Table of Contents

Idaho’s INL Oversight Mission ................................................................. 1
Environmental Surveillance Program ................................................. 1
  Monitoring Results ................................................................................... 2
  Trends ........................................................................................................ 3
  Comparison with DOE Data ................................................................. 3
Air Monitoring ....................................................................................... 3
  Air Monitoring Equipment and Procedures ...................................... 6
  Air Monitoring Results and Trends ................................................. 7
  Particulate Matter in Air ................................................................. 7
  Atmospheric Tritium ........................................................................... 8
  Gaseous Radioiodine .......................................................................... 8
  Precipitation ......................................................................................... 8
  Air Monitoring Verification Results .............................................. 8
  Air Monitoring Impacts and Conclusions ...................................... 9
Radiation Monitoring ........................................................................... 9
  Radiation Monitoring Equipment and Procedures ...................... 10
  Radiation Monitoring Results and Trends .................................. 10
  Radiation Monitoring Verification Results ................................ 11
  Radiation Monitoring Impacts and Conclusions ....................... 12
Water Monitoring ............................................................................... 12
  Water Monitoring Equipment and Procedures .......................... 15
  Water Monitoring Results and Trends ....................................... 18
  Radiological Analytes ...................................................................... 18
  Gross Alpha and Gross Beta Radioactivity .................................. 20
  Gamma-emitting radionuclides ..................................................... 21
  Tritium (³H) ....................................................................................... 21
  Uranium and Transuranic Isotopes ................................................ 23
  Strontium-90 (⁹⁰Sr) ....................................................................... 23
  Technetium-99 (⁹⁹Tc) ..................................................................... 27
  Non-radiological Analytes ............................................................... 29
  Chloride ............................................................................................ 31
  Sulfate ............................................................................................... 32
  Nitrate plus Nitrite .......................................................................... 32
  Chromium ......................................................................................... 32
  Manganese and Iron ....................................................................... 34
  Volatile Organic Compounds ........................................................ 35
  Water Monitoring Verification Results ...................................... 36
  Radiological ...................................................................................... 36
  Non-Radiological .............................................................................. 37
  Water Monitoring Impacts and Conclusions ............................... 38
Terrestrial Monitoring .......................................................................... 39
  Terrestrial Monitoring Equipment and Procedures .................. 39
  Terrestrial Monitoring Results and Trends ................................. 40
  Terrestrial Monitoring Verification Results ............................... 41
  Terrestrial Monitoring Impacts and Conclusions ...................... 41
Quality Assurance for the ESP .............................................................. 41
  Data Assessment Summary .......................................................... 41
Issues and Problems ................................................................. 42
Comparing Data .............................................................................. 42

**Radiological Emergency Response Planning and Preparedness** ........................................ 43
- Non-INL Radiological Activities ................................................................. 43
- Drills and Exercises ............................................................................. 44
- Waste Isolation Pilot Plant Shipment Safety .............................................. 44
- Emergency Response ............................................................................ 44
- Classes and Presentations ...................................................................... 44

**Public Outreach** ............................................................................ 45
- Publications .......................................................................................... 45
- Presentations and Events ...................................................................... 45
- Community Monitoring Network .......................................................... 49
List of Figures

Figure 1. Locations of selected DEQ monitoring sites .................................................................4
Figure 2. Off-site DEQ continuous air monitoring station. .............................................................4
Figure 3. On-site DEQ continuous air monitoring station. ............................................................5
Figure 4. DEQ air monitoring station with a radioiodine sampler, an atmospheric moisture sampler, a precipitation sampler, and two total suspended particulate (TSP) matter samplers. .......................................................................................6
Figure 5. Collecting an electret ionization chamber (EIC) and deploying a new one. ...................10
Figure 6. Locations of HPIC and EIC monitoring sites.................................................................11
Figure 7. Water quality monitoring sites distant from the INL ....................................................14
Figure 8. Water quality monitoring sites on and near the INL ....................................................15
Figure 9. Collecting ground water samples from a well site distant from the INL .......................16
Figure 10. Collecting water samples at an irrigation well. ............................................................17
Figure 11. Water sampling at a distant well in Magic Valley .......................................................17
Figure 12. Tritium (³H) concentrations (pCi/L) over time for facility wells. ...............................22
Figure 13. Tritium concentrations for DEQ sample locations in 2017 ........................................23
Figure 14. ⁹⁰Sr concentrations over time for selected wells near Test Area North (TAN) ..........25
Figure 15. ⁹⁰Sr concentrations over time for selected INL Site wells at INTEC and ATR ..........26
Figure 16. 2017 ⁹⁰Sr concentrations (pCi/L) for DEQ sample locations ....................................27
Figure 17. ⁹⁹Tc concentrations over time for wells at or downgradient of INTEC .........................28
Figure 18. 2017 ⁹⁹Tc concentrations (pCi/L) for DEQ sample locations .................................29
Figure 19. Chloride concentrations for sample location NRF-06 over time .............................31
Figure 20. 2017 chloride concentrations for DEQ sample locations .........................................32
Figure 21. Chromium concentrations (µg/L) over time for selected aquifer wells at ATR and INTEC .................................................................33
Figure 22. 2017 chromium concentrations (µg/L) for DEQ sample locations ............................34
Figure 23. TCE concentrations (µg/L) over time for selected wells located in the medial zone at TAN .................................................................36
Figure 24. DEQ soil sampling locations for 2017 .................................................................40
Figure 25. Water Awareness Poetry Contest 2017 on display at the Idaho Falls Library ..........46
Figure 26. Children enjoying “Edible Aquifer” food activity at Water Festival 2017 ...............46
Figure 27. Children preparing their rain sticks at the Water Festival event 2017 ....................47
Figure 28. Children learning about Macro Invertebrates at the Water Festival event 2017 ....47
Figure 29. Children participating in the Edible Aquifer activity at the 2017 Earth Day event ....48
Figure 30. DEQ staff handing out give-away items at the 2017 Earth Day event ....................48
Figure 31. Community monitoring station at the Fort Hall location ............................................49
List of Tables
Table 1. Gross alpha and beta screening ranges and averages observed by DEQ-INL Oversight Program for 2017. ............................................................................................................ 7
Table 2. Comparison of DEQ suspended particulate matter analysis results for paired samples with ESER and BEA results in 2017. ............................................................................................... 9
Table 3. Comparison of DEQ, ESER and BEA radiation measurements at co-located sites in 2017. (Units in micro-Roentgen per hour or µR/hr) ............................................................................................................ 12
Table 4. Summary of analytical results (pCi/L) for radiological constituents in groundwater in 2017. Surface water and wastewater results are excluded. ........................................... 19
Table 5. Summary of selected non-radiological analytical results for DEQ water samples for 2017. ........................................................................................................................................ 30
Table 6. Radiological results for co-samples collected by DOE and DEQ in 2017. ......................... 37
Table 7. Non-radiological results for co-samples collected by DOE and DEQ in 2017. ............... 38
<table>
<thead>
<tr>
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<th>Acronym</th>
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<tr>
<td>aCi/m³</td>
<td>attocuries per cubic meter</td>
<td>EPA</td>
<td>Environmental Protection Agency</td>
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<td>ARP</td>
<td>Accelerated Retrieval Project</td>
<td>ESER</td>
<td>Environmental Surveillance, Education and Research Program</td>
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<td>AMWTP</td>
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<td>fCi/m³</td>
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<td>Code of Federal Regulations</td>
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<td>Carbon Reduction Reformer</td>
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<td>IWTU</td>
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<td>LLD</td>
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<td>RPD</td>
<td>Relative Percent Difference</td>
<td>VOC</td>
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Idaho’s INL Oversight Mission

For more than half a century, the Idaho National Laboratory (INL) Site, operated by the Department of Energy (DOE) and its contractors, has been the site of research and development of nuclear technology. The work performed at INL addressed the nation’s interests in establishing nuclear reactors as a viable source of energy for civilian and military applications. Beginning in the 1950s, numerous facilities were constructed at INL to study all aspects of the nuclear fuel cycle, including fuel testing, reprocessing, and reactor prototype safety testing. The INL consequently became a site for management of spent reactor fuel (primarily from naval reactors), and radioactive and mixed wastes. Covering almost 900 square miles of the Snake River Plain and located 40 miles west of Idaho Falls, Idaho, the INL was well-suited for these activities. In the late 1980s, environmental management became a major part of the INL’s mission. DOE initiated projects to decontaminate and decommission aging facilities, remove waste, and perform environmental cleanup and restoration.

In 1989, the Idaho Legislature established an INL oversight program to provide citizens with independent information and analysis related to the INL Site. In 2007, legislation was enacted to confirm DEQ as the agency responsible for the INL Oversight Program (DEQ-INL OP), which verifies that INL Site activities are protective of public health and the environment. Our staff has expertise in radiation protection, hydrogeology, engineering, ecology, biology, computer science, education, and communications. We serve our fellow Idahoans by:

- Monitoring the environment on and around the INL Site.
- Preparing for emergencies involving radioactive materials.
- Keeping the public informed about INL Site activities.

