ORAU Team Dose Reconstruction Project for NIOSH Y-12 National Security Complex – Occupational Internal Dose	Document Number: ORAUT-TKBS-0014-5 Effective Date: 02/14/2006 Revision No.: 01 PC-3 Controlled Copy No.: Page 1 of 46
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Document OwnerApproval:Signature on File William E. Murray, TBD Team LeaderDate:04/19/2005Approval:Signature on File Judson L. Kenoyer, Task 3 ManagerDate:04/20/2005	Revision No.: 00
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RECORD OF ISSUE/REVISIONS

ISSUE			
AUTHORIZATION DATE	EFFECTIVE DATE	REV. NO.	DESCRIPTION
03/17/2004	03/17/2004	00	New document to establish TBD for the Y-12 National Security Complex – Occupational Internal Dose. First approved issue. Initiated by William E. Murray.
05/10/2005	05/10/2005	01	First revision to include table numbers and titles in the body of the document. Approved issue of Revision 01. Initiated by William E. Murray.
05/10/2005	10/11/2005	01 PC-1	Page change initiated to incorporate the definition of U.S.C. on page 6 and details for the definition of a DOE facility on page 7. No sections were deleted. First approved page change revision for Revision 01. Initiated by William E. Murray. Approval:
			Signature on File 10/04/2005 Judson L. Kenoyer, Task 3 Manager Signature on File 10/04/2005
			Richard E. Toohey, Project Director Signature on File10/11/2005 James W. Neton, Associate Director for Science
05/10/2005	01/12/2006	01 PC-2	No changes occurred as a result of formal internal review. Page change initiated to make minor changes to text in Section 5.1, page 7. Deletes and adds text about the default value for the Th-228/Th-232 equilibrium ratio for lung counts in Section 5.3, page 31. Adds new Figure 5.4 variation of activity in natural thorium after a single purification on page 31. Also, changes date in C. M. West reference on page 36. No sections were deleted. Training required: As determined by the Task Manager. Initiated by William E. Murray. Approval:
			Signature on File01/11/2006William E. Murray, TBD Team Leader
			Signature on File 01/10/2006 Judson L. Kenoyer, Task 3 Manager
			Signature on File 01/12/2006 Kate Kimpan, Project Director
			Signature on File 01/12/2006 James W. Neton, Associate Director for Science

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05/10/2005	02/14/2006	01 PC-3	Approved page change revision. initiated to incorporate definitions for dose reconstruction for non-pr cancers that are excluded from th 1947 Special Exposure Cohort. A are: page 7, page 9 in Section 5 and 21 in Section 5.2.1.1, and pa Reference Section. No sections Training required: As determined Manager. Initiated by William E. Approval:	and directions resumptive ne 1943 through Affected pages .1, pages 20 ige 36 in the were deleted. d by the Task
			Signature on File William E. Murray, TBD Team Leader	_02/09/2006
			Signature on File Judson L. Kenoyer, Task 3 Manager	02/08/2006
			Signature on File Kate Kimpan, Project Director	02/07/2006
			Signature on File James W. Neton, Associate Director for	02/14/2006 Science

ACRONYMS AND ABBREVIATIONS

AMAD	aerodynamic median activity diameter
Bq	becquerel
CEDE	committed effective dose equivalent
Ci	curie
cpm	counts per minute
d	day
DF	decontamination factor
dpm	disintegrations per minute
DU	depleted uranium (less than 0.72% ²³⁵ U)
EU	enriched uranium
g	gram
hr	hour
HEU	highly enriched uranium (generally between 20% and 90% ²³⁵ U)
HPGe	high-purity germanium (detector)
ICRP in	International Commission on Radiological Protection inches
keV	kilovolt-electron, 1,000 electron volts
kg	kilogram
L	liter
L _C	critical level (commonly 5% chance of a false positive)
L _D	detection level (commonly 5% chance of a false negative)
LEU	low enriched uranium (generally less than 20% ²³⁵ U)
MDA	minimum detectable activity
mg	milligram
min	minute
mL	milliliter
mm	millimeter
MPBB	maximum permissible body burden
MPLB	maximum permissible lung burden
mrad	millirad
mrem	millirem
nCi	nanocurie
NIOSH	National Institute for Occupational Safety and Health
NU	natural uranium
ORAU	Oak Ridge Associated Universities
ORNL	Oak Ridge National Laboratory

pCi	picocurie
ppb	parts per billion
ppm	parts per million

- RWP Radiation Work Permit
- RU recycled uranium
- SECSpecial Exposure CohortSMCIdaho Specific Manufacturing Capability
- U.S.C. United States Code
- VHEU very highly enriched uranium (greater than 90% ²³⁵U)
- wt % weight percent
- yr year
- μCi microcurie μg microgram μm micrometer

5.1 INTRODUCTION

Technical basis documents and site profile documents are general working documents that provide guidance concerning the preparation of dose reconstructions at particular sites or categories of sites. They will be revised in the event additional relevant information is obtained about the affected site(s). These documents may be used to assist the National Institute for Occupational Safety and Health (NIOSH) in the completion of the individual work required for each dose reconstruction.

In this document the word "facility" is used as a general term for an area, building, or group of buildings that served a specific purpose at a site. It does not necessarily connote an "atomic weapons employer facility" or a "Department of Energy [DOE] facility" as defined in the Energy Employees Occupational Illness Compensation Program Act [EEOICPA; 42 U.S.C. § 7384I(5) and (12)]. EEOICPA defines a DOE facility as "any building, structure, or premise, including the grounds upon which such building, structure, or premise is located … in which operations are, or have been, conducted by, or on behalf of, the Department of Energy (except for buildings, structures, premises, grounds, or operations … pertaining to the Naval Nuclear Propulsion Program)" [42 U.S.C. § 7384I(12)]. Accordingly, except for the exclusion for the Naval Nuclear Propulsion Program noted above, any facility that performs or performed DOE operations of any nature whatsoever is a DOE facility encompassed by EEOICPA.

For employees of DOE or its contractors with cancer, the DOE facility definition only determines eligibility for a dose reconstruction, which is a prerequisite to a compensation decision (except for members of the Special Exposure Cohort). The compensation decision for cancer claimants is based on a section of the statute entitled "Exposure in the Performance of Duty." That provision [42 U.S.C. § 7384n(b)] says that an individual with cancer "shall be determined to have sustained that cancer in the performance of duty for purposes of the compensation program if, and only if, the cancer ... was at least as likely as not related to employment at the facility [where the employee worked], as determined in accordance with the [probability of causation] guidelines established under subsection (c)" [42 U.S.C. § 7384n(b)]. Neither the statute nor the probability of causation guidelines (nor the dose reconstruction regulation) define "performance of duty" for DOE employees with a covered cancer or restrict the "duty" to nuclear weapons work.

As noted above, the statute includes a definition of a DOE facility that excludes "buildings, structures, premises, grounds, or operations covered by Executive Order No. 12344, dated February 1, 1982 (42 U.S.C. 7158 note), pertaining to the Naval Nuclear Propulsion Program" [42 U.S.C. § 7384I(12)]. While this definition contains an exclusion with respect to the Naval Nuclear Propulsion Program, the section of EEOICPA that deals with the compensation decision for covered employees with cancer [i.e., 42 U.S.C. § 7384n(b), entitled "Exposure in the Performance of Duty"] does not contain such an exclusion. Therefore, the statute requires NIOSH to include all radiation exposures in its dose reconstructions for employees at DOE facilities, including radiation exposures related to the Naval Nuclear Propulsion Program. As a result, all internal and external dosimetry results are considered valid for use in dose reconstruction. No efforts are made to determine the eligibility of any fraction of total measured exposure for inclusion in dose reconstruction.

At the Y-12 Plant, now the Y-12 National Security Complex, uranium isotopes in various chemical and physical forms have been the primary contributors to internal radiation doses to workers since November of 1943. Y-12 missions involved other radionuclides and, at times, some uranium compounds could have contained impurities with radiological health implications. These implications and default analyses are discussed in this report. However, the primary focus on internal dose control has been on uranium compounds and alloys over a wide range of ²³⁵U enrichment. Therefore, this

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section begins with an overview of concepts that apply broadly to the history of uranium work at Y-12. Attachment 5A summarizes some of the key points from the following sections.

There were programs in the early years of Y-12 operations for which the currently available information is insufficient to provide general guidance for internal dose reconstruction. At the peak of U-235 enrichment operations in 1943-45, there were 1156 "large mass spectrographs," called "calutrons" in operation. This enrichment production technique was shut down in 1946.

It has been determined that employees who were involved in the calutron uranium enrichment process were exposed to levels of airborne uranium products that cannot be determined because of the absence of bioassay data for the period and the lack of air sampling sufficient to develop maximum exposure scenarios.

Such doses can not be estimated based on source term and process information for lack of documentation on the varying levels of enrichment of the source materials and on the production rate of the operations. Furthermore, the manual recycling and cleaning activities are unique and not comparable to any operations for which NIOSH has access to adequate monitoring data.

The occupational environmental internal dose from uranium can be calculated using data provided in the Y-12 National Security Complex – Occupational Environmental Dose (ORAU 2005).

However, the calutrons in Building 9204-3 were left operational for research and development in improving the production of the calutron concept and in separating other element isotopes. Eight calutrons were still operating as late as 1997. This program produced an inventory of 225 isotopes from nearly every element on the periodic chart.

The 86-inch Cyclotron started operation on November 11, 1950, and operated until the early 1980s. This cyclotron is no longer in operation. Polonium isotopes and alpha airborne activity are the mentioned internal dose concerns. The 86-inch cyclotron was later used to create neutron-deficient radionuclides as a part of the R&D program above. There were programs in these R&D efforts, which involved plutonium, and workers could have plutonium bioassay results in their records. When claim information indicates that a Y-12 worker was involved with research activities involving the Calutron, cyclotron (accelerator), fusion work, or plutonium (except in the case of recycled uranium (RU) exposure, which is addressed in this section), consideration must be given to possible exposure to radionuclides other than uranium. (Wilcox et al. 1999)

5.1.1 Uranium Solubility in the Lung

The uranium compounds with which Y-12 has worked ranges from highly soluble to very insoluble. Health physicists have long recognized that nephrotoxicity is the primary hazard associated with soluble uranium compounds (depleted through low enrichment). According to Sterner and Riley (1946), exposures to soluble compounds were monitored from the closing days of World War II by clinical tests of renal function and by fluorometric tests for uranium in urine. However, no urinalysis

data have been found for Y-12 prior to 1948, and this issue will have to be addressed in a revision of this document.

Health physicists also recognized early that the lung is the primary organ of concern for the less soluble uranium compounds. During the greater part of the plant's history, control measures for less soluble compounds were guided by the lung and metabolic models presented in the International Commission on Radiological Protection (ICRP) Publication 2 (ICRP 1960) or by similar predecessor

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models. These models related lung dose to uranium excretion in urine, particularly under chronic exposure conditions.

It is noteworthy that insoluble material as described in ICRP Publication 2 would be classified as moderately soluble type M material in the modern ICRP Publication 66 framework (ICRP 1994). It is also noteworthy that ICRP has recognized that some forms of uranium are less soluble than initially believed. Figure 5-1 shows normalized lung retention for ICRP Publication 2 insoluble material, ICRP Publication 30 (ICRP 1979) class W particles, class Q material as described in the 1990s (Barber and Forrest 1995), and the Publication 66 type M particles. The boxes on the diagram define the modern clearance types F (fast), M (moderate), and S (slow) as a function of lung retention and time as illustrated in Annex D of ICRP Publication 71 (ICRP 1995). The ICRP 2 insoluble model and the class W models fit within the range of type M material.

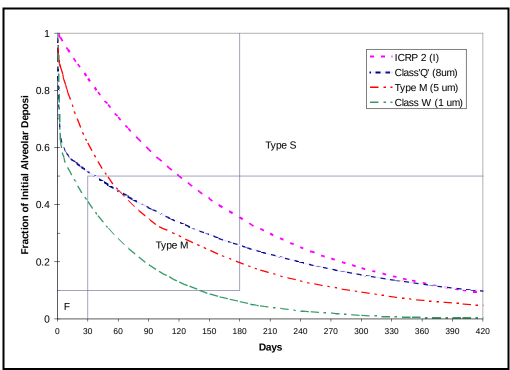


Figure 5-1. Historical solubility types meeting the type M definition.

After *in vivo* lung counting was begun at Y-12 in 1961, it was recognized that a few workers had uranium lung burdens in considerable excess of the amount indicated by urinalysis. Writing in 1969, West and Scott (1969) noted that, "5 exposure cases noted in 1962 and 1963 showed chest burdens of enriched uranium [EU] which exhibited unexpectedly slow decreases after the persons were assigned to non-uranium activities" (West and Scott 1969). The job description of all five individuals was Process Operator (Snapp 2003).

