed) Solution Volume: 300 mL (10:1 volume to surface area ratio, cm) Runs: 1 in each solution Temp: 25°C Duration: 467 days Plateout: Yes, solution samples acidified	Katayama et al. 1980a Katayama and Bradley 1980
K4S, Sweden	Samples: 20 mm long sections of LWR spent fuel elements with cladding Linear Power Rating: 11.2 kw/m and 23.5 kw/m Burnup: 11.5 and 26 MWd/kg U Enrichment: ? Surface Area: ? Mass: 21-22 g with cladding 17-18 g without cladding	 Distilled water pH = ? Swedish groundwater pH = 8,5 ~300 ppm HCO₃ 	 Purpose: Study the effects of linear power rating and different solutions on release rates Method: Modified IAEA, no stirring Container Material: Glass Solution Volume: 500 mL Runs: 4 total; 1 in each solution for each fuel power rating Temp: 60°C Duration: 105 days Plateout: No mention, but solution samples acidified 	Eklund and Forsyth 1978 Forsyth 1983
Whiteshell Nuclear Research Establishment (WNRE), Canada	Samples: 5 cm long sections of CANDU/Pickering fuel elements with cladding Burnup: 7.1 MWd/kg U Enrichment: natural Linear Power Rating: 53 kw/m Mass: 92.02 g Surface Area: 3.2 cm ² (ends of pellet)	 Distilled, deionized water pH = 5,9 air saturated Chlorinated river water pH = 7.7 air saturated 	Purpose: Study the effect of different solutions on release rates Method: Modified LAEA Container Material: Polyethylene Solution Volume: 100 mL Runs: 4 total; 2 in each solution Temp: 25°C Duration: ~900 days Plateout: No mention, but solution samples aridified	Vandergraaf 1980 Vandergraaf et al. 1980

(a) pH values for brine solutions are inexact because it is difficult to accurately measure pH in high ionic strength solutions. ? = Not reported.

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samples acidified

Description of Solution(s)

Laboratory

Los Alamos National Laboratory, USA Samples: H. B. Robinson-II fuel cut into wafers (with cladding) Burnup: 28.0 MHd/kg U 23 Enrichment: 3.10 wt% 235U Mass: 2.6 to 3.7 g Surface Area: 1.36 cm² (ends of wafers)

Description of Spent Fuel Specimen(s)

1) Deionized water pH = 4-5.5 Eh = +0.44 V20% 0₂ bubbled through 2) Deionized water pH = 4-5.5 Eh = -0.12 V5% H₂ bubbled through Purpose: Study the effects of temperature and Eh on release rates Nethod: Modified IAEA Container Material: Glass Solution Volume: 5 mL Runs: 4 total; 2 in each solution Temp raised from 25°C to 70°C after 65 days Temp: 25°C 70°C Duration: ~190 days Plateout: No mention

Experimental Procedure and Conditions

Ogard et al. 1931

References

Norris 1978

Bryant 1979

Norris 1979a

Norris 1979b

Norris 1979c

Norris 1979d

Norris 1980a

Norris 1980b

Norris 1980c

Nak Ridge National Laboratory, USA Samples: Crushed Oconee-1 spent fuel (no cladding) Burnup: 23 MMd/kg H Mass: -10 g Surface Area: 3.8 cm²/g (18 mesh) and 42 cm²/g (200 mesh) 1) Distilled water with $0.30 \text{ M} \text{ H}_3\text{RO}_3 0.04 \text{ M}$ NaOH pH = 8.0(reactor pool cooling water composition) Purpose: Determine release rates in reactor cooling water Method: Modified LAEA Container Material: Glass Solution Volume: 100 mL Runs: 4 total; one of each type of sample (18 mesh & 200 mesh) at low temp. and high temp. Temps: 35°C 100°C Ouration: 280 days Plateout: No mention Mitchell, Goode and Vaughen 1931
TABLE 2. (contd)

Laboratory

WNRE, Canada

Description of Spent Fuel Specimen(s)

Description of Solution(s) 1) Distilled, deionized

Experimental Procedure and Conditions References Purpose: Study the effects of 0₂ pres-

Johnson et al. 1981 Johnson and Joling 1982

Samples: 4-5 cm long sections of CANDU/Pickering fuel elements with cladding Burnup: 7.9 MWd/kg U Enrichment: Natural Mass: ~80-90 g Surface Area: 3.2 cm² (ends of pellets)

water pH = 5-5.6 air saturated 2) Distilled water at 02 pressures ranging from 20 to 700 kPa

- 3) Granite groundwater pH = 6.9 to 7.5 air saturated
- 4) Granite groundwater pH = ~7

at 100 kPa H₂ pressure

Solution Volumes: 100 mL for modified IAEA and one day static tests

sure, temperature, and different

Method: Modified IAEA followed by static

polyethylene for modified IAEA tests

• stainless steel autoclaves for one day

titanium autoclaves for longer static

tests of varying duration

solutions on release rates and total

500 mL for longer static tests

release

Container Materials:

static tests

tests

- Runs: 1) All samples leached by IAEA method in solution 1 for 100-120 days at 25°C
- prior to static testing2) 5 one day static tests in solution 2 at 20, 200, 300, 400, and 700 kPa 02 pressure
- 3) 2 static tests in solution 1 at 150°C for 10 days
- 4) 3 static tests in solution 3 at 150°C for 8 to 28 days
- 5) 1 static test in solution 4 at 150°C for ~20 days Temps: 25°C

- 150°C
- Duration: see "Runs" above Plateout: No mention, but solution samples acidified

TABLE 2. (contd)

Laboratory	Description of Spent Fuel Specimen(s)	Description of Solution(s)	Experimental Procedure and Conditions	References
WNRE, Canada	Samples: ~5 cm long sections of CANDU/Pickering fuel elements with cladding Burnup: 7.9 MWd/kg U Enrichment: Natural Linear Power Ratings: 43 kw/m and 53 kw/m Mass: 70-80 g Surface Area: 3.2 cm ² (ends of pellets)	 Distilled, deionized water pH = 5.8 Swedish groundwater pH = 8.5 Granite groundwater pH = 6.5-7.0 at O₂ concentrations ranging from 1 x 10⁻⁰ to 2.5 x 10⁻⁴ mol/kg Standard Canadian shield saline solution pH = 7.0 	 Purpose: Study the effects of O₂ concentration, linear power rating, and different solutions on release rates Method: Modified IAEA Container Materials: Polypropylene and glass (for U₂ bubbling experiments) Solution Volume: 100 mL Runs: 2 with each solution 1 at each of the O₂ concentrations in solution 1 1 of fuel with each power rating in solution 1 Temp: 25°C Duration: -900 days for each solution ~450 days for O₂ studies Plateout: Yes solution samples acidified 	Johnson 1982 Johnson et al. 1982
КВS, Sweden	Samples: 20 mm long sections of LWR spent fuel elements with cladding Burnup: 42 MWd/kg U Enrichment: ? Mass: ~20 g with cladding ~16 g without cladding Surface Area: ?	 Deionized water pH = 7.0 oxidizing conditions Swedish groundwater pH = 8.0-8.2 ~123 ppm HCO₃ oxidizing conditions Same as (2), but bubbled with H₂ (reducing conditions) Same as (2), but pH adjusted to 2 to 5 	 Purpose: Study the effects of different solutions and different oxidizing/reducing conditions on release rates and total release Method: Modified IAEA and Static Container Material: Pyrex glass Solution Volume: 200 mL Runs: Series of modified IAEA tests in solution 1 Series of modified IAEA tests in solution 2 Series of static tests in solution 2 Series of static tests in solution 4 for 20 days at end of test Series of tests with pre-leach in solution 3 Temp: 20-25°C Duration: 360 to 570 days 	Forsyth 1982 . Forsyth 1983 Forsyth et al. 1984

? = Not reported.

TABLE 2. (contd)

WNRE, Canada	Samples: Several 1 to 5 cm long sections of CANDU fuel elements from Pickering and Bruce reactors (with cladding). Some longer sections of fuel elements	 Distilled, deionized water pH = ? air saturated 	Purpose: Study the rapid release of Cs and I under various conditions Methods: Modified IAEA, continuous flow,	Burns, Moore and Boase 1982
	were also used. Burnup: Pickering: 7.1 to 12.6 MWd/kg U Bruce: 9.6 to 13.8 MWd/kg U Linear Power Rating: Pickering: 29 to 52 kw/m Bruce: 32 to 47 kw/m Enrichment: Natural Mass: 15 to 80 g Surface Area: 3.2 cm ² (ends of pellets)	 2) 0.2 g/L Kl air saturated 3) 0.01 g/L CsCl air saturated 4) 1.0 g/L 1⁻ 0.01 g/L Cs+ air saturated 5) 0.5 g/L NaCl 	 and static Container Material: Plastic at low temp., titanium or stainless steel at high temp. Solution Volumes: Modified IAEA: 100 mL Continuous Flow: 5 mL/min Static: 1 to 7 L Runs: 1) Several modified IAEA tests at 25°C in solutions 1, 2 and 3 for anywhere from 30 to 300 days 2) Some tests involving continuous flow of solution 4 at 25° to 35°C with the flow being through a long section of fuel element 3) Several static tests with solution 1 at 150°C 4) Two static tests at 150°C on whole fuel elements with 1 to 5 mm slit cladding defects in cladding Temps: 25°C 150°C Duration: Few days to 300 days Plateout: No mention 	Johnson et al. 1983 Johnson et al. 1984

? = Not reported.

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TABLE 2. (contd)

Laboratory	Description of Spent Fuel Specimen(s)	Descr	iption of Solution(s)	Experimental Procedure and Conditions		References	
Hanford Engineering Development Laboratory, Lawrence Livermore National Laboratory, USA	Samples: 2 to 5 inch long sections of completely clad spent LWR fuel with cladding defects (with water tight stainless steel caps on the ends of the sections) Fuel: 1) Turkey Point fuel Burnup: 25.5 to 27 MMd/kg U Enrichment: 2.56 wt% ²³⁵ U 2) H. 8. Robinson-II fuel Burnup: 27.7 to 31 MMd/kg U Enrichment: 2.55 wt% ²³⁵ U Types of Defects 1) Undefected rod 2) Two laser drilled holes, 200 µm in diameter 3) One slit defect, 2 to 2.5 cm long by 150 to 200 µm wide 4) Bare fuel plus cladding hulls	1) D 2) J t t	leionized water HH = 6 to 7 I-13 well water from fucca Mountain, Nevada HH = 7.2 to 8.5 during sest -20 ppm HCO3	 Purpose: Study the containment of radionuclides provided by spent fuel cladding Method: Essentially static with some replenishment (semi-static) Container Material: Fused quartz Solution Volume: 250 mL (10 mL samples taken in modified IAEA fashion and volume replenished with fresh solution) Runs: Two of each type of defected Turkey Point fuel specimen in solution 1. Two of each type of defected M. B. Robinson fuel specimen in solution 2. Two of each type of defected Turkey Point fuel specimen in solution 2. Two of each type of defected Turkey Point fuel specimen in solution 2. Two of each type of defected Turkey Point fuel specimen in solution 2. Two of each type of defected Turkey Point fuel specimen in solution 2. Two of each type of defected Turkey Point fuel specimen in solution 2. Temp: ~25°C Duration: 220 to 250 days (some in progress) Plateout: Yes 	Wilson Wilson Wilson Wilson	1983 and Oversby 1984 and Oversby 1985	1984
Pacific Northwest Laboratory (PNL), USA	Samples: Ground LWR spent fuel from Turkey Point Burnup: 25.5 to 27.0 Myd/kg U Enrichment: 2.56 wt% ²⁵⁰ U Mass: 19.26 g	1) 8 p t	asalt groundwater H = 7.2 to 8.0 during est	Purpose: Study release under high temperature, high pressure conditions in basalt groundwater Method: Static Test Container Material: Gold bag (within Dickson rocking autoclave) Solution Volume: ~195 mL Run: 1 with solution 1 at 30 mPa pressure Temp: 200°C Duration: 55 days Plateout: No mention	Schrami and	ke, Simonson Coles 1984	

Laboratory	Description of Spent Fuel Specimen(s)	Description of Solution(s)		Experimental Procedure and Conditions	References	
Pacific Northwest Laboratory (PNL), USA	Samples: Fuel fragments from H. B. Robinson spent fuel Burnup: 28.0 MHd/kg U Enrichment: 2.55 wt% ²³⁵ U Mass: ? Surface Area: ~2 cm ²	1) 2)	Synthetic Permian Basin Brine pH = 4.65 to 6.5(a) air atmosphere Real Brine Final pH = 6.33(a) air atmosphere	Purpose: Study the effect of the presence of iron and of temperature on total release Method: Static Test (MCC-3) Container Material: Quartz Solution Volume: 17 to 30 mL (10:1 Volume/SA ratio) Runs: 1) 5 in solution 1 at 25°C 2) 5 with iron in solution 1 at 25°C 3) 1 in solution 2 at 25°C 4) 2 in solution 1 at 75°C 5) 2 with iron in solution 1 at 75°C Temps: 25°C Durations: 2, 5, 14, 28, and 60 days Plateout: Yes	Gray et al. 1983 Gray and McVay 1: Barner et al. 198 Gray and McVay 198	984 36 36
Pacific Northwest Laboratory (PNL), USA	Samples: Fuel fragments from H. B. Robinson spent fuel Burnup: 28 MWd/kg U Enrighment: 2.55 wt% 230 Mass: ?	1) 2) 3)	Deionized water pH = 5.0 air atmosphere 0.03 M NaHCO ₃ pH = 8.9 air atmosphere WIPP "B" Brine pH = 6.9(a) air atmosphere	Purpose: Study mechanisms of spent fuel matrix dissolution Method: Electrochemical Container Material: Pyrex glass Solution Volume: <250 mL Runs: At least one in each solution Temp: 25°C	Wang and Katayama Wang and Katayama Wang and Katayama Wang and Katayama Wang and Katayama	1980 1981a 1981b 1981c 1982

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(a) pH values for brine solutions are inexact because it is difficult to accurately measure pH in high ionic strength solutions. ? = Not reported.

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TABLE 2. (contd)

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may be of use, however, in observing the formation of alteration phases in a simulated geologic disposal environment and in observing in which phases the various components of the waste form end up after they are released from the waste form.

Table 3 summarizes the studies that have been done on simulated spent fuel.

Laboratory	Description of Spent Fuel Specimen(s)	Description of Sol	tion(s) Experimental Procedure and Conditions	References
Basalt Waste Isolation Project, USA	Samples: crushed simu- lated spent fuel	 Synthetic Grand Basalt Groundwa pH = 6.0 to 10 runs Eh = ? 	e Ronde ter Juring Juring Saving Discrete Static Test Container Material Gold bag (within Dickson rocking autoclave) Solution Volume: 10:1 solution to SSF mass ratio Runs: 2 at 100°C, 30 mPa 1 at 300°C, 30 mPa Duration: 427 to 2514 hours	Apted and Myers 1982 Myers, Apted and Mazer 1984
Temple University, USA	Samples: crushed simu- lated spent fuel with crushed basalt rock also present	 Synthetic Basal Groundwater pH = 6.0 to 10 runs Eh = ? 	 Purpose: Study release under high temperature, high pressure conditions in basalt groundwater in the presence of basalt rock Method: Static Test Container Material: Gold bag (within Dickson rocking autoclave) Solution Volume: Ranging from 12.5:1 to 50:1 solution to SSF mass ratio Runs: 3 at 300°C, 30 mPa and 20:1:1 solution/SSF/basalt ratio 1 at 100°C, 30 mPa, and 20:1:1 ratio 1 at 300°C, 30 mPa, and 50:1:1 ratio 1 at 300°C, 30 mPa, and 50:1:1 ratio 1 at 300°C, 30 mPa, and 50:1:4 ratio 1 at 300°C, 30 mPa, and 50:1:4 ratio 	Apted and Myers 1982 McKeon, Ulmer and Grandstaff 1982 Grandstaff et al. 198 McKeon, Ulmer and Grandstaff 1984 Myers, Apted and Mazer 1984

TABLE 3. Studies of Release from Simulated Spent Fuel

? = Not reported.

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5.0 THEORETICAL STUDIES OF URANIUM AND ACTINIDE SOLUBILITY

Much work has been done to develop methods of predicting the solubility of actinide species (uranium in particular) in solutions of various compositions. This work has provided valuable insights into what factors are important in the dissolution of UO_2 under a wide range of conditions. The work has proven useful in explaining and interpreting results of tests of radionuclide release from UO_2 and spent fuel.

The methods of predicting solubility are all based on fundamental thermodynamic data. These data include standard free energies of formation for each species in the system, and equilibrium constants for each reaction occurring in the system. The reactions include dissolution, complex formation, hydrolysis, dissociation, and redox reactions. Because of the complex chemistry of the actinide elements, this amounts to a great deal of data. Fortunately, several investigators have compiled most of the pertinent data that have appeared in the literature (Smith and Martell 1976; Baes and Mesmer 1976; Rai and Serne 1977; Langmuir 1978; Cleveland 1979; Lemire and Tremaine 1980; Langmuir and Herman 1980; Krupka, Jenne and Deutsch 1983; Lemire 1984).

The thermodynamic data have been used in various ways to arrive at predictions of the equilibrium solubility of uranium and other actinides under various conditions. All of the methods involve the simultaneous solution of the thermodynamic equations that describe the system. This is equivalent to finding the global minimum of the total Gibbs free energy of the system. At equilibrium, all of the expressions describing solubility, complex formation, dissociation, and hydrolysis must be satisfied as well as the Nernst equation for each oxidation or reduction half-reaction as paired with a standard electrode half reaction (Garrels and Christ 1965; Zeleznik and Gordon 1968; van Zeggeren and Storey 1970). Because of the large number of equations that this represents, computer methods are almost always used. Some of the computer codes used to do the complex thermodynamic calculations include SOLMNEQ (Kharaka and Barnes 1973), WATEQ and its later versions (Truesdell and Jones 1974), MINEQL (Westall, Zachary and Morel 1976), PHREEQE (Parkhurst, Thorstenson and Plummer 1980), EQ3/EQ6 (INTERA 1983), MINTEQ (Felmy, Girvin and

Jenne 1984), and EQ3/6 (Wolery et al. 1985). Solution of the system of equations yields the equilibrium concentrations of each species at a given pH, Eh, temperature, and initial composition of solution.