The purpose of this report is to provide a summary of the activities performed by DEQ during 2017. The report is divided into sections covering the Environmental Surveillance Program (ESP), Radiological Emergency Response Planning and Preparedness, and Public Outreach.

Environmental Surveillance Program

DEQ provides independent environmental monitoring of the INL site for the citizens of Idaho through a multifaceted program of environmental media measurements. Measurements are made at locations on and near the INL Site, including population centers close to the INL Site boundary, and at relatively distant locations in southeast and south central Idaho. DEQ scientists use their data to evaluate public and environmental safety, and to verify monitoring of ambient environmental radiation and radioactivity in air, water, soil, and milk performed by DOE contractors. Currently, DOE funds environmental surveillance through contracts with Wastren Advantage, Inc. (WAI), the United States Geological Survey (USGS), Idaho Cleanup Project Core contractor (Fluor LLC), and the prime INL contractor, Battelle Energy Alliance, LLC (BEA). WAI conducts the Environmental Surveillance, Education and Research (ESER) program, which performs environmental surveillance outside the INL site boundary – BEA performs surveillance within the INL site.
In order to present sampling results to the public and interested agencies, DEQ publishes quarterly and annual reports. Each quarterly report contains detailed data and results of the DEQ environmental monitoring program. Annual reports summarize the quarterly data, identify general trends in the concentrations of major contaminants found in and around the INL Site, assess the impacts of DOE operations on the environment, and evaluate the reliability of DOE-contracted monitoring programs.

**Monitoring Results**

In 2017, DEQ conducted monitoring to measure environmental radiation levels and radioactivity in air, water, soil, and milk around the INL Site. Radioactivity levels found in air, soil, and milk samples were typical of background values. Tritium in groundwater was not detected at a concentration above background in the vicinity of the southern INL boundary; however, boundary sites with a history of elevated tritium concentrations were not sampled in 2017. No sites monitored by DEQ exceed federal drinking water standards for tritium. Concentrations of tritium at the INL continue to decline site-wide. No other contaminants attributable to INL Site operations were identified in groundwater samples collected outside of the INL Site.

Environmental measurements made by DEQ within the INL Site in 2017 were consistent with past results. Water samples collected from on-site locations near INL Site facilities identified concentrations of $^{90}$Sr (strontium-90), chloride, manganese, iron, and some volatile organic compounds (VOCs) greater than drinking water standards. These contaminants were found in known INL contaminant plumes and at levels consistent with historic trends for the sampling locations. These water sources are not used by the public or INL Site workers. Other contaminants from historic INL Site operations were identified in water, but at concentrations less than drinking water standards and within expected levels.

Tritium was occasionally detected in atmospheric moisture samples collected from both on-site and off-site monitoring locations. When detected these levels were less than one percent of EPA regulatory limits. Environmental measurements of radioactivity in air and direct radiation were typical of background levels at all sites. Radioactivity in the terrestrial environment and food

---

**Did You Know?**

The amount of radioactivity in the environment is measured using terms that describe how often the material undergoes radioactive decay.

A **curie** is a unit of radioactivity, symbolized as Ci, equal to $3.7 \times 10^{10}$ disintegrations or nuclear transformations per second. This is approximately the amount of radioactivity emitted by one gram (1g) of radium-226. The unit is named after Pierre Curie, a French physicist.

Fractions of curie are typically used to define small amounts of radioactivity. For example:
- **milli** - millicurie is simply one one-thousandth of a curie
- **micro** - microcurie is simply one one-millionth of a curie
- **nano** - nanocurie is simply one one-billionth of a curie
- **pico** - picocurie is simply one one-trillionth of a curie
- **femto** - femtocurie is one-quadrillionth of a curie
- **atto** - attocurie is one-quintillionth of a curie

<table>
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<th>Multiplication Factor</th>
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<td>$0.001 \times 10^{-3}$</td>
<td>milli</td>
<td>m</td>
</tr>
<tr>
<td>$0.000001 \times 10^{-6}$</td>
<td>micro</td>
<td>μ</td>
</tr>
<tr>
<td>$0.000000001 \times 10^{-9}$</td>
<td>nano</td>
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</tr>
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chain remained at background levels, based on soil and milk sampling results.

**Trends**

Results for 2017 monitoring in terrestrial media and air were generally consistent with historic trends. Radiation levels were consistent with historic background measurements. Concentrations of $^{90}$Sr, chloride, manganese, iron, and some VOCs exceeded federal drinking water standards at locations on the INL in 2017. Tritium concentration in groundwater continues to decline. Gross beta radioactivity in groundwater at all locations followed trends for $^{90}$Sr. The concentrations of some contaminants in groundwater (such as gross alpha radioactivity, $^{99}$Tc (technetium-99), and VOCs) showed trends that were not as clearly understood, possibly resulting from changes in INL operations and cleanup efforts. Tritium concentrations in atmospheric moisture remained consistent over time.

**Comparison with DOE Data**

In general, there is satisfactory agreement between the environmental monitoring data reported by DEQ and the DOE. This level of comparability between DEQ and DOE confirms that both programs present reasonable representations of the state of the environment surrounding the INL. This helps to foster public confidence in both the State’s and DOE’s monitoring programs and in the conclusions drawn from their monitoring.

In the pages that follow, the results of DEQ’s monitoring for each type of media (air, radiation, water, soil, and milk) are discussed in greater detail.

**Air Monitoring**

Continuous air monitoring is conducted at 11 locations to monitor concentrations of radionuclides in the atmosphere. These 11 locations include one air monitoring station operated by the Shoshone-Bannock Tribes at Fort Hall, Idaho.

Air monitoring locations (and selected other DEQ monitoring sites) are shown in Figure 1 and continuous air monitoring stations are pictured in Figures 2 and 3.
Figure 1. Locations of selected DEQ monitoring sites.

Figure 2. Off-site DEQ continuous air monitoring station.
Air monitoring stations are segregated into three categories:

- **On-site stations** are located within the INL boundary and include Experimental Field Station, Van Buren Avenue, Highway 20 Rest Area, and Sand Dunes/INL Gate 4.
- **Off-site stations** are located near the INL boundary and include Mud Lake, Monteview, Howe, and Atomic City.
- **Distant background stations** are located at the Craters of the Moon visitor center, Idaho Falls, and Fort Hall. Measurements at distant locations characterize the regional background conditions for comparison with conditions at on- and off-site stations.

Particulate air samples (filters) and radioactive iodine gas samples (charcoal cartridges) are collected weekly to monitor short-term radiological conditions in the environment. Atmospheric moisture is also collected continuously to measure tritium concentrations present in the air. Finally, precipitation samples are collected at six locations to monitor for tritium and gamma-emitting radionuclides that may be present in the environment. A DEQ air monitoring station with all four types of sampling equipment is pictured in **Figure 4**.
Figure 4. DEQ air monitoring station with a radioiodine sampler, an atmospheric moisture sampler, a precipitation sampler, and two total suspended particulate (TSP) matter samplers.

In order to verify results, data collected by DEQ at some air monitoring stations are directly compared to the air monitoring results obtained by the DOE and its contractors at co-located sample sites.

Air Monitoring Equipment and Procedures

Particulate matter is collected on filters using high-volume total suspended particulate (TSP) matter air samplers. The filters are collected weekly and are analyzed for gross alpha and gross beta radioactivity. Air concentrations are calculated based upon the amount of radioactivity on the filter divided by the volume of air that has passed through the filter. Quarterly composite samples of all TSP filters collected from each location are analyzed for gamma-emitting radionuclides. Yearly composite samples of all TSP filters collected from each location are analyzed via radiochemical separation for $^{90}$Sr (strontium-90), $^{241}$Am (americium-241), $^{238}$Pu (plutonium-238), and $^{239/240}$Pu (plutonium-239/240).

Radioactive iodine (radioiodine) samples are collected weekly. Samples are collected by drawing air through a canister filled with activated charcoal, using a low-volume air pump. The activated charcoal contained in the canister traps the radioiodine by adsorption onto its porous surface. Each week, canisters are collected from all 11 air monitoring stations and analyzed together as a group. If radioiodine is detected in this grouping, the canisters are individually analyzed.
Atmospheric moisture is collected by drawing air through a column filled with molecular sieve beads (a desiccant or water-absorbing material). Upon saturation with moisture, the column is removed and the beads are heated, causing them to release their stored moisture. This moisture is then condensed and collected as water and subsequently analyzed for tritium.

Precipitation samples are obtained at six locations using a collection tray that is heated during the winter months. The sample flows from the tray into a 5-gallon container that is collected at the end of each calendar quarter or whenever it is full. The precipitation samples are analyzed for tritium and for gamma-emitting nuclides.