Subsequent investigations suggested that three process combustion ashes from the EU foundry were the most likely exposure materials for these five cases (Steckel and West 1966, p. 31). The particular thermal histories and particle size distributions of these process ashes were considered to be important factors leading to the low solubility in simulated lung fluid (Steckel and West 1966, p. 30).

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Particle sizes tended to be relatively larger (0.86 to 1.6 μ m), and the solubility tended to be lower, for materials processed at higher temperatures.

While the exceptional cases with unusually protracted lung clearance are important, it is more important to note that, for the vast majority of individuals, lung clearance took place in approximate accordance with the ICRP Publication 2 (1960) insoluble model, which fits within the current type M framework.

In 1964, a composite urinary excretion curve was described for 157 workers who had been removed from uranium work because of high uranium urinalysis results (Scott 1964). Figure 5-2 shows this empirical composite curve together with expected excretion curves for types F, M, and S uranium. The latter curves are derived from 5-µm aerodynamic median activity diameter (AMAD) excretion curves (Potter 2002). They were normalized to match the Scott curve at 200 d after removal from uranium work areas. The empirical excretion curve matches the type M curve much better than the type F or S curves. Indeed, the Scott curve fits the type M curve quite well over most of the 420-d time range. These results are consistent with the subsequent conclusions of Rucker et al. (2001) that class W is the appropriate default for uranium at Bechtel Jacobs facilities, including Y-12. Exceptions to this default included *high-fired* uranium oxides and compounds formed by the slow oxidation of uranium metal.

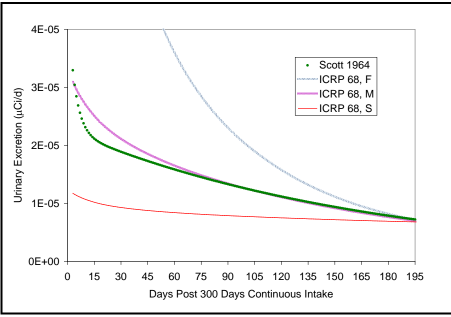


Figure 5-2. Empirical and theoretical uranium excretion after uranium restriction.

Beginning in September 1994, the plant was placed in a stand-down mode. This effectively stopped all routine work (and chronic exposure potential) in the process areas. Only minimal walkthroughs and area checks continued during the stand-down, which lasted until August 1998. The stand-down significantly influenced the available source term for exposures. Before stand-down, the Y-12 program default modeling assumption was class Q (90% Super-W, 10% Y). During the stand-down, the Y-12 program default assumption of chronic exposure was modified to assume acute exposures occurring at the midpoint of a quarterly sampling frequency. No wet chemical operations were performed during this 4-yr period. Therefore, no soluble component was being produced and materials that were stopped in process were allowed to oxidize, which provided a larger component of

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type S material as an exposure source. In August 1998, a partial resumption of activities occurred; wet chemical operations were not restarted and were still not restarted as of October 2003. The predominant material encountered after partial restart in August 1998 is uranium oxide, which fecal sampling has shown to be more consistent with 100% type S material. Based on the changing workplace conditions after partial restart in 1998, the default assumptions were modified to return to chronic exposure but to use type S solubility.

For a workplace as varied as Y-12, it is clear that no single solubility or particle size would apply to all workers. Furthermore, accurate assignment of the uranium lung clearance type to a given bioassay result was considered virtually impossible because of uncertainties about chemical form and limitations of the personnel-tracking system (Barber and Forest 1995, p. 669). As of 2001, Y-12 implemented a radiation work permit (RWP) tracking system that tracks the location of workers, their type of work, and the type of bioassay needed (ORAU 2003, pp. 19 ff.).

Exposure to type M material from 1948 to July 1998 appears to be the more likely absorption type. After July 1998, exposure to absorption type S material is more likely. However, the absorption type can be based on the monitoring data, claimant-favorable assumptions, or both.

5.1.2 Particle Size Information

Several particle size studies in uranium process areas have been conducted at Y-12 (e.g., CCCC 1953; Steckel and West 1966; Barber and Forest 1995; BWXT Y-12 2003a). For different times and different processes, reported particle sizes ranged from less than 1 to over 10 μ m (physical). Steckel and West (1966) reported a positive correlation between uranium oxide particle size and process temperature. Barber and Forest (1995) used an 8- μ m AMAD based on particle size measurements as the basis for the class Q dosimetry system used in the 1990s. (The class Q system is further described as 10% class Y and 90% modified class W. The modification consists of increasing the class W 50-d compartment to 120 d.) In terms of lung deposition and retention, the 8- μ m AMAD class Q material is closer to 5- μ m AMAD type M material than to 1- μ m AMAD class W material, as shown in Figure 5-3.

Since 2000, Y-12 has implemented the latest guidance provided by the ICRP including the ICRP Publication 66 (ICRP 1994) lung model and ICRP Publication 78 (ICRP 1998) methods and models (BWXT Y-12 2003b, p. 89). This is the justification for the change to the 5- μ m AMAD as the default particle size. Y-12 uses the newer weighting factors proposed by the ICRP in its 1990 recommendations (ICRP 1991). For Y-12 dose reconstructions, the default particle size distribution is 5 μ m AMAD.

5.1.3 Uranium Alpha Activity as a Function of Enrichment

At Y-12, uranium enrichments range from depleted uranium (DU) [less than 0.72 wt % ²³⁵U] to very highly enriched uranium (VHEU; greater than 93 wt % ²³⁵U). Section 5.1.4.1 lists the uranium isotopic mixes with recycled uranium (RU) contaminants that were typical of uranium materials at Y-12. The potential dosimetric complexity is simplified by the similarity of the dose conversion factors for the most important uranium isotopes and by the quantities measured in the *in vitro* bioassay and chest-counting programs.

For work with enriched compounds and alloys, urinalysis programs after 1950 reported either total uranium alpha counts or isotopic uranium results, both of which reflect the dosimetric potential of the isotopes generally handled. Early *in vivo* chest-counting results were reported either as micrograms of ²³⁵U or milligrams of ²³⁸U based on whether an exposure was believed to be from general enriched

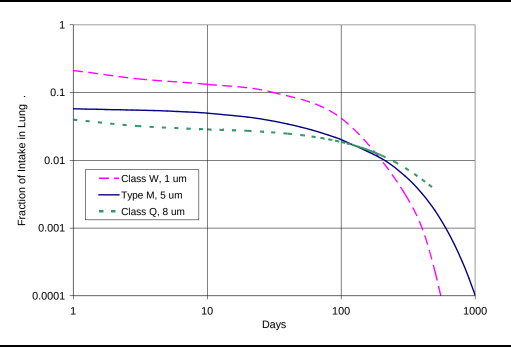


Figure 5-3. Lung deposition and retention of class W, class Q, and type M uranium. (Particle sizes specified are in micrometers AMAD.)

uranium (EU), natural uranium (NU), or DU. Based on these reporting techniques, the following assumptions can be made about data interpretation for the periods from 1950 through 1989 for urinalysis and from 1961 to 1989 for lung counting.

For lung counts a combination of the information in the Type Analysis and Material Type (see Section 5.A.1.3 for details) reporting fields can be used to determine whether the count was believed to be due to NU or DU. For records through 1971, a Type Analysis of 1 with a Material Type of 2 or 3 indicates DU, while a Material Type of 7 indicates NU. For records after 1971, a Type Analysis of 4 with a Material Type of 2 or 3 indicates DU, while a Material Type of 2 or 3 indicates DU, while a Material Type of 2 or 3 indicates DU, while a Material Type of 7 indicates DU, while a Material Type of 7 indicates DU, while a Material Type of 7 indicates NU.

able 5-1. Enforment assumptions for dose assessment for various analytical techniques.							
Analytical technique (Urinalysis 1948–present) (Lung count 1961–1989)	Measurement information	Default enrichment assumption					
Urine by fluorometry (usually for an individual who worked in areas with NU or DU; before 1950, also used for workers in EU areas.	Alpha dpm can be calculated from mass.	Before 1950, 93% enrichment. After 1949, NU.					
Urine by gross alpha counting (usually for an individual who worked in areas with EU).	Alpha dpm is the sum of all uranium alphas.	93% enrichment.					
Urine by alpha spectrometry (10/1989 to present).	Alpha dpm/d is reported isotopically and total uranium is determined by summing isotopic results.						
Lung count	μg U-235	93% enrichment.					
Lung count	μg U-238	NU.					

Table 5-1	Enrichment assum	ptions for dose asses	sment for various a	analytical techniques.

At Y-12, it is current practice to assign dose by applying the dose conversion factor for ²³⁴U to all inferred uranium intakes. This practice is slightly conservative for EU; the resultant overestimate is less than 1% of the effective dose. For DU, the use of the ²³⁴U dose conversion factor leads to an overestimate of 16% for type S uranium and 20% for type M uranium (Eckerman and Kerr 1999, p. 16).

5.1.4 <u>Temporal Pattern of Uranium Exposures</u>

As a rule, routine uranium exposures at Y-12 were considered to be of a chronic nature. For example, in 1958, Patterson stated, "...our interpretation of urinalysis results and our assignment of internal dose assume an exposure under equilibrium conditions of intake and elimination" (Patterson 1958, p. 58). For the stand-down period from September 1994 until August 1998, acute exposures should be assumed to be the more likely mode of exposure. After this time, while complete equilibrium is not expected in modern internal dosimetry models, the presumption of chronic exposure conditions for uranium remains in place as noted in a 2003 report: "The most likely exposure potential for uranium work at the Y-12 Complex is chronic in nature" (BWXT Y-12 2003b, p. 107). Because workers were assigned to various buildings and processes throughout their employment, it is not practical to characterize intakes based on historical operational assignments. The claimant files do not generally contain detailed work assignments, but do contain all bioassay data for each individual worker during the years of operational exposure - primarily in vitro (urine) and in vivo (lung counts) bioassay. Radiological materials, including isotopic mixes and so forth, are not defined by operational area or building in this document, because internal dose reconstruction relies primarily on the recorded bioassay data and the defaults for unmonitored isotopes that were conservatively established on a plant-wide basis (see Table 5-8 later in this document for plant wide RU contaminant defaults).

5.1.5 Other Radionuclides of Concern Including Recycled Uranium Contaminants

In addition to ²³⁴U, ²³⁵U, and ²³⁸U, the following radionuclides were identified in the Y-12 Technical Basis Document for Internal Dosimetry (BWXT Y-12 2003a):

- ³H, ⁹⁰Sr, ⁹⁹Tc
- ²²⁸Th, ²³²Th
- ²³²U, ²³³U, ²³⁶U
- ²³⁸Pu, ²³⁹Pu, ²⁴¹Pu
- ²³⁷Np, ²⁴¹Am

The following radionuclides that could interfere with *in vivo* analysis of uranium and thorium were quantified so that their effects on the spectra could be taken into account:

• ⁴⁰K, ¹³⁷Cs

Other radionuclides addressed included:

- ⁶⁰Co and ⁹⁵Zr/⁹⁵Nb for organizations outside Y-12 (Cofield 1961)
- ²¹⁰Po

Many of these radionuclides are accompanied by progeny in various stages of equilibrium. In addition, operators of the *in vivo* analysis equipment reviewed accumulating spectra for interferences from medical radioisotopes such as ¹³¹I.

5.1.5.1 Unmonitored Radionuclides from Recycled Uranium

One of the primary sources of significant contaminants (including the production of other than the normal uranium isotopes) that were introduced into the Y-12 plant systems were those associated with RU from 1953 until 1999. These contaminants were a result of fission and activation processes of a variety of uranium enrichment isotope mixtures, which were irradiated in production and test reactors. After completing their useful life in the reactors, the unused uranium in the spent fuel elements or targets was recovered in chemical extraction plants and returned to the inventories in the DOE system along with trace quantities of the contaminants (in some instances U-236 was produced in quantities greater than "trace" levels).

The major source of RU contaminants at Y-12 was in the receipt and processing of returned VHEU from SRS and the Idaho Chemical Processing Plant (ICPP). Tables 5-2 and 5-3 document the quantities of uranium and the calculated mass quantities of plutonium, neptunium, and technetium actually delivered to Y-12. Table 5-4 presents a computer-generated listing of the relative production of a few isotopes of concern, following irradiation of VHEU fuel in a high-flux test reactor. The process streams included a spectrum of chemical forms of uranium, including uranyl nitrate $[UO_2(NO_3)_2]$ solutions, uranium trioxide (UO_3) , uranium-aluminum (U-AI) alloy ingots, uranium scrap, uranium tetrafluoride (UF_4) , uranium metal, solvent extraction raffinates, and a variety of secondary process wastes and residues.

