4.0 Radiological Protection and Doses

This section provides information on the Hanford Site radiological program and doses as well as cleanup activities as the U.S. Department of Energy (DOE) progresses toward site closure and the likely transfer of property to other entities. Additional information on radiation, dose rates, and dose terminology can be found in Appendix A, Helpful Information and Appendix B, Glossary.

4.1 External Radiation Monitoring

*CJ Perkins*

External radiation is defined as radiation originating from a source external to the human body. External radiation was monitored at the Hanford Site in relative proximity to known or potential radiation sources. Sources of external radiation at the Hanford Site include waste materials associated with the historical production of plutonium for defense; residual nuclear inventories in former production and processing facilities; radioactive waste handling, storage, and disposal activities; waste cleanup and remediation activities; atmospheric fallout from historical nuclear weapons testing; and natural sources such as cosmic radiation. During any given year, external radiation levels can vary from 15 to 25% at any location because of changes in soil moisture and snow cover (NCRP 1975).

The Harshaw™1 thermoluminescent dosimeter (TLD) system is used to measure external radiation on the Hanford Site. This type of TLD measures very low dose rates only and is not suitable for use for personnel monitoring. This system includes the Harshaw 8800-series dosimeter and the Harshaw 8800 reader. The Harshaw 8800-series environmental dosimeter consists of two TLD-700 chips and two TLD-200 chips and provides both shallow- and deep-dose measurement capabilities using filters in the dosimeter. Data obtained from the two TLD-700 chips were used to determine the average total environmental dose at each location. The two TLD-200 chips were included to determine doses in the event of a radiological emergency and were not used in calculating average total environmental dose. The average daily dose rate was determined by dividing the average total environmental dose by the number of days the dosimeter was exposed. Daily dose equivalent rates (mrem/day) at each location were converted to annual dose equivalent rates (mrem/yr) by averaging the daily dose rates and multiplying by 365 days/yr. The TLDs were positioned approximately 3.3 ft (1 m) above ground and were collected and read quarterly.

Radiation surveys with portable instruments are conducted to monitor and detect contamination and to provide a coarse screening for external radiation fields. The types of areas surveyed included underground radioactive material areas, contamination areas, soil contamination areas, high-contamination areas, roads, and fence lines.

4.1.1 External Radiation Measurements

External radiation fields were monitored in 2016 at 125 locations (Table 4-1) near Hanford Site facilities and operations. The TLD results were used individually or averaged to determine dose rates in a given area for a specific sampling period.

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1 Harshaw is a trademark of Thermo Fisher Scientific, Inc., Waltham, Massachusetts.
During 2016, 10 new TLD monitoring locations were added. These included:

- Three locations added in the 100/600 Areas to replace retired/retiring air sampling locations
- Three locations added at onsite air sampling locations supporting the Waste Treatment Plant (WTP)
- Three locations added at offsite (perimeter) air sampling locations supporting the WTP
- One reference location in Yakima.

<table>
<thead>
<tr>
<th>Location</th>
<th>No. of Dosimeters</th>
</tr>
</thead>
<tbody>
<tr>
<td>100-K Area</td>
<td>14</td>
</tr>
<tr>
<td>100 Area</td>
<td>4</td>
</tr>
<tr>
<td>200-East Area and WTP</td>
<td>45</td>
</tr>
<tr>
<td>200-Westest Area</td>
<td>24</td>
</tr>
<tr>
<td>200-Northorth Area</td>
<td>1</td>
</tr>
<tr>
<td>300 Area</td>
<td>14</td>
</tr>
<tr>
<td>400 Area</td>
<td>7</td>
</tr>
<tr>
<td>618-10 Burial Ground</td>
<td>4</td>
</tr>
<tr>
<td>CVDF</td>
<td>4</td>
</tr>
<tr>
<td>ERDF</td>
<td>3</td>
</tr>
<tr>
<td>IDF</td>
<td>1</td>
</tr>
<tr>
<td>Perimeter</td>
<td>3</td>
</tr>
<tr>
<td>Reference</td>
<td>1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>125</strong></td>
</tr>
</tbody>
</table>

CVDF=Cold Vacuum Drying Facility (100 K Area)
ERDF=Environmental Restoration Disposal Facility (200-Westest Area)
IDF=Integrated Disposal Facility (200-Eastast Area)
TEDF=300 Area Treated Effluent Disposal Facility
WTP= Waste Treatment Plant

In 2015, the Hanford External Dosimetry Program’s (HEDP) laboratory was relocated from its long-time location near the 300 Area to a location between the 200-East and 200-West Areas. This relocation introduced two substantial variances.

One significant difference was the higher background dose rate levels (approximately 50% higher) attributable to elevated radon levels inherent within the new HEDP facility (Figure 4-1). These background values are subtracted from the ambient dose rate levels measured in the environs to determine site-specific dose rate levels above background. This simple calculation, produced artificially, decreased dose rate values for 2015 when compared to previous years’ values.

The second significant difference that occurred during 2015 was the material used for shielding the dosimeters during their residency in the HEDP facility. Prior to 2015, the shielding material used was lead, which is substantially better than the steel shielding used in 2015. The effect of this change was to
expose the dosimeters to the significantly higher background levels, discussed in the previous paragraphs, and ultimately cause a higher dose rate reading.

Due to the complexities and uncertainties imparted on the TLD data by these changes at the HEDP facility, definitive annual data comparisons for the 2016 data are impractical.

The average dose rate levels measured in the operational areas during 2016 were comparable to the previous years’ levels (Figure 4-1). Individual TLD results and detailed maps of monitoring locations are available upon request.

4.1.1.1 100-K Area. As in years past, the 2016 dose rate levels near the load-out area of the 105-KW (reactor) Building, where radioactive contaminated sludge and debris from the cleanout of the 100-K West Basin was transported, were noticeably higher than other TLD locations at 100-K.

4.1.1.2 100 Areas. Dose rates measured along the Columbia River shoreline in the 100-N Area (N Springs) continued to decrease during 2016. Three new locations along the river corridor were established during September to provide continued radiological monitoring at air sampling locations that were retired (100-F and Hanford Townsite), or are expected to be retired in the near future (100-D).

4.1.1.3 200-East Area. Dose rate levels measured during 2016 near the “A” and “C” Tank Farms were higher than other 200-East Area locations. Three new locations in support of the WTP were added at air sampling locations “200 ESE” (N920), “B Pond” (N924), and “WTP New Station” (N584).
4.1.1.4 **200-West Area.** Dose rate levels measured during 2016 near the “S” and “T” Tank Farms and at the Solid Waste Operations Complex were higher than other 200-West Area locations.

4.1.1.5 **200-North.** Dose rates measured in 2016 were low, and all four quarterly measurements were similar.

4.1.1.6 **300 Area.** Dose rate levels measured during 2016 at locations in the southern portion of the 300 Area were slightly higher (5-10%) than at other 300 Area locations.

4.1.1.7 **400 Area.** Dose rates measured in 2016 at all seven monitoring locations were low and similar.

4.1.1.8 **Environmental Restoration Disposal Facility (ERDF).** Dose rates measured in 2016 at all three monitoring locations were low and similar.

4.1.1.9 **618-10 Burial Ground.** Dose rates measured in 2016 at all four monitoring locations were low and similar.

4.1.1.10 **Integrated Disposal Facility.** Dose rates measured in 2016 were low and all four quarterly measurements were similar.

4.1.1.11 **Perimeter Locations.** Three locations (Ringold, west end of Fir Road, and Dogwood Met Tower) were established in January 2016 and all four quarterly measurements were similar to each other and to onsite levels.

4.1.1.12 **Reference Locations.** One new location was added during September to provide a reference (aka, background) dose rate level station at the Yakima airport.

4.1.2 **Waste Disposal Sites Radiological Surveys**

*JW Wilde*

During 2016, 1,149 environmental radiological surveys were reported as performed at active and inactive waste disposal sites and the surrounding terrain to detect and characterize radioactive surface contamination. Radiation surveys with portable instruments are conducted to monitor and detect contamination and to provide a coarse screening for external radiation fields. The types of areas surveyed included underground radioactive material areas, contamination areas, soil contamination areas, high-contamination areas, roads, and fence lines. Vehicles equipped with radiation detection devices, and global positioning systems were used to measure accurately the extent of contamination along ERDF haul routes. Routine radiological survey locations included former waste disposal cribs and trenches, retention basin perimeters, ditch banks, solid waste disposal sites (e.g., burial grounds), unplanned release sites, tank farm perimeters, stabilized waste disposal sites, roads, and firebreaks in and around the Hanford Site operational areas. These sites were posted as underground radioactive material areas, contamination areas, and soil contamination areas.

Underground radioactive material areas are regions where radioactive materials occur below the soil surface. These areas are typically stabilized cribs, burial grounds, covered ponds, trenches, and ditches. Barriers have been placed over the contamination sources to inhibit radionuclide transport to the surface. These areas are surveyed at least annually to assess the effectiveness of the barriers.
Contamination areas and soil contamination areas may or may not be associated with an underground structure containing radioactive material. A breach in the surface barrier of a contaminated underground area may result in the growth of contaminated vegetation. Insects or animals may burrow into the soil and bring contamination to the surface. Vent pipes or risers from an underground structure may be sources of speck contamination (particles with a diameter less than 0.25 in. [0.6 cm]). Areas of contamination not related to subsurface structures can include sites contaminated with fallout from effluent stacks or with materials from unplanned releases (e.g., contaminated tumbleweeds and animal feces).

All contaminated areas may be susceptible to contaminant migration and are surveyed at least annually to assess their current radiological status. In addition, onsite paved roadways on which radioactive materials are transported to ERDF are surveyed annually.

4.2 Potential Radiological Doses

*Perona, RT Ryti, AG Fleury*

Potential radiological doses to the public and biota from Hanford Site operations in 2016 were evaluated to determine compliance with pertinent regulations and limits. Potential sources of radionuclide contamination included gaseous emissions from stacks and ventilation exhausts, contaminated groundwater seeping into the Columbia River, and fugitive emissions from areas of contaminated soil and operating facilities. A summary of the methods and results of the public and biota dose assessments is provided here. Details of the methods used to calculate radiological doses are provided in Appendix D.

The total annual dose to a hypothetical, maximally exposed individual (MEI) in 2016 at the offsite location where projected doses were highest (Horn Rapids Road) was 0.12 mrem (1.2 μSv). This dose is 0.12% of the 100 mrem (1000 μSv) per year public dose limit specified in DOE O 458.1, *Radiation Protection of the Public and the Environment*. For context, a 2009 National Council on Radiation Protection and Measurements report estimated that the overall annual exposure to ionizing radiation for the average American is 620 mrem (6,200 μSv), approximately half of which is related to natural sources and the other half attributable primarily to medical procedures.