All samples collected from DEQ’s air monitoring program are analyzed by the Idaho State University Environmental Monitoring Laboratory (ISU-EML) or its subcontractor(s). Analysis methods used are consistent with industry standards.

**Air Monitoring Results and Trends**

The following sections include monitoring results and trends for air monitoring.

**Particulate Matter in Air**

A total of 559 filters from TSP samplers were collected during 2017. The results from the analyses of off-site location samples were indistinguishable from those of on-site locations. All gross alpha and beta screening results during 2017 were less than the DEQ action levels for prompt response to elevated air screening measurements. Gross alpha/beta results are summarized in Table 1.

**Table 1.** Gross alpha and beta screening ranges and averages observed by DEQ-INL Oversight Program for 2017.

<table>
<thead>
<tr>
<th>DEQ-INL Oversight Program</th>
<th>Gross Alpha Range (fCi/m³)a</th>
<th>Gross Alpha Average (fCi/m³)</th>
<th>Gross Beta Range (fCi/m³)</th>
<th>Gross Beta Average (fCi/m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2017</td>
<td>-0.03 to 4.8</td>
<td>0.9 ± 0.2</td>
<td>3.0 to 109.8</td>
<td>26.9 ± 1.2</td>
</tr>
</tbody>
</table>

a. fCi/m³ – femto(10⁻¹⁵) curies per cubic meter

Radiochemical analysis of the annual TSP filter composite samples resulted in detection of $^{239/240}$Pu at Atomic City: $0.9 ± 1.3$ attocuries¹ per cubic meter (aCi/m³) (MDC 0.8 aCi/m³) and Monteview: $2.0 ± 1.6$ aCi/m³ (MDC 1.9 aCi/m³). These values are within the expected range due to global fallout from historic above-ground nuclear weapons testing. The reported concentration is much less than one percent of the federal regulatory limit for $^{239/240}$Pu of 2000 aCi/m³ (40 CFR 61).

Composites of filters collected using TSP samplers during the course of a calendar quarter are analyzed using gamma spectroscopy. No manmade gamma-emitting radionuclides were detected by DEQ in the quarterly composites of TSP filters.

¹ An attocurie is $10^{-18}$ curies, or 1/1000th of a femtocurie.
Atmospheric Tritium

A total of 126 atmospheric moisture samples were collected in 2017 from 11 monitoring locations and analyzed for tritium. Detectable airborne tritium concentrations are occasionally observed in the environment. The highest airborne tritium concentrations observed by DEQ on the INL in 2017 were $0.88 \pm 0.42 \text{ pCi/m}^3$ at the Experimental Field Station for the time period of August 4 through September 7, $0.73 \pm 0.59 \text{ pCi/m}^3$ at Van Buren Avenue for the time period of June 27 through July 14, $0.73 \pm 0.50 \text{ pCi/m}^3$ at the Big Lost River Rest Area station for the time period of July 18 through August 14, and $0.63 \pm 0.69 \text{ pCi/m}^3$ at the Sand Dunes station for the time period of July 20 through August 10.

All atmospheric tritium measurements for 2017 were much less than one percent of the concentration for compliance with federal regulations (40 CFR 61), 1500 pCi/m$^3$. Tritium levels were at or near background levels at all locations.

Gaseous Radioiodine

No gaseous radioiodine was detected by DEQ in 2017.

Precipitation

No tritium or manmade gamma-emitting radionuclides were detected by DEQ in precipitation samples at any location throughout the year.

Air Monitoring Verification Results

Gross alpha and beta particle results for suspended particulate matter samples from monitoring stations used by DEQ are compared with results from co-located stations operated by the Environmental Surveillance, Education and Research Program (ESER) and by Battelle Energy Alliance (BEA). As a convention, paired sample results are taken to agree if they differ from each other by no more than 20 percent of their average value, or to within 3 times the combined uncertainty of the two measurements. Agreement between 80% of the paired samples is considered to indicate overall statistical agreement of the programs being compared. Another test of agreement is to determine if the conclusions relevant to public health drawn from the results of one program differ from those drawn from the results of another program.

For 2017, over 80% of BEA’s gross alpha and beta particle results were in agreement with DEQ’s results, indicating overall statistical agreement between DEQ’s and BEA’s data sets. (Table 2). Comparisons between DEQ and ESER were not in overall statistical agreement. Variations in sampling schedule, equipment configuration and random uncertainty may contribute to observed differences. It is important to recognize that gross alpha and beta particle measurements are a screening method and do not represent quantitative measurement of specific radionuclides.

The results do agree in the important sense that all measurements from the three monitoring organizations are several orders of magnitude below the most restrictive regulatory limit for radionuclides of concern from the INL. The results from all three monitoring agencies indicate that there is no public health risk.
Table 2. Comparison of DEQ suspended particulate matter analysis results for paired samples with ESER and BEA results in 2017.
(Results are presented as percentage of samples that agree within 20 percent or 3 times the combined uncertainty.)

<table>
<thead>
<tr>
<th>Sampling Agency</th>
<th>ESER WAI&lt;sup&gt;a&lt;/sup&gt;</th>
<th>BEA&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEQ Gross Alpha Analysis</td>
<td>68 %</td>
<td>100%</td>
</tr>
<tr>
<td>DEQ Gross Beta Analysis</td>
<td>44 %</td>
<td>80 %</td>
</tr>
</tbody>
</table>

<sup>a</sup> ESER – Environmental Surveillance, Education and Research [Program], conducted by INL contractor Wastren Advantage, Inc.

<sup>b</sup> BEA – Battelle Energy Alliance, INL prime contractor during 2017.

Comparing tritium sample results among DEQ, ESER, and BEA is problematic because although sampling sites are co-located, samples are not paired or split samples. Each monitoring agency collects its tritium sample when the desiccant material becomes saturated with moisture; therefore the sampling frequency is dependent on the volume of desiccant used and the sampler flow rate resulting in differences and overlaps in sampling schedules throughout the year. Also, most of the results are near or below the MDC, where statistical uncertainties are relatively high. These factors make a direct one-to-one comparison of results not possible. However, all the results agree in that the maximum measured concentrations are about 3 orders of magnitude below the regulatory limit. Results from all three monitoring agencies indicate no public health risk.

**Air Monitoring Impacts and Conclusions**

Based upon 2017 air quality measurements, DEQ concludes that there are no discernable impacts to off-site locations as a result of INL operations. The results of screening analyses performed on particulate filters collected at boundary locations are consistent with the results obtained from background locations. Two specific radionuclide analyses of composite air samples resulted in statistical detections of a human-made radionuclide ($^{239/240}$Pu) at a concentration much less than 1% of the federal standard for members of the public (40CFR61).

Atmospheric moisture and precipitation sampling by all three agencies has occasionally shown detectable quantities of tritium in the environment; however, all detected quantities are well below federal regulatory limits and indicate no risk to public health.

Overall, DEQ and DOE contractor air monitoring results are considered to be in agreement based on (1) direct statistical comparison or, (2) because each organization’s results support the conclusion that environmental concentrations are well below regulatory limits and pose no health concerns for the citizens of Idaho.

**Radiation Monitoring**

Penetrating radiation is naturally present in the environment due to cosmic sources and naturally occurring radioactive materials in rock and soil. Human-made sources include nuclear reactor operations and the residual radioactivity present in soil from historic above-ground testing of nuclear weapons. Radiological conditions on the INL and throughout the eastern Snake River Plain are continuously monitored by DEQ. Penetrating radiation is measured at each of DEQ’s air monitoring stations, at meteorological towers maintained by the National Oceanic and
Atmospheric Administration (NOAA), along roadways that bound or cross the INL, and at background locations far from the INL (Figure 6). Co-located radiation monitoring is conducted by DEQ and DOE contractors at a number of locations. DEQ measurements at these locations are compared with the DOE contractors’ results to determine whether the data are in agreement.

**Radiation Monitoring Equipment and Procedures**

A network of 11 high-pressure ion chambers (HPICs) provides “real-time” monitoring of radiation exposure rates. One of these monitoring stations is owned by the Shoshone-Bannock Tribes at Fort Hall, Idaho. The real-time HPIC measurements are available to the public on the World Wide Web at:


DEQ also uses a network of passive electret ionization chambers (EICs) on and around the INL to measure cumulative radiation exposure over quarterly monitoring periods. The objectives of the DEQ EIC network are to identify baseline (background radiation) levels to use for comparison in the event of an upset condition (accidental release of radioactive material), assess potential dose in the ambient environment, validate dose assessment models, and to verify contractor environmental radiation data. Figure 5 shows a DEQ staff member collecting an EIC for analysis and deploying a new one.