One of the ways of presenting the results of the above computations is with potential-pH diagrams (Skytte-Jensen 1980; Paquette and Lemire 1981; Lemire 1984). These diagrams show the most stable species in the system as a function of pH and Eh. An example of potential-pH diagrams for the UO_2 /water system at different temperatures is given in Figure 1 (from Paquette and Lemire



FIGURE 1. Potential pH Diagrams for the Uranium/Water System at (a) 25° C, (b) 100° C, and (c) 200° C. Dissolved species activity is 10^{-9} . Dotted lines represent upper and lower limits of stability of water (Paquette and Lemire 1981). It was later established that the field shown for U(0H)₅ is unreasonably large.

1981). A full treatment of the basis of potential-pH diagrams is given by Pourbaix (1966) and by Garrels and Christ (1965), and an excellent outline of the general principles as applied to actinide species is given by Paquette and Lemire (1981). Contour plotting of isosolubility lines from potential-pH diagrams can provide information on the solubility of various species at different pH and Eh (Paquette and Lemire 1981).

A shortcoming of potential-pH diagrams is that they provide solubility information on only the most stable species (i.e., those present in highest concentration) as a function of Eh and pH. In many cases, the solubility of the most stable species provides a reasonable approximation to the solubility of all actinide-bearing species (because the other species are present at orders of magnitude lower concentrations), but this is not always the case. For this reason, when actinide solubility is the quantity of interest, most investigators present the results of their calculations as plots or tables of species concentration versus pH or Eh with all the other variables held constant (Goodwin 1980; Bird 1980; Allard 1982, 1983, 1984a; Duffy and Ogard 1982; Rees 1985). Frequently, the pH and Eh of a solution are not independent because a reversible reaction is buffering the redox potential of the solution. In this case, the pH-Eh dependence is described by the Nernst equation for the buffering reaction. An example of a plot of uranium concentration versus pH where the Eh is set by magnetite-hematite (Fe_3O_4 - Fe_2O_3) equilibria is given in Figure 2 (Goodwin 1980).

At least one team of investigators has attempted to reduce the complexity of solving the large system of equations necessary to predict uranium solubility under various conditions (Garisto and Garisto 1985). They have compiled tables of constants that can be plugged into a formula to calculate total uranium solubility as a function of pH, Eh, temperature, and solution composition. Their model has limits of applicability, however, which they carefully define.

Some of the general limitations of thermodynamic calculations for predicting uranium and actinide solubility are listed below:

• The data base of thermodynamic constants is incomplete and, in many cases, the data that are available have a great deal of uncertainty.



FIGURE 2. Maximum Uranium Solubilities in Brine Ground Water. The abscissa represents the Eh-pH equilibrium boundary of the hematitemagnetite reaction (only pH values are shown). The asterisks show what is believed to be the natural Eh-pH conditions of the brine at the three temperatures (Goodwin 1980).

- The thermodynamic calculations predict only equilibrium concentrations--they do not take into account kinetic effects. Some of the geochemical computer codes are capable of calculating reaction kinetics, but much of the kinetic data necessary to do these calculations simply does not exist at the present time. It is possible that under geologic disposal conditions equilibrium will not be attained for thousands of years, and possibly it will never be attained. In this case, the applicability of thermodynamic calculations is limited.
- The thermodynamic calculations do not account for metastable phases. These may be present for indefinite periods of time under geologic disposal conditions.
- The thermodynamic calculations do not account for colloidal species. These have frequently been observed in experimental systems.
- The thermodynamic calculations that have appeared in the literature have dealt almost entirely with actinide-ground-water systems. The situation may be considerably more complex when host rock is included in the system. In this case, thermodynamically stable alteration phases consisting of compounds containing actinides and rock constituents must be considered.

A more complete discussion of these and other limitations is given by Goodwin (1980).

Some of the major conclusions drawn from the theoretical studies of uranium and actinide solubility are:

 The solubility of uranium and the actinides in general is predicted to be lowest in the pH range 4 to 10 in most solutions. At lower pH's, the actinides are apt to be oxidized to more soluble ionic species and/or dissolved by simple acid dissolution. At higher pH's, soluble hydroxyl complexes or other anionic species complexes are more likely to be formed.

- The solubility of uranium and the actinides is predicted to increase with increasing Eh. This is because the higher oxides of the actinides are generally more soluble than the lower oxides. An exception to this is Plutonium (VI), which can form anionic species that coprecipitate with Na⁺ in brines.
- Temperature has very little effect on the solubility of uranium and the actinides. Its effect is due mainly to the way that it affects the equilibrium constants in the system and the pH of the solution.
- The solubility of uranium and the actinides is enhanced by the presence of certain anionic species that form complexes with the actinides. These species include OH⁻, HCO₃⁻, CO₃⁻², F⁻, Cl⁻, SO₄⁻², H₂PO₄⁻², HPO₄⁻², and PO₄⁻³, of which the carbonates and phosphates are the most important.

6.0 <u>RESULTS FROM EXPERIMENTAL STUDIES OF RADIONUCLIDE RELEASE FROM</u> URANIUM DIOXIDE AND SPENT FUEL

Many experimental studies have been performed to investigate the effects of different variables on the release of radionuclides from UO_2 and spent fuel. This section discusses the results of these studies and how well these results agree with what was expected or predicted from theory.

6.1 GENERAL TRENDS OF RELEASE OF VARIOUS RADIONUCLIDES

Essentially all of the experimentalists who have studied radionuclide release from spent fuel have observed different rates of release for different radionuclides from spent fuel (see Table 2 for references). Figure 3 shows the release rates of several elements over a three-year period in deionized water at 25°C as measured by Katayama, Bradley and Harvey (1980a). To some extent, the different studies listed in Table 2 have yielded conflicting results, but a few basic trends have been observed, which are summarized below.

6.1.1 <u>Release of Cesium and Iodine</u>

The fission products cesium-134 (134 Cs), cesium-135 (135 Cs), cesium-137 (137 Cs), and iodine-129 (129 I) all release at very high rates initially and then, after a period of a few days to several weeks, they drop off to rates that are comparable to other fission products. 99 Tc has also been observed to exhibit this behavior, but the overall release of this radionuclide has generally been slightly lower than that for cesium or iodine (Barner et al. 1985; Wilson 1985). Also, there is less data available for 99 Tc. Certain other radionuclides, such as selenium-79 and carbon-14^(a), are also expected to be rapidly released, but there are essentially no data on the release of these elements from spent fuel.

The amount of cesium and iodine released during the initial high release period may be as much as 1 to 5% of the inventory of these elements. The reason for the high initial release of these elements (and for 99 Tc) has generally

⁽a) Carbon-14 is an activation product resulting from irradiation of nitrogen impurities in the fuel and from irradiation of 17 O.



FIGURE 3. Fractional Release Rate of Various Radionuclides from 28.0 MWd/kg U Spent Fuel in Deionized Water at 25°C (Katayama, Bradley and Harvey 1980a)

been attributed to 1) their relatively high solubility and 2) their tendency to accumulate at exposed surfaces of the fuel during irradiation of the fuel. Regarding the second characteristic, these elements are all very insoluble in the solid UO_2 matrix (i.e., they are not easily incorporated into the matrix as oxides or otherwise) and, upon forming, they tend to migrate to grain boundaries or cracks in the fuel and/or down the temperature gradient to the periphery of the fuel. Because these elements are not bound to the UO_2 matrix and because the cracks, grain boundaries, and the outer periphery of the fuel are the first surfaces to come in contact with the solution during radionuclide release testing, these elements are very rapidly released from the fuel.

In addition to being observed in the studies of radionuclide release from spent fuel presented in Table 2, the rapid release of cesium and iodine has been studied by several other investigators (Peehs et al. 1978; Devell and Hesbol 1978; Lorenz et al. 1980; Peehs et al. 1980; Peehs, Kaspar and Neeb 1983). These studies were concerned with the release of cesium and iodine from failed fuel pins (i.e., pins with ruptured cladding) to reactor core water or water storage pools at reactor sites. The results from all of the studies mentioned in this section suggest that the amount of radionuclides that are rapidly released is dependent on conditions that prevailed during irradiation rather than on conditions that prevailed during the release tests themselves.

Some attempts have been made by Canadian investigators to correlate the amount of cesium and iodine released during the initial rapid release period to the amount of the gaseous fission product xenon released from the fuel (Johnson et al. 1983; Johnson et al. 1984). There are reasons to believe that the gap and grain boundary inventories of cesium and iodine are correlated with the noble gas inventory in the gap and grain boundaries. The amount of xenon released (i.e., in the gap) can be predicted using the Canadian computer code, ELESIM, which calculates the release based on the power history of the fuel (Notley 1979). These efforts to correlate cesium and iodine release to xenon release have met with modest success.

6.1.2 Release of Other Fission Products and Actinides

Most fission products (other than cesium and iodine) and actinides (including uranium) all tend to be released from the spent fuel matrix at initially lower rates than either cesium or iodine. This has generally been attributed to the inclusion of these elements in the UO_2 matrix as solid compounds, generally oxides. Once incorporated in the matrix, the migration of these elements is prevented or retarded. It has been widely proposed that the release of these elements is controlled by dissolution of the UO_2 matrix.

Some investigators have argued that fission products and actinides are released from the UO_2 matrix congruently with uranium, but many studies have produced results that suggest that some elements are released at slightly higher rates than uranium (i.e., they are preferentially leached). There is little agreement on the relative order of release rates for the different

elements, but in general, the fission products have been observed to be released slightly faster than the actinides. The latter are released at about the same rate as uranium.

6.2 EFFECT OF SOLUTION COMPOSITION AND CHEMICAL SPECIES ON RADIONUCLIDE RELEASE

The effects of different solution compositions on the release rates of various elements from spent fuel and UO_2 have been investigated in several studies (see Table 2). The most commonly used liquids have been deionized and/or distilled water and solutions that simulate ground waters from proposed geologic repository sites. The latter include brines, granite ground waters, basalt ground waters, and tuff ground waters. A few studies have been done using salt solutions (e.g., sodium chloride and calcium chloride) and carbonate/bicarbonate solutions.

In general, solution composition has been observed to have a greater effect on the release rates of radionuclides that are incorporated in the UO_2 matrix than on the release rates of radionuclides that are present at the exposed surfaces of the fuel (e.g., cesium and iodine). To illustrate this, Figures 4 and 5 show the release rates of 137 Cs and Curium-244 (244 Cm), respectively, in the same five solutions (Katayama, Bradley and Harvey 1980a). Curium is expected to be incorporated in the UO_2 matrix as an oxide, and it is apparent that its release rate is more affected by solution composition than is the release rate of cesium. This suggests that the main effect of solution composition is on the degradation rate of the UO_2 matrix.

The results for uranium (especially in deionized water and carbonate water) are somewhat contradictory, which makes the assertion that solution composition affects UO_2 on matrix dissolution difficult to support. Johnson (1982) observed that uranium dissolved faster from spent fuel in carbonate ground water than in air-saturated distilled water. This is in agreement with experimental studies of unirradiated UO_2 , which show enhanced solubility in carbonate solutions due to the formation of uranyl carbonate complexes (Wang 1981). Eklund and Forsyth (1978), however, used the same carbonate ground water as Johnson and observed no difference between the release rate of uranium



FIGURE 4. Fractional Release of ¹³⁷Cs from 28 MWd/kg U Spent Fuel in Various Solutions at 25°C (Katayama, Bradley and Harvey 1980a)

in distilled water and carbonate water. Katayama, Bradley and Harvey (1980a) observed higher uranium release rates in air-saturated deionized water than in any of the other solutions that they studied, including 0.03 M NaHCO₃.

Very few explanations have been offered for the conflicting uranium behavior in these experiments. Wang and Katayama (1981b) have suggested that radiolysis may have played a role in the Katayama experiments. They note that the pH drop that was observed in all the solutions during the course of the experiments was much greater in deionized water than in the other solutions. They



FIGURE 5. Fractional Release Rate of ²⁴⁴Cm from 28.0 MWd/kg U Spent Fuel in Various Solutions at 25°C (Katayama, Bradley and Harvey 1980a)

attribute this to the radiolysis-induced formation of nitric acid (with nitrogen coming from air). The pH drop, which enhances the solubility of UO_2 , was not so dramatic in the other solutions because they were buffered by other constituents. Johnson et al. (1981) point out that the results of spent fuel radionuclide release experiments with respect to uranium can be affected by the

degree to which uranium precipitates within the experimental system or adsorbs back onto the surface of the spent fuel if these phenomena are not accounted for.

The issue of how solution composition affects radionuclide release from spent fuel will probably not be resolved until there is a better understanding of two factors: 1) how the solutions are affected by the radiation field from spent fuel, and 2) what the disposition of uranium is in different solutions after it has been released from the waste form.

6.3 EFFECT OF TEMPERATURE ON RADIONUCLIDE RELEASE

Temperature is predicted to have very little effect on the solubility of UO_2 , and it has been observed to have little effect on release rates and cumulative release from spent fuel. Slight increases in release rates of certain elements have been observed in systems in which the temperature was increased during the course of experimental runs (Norris 1979b; Johnson et al. 1982). The increased rates, however, rapidly decayed to rates that were the same as or below the rates observed before the temperature was increased. The cumulative release in these experiments was not noticeably affected by the temperature changes, which is in agreement with the work of Barner et al. (1986), who studied cumulative release from spent fuel in brines at $25^{\circ}C$ and $75^{\circ}C$.

6.4 EFFECT OF pH ON RADIONUCLIDE RELEASE

The effect of pH on radionuclide release from spent fuel has not been studied in detail because most spent fuel studies have been done using deionized water or simulated ground waters at their natural pH. Results from experimental studies of UO_2 dissolution and uranium solubility and from the theoretical work discussed in Section 5.0, however, suggest that UO_2 dissolution rates and uranium solubilities should be lowest in the pH range of 5 to 10. All of the solutions that have been used with spent fuel fall into this range.

Figure 6 shows uranium release rates from UO_2 as a function of pH as measured by Thomas and Till (1984). Figure 7 shows the measured solubility of uranium in 0.001 M NaCl as a function of pH (Silva and Yee 1981).



FIGURE 6. Fractional Release Rate of Uranium from Unirradiated U02 Pellets as a Function of pH (Thomas and Till 1984)

6.5 EFFECT OF OXYGEN CONCENTRATION AND EN ON RADIONUCLIDE RELEASE

Several investigators have studied the effect of oxygen concentration and Eh on radionuclide release from spent fuel and UO₂. The results indicate that release rates and total release for most elements are greater in oxidizing solutions than in reducing solutions. This has been attributed to the fact that under oxidizing conditions, the surface of the UO₂ matrix reacts to form higher oxides of uranium, which are much more soluble than UO₂ under most conditions. One of these higher oxide species, the uranyl ion (UO_2^{+2}) complexes



FIGURE 7. Uranium Solution Concentration as a Function of pH at 23°C. Points represent experimental results (Silva and Yee 1981)

readily with many anions. This species and its hydrolysis products are believed to be the most prevalent uranium species in solution under disposal conditions.

The release rate dependency on oxygen concentration is generally less than what would be expected if the dependence were first-order. This could possibly

be attributed to surface oxidation of the spent fuel or UO_2 during sample preparation, which would serve to passivate the surface from further oxidation and consequent dissolution.

Figure 8 shows the release rate of 90 Sr from spent fuel under oxidizing and reducing conditions in deionized water (Norris 1979b). Johnson (1982) observed similar results for 90 Sr in granite ground water. These results are typical for elements that are incorporated in the UO₂ fuel matrix as solid compounds. The results lend credence to the hypothesis that the release of these elements from spent fuel is controlled by the dissolution of UO₂.



FIGURE 8. Fractional Release Rate of ⁹⁰Sr from 28.0 MWd/kg U Spent Fuel Under Reducing and Oxidizing Conditions (Norris 1979b)

Figure 9 shows the release rate of 137 Cs from spent fuel as measured under oxidizing and reducing conditions (Norris 1979b). The fact that there is very little difference between the release rates under the two conditions suggests that the release of cesium is not controlled by the dissolution of UO₂; this is the expected result because cesium does not go into solid solution with UO₂.



FIGURE 9. Fractional Release Rate of 137Cs from 28.0 MWd/kg U Spent Fuel Under Reducing and Oxidizing Conditions (Norris 1979b)

6.6 EFFECT OF FUEL IRRADIATION HISTORY ON RADIONUCLIDE RELEASE

The effect of fuel irradiation history (burnup and linear power rating) on radionuclide release from spent fuel has not been studied in great detail. Katayama and Bradley (1980) conducted modified IAEA tests on fuels with burnups of 9.0, 28.0 and 54.5 MWd/kg U, and they concluded that there was no discernable effect of burnup on long-term release rate. When plotted as fractional release rate versus time, their data suggest that while the two lower burnup fuels released radionuclides at about the same rate, the higher burnup fuel appeared to release them slightly faster. The faster fractional release rate of the higher burnup fuel could have been due to a larger surface area, however, because the reported surface area per unit mass of the higher burnup fuel was about twice that of the two lower burnup fuels. The accuracy of the reported surface areas is questionable, because these values were estimated from pictures taken through a microscope at low magnification.

Katayama and Bradley also observed an accelerated release period that occurred between 600 and 800 days into the test for the 54.5 MWd/kg U fuel. During this period, the release rate of all elements became nearly equal and was anywhere from one to four orders of magnitude higher than the release rates before and after the accelerated period. This phenomenon was observed (in duplicate runs) only in deionized water, and it was not observed at all for fuels of lower burnup. Apparently the higher burnup fuel underwent a change of release mechanism in deionized water that it did not undergo in other solutions (and that was not observed in lower burnup fuel). A full explanation for this has not been offered. Figure 10 shows the data of Katayama and Bradley (1980) for the release rate of 137 Cs from the three different spent fuels. The release rates of other elements exhibited similar behavior.

The Canadians and Swedes have both investigated the effect of linear power rating on the release of radionuclides from spent fuel (Johnson et al. 1982; Eklund and Forsyth 1978). The results indicate that the release of radionuclides that do not go into solid solution with UO_2 (e.g., cesium and iodine) are enhanced by a higher power rating, while the release of radionuclides that go into solid solution with UO_2 are much less affected by power rating. An



FIGURE 10. Fractional Release Rate of ¹³⁷Cs from Spent Fuels with Burnups of 9.0, 28.0, and 54.5 MWd/kg U in Deionized Water at 25°C (Katayama and Bradley 1980). The reported surfacearea-per-unit-mass of the three fuel samples were 1.16, 0.99, and 2.28 cm²/g for the 9.0, 28.0 and 54.5 MWd/kgU fuels, respectively.

exception to this was observed by Eklund and Forsyth (1978) for 90 Sr, which exhibited a higher release rate from fuel with a higher power rating even though strontium is known to form oxides in the UO₂ matrix.