The uranium enrichments ranged from DU, NU, low enriched uranium (LEU - less than 20% ²³⁵U), highly enriched uranium (HEU - from 20 to approximately 93% ²³⁵U), and very highly enriched (VHEU - more than 93% ²³⁵U). The predominant RU and associated contaminants were in HEU and VHEU materials. However, all of the uranium at Y-12 came from other DOE facilities, which had also either generated and/or received RU materials. Therefore, nearly all of the uranium in the DOE facilities contained RU contaminants to varying degrees through being processed in the same equipment, blending with other materials to adjust the degree of enrichment, etc. For example, most of the DU received and processed at Y-12 was seldom chemically processed but was received in forms from which parts were produced by mechanical processing. However, even these materials were received from other plants, such as Fernald, which also had an RU contaminant inventory. Therefore, the DU contaminant levels at Y-12 were inferred from both the Fernald and the Idaho Specific Manufacturing Capability (SMC) projects' recycle reports (ORAUT 2004 and DOE 2000).

The fundamental conclusion from a variety of reports, including a lack of definitive RU contaminant bioassay data, is that for dose reconstruction purposes a default level of RU contaminants should be derived and applied as a percentage increase to the derived uranium intake for each of the four major contaminants. Analytical information derived from a variety of sources allows the calculation or interpolation of the levels of the predominant RU contaminants in the uranium materials received, processed, and handled at Y-12. One of the primary sources of RU contaminant information was the documented report generated as a consequence of a comprehensive study of Recycled Uranium in DOE facilities in 2000. The applicable report for Y-12 was issued in December 2000 and titled Recycled Uranium Mass Balance Project Y-12 National Security Complex Site Report (DOE 2000). It should be noted that there were some recognized inconsistencies in shipping and receipts between the various DOE facilities in the DOE 2000 RU reports. In an attempt to resolve these inconsistencies a summary report in 2003 by the DOE Office of Security, titled Recycled Uranium, United States Production, Enrichment, and Utilization (DOE 2003) was issued though correcting some inconsistencies, there remain significant inconsistencies in the two reports. For purposes of reconstruction of RU contaminant inventories in the Y-12 facilities and internal dose default analyses, the rationale for resolving those inconsistencies is provided herein. One approach to gain an estimate

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of the average contaminant concentrations through the years is to document the total receipts of contaminants and uranium and then to derive the average long-term RU contaminant concentrations.

Tables 5-2 and 5-3 list the documented RU contaminants and their primary source in recycled VHEU. For completeness, other DU receipts are listed by the DOE (2000) report without documenting the associated RU contaminant levels. However, it is known that these uranium receipts also contained low-level RU contaminants, the concentrations of which are discussed later in this report.

Table 5-2. Y-12 Uranium receipts summary –presenting average levels of three predominant recycled uranium contaminants through plant history (1953 to 1999).

		Pu		Np			Тс			
Site	U Kg	g	ppb	pCi/µg U	g	ppm	pCi/µg U	g	ppm	pCi/µg U
Y-12 VHEU Receipts Summary – DOE 2000 RU Report										
SRS	125,161	0.0455	0.36	5.2E-03	3,600	29	2E-02	14,268	114	1.9
ICPP	25,696	0.00124	0.05	7.2E-04	66	3	2E-03	231	9	0.15
EU Total	150,857	0.04674	0.3	4.3E-03	3,666	23	1.6E-02	14,499	96	1.64
DU Receipts S	Summary - D	DOE 2000 R	U Repo	ort						
Hanford	1,502	No Data								
ORGDP	192,836									
PGDP	38,423]								
DU total	232,761									
Grand total	383,618									

a. ICPP = Idaho Chemical Processing Plant; ORGDP = Oak Ridge Gaseous Diffusion Plant; PGDP = Paducah Gaseous Diffusion Plant.

	U		Ρι	1	Np		Тс			
Shipping site	(kg)	g	ppb	pCi/µg U	g	ppm	pCi/µg U	g	ppm	pCi/µg U
SRS	153,000									
ICPP	25,700									
Hanford	4,400									
West Valley	900									
Total	184,000	0.5	2.7	0.039	54	0.003	0.00021	9,100	49.5	0.85

Table 5-3. Total recycled uranium receipts summary.

The results of the DOE Office of Security RU report (DOE 2003) are presented in Table 5-3. As indicated there are significant differences with the results recorded in DOE (2000) report (summarized in Table 5-2) and 5-3. The objective of DOE (2003) was to resolve the differences in the reported shipment amounts from the recycling plants (the "primary sites") with the reported receipts at the first or "primary receiving sites." Also the detail in DOE 2003 related to uranium enrichment and specific RU contaminant concentrations from each site were not provided. (Note the lack of detail in table 5-3.) In examining DOE (2003), the shipments from these "primary receiving sites" (following processing and incorporation of recycled materials into their process streams) to what can be called "secondary" receiving sites were not documented. For example, Y-12 received uranium materials from gaseous diffusion plants, Fernald, and others, which also had RU contaminant inventories. Thus the discrepancies in the current values reported in DOE (2003), while DOE (2000) accounts for "secondary shipments" as well. However, where large differences occur, the most clamant-favorable value is used in deriving the defaults for dose reconstruction, which are generally the results in DOE (2000).

Though plutonium, neptunium, and technetium isotopes were analyzed and documented in all of the facilities as the primary isotopes of concern, there was a significant level of ²²⁸Th detected in nearly all of the facilities at Y-12 and in reported process stream analyses (DOE 2000). To establish the source of this isotope and provide a basis for including it in the RU contaminants to be considered in the internal dose reconstruction, Tables 5-4 and 5-5 are included to demonstrate the production of this

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isotope in VHEU fuels and the relative/effective "enrichment" during chemical reprocessing. The values in Table 5-4 were derived for purposes of the DOE (2000) Recycled Uranium report at the ICPP. The ORIGIN2 computer program results illustrate the production of isotopes during use of VHEU fuels (D.R.Wenzel 2000 and D.R.Wenzel "specific tables" of isotopic production---"). The isotopes of interest for purposes of this illustration are both the uranium isotopes and the plutonium, neptunium, technetium, and thorium. Thorium-228 is a contaminant that became an isotope of concern at Y-12, which is the primary receiving site of recycled VHEU fuel materials.

		Activity	Activity	Percent weight		Activity ratio
Isotope	Half-life (yr)	(µCi/µg Total U)	(µCi/g Total U)	g/100 g U ^a	ppm U	(isotope/U)
Tc-99	2.11E+05	3.8E-04	3.8E+02	2.2E+00	2.2E+04	24
Th-227	18.72 d	3.3E-11	3.3E-05	1.1E-13	1.1E-09	
Th-228	1.913	1.6E-07	1.6E-01	1.9E-08	1.9E-04	0.01
Th-229	7.34E+03	1.4E-12	1.4E-06	6.7E-10	6.7E-06	
Th-230	7.7E+04	1.9E-11	1.9E-05	9.6E-08	9.6E-04	1.2E-06
Th-231	25.52 hr	1.7E-06	1.7E+00	3.2E-10	3.2E-06	0.1
Th-232	1.41E+10	3.5E-15	3.5E-09	3.2E-06	3.2E-02	2.2E-10
Total			1.9			
U-232	6.89E+01	2.7E-07	2.7E-01	1.3E-06	1.3E-02	
U-233	1.59E+05	2.6E-10	2.6E-04	2.7E-06	2.7E-02	
U-234	2.46E+05	6.6E-07	6.6E-01	1.0E+00	1.0E+04	
U-235	7.04E+08	1.7E-06	1.7E-00	7.8E+01	7.8E+05	
U-236	2.34E+07	1.3E-05	1.3E+01	2.0E+01	2.0E+05	
U-237	6.75 d	3.6E-08	3.6E-02	4.4E-11	4.4E-11	
U-238	4.47E+09	5.6E-09	5.6E-03	1.7E+00	1.7E+04	
Total			15.7			
Np-236	1.15E+05	9.4E-11	9.4E-05	7.2E-07	7.2E-03	
Np-237	2.14E+06	9.1E-06	9.1E+00	1.3E+00	1.3E+04	0.6
Np-238	2.117 d	9.2E-10	9.2E-04	3.6E-13	3.6E-09	
Np-239	2.355	4.3E-08	4.3E-02	1.9E-11	1.9E-07	
Total			9.1			
Pu-236	2.851	3.6E-06	3.6E+00	6.8E-07	6.8E-03	0.2
Pu-238	87.74	3.6E-02	3.6E+04	2.1E-01	2.1E+03	2.3E+03
Pu-239	2.41E+04	1.9E-05	1.9E+01	3.1E-02	3.1E+02	1.2
Pu-240	6.54E+03	1.5E-05	1.5E+01	6.4E-03	64	1.0
Pu-241	14.4	1.5E-03	1.5E+03	1.4E-03	14	1.0E+02
Pu-242	3.73E+05	8.8-E-09	8.8E-03	2.3E-04	2.3	5.6E-04
Total			4.6E+04			
Am-241	4.32E+02	9.6E-06	9.6E+00	2.8E-04	2.8	0.6

Table 5-4. Primary contaminants (at 3-yr decay) in VHEU fuels and of concern in internal dose reconstruction.

a. There were basically three types of fuel elements with varying enrichments (both at the start and after burn-up). The typical after burn-up enrichments were 78.21% for aluminum-clad, 78.42% for zirconium-clad, and 51.9% for stainless-steel fuel elements. Values represent the 78% after burn-up enrichments as representative, with typical 1%+ for U-234.

Table 5-5. Comparison of the production of three selected isotopes in VHEU spent fuel with levels reported in HEU shipments to Y-12.

Isotope	μCi/g U product from SRS	μCi/g U product from ICPP	μCi/g U in high burn-up VHEU reactor fuel ^b	Approximate Derived Decontamination Factor (DF)
Th-228	0.01 – 0.03	NA ^a	0.16	10
Np-237	0.01 – 0.02	1E-03	9.1	1E+03–1E+04
Pu-238	4E-03	3E-04	3.6E+04	1E+07–1E+08

a. There were no analytical data for Th-228 reported in the ICPP Y-12 receipt reports. The DF reported is interpolated from Y-12 process stream analytical data. These values were taken directly from column 4 in table 5-4 above.

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Comparing the concentrations in the spent fuels with the reported concentrations in the product received at Y-12 provides a rough indication of the decontamination factors (DF) of the chemical extraction processes. Reference to these DFs in Table 5-5 and the average contaminant concentrations in Table 5-6 indicate that plutonium is the most efficiently removed with an effective DF of approximately 1×10^7 to 10^8 , with a neptunium DF of approximately 1×10^4 , while thorium appears to follow the chemistry of uranium more closely, resulting in a DF of only 10, which in turn results in a relative "enrichment" of ²²⁸Th in the VHEU product received at Y-12. Although ²²⁸Th has not been considered an RU contaminant of concern in other DOE facilities and was not routinely analyzed, there were sample analyses of contaminants in surface smears and wipes in most of the Y-12 plant facilities indicate a predominance of ²²⁸Th as a contaminant. The uranium to non-uranium alpha ratios range from 4 to more than 1×10^4 , which is not useful for dose reconstruction. However, these data do indicate that ²²⁸Th is one of the primary contaminants at Y-12, which in turn provides a technical base for inclusion of ²²⁸Th in the list of RU contaminant isotopes.

Table 5-6. Ranges of reported contaminant values in the main processing facilities (summary of Tables B-3 and B-4 of).^a

Contaminant	Range	Ave. of range	Ave. of range minus the high	Raffinate values ^b
Pu	0.11 - 4.5 ppb	0.5 ppb	0.25 ppb	62 ppb
Np	4.7 – 346 ppm	30.6 ppm	20.8 ppm	2,980 ppm
Тс	0.13 – 211 ppm	83 ppm	66 ppm	641 ppm

a. The highest values reported in the processing facilities were in just one of 17 process streams - that being the solvent extraction purification of UN. Uranium levels are still significant in this stream, which indicates a marked enrichment of contaminants relative to U in this stream.

b. The raffinate streams are depleted in uranium to a range of 5 ppm U levels in the sludge. Therefore the parts per million of the RU contaminants relative to U are not meaningful (i.e., total Pu – maximum 30 and average 21 pCi/g net sludge, Np – maximum 12 and average 8 pCi/g sludge, Tc – maximum 1.2E+4 and average 3.7E+3 pCi/g sludge).

Uranium has always been the dominant contributor to collective internal dose at Y-12. Monitoring for other radionuclides has been performed on a limited basis. Both the relatively small concentrations and the difficulty of analyses have contributed to the lack of data. The isotopic characterization of the large variety of uranium enrichments/contaminants is illustrated in Table 5-7, which is an average representation of six specific generalized enrichment types handled at Y-12. The uranium to non-uranium alpha ratios in Table 5-7 are in the same general range as those measured in surface contamination in the plant facilities after years of operation.