The offsite MEI dose is one of eight radiological impacts of Hanford Site operations that are assessed in this chapter:

- Dose to a hypothetical, maximally exposed individual (MEI) at an offsite location, evaluated by using a multimedia pathway assessment DOE O 458.1 (Section 4.2.1)
- Collective dose to the population residing within 50 mi (80 km) of Hanford Site operation areas (Section 4.2.2)
- Dose for air pathways calculated using regulation-specified U.S. Environmental Protection Agency (EPA) methods for comparison to the *Clean Air Act* standards in 40 CFR 61, Subpart H, “National Emission Standards for Emissions of Radionuclides Other than Radon from Department of Energy Facilities” (Section 4.2.3)
• Dose from recreational activities, including hunting and fishing (Section 4.2.4.1)
• Dose to a worker consuming drinking water on the Hanford Site (Section 4.2.4.2)
• Dose to a visitor of the Manhattan Project National Historical Park (Section 4.2.4.3)
• Dose from non-DOE industrial sources on and near the Hanford Site (Section 4.2.5)
• Absorbed dose received by biota exposed to radionuclide releases to the Columbia River and to radionuclides in onsite surfacewater bodies (Section 4.2.6).

Radiological dose assessments related to environmental releases are ideally based on direct measurements of radionuclide concentrations in specific exposure media; however, amounts of many radioactive materials released to the Columbia River or the atmosphere from Hanford Site sources are too small to be measured in environmental media after they are dispersed in the offsite environment. For the radionuclides present in measurable amounts, it can be difficult to distinguish the small contribution of Hanford Site sources from contributions caused by fallout from historical nuclear weapons testing and naturally occurring radionuclides such as uranium and its decay products. As a result, computer models are employed to calculate offsite radionuclide concentrations based on measured and estimated releases. In specific instances, such as ambient air measurements of tritium at onsite and offsite locations near the 300 Area, radionuclide concentrations may be distinguishable from background levels; these measurements are used to support the dose assessment.

Calculations of radiation dose require the use of biological and radiological models of the behavior of radioactive material in the human body. Scientific understanding of these processes has improved over time. In the 1960s, the annual environmental reporting at the Hanford Site used the recommendations and methodologies of the International Convention on Radiological Protection (ICRP) Publication 2 (Permissible Dose for Internal Radiation [ICRP 1959]). In the 1970s, the annual reports began to follow the newer recommendations in ICRP Publication 26 (ICRP 1977) and Publication 30 (Limits for Intakes of Radionuclides by Workers, Part 1 [ICRP 1979a] and Supplement to Part 1 [ICRP 1979b]) incorporated in the dose factors from the EPA in Federal Guidance Reports 11 and 12 (EPA 1988 and EPA 1993, respectively). The GENII Version 1 computer code at the Hanford Site beginning in 1988 used ICRP Publications 26 and 30 as well as EPA dose factors. The GENII Version 2 computer code for the annual report dose calculations beginning in 2009 uses ICRP Publication 60 methods (ICRP 1991) and updated EPA dose factors (EPA 1999).

Offsite dose for an MEI (Section 4.2.1) and collective dose for population residing within 50 mi (80 km) of Hanford Site operation areas (Section 4.2.2) are calculated separately for liquid releases to the Columbia River and stack air emissions. Radiological doses from the water pathways are calculated based on differences in radionuclide concentrations between upstream and downstream sampling points on the Columbia River. Although the downstream minus upstream radionuclide concentrations potentially include groundwater-related contributions from other operating areas, they have been assigned to the 200 Areas for tabulation of radiological dose. No direct discharge of radioactive materials from the 100 or 300 Areas to the Columbia River was reported during 2016. Radiological doses from the air pathways are calculated based on stack emissions measurements from approximately 60 emission points in Hanford Site operation areas.
Columbia River shoreline spring and seep water containing radionuclides is known to enter the river along the portion of the Hanford Site shoreline extending from the 100-BC Area downstream to the 300 Area. Tritium and uranium isotopes were measured in the Columbia River downstream of the Hanford Site (Richland Pumphouse station, HRM 46.4) in 2016 at low concentrations that were nevertheless greater than upstream (Priest Rapids Dam station) levels (Appendix D). Radioactive air emissions are discussed in Section 6.1 and summarized in Table 6-1. For the GENII Version 2.10.1 (PNNL-14583) calculations supporting this dose assessment, ingrowth of short-lived radioactive progeny during environmental transport was calculated to develop a complete set of radionuclide release estimates. Details on the development of air pathway and water pathway radioactive release estimates are provided in Appendix D.

4.2.1 Maximally Exposed Individual Dose (Offsite Resident)
The MEI is a hypothetical person whose location and lifestyle are such that it is unlikely any actual member of the public would have received a higher radiological dose from Hanford Site releases during 2016. This individual’s exposure pathways were chosen to maximize the combined doses from all reasonable environmental routes of exposure to radionuclides in Hanford Site liquid effluents and air emissions using a multimedia pathway assessment (DOE O 458.1, Section 4.e). In reality, such a combination of maximized exposures to radioactive materials is highly unlikely to apply to any single person. The individual pathway dose calculations themselves also incorporate conservative assumptions intended to ensure that modeled concentrations of radionuclides in exposure media and resulting doses are protective. For these reasons, the dose assessment results for the MEI represent a hypothetical upper bound of potential individual dose rather than an anticipated dose to an actual individual.

The location of the hypothetical MEI varies depending on the relative contributions of radioactive air emissions and liquid effluent releases from Hanford Site operational areas. Four offsite locations were evaluated to determine the location of the offsite MEI (Figure 4-2). The Ringold location receives maximal air pathway impacts from the 200 Areas. Depending on annual differences in the prevailing wind direction, either the Sagemoor or Horn Rapids Road location may receive maximal air pathway impacts from the 300 Area. A small population of West Pasco residents obtain their drinking water from the Riverview location via a community water system; the domestic drinking water pathway is applied to that location. Residences in the vicinity of Horn Rapids Road receive drinking water from the City of Richland, which has an intake downstream of the Hanford Site; the domestic drinking water pathway is also applied here. Both Riverview and Horn Rapids Road are locations where Columbia River water is withdrawn for irrigation.

Dose calculations for 2016 releases indicate that the MEI is located in the vicinity of the Pacific Northwest National Laboratory (PNNL) Laboratory Support Warehouse, an offsite business just to the south of the Hanford Site 300 Area at 638 Horn Rapids Road. For the Horn Rapids Road receptor dose calculations, the radiological dose was modeled using the aforementioned Columbia River and air emissions data for the following exposure routes:

- Inhalation and external radiation exposure related to airborne radionuclides
- External radiation exposure and inadvertent soil ingestion for radionuclides deposited on the ground
- Ingestion of domestic drinking water from the Columbia River
• Ingestion of locally grown food products irrigated with Columbia River water and/or containing radionuclides deposited from the air

• External radiation exposure to radionuclides in Columbia River water and sediments near the Hanford Site during recreational activities (i.e., fishing, boating) and inadvertent ingestion of water while swimming

• Consumption of locally caught Columbia River nonmigratory fish.

A graphical depiction of the conceptual site model showing all potentially complete exposure pathways for the Horn Rapids Road MEI evaluated using GENII Version 2.10.1 (PNNL-14583) is provided in Figure 4-3. Additional information related to selection of the MEI location for releases is included in Appendix D. Exposure variable input values related to residency and recreational exposure times, intake rates for water, foods, other media, and agricultural pathway assumptions for the MEI are provided in Appendix D.

The total dose to the MEI at Horn Rapids Road in 2016 was calculated to be 0.12 mrem (1.2 μSv)/yr (Table 4-2; Figure 4-4). This dose is 0.12% of the 100 mrem (1000 μSv) per year public dose limit specified in DOE O 458.1 and 0.48% of the 25-mrem (250-μSv)/yr threshold where a supplemental assessment of dose to the lens of the eye, skin, and extremities is required. Air pathway contributions from sources in the 300 Area contributed approximately 83% of the total dose of 0.12 mrem (1.2 μSv)/yr, with the remaining dose related to water pathway exposures.

The primary radionuclides and exposure pathways contributing to the modeled MEI dose are as follows:

• **Air Releases.** Consumption of food products containing tritium released from the 300 Area contributed approximately 72% of the total air pathways dose of 0.10 mrem (1.0 μSv)/yr.

• **Water Releases.** Consumption of fish from the Columbia River contributed 0.014 mrem (0.14 μSv) or 54% of the total water pathways dose of 0.026 mrem (0.26 μSv)/yr. Consumption of food grown using Columbia River water withdrawn downstream from the Hanford Site contributed approximately another 32% of the total water pathways dose, and drinking water ingestion contributed the remaining 14%. Isotopes of uranium and their progeny, particularly uranium-234 and uranium-238, contribute approximately 97% of the total water pathways dose of 0.026 mrem (0.26 μSv)/yr. Most of the remaining 3% of the water pathways dose is related to tritium.
Figure 4-2. Locations Evaluated for Onsite and Offsite Receptors.
Figure 4-3. Conceptual Site Model of Exposure Pathways Evaluated in Dose Calculations (Horn Rapids Road MEI).
Table 4-2. Pathway Doses for the Hypothetical MEI Residing at Horn Rapids Road.

<table>
<thead>
<tr>
<th>Release Type</th>
<th>Exposure Pathway</th>
<th>Dose Contributions from Operational Areas (mrem)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Pathway Total</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>100 Area</td>
<td>200 Areas</td>
</tr>
<tr>
<td>Air</td>
<td>Food Ingestion</td>
<td>4.5E-07</td>
<td>1.4E-04</td>
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<tr>
<td></td>
<td>Inhalation</td>
<td>3.3E-06</td>
<td>4.5E-05</td>
</tr>
<tr>
<td></td>
<td>External, Soil Ingestion</td>
<td>4.2E-09</td>
<td>1.6E-07</td>
</tr>
<tr>
<td></td>
<td>Subtotal Air</td>
<td>3.8E-06</td>
<td>1.8E-04</td>
</tr>
<tr>
<td>Water</td>
<td>Irrigation (food and soil ingestion; external)</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.0083&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Drinking Water Ingestion</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.0040&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Recreation (river water, sediments; external, ingestion)</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;</td>
<td>5.0E-05&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Fish Ingestion</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.014&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Subtotal Water</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.026</td>
</tr>
<tr>
<td></td>
<td>Air + Water Total</td>
<td>3.8E-06</td>
<td>0.026</td>
</tr>
</tbody>
</table>

<sup>a</sup>To convert mrem to International System dose units (μSv), multiply by 10.

<sup>b</sup>No measured releases; the last 100 Area NPDES-permitted outfall (1908-K Outfall) ceased releases in March 2011.

<sup>c</sup>Integrates releases from all operational areas based on difference between down and upstream Columbia River radionuclide concentrations.

NA = Not applicable. All liquid discharges reflected in the difference between upstream and downstream radionuclide concentrations are assigned to the 200 Areas.