**Figure 5. Collecting an electret ionization chamber (EIC) and deploying a new one.**

**Radiation Monitoring Results and Trends**

During the course of 2017, EIC and HPIC measurements performed at locations on the INL were similar to those at off-site monitoring locations and were consistent with expected background radiation exposure associated with cosmic, naturally occurring terrestrial, and human-made sources.
Radiation Monitoring Verification Results

DEQ uses EICs at several locations where DOE contractors monitor radiation using optically stimulated luminescent dosimeters (OSLD). Results of the contractors’ and DEQ’s measurements are used to determine the comparability of the organizations’ ambient penetrating radiation measurement programs. During 2017, 100% of BEA’s annual average OSLD dosimeters and 90% of ESER’s annual average OSLD measurements were in statistical agreement with DEQ’s measurements at co-located EIC sites (Table 3), meeting the program’s objectives.
Table 3. Comparison of DEQ, ESER and BEA radiation measurements at co-located sites in 2017. (Units in micro-Roentgen per hour or µR/hr)

<table>
<thead>
<tr>
<th>Statistical Measure</th>
<th>DEQ (µR/hr)</th>
<th>ESER(^a) (µR/hr)</th>
<th>DEQ(^b) (µR/hr)</th>
<th>BEA(^b) (µR/hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>12.3</td>
<td>13.1</td>
<td>11.7</td>
<td>13.7</td>
</tr>
<tr>
<td>Median</td>
<td>12.0</td>
<td>12.9</td>
<td>12.3</td>
<td>14.4</td>
</tr>
<tr>
<td>Standard Deviation</td>
<td>1.2</td>
<td>1.2</td>
<td>1.9</td>
<td>2.4</td>
</tr>
<tr>
<td>Minimum</td>
<td>11.1</td>
<td>11.7</td>
<td>6.4</td>
<td>7.1</td>
</tr>
<tr>
<td>Maximum</td>
<td>15.1</td>
<td>16.0</td>
<td>13.4</td>
<td>15.8</td>
</tr>
<tr>
<td>Average % difference</td>
<td>-6%</td>
<td></td>
<td>-15%</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) ESER – Environmental Surveillance, Education and Research [Program], conducted by INL contractor Wastren Advantage, Inc.

\(^b\) BEA – Battelle Energy Alliance, INL prime contractor during 2017.

\(^c\) Each organization’s dataset is reviewed to ensure that it supports a valid test of comparability of measurements.

Radiation Monitoring Impacts and Conclusions

Based upon radiation measurements made by DEQ, there were no discernable impacts from INL operations in 2017. Measurements on the INL are comparable to those at background locations. Quarterly averaged HPIC and EIC exposure measurements during 2017 met DEQ’s criterion for agreement. The results from all three monitoring agencies indicate no public health risk from environmental ambient penetrating radiation from both natural and human-made sources.

**Water Monitoring**

During 2017, the DEQ collected water samples from 62 sites within and downgradient of the INL.\(^2\) Water samples are used to identify INL-related impacts to the eastern Snake River Plain aquifer (ESRPA), evaluate trends of known INL contaminants and other general groundwater quality indicators, and verify DOE and USGS monitoring results. Samples are collected from groundwater, surface water, and wastewater, with the vast majority being from groundwater. Most sites sampled by DEQ are sampled concurrently (“co-sampled”) with a DOE contractor (Fluor, BEA, or WAI) or the USGS. DEQ annually compares its own analytical results with those obtained by co-samplers to evaluate consistency.

Of the sites sampled in 2017, 60 are groundwater locations (wells and springs), one is a surface water location (stream), and one is a wastewater location (Figures 7 and 8). Each water-monitoring site is categorized as upgradient, facility, boundary, distant, surface water, or wastewater. Upgradient sites are situated north and northeast of INL facilities and have not been affected by INL operations. Facility sites are near facility complexes within the INL, including the Idaho Nuclear Technology and Engineering Center (INTEC), the Advanced Test Reactor Complex (ATR), Test Area North (TAN), the Radioactive Waste Management Complex (RWMC), the Central Facilities Area (CFA), and the Naval Reactors Facility (NRF). Many facility sites are in areas of known contamination and are sampled to monitor trends of specific contaminants.

\(^2\) In most years, 80+ locations are sampled. Fewer locations were sampled in 2017 due to staff turnover.
contaminants. Boundary sites are on or near the southern boundary of the INL, downgradient of potential sources of INL contamination. Distant sites are farther downgradient of the INL, primarily in the Magic Valley, and include wells and springs used for agricultural, municipal, domestic, and industrial purposes. Surface water and wastewater samples are collected from locations within and upgradient of the INL. All wells sampled in 2017 penetrate the aquifer except for eight shallow wells at ATR that sample groundwater from a perched zone overlying the aquifer.

Samples collected from water-monitoring sites are analyzed for radiological and non-radiological constituents, many of which are present in the aquifer both naturally and as a result of INL operations. All locations are sampled for gross alpha and gross beta radioactivity, gamma-emitting radionuclides, tritium, common ions, trace metals, and nutrients. Selected sites are also sampled for specific radionuclides—including uranium isotopes ($^{234}$U, $^{235}$U, and $^{238}$U), plutonium isotopes ($^{238}$Pu, $^{239/240}$Pu), americium-241 ($^{241}$Am), strontium-90 ($^{90}$Sr), and technetium-99 ($^{99}$Tc)—and/or volatile organic compounds (VOCs) based on past and present INL operations or a history of elevated concentrations. If unexpected levels of radioactivity are detected in gross measurements, additional samples may be collected and analyzed for specific radionuclides.
Figure 7. Water quality monitoring sites distant from the INL.
Most groundwater samples were collected from wells equipped with submersible pumps. Wells fitted with Westbay™ multilevel sampling systems (referred to in prior reports as “Westbay wells”) were not sampled in 2017. Groundwater samples from springs and the surface water sample were collected as grab samples from the water source. The wastewater sample was collected as a 24-hour composite prepared by BEA.

Wells co-sampled with Fluor or the USGS, including all facility and boundary wells and most upgradient wells, were purged and stabilized by the co-sampler according to stabilization criteria published in the co-samplers’ field sample plans. Wells sampled independently by DEQ or co-sampled with WAI, including all distant wells and one upgradient well, were purged and stabilized by DEQ. See Figures 9, 10, and 11. DEQ considers a well to be stable when three consecutive water-quality readings at 3-to-5-minute intervals are within 0.1 units for pH, 3% for specific conductance, 3% for temperature, and 10% for dissolved oxygen.

Water samples were collected, handled, and preserved using standard DEQ sampling procedures. Trace metals and nutrients samples were filtered, and samples for gross alpha, gross beta,
Gamma-emitting nuclides, uranium isotopes, plutonium isotopes, americium-241, strontium-90, trace metals, and nutrients were preserved with acid immediately after sample collection. VOCs were collected in vials already containing acid.

Radiological analyses were performed by ISU-EML or its subcontractor(s). Non-radiological analyses were conducted by the Idaho Bureau of Laboratories in Boise. Common ions analyzed are calcium, magnesium, sodium, potassium, chloride, fluoride, sulfate, and total alkalinity. Nutrients analyzed are total nitrate plus nitrite and total phosphorus. Trace metals analyzed are arsenic, barium, chromium, iron, manganese, lead, selenium, and zinc. Samples from selected sites were also analyzed for volatile organic compounds (VOCs).

Laboratory methods used for all analyses were consistent with industry standards for drinking water samples.

Figure 9. Collecting ground water samples from a well site distant from the INL.
Figure 10. Collecting water samples at an irrigation well.

Figure 11. Water sampling at a distant well in Magic Valley.
Water Monitoring Results and Trends

A summary of analyte concentrations measured at upgradient, facility, boundary, and distant monitoring sites is presented below. Analytical results from several sample locations with histories of high concentrations are examined more closely to identify current trends. Results for all environmental surveillance samples collected by DEQ are available in quarterly data reports on the DEQ website http://www.deq.idaho.gov/inl-oversight/monitoring/reports.aspx.