Some of the observed dependence of release rates on linear power rating may be caused by differences in the surface area of the various fuel samples used in the experiments. Unfortunately, neither the Canadians nor the Swedes reported the surface areas of their samples. The fact that the release of

radionuclides that do not go into solid solution with UO_2 is dependent on power rating, however, can also be explained by the tendency of these elements to migrate down the radial temperature gradient of the fuel during irradiation. A fuel with a higher power rating generally has a larger radial temperature gradient, which serves to enhance the migration of elements such as cesium and iodine to the grain boundaries and exposed surfaces of the fuel where they can be readily dissolved. Figure 11 shows the release rate of 137 Cs from Canadian spent fuels of two different power ratings (Johnson et al. 1982).



FIGURE 11. Fractional Release Rate of 137Cs from Spent Fuels with Moderate and High Power Ratings (Johnson et al. 1982). Surface areas were not reported.

6.7 CREDIT FOR CLADDING CONTAINMENT

A series of studies has been conducted at Westinghouse Hanford Company with the intent of showing that the cladding that surrounds spent fuel rods can significantly hinder the release of radionuclides from the spent fuel, even when the cladding is defected (Wilson and Oversby 1984, 1985; Wilson 1985). Defects in the cladding can occur during reactor operation or during postirradiation handling. This work is being done for the tuff repository project, which is managed by the Nevada Nuclear Waste Storage Investigations (NNWSI). Dissolution tests have been performed on sections of clad spent fuel rods that have been capped on the ends with stainless steel fittings. The specimens have had cladding in the following conditions:

- No defects (cladding completely intact)
- Two 0.2-mm laser-drilled holes in the cladding
- One slit defect cut in the cladding, 2 to 2.5 cm long by 0.15 to 0.2 mm wide
- Cladding completely removed.

The results of these tests have indicated that the defected cladding does hinder the release of uranium and plutonium to the test solutions, but that it has little effect on the release of cesium to the solutions (except when the cladding is intact, in which case essentially no cesium is released). The defected cladding provided retardation of the release of cesium, but after 30 days the amount of cesium in solution was about the same as in the cases where the cladding was completely removed. The tests simulated low repository flow rates; each sample of solution. Figures 12 and 13 show the fraction of total uranium and cesium inventory in solution at each sampling time for each of the four types of specimens. Well water from the Nevada repository site was used as the solution in these tests.

The main problem with taking credit for cladding containment in predicting the release of radionuclides from spent fuel is that it is very difficult to characterize the disposal condition of the cladding. There is currently no way of predicting a priori what types of defects the cladding will have or what



FIGURE 12. Fraction of Uranium Inventory in Solution for 31 MWd/kg U PWR Spent Fuel with Various Cladding Defects in Yucca Mountain Well Water at 25°C (Wilson and Oversby 1985)

percentage of fuel rods will have these defects. For this reason, most performance assessment studies of spent fuel waste packages have taken the conservative approach of assuming that all cladding has failed and that the waste form is essentially bare fuel.





6.8 MECHANISMS OF OXIDATION AND DISSOLUTION

Because the solubility of UO_2 in neutral and alkaline solutions is much lower under reducing conditions than under oxidizing conditions, it is widely believed that the dissolution of UO_2 involves first its oxidation and then the dissolution of the oxidized species.

Mechanisms of the oxidation and dissolution of spent fuel have been postulated mainly by experimentalists who have conducted electrochemical dissolution

studies on UO_2 . Wang (1981) proposed a reaction scheme for the dissolution of UO_2 that is summarized in Figure 14. This scheme involves the oxidation of UO_2 to UO_{2+x} species (where $0 < x \leq 1$) followed by the formation of uranyl ions (UO_2^{+2}) . The uranyl ions are the species that actually go into solution. Wang's reaction scheme accounts for the formation of a UO_3 hydrate film on the surface of the UO_2 , which has been observed in other studies (Sunder et al. 1982). This suggests that UO_2 may continue to dissolve and precipitate even after the solubilities of the uranyl ion and other soluble uranium species have been reached. The end result of this is that radionuclides contained in the UO_2 matrix that are more soluble than UO_2 may continue to be released even after the solution is saturated with respect to uranium.

A team of Canadian investigators conducted studies combining electrochemical techniques with X-ray photoelectron spectroscopy, which allowed identification of different oxides of uranium on the surface of the UO_2 (Sunder et al. 1981, 1982, 1983; Shoesmith et al. 1983, 1984). Using these techniques, the Canadians were able to follow the oxidation of UO_2 through various stages and determine at approximately which stage dissolution began in their experiments. Their results suggest that dissolution does not become prevalent until the surface of the UO_2 has been oxidized to a composition corresponding to $UO_{2.33}$ (U_3O_7). They have proposed that the uranyl ion is the active intermediate in the dissolution of uranium. Figure 15 summarizes the various stages of UO_2 oxidation that they believe occurred under neutral and alkaline conditions in their experiments.

The results of the electrochemical/surface analysis studies discussed above suggest that spent fuel should be much more resistant to dissolution in reducing environments than in oxidizing environments, and that dissolution should be minimized under conditions where UO_2 is not oxidized beyond the $UO_{2.33}$ state. Also, since the uranyl ion readily complexes with a number of anions (OH⁻, HCO₃⁻, CO₃⁻², F⁻, Cl⁻, SO₄⁻², H₂PO₄⁻², HPO₄⁻², and PO₄⁻³), the dissolution of UO₂ can be expected to be enhanced by the presence of these anions.



Reactions

(1)	Surface Uxidation (several angstroms)	
	$UO_2 + x/2 O_2 \longrightarrow UO_{2+x}$	0 < x ≤ 1
(2)	Oxidation-Dissolution	
	$UO_{2+x} + 2H^+ + (1-x)/2 O_2 \longrightarrow UO_2^{+2} + H_2O_1$	pH < 4
	$UO_{2+x} + H^+ + (1-x)/2 O_2 \longrightarrow UO_2(OH)^+,$	4 ≤ pH ≤ 7
	$UO_{2+x} + H_{2}O + (1-x)/2 O_{2} \longrightarrow UO_{2}(OH)_{2}^{0}$	pH ≥ 7
(3)	Transport	
(4)	HydrolysisFilm Formation, 25 to 75°C	
	$UO_2^{+2} + 3H_2O \longrightarrow UO_3 \cdot 2H_2O + H^+,$	pH ≤ 4
	$UO_2(OH)^+ + 2H_2O \longrightarrow UO_3 \cdot 2H_2O + H^+,$	4 ≤ pH ≤ 7
	$UO_2(OH)_2^0 + H_2O \longrightarrow UO_3 \cdot 2H_2O$,	pH ≥ 7
(5)	HydrolysisCrystal Growth, 150°C	
	$UO_2^{+2} + 2H_2O = UO_2(OH)_2 + 2H^+,$	pH ≤ 4
	$UO_2(OH)^+ + H_2O \implies UO_2(OH)_2 + H^+,$	4 ≤ pH ≤ 7
	$UO_2(OH)_2^0 \longrightarrow UO_2(OH)_2$	pH ≥7

<u>FIGURE 14</u>. Dissolution Mechanisms for UO_2 in Deionized Water (Wang 1981)



FIGURE 15. Reaction Scheme for the Anodic Oxidation of UO_2 (Sunder et al. 1982)

6.9 EFFECT OF RADIOLYSIS ON RADIONUCLIDE RELEASE

The effect of radiolysis on radionuclide release from spent fuel has been investigated from a theoretical standpoint, but very few experimental studies have been done to date. Alpha-radiolysis of water is expected to produce hydrogen peroxide (H_2O_2) and hydrogen (H_2) within a 30 µm layer surrounding the spent fuel surface (the alpha-particles are expected to penetrate the water only to this extent) (Draganic and Draganic 1971; Christensen and Bjergbakke 1982; Simonson and Kuhn 1984; Gray and Simonson 1985). Because hydrogen is relatively inert (in the absence of catalysts) at the temperatures expected under disposal conditions, and because hydrogen peroxide is a strong oxidizing agent, radiolysis is predicted to produce an oxidizing environment in the vicinity of the spent fuel surface. This oxidizing environment is predicted to enhance the dissolution of the spent fuel, which should also enhance the release of radionuclides from the spent fuel (Grenthe, Puigdomenech and Bruno 1983; Neretnieks 1984).

Other effects of radiolysis on spent fuel dissolution have also been investigated (Christensen and Bjergbakke 1984; Christensen 1984). If gaseous nitrogen or certain organic species are present in the radiation field, the formation of nitric or organic acids could result (Rai et al. 1980; Christensen 1984). This would drop the pH in the vicinity of the spent fuel surface, which would increase the solubility of UO_2 in the ground water. Beta-radiolysis may have a beneficial effect on the solution environment surrounding the spent fuel. Christensen and Bjergbakke (1984) have predicted that beta-radiolysis may help recombine some of the products of alpha-radiolysis, although this effect would probably be most prevalent outside the 30 μ m layer in which the alpha-radiolysis products are formed.

In a recent study done by the Canadians, a UO₂ electrode was exposed to an alpha source at various distances to get an idea of how radiolysis would affect the oxidizing potential of the electrode (Bailey, Johnson and Shoesmith 1985). It was found that the electrode potential was unaffected until the alpha source was within about 30 μ m of the surface, at which time the potential jumped to a very oxidizing value. Some surface oxidation of the electrode was observed, but the extent of this was not determined. The experimentalists believed that the main reaction occurring was the decomposition of hydrogen peroxide to water and O₂.

Differences between the dissolution rates of UO_2 and spent fuel under similar experimental conditions have been observed in some studies (Barner et al. 1985; Myers, Apted and Mazer 1984; Schramke, Simonson and Coles 1984). Although one would expect the dissolution rate to be greater from the spent fuel due to radiolysis effects, the results have been mixed. Barner et al. (1985) observed that the total release of uranium from spent fuel in brine was over 100 times greater than the uranium released from UO_2 in the same solution. Results from Myers et al. (1984) and Schramke, Simonson and Coles (1984), however, indicate that the uranium released from simulated spent fuel in basalt ground water was almost an order magnitude greater than the release from spent fuel. Clearly radiolysis cannot be used to explain these results, because the simulated spent fuel does not have a radiation field associated with it.

6.10 EFFECT OF WASTE PACKAGE CONSTITUENTS ON RADIONUCLIDE RELEASE

The effect of the presence of waste package constituents such as iron, zircaloy, and packing materials (e.g., bentonite) on radionuclide release from spent fuel is of interest because it is widely believed that these constituents will help retard the release of radionuclides from waste packages. In one of the few experimental studies that have been done in this area, Barner et al. (1985) have conducted tests on UO_2 and spent fuel in brines in the presence of iron and oxidized zircaloy.

In these tests, uranium, plutonium, and technetium were all observed to precipitate in the presence of iron in brine solutions. The iron did not affect the total release of these elements, but it drastically reduced the concentrations of these elements in the brine. Cesium did not exhibit this behavior in the presence of iron. Another finding was that the presence or absence of oxidized zircaloy appeared to have no affect on the disposition of any of the radionuclides in brine.

The fact that the presence of iron caused precipitation of the actinides and technetium can be attributed to two factors: 1) the reducing characteristics of ferrous iron (Fe^{+2}), which reduce the radionuclides to less soluble oxidation states, and/or 2) the selective sorption of the multivalent ions onto the iron or container walls. These characteristics make iron (particularly in lower oxidation states) a desirable component of canisters and of packing materials such as bentonite. (The net effect of the iron is to retard the migration of radionuclides by dropping them out of solution.) The ferrous iron that naturally occurs in basalt is expected to have similar beneficial effects on the performance of the waste package. The iron in basalt may explain why the release of uranium from spent fuel was no greater than the release of uranium from simulated spent fuel in the presence of basalt in the Myers, Apted and Mazer (1984) and the Schramke, Simonson and Coles (1984) experiments, even though radiolysis was occurring in the case of spent fuel (i.e., the oxidants formed by radiolysis may have been reduced by the iron). The fact that the disposition of cesium is not affected by the presence of iron can be attributed to its high solubility and lack of redox behavior.
From the oxidized zircaloy results, it can be concluded that zircaloy does not appreciably reduce the radionuclides and that the radionuclides do not adsorb to the surface of the zircaloy. Zircaloy, therefore, can not be expected to beneficially affect the performance of the waste package other than to provide physical containment of the radionuclides prior to cladding failure (i.e., rupture).

6.11 SORPTION, PRECIPITATION, COLLOIDS, AND THE FORMATION OF ALTERATION PHASES

Probably the most complex and least understood aspect of radionuclide release from spent fuel (and the dissolution of UO_2) is the disposition of the radionuclides after being released from the spent fuel. Once released, the radionuclides are free to interact with waste package constituents (see Section 6.10) and the host rock. Various radionuclides have been observed to 1) adsorb to solid surfaces, 2) precipitate from solution (sometimes only on certain surfaces), 3) combine with constituents in solution or in the host rock to form thermodynamically stable phases, and 4) form colloids with other species in solution. The first three effects are generally beneficial to the performance of the spent fuel waste package, but the problem of how to quantify them so that credit can be taken for them remains unsolved. Indeed these effects may be responsible for some of the disagreement in the release rate data from different radionuclide release experiments.

"Plateout", a term used to describe the sorption or precipitation of an element onto a solid surface, has been observed in many of the radionuclide release experiments described in this report. Differences between the activities of radionuclides in filtered and unfiltered solutions have also been observed, suggesting the formation of filterable colloids or suspended precipitates containing radionuclides. Unfortunately, plateout inventory and colloidal inventory have not always been accounted for in experiments or in the reporting of results, which makes comparison between experiments difficult. In some cases, the plated out and colloidal material have been combined with the material in solution to give the total release from the waste form. In other

cases, the amount of material accumulated on each surface and in the solution as both dissolved species and filterable species has been meticulously measured and separately reported.

Some general observations regarding the disposition of radionuclides after being released from spent fuel are listed below. (These are in addition to the observations made by Barner et al. (1985) in their study of the effects of iron and zircaloy.)

- 1. Uranium has been observed to plate out on container walls in a number of spent fuel release tests (see Table 2).
- Uranium, plutonium, americium, and curium have all been observed to plate out on quartz in tuff ground water (Wilson and Oversby 1985). Technetium and cesium plated out to a much lesser extent in this solution.
- 3. The alteration phases coffinite $(U(SiO_4)_{1-x}(OH)_{4x})$ and weeksite and/or boltwoodite (hydrated potassium-uranium-silicate phases) have been observed to form in autoclave tests on simulated spent fuel at 300° C in the presence of basalt rock (Myers, Apted and Mazer 1984). Coffinite is predicted to be the most stable phase in the simulated spent fuel/basalt system, but weeksite/boltwoodite was observed in greater quantities, suggesting that it may be an intermediate phase (Myers, Apted and Mazer 1984).

The formation of stable alteration phases is a thermodynamically driven process that is summarized by the expression (Giggenbach 1981):

Initial Solution + Primary (unstable) Phases ---> Final Solution + Secondary (stable) Phases

The reactions represented by this expression may be difficult to characterize because the kinetics can be very slow and because the reactions may proceed through a series of metastable intermediates that persist for indefinite periods of time. Excellent discussions of some of the thermodynamic and kinetic considerations that are important in predicting radionuclide release from waste forms are given by Apted (1982) and Myers, Apted and Mazer (1984). These discussions reference a number of studies of interactions of nuclear waste with surrounding rock and of fundamental geothermal equilibria (Garrels and Christ 1965; Helgeson 1968; Mottl and Holland 1978; Berner 1978; McCarthy et al. 1979; Seyfried and Bischoff 1979 and 1981; Giggenbach 1981; Wolery 1981; Dibble and Tiller 1981; Komarneni 1981; Freeborn et al. 1982).

Radionuclide sorption and colloid formation studies were not extensively reviewed for this report. We refer the reader to the Scientific Basis for Nuclear Waste Management series (Materials Research Society symposia proceedings) as a starting point for obtaining information on the progress of this work. ,

7.0. CHARACTERISTICS OF SPENT FUEL PERTINENT TO RADIONUCLIDE RELEASE

The characteristics of spent fuel expected to be important to the release of radionuclides from spent fuel are (Jenson 1982):

- the surface area of the spent fuel exposed to the dissolving solution
- the residence locations of the fission products and actinides and the phases that they exist in
- the grain boundary conditions of the fuel
- the surface oxidation state of the fuel.

The surface area is a function mainly of the irradiation and thermal history of the fuel. Thermal cycling and the development of steep axial and radial thermal gradients during irradiation cause the fuel to crack, which generates more surface area.

The importance of the residence locations of the radionuclides and the phases that they exist in was discussed in Section 6.1. Radionuclides that form compounds (primarily oxides) that are soluble in the UO_2 matrix generally remain in the UO_2 matrix during irradiation. Nuclides that are not soluble in the UO_2 matrix tend to migrate down the temperature gradient to the fuel periphery or grain boundaries, making them more accessible to solutions. Exceptions to this have been observed, however (see some of the references in Appendix B).

The grain boundary conditions of the fuel are important because the grain boundaries may contain significant inventories of radionuclides that are not soluble in the UO_2 matrix. If the grains are tightly bound together, release of the radionuclides from the grain boundaries may be restricted; but if the grains are loosely bound and there is sufficient space between them to allow solution access, release may occur quite rapidly.

The oxidation state of the surface of the fuel is expected to be important to radionuclide release because UO_2 dissolution is believed to proceed by a mechanism that involves oxidation of the UO_2 prior to dissolution (see Section 6.8). The presence of an oxidized surface layer on the fuel may speed up

dissolution under oxidizing conditions because the fuel will already be oxidized to some extent before contacting the solution. Under reducing conditions, the oxidized surface layer may serve to passivate the surface of the fuel, thereby hindering release. Oxidation of the surface of the fuel can occur during pool storage or dry storage if the fuel cladding has been ruptured.

As part of this literature survey, a great deal of published material on the characterization of spent fuel has been compiled for the informal library of the Waste Package Program (WPP) modeling task at PNL. An annotated bibliography of these materials is given in Appendix B for the reader who wants more information concerning the characterization of spent fuel.

8.0 STATUS OF MODELING OF RADIONUCLIDE RELEASE FROM SPENT FUEL

Because so many aspects of radionuclide release from spent fuel are not yet fully understood, modeling of radionuclide release from spent fuel has usually been discussed in general terms rather than mathematical terms (e.g., Liebetrau and Apted 1985). Stochastic models of release from spent fuel for use in performance assessment studies have been proposed (Goodwin, Johnson and Wuschke 1983; Goodwin, Lemire and Johnson 1982), but because these models are parametric in nature, they do not properly describe the mechanistic processes of release.