Figure 4-4. Total Dose for the Hypothetical MEI Over Time.

NOTE: 2016 doses calculated using GENII v2.10.1
4.2.1.1 MEI Dose Discussion. The MEI dose in 2016 of 0.12 mrem (1.2 μSv) is below the 0.21 mrem (2.1 μSv) MEI dose calculated in 2015 (DOE/RL-2016-33, Hanford Site Environmental Report for Calendar Year 2015). The difference between the 2016 and 2015 dose estimates is mostly attributable to larger 2015 releases of tritium and radon-220 from the 300 Area, and higher 2015 downstream concentrations of uranium-238 in the 2015 water pathways dose calculations.

The MEI dose estimate incorporates a number of conservative assumptions to ensure that pathway doses are protective and therefore calculated doses are likely to be overestimated. In the air pathways calculations, gross alpha and gross beta radiation measurements in stack emissions from the 100, 200, and 300 Areas were protectively added to the measured emissions of plutonium-239/-240 (an alpha-emitting radionuclide related to Hanford operations) and cesium-137 (a beta-emitting radionuclide related to Hanford operations), respectively. The actual measured total air releases of plutonium-239/-240 and cesium-137 are a small fraction (23 and 11%, respectively) of assumed releases that include the contribution of gross radioactivity values. Although gross alpha and gross beta levels in stack emissions are similar to ambient air background levels, the addition of these values ensures that possible contributions from any unmeasured operations-related radionuclides are protectively incorporated in the estimated doses.

In the irrigation pathways calculations, all produce eaten by the MEI was protectively assumed to be locally grown and originate from areas irrigated with Columbia River water. For the fish consumption pathway, near-shore water samples were protectively used to represent Columbia River water generally, and it was assumed that all fish consumed by the MEI are resident species rather than anadromous fish, such as salmon or steelhead. Because anadromous fish spend most of their lives in the ocean they would have a much lesser exposure to contaminants associated with the Hanford Reach compared to species that spend their entire lives in the Hanford Reach, such as carp and bass.

Because releases of tritium from the 300 Area are the major source of calculated Hanford-related radiological dose for the hypothetical MEI, modeled annual-average tritium concentrations at locations near the 300 Area were compared to concentrations based on air monitoring station samples. Figure 4-5 shows the 2016 modeled annual average air concentrations of tritiated water vapor (HTO) at the Horn Rapids Road MEI location and 2016 annual averages based on measured values at two offsite locations south and east of the 300 Area (Battelle Complex and Byers Landing) and two onsite locations near the southern border of the 300 Area northeast of the Horn Rapids Road MEI location. Measured monthly tritium concentrations vary substantially at each monitoring location. The 95% upper and lower confidence intervals of the annual average values are shown on Figure 4-5 in addition to the annual average.
Figure 4-5. Comparison of Measured and Modeled Tritium Air Concentrations Near the 300 Area.
NOTE: Error bars are 95% confidence intervals of the mean.

The modeled annual-average tritium concentration at the Horn Rapids Road MEI location is above the range of the 95% upper and lower confidence interval of the mean of the measured values at the two nearby offsite monitoring locations and the 300 South Gate onsite location. The Horn Rapids Road modeled annual-average tritium concentration is approximately equal to the 95% upper confidence interval of the mean of the measured values at the 300 South West monitoring station. A relationship between 300 Area monthly tritium air emissions and onsite 300 Area ambient air concentrations in 2006 was shown by Barfuss (2007), but there was little correlation of monthly emissions and air concentrations for a combined group of four nearby offsite monitoring locations. Figure 4-5 shows that the modeled MEI tritium air concentration is higher than the upper 95% confidence interval annual-average tritium concentrations measured at Battelle Complex (near the Horn Rapids Road MEI Location) and Byers Landing (near the Sagemoor MEI location) stations. This suggests that modeled tritium air concentrations may overestimate actual annual-average levels at these offsite locations.

Note that exact correspondence between modeled and measured annual average values would not be expected because the episodic nature of HTO releases is not captured in the GENII air dispersion modeling, which assumes a constant rate of HTO emissions. However, the modeled tritium values do not account for regional background levels of tritium, which would add between 1.5 and 4 pCi/m³ to the modeled values (Figure 11 in Barfuss 2007) and exacerbate the difference between modeled and measured tritium values at offsite locations near the 300 Area.

Samples of locally raised foodstuffs were collected in 2016 from four locations including the Sagemoor, Riverview, Sunnyside, and East Wahluke areas. Sampled foodstuffs included fruits (apricots, melons, and tomatoes), leafy vegetables, potatoes, corn, milk, and wine. Gamma-emitting radionuclides and
strontium-90 were measured in all foodstuffs, and tritium was measured in tomatoes and milk. Gamma-emitting radionuclides and tritium were measured in wine. Carbon-14 was measured in melons, corn, and leafy vegetables. Measured concentrations of the Hanford-related radionuclides carbon-14, cesium-137, and tritium in foods were compared to modeled concentrations calculated for the MEI receptor using the GENII computer code. These comparisons encompassed fruits, leafy vegetables, grain (corn), root vegetables (potatoes), and milk. Modeled concentrations of carbon-14, tritium and cesium-137 are related primarily to air emissions, whereas modeled concentrations of strontium-90 are related to irrigation with Columbia River water. The following observations are drawn from the comparisons:

- Carbon-14 was detected in only one of the 14 samples of melons, corn, and leafy vegetables collected from the Sagemoor, Riverview, Sunnyside, and East Wahluke areas. This measured concentration of 0.592 pCi/g in corn from Sunnyside was up to about one-third higher than the nondetect values (0.436, 0.454, and 0.480 pCi/g) reported for the three other corn samples. The modeled carbon-14 concentration in grain grown at the MEI location of Horn Rapids Road is approximately 1E-05 pCi/g, reflecting how naturally-occurring levels of carbon-14 in the environment are far higher than worst-case levels related to the carbon-14 stack emissions of 0.00012 Ci in 2016 (see Table D-2.)

- Cesium-137 was not detected in any food sample. Analytical detection limits were approximately a factor of 1,000 to 10,000 times larger than the worst-case modeled concentrations but comparable to or below environmental surveillance project dose-based reporting limits (DOE/RL-91-50) calculated using a 1 mrem (10 µSv)/yr threshold. Cesium-137 routine air releases are far below levels of detection or radiological concern in foodstuffs.

- Strontium-90 was detected only in two leafy vegetable samples from the Riverview and East Wahluke areas. The Riverview and East Wahluke areas had strontium-90 concentrations of 0.01 pCi/g and 0.007 pCi/g, respectively. Strontium-90 was not elevated in downstream Columbia River water samples in 2016 and therefore was not included in the water pathways dose calculations. Low levels of strontium-90 in the environment are widespread due to past above-ground weapons testing, and measured concentrations in leafy vegetables from the Sagemoor, Sunnyside, and East Wahluke areas are consistent with trends based on observations in offsite vegetation samples (PNNL-20577, Radionuclide Concentrations in Terrestrial Vegetation and Soil Samples On and Around the Hanford Site, 1971 Through 2008). Detected concentrations of approximately 0.01 pCi/g in leafy vegetables are almost 100 times below the environmental surveillance project dose-based reporting limit (DOE/RL-91-50).

- Tritium was measured in samples of tomatoes from the Sunnyside and Riverview areas but was not detected at either location with analytical detection limits of approximately 0.1 and 0.2 pCi/g, respectively. Tritium was detected in samples of milk at average concentrations of approximately 32 pCi/L (Sunnyside), 30 pCi/L (East Wahluke), and 38 pCi/L (Sagemoor). The relative tritium concentrations in milk from these three locations is consistent with expectations for air dispersion from releases in the 300 Area. However, these concentrations are well below the modeled worst-case tritium concentration in milk for cows grazing at the MEI location of Horn Rapids Road (approximately 480 pCi/L) and far less than the environmental surveillance project dose-based reporting limit of 17,000 pCi/L (DOE/RL-91-50).
4.2.2 Collective Dose

Collective dose is defined as the sum of doses to all individual members of the public within a defined distance of a specific release location. The regional collective dose from 2016 Hanford Site operations was estimated by calculating the radiological dose to the population residing within a 50-mi (80-km) radius of onsite operating areas (DOE O 458.1; Section 4.e(d)). The collective doses reported are based on regional population data from the 2010 census, as described in Appendix D.

The conceptual site model of potentially complete exposure pathways for the Horn Rapids Road MEI shown in Figure 4-3 is also applicable to the collective dose calculations. Like the Horn Rapids Road MEI, the collective dose calculation also incorporates the drinking water exposure pathway because the cities of Richland and Pasco obtain all or part of their municipal water directly from the Columbia River downstream from the Hanford Site, and the City of Kennewick obtains its municipal water indirectly from wells adjacent to the river. A primary distinction between the MEI and collective dose calculations is the use of population-average values for certain exposure variables in place of reasonable upper bound values. Exposure variable input values related to residency and recreational exposure times, intake rates for foods and other media, and agricultural pathway assumptions for the collective dose calculations are provided in Appendix D. The air pathways collective dose calculations employ population data from the 2010 census broken out according to direction and distance to coincide with air dispersion and deposition modeling conducted within the GENII Version 2.10.1 computer code (PNNL-14583).

The annual collective dose is reported in units of person-rem (person-sievert), which is the sum of doses to all individual members of the exposed population. The total collective dose calculated for this population in 2016 was 1.2 person-rem (0.012 person-Sv)/yr (Table 4-3), below the collective dose calculated in 2013-2015 and approximately equal to that calculated in 2012 (Figure 4-6). Air pathway contributions from releases in the 300 Area contributed approximately 60%, and water pathway contributions assigned to the 200 Areas contributed approximately 40% to the total collective dose of 1.2 person-rem (0.012 person-Sv) in 2016.

The primary radionuclides and exposure pathways contributing to the collective dose are as follows.

- **Air Releases.** Consumption of food products grown downwind of the 300 Area contributed approximately 67% of the of the air pathways collective dose of 0.72 person-rem (0.0072 person-Sv). The remaining air pathways collective dose is primarily related to inhalation. About 60% of these food and inhalation air pathways doses are due to releases of tritium from the 300 Area. Approximately another 20% of the total air pathways collective dose is associated with inhalation of the radioactive progeny of radon-220 released from the 300 Area. Air releases from the 100, 200, and 400 Areas had negligible contributions to the air pathways collective dose.

- **Water Releases.** Consumption of drinking water drawn from the Columbia River downstream of the Hanford Site contributed approximately 97% of the total water pathways collective dose of 0.45 person-rem (0.0045 person-Sv). Two isotopes of uranium (-234 and -238) and their progeny from releases assigned to the 200 Areas were the largest contributors (approximately 88%) to the drinking water collective dose.