From the values in Table 5-6, it is evident that the maximum of the ranges in the plant tend to run a factor of approximately 10 above the average of the range for both plutonium and neptunium, with technetium at a factor of 3. A factor of 10 is also applied to the thorium levels. The defaults in Table 5-8 were chosen consistent with all of the information discussed above with the objective of representing the maximum values of the documented RU contaminant levels in the Y-12 process streams and thus represent claimant-favorable values, and which can apply on a plant–wide basis. The application of the default level should be based upon the indication by the bioassay as to the likelihood of VHEU/HEU or LEU/NU/DU. The appropriate default ratio should be chosen based on the ²³⁵U or ²³⁸U indicated in the bioassay results. For each microgram or pCi of uranium intake calculated based on bioassay data in the claimant files, an additional intake of each of the four RU contaminants should be added.

From a practical and simplified approach, when the results in the claimant files are given in mass units of uranium per liter or lung count, column 3 values can be used to add the RU contaminant activity to

	Activity		Activity		
Uranium Isotope	(Ci/g)	Mass fraction	(pCi/µg total U)	RU Cont.	Activity (pCi/µg total U)
High burn-up (~50%	b) recycled VHEU from	n SRS			
233	9.66E-03	1.00E-04	9.660E-01	Pu-238	6.20E-03
234	6.22E-03	1.28E-02	7.962E+01	Pu-239	1.00E-04
235	2.16E-06	5.22E-01	1.128E+00	Np-237	2.00E-03
236	6.50E-05	2.92E-01	1.898E+01	Tc-99	1.90E+00
238	3.36E-07	1.73E-01	5.796E-02	Th-228	3.17E-02
Total		9.99E-01	1.007E+02	Total alpha	4.00E-02
dpm/µg total U			2.24E+02	α dpm/µg total U	8.87E-02
U a:Non U a				2.52E+03	
	%) recycled VHEU fro	om ICPP		2:022:00	
233	9.66E-03	3.000E-06	2.90E-02	Pu-238	2.00E-03
234	6.22E-03	1.000E-02	6.22E+01	Pu-239	3.30E-05
235	2.16E-06	7.800E-01	1.68E+00	Np-237	2.00E-03
236	6.50E-05	2.000E-01	1.30E+01	Tc-99	1.50E-01
238	3.36E-07	1.700E-02	7.69E-01	Th-228	5.00E-02
Total	5.50⊑-07	1.007E+00	7.69E+01	Total α	5.40E-02
dpm/µg total U		1.007 L+00	1.71E+02	α dpm/µg total U	1.20E-02
U α :Non U α			1.7 12+02	1.4E+03	1.202-01
Weapons-grade Ora				1.405	
233	9.66E-03	0.000 - 00	0.000,000	Pu-238	2.00E-02
		0.000E+00	0.00E+00	Pu-238 Pu-239	
234	6.22E-03	1.100E-02	6.84E-01		2.00E-02
235	2.16E-06	9.300E-01	2.01E+00	Np-237	6.00E-02
236	6.50E-05	7.500E-03	4.88E-01	Tc-99	5.00E-02
238	3.36E-07	6.000E-02	2.02E-02	Th-228	5.00E-02
Total		1.009E+00	7.09E+01	Total α	1.50E-01
dpm/µg total U			1.57E+02	α dpm/µg total U	3.33E-01
U α:Non U α				4.73E+02	
Recycled NU	-				
233	9.66E-03	0.000E+00	0.00E+00	Pu-238	0.00E+00
234	6.22E-03	5.400E-05	3.36E-01	Pu-239	3.00E-06
235	2.16E-06	7.200E-03	1.56E-02	Np-237	2.40E-05
236	6.50E-05	0.000E+00	0.00E+00	Tc-99	2.00E-04
238	3.36E-07	9.927E-01	3.34E-01	Th-228	3.00E-05
Total			6.85E-01	Total alpha	5.70E-05
dpm/µg total U			1.52E+00	α dpm/µg total U	1.27E-04
U α:Non U α				1.20E+04	
Recycled LEU (2%)					
233	9.66E-03	0.000E+00	0.00E+00	Pu-238	0.00E+00
234	6.22E-03	2.000E-04	1.24E+00	Pu-239	2.00E-04
235	2.16E-06	2.000E-02	4.32E-02	Np-237	2.00E-04
236	6.50E-05	0.000E+00	0.00E+00	Tc-99	9.20E-02
238	3.36E-07	9.798E-01	3.29E-01	Th-228	3.00E-04
Total			1.62E+00	Total alpha	7.00E-04
dpm/µg total U			3.59E+00	α dpm/µg total U	1.55E-03
U a:Non U a				2.31E+03	
Recycled DU				21012100	
233	9.66E-03	0.000E+00	0.00E+00	Pu-238	0.00E+00
234	6.22E-03	1.000E-05	6.22E-02	Pu-239	3.70E-06
235	2.16E-06	2.000E-03	4.32E-03	Np-237	2.70E-05
236	6.50E-05	3.000E-05	1.95E-04	Tc-99	4.00E-04
238	3.36E-07	9.980E-01	3.35E-01	Th-228	4.00E-04
Total	5.502-07	9.900L-01	4.02E-01	Total alpha	7.07E-05
dpm/µg total U			8.93E-01	α dpm/µg total U	1.57E-04
		1	0.305-01		1.0/E-04

Table 5-7. Typical uranium enrichment materials at Y-12 with calculated and inferred RU contaminant
levels.

a.

Values derived from a number of sources and should be considered as largely average values. The recycled DU, recycled NU, and recycled LEU values were derived from Fernald data because this site both supplied and received b. uranium to and from all the other DOE sites, including Y-12. In addition, the recycled DU information was verified by the analytical data reported by the Idaho SMC facility for DOE (2000).

c.

The weapons-grade Oralloy VHEU and other burn-up example VHEU were derived from Y-12 data and ICPP reports. The Oralloy RU contaminant data was presented in the reference reports in very general terms (i.e., <0.1 µCi/g U for total Np and Pu d. and <0.05 µCi/g for "Any other radionuclide").

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Table 5-8. R	Table 5-8. Recommended defaults based on upper levels of expected ranges.				
Isotope	Default levels in pCi/µg U		Default levels in pCi/pCi total U ^c		
	VHEU & Oralloy	LEU, NU, & DU	VHEU & Oralloy	LEU, NU, & DU	
Tc-99	4	0.3	4E-02	0.15	
Th-228 ^a	0.5	0.003	5E-03	1.5E-03	
Np-237	0.6	0.003	6E-03	1.5E-03	
Pu-238 ^b	0.2		2E-03		
Pu-239 ^b	0.2 (Oralloy only)	0.003	2E-03 - Oralloy	1.5E-03	
Total non U	1.3	0.01			
Total U	75	2			

Table 5-8. Recommended defaults based on upper levels of expected ranges.

a. The Th-228 default is justified in concept due to the demonstrated existence of general plant-wide contamination and further indicated by the demonstrated production in VHEU fuels, which in turn is further validated in the process stream analyses documented in DOE (2000).

b. VHEU was used in the metal form at Y-12 and called Oralloy. Plutonium analyses for Oralloy were not isotope specific. For Oralloy VHEU only, Pu-239 is the chosen isotope because the total quantities of recycled VHEU (in which Pu-238 was the predominant plutonium isotope) returned to the gaseous diffusion plants was miniscule compared to the LEU returns.

c. Reference to Table 5-6 indicates activity in picocuries per gram of total U for VHEU ranges from 71 to 101; 100 is used for simplicity. For the LEU, NU, and DU, the picocuries per gram of total U ranges from 0.9 to 1.6; 2 pCi/g total U is used for simplicity.

the intake. However, when the bioassay units are in uranium alpha activity per unit urine volume and/or U isotope activity units, the values in column 4 can be used. Again, Table 5-8 values are maximum values expected (and claimant-favorable), while average values are basically a factor of 10 less than the values for plutonium, neptunium, and thorium and a factor of 3 less for technetium. If a best estimate dose reconstruction is needed, average values (a factor of 10 less than the values in Table 5-8 for plutonium, neptunium, and thorium and a factor of 3 less for technetium) should be used.

Reference to Table 5-8 with relationship to the plutonium contaminant, the recommended isotope Pu-239 should be assumed for Oralloy only, since Oralloy used at Y-12 did not come directly from the primary RU generating sites, i.e. it was enriched at the GDP facilities in which Pu-239 was dominant compared to Pu-238.

It is important to note that the existence of air contamination and personnel exposures from ²³²Th operations is not addressed in this section. This section deals only with RU contaminants.

5.2 *IN VITRO* MINIMUM DETECTABLE ACTIVITIES, COUNTING METHODS, AND REPORTING PROTOCOLS

5.2.1 In Vitro Urine Analysis

5.2.1.1 Coverage

Uranium enrichment activities began in the fall of 1943. A uranium urinalysis program based on fluorometry was used in conjunction with medical examinations to monitor for kidney damage from exposure to soluble uranium compounds (Sterner and Riley 1946).

It has been determined that employees who were involved in the calutron uranium enrichment process were exposed to levels of airborne uranium products that cannot be determined because of the absence of bioassay data for the period and the lack of air sampling sufficient to develop maximum exposure scenarios.

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Such doses can not be estimated based on source term and process information for lack of documentation on the varying levels of enrichment of the source materials and on the production rate of the operations. Furthermore, the manual recycling and cleaning activities are unique and not comparable to any operations for which NIOSH has access to adequate monitoring data. The occupational environmental internal dose from uranium can be calculated using data provided in the Y-12 National Security Complex – Occupational Environmental Dose (ORAU 2005).

The earliest fluorometry data was dated in 1948. In 1950 workers in production areas were placed on a uranium fluorometry urinalysis program for the purpose of estimating internal exposure. The program was expanded to include certain maintenance workers in 1954 (McLendon 1960). Y-12 currently requires workers with a potential for internal exposures in excess of 100 mrem/yr committed effective dose equivalent (CEDE) to participate in the bioassay program (BWXT Y-12 2003b, p. 23). Until September 1989, routine urinalysis focused on EU, DU, NU, tritium, and plutonium. Beginning in October 1989, uranium results were no longer classified as EU, DU, or NU. Rather, uranium results were reported as isotopic results based on alpha spectrometry analysis. Analyses for other radionuclides were performed on an as-needed basis.

5.2.1.2 Sample Collection

Sample Volumes

For most of the plant's history, the primary urine collection method was a spot sample submitted Monday morning before entering the work area. That is, routine samples were submitted after a minimum of a 48-hr absence from the work area. [In the July 1 to December 31, 1951, Health Physics Progress Report (UCC 1952, p. 25), it was stated that Friday evening samples would be discontinued in favor of Monday morning samples.] The fraction of the daily void volume was estimated on the basis of the time between the sample void and the previous void. This *rate method* of estimating daily void volumes was used explicitly in the calculation of the daily radionuclide excretion.

For example, the following formula was given for calculating EU excretion by electrodeposition and gross alpha counting (UCC 1965, p. 25):

$$dpm/d = 8 \times (AP \div Eff) \times (Vol \div Tl)$$
(3.1)

where:

AP = Average number of counts in 30 min on a plate (disk)

- *Eff* = Uranium recovery efficiency (%)
- *Vol* = Volume of total void (ml)
- TI = Time between sample void and previous void (hr)
- 8 = A constant incorporating time, count, and volume constants, including the 20-ml electroplating aliquot volume. [There is no information in UCC (1965) on what the constant includes or what the daily excretion was.]

The term (*Vol* \div *TI*) also appears in the corresponding equation for fluorometric determinations of NU and DU (UCC 1965, p. 27). The use of the rate method to estimate daily urinary excretion (and hence, radionuclide elimination) contributed to the uncertainty associated with any given measurement; and the corresponding detection level is discussed below.

Since 1989, routine samples have been collected over a 24-hr period, typically while the employee was on a scheduled break from the workplace. Many workers have elected to submit a *simulated* 24-hr sample. This sample is obtained by collecting the last void in the evening (before retiring), any

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urine excreted during the night, and the first void the following morning. This procedure is repeated for two consecutive nights (BWXT Y-12 2003b, p. 70).

Routine Urine Sample Frequency

Urine samples were collected monthly in 1950, and weekly collection for some employees began in 1951. By 1963, health physics personnel were basing the frequency of participation in the urinalysis program for each department on the most recent urinalysis results of that department. The 1963 edition of the *Y-12 Radiation Safety Manual* (UCC 1963a) stated that the frequency of participation schedules was reviewed monthly and adjusted semiannually to meet the following criterion: "Sample at the frequency necessary to assure, with at least 95% confidence, that 95% of the individuals in a department have a quarterly average below the plant action limit." This criterion was used within the limitations of a maximum frequency of once per week and a minimum frequency of once per quarter. Since the late 1980s, most personnel have been on a quarterly frequency.