Goodwin (1984), Allard (1984a and b), Skytte-Jensen (1984), and Johnson, Garisto and Stroes-Gascoyne (1985) have all discussed some of the complexities of modeling radionuclide release from spent fuel. In an early modeling attempt, Kuhn (1981) assumed (for computational purposes) that the release of uranium from spent fuel was controlled by uranium solubility and that all radionuclides were released congruently with uranium. Many investigators now believe that oxidation of the UO₂ matrix (as a precursor to dissolution) is the controlling process for uranium release, and it is widely recognized that the fission products 134 Cs, 137 Cs and 129 I tend to release more rapidly from spent fuel than other radionuclides (99 Tc may also be a rapidly releasing radionuclide). Long-term release, however, may still be controlled by thermodynamics if precipitates or stable alteration phases containing radionuclides form in the system.

Johnson and Joling (1984) have used a probability distribution function to predict the instant release fraction of cesium and iodine from the fuelcladding gap and grain boundaries of Canadian (CANDU) fuel. Their work suggests that the inventory of radionuclides at the grain boundaries is far more significant than the inventory in the gap. This point is illustrated graphically in Figure 16 (Johnson, Garisto and Stroes-Gascoyne 1985), which shows the release rate of radionuclides from the gap, grain boundaries, and grains of spent fuel as a function of time. It is assumed here that the UO_2 grains dissolve at a constant rate and that the radionuclides contained in the grains are released congruently from them. A major uncertainty that needs to be



FIGURE 16. Radionuclide Release from UO₂ Fuel in Water, Oxidizing Conditions (Johnson, Garisto and Stroes-Gascoyne 1985)

investigated is how quickly the radionuclides at the grain boundaries will become exposed to solution so that the soluble constituents can be dissolved.

Modeling of the release of radionuclides from spent fuel grains (i.e., the UO_2 matrix) is still in the conceptual stage. Since spent fuel is primarily UO_2 and hydrates slowly if not oxidized, the degradation of spent fuel is expected to be essentially the rate of oxidation of UO_2 . The resulting oxidized matrix hydrates and releases its radionuclides to varying extents. It appears that most radionuclides contained in spent fuel, including uranium, release roughly in proportion to the square root of time. This behavior is probably a result of release from the grain boundaries of the fuel (see Figure 16). The effect of an increased uranium concentration in the solution on the degradation rate of spent fuel is not known.

The effects of alpha-radiolysis on the release of radionuclides from spent fuel are not clearly understood. Some investigators believe that alpharadiolysis can promote the oxidation of spent fuel through the radiolytic

production of H_2O_2 and other oxidizing species in the solution adjacent to the surface of the spent fuel. Under conditions of low, buffered Eh, however, the radiolytically produced oxidants may be quickly reduced to species that do not affect the spent fuel. Even if radiolysis is capable of generating a locally oxidizing environment near the surface of spent fuel in an overall reducing environment, many of the radionuclides that are released as a result of being oxidized to more soluble valence states might revert to lower valence states and precipitate once they are transported out of the oxidizing environment near the surface.

The formation and degradation of alteration phases in the waste package environment is poorly understood. The rates of formation of these phases, the intermediates involved in this process, and the characteristics of the phases need to be studied in more detail before alteration phases can be accounted for in models.

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9.0 CONCLUSIONS REGARDING MODELING OF RADIONUCLIDE RELEASE FROM SPENT FUEL

Based on what has been learned in studies of radionuclide release from spent fuel and UO_2 , a complete model for the release of radionuclides from spent fuel should include:

- An instant release submodel describing the rate of release of radionuclides that have accumulated in the fuel-cladding gap and at the grain boundaries of the fuel.
- A kinetic submodel describing the rate of degradation (i.e., oxidation and dissolution) of the spent fuel matrix, which describes the rate at which many radionuclides become unbound from the spent fuel and available for transport.
- A post-release submodel, which describes the accumulation of radionuclides in solution, the formation of precipitates, colloids, and alteration phases, and the constraints on these processes.
- A mass transfer submodel, which describes the transport of the radionuclides away from the spent fuel according to the concentration of radionuclides in solution.

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APPENDIX A

DIGITIZED DATA SETS OF SPENT FUEL RELEASE

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APPENDIX A

DIGITIZED DATA SETS OF SPENT FUEL RELEASE

The data contained in the following tables have been digitized from graphs appearing in documents that report results of spent fuel radionuclide release experiments. References, graph symbols, labels, and units are included with each set of data. Conditions under which the data were taken are described in the references; brief descriptions of the experimental conditions are given in Table 2. Data that have already been tabulated in the literature have not been included in this appendix. The tables in this appendix are organized in chronological order; the earliest data appear first, and the most recent data appear last.

The digitizing system used an HP plotter and sight pen in conjunction with a program written in IBM Advanced Basic. Duplicate measurements have confirmed that the system reads data from the graphs within plus or minus 10%.

DIGITIZED DATA SETS

REFERENCE: EKLUND AND FORSYTH 1978, FIGURE 3

FIGURE TITLE: 'UTLAKNINGSHASTIGHET FOR URANIUM'

.

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 2

LABEL= GROUND WATER 11.2 kW/m

SYMBOL= FILLED CIRCLE

С	UR VI	E 1	

LABEL= DISTILLED WATER 11.2 kW/m SYMBOL= OPEN CIRCLE

X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
2	3.4E-06	2	3.1E-06
3	3.1E-06	3	2.4E-06
6	1.2E-06	6	1.4E-06
11	1.1E-06	11	1.5E-06
21	9.0E-07	21	1.1E-06
35	7.1E-07	35	1.0E-06
56	5.7E-07	56	8.1E-07
86	4.7E-07	86	6.7E-07

CURVE 3

LABEL= GROUND WATER 23.5 kW/m SYMBOL= FILLED TRIANGLE

X COORDINATE	Y COORDINATE
2	8.3E-05
3	3.4E-06
6	8.2E-07
11	4.7E-07
21	4.7E-07
35	5.1E-07
55	3.1E-07
86	3.0E-07

Y

CURVE 4

LABEL= DISTILLED WATER 23.5 kW/m SYMBOL= OPEN TRIANGLE

X COORDINATE	Y COORDINATE
2	6.4E-05
3	3.4E-06 1.4E-06
21	1.5E-06 1.3E-06
35 55	1.2E∸06 8.7E-07
86	6.6E-07

REFERENCE: EKLUND AND FORSYTH 1978, FIGURE 4

FIGURE TITLE: 'UTLAKNINGSHASTIGHET FOR ⁹⁰Sr'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES	LABEL	IS	'FRACTIONAL
			RELEASE
			RATE
			(1/DAYS)'

CURVE 1 LABEL= DISTILLED WATER 11.2 kW/m SYMBOL= OPEN CIRCLE		CURVE 2		
		LABEL= GROUND WATER 11.2 kW SYMBOL= FILLED CIRCLE		
X	Y	X	Y	
COORDINATE	COORDINATE	COORDINATE	COORDINATE	
2	3.3F-05	2	3.7F-06	
3	3.7E-06	6	2.5E-06	
6	1.0E-06	11	6.8E-07	
11	9.3E-07	20	4.4E-07	
20	4.7E-07	33	3.2E-07	
33	4.8E-07	52	3.4E-07	
52	3.4E-07	82	2.4E-07	
81	2.4E-07			

CURVE 3

LABEL= DISTILLED WATER 23.5 kW/m SYMBOL= OPEN TRIANGLE

UURVE (4
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LABEL= GROUND WATER 23.5 kW/m SYMBOL= FILLED TRIANGLE

X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
2	1.0E-04	2	1.3E-04
6	3.8E-06	· <u>3</u>	1.2E-05
10	3.1E-06	6	9.0E-06
20	2.5E-06	11	9.8E-06
33	1.9E-06	20	7.4E-06
53	1.2E-06	33	2.4E-06
82	7.7E-07	52	1.5E-06
		82	6.7E-07
DRAWING NO. = TP	M-BL-133		

REFERENCE: EKLUND AND FORSYTH 1978, FIGURE 5

FIGURE TITLE: 'UTLAKNINGSHASTIGHET FOR ¹³⁷Cs'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1 LABEL= DISTILLED WATER 11.2 kW/m SYMBOL= OPEN CIRCLE		CURVE 2			
		LABEL= GROUND WATER 11.2 kW/m SYMBOL= FILLED CIRCLE			
	X COORDINATE	Y COORDINATE	X <u>COORDINATE</u>	Y COORDINATE	
	1	1.1E-04 6.1E-06	1	1.2E-04	
	5	3.3E-06 2.5E-06	5	2.3E-06	
	20 33	1.9E-06 1.6E-06	20 33	1.0E-06	
	52 82	1.4E-06 1.3E-06	52 82	1.5E-06 1.8E-06	

CI	ID	VC	2
ີບເ	ж	V C.	ు

LABEL= DISTILLED WATER 25.5 kW/m SYMBOL= OPEN TRIANGLE

CI	JR	VE	4
\mathbf{u}	<i>J</i> IN		

LABEL= GROUND WATER 23.5 kW/m SYMBOL= FILLED TRIANGLE

Х	Y	Х	Y
COORDINATE	COORDINATE	COORDINATE	COORDINATE
1	0 OF 01	1	0 05 04
1	5.92-04	1	9.95-04
Z	5.UE-U4	2	4.8E-04
5	3.4E-04	5	3.1E-04
10	1.9E-04	10	2.1E-04
20	2.7E-05	20	7.6E-05
33	7.8E-06	33	1.5E-05
52	3.9E-06	53	3.0E-06
82	3.5E-06	82	• 4E 06

REFERENCE: VANDERGRAAF 1980, p. 44

FIGURE TITLE: 'DISSOLUTION RESULTS FOR URANIUM AS A FUNCTION OF TIME'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

	C	URVE	1		
LABEL= DIST SYMBOL= OPE	ILLED WATE N CIRCLE	R			
X COORDINATE	Y COORDINAT	<u>E</u>		X COORDINATE	Y COORDINATE
16	1.2E-05			440	1.4E-07
25	6.3E-05			460	1.9E-07
87	7.9E-06			460	1.4E-07
93	3.6E-06			770	5.3E-07
140	3.1E-06			790	1.3E-06
200	8.2E-05			790	2.0E-07
290	3.6E-05			810	3.6E-06
290	2.2E-06			810	3.6E-07
300	1.4E-06			820	2.5E-06
300	3.9E-05			820	1.8E-07
310	1.8E-04			840	3.3E-06
300	9.4E-07			840	1.4E-07
330	7.1E-07			860	2.4E-06
350	5.3E-06			860	1.8E-07
350	3.2E-06			870	6.5E-07
360	2.8E-06			870	9.0E-08
400	6.9E-07			890	6.6E-07
400	4.5E-07			890	5.6E-07
420	1.0E-05				

REFERENCE: VANDERGRAAF 1980, p. 44 (contd)

CURVE 2

LABEL= CHLORINATED RIVER WATER SYMBOL= FILLED TRIANGLE

X	Y	Х	Ŷ
COORDINATE	COORDINATE	COORDINATE	COORDINATE
1	1.0E-04	403	6.5E-06
1	5.9E-05	403	3.1E-06
8	7.5E-05	422	3.5E-06
16	1.5E-05	422	2.4E-06
65	1.1E-05	439	1.8E-06
79	9.2E-06	439	1.5E-06
94	1.0E-04	454	1.7E-06
141	2.1E-05	454	1.3E-06
202	1.0E-05	768	2.9E-06
290	1.0E-05	768	2.3E-06
290	8.7E-06	791	7.1E-06
297	5.0E-03	791	4.8E-06
298	7.7E-05	811	3.0E-06
301	3.0E-05	822	4.4E-06
301	1.3E-05	839	7.9E-06
316	1.3E-05	839	5.4E-06
312	9.1E-07	856	6.0E-06
335	6.0E-06	856	4.4E-06
337	4.9E-06	868	1.2E-05
346	1.1E-05	867	4.8E-06
346	5.8E-06	887	8.2E-06
363	6.4E-06	887	4.1E-06
363	5.4E-06		

REFERENCE: VANDERGRAAF 1980, p. 38

FIGURE TITLE: 'LEACHING RESULTS FOR ¹³⁷Cs AS A FUNCTION OF TIME'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'Bq (1/DAYS)'

CURVE 1

LABEL=	DISTIL	LED	WATER
SYMBOL=	OPEN	CIRC	CLE

Х	Y	Х	Y	Х	Y
COORDINATE	COORDINATE	COORDINATE	COORDINATE	COORDINATE	COORDINATE
0	1.1E+09	90	1.1E+06	368	6.5E+04
0	7.3E+08	90	8.5E+05	383	1.1E+05
0	3.7E+08	96	9.5E+05	408	1.5E+05
0	2.3E+08	96	1.4E+06	428	7.3E+04
0	1.0E+08	135	8.7E+05	428	2.5E+04
0	6.4E+07	136	6.3E+05	443	2.3E+05
10	3.3E+07	144	1.0E+06	443	1.7E+05
10	2.2E+07	145	7.4E+05	455	1.5E+05
12	2.1E+07	193	8.7E+05	455	6.3E+04
12	1.4E+07	193	3.9E+05	500	8.1E+04
17	1.5E+07	205	3.8E+06	500	4.6E+04
21	1.3E+07	205	8.7E+05	561	9.6E+04
21	1.1E+07	286	4.1E+05	561	6.4E+04
25	1.2E+07	287	1.4E+05	617	7.2E+04
29	1.0E+07	293	8.6E+05	618	6.0E+04
29	9.7E+06	293	4.5E+05	775	8.2E+04
33	9.6E+06	303	5.6E+05	800	8.4E+04
32	8.7E+06	303	3.9E+05	800	5.2E+04
38	7.5E+06	306	6.8E+05	819	1.3E+05
51	5.3E+06	306	3.5E+05	819	1.1E+05
51	4.2E+06	315	1.1E+06	832	1.6E+05
58	3.9E+06	315	3.1E+05	832	5.1E+4
58	2.3E+06	321	7.3E+05	848	3.4E+4
67	3.2E+06	323	2.8E+05	848	2.0E+4
67	2.1E+06	338	6.4E+05	866	1.0E+5
75	1.9E+06	338	2.1E+05	864	8.7E+4
74	1.3E+06	350	3.6E+05	876	7.9E+4
78	3.3E+06	350	7.3E+05	878	3.5E+4
78	2.4E+06	366	2.5E+05	905	9.8E+4
				905	8.7E+4

REFERENCE: VANDERGRAAF 1980 p. 38 (contd)

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	CURVE	2		
LABEL= TAP WATER SYMBOL= FILLED TRIANGLE				
X COORDINA	Y TE COORDINATE	X COORDINAT	Y <u>E COORDINATE</u>	
COORDINA 0 0 0 0 0 0 10 10 10 10 10 14 20 30 38 50 58 56 67 67 67 73 73 73 73 73 73 73 91 91 95 95	$\begin{array}{c} \underline{\text{TE} \ \text{COORDINATE}} \\ 6.1E+08 \\ 5.0E+08 \\ 2.0E+08 \\ 1.8E+08 \\ 1.8E+08 \\ 1.1E+08 \\ 7.0E+07 \\ 2.9E+07 \\ 1.6E+07 \\ 1.6E+07 \\ 1.2E+07 \\ 1.0E+07 \\ 8.4E+06 \\ 8.4E+06 \\ 5.6E+06 \\ 5.6E+06 \\ 5.6E+06 \\ 5.6E+06 \\ 5.6E+06 \\ 5.6E+06 \\ 1.6E+06 \\ 1.6E+06 \\ 3.4E+06 \\ 2.9E+06 \\ 1.6E+06 \\ 2.0E+06 \\ \end{array}$	COORDINAT 306 315 322 322 339 351 366 366 382 384 407 407 427 427 427 427 427 427 443 456 456 499 499 561 561 617 617	E COORDINATE 4.6E+05 4.3E+05 5.0E+05 4.2E+05 5.3E+05 4.3E+05 4.3E+05 3.2E+05 3.7E+05 3.3E+05 4.0E+05 6.1E+04 3.6E+05 2.9E+05 3.2E+05 3.0E+05 2.6E+05 3.0E+05 3.1E+05 2.8E+05 3.4E+05 3.4E+05 2.8E+05 3.0E+05 3.4E+05 2.8E+05 3.0E+05 2.4E+05	
137 137 143 192 193 204 204 204 287 287 294 294 302 306	2.0E+06 1.0E+06 8.3E+05 6.6E+05 6.0E+05 3.0E+07 6.0E+05 6.4E+05 5.4E+05 5.3E+05 6.6E+05 5.4E+05 5.4E+05 5.4E+05	774 774 798 798 833 831 865 865 877 877 904 904	2.4E+05 2.8E+05 2.9E+05 2.9E+05 1.7E+05 8.2E+05 1.5E+05 1.6E+05 1.6E+05 1.0E+05 1.3E+05 1.0E+05	
REFERENCE: VANDERGRAAF 1980 p. 34

FIGURE TITLE: 'LEACHING RESULTS FOR ⁹⁰Sr AS A FUNCTION OF TIME'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'Bq (1/DAYS)'

	CUR	<u>RVE 1</u>	
LABEL= DIST SYMBOL= OPE	ILLED WATER N CIRCLE		
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
0 13 22 90 140 199 290 289 300 298 300 298 300 303 309 311 316 317	7.5E+05 5.2E+05 5.2E+05 7.7E+04 4.9E+06 3.0E+05 4.8E+04 7.8E+03 5.1E+04 6.8E+03 3.6E+04 5.4E+03 1.2E+05 6.9E+03 4.9E+04 4.7E+03	402 423 424 450 452 496 496 556 556 614 612 769 792 795 814 814	1.0E+03 7.7E+03 1.1E+03 7.2E+03 1.0E+03 3.6E+03 6.5E+02 3.2E+03 1.1E+02 8.9E+03 2.3E+03 1.2E+02 1.1E+03 3.2E+02 1.1E+03 3.2E+02 1.8E+03 7.3E+02
317 333 335 345 345 362 362 378 380 402	4.7E+03 4.3E+04 4.0E+03 2.0E+04 3.0E+03 2.2E+04 2.2E+03 1.9E+04 1.0E+03 6.2E+03	814 825 825 842 842 858 858 858 868 870	7.3E+02 1.3E+03 9.3E+02 1.9E+03 2.7E+02 1.8E+03 5.3E+02 1.0E+03 3.3E+02