The collective dose in 2016 of 1.2 person-rem (0.012 person-Sv) is below the 1.7 person-rem (0.017 person-Sv) collective dose calculated in 2015 (DOE/RL 2016 33) and approximately equal to
the 2012 collective dose. There is no specific collective dose metric analogous to the 100 mrem (1,000 mSv) per year public dose limit for individual exposures described in Section 4.2.

### Table 4-3. Collective Pathway Doses within a 50-mi (80-km) Radius.

<table>
<thead>
<tr>
<th>Release Type</th>
<th>Exposure Pathway</th>
<th>Dose Contributions from Operational Areas, person-rem&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>Food Ingestion</td>
<td>7.9E-05, 0.013, 0.48, 4.4E-05, 0.50</td>
</tr>
<tr>
<td></td>
<td>Inhalation</td>
<td>0.0012, 0.007, 0.21, 1.8E-04, 0.22</td>
</tr>
<tr>
<td></td>
<td>External, Soil Ingestion</td>
<td>9.3E-07, 1.3E-05, 0.0027, 3.7E-07, 0.0027</td>
</tr>
<tr>
<td></td>
<td><strong>Subtotal Air</strong></td>
<td>0.0012, 0.020, 0.7, 2.2E-04, 0.72</td>
</tr>
<tr>
<td>Water</td>
<td>Irrigation (food and soil ingestion; external)</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;, 0.0086&lt;sup&gt;c&lt;/sup&gt;, NA, NA, 0.009</td>
</tr>
<tr>
<td></td>
<td>Drinking Water Ingestion</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;, 4.9E-04&lt;sup&gt;c&lt;/sup&gt;, NA, NA, 4.9E-04</td>
</tr>
<tr>
<td></td>
<td>Recreation (river water, sediments; external, ingestion)</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;, 0.0053&lt;sup&gt;c&lt;/sup&gt;, NA, NA, 0.0053</td>
</tr>
<tr>
<td></td>
<td>Fish Ingestion</td>
<td>NA&lt;sup&gt;b&lt;/sup&gt;, 0.44&lt;sup&gt;c&lt;/sup&gt;, NA, NA, 0.44</td>
</tr>
<tr>
<td></td>
<td><strong>Subtotal Water</strong></td>
<td>NA, 0.45, NA, NA, 0.45</td>
</tr>
<tr>
<td></td>
<td>Air + Water Total</td>
<td>0.0012, 0.47, 0.7, 2.2E-04, 1.2</td>
</tr>
</tbody>
</table>

<sup>a</sup> To convert person-rem to International System dose units (person-Sv), divide by 100.

<sup>b</sup> No measured releases; the last 100 Area NPDES-permitted outfall (1908-K Outfall) ceased releases in March 2011.

<sup>c</sup> Integrates releases from all operational areas based on difference between down- and upstream Columbia River radionuclide concentrations.

NA = not applicable. All liquid discharges reflected in difference between up- and downstream radionuclide concentrations assigned to 200 Areas.

![Figure 4-6. Collective Total Dose within a 50-mi (80-km) Radius.](image)

**4.2.3 Compliance with Clean Air Act Standards**

Historically at the Hanford Site, there has been one primary expression of radiological risk to an offsite individual; however, the MEI dose is currently calculated by two different methods in response to two different requirements. One MEI dose computation is required by DOE O 458.1 and is calculated using...
the GENII computer code as described in Section 4.2.1 of this report. This calculation considers all potential environmental exposure pathways (e.g., from releases to both air and water) that maximize a hypothetical offsite individual’s exposure to the Hanford Site’s radiological liquid effluents and air emissions. A second estimate of MEI air pathways dose is required by the Clean Air Act and must be calculated using an EPA air dispersion and dose modeling computer code (CAP-88; EPA 2000) or other methods accepted by the EPA under the Clean Air Act to demonstrate compliance with 40 CFR 61, Subpart H requirements. This regulation specifies that no member of the public shall receive a dose greater than 10 mrem (100 μSv)/yr from exposure to airborne radionuclide emissions (other than radon) released at DOE facilities. The Hanford Site stack emissions and emissions from diffuse and unmonitored sources (e.g., windblown dust) are considered in the offsite dose for the Clean Air Act and are based solely on an airborne radionuclide emissions pathway.

The assumptions embodied in the CAP-88 computer code differ slightly from standard air pathways assumptions used with the GENII computer code; therefore, air-pathway doses calculated by the two codes may differ somewhat. In principle, the MEI for air pathways assessed under 40 CFR 61, Subpart H may be evaluated at a different location from the all-pathways MEI if dose from the water pathways exceeds that from air pathways (Appendix D).

The Clean Air Act regulation also requires that an annual report for each DOE facility be submitted to EPA that supplies information about atmospheric emissions for the preceding year and any potential contributions to offsite dose. For more detailed information about 2016 air emissions at the Hanford Site, refer to DOE’s report to EPA (DOE/RL-2016-10).

4.2.3.1 Dose from Stack Emissions to an Offsite Maximally Exposed Individual. Using CAP-88, the offsite MEI for air pathways in 2016 was at the PNNL Richland Campus’ Laboratory Supply Warehouse, an offsite business located in north Richland, Benton County, Washington, directly south of the Hanford Site 300 Area and proximal to the Horn Rapids Road MEI location (Figure 4-2). The potential air pathway dose from stack emissions to an MEI at that location calculated using the CAP-88 computer code was determined to be 0.038 mrem (0.38 μSv)/yr, less than 1% of the EPA standard of 10 mrem (100 μSv)/yr. The CAP-88 result is approximately one-half of the air pathway dose of 0.10 mrem (1.0 μSv) for stack emissions calculated with GENII (Table 4-2).

Dose related to radon-220 is not included in the dose calculated for EPA compliance in 40 CFR 61, Subpart H but is regulated by the 10-mrem (100-μSv)/yr standard established in WAC 246-247, “Radiation Protection – Air Emissions.” A release of 178 curies of radon-220 was calculated from engineering estimates for stack emissions from the 325 Building in the 300 Area. A radon-220 dose of 0.026 mrem (0.26 μSv)/yr was calculated using the CAP-88 computer code for the Laboratory Supply Warehouse MEI, far below the WAC 246-247 standard. The sum of MEI dose for radon-220 and dose calculated for compliance with 40 CFR 61, Subpart H using the CAP-88 computer code is approximately 0.064 mrem (0.64 μSv), which is about 60% of the total Horn Rapids Road air pathways MEI dose of 0.10 mrem (1.0 μSv) calculated using the GENII computer code.
4.2.3.2 Dose from Diffuse and Fugitive Radionuclide Emissions to an Offsite Maximally Exposed Individual. The December 15, 1989, revisions to 40 CFR 61, Subpart H required DOE facilities to estimate the dose to a member of the public for radionuclides released from all potential sources of airborne radionuclides. DOE and EPA interpreted the regulation to include diffuse and fugitive (nonpoint source) emissions, as well as emissions from monitored point sources (i.e., stacks) described in Section 4.2.3.1. EPA has not specified or approved standardized methods to estimate diffuse airborne emissions because of the wide variety of sources at DOE sites. The method developed at the Hanford Site to estimate potential diffuse emissions is based on environmental monitoring measurements of airborne radionuclides at the site perimeter (DOE/RL-2016-10). Modeled contributions from monitored stack emissions and contributions from background levels of radionuclides are subtracted from perimeter ambient air concentrations measured for each radionuclide and positive differences are attributed to a virtual fugitive source located near the center of the Hanford Site.

The Laboratory Supply Warehouse location immediately south of the 300 Area was chosen for purposes of demonstrating compliance with the MEI dose standard for diffuse and fugitive emissions (DOE/RL-2016-10). The estimated dose from diffuse emissions to an MEI was calculated using the CAP-88 computer code to be 0.0060 mrem (0.060 μSv)/yr. Therefore, the potential combined dose from stack emissions, radon-220 emissions, and diffuse emissions (excluding radon) during 2016 at the Laboratory Supply Warehouse location was 0.070 mrem (0.70 μSv)/yr, far below the 10 mrem (100 μSv) per year federal and state standards described above.

4.2.3.3 Maximum Dose to Non-U.S. Department of Energy Workers at the Hanford Site. DOE allows private businesses to locate their activities and personnel on some regions of the Hanford Site. The EPA Region 10 Office and the Washington State Department of Health provided guidance to the U.S. Department of Energy, Richland Operations Office (DOE-RL) that, when demonstrating compliance with 40 CFR 61 standards, it should evaluate potential doses to non-DOE employees who work at facilities within the Hanford Site but who are not under direct DOE control. This situation has created the need to calculate a maximum dose for an onsite individual employed by a non-DOE business who works within the boundary of the Hanford Site.

Doses to members of the public employed at non-DOE facilities at locations outside access-controlled areas on the Hanford Site (those requiring DOE-access authorization for entry) were evaluated in the 2016 EPA air emissions report (DOE/RL-2016-10) as possible MEI locations. Included in these locations were the Columbia Generating Station operated by Energy Northwest and Laser Interferometer Gravitational Wave Observatory (LIGO) operated by the University of California (Figure 4-2). The non-DOE worker dose due to stack emissions at these facilities was calculated using the CAP-88 computer code assuming full-time occupancy because EPA guidance does not allow for adjustment of such doses to account for less than full-time occupancy. The highest estimated dose to a member of the public from fugitive emissions was at LIGO. The total dose attributable to 2016 stack emissions, fugitive source emissions, and radon-220 at LIGO was calculated using CAP-88 to be 0.026 mrem (0.26 μSv; DOE/RL-2016-10). Even assuming that a LIGO employee is continuously present, the estimated total dose to non-DOE onsite workers in 2016 was lower than the 0.070 mrem (0.70 μSv)/yr total dose calculated with CAP-88 to an offsite MEI at the Laboratory Supply Warehouse.

4.2.4 Special Case Dose Estimates
The exposure assumptions used to calculate the dose to the MEI were selected to provide a scenario yielding a reasonable upper bound dose estimate. The MEI dose calculations are based on
measurements of radionuclide releases from stack emissions (air pathways) and differences between
downstream and upstream radionuclide concentrations in the Columbia River (water pathways),
followed by modeling of environmental transport related to a number of different exposure pathways
(Figure 4-3). Exposure pathways using other radionuclide measurements also exist that could have
resulted in radiological exposures. Three such scenarios include an outdoor recreationalist who
consumed meat from contaminated wildlife that migrated from the Hanford Site; an individual who
drank water from one of four DOE-owned water treatment facilities at the Hanford Site; and a visitor to
the Manhattan Project National Historical Park. The potential doses resulting from these scenarios are
examined in the following sections.

4.2.4.1 Outdoor Recreationalist Dose. Wildlife has access to Hanford Site areas that are
contaminated with radioactive materials and have the potential to acquire radioactive contamination
and migrate offsite. Wildlife sampling was conducted at the Hanford Site to estimate radionuclide tissue
concentrations in animals from the site that could potentially have been hunted offsite. An outdoor
recreationalist is also potentially exposed to contaminated soil and sediment along the river corridor if
they access this area from the Columbia River.