The current practice is to schedule samples based on the RWP usage rather than a default frequency. Only if an RWP was used will a person be scheduled for a sample. If the RWP specified a fecal sample, a paired set of urine and fecal samples is scheduled approximately 60 d after entry date. If the RWP specified urine only, a urine sample is scheduled approximately 90 d after entry date.

5.2.1.3 Minimum Detectable Activities

Normal and Depleted Uranium in Urine by Fluorometry (1945 to 1989) and Enriched Uranium (1948 to 1950)

In fluorometry, the visible radiation emitted from a doped, sodium-fluoride bead illuminated by ultraviolet radiation is measured. Poisson counting statistics used in nuclear particle counting do not apply to this procedure. The detection limits were historically determined by testing the performance of a particular configuration against standards of known content. Early Health Physics Progress Reports give the minimum detectable limit as 5 ppb, which is a minimum detectable activity of 7 μ g/d for a nominal 1.4 L/d urine excretion rate (UCC 1949, 1950). In 1952, the instrument sensitivity and sample volumes indicated an MDA of 7 μ g/d for 1.4 L/d of urine excreted (CCCC 1953, p. 28).

The activity was calculated from the fluorometric mass reading for NU using a specific activity of 1.55 dpm/µg. For EU (93 wt % 235 U), a specific activity of 150 dpm/µg was used. The corresponding limits of detection are 11 dpm/d for NU and 1,100 dpm/d for EU (93 wt % 235 U). At times, procedures called for the direct conversion of fluorometer current into units of disintegrations per minute, implicitly omitting the mass calculation. Whether the results were expressed in terms of mass or activity, the method remained essentially stable until replaced by long-duration alpha spectroscopy in 1989. Therefore, from 1945 through 1989, the MDA was <u>7 µg/d or 11 dpm/d</u> for NU. From 1948 to 1950, the MDA was <u>7 µg/d or 1,100 dpm/d</u> for EU (93 wt % 235 U). Additional variation resulted from individual urine excretion volumes. Given the limitations of the rate method of estimating daily urine volumes, uncertainty in the excretion volume is likely to contribute significantly to the uncertainty associated with the detection limit of a single measurement.

Enriched Uranium in Urine by Gross Alpha Counting (1950 to September 1989)

Determination of the MDAs for this method is complicated by the standard counting method used. From the mid-1950s to 1989, each sample consisted of two silver disks, each containing uranium electroplated from separate 20-ml raw urine aliquots drawn from the container submitted by the worker. Each disk was counted twice (on two different proportional counters) for 30 min per count. If the two results from a single disk did not agree within tabulated limits, a third count was made (UCC 1966, p. 6) and the two most concordant counts were used. If the average results of the two disks from the same sample did not agree within specified limits, then two more plates were prepared,

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volume permitting (UCC 1966, p. 6). The expression of potentially censored data of this sort in terms of formal detection limits is not straightforward, and no detailed analysis of the statistics of this process has yet been found. However, an approximate indicator of the detection limit can be determined from published values, if issues about statistical independence are set aside.

The background count rate for the proportional counters was reported as 0.12 cpm in 1963 (UCC 1963a, p. 38). Counters were not used if the background was greater than 5 counts in 30 min (Hamrick 1958, p. 7). For a disk counted for 120 min (in fact, two disks were counted for 60 min each), assuming a well-known background and alpha of 0.05, the detection level L_{D} (Currie 1968) is 0.13 cpm. For 1,400 ml/d urine output, a nominal 0.5 cpm/dpm counting efficiency, and an average uranium recovery of 40% as reported by Patterson (1958), the $L_{\rm D}$ would be 46 dpm/d. With an increase in average uranium recovery to 73% (UCC 1965 p. 33), L_D would be reduced to 25 dpm/d. In reviewing historical gross alpha urinalysis data, Barber and Forest (1995) report a decision level of approximately 20 dpm/d. Average uranium recoveries between 1958 and 1965 have not been identified, nor is it clear that the statistical convention used by Barber and Forest was identical to that used here. It is provisionally assumed that L_{D} was 46 dpm/d before 1965 and 25 dpm/d after 1965. However, given the limitations of the rate method of estimating daily urine volumes, uncertainty in the excretion volume is expected to add substantially to the uncertainty associated with the detection limit of a single measurement. Because recoveries were based on batch rather than individual measurements, uncertainties in recovery would also add to the uncertainty of the detection limit of a single measurement.

Uranium in Urine by Alpha Spectrometry (October 1989 to Present)

The present alpha spectrometry system employs large urine aliquots, chemical separation of uranium, sample- specific tracers to determine recovery, and long counting times on low-background detectors. The resultant L_D is approximately 0.15 dpm/d (BWXT Y-12 2002, p. 15). The L_D varies with the uranium recovery of a particular sample, and is reported with the sample result.

Tritium in Urine by Gas Counting

Throughout most of the plant's history (the earliest recorded data was from 1957), tritium was counted by reacting urine with calcium to evolve hydrogen gas, which was counted in an ion chamber. The MDA for this method has not been identified to date. However, the MDA was almost certainly far below the plant action level of 0.25 µCi/ml (8 × 10⁸ dpm/d) (UCC 1957, p. 13). Results are reported in microcuries per milliliter.

Tritium in Urine by Liquid Scintillation

The current system is to place 5 ml of middle distillate from a urine sample in liquid scintillation fluid for counting. Assuming typical method efficiencies, the MDA is 2,000 dpm/d.

Plutonium in Urine by Gross Alpha Counting

The plutonium in urine procedure used before 1988 involved chemical separation from a 24-hr void or a simulated 24-hr sample before gross alpha counting. The detection limit has not yet been identified, but was certainly far below the corresponding limit for uranium in urine, which was based on 20-ml aliquots. If the counting method was identical (same count time of 60 min), chemical recovery (40% to 73%), and counting efficiency (0.5 cpm/dpm) to the early uranium-counting method, the MDA for a 24-hr plutonium sample (less 40 ml) would range from about 0.7 to 1.3 dpm/sample.

Plutonium in Urine by Alpha Spectrometry (1988 to Present) The *a priori* L_D for both ²³⁸Pu and for ^{239,240}Pu is <u>0.025 dpm/sample</u>, or 0.025 dpm/d, assuming a full 24-hr void volume. Actual sample-specific MDAs are included in the analysis reports.

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Other Actinides in Urine and Feces

Table 5-9 lists the analytical laboratory L_D values used since about 1988 (BWXT Y-12 2003b, p. 107).

Table 5-9. L _D values.		
Radionuclide	L _D (dpm/sample)	
²⁴¹ Am	0.05	
²²⁸ Th	0.15	
²³² Th	0.07	
²³⁷ Np	0.1	

Uranium in Feces

Fecal sampling was started in 1998 when the presence of a less soluble component was identified as a result of the stand-down. In 2002, out of 2,800 participants in the bioassay program, 700 Y-12 workers participated in the fecal-sampling program (BWXT Y-12 2002, p. 7). Participation in the fecal-sampling program is based on the potential for significant exposures to insoluble uranium [i.e., workers who are expected to have an internal exposure to type S uranium and there is a potential to exceed 100 mrem (BWXT Y-12 2003b, p. 80)]. At present, the analytical chemistry division cites the value of 0.15 dpm/sample as a representative MDA for fecal analysis (BWXT Y-12 2002, p. 15). Samples usually consist of a single void. The results in the claim files are assumed to be in units of activity per sample unless otherwise indicated (BWXT Y-12 2003c, p. 7).

5.2.2 In Vitro Methods for Individual Radionuclides

The following section discusses the *in vitro* methods for specific radionuclides in urine. (Section 5.2.3 discusses analysis of fecal samples.)

5.2.2.1 In Vitro Bioassay for Uranium

Overview

The *in vitro* bioassay program for uranium at Y-12 can be divided into four eras. First, from 1943 to 1947, limited monitoring occurred, for which the data cannot be retrieved. Second, from 1948 to 1950, fluorometric analyses of urine and blood occurred as part of general medical surveillance to prevent kidney damage from exposure to soluble uranium compounds. Third, from 1950 to 1989, collection of routine samples based on uranium exposure potential was initiated. During this third phase, fluorometric analyses were performed on samples submitted by workers in NU and DU areas. Electrodeposition of uranium, followed by gross alpha counting, was used for samples submitted by workers in EU areas. The primary goal for the EU analyses was to control lung doses from insoluble compounds under the assumption that such materials cleared from the lung with a 120-d half-life. During this lengthy era there were modifications in procedures, but the basic approach remained the same. Fecal sampling was used for some follow-up investigations. Fourth, after October 1989, routine 24-hr urine samples were collected. The uranium was extracted by chemical separation and ion exchange, and the extract was counted by alpha spectroscopy for an extended period (1,000 min/sample). Since 2000, fecal samples have been submitted routinely for individuals working with largely insoluble forms of uranium.

Uranium Analysis by Fluorometry (1948 to 1989)

Uranium processing began in the fall of 1943. Y-12 technical reports describe the use of fluorometry for uranium detection as early as 1944 (Van Wazer and Reiss 1944). By 1946, fluorometric analyses for uranium in urine and blood were in general use to supplement clinical surveillance for soluble uranium exposures (Sterner and Riley 1946). Fluorometric uranium urinalysis data from the

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Tennessee Eastman Corporation era (1943 to 1947) were discussed in a 1981 mortality study of plant personnel (Polednak and Frome 1981).

Early in the Union Carbide management of Y-12, the fluorometric method was reassessed. Extraction methods were used starting in 1949 and 1950 (CCCC 1949, p. 7; UCC 1950, p. 14). By the first half of 1952, a technique involving small (0.2-ml) aliquots of raw urine had been put in place (CCCC 1953, p. 28). However, after this change in technique was instituted the urinary uranium concentrations increased greatly as shown in Table 5-7. This raises the concern that the technique used before May of 1952 could have underestimated the urinary uranium concentrations. (See Section 5.4.2 for discussion of missed doses.)

Fluorometric urinalysis continued to be used for NU and DU until 1989, when alpha spectroscopy began to be used. Fluorometry yielded results in mass units (e.g., μ g/L), but results were often converted to disintegrations per minute per day with the assumed specific activity of 1.55 dpm/µg for NU. The fluorometric technique had an industry standard sensitivity of about 5 ppb (5 µg/L) (e.g., UCC 1949, p. 7).

Uranium Analysis by Electrodeposition and Alpha Counting (1950 to 1989)

The primary objective of this procedure was to prevent EU lung burdens in excess of the prevailing limit (0.017 μ Ci) corresponding to an ICRP 2 (ICRP 1960) calculated lung dose of 15 rem/yr (using a quality factor of 10). Samples from workers in EU areas were analyzed by electrodeposition of uranium onto silver discs, which were then counted for gross alpha activity in a proportional counter. Development of the gross alpha counting method was reported to be underway in early 1949 (CCCC 1949, p. 7). At first, samples were subjected to an acid digestion step before electrodeposition. In 1951, methodological problems leading to underestimates of a factor of 2 were noted and corrected (UCC 1952, p. 53). The information available was not sufficient to determine if the records themselves were corrected. In the mid-1950s, the acid digestion step was discontinued altogether due to reported contamination problems, as well as for logistical considerations (Patterson 1958, p. 34).

From the mid-1950s, two 20-ml aliquots of raw urine from each sample were placed in electrodeposition cells, which produced two discs per sample. Each disc was counted once on one proportional counter for 30 min and a second time on a different counter. The results were then averaged. The background count rate was reported as 0.12 cpm in 1963 (UCC 1963a, p. 38). The expected net count rate for a person excreting at the action level of 70 dpm/d was 0.2 cpm, given a urine output of 1.4 L/d, a nominal 0.5 dpm/cpm counting efficiency, and a uranium recovery of 40%. Because each disk was counted twice for 30 min each, the expected net count for the two counts on each disk under these conditions was 12 counts, over a background of 7.2 counts.

Recoveries were estimated on the basis of spiked samples that accompanied each counting run. The typical spike contained relatively little activity. Recovery rates from raw urine tended to be around 40% in the late 1950s (Patterson 1958), but they improved over time: In 1965, the average recovery for electroplating was 73% (UCC 1965, p. 33). Method limitations are discussed in a number of reports, including Johnson et al. (1959) and UCNC (1959). Recovery rates were found to vary with pH, drying procedure, and uranium content, with rates decreasing for uranium concentrations below the plant action level. The precision of any one sample was acknowledged to be relatively low:

Further, realizing the over-all lack of precision in any one sample result, we normally recommend restriction only on the basis of the 13-week or quarterly cumulative internal dose as indicated by from 2 to 13 samples. (Patterson 1958, p. 57)

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The precision of an individual result in this type of procedure is poor and for this reason little significance is attached to individual results, particularly those below the "Plant Action Limit (PAL)" level. (UCC 1966).