Radiological Analytes

DEQ samples all water monitoring locations for gross alpha and gross beta radioactivity, gamma-emitting radionuclides, and tritium. Selected locations are also sampled for specific radionuclides. Concentrations of radiological analytes measured in 2017 were generally consistent with those measured in previous years. Results are summarized in Table 4. Significant findings for each radiological analyte are discussed below.
### Table 4. Summary of analytical results (pCi/L) for radiological constituents in groundwater in 2017. Surface water and wastewater results are excluded.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Upgradient</th>
<th>Facility</th>
<th>Boundary</th>
<th>Distant</th>
<th>Background¹</th>
<th>Drinking Water Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min</td>
<td>Med</td>
<td>Max</td>
<td>Min</td>
<td>Med</td>
<td>Max</td>
</tr>
<tr>
<td>Gross Alpha</td>
<td>&lt;MDC</td>
<td>1.7</td>
<td>2.8 ± 1.4</td>
<td>&lt;MDC</td>
<td>2.6</td>
<td>10.8 ± 2.3</td>
</tr>
<tr>
<td>Gross Beta²</td>
<td>1.5 ± 0.8</td>
<td>3.5</td>
<td>7.4 ± 1.1</td>
<td>2.1 ± 1.0</td>
<td>8.6</td>
<td>1117 ± 15</td>
</tr>
<tr>
<td>¹³⁷Cs</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>17.0 ± 2.6</td>
</tr>
<tr>
<td>Tritium³</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>715</td>
<td>3170 ± 260</td>
</tr>
<tr>
<td>²³⁴U</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>1.37 ± 0.34</td>
<td>1.65</td>
<td>8.5 ± 1.5</td>
</tr>
<tr>
<td>²³⁶U</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>&lt;MDC</td>
<td>0.09</td>
<td>0.44 ± 0.16</td>
</tr>
<tr>
<td>²³⁸U</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>0.59 ± 0.19</td>
<td>0.82</td>
<td>1.26 ± 0.30</td>
</tr>
<tr>
<td>²³⁹,²⁴⁰Pu</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
</tr>
<tr>
<td>²³⁹Pu</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
</tr>
<tr>
<td>²⁴¹Am</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
<td>&lt;MDC</td>
</tr>
<tr>
<td>⁹⁰Sr</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>&lt;MDC</td>
<td>0.44</td>
<td>398 ± 94</td>
</tr>
<tr>
<td>⁹⁹Tc</td>
<td>0.4 ± 0.2</td>
<td>0.4</td>
<td>1.4 ± 0.2</td>
<td>1.0 ± 0.2</td>
<td>2.95</td>
<td>278.3 ± 1.5</td>
</tr>
</tbody>
</table>

Uncertainties are reported at 2σ. Abbreviations: pCi/L, picocuries per liter; MDC, minimum detectable concentration, NS, not sampled.

¹ Background levels for gross alpha, gross beta, and ¹³⁷Cs are derived from over 20 years of DEQ groundwater monitoring in the ESRPA. Background levels for ³⁵H, uranium isotopes, plutonium isotopes, ⁹⁰Sr, and ⁹⁹Tc are based on the minimum and median values reported for western tributary water in Table 1 in Bartholomay and Hall (2016; DOE/ID 22237). Background concentrations depend on local geology and proximity to surface water recharge locations. Concentrations for sites not influenced by INL activities may still be higher than the given background ranges.

² Gross beta as ¹³⁷Cs.

³ Results for tritium are from the standard analysis method, with an MDC of approximately 130 pCi/L.

⁴ The federal drinking water standard is expressed as a cumulative annual dose of 4 millirem/year. This value was converted to a specific concentration (pCi/L) for each analyte.
Gross Alpha and Gross Beta Radioactivity

Radionuclide contributors of alpha and beta activity are present in the aquifer naturally and as a result of past INL operations. Gross alpha and gross beta analyses measure radioactivity contributed by all alpha and beta emitters in a water sample (excluding radon and tritium). These analyses are used for screening purposes only and do not yield quantitative measurements of specific radionuclides.

The primary natural sources of alpha radioactivity in groundwater and surface water are uranium and thorium, and the primary natural sources of beta radioactivity are potassium-40 and beta-emitting daughter products of naturally occurring uranium and thorium. All of these nuclides are present in the bedrock and sediments of the eastern Snake River Plain at low concentrations, and their presence in groundwater contributes to a low but measureable level of radioactivity in the aquifer, defined as background. Background concentrations of gross alpha and gross beta radioactivity, derived from over 20 years of DEQ data collected from ESRPA locations not affected by INL activities, are given as ranges in Table 4.

Gross alpha levels observed at most locations in 2017 were within the background range defined by DEQ and can be attributed to natural sources. Levels slightly above background were measured in the aquifer at TAN, CFA, and distant well MV-43, and in perched groundwater at ATR. The highest gross alpha concentration measured was 10.8 ± 2.3 pCi/L at ATR well PW-9. The EPA maximum contaminant level (MCL) is 15 pCi/L.

Gross beta levels exceeded background at several facility locations and one distant location (MV-43). Most exceedances were at TAN, where the highest concentration was 1117 ± 15 pCi/L at TAN-37A (‘A’ denotes the shallowest sampling depth, 240 feet below the surface, in well TAN-37), and at INTEC, where the highest concentration was 180 ± 3 pCi/L at ICPP-2020. The high level of gross beta activity at TAN is due to a high 90Sr concentration, discussed below. The MCL for beta activity is 4 mrem/year, which is equivalent to 8 pCi/L if the source is 90Sr, 900 pCi/L if 99Tc, and 20,000 pCi/L if tritium.

The above-background concentrations of gross alpha (8.6 ± 2.9 pCi/L) and gross beta (11.4 ± 2.4 pCi/L) radioactivity measured at distant location MV-43 are best explained by elevated concentrations of solutes in this well (see tables 16, 17, and 18 in DEQ’s “Environmental Surveillance Program Quarterly Data Report, October - December 2017”). MV-43 is in an irrigated agricultural area, and solute concentrations are high probably due to evaporation of irrigation waters prior to aquifer recharge. Alpha-emitting uranium and thorium and their beta-emitting daughter products were not measured at this location, but their concentrations are likely elevated proportionally to other solutes.
**Gamma-emitting radionuclides**

The only gamma-emitting radionuclide reported in 2017 is cesium-137. $^{137}$Cs has historically been detected at low concentrations at TAN and INTEC. In 2017, $^{137}$Cs was detected at a single location: TAN-37A ($17 \pm 3$ pCi/L). Prior measurements of $^{137}$Cs at TAN-37A from 2007 to 2016 ranged from $3.6 \pm 2.0$ pCi/L to $11.7 \pm 2.3$ pCi/L, with the previous maximum value reported in 2014. The MCL for $^{137}$Cs is 200 pCi/L.

**Tritium ($^3$H)**

Tritium in ESRPA groundwater comes from natural sources, twentieth-century nuclear weapons tests (referred to below as “bomb-pulse tritium”), and past INL waste disposal practices. Natural tritium, produced primarily by the interaction of atmospheric nitrogen with cosmic rays, and bomb-pulse tritium are incorporated in groundwater through surface recharge, resulting in a low background concentration of tritium in young groundwater that decreases with residence time in the aquifer. Groundwater locations close to areas of surface recharge may have background tritium concentrations that are higher than the given background range, whereas groundwater that is distant from surface recharge, such as near the center of the eastern Snake River Plain, are more likely to have background tritium concentrations near zero. As surface-water concentrations of bomb-pulse tritium decrease over time due to radioactive decay, the upper end of the background range decreases. The range of background concentrations of tritium typically observed in the ESRPA is given in Table 4.

Tritium was introduced to the aquifer at concentrations well above the background range by past INL waste disposal practices, including the use of wastewater injection wells and percolation ponds at ATR, INTEC, and TAN (DOE/ID-22242). Tritium concentrations once exceeded the MCL of 20,000 pCi/L at some wells in these areas; however, over the past two decades, concentrations have declined significantly due to radioactive decay and dilution.

In 2017, elevated tritium concentrations were measured in facility wells at ATR, INTEC, TAN, CFA, and RWMC, consistent with previous years. The highest concentration measured in the aquifer at each of these facility complexes was:

- **ATR** $\rightarrow$ $2500 \pm 240$ pCi/L at USGS-065
- **CFA** $\rightarrow$ $3170 \pm 260$ pCi/L at CFA-1
- **INTEC** $\rightarrow$ $2400 \pm 260$ pCi/L at USGS-067
- **TAN** $\rightarrow$ $1330 \pm 210$ pCi/L at TAN-29
- **RWMC** $\rightarrow$ $530 \pm 180$ pCi/L at USGS-087

Tritium was also detected well above background levels in perched groundwater at ATR, with a maximum concentration of $2940 \pm 290$ pCi/L. **Figure 12** shows tritium trends for selected wells at ATR, INTEC, and RWMC. Overall, tritium concentrations in facility wells in 2017 were consistent with previous years and continue to decline gradually.

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3 Well TRA-07, which typically has the highest aquifer tritium concentration at ATR, was not sampled in 2017. TRA-07 will be sampled in 2018.
Tritium was not detected above background in any boundary well. In previous years, tritium has been detected in the vicinity of the southern INL boundary at concentrations sometimes exceeding 1,000 pCi/L. The absence of detections in 2017 is because 1) Westbay wells, in which the highest boundary-area concentrations of tritium have been detected in past years, were not sampled in 2017, and 2) enhanced tritium analyses, for which the detection limit is an order of magnitude lower than the standard tritium analyses, have not yet been completed by the lab for most boundary wells sampled in 2017. Westbay wells will again be sampled in the second quarter of 2018, and the results of enhanced tritium analyses for 2017 samples will be reported in future quarterly monitoring reports as they become available.