Concentrations of radionuclides measured in soil (cesium-137, plutonium-238, plutonium-239/-240, and
strontium-90) at far field sampling locations are not readily distinguishable from background levels, and
soil concentrations are less susceptible to yearly variation than sediment and wildlife. An evaluation of
radionuclide soil concentrations and trends over time is provided in PNNL-20577. Review of the 2016
sediment data indicates that concentrations of key radionuclides frequently detected in sediment
(including cesium-137, plutonium-239/-240, and uranium isotopes) have approximately equal
concentrations at upstream (Priest Rapids Dam) and downstream (McNary Dam) locations. Also,
sediment concentrations at the dam locations are generally as large as or larger than concentrations at
slough locations along the Hanford Site near White Bluff and the Hanford Townsite. The 2016 sediment
data do not indicate the presence of a Hanford contribution to sediment radionuclide concentrations.
Therefore, the screening assessment of outdoor recreationalist dose will focus on wildlife samples
obtained in 2016.

Gamma-emitting radionuclides were analyzed in muscle tissue samples collected in 2016 from elk, mule
deer, and quail. In addition to muscle tissue, samples of liver tissue were obtained from elk and mule
deer and analyzed for gamma-emitting radionuclides and isotopic plutonium. Bone samples were also
collected from elk, mule deer, and quail and analyzed for strontium-90, a radionuclide that accumulates
in bone tissue. For estimating dose from ingestion of game meat, radionuclide concentrations in muscle
tissue are most applicable. The only radionuclide detected in the muscle and liver tissue of any animal
was potassium-40, a naturally occurring primordial radioisotope not of Hanford Site origin.

Fillet tissue and carcass samples were obtained from carp and bass in two river sections of the Hanford
Reach in 2016. Fillet samples were analyzed for gamma-emitting radionuclides, tritium, strontium-90,
and isotopes of plutonium and uranium. Carcass samples were only analyzed for strontium-90. Detected
radionuclides in fillet samples were limited to potassium-40, uranium-234, uranium-235, and
uranium-238. Potassium-40 is a naturally occurring radionuclide that is not of Hanford Site origin.
However, uranium isotopes are associated with Hanford Site operations.

Uranium-234 was detected in three carp fish fillet samples from the 100 Area, three fillet samples from
the 300 Area, and one fillet sample from the reference area. Uranium-235 was detected in two carp fish
fillet samples from the 300 Area and two fillet samples from the reference area. Uranium-238 was detected in two carp fish fillet samples from the 100 Area, two fillet samples from the 300 Area, and two fillet samples from the reference area. Both average and maximum isotopic uranium concentrations detected in the reference area carp fillet samples were higher than the values detected in the 100 Area fillet samples. Average reference area carp fillet concentrations of uranium-234 and uranium-235 were higher than those from 300 Area fillet samples, but uranium-238 concentrations were lower than those from the 300 Area fillet samples. In terms of maximum values, the reference area fillet samples were all lower than the 300 Area fillet samples. As a result of the variability in the isotopic uranium concentrations in carp fish fillets between the 300 Area and the reference area, the potential radiation dose from consumption of carp fish fillets with isotopic uranium concentrations were examined for the 300 Area.

These uranium-234 and uranium-235 results for carp are similar to the respective uranium isotopes in whitefish fillet samples collected in 2015. The uranium-238 results for carp are similar to sampling results for uranium-238 in carp fillet samples collected in 2014, where concentrations were observed to increase in carp fillet samples with downstream distance from an upstream reference area from the 100 Area to the 300 Area. Differences in sampling locations and species may explain these differences between 2014 carp and 2015 whitefish fish fillet results; Hanford Site uranium releases to the Columbia River estimated from downstream and upstream river concentrations were approximately equivalent in these 2 years.

Isotopic uranium concentrations in bass fish fillet samples were detected in the 100 Area, 300 Area, and reference area. One bass fillet sample was collected for each area and uranium-234, uranium-235, and uranium-238 were all found within each sample. Uranium-235 concentrations detected in the reference area bass fillet samples were all higher than the values detected in the 100 and 300 Area fillet samples. Uranium-234 and uranium-238 concentrations detected in the reference area bass fillet samples were all lower than the values detected in the 100 and 300 Area fillet samples. These uranium isotope results for bass are similar to sampling results for uranium-234 and uranium-238 in carp fillet samples collected in 2014. The potential radiation dose from consumption of bass fish fillets with the measured isotopic uranium concentrations were examined for the 100 and 300 Areas.

The potential radiation dose received from consumption of fish fillets with isotopic uranium concentrations measured in bass in 2016 would be negligible. Assuming annual fish consumption of 88 lb (40 kg) for an MEI (Table D-4), the annual radiation dose related to fish ingestion for bass that contains isotopic uranium is estimated to be 0.17 mrem (1.7 μSv) in the 100 Area and 0.16 mrem (1.6 μSv) in the 300 Area.

The potential radiation dose received from consumption of fish fillets with average isotopic uranium concentrations measured in carp from the 300 Area in 2016 would also be negligible. Assuming annual fish consumption of 88 lb (40 kg) for an MEI (Table D-4), the annual radiation dose related to fish ingestion for carp is estimated to be 0.064 mrem (0.64 μSv).

The dose estimate for carp ingestion was derived using the average value from 300 Area fillet samples for each isotopic uranium concentration and an ingestion dose factor of 1.8 × 10⁻⁴ mrem/pCi (4.9 × 10⁻² μSv/Bq) for uranium-234, 1.7 × 10⁻⁴ mrem/pCi (4.6 × 10⁻² μSv/Bq) for uranium-235, and 1.7 × 10⁻⁴ mrem/pCi (4.6 × 10⁻² μSv/Bq) for uranium-238 from ICRP Publication 72 (ICRP 1995) in the following manner:
The dose estimate for ingestion of bass fillets was derived using the measured value for each isotopic uranium concentration and an ingestion dose factor of $1.8 \times 10^{-4}$ mrem/pCi (4.9 $\times 10^{-2}$ μSv/Bq) for uranium-234, $1.7 \times 10^{-4}$ mrem/pCi (4.6 $\times 10^{-2}$ μSv/Bq) for uranium-235, and $1.7 \times 10^{-4}$ mrem/pCi (4.6 $\times 10^{-2}$ μSv/Bq) for uranium-238 from ICRP Publication 72 (ICRP 1995) in the following manner:

- **100 Area**

$$((0.011 \text{ pCi uranium-234/g } \times 1.8 \times 10^{-4} \text{ mrem/pCi}) + (0.0060 \text{ pCi uranium-235/g } \times 1.7 \times 10^{-4} \text{ mrem/pCi}) + (0.0069 \text{ pCi uranium-238/g } \times 1.7 \times 10^{-4} \text{ mrem/pCi})) \times 40 \text{ kg/yr } \times 1,000 \text{ g/kg} = 0.17 \text{ mrem (1.7 μSv)/yr}$$

- **300 Area**

$$((0.011 \text{ pCi uranium-234/g } \times 1.8 \times 10^{-4} \text{ mrem/pCi}) + (0.0047 \text{ pCi uranium-235/g } \times 1.7 \times 10^{-4} \text{ mrem/pCi}) + (0.0069 \text{ pCi uranium-238/g } \times 1.7 \times 10^{-4} \text{ mrem/pCi})) \times 40 \text{ kg/yr } \times 1,000 \text{ g/kg} = 0.16 \text{ mrem (1.6 μSv)/yr}$$

### 4.2.4.2 Hanford Site Drinking Water Dose.

Drinking water was sampled and analyzed for tritium, strontium-90, gross alpha radiation, and gross beta radiation during 2016 in accordance with applicable regulations (40 CFR 141); water samples were collected from the 100-K Area, 200-West Area, and three sources in the 400 Area (a primary well and two emergency backup wells). The water supply for the 100-K and 200-West Areas is the Columbia River, whereas the primary and backup water supplies for the 400 Area are groundwater wells (see Section 7.1).

A comparison of analytical results for the 100-K, 200, and 400 Areas drinking water samples to state and federal standards is provided in Section 7.1. Tritium and strontium-90 are both man-made soluble beta radiation emitters; there are also naturally occurring beta emitters in the uranium, actinium, and thorium decay series. Potential onsite drinking water dose from Hanford-related beta-emitting radionuclides is addressed in this section by evaluating drinking water data for tritium and strontium-90.

Strontium-90 was analyzed in one sample from each of the five drinking water sources in 2016 and was not identified above its analytical detection limit in any drinking water sample. Tritium was analyzed in one sample from both the 100-K and 200-West Areas and was not detected above its analytical detection limit in either sample. Tritium was detected in all four drinking water samples collected from the primary drinking water sources for the 400 Area (well P-16) and also in two samples from backup wells P-14 and P-15. Based on the four quarterly samples from the primary well, the annual average 400 Area drinking water tritium concentration was 2,223 pCi/L (82 Bq/L). Assuming a consumption rate of 0.26 gal (1 L)/day for 250 working days at the Fast Flux Test Facility in the 400 Area, the potential annual worker dose in 2016 would be approximately 0.037 mrem (0.37 μSv). The single tritium samples collected at each of the backup wells are independently assessed for worker dose because it is unlikely that both backup wells would be active at the same time and the water supply blended. The drinking water tritium concentration at backup well P-14 was 12,000 pCi/L. Based on this single measurement, an annual worker drinking water dose for water obtained exclusively from the backup P-14 well would be 0.20 mrem (2.0 μSv). The drinking water tritium concentration at backup well P-15 was 2320 pCi/L.
resulting in an annual worker drinking water dose of 0.039 mrem (0.39 μSv). These estimates are well below EPA’s drinking water dose limit of 4 mrem (40 μSv)/yr for beta-emitting radionuclides in public drinking water supplies.

The dose estimate for the primary 400 Area drinking water source was derived using a tritium ingestion dose factor of $6.7 \times 10^{-8}$ mrem/pCi (1.8 × 10⁻⁵ μSv/Bq) from ICRP Publication 72 (ICRP 1995) in the following manner:

$$2,223 \text{ pCi tritium/L} \times 1 \text{ L/day} \times 250 \text{ d/year} \times 6.7 \times 10^{-8} \text{ mrem/pCi} = 0.037 \text{ mrem/yr}$$

4.2.4.3 Manhattan Project National Historical Park Visitor Dose. The Manhattan Project National Historical Park at Hanford includes guided tours of the B Reactor as well as access to several pre-Manhattan Project locations, two of which (Hanford Townsite and White Bluffs Bank) are situated to the east of the 100-K and the 200 Areas. These historical locations are geographically closer to these air emissions sources than the offsite MEI locations evaluated in Section 4.2.1. However, unlike an offsite residential MEI receptor, visitors to these locations would not be exposed from agricultural and drinking water exposure pathways, nor would they be continually exposed over the course of a year, as might be anticipated for some residents. For these reasons, potential doses at these locations are likely to be considerably below those calculated for the hypothetical offsite MEI.