These limitations notwithstanding, the salient point is that, when *in vivo* counting became routine in the early 1960s, very few additional workers were found to require restriction from uranium areas (e.g., UCC 1963b, p. 6). The *in vivo* monitoring frequency was determined by urinalysis results and ranged from monthly to once each 18 months (UCC 1963b, p. 1). Almost all of the restrictions that did take place would have occurred on the basis of the urinalysis program alone. In those cases in which restriction was based on *in vivo* analysis alone, lung retention times were often observed to be considerably longer than the assumed 120-d lung half-life on which the urinalysis program was based. In other words, the problem was largely due to the inapplicability of the lung model for some materials and individuals, rather than a failure of the urinalysis program.

Uranium-233 by Gross Alpha Counting

In early 1962, Y-12 undertook the fabrication of metallic ²³³U from a nitrate solution (²³³U uranyl nitrate) in Building 9205 (West and Roberts 1962). In recognition of the high specific activity of this material, and the potential for gamma radiation from the progeny of contaminant ²³²U, operations were doubly contained. Pilot runs were made with less hazardous materials to test the protective measures. As part of these precautions, eight workers submitted urine samples for analysis by electrodeposition and gross alpha counting as described above, with the same recoveries and detection efficiencies. Ratios of ²³³U to ²³²U varied with operation and time. Table 5-10 list ratios for various exposure scenarios. In addition to absorption types M and S, exposure to type F material is considered likely in this process. Uranium-232 has the larger dose conversion factor.

Table 5-10. 200 to 2020 ratios.	
Process	U-233:U-232 activity ratio
Reduction of UF ₄	1.3
Casting of metal-crucible skull	15–32
Machining final uranium component	160

Fable 5-10. ²³³U to ²³²U ratios.

Due to uncertainty in the process, claimant-favorable assumptions should be made about solubility and uranium activity ratios.

Uranium Analysis by Alpha Spectrometry (1989 to Present)

Up to 2 L of urine, along with a 232 U tracer, are treated with nitric acid (HNO₃) and hydrogen peroxide (H₂O₂). The sample is wet-ashed, and the uranium is coprecipitated with calcium oxalate (CaC₂O₂). After dissolving the precipitate in hydrochloric acid (HCl), the uranium is further separated by ion exchange chromatography. The uranium is eluted from the column with a solution of dilute HCl to which titanous chloride (TiCl₃) has been added to reduce actinides that could be in an elevated oxidation state. The final fraction of the eluate is first treated with ascorbic acid to reduce the presence of any ferric iron and next with hydrofluoric acid (HF). Then the uranium isotopes are coprecipitated on neodymium fluoride (NdF₃). The NdF₃ is caught on a 0.1-mm filter, rinsed, dried, and mounted on a planchet for alpha spectrometry (BWXT Y-12 2003b). Typical recoveries are about 85%. Samples are typically counted for 16 hr on a passivated implanted planar silicon detector.

5.2.2.2 In Vitro Bioassay for Plutonium

Plutonium in Urine by Gross Alpha Counting

In 1957, certain workers in the Special Testing Department were tested for plutonium intake by urinalysis. Because large urine volumes were required, these employees submitted 24-hr samples through the use of take-home kits. The sampling frequency was monthly (UCC 1957, p. 38). After two 20-ml aliquots were removed for uranium analysis by electrodeposition, the balance of the sample was treated by chemical separation, precipitation, and evaporation onto a stainless-steel planchet for alpha counting. Health physics personnel converted the reported activity rate (disintegrations per minute per 24-hr void) to millirem per day to the critical organ.

Uranium, Plutonium, and Americium by Alpha Spectrometry (October 1989 to Present)

Up to 2 L of urine, along with ²³²U, ²⁴²Pu, and ²⁴³Am tracers, are treated with HNO₃ and H₂O₂. The sample is wet ashed, and the uranium, plutonium, and americium are coprecipitated with CaC₂O₂. After dissolving the precipitate in HCI, the uranium, plutonium, and americium are further separated by the use of two ion exchange columns and an additional oxalate precipitation. The uranium, plutonium, and americium in the final fractions are coprecipitated with NdF₃. The NdF₃ is caught on a 0.1-mm filter, rinsed, dried, and mounted on a planchet for alpha spectrometry (BWXT Y-12 2003b). The same procedure is used for isotopic thorium.

5.2.2.3 In Vitro Bioassay for Tritium

As of 1957, personnel engaged in processing materials with a potential for tritium contamination submitted three urine samples per month. Samples were submitted at the same stations used for uranium and plutonium samples. The samples were reacted to evolve hydrogen gas, which was then counted for beta activity in an ionization chamber. Results were reported in microcuries per milliliter. (UCC 1957, pp. 38–39)

Tritium is currently measured by liquid scintillation counting. An aliquot of the urine sample is distilled in a heating mantle, and the middle fraction of the distillate is collected. A 5-ml portion of the collected distillate is mixed with liquid scintillation cocktail. The beta activity of the tritium is then measured by liquid scintillation counting (BWXT Y-12 2003b). Exposure should be assumed to be due to HTO.

5.2.2.4 In Vitro Bioassay for Other Radionuclides

The following alpha spectrometry procedures described in Section 3.2 of BWXT Y-12 (2003b) have been in use since 1989.

Neptunium-237 in Urine by Alpha Spectrometry

In this procedure, ²³⁹Np is used as an internal yield monitor. Americium-243, which is in equilibrium with its daughter, ²³⁹Np, is added during the sample preparation process. The separation time of ²³⁹Np from its parent, ²⁴³Am, is recorded (so that the count can be corrected for decay), and the beta activity is measured to calculate the radiochemical yield. The tracer is added to the urine sample (or fecal material) along with concentrated HNO₃ and H₂O₂, and the sample is wet ashed. Neptunium is coprecipitated with CaC₂O₂ and mixed hydroxides. After dissolving the precipitate in concentrated HNO₃ and ashing in a muffle furnace, the oxidation state of neptunium is separated from other actinides by the anion exchange method. Neptunium is eluted with 0.5-molar HCl solution containing TiCl₃. The eluate is collected, and neptunium is coprecipitated on NdF₃ and filtered on 0.1-mm filter paper. The NdF₃ is rinsed, dried, and mounted on a planchet for beta counting and alpha spectrometry.

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Isotopic Thorium (²²⁸**Th,** ²³⁰**Th, and** ²³²**Th) by Alpha Spectrometry** Up to 2 L of urine along with a ²²⁹Th tracer is treated with HNO₃ and H₂O₂. The sample is wet ashed, and the thorium is coprecipitated with CaC_2O_2 . After dissolving the precipitate in HNO₃ and mixing with methanol, the thorium is further separated by ion exchange chromatography. The thorium is eluted from the column with 8-molar HCI. Hydrofluoric acid (HF) is added, and the thorium isotopes are coprecipitated with NdF₃. The NdF₃ is caught on a 0.1-mm filter, rinsed, dried, and mounted on a planchet for alpha spectrometry. The same procedure is used for uranium, plutonium, and americium determinations. This method accommodates substantial quantities of common interferences, such as iron, aluminum, titanium, zirconium, and uranium. However, plutonium can present some interference. If plutonium is expected, an alternate method should be chosen (BWXT Y-12 2003b).

5.2.3 **Fecal Sample Analysis**

Fecal samples have long been used in follow-up investigations when urinalysis or in vivo measurements indicate the likelihood of a substantial intake. West and Scott (1966) describe such an investigation in the 1960s in which fecal, urine, and in vivo measurements were used jointly to investigate clearance. Fecal sampling was discontinued several times in the past, but was reinstated in 1998 due to changes in workplace exposure conditions. At first, fecal sampling was used for a limited number of workers in Building 9212 but was expanded to include Building 9215 in 1999 and to other areas of the plant in 2000 and 2001 (BWXT Y-12 2002, p. 6).

If a work area has been determined to have predominantly insoluble airborne uranium, it is current practice for workers with a moderate to high exposure potential (CEDE greater than 100 mrem) to submit both urine and fecal samples. The relative elimination by urinary and fecal pathways is used to determine the solubility mixtures for each individual. For combined urine and fecal monitoring, the routine sampling interval is 53 d (BWXT Y-12 2002, p. 14) with 24-hr fecal collections preferred. When 24-hr fecal samples are not available, the results can be normalized using the ratio of the Reference Man excretion rate of 135 g/d to the mass of the submitted sample. The Y-12 practice was to routinely normalize results to Reference Man. These normalized results or actual individual daily excretion results can be used to reconstruct dose.

5.2.4 Interferences

As investigation levels decreased over time, the contribution of natural background to worker uranium excretion assumed greater importance. Since 1989, Y-12 has corrected the measured uranium excretion values for background uranium interference when performing dose assessments. However, the uranium results reported in response to NIOSH requests for dose records have not been corrected for dietary uranium.

Y-12 used the following methods to correct sample results reported after 1989 for dietary levels of uranium. The range of background uranium values expected to be observed in urine samples is 0.14 to 0.57 dpm and the range of background uranium values expected to be observed in the fecal samples is 2.32 to 2.75 dpm (BWXT Y-12 2003b, p. 75). Therefore, at Y-12, the reference point is established at the midpoint of the quoted ranges of values for urine (0.35 dpm/d) and fecal samples (2.5 dpm/sample). The ratios between ²³⁴U and ²³⁸U can also provide information to help characterize the sample. Natural background uranium typically has a ²³⁴U to ²³⁸U ratio of approximately 1:1. EU also has a characteristic ratio, typically greater than 3:1. DU, on the other hand, typically has a characteristic ratio of less than 0.9:1.

Based on the above information, the background uranium corrections used at Y-12 are as follows:

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For bioassay sample results less than the detection limit for the analytical technique, no action is required. Note, that corrections for samples from the early days when MDAs were significantly larger are not required.

To adjust results for background, the ²³⁴U-to-²³⁸U ratios are reviewed for each sample to determine if the sample is consistent with EU or DU. Conservative ratios are applied in this determination: All samples with ²³⁴U-to-²³⁸U ratios greater than 1:1 are considered to be consistent with EU, and all samples with ²³⁴U-to-²³⁸U ratios less than 1:1 are considered to be consistent with DU.

The background contribution is subtracted from EU samples as follows:

- a. Assume that the ratio of ²³⁴U to ²³⁸U is 1:1 for dietary uranium.
- b. Determine if the ²³⁸U result exceeds its corresponding critical level value.
- c. If the ²³⁸U result is greater than the critical level, then subtract two times that ²³⁸U result from the total uranium result.
- d. If the ²³⁸U result is less than its corresponding critical level value, do not subtract a dietary component.

For DU samples (e.g., those with a ²³⁴U-to-²³⁸U ratio less than 1:1), the following steps are used to determine the dietary correction for DU samples:

a. The following equations are solved to determine an equation for the occupational ²³⁸U content and the occupational ²³⁴U content in a DU sample:

$$\frac{^{234}U_{diet}}{^{238}U_{diet}} = 1$$
(3-2)

$$\frac{^{234}U_{occ}}{^{238}U_{occ}} = 0.2$$
(3-3)

$${}^{238}U_{actual} = {}^{238}U_{diet} + {}^{238}U_{occ}$$
(3-4)

$$^{234}U_{actual} = ^{234}U_{diet} + ^{234}U_{occ}$$
(3-5)

b. Using Equation 3-5, and substituting the ${}^{238}U_{diet}$ for the ${}^{234}U_{diet}$ based on Equation 3-2 and substituting 0.2 (${}^{238}U_{occ}$) based on Equation 3-3 for the ${}^{234}U_{occ}$, the following relationship results:

$$^{234}U_{act} = ^{238}U_{diet} + 0.2(^{238}U_{occ})$$
(3-6)

Because the ${}^{238}U_{diet}$ can be replaced with ${}^{238}U_{actual} - {}^{238}U_{occ}$ derived from Equation 3-4, Equation 3-5 can be solved to determine the occupational quantity of 238 U. The resultant equation for determining the occupational quantity of 238 U is:

$$\frac{{}^{234}U_{act} - {}^{238}U_{act}}{-0.8} = {}^{238}U_{occ}$$
(3-7)

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c. Once the occupational ²³⁸U quantity is known, the occupational quantity of ²³⁴U can be derived using Equation 3-3:

$$^{234}U_{occ} = 0.2(^{238}U_{occ}) \tag{3-8}$$

- d. Ensure that both the ²³⁴U and ²³⁸U results exceed their corresponding critical level values. If both of these results are not greater than their corresponding critical level values, then a background determination cannot be made. In this case, it is assumed that all of the activity observed in the total uranium sample result will contribute to occupational dose.
- e. Use either Equation 3-7 or 3-8 to determine the occupational ²³⁸U component or the occupational ²³⁴U component, respectively. For ease of reference, the ²³⁸U component was chosen for the remaining steps.
- f. Subtract the occupational component of the sample from the corresponding observed component to determine the dietary contributor:

$${}^{238}U_{act} - {}^{238}U_{occ} = {}^{238}U_{diet}$$
(3-9)

g. Once the dietary component has been calculated, subtract twice this dietary value from the total uranium result to remove the background:

$$U_{tot} - 2\binom{238}{U_{diet}} = U_{totalocc}$$
(3-10)

5.3 *IN VIVO* MINIMUM DETECTABLE ACTIVITIES, COUNTING METHODS, AND REPORTING PRACTICES

5.3.1 Whole-Body Counting

Whole-body counting was not routinely practiced at Y-12. The primary *in vivo* detection method was chest counting, as described below.