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CURVE 2

LABEL = CHLORINA	TED RIVER	WATER
SYMBOL= FILLED	TRIANGLE	

Х	Ŷ	X	Ŷ
COORDINATE	COORDINATE	COORDINATE	COORDINATE
0	7.2E+06	378	1.7E+04
0	2.5E+06	403	1.7E+04
5	3.0E+05	403	1.5E+04
14	2.3E+05	421	1.5E+04
22	2.7E+05	421	1.3E+04
62	1.1E+05	439	1.0E+04
74	1.0E+05	438	9.4E+03
91	1.9E+05	450	1.0E+04
138	5.8E+04	496	1.2E+04
200	3.0E+04	556	1.2E+04
289	3.0E+04	556	1.0E+04
289	2.6E+04	597	5.7E+05
296	1.8E+05	611	1.0E+04
301	5.2E+04	769	9.9E+03
301	3.4E+04	769	7.7E+03
295	3.7E+06	792	1.4E+04
308	2.1E+04	792	1.0E+04
317	4.2E+04	813	1.2E+04
317	2.1E+04	813	8.9E+03
335	2.6E+04	824	1.0E+04
333	2.1E+04	841	8.2E+03
345	2.4E+04	841	5.7E+03
361	1.6E+04	857	5.5E+03
378	2.0E+04	869	8.3E+03
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REFERENCE: VANDERGRAAF 1980, p. 41

FIGURE TITLE: 'LEACHING RESULTS FOR 239Pu AND 240Pu AS A FUNCTION OF TIME'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'Bq (1/DAYS)'

CI	UR	VE.	- 1

LABEL= DISTILLED WATER SYMBOL= OPEN CIRCLE

X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
7	1.2E+02	429	7.7E+00	827	2.7E+00
25	9.5E+02	444	1.2E+01	845	7.4E+00
31	1.8E+03	444	6.5E+00	843	2.5E+00
209	4.8E+02	461	1.1E+01	860	1.1E+01
296	4.8E+02	772	1.3E+01	859	3.1E+00
304	5.6E+02	797	1.4E+01	873	4.6E+00
317	1.1E+03	797	1.2E+01	871	3.9E+00
325	7.5E+02	816	8.2E+00	899	1.4E+01
342	1.9E+01	816	6.3E+00	899	1.1E+01
385	1.9E+02	827	3.2E+00		

CURVE 2

LABEL = CHLORINATED RIVER WATER SYMBOL= FILLED TRIANGLE

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X	Y X	Y	X	Y
COORDINATE COORD	DINATE COORDI	NATE COORDINATE	COORDINAT	TE COORDINATE
7 1.88 17 1.68 27 8.88 33 1.58 75 2.48 00 2.38 150 4.78 210 1.98 297 2.18 302 6.38 305 1.68 308 5.68	+03 310 +03 316 +01 324 +03 342 +02 353 +03 409 +02 409 +02 430 +02 446 +04 461 +03 772 +02 797	3.1E+02 1.8E+02 3.1E+02 3.0E+04 1.1E+02 5.0E+01 2.2E+01 5.7E+01 5.6E+01 3.1E+00 3.5E+00 4.1E+00 2.2E+01	797 816 827 827 845 844 860 859 871 871 900 900	1.6E+01 1.3E+01 1.5E+01 8.9E+00 1.2E+01 7.7E+00 4.9E+00 1.9E+00 3.8E+01 3.3E+00 1.5E+01 1.5E+00

REFERENCE: VANDERGRAAF 1980, p. 39

FIGURE TITLE: 'LEACHING RESULTS FOR ¹⁴⁴Ce AS A FUNCTION OF TIME'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'Bq (1/DAYS)'

CURVE 1					
LABEL= D SYMBOL= (ISTILLED WATER OPEN CIRCLE				
X COORDINA	Y TE COORDINATE	X COORDINA	Y TE COORDINATE	X COORDINA	Y TE COORDINATE
1 1 8 8 8 8 14 14 14 14 25 19 19 23 26 30 30	1. 3E +06 3. 6E +05 3. 3E +04 1. 1E +04 8. 3E +03 5. 9E +03 4. 2E +04 3. 6E +04 3. 6E +04 6. 1E +01 3. 6E +04 2. 1E +04 1. 8E +04 2. 6E +04 2. 5E +04 2. 0E +04	55 55 64 64 70 70 78 78 82 82 96 96 100 138 138 138 146	1.0E+05 1.0E+04 2.0E+04 1.6E+04 7.4E+04 4.6E+04 1.0E+04 5.8E+03 1.5E+04 1.0E+04 1.0E+04 9.4E+03 1.0E+04 3.3E+03 2.9E+03 9.3E+03	304 307 307 314 316 406 406 426 426 426 443 443 458 458 458 499 499 558	3.3E+03 2.0E+04 2.6E+03 1.0E+05 6.4E+03 2.2E+02 1.3E+02 4.2E+02 2.6E+03 1.1E+02 6.4E+02 1.0E+03 4.2E+02 1.6E+02 1.6E+02 6.7E+01 2.7E+02
35 35 39 39 34 43 43	2.6E+04 1.4E+04 1.3E+04 2.9E+04 2.7E+04 1.5E+04 3.4E+04 2.3E+04	146 193 195 207 286 294 294 304	6.7E+03 4.3E+04 2.7E+03 5.9E+03 1.3E+03 7.5E+03 4.6E+03	558 615 614 769 793 814 814	2.0E+02 3.4E+02 1.3E+02 2.4E+02 1.1E+02 1.2E+02 5.0E+01

REFERENCE: VANDERGRAAF 1980, p. 39 (contd)

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CURVE 2

LABEL = CHLORINATED RIVER WATER SYMBOL = FILLED TRIANGLE

X	Ŷ	Х	Y
COORDINATE	COORDINATE	COORDINATE	COORDINATE
3	7.2E+05	145	1.2E+04
3	1.2E+05	195	1.6E+03
7	2.5E+05	195	7.6E+04
7	1.1E+05	205	9.1E+02
7	8.3E+04	205	7.1E+03
7	8.1E+03	288	7.9E+02
7	7.4E+03	288	3.7E+02
13	9.1E+04	295	5.4E+03
18	1.2E+05	295	3.2E+03
18	7.9E+04	302	3.2E+04
20	2.8E+04	302	2.2E+04
20	1.8E+04	306	5.2E+04
15	1.6E+04	307	4.8E+03
23	1.0E+04	315	1.5E+04
32	3.3E+04	322	2.9E+03
32	1.2E+04	338	1.7E+03
35	6.3E+03	406	2.5E+03
38	1.2E+04	406	1.3E+03
63	1.2E+04	427	4.2E+03
64	9.0E+03	428	1.2E+03
71	2.8E+04	443	3.3E+03
71	6.4E+03	443	1.4E+03
71	2.8E+04	459	1.2E+03
79	5.1E+03	459	9.4E+04
83	7.1E+03	458	1.2E+03
83	4.3E+03	498	3.1E+02
94	5.9E+03	498	2.0E+02
94	4.9E+05	559	1.5E+02
98	1.2E+04	613	1.9E+02
98	1.6E+03	613	1.4E+02
138	1.0E+04	792	2.9E+02
139	7.1E+02	794	1.6E+02
148	1.7E+04	808	1.1E+02

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 11

FIGURE TITLE: 'LEACH RATES FOR URANIUM'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'LEACH RATE'

CURVE 1 LABEL= 85-18 URANIUM SYMBOL= OPEN SQUARE		CURVE 2 LABEL= 85-200 URANIUM SYMBOL= FILLED CIRCLE		
COORDINATE	COORDINATE	COORDINATE	COORDINATE	
1	6.2E-06	2	6,2E-07	
2	5.6E-06	7	5.2E-07	
4	4.3E-06	11	4.8E-07	
7	2.8E-06	14	4.2E-07	
11	2.3E-06	15	3.3E-07	
14	2.1E-06	18	2.5E-07	
15	1.5E-06	22	2.1E-07	
18	9.3E-07	25	2.0E-07	
22	5.8E-07	28	1.8E-07	
25	5.5E-07	33	1.4E-07	
28	5.4E-07	40	1.2E-07	
33	4.7E-07	48	1.1E-07	
40	3.9E-07	57	7.9E-08	
48	3.5E-07	75	5.3E-08	
57	3.0E-07	100	4.2E-08	
75	2.1E-07	130	3.8E-08	
100	1.4E-07	160	3.1E-08	
130	1.0E-07	210	2.0E-08	
160	7.0E-08	280	1.4E-08	
210	4.1E-08			
280	2.1E-08			

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REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 11 (contd)

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CURVE 3

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CURVE 4

LABEL=	100-18	URANIUM
SYMBOL=	OPEN C	CIRCLE

LABEL=	100-200	URANIUM
SYMBOL=	FILLED	SQUARE

X COORDINATE	Y COORDINATE
1 5 11 14 15 18 26 33 40 48 57 75 100 130 160 210 280	2.4E-06 8.6E-07 4.5E-07 3.0E-07 1.9E-07 1.4E-07 1.2E-07 9.4E-08 7.8E-08 7.6E-08 6.4E-08 3.9E-08 2.2E-08 1.8E-08 1.8E-08 1.3E-08 5.6E-09 2.9E-09

X	Ŷ
COORDINATE	COORDINATE
1	5.6E-07
5	5.1E-07
11	3.5E-07
14	2.6E-07
15	2.2E-07
18	2.0E-07
22	2.1E-07
23	2.5E-07
26	2.3E-07
29	1.6E-07
33	8.5E-08
40	5.2E-08
48	4.4E-08
57	4.1E-08
75	3.4E-08
100	2.2E-08
130	1.6E-08
160	1.3E-08
210	1.2E-08
280	1.2E-08

DRAWING NO. = ORNL-DWG-80-1454

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 14 FIGURE TITLE: 'LEACH RATES FOR 144Ce'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'LEACH RATE'

CURVE	<u> 1 </u>	CURV	Ξ 2
LABEL= 85-18 144 SYMBOL= OPEN SQUA	Ce ARE	LABEL= 85-200 ^{14.} SYMBOL= FILLED C	⁴ Ce IRCLE
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
1 4 5 8 11 15 21 25	2.3E-06 2.4E-06 2.2E-06 1.3E-06 5.4E-07 1.9E-07 8.9E-08 6.5E-08	4 6 7 9 10 15 21 25 30 38 45 55 72 97	2.4E-07 2.2E-07 1.5E-07 9.0E-08 5.7E-08 4.0E-08 2.8E-08 2.2E-08 2.2E-08 2.3E-08 2.3E-08 1.8E-08 1.8E-08 1.1E-08 9.4E-09
		162 211 278	1.2E-08 1.2E-08 1.4E-08 1.4E-08

A.16

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 14 (contd)

CURVE	E 3		4
LABEL= 100-18 ¹⁴⁴ SYMBOL= OPEN CIRC	⁴ Ce CLE	LABEL= 100-200 ¹⁴ SYMBOL= FILLED SC	¹⁴ Ce QUARE
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
0	4.9E-06	2	1.0E-06
4	2.9E-06	5	1.0E-06
6	1.3E-06	9	1.0E-06
7	8.1E-07	10	5.7E-07
10	6.2E-07	14	2.4E-07
11	5.0E-07	16	1.6E-07
14	3.1E-07	18	1.3E-07
18	1.5E-07	19	1.2E-07
20	1.1E-07	25	1.1E-07
23	1.0E-07	28	6.6E-08
24	7.3E-08	32	3.9E-08
28	4.6E-08	39	3.5E-08
40	3.0E-08	51	4.1E-08
73	2.5E-08	72	3.8E-08
25	2.4E-08	99	2.1E-08
		137	1.1E-08
		189	8.4E-09

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DRAWING NO. ORNL-DWG-80-1456

A.17

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 15 FIGURE TITLE: 'LEACH RATES FOR ¹⁵⁴Eu'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'LEACH RATE'

CURVE 1	CURVE 2
LABEL= 85-18 ¹⁵⁴ Eu	LABEL= 85-200 ¹⁵⁴ Eu
SYMBOL= OPEN SQUARE	SYMBOL= FILLED CIRCLE

X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
2	6.2E-06	.5	1.3E-06
8	6.2E-06	7	8.6E-07
11	6.3E-06	9	3.6E-07
		10	2.1E-07
		20	1.3E-07
		48	4.7E-08
		91	1.9E-08
	•	127	1.5E-08
		161	1.5E-08
		210	1.4E-08
		277	1.4E-08

CURVE 3

LABEL= 100-18 ¹⁵⁴Eu SYMBOL = OPEN CIRCLE

Х	Y
COORDINATE	COORDINATE
3	3.0E-06
5	1.8E-06
8	1.2E-06
10	9.5E-07
10	6.3E-07
13	4.4E-07
19	2.7E-07
27	1.2E-07
41	5.2E-08
65	2.9E-08

CU	IR	VE	4
CU	IK	٧Ŀ	- 4

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LABEL= 100-200 ¹⁵⁴Eu SYMBOL= FILLED SQUARE

Х	Y
COORDINATE	COORDINATE
3	1.2E-06
4	9.3E-07
9	6.7E-07
10	3.7E-07
12	2.5E-07
14	2.2E-07
16	1.6E-07
19	1.2F-07
21	1.1F-07
30	9 OF -08
38	5 9F-08
51	
71	4.0E-00
/1	3.2E-08
98	1.4E-08
136	1.0E-08
188	9.6E-09

DRAWING NO. = ORNL-DWG-80-1457

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 17

FIGURE TITLE: 'LEACH RATES FOR 137Cs'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'LEACH RATE'

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CURVE 1 CUR		<u>/E 2</u>	
LABEL= 85-18 137 SYMBOL= OPEN SQU	7CsLABEL= 85-200137CsJARESYMBOL= FILLED CIRCLE		7 _{Cs} IRCLE
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
1 3 6 7 9 10 12 14 16 19 21 24 25 29 34 40 48 58 75 100 129	2.7E-04 9.0E-05 3.7E-05 1.8E-05 1.0E-05 8.6E-06 7.2E-06 6.2E-06 3.5E-06 2.7E-06 2.3E-06 1.9E-06 1.6E-06 1.6E-06 1.4E-06 1.2E-07 6.3E-07 5.1E-07 4.7E-07	2 5 7 9 10 11 14 16 19 21 24 25 29 34 40 48 58 75 100 129 162	6.3E-05 2.0E-06 1.1E-06 8.3E-07 7.3E-07 7.0E-07 6.3E-07 5.9E-07 5.9E-07 3.6E-07 3.6E-07 3.6E-07 3.6E-07 3.5E-07 3.5E-07 1.7E-07 1.4E-07 1.6E-07 1.6E-07
210 278	4.3E-07 2.6E-07 1.1E-07	210 278	1.6E-07 1.6E-07

A.19

REFERENCE: MITCHELL	, GOODE	AND	VAUGHEN	1981,	p.	17	(contd))
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CURVE	E 3 ·	CURV	E 4
LABEL= 100-18 13 SYMBOL= OPEN CIRC	7 _{Cs} CLE	LABEL= 100-200 1: SYMBOL= FILLED S	37 _{Cs} QUARE
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
1 4 7 9 10 12 15 16 21 24 26 29 34 41 48 58 75 100	3.9E-04 7.0E-05 9.8E-06 6.1E-06 4.4E-06 3.9E-06 2.5E-06 2.6E-06 2.6E-06 1.4E-06 1.0E-06 8.8E-07 8.3E-07 6.8E-07 3.6E-07	2 3 3 5 5 7 9 11 12 14 19 22 24 26 29 34 41 48	1.1E-05 7.2E-06 4.5E-06 2.4E-06 2.1E-06 1.7E-06 1.3E-06 1.0E-06 8.2E-07 7.0E-07 5.9E-07 6.5E-07 6.8E-07 6.8E-07 4.7E-07 3.0E-07 2.5E-07 2.5E-07
163 211 278	3.8E-07 3.7E-07 2.6E-07 1.3E-07	58 75 100 162 211 278	2.4E-07 2.1E-07 1.9E-07 1.3E-07 1.1E-07 9.9E-08

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 18 FIGURE TITLE: 'LEACH RATES FOR ¹²⁹I'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'LEACH RATE'

CURVE 1 LABEL= 85-18 ¹²⁹ I SYMBOL= OPEN SQUARE		CURVE 2 LABEL= 85-200 ¹²⁹ I SYMBOL = FILLED CIRCLE		
COORDINATE	COORDINATE	COORDINATE	COORDINATE	
0	3.8E-04	0	1.4E-04	
9	3.5E-04	9	4.1E-05	
12	5.2E-04	12	6.1E-05	
100	3.2E-06	100	4.1E-07	

CU	IR VE	E 3
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1.0E-06

1.1E-06

1.1E-06

LABEL= 100-18 129_I SYMBOL= OPEN CIRCLE

131

163

280

Х	Y
COORDINATE	COORDINATE
0	5.5E-04
10	3.7E-04
12	5.3E-04
100	3.4E-06
280	1.0E-06

CU	R	VE	- 4
			-

1.6E-07

4.0E-07

2.6E-07

LABEL= 100-200 ¹²⁹I SYMBOL= FILLED SQUARE

131

163

280

X	Y
COORDINATE	COORDINATE
0	1.8E-04
1	4.7E-05
10	3.9E-05
12	5.7E-05
100	4.7E-07
280	1.4E-07

DRAWING NO. = ORNL-DWG-80-1450

REFERENCE: MITCHELL, GOODE AND VAUGHEN, p. 21 FIGURE TITLE: 'LEACH RATES FOR 125Sb'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'LEACH RATE'

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CUR	VE 1	CURVE 2	
LABEL= 85-18 125 SYMBOL= OPEN SQU	Sb ARE	LABEL= 85-200 125 SYMBOL= FILLED C	⁵ Sb IRCLE
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
0 2 4 6 7 9 11 13 15 17 20 22 24 28 33	1.5E-04 7.1E-05 3.4E-05 2.4E-05 2.2E-05 2.0E-05 1.7E-05 1.4E-05 1.1E-05 9.2E-06 8.4E-06 7.8E-06 6.8E-06 5.5E-06 4.9E-06	1 4 6 7 9 11 13 15 18 20 22 25 27 33 40	1.7E-05 1.6E-05 1.2E-05 8.6E-06 6.2E-06 4.4E-06 3.0E-06 1.8E-06 1.5E-06 1.3E-06 1.3E-06 1.3E-06 1.3E-06 1.3E-06 1.3E-06 1.3E-06 1.3E-06 1.3E-06
39 47 56 73 98 127 161 210 278	4.8E-06 4.8E-06 4.8E-06 4.1E-06 3.0E-06 2.4E-06 1.7E-06 1.0E-06 6.4E-07	47 57 73 98 129 162 210 277	1.1E-06 7.7E-07 6.3E-07 6.3E-07 6.5E-07 6.9E-07 7.0E-07 7.0E-07

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 21 (contd)