Inhalation dose related to 100-K and 200 Areas stack emissions was calculated for a hypothetical individual at the Hanford Townsite and White Bluffs Bank locations using the GENII Version 2.10.1 computer code. Although Historical Park visitors would be present only briefly and on a single occasion at these locations, individuals conducting tours could be present for greater lengths of time. Additionally, these locations are adjacent to the Columbia River where recreationalists might be exposed while boating, fishing, or engaging in other activities. For this screening calculation, continuous exposure at the Hanford Townsite and White Bluffs Bank locations was assumed. The results of these dose calculations are presented in Table 4-4.

<table>
<thead>
<tr>
<th>Release Type</th>
<th>Exposure Pathway</th>
<th>Location</th>
<th>Dose Contributions from Operational Areas, mrem²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>Inhalation</td>
<td>Hanford Townsite</td>
<td>4.6E-05</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8.1E-05</td>
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<td></td>
<td>1.3E-04</td>
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<td></td>
<td></td>
<td>White Bluffs Bank</td>
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<td>1.6E-04</td>
</tr>
</tbody>
</table>

*To convert mrem to International System dose units (μSv), multiply by 10.*

Radiological doses assuming continuous inhalation exposure at either the Hanford Townsite or White Bluffs Bank locations are far below the hypothetical offsite MEI air pathways dose of 0.10 mrem (1.0 μSv; Table 4-2) at Horn Rapids Road.

4.2.5 Doses from Non-U.S. Department of Energy Sources
Doses from non-DOE sources were not quantified in 2016 because the MEI dose of 0.12 mrem (1.2 μSv)/yr from DOE-related sources (Section 4.2.1) was far below the threshold of 25 mrem.
(250 μSv)/yr at which the contribution of non-DOE sources must be included. DOE O 458.1 paragraph 4.e(1)(c) states that dose evaluations to demonstrate compliance with the public dose limit must include:

> [t]he dose to members of the public from DOE-related exposure sources only, if the projected DOE-related dose to the representative person or MEI is 25 mrem (250 μSv) in a year or less. If the DOE-related dose is greater than 25 mrem (250 μSv) in a year, the dose to members of the public must include major non-DOE sources of exposure and dose from DOE-related sources.

Before it was superseded by the release of DOE O 458.1 in 2011, DOE O 5400.5 provided the applicable requirements for radiation protection of members of the public. Chapter II, Paragraph 7 of DOE O 5400.5, Chg 2 has a reporting requirement for a combined dose due to DOE and other manmade sources. Therefore, Hanford Site environmental reports prior to 2011 routinely evaluated dose contributions from various non-DOE industrial sources of radiation exposure on or near the Hanford Site. These included a commercial, low-level radioactive waste burial ground at the Hanford Site operated by the Washington State Department of Ecology; a nuclear power-generating station at the Hanford Site operated by Energy Northwest; a nuclear-fuel production plant operated near the site by AREVA NP, Inc.; a commercial, low-level radioactive waste treatment facility operated near the site by Perma-Fix Northwest, Inc.; and a commercial decontamination facility operated near the site by Perma-Fix Northwest, Inc. (Figure 4-2). The total individual dose from non-DOE source activities in 2010 was conservatively estimated at about 0.004 mrem (0.04 μSv)/yr (PNNL-20548).

### 4.2.6 Dose to Non-Human Biota

Dose assessments for non-human biota evaluate the potential for exposures from Columbia River sediment and water, soils (near facilities), and exposures associated with West Lake. Upper estimates of the radiological dose to aquatic organisms were made in accordance with the DOE O 458.1 requirement for management and control of liquid discharges and air emissions. The current dose limit for aquatic animal organisms is 1 rad (10 milligray [mGy]) per day. Rad is a unit of absorbed dose of ionizing radiation equal to an energy of 100 ergs/g of irradiated material. In addition to the dose limit for aquatic organisms, there is a dose limit for riparian or terrestrial wildlife of 0.1 rad (1 mGy)/day.

Concentration guides for assessing doses to biota are very different from the DOE-derived concentration standards used to assess radiological doses to humans. A tiered approach is used to estimate radiological doses to aquatic and terrestrial biota. This method uses the RESidual RADIOactive (RESRAD)-BIOTA computer code ([DOE/EH-0676, User’s Guide, Version 1, RESRAD-BIOTA: A Tool for Implementing a Graded Approach to Biota Dose Evaluation; DOE-STD-1153-2002, A Graded Approach for Evaluating Radiation Doses to Aquatic and Terrestrial Biota](#)) to compare radionuclide concentrations measured by routine monitoring programs to a set of biota concentration guides.

Biota concentration guides are the soil, water, or sediment concentrations of a radionuclide that would produce 1 rad (10 mGy)/day for aquatic biota or terrestrial plants or 0.1 rad (1 mGy)/day for riparian or terrestrial wildlife. For samples containing multiple radionuclides, a sum of fractions is calculated to account for the contribution to dose from each radionuclide relative to the dose limit. If the sum of fractions exceeds 1.0, then the dose limit has been exceeded. If the initial estimated screening value (Tier 1) exceeds the guideline (sum of fractions more than 1.0), additional screening calculations are performed (Tier 2 or Tier 3) to evaluate more accurately exposure of the biota to the radionuclides. The
process may culminate in a site-specific assessment requiring additional sampling and study of exposure. Biota-dose screening assessments were conducted using surveillance data collected in 2016 from on and around the Hanford Site.

Researchers used the RESRAD-BIOTA computer code to evaluate potential effects on biota from the maximum concentrations of radionuclides measured in Columbia River sediment and water as tabulated in Appendix C. The detected radionuclides evaluated across all locations in the Columbia River sediment and water biota dose assessment are carbon-14, cobalt-60, cesium-137, plutonium-239/-240, strontium-90, tritium, uranium-234, uranium-235, and uranium-238. Beryllium-7 was detected but is of cosmogenic origin and is not associated with the Hanford Site. Potassium-40 was also detected in sediments upstream, onsite, and downstream of the Hanford Site. Potassium-40 is a naturally occurring radionuclide and is not associated with releases from the reactors or any groundwater plumes entering the Columbia River. Therefore, dose associated with potassium-40 is not included in the biota dose assessment. Most of the locations located on the Columbia River had samples collected from riverbank springs or seeps that carry groundwater contaminants into the Columbia River. Concentrations in springs or seeps are greater than those observed in the river water; therefore, the dose assessment results for these discrete areas of elevated concentrations are protective relative to the potential for impacts on populations of biota in the Columbia River. For an initial screen of ecological populations, the sediment and water data were split into five subareas: upstream, 100 Area, Hanford Townsite, 300 Area, and downstream and the maxima concentrations evaluated in these locations. If risks to biota were identified in the initial screen, then further assessments using average concentration over smaller spatial units would be evaluated. The results of the screening calculations listed in Table 4-5 show the concentrations in all Columbia River sediment and water samples passed the Tier 1 screen and indicate that the calculated doses were below dose limits (sum of fractions less than one). Most of the estimated dose in the 100 Area is from carbon-14 (70%) and strontium-90 (26%) and dose in the 300 Area is basically entirely associated with uranium isotopes. Biota doses upstream at the Hanford Townsite and downstream were all similar and likely related to background concentrations in water and sediment. Further documentation of the Columbia River biota dose calculations is provided in Appendix D.

Biota dose calculations also were completed for West Lake, located on the Central Plateau of the Hanford Site. West Lake is a vernal pool or ephemeral wetland that fills with water during the winter and generally becomes smaller or dries up entirely in other seasons. West Lake is part of the 200 Areas Unplanned Release Waste Group Operable Unit (200-UR-1 Operable Unit), and is planned for supplemental characterization (DOE/RL-2009-121, Sampling and Analysis Plan for the West Lake Site). The results of these planned investigations will be presented in the appropriate Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA) remedial action document for the 216-N-8 waste site. In parallel with these planned CERCLA studies, this program has been collecting sediment data annually. In addition, other media (water and biota) have been collected from West Lake on a less regular schedule. Both sediment and water samples were collected in 2016 and data tabulated (Appendix C, Tables C-2, C-3, and C-4).

The results of the 2016 screening calculations listed in Table 4-6 show the West Lake sediment and water concentrations failed the Tier 1 and 2 screens. The Tier 1 screen was based on the maximum concentration, and the Tier 2 screen was based on the average concentrations of five water and three sediment samples. The estimated biota dose for Tiers 1 and 2 was almost entirely due to the measured concentration of uranium in water and the assumed potential for uptake from water to aquatic biota using a default bioaccumulation factor.
Table 4-5. Estimated Doses to Biota Associated with Columbia River Sediment and Water. 

<table>
<thead>
<tr>
<th>Location</th>
<th>Media Sampled for Key Radionuclidesb</th>
<th>Tier 1 Screen Sum of Fractionsc</th>
<th>2015</th>
<th>2016</th>
<th>Pass or Fail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upstream</td>
<td>Sediment, Water</td>
<td></td>
<td>0.015</td>
<td>0.018</td>
<td>Pass</td>
</tr>
<tr>
<td>100 Area</td>
<td>Sediment, Water</td>
<td></td>
<td>0.64</td>
<td>0.71</td>
<td>Pass</td>
</tr>
<tr>
<td>Hanford Townsite</td>
<td>Sediment, Water</td>
<td></td>
<td>0.016</td>
<td>0.014</td>
<td>Pass</td>
</tr>
<tr>
<td>300 Area</td>
<td>Water</td>
<td></td>
<td>0.30</td>
<td>0.25</td>
<td>Pass</td>
</tr>
<tr>
<td>Downstream</td>
<td>Sediment, Water</td>
<td></td>
<td>0.016</td>
<td>0.015</td>
<td>Pass</td>
</tr>
</tbody>
</table>

a Using RESRAD-BIOTA 1.8 computer code, a screening method to estimate radiological doses to aquatic and riparian biota. 
b A sum of fractions is calculated to account for the contribution to dose from each radionuclide. If the sum of fractions exceeds 1.0, then the dose guideline has been exceeded and further screening (Tier 2 or 3) is required. The sum of fractions has been rounded to two figures with a maximum of three decimal points. Maximum concentrations and the Biota Concentration Guides are presented in Appendix D. 
c The biota dose assessment requires concentration data for both sediment and water. If one of these media is not measured then it is estimated by using the default water to sediment partition coefficient. If water was measured, then sediment was estimated from water and if sediment was measured then water was estimated from sediment. In some cases where both sediment and water were measured a radionuclide was only measured in one medium (e.g., tritium in water), and the concentration for that radionuclide in the other medium was estimated. See Appendix D for details on what was measured.

Table 4-6. Estimated Doses to Biota Associated with West Lake. 