Tritium concentrations were within the background range or below lab detection limits for all distant, upgradient, surface water, and wastewater locations.

**Figure 13** shows a concentration map of all tritium measurements in 2017.

![Figure 13](image)

**Figure 12.** Tritium ($^3$H) concentrations (pCi/L) over time for facility wells.
Uranium and Transuranic Isotopes

Selected locations at TAN, ATR, and INTEC were sampled for uranium and plutonium isotopes and $^{241}$Am in 2017. Plutonium isotopes and $^{241}$Am were not detected at any location. Uranium isotopes ($^{234}$U, $^{235}$U, and $^{238}$U) were detected in the aquifer at concentrations above background in eight wells: four at INTEC, three at TAN and one at ATR. The highest concentrations of all three uranium isotopes were found at TAN-28 (see maximum concentrations in Table 4). Elevated uranium concentrations in the groundwater at TAN, ATR, and INTEC have been previously identified and are attributed to past waste disposal practices.

Strontium-90 ($^{90}$Sr)

Strontium-90 is one of the two main sources of above-background levels of gross beta radioactivity (the other is $^{99}$Tc, discussed in the next section). Past waste disposal practices and spills have resulted in elevated concentrations of $^{90}$Sr in the aquifer at TAN and INTEC and in
perched groundwater at INTEC and ATR. Concentrations of $^{90}\text{Sr}$ above the MCL of 8 pCi/L are typically measured in wells at each of these facility complexes.

In 2017, the highest $^{90}\text{Sr}$ concentrations continue to be measured at and near TAN, with a maximum concentration of $398 \pm 94$ pCi/L at TAN-37A. $^{90}\text{Sr}$ concentrations over time for all TAN wells sampled by DEQ are shown in Figure 14. The concentration of $^{90}\text{Sr}$ in the groundwater at TAN is directly affected by in situ bioremediation (ISB) treatment of the TAN TCE plume (see discussion in the “Volatile Organic Compounds” below). Injections of whey and sodium lactate into the aquifer from 1999 to 2012 increased calcium and magnesium concentrations in the groundwater, resulting in increased competition for adsorption sites on aquifer minerals and consequent displacement of strontium cations into the groundwater. Injections were stopped in 2012 to allow redox conditions in the aquifer to return to their pre-treatment state. Theoretically, $^{90}\text{Sr}$ concentrations should decrease as added calcium and magnesium cations diffuse and disperse; this may explain the $^{90}\text{Sr}$ trend at TAN-37A, where Sr-90 concentrations have been consistently lower post-2012 than they were pre-2012 (see Figure 14). In January 2016, injections of an oil-based amendment were started at TAN-2272 to treat a residual TCE source in the vicinity of TAN-28 (downgradient of TAN-2272). Injections at this location may affect $^{90}\text{Sr}$ concentrations at nearby well TAN-2271 and downgradient wells TAN-37A, TAN-28, and possibly TAN-29 over the coming years.

$^{90}\text{Sr}$ concentrations above the MCL were also measured in the aquifer at INTEC and in perched groundwater at ATR. Figure 15 shows $^{90}\text{Sr}$ concentrations over time for aquifer wells at INTEC (USGS-047, USGS-067, ICPP-2020, USGS-085, USGS-112) and a perched groundwater well at ATR (USGS-055). The highest concentration measured in the aquifer at INTEC was $16 \pm 4$ pCi/L at USGS-047. The highest concentration measured in perched groundwater at ATR was $29 \pm 7$ pCi/L at USGS-055. No $^{90}\text{Sr}$ was detected in the aquifer at ATR. All concentrations measured in 2017 were consistent with previous years.

A concentration map of all locations sampled for $^{90}\text{Sr}$ in 2017 is shown in Figure 16.

---

4 The Idaho Cleanup Project contractor has detected high concentrations of $^{90}\text{Sr}$ in perched groundwater overlying the aquifer at INTEC. DEQ does not currently sample perched groundwater at INTEC.

5 TAN-2271 and TAN-2272, which had the maximum $^{90}\text{Sr}$ concentrations in 2015 and 2016, were not sampled in 2017.
Figure 14. $^{90}$Sr concentrations over time for selected wells near Test Area North (TAN).
Figure 15. $^{90}$Sr concentrations over time for selected INL Site wells at INTEC and ATR.
Figure 16. 2017 $^{90}$Sr concentrations (pCi/L) for DEQ sample locations.

Technetium-99 ($^{99}$Tc)

$^{99}$Tc has been introduced to the aquifer by leaks and spills at INTEC, including the inadvertent release of 18,600 gallons of sodium-bearing waste during a transfer between underground storage tanks at INTEC in 1972. Figure 17 shows $^{99}$Tc concentrations over time for selected INL wells located at or downgradient of INTEC. In 2017, all $^{99}$Tc detections remained well below the MCL of 900 pCi/L. The highest concentrations measured by DEQ in 2017 continue to be at USGS-052 (278.3 ± 1.5 pCi/L), ICPP-2020 (236.8 ± 1.4 pCi/L), and USGS-067 (110.5 ± 1.0 pCi/L). All other $^{99}$Tc detections in 2017 were below 10 pCi/L and were consistent with measurements in previous years. Figure 18 shows a concentration map of all $^{99}$Tc sample locations in 2017.
Figure 17. $^{99}$Tc concentrations over time for wells at or downgradient of INTEC.
Figure 18. 2017 $^{99}$Tc concentrations (pCi/L) for DEQ sample locations.

**Non-radiological Analytes**

DEQ samples all water monitoring locations for common ions, nutrients, and dissolved trace metals. Selected locations are also sampled for VOCs. Elevated concentrations of these constituents are present in the groundwater at some locations as a result of past INL waste disposal practices.

Concentrations of non-radiological analytes measured in 2017 were generally consistent with those measured in previous years. Results are summarized in Table 5. Analytes that exceeded drinking water standards in 2017 or in the recent past, which include chloride, sulfate, nitrate plus nitrite, chromium, manganese, iron, and certain VOCs, are discussed in greater detail below.
Table 5. Summary of selected non-radiological analytical results for DEQ water samples for 2017.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Upgradient</th>
<th>Facility</th>
<th>Boundary</th>
<th>Distant</th>
<th>Background&lt;sup&gt;1&lt;/sup&gt;</th>
<th>Drinking Water Standard&lt;sup&gt;2&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min</td>
<td>Median</td>
<td>Max</td>
<td>Min</td>
<td>Median</td>
<td>Max</td>
</tr>
<tr>
<td>Common Ions (mg/L)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alkalinity (as CaCO&lt;sub&gt;3&lt;/sub&gt;)</td>
<td>92</td>
<td>140</td>
<td>163</td>
<td>111</td>
<td>146</td>
<td>587</td>
</tr>
<tr>
<td>Calcium</td>
<td>8.9</td>
<td>40</td>
<td>50</td>
<td>34</td>
<td>63</td>
<td>130</td>
</tr>
<tr>
<td>Chloride</td>
<td>4.97</td>
<td>9.56</td>
<td>49.3</td>
<td>12.7</td>
<td>21.6</td>
<td>462</td>
</tr>
<tr>
<td>Fluoride</td>
<td>&lt;DL&lt;sup&gt;3&lt;/sup&gt;</td>
<td>0.502</td>
<td>0.701</td>
<td>&lt;DL</td>
<td>0.223</td>
<td>0.740</td>
</tr>
<tr>
<td>Magnesium</td>
<td>2.8</td>
<td>16</td>
<td>18</td>
<td>12</td>
<td>18</td>
<td>55</td>
</tr>
<tr>
<td>Potassium</td>
<td>1.4</td>
<td>3.0</td>
<td>5.9</td>
<td>2.0</td>
<td>2.9</td>
<td>12</td>
</tr>
<tr>
<td>Sodium</td>
<td>8.4</td>
<td>14</td>
<td>31</td>
<td>8.8</td>
<td>16</td>
<td>180</td>
</tr>
<tr>
<td>Sulfate</td>
<td>8.51</td>
<td>24.1</td>
<td>40.5</td>
<td>17.4</td>
<td>40.4</td>
<td>200</td>
</tr>
<tr>
<td>Nutrients (mg/L)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Nitrate plus Nitrite</td>
<td>0.50</td>
<td>0.76</td>
<td>2.5</td>
<td>0.01</td>
<td>1.8</td>
<td>6.5</td>
</tr>
<tr>
<td>Total Phosphorus</td>
<td>0.0073</td>
<td>0.014</td>
<td>0.018</td>
<td>0.005</td>
<td>0.029</td>
<td>0.70</td>
</tr>
<tr>
<td>Trace Metals (μg/L)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arsenic</td>
<td>&lt;DL</td>
<td>2.3</td>
<td>4.2</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>9.5</td>
</tr>
<tr>
<td>Barium</td>
<td>45</td>
<td>71</td>
<td>85</td>
<td>27</td>
<td>80</td>
<td>460</td>
</tr>
<tr>
<td>Chromium</td>
<td>1.6</td>
<td>3.5</td>
<td>5.5</td>
<td>&lt;DL</td>
<td>9.5</td>
<td>79</td>
</tr>
<tr>
<td>Iron</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>43</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>5600</td>
</tr>
<tr>
<td>Lead</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>1.4</td>
</tr>
<tr>
<td>Manganese</td>
<td>&lt;DL</td>
<td>2.2</td>
<td>8.0</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>1600</td>
</tr>
<tr>
<td>Selenium</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>2.9</td>
</tr>
<tr>
<td>Zinc</td>
<td>&lt;DL</td>
<td>10</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>&lt;DL</td>
<td>10</td>
</tr>
</tbody>
</table>

<sup>1</sup> Background concentrations depend on local geology. Concentrations for sites not influenced by INL activities may still be higher than the given background ranges. Sources for background ranges are: <sup>1</sup> Bartholomay and Hall, 2016 (DOE/ID-22237); <sup>2</sup> Knobel and others, 1999 (DOE/ID-22164); <sup>3</sup> Knobel and others, 1992; <sup>4</sup> DEQ data compiled from distant, boundary, and surface water sites in previous years.