CURV	CURVE 3 CURVE 4	
LABEL= 100-18 12 SYMBOL= OPEN CIR	⁵ Sb CLE	LABEL= 100-200 ¹²⁵ Sb SYMBOL= FILLED SQUARE
X COORDINATE	Y COORDINATE	X Y COORDINATE COORDINATE
0 2 3 4 5 7 9 11 14 16 18 20 22 25 29 33 40 47	1.9E-04 5.4E-05 3.2E-05 2.3E-05 1.7E-05 1.8E-05 1.8E-05 1.6E-05 1.1E-05 8.7E-06 6.6E-06 5.9E-06 5.4E-06 3.4E-06 2.1E-06 1.4E-06	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
47 57 74	6.3E-07 5.1E-07	128 2.0E-07 162 2.0E-07
99 128 162	5.0E-07 5.5E-07 6.2E-07	210 2.0E-07 278 2.0E-07
210 277	6.6E-07 6.7E-07	DRAWING NO. = ORNL-DWG-80-1452

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 22 FIGURE TITLE: 'LEACH RATES FOR 106Ru'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'LEACH RATE'

CURVE 1 LABEL= 85-18 ¹⁰⁶ Ru SYMBOL= OPEN SQUARE		CURVE	CURVE 2	
		LABEL= 85-200 100 SYMBOL= FILLED C	LABEL= 85-200 ¹⁰⁶ Ru SYMBOL= FILLED CIRCLE	
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	
4 6 8 11 12 15 18 22 25 26 30 35	4.0E-06 3.1E-06 1.9E-06 7.7E-07 5.3E-07 3.7E-07 1.4E-07 1.0E-07 9.3E-08 9.0E-08 8.3E-08 6.8E-08	4 6 8 10 11 12 15 18 20 21 25 26 30 30 35 42 49 58 75 100 129 162 210 277	6.7E-07 6.3E-07 5.0E-07 3.3E-07 1.9E-07 1.0E-07 5.2E-08 2.7E-08 2.0E-08 2.0E-08 2.4E-08 3.1E-08 3.5E-08 3.5E-08 3.3E-08 2.7E-08 2.7E-08 2.0E-08 1.8E-08 1.8E-08 1.8E-08 1.9E-08 2.2E-08 2.3	

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 22 (contd)

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CUR	VE 3	CURVE 4
LABEL= 100-18 ¹⁰⁰ SYMBOL= OPEN CIRC	⁶ Ru CLE	LABEL= 100-200 ¹⁰⁶ Ru SYMBOL= FILLED SQUARE
X COORDINATE	Y COORDINATE	X Y COORDINATE COORDINATE
2	7.3E-06	4 1.2E-06
6	3.4E-06	8 1.2E-06
8	1.54E-06	10 8.8E-07
10	1.0E-06	13 4.7E-07
11	7.5E-07	15 2.9E-07
12	5.1E-07	17 2,5E-07
15	3.0E-07	20 2.5E-07
17	2.4E-07	22 2,9E-07
20	1.6E-07	27 2,5E-07
22	1.6E-07	29 1.4E-07
25	1.5E-07	34 7.8E-08
26	1.2E-07	40 6.1E-08
29	6.0E-08	52 6,6E-08
34	3.4E-08	72 3.7E-08
40	3.1E-08	100 1.1E-08
52	2.7E-08	137 6.0E-09
72	2.0E-08	189 6.1F-09
100	1.7E-08	
. 130	1.7E-08	

DRAWING NO. = ORNL-DWG-80-1469

REFERENCE: MITCHELL, GOODE AND VAUGHEN 1981, p. 23

FIGURE TITLE: 'LEACH RATES FOR 90Sr'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'LEACH RATE'

CUF	RVE 1	CURV	/E 2	CUR	VE 3
LABEL = 85- SYMBOL = OF	-18 ⁹⁰ Sr PEN SQUARE	LABEL= 85- SYMBOL= FI	-200 ⁹⁰ Sr LLED CIRCLE	LABEL= 10 SYMBOL= 0	0-18 ⁹⁰ Sr PEN CIRCLE
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	X COORDINAT	Y E COORDINATE
1 3 5 9 12 17 25 38 54 73 100 129 169 226	6.3E-06 4.1E-06 2.6E-06 1.7E-06 1.3E-06 1.0E-06 4.8E-07 1.9E-07 9.7E-08 5.1E-08 2.7E-08 2.0E-08 1.7E-08 1.5E-08	1 3 7 12 17 25 38 54 73 100 129 169 226	2.1E-07 2.0E-07 1.6E-07 1.4E-07 9.4E-08 4.2E-08 2.4E-08 1.8E-08 1.1E-08 5.7E-09 4.2E-09 4.2E-09 4.3E-09	1 3 6 7 10 12 17 25 38 54 73 100 129 169 226	4.1E-06 3.5E-06 2.3E-06 1.3E-06 9.5E-07 7.6E-07 4.6E-07 3.2E-07 2.9E-07 1.6E-07 5.4E-08 2.8E-08 2.7E-08 2.7E-08
		CUR	/E 4		
	LABEL= 100 SYMBOL= FI	0-200 ⁹⁰ Sr LLED SQUARE			
	X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	
	0 3 6 7 10 13 17 25	4.2E-07 4.2E-07 4.2E-07 3.3E-07 1.8E-07 1.1E-07 8.4E-08 5.4E-08	38 54 73 100 129 169 226	2.5E-08 1.2E-08 8.3E-09 8.1E-09 8.1E-09 7.2E-09 5.9E-09	

DRAWING NO. = ORNL-DWG-80-1470

REFERENCE: JOHNSON et al. 1981, p. 23

'FRACTIONAL RELEASE OF ¹³⁷Cs FROM FUEL DURING 25C LEACHING FIGURE TITLE: PERIOD'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

> 1.5E-06 1.1E-06

CURVE 1 LABEL= RUN #1 SYMBOL= OPEN SQUARE		CURVE 2 LABEL= RUN #3 SYMBOL= INVERTED TRIANGLE		
. 9	1.3E-04	2	9.8E-04	
22	8.2E-06	9	2.0E-04	
39	6.2E-06	15	5.3E-05	
59	3.4E-06	29	1.2E-05	
72	2.1E-06	59	4.1E-06	

CURVE 3

1.6E-06

LABEL = RUN #4 SYMBOL= OPEN TRIANGLE

92

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X COORDINATE	Y COORDINATE
2	2.5E-04
8	7.3E-05
15	1.7E-05
29	5.4E-06
59	3.0E-06
88	1.7E-06
110	6.1E-07

CURVE 4

LABEL= RUN #5 SYMBOL= OPEN CIRCLE

89

110

X	Y
COORDINATE	COORDINATE
2	2.7E-03
8	3.6E-04
16	8.6E-05
29	1.8E-05
59	5.4E-06
89	2.2E-06
120	1.5E-06

'FRACTIONAL RELEASE RATES OF RADIONUCLIDES IN DEIONIZED DISTILLED WATER (LEACHANT A)' FIGURE TITLE:

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X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1		CURVE 2		CURVE 3		
LABEL = 137_{CS} SYMBOL = OPEN		LABEL= ⁹⁰ SYMBOL= 0	LABEL= ⁹⁰ Sr SYMBOL= OPEN SQUARE		LABEL= U SYMBOL= OPEN CIRCLE	
		Х	Y	Х	Y	
Х	Y	COORDINAT	E COORDINATE	COORDINATE	COORDINATE	
COORDIN	ATE COORDINATE		···· <u>-···············</u> ·			
		0	8.0E-06	181	1.7E-09	
3	9.9E-04	7	9.2E-06	214	1.3E-09	
6	3.0E-04	5	7.6E-06	275	9.0E-10	
9	1.1E-04	11	6.5E-06	337	2.9E-09	
15	1.5E-05	11	5.5E-06	395	1.0E-09	
17	6.4E-06	21	4.9E-06	458	2.0E-09	
29	5.7E-06	27	4.0E-06	517	1.8E-09	
35	4.6E-06	33	3.4E-06	579	1.1E-09	
43	4.0E-06	41	3.3E-06	673	7.7E-10	
54	2.2E-06	56	2.8E-06	783	1.0E-09	
84	1.2E-06	82	1.6E-06	812	4.2E-10	
114	1.1E-06	115	1.6E-06	823	1.0E-08	
148	9.5E-07	149	1.5E-06			
181	8.2E-07	181	1.1E-06			
207	9.0E-07	212	6.5E-07			
274	3.2E-07	272	3.9E-07			
335	4.0E-07	336	2.6E-07			
398	4.4E-07	397	1.6E-07			
459	4.8E-07	459	9.3E-08			
516	4.7E-07	518	8.5E-08			
579	3.6E-07	580	8.5E-08			
666	2.1E-07	668	2.9E-08			
779	3.1E-07	779	2.0E-08			
809	5.5E-07	817	1.7E-08			
820	1.2E-06	820	4.5E-08			
911	5.0E-07					

REFERENCE: JOHNSON 1982, p. 18 (contd)

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CURVE 4		CURVE 5		
LABEL= ⁹⁹ Tc		LABEL= 239+240 _{Pu}		
SYMBOL= X		SYMBOL= FILLED CIRCLE		
X	Y	X	Y	
COORDINATE	COORDINATE	COORDINATE	COORDINATE	
578 669 815 821	1.0E-07 4.7E-08 8.8E-08 2.1E-07	0 4 7 13 22 28 35 58 84 113 145 183 213 273 335 398 456 517 575 670 780 819 820	1.5E-06 5.1E-07 5.8E-07 5.1E-07 4.8E-07 5.7E-07 3.4E-07 3.5E-07 3.5E-08 3.5E-08 4.3E-08 6.5E-08 4.3E-08 9.5E-09 1.7E-08 2.1E-08 3.2E-08 3.2E-08 1.4E-08 1.6E-08 1.6E-08 1.3E-08 3.8E-08 3.8E-08	

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A.29

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'FRACTIONAL RELEASE RATES OF RADIONUCLIDES IN KBS GROUNDWATER (LEACHANT B)' FIGURE TITLE:

•

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1		CURVE 2		CURVE 3	
LABEL= ¹³⁷ Cs SYMBOL= OPEN TRIANGLE		LABEL= ⁹⁰ Sr SYMBOL= OPEN SQUARE		LABEL= URANIUM SYMBOL= OPEN CIRCLE	
X COORDINA	Y TE COORDINATE	X COORDINAT	Y E COORDINATE	X COORDINATE	Y COORDINATE
0 0 4 13 22 25 33 83 114 181 212 277 332 404 461 524 584 676 787 818 824	9.6E-04 2.3E-04 4.3E-05 1.5E-05 7.6E-06 6.0E-06 3.7E-06 3.7E-06 3.2E-06 1.6E-06 8.9E-07 4.6E-07 6.7E-07 3.2E-07 3.2E-07 3.6E-07 8.3E-07 5.4E-07 1.9E-07 2.8	0 5 7 13 20 30 44 58 89 118 148 148 183 216 277 333 401 463 523 584 677 787	7.1E-06 3.3E-06 2.3E-06 1.2E-06 9.7E-07 8.3E-07 5.5E-07 3.8E-07 3.8E-07 3.6E-07 2.6E-07 2.7E-07 1.2E-07 8.3E-08 9.5E-08 7.8E-08 9.1E-08 3.6E-07 2.4E-07 6.4E-08 9.2E-08	184 218 277 335 402 464 522 588 676 789 818 826	9.2E-08 6.8E-08 7.0E-08 7.7E-08 6.3E-08 7.6E-08 4.1E-07 6.9E-08 5.4E-08 1.7E-08 3.9E-08 4.4E-08
919	2.5E-07	821 827	9.6E-08 1.2E-07		

REFERENCE: JOHNSON 1982, p. 19 (contd)

CURVE 4		CUR	CURVE 5		
LABEL = ⁹⁹ Tc		LABEL= ²³⁹⁺²⁴⁰ Pu			
SYMBOL = X		SYMBOL= .			
X	Y	X	Y		
COORDINATE	COORDINATE	COORDINATE	COORDINATE		
585 676 821 826	9.9E-08 8.8E-08 6.5E-08 5.2E-08	2 5 7 17 22 30 34 45 57 89 117 154 186 217 335 402 464 524 586 678 789	1.1E-06 1.2E-07 3.1E-07 6.7E-08 7.4E-08 4.8E-08 3.5E-07 2.7E-08 1.1E-07 5.9E-08 3.5E-08 3.5E-08 3.7E-08 3.7E-08 5.5E-08 2.0E-08 2.8E-08 2.8E-08 7.5E-09 5.2E-08 6.1E-09 3.2E-09 3.1E-09 3.1E-09		

FIGURE TITLE: 'FRACTIONAL RELEASE RATES OF RADIONUCLIDES IN GRANITE GROUNDWATER (LEACHANT C)'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

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CURVE	1

LABEL=	137 _{Cs}	
SYMBOL=	OPEN	TRIANGLE

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Х	Y
COORDINATE	COORDINATE
0	4.5E-04
2	1.6E-04
5	6.9E-05
13	1.5E-05
21	3.7E-06
26	2.9E-06
35	2.5E-06
43	1.8E-06
55	1.7E-06
83	1.3E-06
116	9.5E-07
181	8.6E-07
217	3.6E-07
276	5.3E-07
338	6.6E-07
401	5.1E-07
461	4.7E-07
523	5.3E-07
583	3.8E-07
672	2.8E-07
784	3.2E-07
816	2.7E-07
821	2.8E-07
914	3.3E-07

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LABEL= ⁹⁰Sr SYMBOL= OPEN SQUARE

X	Y
COORDINATE	COORDINATE
0	9.1E-06
2	5.7F-06
5	4 8F_06
12	3 7E-06
20	2 4F_06
29	2.5E-06
20	2.JE-00
20	1 65 06
13	1 25 06
43	1. JE -00
50	1.25-00
8/	9.3E-07
110	6.5E-0/
151	4.8E-07
186	3.8E-07
217	2.0E-07
278	1.5E-07
339	2.3E-07
402	1.7E-07
462	1.5E-07
525	1.8E-07
583	1.1E-07
675	9.3E-08
784	1.2E-07
815	1.3E-07
825	1.6E-07

REFERENCE: JOHNSON 1982, p. 20 (contd)

CURVE 3	CURVE 4	CURVE 5
LABEL= URANIUM	LABEL= ⁹⁹ Tc	LABEL= ²³⁹⁺²⁴⁰ Pu
SYMBOL= OPEN CIRCLE	SYMBOL= X	SYMBOL= FILLED CIRCLE
X Y	X Y	X Y
COORDINATE COORDINATE	COORDINATE COORDINATE	COORDINATE COORDINATE
182 9.5E-08 214 4.7E-08 278 3.6E-08 334 4.0E-08 397 2.9E-08 461 3.1E-08 524 5.5E-08 583 2.2E-08 671 3.9E-08 786 5.7E-09 820 1.4E-08 826 4.4E-08	581 2.5E-07 674 1.2E-07 814 1.6E-07 824 3.3E-08	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

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'FRACTIONAL RELEASE RATES OF RADIONUCLIDES IN STANDARD CANADIAN SHIELD SALINE SOLUTION (LEACHANT D)' FIGURE TITLE:

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

5.9E-08

CURVE 1 LABEL = ¹³⁷ Cs SYMBOL = OPEN TRIANGLE		CURVE 2		
		LABEL= ⁹⁰ Sr SYMBOL= OPEN SQUARE		
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	
0 4 13 19 28 32 41 117 182 213 272 338 400 461	2.5E-04 8.0E-05 4.3E-05 1.1E-05 3.1E-06 1.7E-06 1.3E-06 1.2E-06 7.3E-07 8.7E-07 8.7E-07 8.7E-07 1.3E-07 2.3E-07 1.6E-07	3 3 10 13 22 28 35 44 55 84 117 149 183 216 338	7.4E-06 3.4E-06 2.7E-06 2.3E-06 2.5E-06 2.3E-06 1.9E-06 1.4E-06 1.1E-06 5.4E-07 4.6E-07 6.1E-07 2.7E-07 3.7E-07	
461 518 580 672 781 817 823 914	1.6E-07 2.2E-07 1.9E-07 5.4E-07 5.3E-07 1.3E-07 1.0E-07 1.2E-07	338 399 459 521 583 673 782 816 822	3.5E-08 2.6E-08 5.0E-08 8.9E-08 1.9E-08 1.1E-07 7.6E-08 7.4E-08 5.9E-08	

CURVE 3

LABEL=	⁹⁹ Tc
SYMBOL=	X

X COORDINATE	Y COORDINATE
580 672	3.6E-07
785	1.8E-07
816 825	1.5E-07
672 785 816 825	2.7E-07 1.8E-07 1.5E-07 9.0E-08

A.34

FIGURE TITLE: 'FRACTIONAL RELEASE RATES OF ¹³⁷Cs UNDER CONTROLLED OXYGEN. 'PRESSURE CONDITIONS'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

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CURVE 1 CUF		VE 2	CUR	CURVE 3	
LABEL= 2 SYMBOL=	.5E-04 OPEN CIRCLE	E-04 EN CIRCLE EN CIRCLE EN CIRCLE EN CIRCLE EN CIRCLE		LABEL= 1.0E-06 SYMBOL= OPEN TRIANGLE	
X COORDINA	Y TE COORDINATE	X COORDINAT	Y <u>E COORDINATE</u>	X COORDINATE	Y COORDINATE
0	4.4E-04	0	2.3E-04	0	6.4E-04
2	1.6E-05	3	1.8E-05	3	3.5E-05
4	5.5E-06	5	6.2E-06	5	1.1E-05
11	4./E-06	11	1.8E-06	10	3.6E-06
26	1.9E-06	26	8.0E-07	27	1.6E-06
47	2.1E-06	46	9.7E-07	47	4.1E-07
75	1.7E-06	76	6.3E-07	77	5.0E-07
105	1.0E-06	107	3.7E-07	107	3.1E-07
136	5.9E-07	135	2.5E-07	135	5.0E-07
167	4.2E-07	166	3.5E-07	166	3.1E-07
194	2.4E-07	195	3.7E-07	194	2.8E-07
224	3.7E-07	224	3.0E-07	225	2.6E-07
253	2.5E-07	253	2.2E-07	255	3.3E-07
287	2.1E-07	290	1.2E-07	287	3.9E-07
313	2.4E-07	314	1.2E-07	313	3.6E-07
346	2.6E-07	346	1.0E-07	346	2.1E-07
374	2.3E-07	375	7.5E-08	376	1.8E-07
	-	404	1.5E-07	403	1.8F-07
		435	1.0E-07	434	1.9E-07

[02]mol*kg e-01

'FRACTIONAL RELEASE RATES OF ⁹⁰Sr UNDER CONTROLLED OXYGEN PRESSURE CONDITIONS' FIGURE TITLE:

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1		CURVE 2		CUR	CURVE 3	
LABEL= 2 SYMBOL=	-5E-04 OPEN CIRCLE	LABEL= 6. SYMBOL= (.1E-06 DPEN SQUARE	LABEL= 1. SYMBOL= 0	0E-06 PEN TRIANGLE	
X COORDINA	Y TE COORDINATE	X COORDINAT	Y TE COORDINATE	X COORDINAT	Y E COORDINATE	
5	3.5E-05	5	2.7E-05	5	3.1E-05	
7	3.5E-05	7	5.8E-06	7	9.0E-06	
11	6.6E-06	11	3.4E-06	11	4.5E-06	
17	8.3E-06	16	1.4E-06	15	1.2E-06	
32	4.0E-06	31	3.1E-07	29	2.1E-07	
52	3.2E-06	52	3.6E-07	52	1.2E-07	
78	2.5E-07	75	5.0E-07	75	5.5E-08	
110	2.2E-07	110	3.6E-07	110	3.2E-08	
140	1.3E-07	140	2.4E-07	140	7.1E-08	
170	1.6E-07	170	2.2E-06	170	2.3E-07	
200	1.4E-07	200	2.4E-07	200	2.5E-08	
230	1.0E-07	230	1.2E-07	230	2.8E-08	
250	8.7E-08	250	1.0E-07	250	3.1E-08	
290	1.2E-07	290	4.1E-08	290	2.2E-08	
320	1.1E-07	320	3.6E-08	320	2.0E-08	
340	1.1E-07	340	3.7E-08	340	1.5E-08	
370	1.1E-07	370	3.9E-08	370	1.2E-08	
		400	5.0E-08	400	2.1E-08	
		430	2.7E-08	430	1.4E-08	

FIGURE TITLE: 'FRACTIONAL RELEASE RATES FOR ⁹⁹Tc UNDER CONTROLLED OXYGEN PRESSURE CONDITIONS'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 2

CURVE 1

LABEL= 2.5E-04

SYMBOL= OPEN CIRCLE

LABEL= 6.1E-06 SYMBOL= OPEN SQUARE

X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
82	5.9E-07	194	1.2E-07
107	7.8E-07	227	1.0E-07
137	2.1E-07	257	8.3E-08
167	1.6E-07	289	4.3E-08
228	1.2E-07	317	4.2E-08
255	1.1E-07	349	6.4E-08
290	1.1E-07	379	4.2E-08
318	1.1E-07	410	4.1E-08
349	1.2E-07	441	8.0E-08
378	1.1E-07		

CURVE 3

LABEL= 1.0E-06 SYMBOL= OPEN TRIANGLE

X	Y
COORDINATE	COORDINATE
198	5.0E-08
229	3.2E-08
256	4.2E-08
290	5.9E-08
319	3.1E-08
349	3.0E-08
378	4.2E-08
410	4.0F-08
439	4.0F-08

[02]mol*kg e-01

FIGURE TITLE: 'FRACTIONAL RELEASE RATES OF URANIUM UNDER CONTROLLED OXYGEN PRESSURE CONDITIONS'

X-AXES LABEL IS 'TIME (DAYS)'

437

8.4E-09

CURVE 1		CURVE 2		CURVE 3	
LABĖL= 2 Symbol=	•5E-04 OPEN CIRCLE	LABEL= 6 SYMBOL= (.1E-06 DPEN SQUARE	LABEL= 1. SYMBOL= (.0E-06 DPEN TRIANGLE
X COORDINA	Y TE COORDINATE	X COORDINA	Y TE COORDINATE	X COORDINAT	Y TE COORDINATE
2	2.5E-06	0	3.9E-06	4	2.0E-06
3	8.0E-07	2	6.9E-07	2	7.0E-07
11	8.1E-07	11	5.7E-07	3	5.7E-07
15	2.4E-07	15	1.0E-07	11	6.8E-07
29	8.3E-08	31	4.4E-08	15	8.2E-08
51	8.3E-08	51	3.4E-08	30	3.6E-08
75	1.9E-08	105	7.5E-08	51	2.6E-08
109	1.6E-08	133	5.7E-08	75	2.4E-08
132	3.4E-08	167	9.4E-08	105	2.3E-08
166	2.7E-08	194	3.7E-08	133	1.6E-08
193	2.8E-08	228	2.5E-08	167	2.0E-08
228	2.9E-08	255	-3.7E-08	194	6.3E-09
255	1.2E-08	286	3.5E-08	228	9.4E-09
284	1.1E-08	315	2.6E-08	255	1.6E-08
314	2.2E-08	348	2.1E-08	288	1.4E-08
348	2.5E-08	375	1.8E-08	315	1.3E-08
375	2.1E-08	405	9.6E-09	351	1.0E-08
		438	1.6E-08	380	1.6E-09
				408	2.4E-09

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

REFERENCE: JOHNSON et al. 1982, p. 246

FIGURE TITLE: 'FRACTIONAL RELEASE RATES OF ²³⁹⁺²⁴⁰Pu FROM IRRADIATED FUEL AT VARIOUS OXYGEN CONCENTRATIONS AT 25°C'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1		CURV	CURVE 2		
LABEL= 2.5E-04		LABEL= 6.1E-04			
SYMBOL= X		SYMBOL= OPEN CIRCLE			
X	Y	X	Y		
COORDINATE	COORDINATE	COORDINATE	COORDINATE		
3	1.0E-06	3	5.0E-07		
4	1.3E-07	4	8.0E-08		
7	5.6E-07	7	6.3E-08		
14	2.7E-07	14	3.1E-08		
27	6.0E-08	27	1.9E-08		
49	4.1E-08	49	3.3E-08		
75	5.7E-08	75	1.5E-08		
100	4.4E-08	100	3.3E-08		
130	3.0E-08	130	2.1E-09		

CURVE 3

LABEL=	1.0E-06	
SYMBOL=	FILLED	TRIANGLE

X COORDINATE	Y COORDINATE
3	3.2E-07
4	5.3E-08
7	3.1E-08
14	2.4E-08
27	1.5E-08
49	7.5E-09
75	1.1E-08
100	5.6E-09
130	2.6E-09

[02]mol*kg e-01

A.39

REFERENCE: JOHNSON et al. 1982, p. 247

'FRACTIONAL RELEASE RATES OF $^{137}\mbox{Cs}$ For fuels of moderate and high power ratings' FIGURE TITLE:

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 2

LABEL= 43 kW/m

SYMBOL= OPEN CIRCLE

c_{1}	D.	VE.	1
60	714	¥C.	1

LABEL= 53 kW/m SYMBOL= FILLED CIRCLE

X	Y	X	Ŷ
CUURDINATE	COURDINATE	COORDINATE	COORDINATE
6	1.5E-02	6	9.2E-04
7	5.5E-03	7	3.1E-04
9	1.1E-03	9	9.6E-05
13	5.2E-04	14	1.5E-05
18	2.3E-04	22	6.0E-06
25	2.0E-04	29	5.2E-06
36	1.4E-04	35	4.5E-06
51	8.1E-05	42	3.8E-06
74	3.0E-05	54	2.2E-06
96	2.1E-05	83	1.1E-06
138	1.4E-05	112	1.0E-06
189	1.0E-05	176	8.2E-07
243	4.7E-06	203	8.5E-07
285	7.1E-06	264	3.3E-07
		907	3.3E-07

REFERENCE: JOHNSON et al. 1983, p. 473

FIGURE TITLE: 'GAS RELEASE AND CUMULATIVE ¹³⁷Cs LEACHING DATA FOR PICKERING BUNDLE 15527C'

X-AXES LABEL IS 'TIME (DAYS)'

Y-AXES LABEL IS 'PERCENT RELEASED'

CURVE 1		CURVE 2		
LABEL = CURVE 1 SYMBOL = X		LABEL= CURVE 2 SYMBOL= X		
X	Y	X	Y	
COORDINATE	COORDINATE	COORDINATE	COORDINATE	
1 3 11 20 35 60 97 274	0.78 1.2 1.9 2.2 2.5 2.8 2.9 3.1	1 3 4 11 21 36 60 97 277	0.63 1.0 1.2 1.8 2.1 2.3 2.6 2.8 3.0	
CURVE 3		CURVE 4		
LABEL= CURVE 3		LABEL= CURVE 4		
SYMBOL= OPEN CIRCLE		SYMBOL= OPEN CIRCLE		
X	Y	X	Y	
COORDINATE	COORDINATE	COORDINATE	COORDINATE	
1	0.80	1	1.5	
3	1.2	3	2.1	
4	1.5	4	2.3	
11	2	10	2.7	
20	2.4	20	3.0	
35	2.7	34	3.3	
60	2.9	58	3.5	
96	3.1	93	3.7	
274	3.4	280	3.9	

REFERENCE: JOHNSON et al. 1984, p. 5

'PERCENTAGE ^{137}Cs AND ^{129}I RELEASED FROM BRUCE F21271C OUTER FUEL-ELEMENT SECTIONS LEACHED IN DEIONIZED DISTILLED WATER AT 25°C WITH NO CARRIER PRESENT' FIGURE TITLE:

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'INVENTORY RELEASED'

CURVE 1 LABEL= I WITH ATTACHED SHEATH SYMBOL= FILLED CIRCLE		CURVE 2 LABEL= I WITH CUT OPEN SHEATH SYMBOL= OPEN CIRCLE	
COORDINATE	COORDINATE	COORDINATE	COORDINATE
0	1.2E-02	0	3.1E-02
1	2.8E-02	1	5.3E-02
2	3.3E-02	2	4.7E-02

CURVE 3

LABEL= ¹³⁷Cs with attached sheath SYMBOL= FILLED CIRCLE

CURVE 4

LABEL= ¹³⁷Cs WITH SHEATH CUT OPEN SYMBOL= OPEN CIRCLE

Х	Y	X	Y
COORDINATE	COORDINATE	COORDINATE	COORDINATE
0	0.12	1	1.4
1	0.52	1	1.8
1	1.7	2	2.1
2	1.8		

REFERENCE: JOHNSON et al. 1984, p.6

'PERCENTAGE ¹³⁷Cs AND ¹²⁹I RELEASED FROM PICKERING A13894W OUTER FUEL ELEMENT SECTIONS (IN DEIONIZED DISTILLED WATER AT 25°C WITH 0.2 g/L KI CARRIER)' FIGURE TITLE:

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'INVENTORY RELEASED (%)'

CURVE 1 LABEL = ¹²⁹ I SYMBOL = OPEN CIRCLE		CURVE	CURVE 2	
		LABEL= ¹³⁷ Cs SYMBOL= OPEN DIAMOND		
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	
0	5.4E-02	0	0.58	
0	6.4	• 0	2.1	
1	9.4	1	3.6	
3	10	2	3.9	
4	11	3	3.9	

REFERENCE: WILSON AND OVERSBY 1984, p. 8

FIGURE TITLE: 'URANIUM IN SOLUTION'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1 LABEL= BARE HBR FUEL SYMBOL= OPEN CIRCLE		CURV	CURVE 2	
		LABEL= SPLIT DEFECTS SYMBOL= OPEN SQUARE		
X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE	
1	4.5E+01	2	8.2E-03	
5 16	0.6E-01 0.1E-01	6 14	3.9E-02 2.7E-02	
30 59	0.3E-01 0.2E-01	29 59	2.7E-02 2.3E-02	
09 119 150	0.4E-01 0.5E-01	90 120 149	2.9E-02 3.9E-02 2.2E-02	
179	0.5E-01	179	3.3E-02	

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LABEL= HOLES DEFECTS SYMBOL= OPEN TRIANGLE

~ X	Ŷ
COORDINATE	COORDINATE
2	4.0E-03
6	2.0E-03
15	6.0E-03
30	6.0E-03
59	4.1E-03
90	3.7E-03
119	5.8E-03
148	1.7E-02
178	5.4E-03

CURVE 4

LABEL = UNDEFECTED SYMBOL= FILLED CIRCLE

Х	Y
COORDINATE	COORDINATE
2	2.8E-03
6	2.0E-03
14	1.9E-03
30	1.8E-03
61	3.0E-03
89	1.7E-03
18	1.9E-03
49	5.7E-03
78	2.7E-03
REFERENCE: WILSON AND OVERSBY 1984, p. 8 FIGURE TITLE: '239+240Pu IN SOLUTION'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1 LABEL= BARE FUEL SYMBOL= OPEN CIRCLE		CURVE 2 LABEL= SPLIT DEFECTS SYMBOL= OPEN SQUARE		
3 6 16 30 61 90 119 150 179	1.4E+03 8.0E+02 1.2E+03 1.4E+03 1.0E+03 9.7E+02 9.4E+02 7.4E+02 9.0E+02	2 7 15 29 60 89 118 149 178	2.8E+00 5.5E+00 7.5E+00 2.6E+01 3.8E+01 5.9E+01 1.0E+02 5.9E+01 7.7E+01 7.7E+01	

CURVE 3

LABEL = HOLE DEFECTS SYMBOL= OPEN TRIANGLE

LABEL=	UNDEFEC	TED
SYMBOL=	FILLED	CIRCLE

CURVE 4

X COORDINATE	Y COORDINATE	X COORDINATE	Y COORDINATE
1	1.4E+00	2	0.8E+00
6	1.4E+00	5	0.8E+00
15	6.2E+00	15	1.1E+00
31	1.3E+01	29	4.2E+00
59	1.5E+01	61	6.5E+00
90	1.8E+01	90	8.6E+00
120	1.9E+01	120	1.0E+01
150	1.9E+01	148	1.0E+01
177	1.9E+01	178	1.1E+01

REFERENCE: WILSON AND OVERSBY 1984, p. 8 FIGURE TITLE: '137Cs IN SOLUTION'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1		CURVE	CURVE 2		
LABEL= BARE FUEL		LABEL = SLIT DEFECTS			
SYMBOL = OPEN CIRCLE		SYMBOL = OPEN SQUARE			
X	Y	X	Y		
COORDINATE	COORDINATE	<u>COORDINATE</u>	COORDINATE		
1	9.9E+06	1	2.4E+04		
4	1.6E+07	5	4.9E+05		
13	2.0E+07	13	1.5E+06		
29	2.1E+07	29	3.4E+06		
60	1.9E+07	59	5.9E+06		
89	1.2E+07	90	8.7E+06		
119	5.0E+06	119	9.4E+06		
149	2.5E+06	149	1.0E+07		
178	1.4E+06	179	1.0E+07		
CURVE	3	CURVE	4		
LABEL= HOLE DEFECTS		LABEL= UNDEFECTED			
SYMBOL= OPEN TRIANGLE		SYMBOL= FILLED CIRCLE			
X COORDINATE 1 5 14 29 59 89 120 149 178	Y COORDINATE 1.3E+04 2.4E+04 1.1E+06 2.7E+06 4.2E+06 5.0E+06 4.2E+06 4.9E+06 4.9E+06	X COORDINATE 1 5 14 30 59 90 119 149 149 180	Y <u>COORDINATE</u> 2.9E+03 2.9E+03 3.5E+03 3.2E+03 3.5E+03 3.5E+03 3.5E+03 3.5E+03 3.8E+03		

REFERENCE: WILSON AND OVERSBY 1985, p. 3

FIGURE TITLE: 'URANIUM IN UNFILTERED SOLUTION'

X-AXES LABEL IS 'TIME (DAYS)' Y-AXES LABEL IS 'FRACTIONAL RELEASE RATE (1/DAYS)'

CURVE 1		CURVE	CURVE 2		
LABEL= BARE HBR I SYMBOL= FILLED C	FUEL IRCLE	LABEL= SLIT DEFEC SYMBOL= OPEN SQUA	CT ARE		
Х	Y	Х	Y		
COORDINATE	COORDINATE	COORDINATE	COORDINATE		
3	4.1	3	2.8E-02		
7	4.4	· 7	5.0E-02		
10	4.6	21	5.3E-02		
21	3.8	30	6.1E-02		
30	3.6	63	6.8E-02		
63	2.3	120	6.5E-02		
120	1.5	182	8.7E-02		
181	1.3	223	9.5E-02		
223.0742	1.144536				

CURVE 3	CURVE 4
LABEL= HOLE DEFECTS	LABEL= UNDEFECTED
SYMBOL= OPEN TRIANGLE	SYMBOL= OPEN CIRCLE
X Y	X Y
COORDINATE COORDINATE	COORDINATE COORDINATE
3 7 15-03	3 1 55-02

3	/.IE-03	3	1.5E-02
7	8.9E-03	7	2.0E-02
21	1.0E-02	21	2.2E-02
31	6.7E-03	30	1.9E-02
64	6.7E-03	64	1.6E-02
121	4.6E-03	121	1.4E-02
182	4.6E-03	182	1.4E-02
223	5.5E-03	223	1.1E-02

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APPENDIX B

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ANNOTATED BIBLIOGRAPHY OF LITERATURE ON SPENT FUEL/U02 CHARACTERIZATION

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APPENDIX B

ANNOTATED BIBLIOGRAPHY OF LITERATURE ON SPENT FUEL/UO2 CHARACTERIZATION

Due to the heterogeneous nature of spent nuclear fuel, it is critical to have accurate, detailed information on the characteristics of the materials being used to evaluate the performance of spent fuel as a potential waste form. This appendix presents the literature compiled for the informal Waste Package Program (WPP) library at PNL on characterization and the properties of spent fuel and UO_2 . The annotations are either the abstract or a synopsis of the publication. Exhaustive reviews of the literature in this area were not conducted; the intent of this appendix is to present the available literature.

This appendix is organized with groupings of the various publications into subject areas. The sections address subject areas of general information on spent fuel characterization (including general reference material), experimental methods of characterization, data on destructive and nondestructive testing of spent fuel, and modeling efforts related to spent fuel characterization. Literature references are also included for fuel restructuring, radionuclide redistribution within the spent fuel, and out-of-reactor fuel performance. These last three subject areas are given without individual summaries of the articles.

GENERAL INFORMATION ON SPENT FUEL CHARACTERIZATION

Almassy, M. Y. and R. E. Woodley, August 1972. <u>Possible Effects of UO₂ Oxida-</u> <u>tion on Light Water Reactor Spent Fuel Performance in Long-Term Geologic Dis-</u> <u>posal</u>. HEDL-TC-1502, Rev. 1. Hanford Engineering Development Laboratory, Richland, WA.

A technical assessment of published literature representing the current level (August 1982) of understanding of spent fuel characteristics and conditions that may degrade pellet integrity during a geologic disposal sequence. The main emphasis is the identification of the main modes of UO₂ degradation in a geologic setting (69 references).

Barner, J. O. 1984. <u>LWR Spent Fuel Approved Testing Materials for Radionu-</u> <u>clide Release Studies</u>. PNL-4686, UC-70, Pacific Northwest Laboratory, Richland, WA.