<table>
<thead>
<tr>
<th>Tier</th>
<th>Exposure Assumptions</th>
<th>Sum of Fractionsb</th>
<th>2015</th>
<th>2016</th>
<th>Pass or Fail</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Maximum Sediment, Water Concentration and Default Bioaccumulation</td>
<td>16</td>
<td>115</td>
<td></td>
<td>Fail</td>
</tr>
<tr>
<td>2</td>
<td>Average Sediment, Water Concentration and Default Bioaccumulation</td>
<td>3.7</td>
<td>41</td>
<td></td>
<td>Fail</td>
</tr>
<tr>
<td>3</td>
<td>Average Sediment, Water Concentration and Site-specific Bioaccumulation</td>
<td>0.05</td>
<td>0.49</td>
<td></td>
<td>Pass</td>
</tr>
</tbody>
</table>

a Using RESRAD-BIOTA 1.8 computer code, a screening method to estimate radiological doses to aquatic and riparian biota. 
b A sum of fractions is calculated to account for the contribution to dose from each radionuclide. If the sum of fractions exceeds 1.0, then the dose guideline has been exceeded and further screening (Tier 2 or 3) is required.

The RESRAD-BIOTA default bioaccumulation factor for uranium isotopes from water to aquatic biota is 1,000. This means that the concentration in tissues would be 1,000 times that measured in water. Site-specific data from West Lake support a much lower uranium bioaccumulation factor. Aquatic biota (only brine flies have been sampled, and they are also the most relevant organisms) and water were sampled concurrently in 2000 and 2007 (PNNL-13487, Hanford Site Environmental Report for Calendar Year 2000; DOE/RL-2007-50, Central Plateau Ecological Risk Assessment Data Package Report). The maximum concentration of any of the uranium isotopes in brine flies was 0.77 pCi/g for uranium-233/234 in 2007. The minimum uranium-233/234 water concentration was 940 pCi/L in 2007. The bioaccumulation factor is calculated by dividing the biota concentration (in pCi/g) by the water concentration (in pCi/ml); therefore, the maximum bioaccumulation factor for uranium would be less than one. A bioaccumulation factor of one was used for the Tier 3 biota dose calculation as a somewhat
protective measure of site-specific uranium uptake into the food chain. The Tier 3 biota dose
calculations resulted in sum of fractions less than one, indicating that the calculated doses were below
dose limits related to the biota concentration guides. This result was similar to those calculated for
2014, but the 2016 doses were about 10 times greater than those calculated for 2015 (Table 4-6). The
reason is that the maximum concentrations in West Lake pond water samples varied quite widely, and
isotopic uranium is typically detected in West Lake pond water. The isotopic ratios of uranium indicate a
natural source (PNL-7662). The last 3 years of concentrations were 2014 (uranium-234 at 6,580 pCi/L,
uranium-235 at 248 pCi/L, uranium-238 at 6,380 pCi/L); 2015 (uranium-234 at 1,650 pCi/L, uranium-235
at 87.1 pCi/L, uranium-238 at 1,570 pCi/L); and 2016 (uranium-234 at 10,700 pCi/L, uranium-235 at
43.5 pCi/L, uranium-238 at 13,700 pCi/L). The maximum concentration measured in 2016 was about
8 times greater than that measured in 2015. Further documentation of the West Lake biota dose
calculations, including the Tier 3 Biota Concentration Guides, is provided in Appendix D.

Biota dose calculations were implemented for terrestrial biota based on exposures to soils collected on
the Hanford Site. The RESRAD-BIOTA computer code evaluates potential effects on biota from the
maximum concentrations of radionuclides measured in near field soil samples as tabulated in
Appendix C. The radionuclides evaluated in soil are cesium-137, plutonium-238, plutonium-239/240,
strontium-90, uranium-234, uranium-235, and uranium-238. The results of 2016 screening calculations
listed in Table 4-7 show the near-field soil concentrations passed the Tier 1 screen, based on the
maximum concentration. Basically, the entire estimated dose for near-field locations is from cesium-137
(84%) and strontium-90 (16%). See PNNL-20577 for a long-term trend in soil concentrations and
associated biota doses on and off the Hanford Site.

| Table 4-7. Estimated Doses to Terrestrial Biota Associated with On- and Offsite Soila. |
|------------------------------------------|------------------------------------------|------------------------------------------|
| Location                  | Tier 1 Screen Sum of Fractionsb          | Pass or Fail                             |
|                          | 2015b                                   | 2016                                    |
| Near field               | 0.72                                    | 0.57                                    | Pass                                    |
| Far field                | 0.024                                   | Not measuredc                           | --                                      |

a Using RESRAD-BIOTA 1.8 computer code, a screening method to estimate radiological doses to aquatic and riparian
biota.
b A sum of fractions is calculated to account for the contribution to dose from each radionuclide. If the sum of fractions
exceeds 1.0, then the dose guideline has been exceeded and further screening (Tier 2 or 3) is required. The sum of
fractions has been rounded to two figures with a maximum of three decimal points. Maximum concentrations and the
Biota Concentration Guides are presented in Appendix D.
c Far field soil samples are collected approximately every 3 to 5 years and are planned for collection in 2018.

In addition to the dose assessments related to soils, sediments, and water, there are also fish and
wildlife collected from the Hanford Site and reference locations. Although none of the biota dose
assessments (except for West Lake) required any additional tiers of analyses, these supplemental
calculations characterize more realistic doses based on measured concentrations. Dose to aquatic
animals based on the maximum concentrations of strontium-90 (0.113 pCi/g), uranium-234
(0.0111 pCi/g), uranium-235 (0.00741 pCi/g), and uranium-238 (0.00693 pCi/g), in fish was
0.0001 rad/day. Internal dose to terrestrial plants based on the maximum concentrations of cesium-137
(0.13 pCi/g), plutonium-238 (0.00046 pCi/g), plutonium-239/240 (0.021 pCi/g), strontium-90 (1.3 pCi/g)
uranium-234 (0.14 pCi/g), uranium-235 (0.081 pCi/g), and uranium-238 (0.12 pCi/g), in plants was
0.002 rad/day. Dose to terrestrial animals based on the maximum concentration of strontium-90
(0.163 pCi/g) in deer bone was 0.000009 rad/day. Using the measured tissue data leads to lower doses then using the default bioaccumulation information assumed in the Tier 1 RESRAD-BIOTA calculations.

4.2.7 Radiological Dose in Perspective

The dose for the MEI in 2016 was 0.12 mrem (1.2 μSv; Section 4.2.1). The average individual dose from Hanford Site operations in 2016, based on the 50-mi (80-km) radius population exposed to air emissions and the Tri-Cities populations exposed to water pathways releases to the Columbia River, was approximately 0.0042 mrem (0.042 μSv). To place the MEI and average individual estimated doses into perspective, the estimated doses may be compared with doses received from other routinely encountered sources of radiation. The National Council on Radiation Protection and Measurement report Ionizing Radiation Exposure of the Population of the United States (NCRP 2009) estimated that the overall average exposure to ionizing radiation for the average American is 620 mrem (6,200 μSv) per year. Approximately 50% of the 620 mrem (6,200 μSv)/yr average annual dose is related to natural sources, with the remaining 50% attributable primarily to medical procedures.

The most relevant radiation sources for comparison to doses received from environmental media include natural terrestrial and cosmic background radiation, and inhalation of naturally occurring radon (Figure 4-7). Average annual individual background dose related to terrestrial radiation (19 mrem [190 μSv]), cosmic background radiation (30 mrem [300 μSv]), and radon (radon-222) and thoron (radon-220) gases (230 mrem [2,300 μSv]) are shown relative to Hanford Site operational doses in Figure 4-8. The calculated radiological doses from Hanford Site operations in 2016 were a small percentage of national average annual doses from these natural background sources. Note that annual dose is shown on a linear scale in Figure 4-8, and Hanford-related doses are too small to be observed. For example, the national annual average radiation dose from natural terrestrial sources (approximately 19 mrem [190 μSv]) is approximately 160 times larger than the 2016 Hanford Operations dose to the MEI receptor (0.12 mrem [1.2 μSv]).

Scientific studies (Health Risks from Exposure to Low Levels of Ionizing Radiation, BEIR VII Phase 2 [National Research Council 2006]) have been performed to estimate the possible risk from exposure to low levels of radiation. These studies provide information to government and scientific organizations for use in recommending radiological dose limits and standards for public and occupational safety.

Although no increase in the incidence of health effects from low doses of radiation actually has been confirmed by the scientific community, regulatory agencies cautiously assume that the probability of these types of health effects occurring due to exposure to low doses (down to zero dose) is the same per unit dose as the health effects observed after an exposure to much higher doses (e.g., in atomic bomb survivors, individuals receiving medical exposure, or, historically, painters of radium dials). This concept is known as the “linear no-threshold” hypothesis. Under these assumptions, public exposure to radiation from current Hanford Site releases, exposure to natural background radiation (which is hundreds of times greater), and exposure to very high levels of radiation each increases an individual’s probability or chance of developing a detrimental health effect (primarily cancer) proportional to the dose received.

Scientists do not fully agree on how to translate the available epidemiological data on health effects from high radiological doses into the numerical probability (risk) of detrimental effects from low radiological doses (UNSCEAR 2012, Biological Mechanisms of Radiation Actions at Low Doses). Some scientific studies have indicated that low radiological doses may result in beneficial rather than adverse
effects (Calabrese 2009). Because cancer is a common disease in the general population and may be attributable to many other causes besides radiation (e.g., genetic defects, natural and man-made chemicals, natural biochemical body reactions), some scientists doubt that the risk from low-level radiation exposure can be proven conclusively. In developing Clean Air Act regulations, EPA used a probability of approximately 4 per 10 million ($4 \times 10^{-7}$) for the risk of developing a fatal cancer after receiving a dose of 1 mrem (10 μSv; EPA 1989). Additional data support the reduction of even this small risk value, possibly to zero, for certain types of radiation when the dose is spread over an extended time (National Research Council 2006). Guidance from the Interagency Steering Committee on Radiation Standards (ISCORS 2002) recommends that agencies assign a risk factor of 6 per 10 million ($6 \times 10^{-7}$) for developing a fatal cancer after receiving a dose of 1 mrem (10 μSv).

One approach for providing perspective on calculated risks related to low-dose radiation exposures is to compare them to risks involved in other typical activities. Table 4-8 compares the estimated risks from various radiological doses to the risks of some activities encountered in everyday life.
Figure 4-8. Radiological Doses from Hanford Site Operations Compared to Annual Average from Natural Sources.

Table 4-8. Estimated Risk from Various Activities and Exposures.