<sup>2</sup> Primary standard (MCL) unless otherwise noted. National Primary Drinking Water Regulations are legally enforceable standards that apply to public water systems. Maximum Contaminant Levels (MCLs) are the highest level of a contaminant that is allowed in the drinking water. Secondary standards (SMCLs) are designated with *. Secondary Drinking Water Regulations are non-enforceable guidelines regulating contaminants that may cause aesthetic effects (such as taste, odor, or color) in drinking water. EPA recommends but does not require that water systems comply with SMCLs.

<sup>3</sup> DL=Detection Limit.
Chloride

Chloride has been introduced to the ESRPA at the INL by the discharge of wastewater to the aquifer through injection wells and infiltration ponds. The primary source of chloride in INL wastewater includes the use of sodium chloride to regenerate water softeners. TAN, INTEC, ATR, CFA, and NRF all have wells with chloride concentrations above background. Only one well monitored by DEQ, NRF-06 at NRF, has had chloride concentrations above the secondary maximum contaminant level (SMCL) of 250 mg/L. NRF-06 is located near the NRF industrial waste ditch, in which wastewater from water softeners is discharged. The chloride concentration measured in NRF-06 in 2017 was 462 mg/L, consistent with concentrations measured at this location since 2004 (Figure 19).

Most distant wells in the Magic Valley also had elevated chloride concentrations in 2017, with a maximum concentration of 79.5 mg/L at MV-43. High chloride concentrations in the Magic Valley are most likely tied to agriculture in the region, as salts in irrigation waters are concentrated by evaporation prior to recharging the aquifer.

A chloride concentration map for all 2017 sample locations is shown in Figure 20.

![Chloride Concentration Map](image)

Figure 19. Chloride concentrations for sample location NRF-06 over time.
Sulfate

No location sampled in 2017 had sulfate above the SMCL of 250 mg/L. The maximum sulfate concentration measured was 200 mg/L in USGS-062, a perched groundwater well at ATR.

Nitrate plus Nitrite

In 2017, one well exceeded the MCL of 10 mg/L for nitrate plus nitrite: MV-43 had a nitrate plus nitrite concentration of 35 mg/L. MV-43 is an irrigation well in the Magic Valley. The high nitrogen concentration is almost certainly related to the widespread use of fertilizers in this area. No wells at or near the INL exceeded the MCL for nitrate plus nitrite in 2017.

Chromium

Chromium was used to prevent corrosion in industrial water systems at the INL until the early 1970s. Disposal practices at that time allowed chromium-contaminated water to percolate down to groundwater from injection wells, open disposal ponds, and ditches, resulting in elevated chromium concentrations in some monitoring wells. In 2017, chromium concentrations were below the MCL of 100 µg/L at all locations sampled by DEQ, with a maximum concentration of 79 µg/L in ATR aquifer well USGS-065.
Results for wells that have historically had high levels of chromium are shown in \textbf{Figure 21}. A concentration map for all locations sampled in 2017 is shown in \textbf{Figure 22}.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{chromium_concentrations.png}
\caption{Chromium concentrations (µg/L) over time for selected aquifer wells at ATR and INTEC.}
\end{figure}
Manganese and Iron

Three wells, all at TAN, exceeded the SMCL for manganese (50 μg/L) during the 2017 sample season. The maximum manganese concentration measured was 1600 μg/L at TAN-37A. Manganese concentrations in TAN-28 (1200 μg/L) and TAN-29 (180 μg/L) increased from respective concentrations of 310 μg/L and 34 μg/L in 2016. PW-9, a perched-groundwater well at ATR that had a manganese concentration of 150 μg/L in 2016, had only 26 μg/L of manganese in 2017.

Well TAN-37A also exceeded the SMCL for iron (300 μg/L) in 2017, with a concentration of 5,600 μg/L. No other location had an iron concentration at or near the SMCL.

Elevated concentrations of manganese and iron in the groundwater at TAN are consistent with reducing conditions created by in-situ bioremediation (ISB) as part of the clean-up action for volatile organic compounds (see next section). ISB injections were restarted at a new location, TAN-2272, in January 2016. Manganese and iron concentrations are expected to increase and/or remain high at locations near and downgradient of TAN-2272—including TAN-2271, TAN-37A, TAN-28, and TAN-29—as long as injections continue. Concentrations of manganese and iron in TAN-37A, TAN-28, and TAN-29 all increased from 2016 to 2017. TAN-2272 and TAN-2271 were not sampled in 2017.
**Volatile Organic Compounds**

The primary volatile organic compound (VOC) contamination at the INL is located at and downgradient of TAN, where a plume originating at a former wastewater injection well extends to the east and south. The plume is characterized by high concentrations of trichloroethene (TCE) and its degradation products (cis-1,2-dichloroethene [cis-DCE], trans-1,2-dichloroethene [trans-DCE], and vinyl chloride [VC]) and lower concentrations of tetrachloroethene (PCE). The plume has been divided into three regions based on TCE concentrations reported in 1997 (INEEL/EXT-97-00931), and a different remediation strategy was chosen for each region in a 2001 Record of Decision Amendment (DOE/ID-10139):

- The hot spot (>20,000 μg/L TCE) covers a small area immediately surrounding the former injection well. The remediation strategy here has been in situ bioremediation (ISB), which involved repeated injection of a carbon source (whey and sodium lactate) into the aquifer to promote anaerobic reduction of chlorinated ethenes in the aquifer. Injections began in 1999 and were halted in 2012.
- The medial zone (1,000 to 20,000 μg/L TCE) extends about 1500 feet east-southeast from the hot spot as a narrow lobe. The remediation strategy here is to pump, treat, and reinject groundwater.
- The distal zone (5 to 1,000 μg/L TCE) surrounds the medial zone as a much larger lobe that extends about 900 feet west and 1.7 miles southeast of the hot spot. The remediation strategy here is monitored natural attenuation.

In July 2012, ISB injections were suspended indefinitely in order to initiate the rebound test—a multi-year pause in ISB treatment to evaluate residual VOC contamination in the aquifer once background groundwater conditions returned. In January 2016, ISB injections commenced at TAN-2272, a new well installed in 2015, to treat an apparent residual TCE source in the vicinity of TAN-28. A partial ISB rebound test continues in the vicinity of the original hot spot.

In 2017, DEQ sampled three wells in the medial zone east of the pre-2012 ISB treatment area (TAN-28, TAN-29, TAN-37A). Four VOCs were detected at concentrations above the MCL in TAN wells: TCE (MCL = 5 μg/L) at TAN-28, TAN-29, and TAN-37A; PCE (MCL = 5 μg/L) at TAN-29; cis-DCE (MCL = 70 μg/L) at TAN-29; and VC (MCL = 2 μg/L) at TAN-28, TAN-29, and TAN-37A. **Figure 23** shows TCE concentration trends for TAN-28, TAN-29, and TAN-37A. TCE concentrations in TAN-28 and TAN-29 have varied widely over time, probably as a result of intermittent changes in groundwater chemistry due to ISB injections as well as seasonal changes in groundwater flow (DOE/ID-11444), but clearly remained high throughout the rebound test. In 2017, TCE concentrations in both wells (308 μg/L at TAN-28, 591 μg/L at TAN-29) as well as TAN-37A (5.75 μg/L) were significantly lower than in 2016 but still within the range measured at these locations over the past 15 years.