Criteria are defined for the selection of light water reactor (LWR) spent fuels for use as MCC-Approved Testing Materials (ATMs) in radionuclide dissolution and interaction studies. Fuel-related characteristics affecting the release of radionuclides from spent fuel are reviewed and their pertinence evaluated. ATM spent fuel criteria are defined and classes of ATM spent fuels are determined. The available inventory of governmentowned LWR spent fuel is identified and current plans for acquisition by the MCC are summarized. The characterization data to be supplied with the spent fuel ATMs are also described (11 references).

Belle, J., ed. 1961. <u>Uranium Dioxide: Properties and Nuclear Applica-</u> <u>tions</u>. Naval Reactors, Division of Reactor Development, Unites States Atomic Energy Commission, U.S. Government Printing Office, Washington D.C.

A textbook presentation of UO_2 with regard to the physical properties of UO_2 and the detailed phase relationships in the uranium-oxygen system, solid state reaction and the oxidation and corrosion behavior of uranium dioxide, and the irradiation behavior and performance as a fuel. Numerous references to the literature prior to 1961.

Funk, C. W. and L. D. Jacobson, May 1978. <u>Inventory and Characterization of</u> <u>Spent LWR Fuel</u>. HEDL-TME 77-82, Hanford Engineering Development Laboratory. Richland, WA.

An inventory of spent LWR fuel discharged from reactors and stored in water basins up to 1976. Defective fuel is identified and categorized, and the trends of failed fuel are evaluated for their impact on the operation of receiving facilities. Possible failure mechanisms that lead to the release of radioactivity have been identified. The sources and levels of activity are estimated for typical LWR spent fuel assemblies (38 references).

Greene, E. M. 1980. <u>Spent Fuel Data for Waste Storage Programs</u>. HEDL-TME 79-20, Hanford Engineering Development Laboratory, Richland, WA.

A compilation of data on LWR spent fuel for dissemination to participants in DOE-sponsored waste storage programs. Included are mechanical descriptions of the existing major types of LWR fuel assemblies, fission product inventories for spent LWR fuel calculated by the ORIGEN code, decay heat data, and inventories of LWR spent fuel currently in storage, with projections of future quantities (25 references).

Houston, M. D. 1978. "Properties of Spent Fuel." In <u>Proceedings of the Con-</u> <u>ference on High-Level Radioactive Solid Waste Forms</u>, NUREG/CP-0005, Denver, CO, Nuclear Regulatory Commission, Washington, DC., p. 561-589. The properties of spent fuel from current commercial nuclear power reactors are discussed in this review of the literature by the NRC. The major topics are as follows: fission product formation, fission product relocation and release, pellet cracking and relocation, fuel-to-cladding interaction, and Zircaloy growth and embrittlement. The chemical and physical properties of reactor fuel after achieving its designed irradiation exposure are described. The integrity of spent fuel rods is discussed as it relates to short- and long-term storage (39 references and panel discussion).

Johnson, L. H. and J. L. Crosthwaite, November 1984. <u>Fuel Characterization</u> <u>Research for the Canadian Nuclear Fuel Waste Management Program</u>. AECL-8375. Whiteshell Nuclear Research Establishment, Atomic Energy of Canada Limited. Pinawa, Manitoba, Canada.

An outline of the fuel characterization work being performed in support of the Canadian Nuclear Fuel Waste Management Program. A general evaluation of CANDU fuel characteristics is given with regard to its use as a waste form (29 references).

Olander, D. R., 1976. <u>Fundamental Aspects of Nuclear Reactor Fuel Elements</u>, TID-26711-P1, Technical Information Center, Energy Research and Development Administration.

Covers the general knowledge needed to understand the subsequent discussions of the properties and behaviors of oxide nuclear fuel. The book includes references to advanced fuels as well as a companion volume <u>Solution to Problems</u> which contains solutions to many of the problem sets contained in the text (numerous literature references).

Rothman, A. J. 1984. <u>Potential Corrosion and Degradation Mechanisms of</u> <u>Zircaloy Cladding on Spent Nuclear Fuel in a Tuff Repository</u>. UCID-20172, Lawrence Livermore National Laboratory, Livermore, CA.

A literature review of the potential corrosion and degradation processes applicable to Zircaloy cladding on spent nuclear fuel in a tuff repository. The author concludes that failure due to oxidation of the Zircaloy is not credible, mechanical overload is not a problem, and hydride cracking is not anticipated for the bulk of spent fuel pins. Also, it is concluded that insufficient information exists to evaluate stress corrosion cracking, and some experimental confirmation of crack depths and the effect of slow cooling on the formation of radial hydrides is needed (89 references).

Sutherland, S. H. and D. E. Bennett, September 1979. <u>Defense High-Level Waste</u> and Spent Fuel Characterization for Geologic Waste Repositories. SAND79-0172, Sandia Laboratories, Albuquerque, NM. Results are given for the SANDIA-ORIGEN calculated thermal output and radionuclide content for one spent fuel type from a pressurized water reactor and five spent fuel types from a boiling water reactor (7 references).

Technical Reports Series No. 39, <u>Thermodynamic and Transport Properties of Ura-</u><u>nium Dioxide and Related Phases</u>. Report of the Panel on Thermodynamic and Transport Properties of Uranium Dioxide and Related Phases held in Vienna March 16-20, 1964.

A summary report detailing the knowledge as of 1964 of the structure of UO_2 , thermodynamics, surface and oxidation properties, physical properties, practical implications of the thermodynamic and transport properties, and conclusions (184 references and 12 reports reviewed).

Woodley, R. E., October 1983. <u>The Characteristics of Spent LWR Fuel Relevant</u> to Its Storage in Geologic Repositories. HEDL-TME 83-28, Hanford Engineering Development Laboratory, Richland, WA.

A summary of data on spent LWR fuel (Turkey Point). Fuel rod cladding integrity and conditions within fuel rods established during reactor residence as they affect storage in geologic repositories are emphasized.

EXPERIMENTAL METHODS OF CHARACTERIZATION

Davis, R. B. (HEDL), V. Pasupathi, D. E. Stellrecht (BCL), R. L. Fish (HEDL). 1980. "Remote Characterization of Spent LWR Fuel for Geologic Disposal Demonstrations," <u>Spent-Fuel Handling and Storage</u>. Trans. of Am. Nuc. Soc., Vol. 34, p. 838-839.

A discussion of the remote handling and examination of spent fuel conducted to determine the effect of geologic disposal demonstration tests on the ability of spent fuel to contain radionuclides (1 reference).

De Raedt, Ch., J. L. Genicot, L. Leenders, October 1-5, 1979, <u>Dosimetry Methods</u> for Fuels, <u>Cladding and Structural Materials</u>. <u>Proceedings of the Third ASTM-</u> <u>Euratom Symposium on Reactor Dosimetry</u>, Eur 6813 EN-FR, Ispra (Varese), Italy.

Brief discussions of the experimental techniques for nondestructive testing of nuclear fuel assemblies. The aim of the examinations is to obtain information on burnup, power distribution, fast neutron fluence, and gadolinium burnup. The techniques discussed are gamma spectroscopy in hot cells and underwater in storage pools, and neutron radiography (3 references). Fish, R. L., R. B. Davis, V. Pasupathi (BCL), and R. W. Klingensmith (BCL). March 1980. <u>Spent Fuel Characterization for the Commercial Waste and Spent</u> <u>Fuel Packaging Program</u>. HEDL-TC-1384, Hanford Engineering Development Laboratory, Richland, WA.

A discussion of the rationale for spent fuel characterization and 20 detailed procedures for examination of spent nuclear fuel (5 references).

Jenson, E. D., September 1982. <u>Analytical Techniques for Characterization of</u> <u>Spent LWR Fuels with Respect to Properties Pertinent to Leaching and Dissolu-</u> tion. HEDL-7206, Hanford Engineering Development Laboratory, Richland, WA.

The report identifies the following as key characterization needs with respect to understanding leaching and dissolution behavior from spent LWR fuel: 1) the surface area exposed to water, 2) grain boundary conditions, 3) nature of fission product phases, 4) residence position of fission products, and 5) surface oxidation state.

The techniques to evaluate these five characteristics are discussed. The necessary precision of the measurements and limitations of the equipment are compared (11 references).

Post-irradiation Examination. Proceedings of the Conference held May 13-16, 1980 in Grange-over-Sands. British Nuclear Energy Society, London, England.

This volume contains several articles in the following subject areas:

- achievements and aims of post-irradiation examination
- new and improved nondestructive techniques
- data handling
- destructive techniques
- recent operational and management experience with established postirradiation examination facilities
- recently built, commissioned or refurbished facilities
- future trends.

It also contains brief papers on nondestructive techniques and destructive techniques. The papers cover the European and U.S. programs in these areas to the date of the conference.

Untermyer, S., II, January 1983. <u>Development and Test of Methods for the Non-</u> <u>destructive Assay of Spent-Fuel Assemblies</u>. NP-2812, Research Project 1578-1. Prepared by National Nuclear Corporation, Mountain View, CA.

Several methods were tested for the nondestructive assay of irradiated nuclear fuel assemblies stored in water-filled pools: 1) burnup measurement by neutron emission, 2) residual fissile measurement by multiplication change with boron displacement, and 3) residual fissile measurement by neutron-source multiplication. It was found that both burnup and

residual fissile content could be measured through counting neutron emission from the fuel, both with and without a neutron source.

SPECIFIC DATA FROM TURKEY POINT AND H. B. ROBINSON FUEL CHARACTERIZATION

Atkin, S. D. 1981. Destructive Examination of 3-Cycle LWR Fuel Rods from Turkey Point Unit 3 for the Climax - Spent Fuel Test. HEDL-TME 80-89, Hanford Engineering Development Laboratory, Richland, WA.

The destructive examination of five LWR rods from the Turkey Point Unit 3 reactor are presented. The examinations included fission gas collection and analyses, burnup and hydrogen analyses, and a metallographic evaluation of the fuel, cladding, oxide, and hydrides. The rods exhibited a low fission gas release with all other results appearing representative for pressurized water reactor fuel rods with similar burnups (28 GWd/MTU) and operating histories (11 references).

Barner, J. O. 1984. <u>Characterization of LWR Spent Fuel MCC-Approved Testing</u> Material - ATM-101. PNL-5109, Pacific Northwest Laboratory, Richland, WA.

Characterization of the MCC testing material ATM-101 (i.e. H. B. Robinson spent fuel rods) is provided in the following areas:

- reactor, assembly, and rod descriptions
- assembly B0-5 irradiation history
- a description of unusual incidents that occurred to the rods
- fission gas release measurements
- results of ceramography/metallography examinations
- fuel burnup measurement results and correlations
- results of gamma scanning
- calculated values of the radionuclide inventory
- results of radionuclide chemical analyses.

Results from other studies and the distribution of ATM-101 is included. Intended to be a "living" document; will be updated as new information becomes available (6 references).

Davis, R. B. and V. Pasupathi. 1981. <u>Data Summary Report for the Destructive</u> <u>Examination of Rods G7, G9, J8, I9, and H6 from Turkey Point Fuel Assembly</u> <u>B17</u>. HEDL-TME 80-85, Hanford Engineering Development Laboratory, Richland, WA.

Destructive examination results of five spent fuel rods from a Turkey Point Unit 3 pressurized water reactor. Examinations included fission gas analysis, cladding hydrogen content analysis, fuel burnup analysis, metallographic examinations, autoradiography and shielded electron microprobe analysis. All rods were found to be of sound integrity with an average burnup of 27 GWd/MTU and a 0.3% fission gas release (3 references).

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Davis, R. B. 1981. <u>Pre-Test Nondestructive Examination Data Summary Report on</u> <u>Turkey Point Spent Fuel Assemblies DO1, DO4 and DO6 for the Climax--Spent</u> <u>Fuel Test. HEDL-TME 80-83, Hanford Engineering Development Laboratory,</u> <u>Richland, WA.</u>

Fuel assembly sip testing concluded no leaking rods were among the thirteen fuel assemblies (Turkey Point fuel) included in the Climax--Spent Fuel Test. Detailed nondestructive examination of three (DO1, DO4, and DO6) of the thirteen assemblies is presented (5 references).

Davis, R. B. 1980. Data Report for the Nondestructive Examination of Turkey <u>Point Spent Fuel Assemblies B02, B03, B17, B41 and B43. HEDL-TME 79-68,</u> Hanford Engineering Development Laboratory, Richland, WA.

Fuel assembly sip testing concluded that assembly B17 (Turkey Point fuel) had no leaking rods. Detailed nondestructive examination of five (B02, B03, B17, B41 and B43) of the thirteen assemblies mentioned in the previous article is presented (no references).

- Wilson, C. N. 1985. <u>Microstructural Characteristics of PWR Spent Fuel Rela-</u> <u>tive to its Leaching Behavior</u>. HEDL-SA-3313, Presented at The American Ceramic Society 87th Annual Meeting, Cincinnati, OH.
 - Microstructural, compositional and thermochemical properties of H. B. Robinson spent fuel are discussed relative to its potential performance as a high-level nuclear waste form under proposed tuff repository conditions. Pre- and post-test microstructural characterization data and selected summary radionuclide release data are presented (7 references).

MODELING ACTIVITIES PERTINENT TO SPENT FUEL CHARACTERIZATION

Edlund, Ove. 1983. Calculation of Activity Content and Related Properties in <u>PWR and BWR Fuel Using ORIGEN 2</u>. Studsvik Arbetsrapport--Technical Report, 83-03-07, NW-82/191, Nykoping, Sweden.

This report lists the conditions for calculations of the core inventory for a PWR and BWR. The calculations have been performed using the computer code ORIGEN 2. The amount (grams), the total radioactivity (bequerels), the thermal power (watts), the radioactivity from alpha decay (bequerels), and the neutron emission (neutrons/sec) from the core after the last burnup are determined.

All parameters are calculated as a function of burnup and the natural decay, the latter over a period of 0 - 1.0E07 years. The calculations are performed for 68 heavy nuclides, 60 daughter nuclides, to the heavy nuclides with atomic numbers under 92, 852 fission products, and 7 light nuclides (18 references).

Fields, S. R. 1982. SAM, a Computer Model to Determine the Effective Surface Area of a Spent Fuel Pellet Immersed in Water. HEDL-7208, Hanford Engineering Development Laboratory, Richland, WA.

A mathematical simulation model, SAM (Surface Area Model), was developed to determine the effective surface area of a cracked, porous spent fuel pellet immersed in water. The primary immediate application of SAM is to determine the "time zero" or initial effective surface area of spent fuel, in underground storage, available for contact with ground water after a hypothetical breach of the waste package (3 references).

Notley, M. J. F. 1979. "ELESIM: A Computer Code for Predicting the Performance of Nuclear Fuel Elements." Nuclear Technology 44:455.

The ELESIM code models a single fuel element in a one-dimensional axisymmetric manner. The constituent subroutines are physically based (rather than empirical) models, and include such phenomena as fuel-to-sheath heat transfer; temperature and porosity dependence of fuel thermal conductivity; burnup-dependent neutron flux depression; burnup- and microstructuredependent fission product gas release; fuel thermal expansion, swelling and densification; and stress-, dose-, and temperature-dependent agreement with experimental data (25 references).

FUEL RESTRUCTURING AND RADIONUCLIDE REDISTRIBUTION

The informal WPP library at PNL contains the following articles which present several aspects of fuel restructuring and radionuclide redistribution within spent fuel. These articles are of particular importance in developing models, such as those described in the last section, of the restructuring and redistribution in terms of easily measured characteristics of the fuel (i.e. burnup, linear power, peak temperatures.) Without the ability to model these phenomena, the scope of the characterization would have to be greatly increased to properly account for the many different fuel types and fuel histories.

Baker, C. 1976. "The Fission Gas Bubble Distribution in Uranium Dioxide from High Temperature Irradiated SGHWR Fuel Pins." <u>Journal of Nuclear Materi-</u> als. 66:283-291.

Adamson, M. G. and S. Vaidyanathan (G.E. Sunnyvale). 1981. "Mechanistic Models for Cesium Thermomigration and Cesium-Fuel Chemomechanical Interactions in Mixed-Oxide Fuel Pins." <u>Transactions of the American Nuclear Society</u>. 38:289-291.

- Besmann, T. E., and T. B. Lindemer. 1977. "Chemical Thermodynamics of the System Cs-U-Zr-H-I-O in the Light Water Reactor Fuel-Cladding Gap." <u>Nuclear</u> Technology. 40:297.
- Bramman, J. I., R. M. Sharpe, D. Thom and G. Yates. 1968. "Metallic Fission-Product Inclusions in Irradiated Oxide Fuels." Journal of Nuclear Materials. 25:201-215.
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- Bray, L. A., L. L. Burger, L. G. Morgan, and D. L. Baldwin. 1983. <u>Thermal</u> <u>Release of Volatile Fission Products from Irradiated Nuclear Fuel</u>. <u>PNL-4488</u>, Pacific Northwest Laboratory, Richland, WA.
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- Forsberg, K. and A. R. Massih. 1985. "Fission Gas Release Under Time-Varying Conditions." Journal of Nuclear Materials. 127:141-145.
- Friskney, C. A. and J. A. Turnbull. 1979. "The Characteristics of Fission Gas Release from Uranium Dioxide During Irradiation." <u>Journal of Nuclear Materi-</u> als. 79:184-198.
- Guardini, S. and G. Guzzi. 1981. "Post Irradiation Analysis of BWR and PWR Fuel Samples: Experimental Results and Their Interpretation." "Nuclear Materials Management." In <u>Proceedings of the Institute of Nuclear Materials</u> <u>Management Conference</u>, San Francisco, CA. pp. 90-94.

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- Lorenz, R. A., J. L. Collins, A. P. Malinauskas, O. L. Kirkland, and R. L. Towns. 1980. <u>Fission Product Release From Highly Irradiated LWR</u> <u>Fuel</u>. NUREG/CR-0722 ORNL/NUREG/TM-287/R1, Oak Ridge National Laboratory, Oak Ridge, TN.
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OUT-OF-REACTOR STORAGE EXPERIENCE

The informal WPP library at PNL contains the following articles on the behavior of spent fuels in dry and wet (pool) storage. These articles may be of particular importance if it is deemed necessary to predict the integrity of the spent fuel cladding (i.e. a barrier to the release of radionuclides) over a geologic time period.

Bailey, W. J., and A. B. Johnson, Jr. 1984. Surveillance of LWR Spent Fuel in Wet Storage. EPRI NP-3765, Electric Power Research Institute, Palo Alto, CA.

- Bailey, W. J. and A. B. Johnson, Jr. 1983. <u>Wet Storage Integrity Update</u>. PNL-4726, Pacific Northwest Laboratory, Richland, WA.
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