5.3.2 Chest Counting

The Y-12 *in vivo* chest-counting facility was developed in the late 1950s and was put into routine use in 1961. The original facility is described in Cofield (1959) and Cofield (1960). A 9- by 4-in. sodium iodide (thallium doped) [Nal(TI)] crystal was placed over the chest of a subject reclining in a cot in a shielded room. Subjects showered, shampooed, and changed into a clean garment before counting. They were then surveyed for any remaining surface contamination before entering the counting room. The normal counting time was 20 min.

Incremental improvements in the system were made over time. A second 9- by 4-in. Nal(TI) crystal was placed under the subject's back in 1963 or 1964. It was found that this position provided greater counting efficiency than the position over the chest. A second advantage was that skin contamination seldom occurred on the back. The ratio of the count rates from the two crystals could be used to identify surface contamination missed by the survey meter. In 1965, two 5-in. Nal(TI) crystals were added, positioned at the sides under the arms. In 1985, initial development of a high-purity germanium (HPGe) counting system was begun. The new system began routine operation in a new counting room in June 1992.

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5.3.2.1 Chest Counting for Uranium

With the Nal scintillation system used for much of the plant's history, the amount of uranium deposited in the lungs was inferred from the worker's spectrum by use of a prediction equation. This included the subject's weight and constants derived from the spectra of control subjects. Compton scatter from natural ⁴⁰K and fallout ¹³⁷Cs in the body was a particularly important contributor to background in the ²³⁵U region of interest around 186 keV and the ²³⁸U region of interest around 93 keV. The ¹³⁷Cs contribution to the regions of interest changed over time as fallout levels varied, requiring frequent updates to the control subject spectrum library.

Measured lung burdens were originally expressed in micrograms of ²³⁵U for EU and milligrams of ²³⁸U for NU or DU. The original limits of detection for ²³⁵U and ²³⁸U were reported as 130 µg and 13.5 mg, respectively (Cofield 1959). Improvements in hardware and data reduction procedures led to lower detection limits, as indicated in Tables 5-11 and 5-12 below. Limits are given in terms of both mass units and of activity (nanocurie and dpm), with the published value in regular type and the converted value in italics. When known, the reporting convention used to define the detection limit is indicated in the tables. The alternative definitions listed by King and Barclay (1983) illustrate the importance of this factor.

Due to the uncertainty about actual methods for determining the lung counting detection limits, for dose reconstruction purposes and based on review of the Y-12 data, the ²³⁵U and ²³⁸U lung count detection thresholds are assumed to be 130 μ g and 13.5 mg, respectively, through 1990. Beginning in 1991, the critical level or MDA is supplied with the individual results.

Year	Detectors	U-235 (µg)	U-235 (nCi)	U-235 (dpm)	Reporting convention	Reference
1959	One 9-in. Nal	130	0.28	620		Cofield 1959
1963	Two 9-in. Nal	96	0.21	470		Scott and West 1967
1965	Two 9-in., 2 5-in. Nal	72	0.16	360		Scott and West 1967
1975	Two 9-in., 2 5-in. Nal	70	0.15	330		Scott and West 1975
1983	Two 9-in., 2 5-in. Nal	62	0.13	290	1.96 σ _{вкс}	King and Barclay 1983
1983	Two 9-in., 2 5-in. Nal	68.5	0.15	330	L _C ^a	King and Barclay 1983
1983	Two 9-in., 2 5-in. Nal	137	0.30	670	L _D ^a	King and Barclay 1983
1990	Two 9-in., 2 5-in. Nal	70	0.15	330	b	Barber and Forest 1995
6/1992	HPGe	46	0.10	220	с	BWXT Y-12 2003b

Table 5-11. Reported lung counting detection limits for ²³⁵U.

a. As defined by Currie (1968) with $\alpha = \beta = 0.05$, paired observations.

b. Type I error = 5%.

c. Individual MDA or critical level supplied with analytical results.

Table 5-12. Reported lung counting detection limits for ²³⁸U.

Year	Detectors	U-238 (mg)	U-239 (nCi)	U-238 (dpm)	Reporting convention	Reference
1959	One 9-in. Nal	13.5	4.5	10,000		Cofield 1959
1965	Two 9-in., 2 5-in. Nal	6.5	2.2	4,900		Scott and West 1967
1983	Two 9-in., 2 5-in. Nal	5	1.7	3,800	1.96 σ _{BKG}	King and Barclay 1983
1983	Two 9-in., 2 5-in. Nal	7.2	2.4	5,300	L _C ^a	King and Barclay 1983
1983	Two 9-in., 2 5-in. Nal	14.4	4.8	11,000	L _D ^a	King and Barclay 1983
1991	HPGe	4.5	1.5	3,300	b	BWXT Y-12 2003b

a. As defined by Currie (1968) with $\alpha = \beta = 0.05$, paired observations.

b. Individual MDA or critical level supplied with analytical results.

No description of the conversion count rate to activity to mass has been found. The analyte reported was based on the area in which the employee worked. Individuals working in NU or DU areas had results reported as ²³⁸U, and workers in enriched areas had results reported as ²³⁵U.

Claimant-favorable assumptions should be based on conversions of 93% enrichment for ²³⁵U and natural isotopic abundances for ²³⁸U.

5.3.2.2 Chest Counting for Other Actinides

Thorium

The *in vivo* lung count was the only monitoring technique for monitoring thorium exposure in the body during the plant's first decades. Thorium lung activity was inferred from ²²⁸Ac and/or ²¹²Pb lung activity. Thorium lung counting was conducted from 1958 to 1984 with routine lung counts, scheduled at approximately six-month intervals, starting in 1961.

The interpretation of thorium spectra is complicated by the tendency of decay chain members to become separated during operations, particularly those involving heat. The boiling point of radium is below the melting point of thorium. The ²²⁸Ac gamma radiation in an *in vivo* count could be an indication of a ²²⁸Ra intake or an intake of the entire ²³²Th chain. The 239-keV gamma line from ²¹²Pb, a progeny of ²²⁸Th, is a more reliable indicator of ²³²Th, as the thorium isotopes would remain together during processing. However, if the ²²⁸Ra is lost, ²¹²Pb decreases with the 1.9-yr half-life of its parent, ²²⁸Th, moderated by the ingrowth of new ²²⁸Ra with its 5.7 yr half-life. This complicated disequilibrium

pattern operates both before and after chain members are taken into the body. Some knowledge of the likely state of equilibrium is therefore necessary to translate observed activities into dose.

The maximum permissible lung burden (MPLB) of ²³²Th (corresponding to 15 rem/yr to the lung) varied markedly with the ratio of ²²⁸Th to ²³²Th. For full equilibrium (ratio = 1) the MPLB was calculated to be 2.9 nCi of ²³²Th. For a ratio of 0.1, the lung burden was 12 nCi (West 1965, p. 22). For a ratio of 0.8, an MPLB of 3.2 nCi is indicated. For this ratio, with a ²²⁸Ra-to-²³²Th ratio of 0.6, an MDA of 0.2 lung burdens, <u>or 0.6 nCi</u>, is indicated (West 1965, p. 26). In mass units, this is <u>5.5 mg</u>.

It was reported in 1965 that thorium at Y-12 was processed less than 1 yr after purification by the supplier and as a consequence had only about 10% as much ²²⁸Ra as ²²⁴Ra (West 1965, p. 18). This means that the maximum dose conversion factor per milligram of ²³²Th would be less than that for ²³²Th in full equilibrium with its progeny. For calculating intakes of Th-228 and Th-232 from lung counts, a default assumption must be made for the ratio of equilibrium of Th-228 to Th-232. This ratio will be used to determine the intake of Th-228 from the intake of Th-232 calculated from the lung count results (assumed to be Th-232). As can be seen in Figure 5-4, the activity of Th-228 after initial purification declines over the first five years due to the decay of the original Th-228 in the mixture, then slowly builds back up again due to the Th-228 resulting from the decay of Th-232.

Since the actual time from initial purification is unknown and will change over time, an assumption of an average of 80% equilibrium of Th-228 to Th-232 should be assumed. This is a reasonable assumption since the actual equilibrium percentage ranges from 100% at initial purification to below 60% and then slowly rises again. The drop to 80% occurs within the first six months and continues dropping, over the first year, so the assumption of 80% is consistent with the assumption the thorium was processed within a year of purification. This translates to a ratio of Th-228/Th-232 of 0.8.

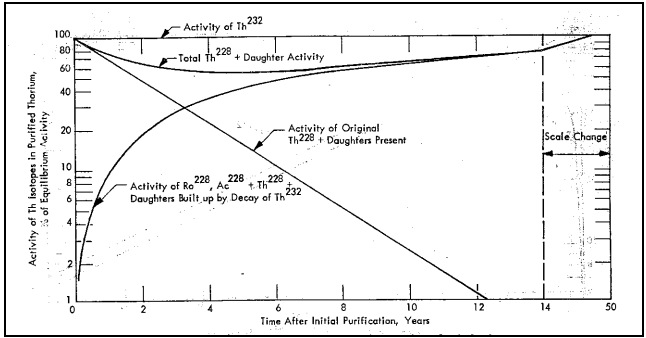


Figure 5-4. Variation of activity in natural thorium after a single purification.

Neptunium

At the time the *in vivo* system was put into routine service in 1961, the reported detection limit for ²³⁷Np without progeny radiation was 2.7 nCi (Cofield 1961). For ²³⁷Np in full equilibrium with ²³³Pa, the detection limit was reported as 0.255 nCi (Cofield 1961).

5.3.2.3 Chest Counting for Other Radionuclides

At the time the *in vivo* system was put into routine service in 1961, the reported detection limit for ${}^{60}Co$ was 0.66 nCi. For ${}^{95}Zr$ in transient equilibrium with ${}^{95}Nb$, the reported detection limit was <u>1 nCi.</u> Although bremsstrahlung counting could have been done for ${}^{99}Tc$, no information is available about the sensitivity of the technique.

5.4 UNMONITORED DOSE (NOT COMPLETE FOR ALL OPERATIONS AND/OR RADIONUCLIDES)

There are two eras of operations at Y-12 for which sufficient monitoring information might not be available in the worker records. From 1943 to 1947, limited or no monitoring was performed. No data have been found for this period. The primary site activities for this period were uranium enrichment by the Calutron process. From 1948 to 1950, fluorometric analyses of urine and blood were conducted as part of general medical surveillance to prevent kidney damage from exposure to soluble uranium compounds; these data cannot be found at this time. Coworker data will be applied to this time period.