<table>
<thead>
<tr>
<th>Activity or Exposure Per Year</th>
<th>Risk of Fatality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Home accidents</td>
<td>$100 \times 10^{-6}^a$</td>
</tr>
<tr>
<td>Firearms (sporting accidents)</td>
<td>$10 \times 10^{-6}^a$</td>
</tr>
<tr>
<td>Flying as an airline passenger (cross-country roundtrip – accidents)</td>
<td>$8 \times 10^{-6}^a$</td>
</tr>
<tr>
<td>Recreational boating (accidents)</td>
<td>$6 \times 10^{-6}^a$</td>
</tr>
<tr>
<td>Riding or driving 300 mi (483 km) in a passenger vehicle</td>
<td>$2 \times 10^{-6}^a$</td>
</tr>
<tr>
<td>Dose of 1 mrem (10 μSv) for 70 yrs</td>
<td>0 to $0.6 \times 10^{-6}^b$</td>
</tr>
<tr>
<td>Natural background radiological dose (310 mrem [3,100 μSv]) for 70 yrs</td>
<td>0 to $200 \times 10^{-6}^b$</td>
</tr>
<tr>
<td>Dose to hypothetical MEI (2016 rate) of 0.12 mrem (1.2 μSv)/yr living near Hanford Site for 70 yrs</td>
<td>0 to $0.1 \times 10^{-6}^b$</td>
</tr>
</tbody>
</table>

---

4.3 Radiological Clearance of Hanford Site Property

*W. Boyd*

Principal requirements for the control and clearance of DOE property containing residual radioactivity are found in DOE O 458.1. These requirements are designed to ensure the following:

- Property is evaluated; radiologically characterized; and, where appropriate, decontaminated before release
- Residual radioactivity level in property to be released is as near background levels as reasonably practicable as determined through DOE’s as low as reasonably achievable process requirements and authorized limits

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\(a\) Real actuarial values.
\(b\) Upper bound calculated using $6 \times 10^{-7}$ risk of developing a fatal cancer after receiving a 1 mrem (10 μSv) dose (ISCORS 2002).
• All property releases are appropriately certified, verified, documented, and reported; public participation needs are addressed; and processes are in place to maintain appropriate records.

4.3.1 Radiological Clearance for Potentially Contaminated Personal Property with Hard-to-Detect Radionuclides

In the process of performing environmental remediation or related support activities, Hanford Site contractors encounter a wide variety of contaminated personal property, including consumables, office items, tools and equipment, and debris. Over 19,000 items of personal property were cleared from radiological areas on the Hanford Site; however, the majority of the items did not leave the Hanford Site. The personal property items primarily consisted of small items such as flashlights, hard hats, radios, cameras, pens, pencils, respiratory protection, radiological control instruments, and industrial hygiene instruments. All of these items met DOE O 458.1 clearance criteria and, therefore, did not require additional radiological controls post-survey. In January 2000, DOE issued a moratorium prohibiting the release of volume-contaminated metals and subsequently suspended the release of metals for recycling purposes from DOE radiological areas in July 2000. As a result, no volume of contaminated metals or metals for recycling purposes were released from Hanford in 2016.

Final disposition of potentially contaminated personal property with hard-to-detect radionuclides depends on whether the property is considered radiologically contaminated, and whether the disposal of such property is subject to CERCLA requirements. Radiologically contaminated property is disposed at ERDF if subject to CERCLA requirements, and if not at the Central Waste Complex in the 200-West Area. Personal property that has contamination levels below approved DOE control and clearance guidelines (DOE O 458.1) are considered for release if the property can be reused. Hanford Site contractors routinely encounter a wide variety of radionuclide mixtures ranging from essentially pure plutonium to fission and activation products. Included in these fission and activation products are low-energy beta emitters, such as carbon-14, iron-55, nickel-59, nickel-63, selenium-79, technetium-99, palladium-107, and europium-155 that are difficult or impossible to detect with routine field-survey methods (i.e., hard-to-detect radionuclides).

Traditionally, field detectable or easy-to-detect radionuclides have been used as an analog for the entire mixture of radionuclides encountered during work activities. The control and release criteria (DOE O 458.1) have been adjusted downward to account for the portion of the activity that is not detectable by field survey methods. As the ratio of hard-to-detect radionuclides to easy-to-detect radionuclides increases, the criteria are reduced to a point where the adjusted limits are difficult or impossible to verify with field survey instruments. Decades of radioactive decay have reduced the contributions of easy-to-detect radionuclides to such low levels that current control and release methodologies are no longer sufficient for verifying that contaminant levels comply with the existing approved DOE property release guidelines in DOE O 458.1.

Accordingly, a request to DOE in May 2006 was submitted by Washington Closure Hanford (WCH) (DOE contractor for the River Corridor Closure Contract) to increase the release criteria (authorized limits) for hard-to-detect radionuclides. The requested authorized limits would apply only to beta-gamma surface contamination on potentially contaminated equipment and materials, and exclude volumetric contamination (contamination that is distributed throughout the volume of the property), contamination in or on persons, unrestricted release of metals, and alpha-surface contamination. Detailed radiological analyses were performed to demonstrate these authorized limits would be
protective of human health and the environment. Based on these analyses, the authorized limits would result in a dose of less than 1 mrem (10 mSv) in any year to the MEI and a collective dose of less than 10 person-rem (0.1 person-Sv) to any exposed population. These authorized limits (Table 4-9) were reviewed by DOE-RL and U.S. Department of Energy, Headquarters (DOE-HQ) personnel and approved for use by WCH in May 2007. In 2008, DOE-RL provided conditional approval to CH2M Plateau Remediation Company (CHPRC) and Fluor Hanford, Inc. to use these hard-to-detect authorized limits. In addition to this request, in 2013 CHPRC requested and was approved an authorized limit to apply the general beta-gamma limits to the low energy beta emitter, plutonium-241 (1,000 dpm/100 cm² removable limit and 5,000 dpm/100 cm² total contamination limit). In June 2009, Washington River Protection Solutions submitted a request to the U.S. Department of Energy, Office of River Protection (DOE-ORP) for approval to use these hard-to-detect authorized limits. DOE-ORP provided conditional approval for this request in June 2009. Mission Support Alliance submitted a request to DOE-RL in October 2009 for approval to use these hard-to-detect authorized limits. DOE-RL provided conditional approval for this request in November 2009.

Table 4-9. Approved Release Criteria (Authorized Limits) for Select Hard-to-Detect Radionuclides\(^a\) for Residual Beta-Gamma Surface Contamination.

<table>
<thead>
<tr>
<th>Average</th>
<th>Maximum</th>
<th>Removable</th>
</tr>
</thead>
<tbody>
<tr>
<td>50,000 dpm/100 cm(^2)</td>
<td>150,000 dpm/100 cm(^2)</td>
<td>10,000 dpm/100 cm(^2)</td>
</tr>
</tbody>
</table>

\(^a\) Carbon-14, iron-55, nickel-59, nickel-63, selenium-79, technetium-99, palladium-107, and europium-155

4.3.2 Granular Activated Carbon for Offsite Shipment and Regeneration Radiological Clearance

Carbon tetrachloride was found in the unconfined aquifer beneath the 200-Westest Area in the mid-1980s. Groundwater monitoring indicated the carbon tetrachloride plume was widespread and concentrations were increasing. An expedited response action was initiated in 1992 to extract carbon tetrachloride from the vadose zone in the 200-ZP-2 Operable Unit, currently designated as the 200-PW-1 Operable Unit, in the 200-Westest Area. The 200-PW-1 Operable Unit soil-vapor extraction system includes vapor-phase granular activated carbon canisters to remove carbon tetrachloride from the extracted vapors prior to discharge. This facility was in full operation by 1995.

Workers installed a groundwater pump-and-treat system in 1996 in a second operable unit (200-ZP-1 Operable Unit) to treat contaminated groundwater in the unconfined aquifer. The system includes an air-stripping unit that volatilizes carbon tetrachloride in the groundwater and then discharges the carbon tetrachloride vapors through granular activated carbon canisters that are identical to the large, carbon-steel granular activated carbon canisters in the 200-PW-1 Operable Unit soil-vapor extraction system.

Each of these systems uses granular-activated carbon canisters to capture the volatile organic compounds removed during the extraction process. When a granular-activated carbon canister has reached volatile organic compound saturation, it is removed from the system and the granular-activated carbon is prepared for shipment to an offsite facility for regeneration and reuse. Regeneration of the granular-activated carbon requires heating it in a hearth furnace to remove the captured volatile organic compounds.
Based on past Hanford Site activities, and the results of characterization sampling, this granular-activated carbon could contain residual radioactivity. Characterization sampling results were used to determine specific radionuclides of concern for this residual radioactivity. For any potential residual radioactivity, DOE O 458.1 requires that the residual radioactivity not exceed established guidelines or that radiological release criteria (i.e., authorized limits) be developed and submitted to the applicable DOE field office. Following review by DOE-RL and DOE-HQ personnel in October 2008, approved authorized limits for offsite shipment and regeneration of granular-activated carbon was approved for use by CHPRC.

In anticipation of placing the new 200-West Area Pump-and-Treat facility online, increasing the volume of spent granular-activated carbon being sent offsite, a request to modify the authorized limits was made by CHPRC and approved by DOE in October 2010 (Table 4-10). This modification to the authorized limits does not change the expected dose to the public. Approximately 98,000 lb (44,400 kg) of granular-activated carbon was shipped offsite in 2016 for regeneration.

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>Authorized Limit (pCi/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Americium-241</td>
<td>29</td>
</tr>
<tr>
<td>Carbon-14</td>
<td>3,000</td>
</tr>
<tr>
<td>Cesium-137</td>
<td>80</td>
</tr>
<tr>
<td>Cobalt-60</td>
<td>21</td>
</tr>
<tr>
<td>Europium-152</td>
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</tr>
<tr>
<td>Europium-154</td>
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</tr>
<tr>
<td>Europium-155</td>
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</tr>
<tr>
<td>Iodine-129</td>
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</tr>
<tr>
<td>Neptunium-237</td>
<td>50</td>
</tr>
<tr>
<td>Nickel-63</td>
<td>100</td>
</tr>
<tr>
<td>Plutonium-238</td>
<td>26</td>
</tr>
<tr>
<td>Plutonium-239</td>
<td>24</td>
</tr>
<tr>
<td>Plutonium-240</td>
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</tr>
<tr>
<td>Protactinium-231</td>
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<tr>
<td>Selenium-79</td>
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<tr>
<td>Strontium-90</td>
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<tr>
<td>Technetium-99</td>
<td>500</td>
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<tr>
<td>Thorium-232 plus progeny</td>
<td>6</td>
</tr>
<tr>
<td>Tritium</td>
<td>300,000</td>
</tr>
<tr>
<td>Uranium-234</td>
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</tr>
<tr>
<td>Uranium-235</td>
<td>100</td>
</tr>
<tr>
<td>Uranium-238 plus short-lived progeny</td>
<td>100</td>
</tr>
</tbody>
</table>

4.3.3 Tri-Cities Development Council Land Conveyance
There were no land conveyances in 2016.