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GLOSSARY

activity median aerodynamic diameter (AMAD)

The diameter of a sphere having a density of 1 gram per cubic centimeter and the same terminal settling velocity in air as that of the aerosol particle whose activity is the median for the entire aerosol.

acute exposure

Exposure of short duration.

background (radiation)

Radiation from:

- Naturally occurring radioactive materials that have not been technologically enhanced
- Cosmic sources
- Global fallout as it exists in the environment (from the testing of nuclear explosive devices)
- Radon and its progeny in concentrations or levels existing in buildings or the environment that have not been elevated as a result of current or previous activities
- Consumer products that contain nominal amounts of radioactive material or produce nominal amounts of radiation.

bioassay

The determination of kinds, quantities, or concentrations and, in some cases, locations of radioactive material in the human body, whether by direct measurement or by analysis and evaluation of radioactive materials excreted or removed from the human body. Also called *radiobioassay*.

chronic exposure

Exposure of long duration, delivered by fractionation or protraction.

class D, W, or Y material

ICRP 30 classifies inhaled radioactive materials as type D, W, or Y (days, weeks, or years) depending on their retention times in the pulmonary region. Class D materials have a pulmonary half-life of less than 10 d; W materials have a half-life from 10 to 100 d; and Y materials have a half-life greater than 100 d. The times actually used for these classes of materials in ICRP 30 in the calculation of annual limits on intake are 0.5 d, 50 d and 500 d for class D, W, and Y materials, respectively. Note: A class Y uranium has been defined as having an effective pulmonary half-life of 100 d and the systemic distribution, retention, and excretion parameters of class W uranium.

class (type) Q system

A class Q dosimetry system uses an $8-\mu$ m AMAD as a basis. The class Q dosimetry system is described as 10% class Y and 90% modified class W (the modification being an increase of the class W 50-d compartment to 120 d).

committed effective dose equivalent (CEDE)

The sum of the committed dose equivalents to various tissues or organs in the body for 50 yr after intake. CEDE is expressed in units of rem (or sievert).

depleted uranium (DU)

A byproduct of the enrichment process, depleted uranium has a reduced abundance of ²³⁵U relative to the 0.72% abundance in natural uranium.

dose

A general term for absorbed dose, dose equivalent, effective dose equivalent, committed dose equivalent, committed effective dose equivalent, or total effective dose equivalent.

enriched uranium (EU)

Uranium with an increased abundance of ²³⁵U. Enriched uranium varies from greater than the natural ²³⁵U abundance of 0.72% up to almost 100%. Enriched uranium can also contain increased abundances of ²³⁴U, ²³⁶U, ²³²U, ²³³U and others.

exposure

(1) The general condition of being subjected to radiation, such as by exposure to radiation from external sources or to radiation sources inside the body. (2) In this document, exposure does not refer to the radiological physics concept of charge liberated per unit mass of air. The product of exposure time to a radioactive aerosol and the average concentration during exposure, divided by the value of the derived air concentration for the radioactive material in question (expressed in derived air concentration-hours).

intake

The amount of radionuclide (mass or activity) taken into the body by inhalation, absorption through the skin, injection, ingestion, or through wounds.

in vitro measurement

Measurements to determine the presence of or to estimate the amount of radioactive material in the excreta or in other biological materials removed from the body.

in vivo measurement

The measurements of radioactive material in the human body using instrumentation that detects radiation emitted from the radioactive material in the body.

maximum permissible lung burden (MPLB)

During the 1970s the occupational limit for plutonium was expressed in terms of a quantity of plutonium that could be present in the chest at any given time, equal to $16 \ \mu$ Ci (0.25 μ g)²³⁹Pu.

minimum detectable amount or activity (MDA)

The smallest amount (mass) or activity (per mass or per sample) of an analyte in a sample that will be detected with a probability ß of nondetection (type II error) while accepting a probability of erroneously deciding that a positive (nonzero) quantity of analyte is present in an appropriate blank sample (type I error).

minimum detectable concentration (MDC)

The minimum detectable amount expressed in units of concentration.

monitoring

The measurement of radiation levels, airborne radioactivity concentrations, radioactive contamination levels, quantities of radioactive material, or individual doses.

natural uranium

The mixture of uranium as it is found in nature (i.e., 99.27% 238 U, 0.72% 235 U, and 0.0054% 234 U by weight). The alpha activity comes mostly, and in about equal amounts, from 238 U and 234 U. The specific activity of this mixture is 2.6 x 10⁷ Bq/kg (0.7 pCi/g).

nephrotoxicity

Poisoning of the kidney.

quality factor

A modifying factor used to calculate the dose equivalent from the absorbed dose.

recycled uranium (RU)

Uranium recovered from spent fuel from fission reactors.

routine bioassay monitoring

Any bioassay measurement made on a predetermined, periodic schedule, to establish a worker's internal exposure status relative to previous periods of time.

sensitivity (detection sensitivity)

The minimum amount of contaminant (or dose) than can be repeatedly measured (or calculated) by a particular analysis. See *minimum detectable amount*.

type F, M, or S material

ICRP 66 classifies inhaled radioactive materials as F, M, or S (fast, medium, or slow) depending on their retention time in the pulmonary region. These designations are similar to the clearance classes D, W, and Y, but refer strictly to the rate of absorption from the lungs to the blood.

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Γ

ATTACHMENT 5A OCCUPATIONAL INTERNAL DOSE FOR MONITORED WORKERS

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5A.1 CONVENTIONS USED IN INTERNAL DOSE REPORTS

The following is adapted from Souleyrette (2003).

5A.1.1 Y-12 Urinalysis Data through September 1989

<u>Dept</u> – Department number of employee at the time of monitoring.

Badge/Old SSN – Employee identification number.

Void Date – Date sample was voided.

Program Code

- 1 = EU urinalysis by gross alpha
- 2 = NU or DU by fluorophotometry (used for EU before 1950)
- 6 = control sample
- 9 = (may have been used for a limited time to indicate experimental analyses methods).

Volume – volume of sample in ml.

<u>Time</u> – Time between last void and sample void. Used to normalize urine result to 24-hr void based on individual employee daily excretion rate. (A zero time in conjunction with a large void volume and program code 9 probably indicates an experimental measurement technique.)

Uranium Result (dpm) – The normalized urine result in disintegrations per minute per day.

<u>Background Value</u> – Background was stored for later results. The column Uranium Result (dpm) is background corrected already.

5A.1.2 Y-12 Urinalysis Data since October 1989

<u>Dept</u> - Department number of employee at the time of monitoring.

Badge - Employee identification number.

Sample Number – Laboratory sample identifier.

Sample Date - Date sample was voided.

Sample Type

 $\begin{array}{l} \mathsf{A} = \mathsf{Alpha \ spec} \\ \mathsf{F} = \mathsf{fecal} \\ \mathsf{U} = \mathsf{urine} \\ \mathsf{S} = \mathsf{Start} \ \mathsf{or} \ \mathsf{stop} \ \mathsf{date} \end{array}$

(Note: For urinalysis, A is expected because all urinalyses were done this way since October 1989. The other entries are for database flexibility.)

Reason – Reason for scheduling a sample:

- 1S = acute exposure
- 2S = follow-up sample to an acute exposure
- 3 = routine (chronic) exposure
- 2R = follow-up to routine (chronic) exposure
- 4 = background sample
- 5 = baseline sample
- 6 = random sample
- 8 = start date of exposure *
- 9 = stop date of exposure *
- 0 = acute exposure
- * (These should appear only in separate records introducing a series related to a chronic intake study).

<u>Retention Class</u> – Assigned retention class: ICRP-30 classes D, W, and Y. (Class Q indicates 90% class Super–W (120-d lung retention) and 10% class Y, 8 µm AMAD.)

<u>Volume (ml)</u> or <u>Weight (g)</u> – sample volume (urine) or weight (fecal).

<u>Total Uranium Result</u> (dpm) - This is the normalized total uranium urine result in disintegrations per minute per day.

Type Assimilation

H = inhalation

- I = ingestion
- J = injection

Incident Date – Date of documented radiological exposure incident, if any.

Incident Time – Time of documented radiological exposure incident, if any.

5A.1.3 In Vivo Data

<u>Dept</u> – Department number of employee at the time of monitoring.

Run Date – Date subject counted.

Run No. – Laboratory identifier of counting session.

Weight (lbs.) – Subject weight in pounds.

<u>Chest Thickness</u> (in.) – Subject chest wall thickness in inches (could be in tenths of inches for some years).

<u>Surface Contamination Code</u> – A six-digit code indicating whether or not surface contamination was detected.

• If the first position is a 1, there was no surface contamination found, and the remaining five digits should be zeros.

• If the first digit is a 2 or a 3, there was surface contamination found and the remaining five digits give the survey meter reading of the contamination.

<u>Type Analysis</u> – Code indicating the activity regions analyzed. There are two distinct sets of codes depending on the date of the count.

From October 1962 to 1971, the codes are:

- 0 = Background run
- 1 = Uranium
- 2 = New hires/control counts
- 3 = Thorium
- 4 = Neptunium
- 5 = Miscellaneous
- 6 = Second thorium count.

Since 1972, the codes are:

 $1 = {}^{235}U$ 3 = Thorium $4 = {}^{238}U \text{ and } {}^{235}U$ $6 = Thorium \text{ and } {}^{235}U \text{ (first thorium count)}$ $7 = {}^{235}U \text{ and Thorium (second thorium count)}.$

Material Type – This code provides additional details for cases where Type Analysis = 0, 1, 2, or 3.

For counts after 1971, where Type Analysis is 0:

00 = Background run 01 = Sensitivity Check.

For Type Analysis of 1 (any year) or Type Analysis of 2 (after 1971):

01 = Unknown 02 = DU 03 = DU 07 = Normal uranium 13 = EU 19 = EU.

For Type Analysis of 3 (any year):

- 00 = Thorium (first exam)
- 02 = Thorium (second or later exam).

<u>U-235</u> (μ g) – ²³⁵U activity detected in chest cavity. If the count is invalid, an '&' appears as the first character.

<u>Thorium</u> (mg) $-^{232}$ Th activity calculated from the daughter isotopes detected in chest cavity.

<u>U-238</u> (mg) $-^{234}$ Th activity detected in chest cavity.

<u>Tc-99</u> (μ Ci) - ⁹⁹Tc activity detected in chest cavity.

<u>Current Dept</u> – Additional instance of employee department number used historically for reporting purposes.

5A.2 URANIUM SOLUBILITY IN THE LUNG

For a workplace as varied as Y-12, it is clear that no single solubility type or particle size distribution applies to all workers. Furthermore, accurate assignment of the uranium lung clearance type to a given bioassay result is considered virtually impossible because of uncertainties about chemical form and limitations of the personnel-tracking system (Barber and Forest 1995, p. 669). Exposure to type M material from 1948 to June 1998 appears to be the most likely absorption type. After June 1998, exposure to absorption type S material is more likely. However, the absorption type can be based on the monitoring data or claimant-favorable assumptions.

5A.3 IN VITRO DETECTION LIMITS

Tables 5A-1 and 5A-2 summarize information developed in Section 5.2. The tabulated values for urinalysis results represent laboratory detection limits L_D and do not include uncertainties introduced by sample collection or conversion from submitted volumes to daily void volumes. As noted in Section 5.2, L_D values for some historical techniques remain to be identified, and will be reported in subsequent revisions as available.

Method	Period	Detection limit (mass)	Detection limit (activity)	Convention
Fluorometry	1950–9/1989	7 µg/d	11 dpm/d	
Gross alpha	1950–1964		47 dpm/d	L _D
Gross alpha	1965–9/1989		26 dpm/d	L _D
Alpha spectrometry	10/1989-present		0.15 dpm/d	L _D

Table 5A-1. Uranium urinalysis detection limits.

Table 5A-2.	Other	in vitro	detection	limits.
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		Detection limit
Method	Period	(activity)
Liquid scintillation	10/1988-present	2,000 dpm/d ^a
Alpha spectrometry	10/1989-present	0.025 dpm/sample
Alpha spectrometry	10/1989-present	0.050 dpm/sample
Alpha spectrometry	10/1989-present	0.150 dpm/sample
Alpha spectrometry	10/1989-present	0.070 dpm/sample
Alpha spectrometry	10/1989-present	0.100 dpm/sample
	Liquid scintillation Alpha spectrometry Alpha spectrometry Alpha spectrometry Alpha spectrometry	Liquid scintillation10/1988-presentAlpha spectrometry10/1989-presentAlpha spectrometry10/1989-presentAlpha spectrometry10/1989-presentAlpha spectrometry10/1989-presentAlpha spectrometry10/1989-present

a. Estimate.

5A.4 IN VIVO DETECTION LIMITS

From the review of Y-12 documents and conversations with present and former Y-12 staff, the vast majority of site operations from 1943 to the present involved uranium in a variety of chemical forms and degree of enrichment. Therefore, it is concluded that the primary internal radiation exposure to Y-12 workers was from uranium. However, the internal dosimetry program has included limited monitoring for ¹³⁷Cs, ⁹⁹Tc, thorium, plutonium, ²²⁸Ac, and tritium, among other radionuclides. There are difficulties in interpreting these measurement data that cannot be resolved satisfactorily at this time. These issues should not have a large impact on the worker's internal dose because exposure to uranium is the source of the greatest part of the internal dose.

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Table 5A-3 summarizes information developed in Section 5.3. Detection limits for some historical techniques remain to be identified, and will be reported in subsequent revisions as available.

Radionuclide /solubility type	Reported limit (nCi)	Reported limit (dpm)	Citation year
Th-232/M, S	0.6	1,300	1965
Np-237/M	2.7	5,900	1961
Np-237/M–Pa-233/M, S	0.255	560	1961
Co-60/M, S	0.66	1,500	1961
Zr-95/M, S–Nb-95/M, S	1.0	2,200	1961

Table 5A-3. Reported *in vivo* detection limits for other radionuclides.

Although internal monitoring has existed from the earliest days, the data from these measurements are not available for years before 1950. Guidance for the period from 1948 to 1950 is provided, but it is not feasible to provide any guidance for exposures before 1948. This will be provided in revisions to this document.

For dose reconstruction purposes and based on review of the Y-12 data, the 235 U and 238 U lung count detection thresholds are assumed to be 130 µg and 13.5 mg, respectively, through 1990. Beginning in 1991, the critical level or MDA is supplied with the individual results.