Other VOC detections in 2017 were at RWMC, where TCE, carbon tetrachloride (MCL = 5 μg/L), and/or chloroform (MCL = 70 μg/L) were detected in three wells. Only one of these detections—carbon tetrachloride in well RWMC Production (5.93 μg/L) was above the MCL. The VOC detections at RWMC are consistent with historical observations.
Water Monitoring Verification Results

DEQ collects water samples at the same time and location as DOE contractors or the USGS and verifies that analytical results from co-sampled locations are consistent. The DEQ sampling verification program is designed to co-sample at approximately 10% of all DOE sample locations for selected analytes. In the event that a significant difference is found between DEQ results and those of the co-sampler, each result is scrutinized individually to ascertain the cause of the difference. Some differences between results are expected due to natural variability in the media being sampled, random errors in the measurements, and systematic differences in how the samples are collected, handled and analyzed. DEQ sets a goal of at least 80 percent of the results from co-sampled locations for each analysis passing the comparison criteria outlined in the Quality Assurance section.

Radiological

A summary of the sample-by-sample comparison of DEQ and DOE/USGS radiological results is presented in Table 6. Most results were in agreement, with at least 80 percent of results for co-sampled pairs passing comparison criteria for all analyses except gross beta radioactivity. The
reason for the differences in gross beta results is unknown at present, but it is notable that for all samples pairs failing to pass the comparison criteria, the result obtained by DEQ was larger than the result obtained by the co-sampler, suggesting a systematic bias. This issue will be investigated further in the coming year.

Table 6. Radiological results for co-samples collected by DOE and DEQ in 2017.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Number of co-sampled pairs in 2017</th>
<th>Percent of co-sampled pairs passing criteria in 2017</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gross alpha</td>
<td>30</td>
<td>97</td>
</tr>
<tr>
<td>Gross beta</td>
<td>30</td>
<td>57</td>
</tr>
<tr>
<td>$^{137}$Cs</td>
<td>22</td>
<td>91</td>
</tr>
<tr>
<td>$^3$H</td>
<td>54</td>
<td>98</td>
</tr>
<tr>
<td>$^{80}$Sr</td>
<td>26</td>
<td>96</td>
</tr>
<tr>
<td>$^{99m}$Tc</td>
<td>5</td>
<td>80</td>
</tr>
<tr>
<td>$^{234}$U</td>
<td>5</td>
<td>100</td>
</tr>
<tr>
<td>$^{235}$U</td>
<td>5</td>
<td>100</td>
</tr>
<tr>
<td>$^{238}$U</td>
<td>5</td>
<td>100</td>
</tr>
<tr>
<td>$^{239}$Pu</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>$^{239/240}$Pu</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>$^{241}$Am</td>
<td>2</td>
<td>100</td>
</tr>
</tbody>
</table>

**Non-Radiological**

A summary of the sample-by-sample comparison of DEQ and DOE/USGS non-radiological results for 2017 is presented in Table 7. Nearly all results were in agreement, with at least 80 percent of results for co-sampled pairs passing comparison criteria for all analyses except VOCs. For all but one pair of VOCs analyses, including all pairs for which Fluor was the co-sampler, DEQ’s result was higher than that of the co-sampler.
Table 7. Non-radiological results for co-samples collected by DOE and DEQ in 2017.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Number of co-sampled pairs in 2017</th>
<th>Percent of co-sampled pairs passing criteria in 2017</th>
</tr>
</thead>
<tbody>
<tr>
<td>Common Ions/Nutrients</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alkalinity</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
<td>Calcium</td>
<td>11</td>
<td>100</td>
</tr>
<tr>
<td>Chloride</td>
<td>38</td>
<td>100</td>
</tr>
<tr>
<td>Fluoride</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Magnesium</td>
<td>11</td>
<td>100</td>
</tr>
<tr>
<td>Potassium</td>
<td>11</td>
<td>100</td>
</tr>
<tr>
<td>Sodium</td>
<td>40</td>
<td>100</td>
</tr>
<tr>
<td>Sulfate</td>
<td>38</td>
<td>100</td>
</tr>
<tr>
<td>Total Nitrate plus Nitrite</td>
<td>31</td>
<td>100</td>
</tr>
<tr>
<td>Total Phosphorus</td>
<td>21</td>
<td>81</td>
</tr>
<tr>
<td>Trace Metals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arsenic</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>Barium</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>Chromium</td>
<td>28</td>
<td>93</td>
</tr>
<tr>
<td>Iron</td>
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<td>100</td>
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<tr>
<td>Lead</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>Manganese</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>Selenium</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>Zinc</td>
<td>7</td>
<td>100</td>
</tr>
<tr>
<td>VOCs(^{1})</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8 VOC analytes</td>
<td>42</td>
<td>79</td>
</tr>
</tbody>
</table>

\(^{1}\)VOCs were analyzed by DEQ at six co-sampled locations. DEQ's results were compared with those of the co-sampler for analytes that were detected in at least one sample.

Water Monitoring Impacts and Conclusions

DEQ sample results are mostly in agreement with those reported by DOE contractors and the USGS. Results of DEQ water monitoring have identified contamination in the Eastern Snake River Plain Aquifer as a result of historic waste disposal practices at the INL. Specifically:

- Concentrations of \(^{90}\)Sr, chloride, manganese, iron, and some VOCs exceeded federal drinking water standards (MCLs or SMCLs) at some sites on the INL in 2017. These sites are not used for drinking water.
- The concentration of total nitrate plus nitrite exceeded the MCL at one location in the Magic Valley, distantly downgradient from the INL. This exceedance may be attributed to agricultural impacts.
- Tritium was not detected at a concentration above background in the vicinity of the southern INL boundary; however, boundary sites with a history of elevated tritium concentrations were not sampled in 2017. No sites monitored by DEQ exceed federal drinking water standards for tritium. Concentrations of tritium at the INL continue to decline site-wide.
- Concentrations for other INL contaminants in water remain constant or continue to decrease at most locations as a result of changes in waste disposal practices. Chromium concentrations remained below the 100 μg/L MCL at all sites sampled by DEQ in 2017.
- INL impacts to the aquifer are not identifiable in water samples collected at sites distant from the INL.
**Terrestrial Monitoring**

Terrestrial monitoring is performed by measuring radionuclide accumulations in soil to help assess long-term trends of radiological conditions in the environment on and around the INL. Monitoring of milk samples is performed to indirectly verify the presence or absence of atmospheric radioiodine deposited in the terrestrial environment on and near the INL. Some of these data are also used to determine whether the monitoring results obtained by the DOE and its contractors were consistent with the soil and milk sampling results obtained by DEQ for these same locations.

**Terrestrial Monitoring Equipment and Procedures**

DEQ uses a combination of *in-situ* gamma spectrometry and physical soil samples to monitor concentrations of gamma-emitting radionuclides in soil at DEQ air monitoring stations and selected soil sampling sites on and around the INL (2017 soil sampling sites are shown in Figure 24). A portable gamma radiation detector was used in the field to collect surface gamma radiation measurements. These *in-situ* sampling measurements were then used to identify radionuclides present and to estimate soil radioactivity concentrations. No physical soil samples were collected during 2017.

DEQ collected milk samples from distribution centers where milk was received and from individual dairies in southern and southeastern Idaho. Milk sampling locations are shown in Figure 1. Raw milk samples were collected from trucks arriving at the distribution centers from each region of interest. For the independent cow and goat dairies, DEQ personnel drop off empty sample containers that are filled by the owner/operator of the dairy. The samples are picked up within 1-2 days of collection.

Two DEQ milk samples were collected and split by a DOE contractor each month. One half of the split samples were analyzed by DOE and the other half were submitted to DEQ for analysis. DEQ used the analysis results from these split samples to verify the DOE contractor’s milk sampling results and conclusions.
Terrestrial Monitoring Results and Trends

Monitoring concentrations of gamma-emitting radionuclides in surface soil provides insight to the transport, deposition, and accumulation of radioactive material in the environment as a result of INL operations and historic atmospheric testing of nuclear weapons. During 2017, DEQ made in-situ gamma spectrometry measurements to estimate accumulations of gamma-emitting radionuclides in surface soil at 34 locations. Of the 34 measurements, Cesium-137 ($^{137}$Cs) was the only man-made radionuclide that was detected. The average $^{137}$Cs value for in-situ measurements was 0.14 picocuries per gram (pCi/g) with a minimum value of 0.05 pCi/g and a maximum of 0.26 pCi/g. All results were well below the recommended federal screening limit for surface soil of 6.8 pCi/g of Cesium-137 (NCRP Report 129).

Milk sampling is conducted by DEQ to determine whether radioiodine is present or absent in the food supply. Radioiodine is produced in relatively large quantities during fission reactions (e.g., in nuclear reactors). The chemical nature of iodine makes it mobile under normal conditions. Gaseous radioiodine can be dispersed through the atmosphere and carried along with the wind until it is deposited on plants. Dairy cows and goats that graze on radioiodine-contaminated pasture or feed will accumulate iodine in the milk they produce. Drinking this milk could lead to an accumulation of radioiodine in the thyroid gland and a greater risk of thyroid cancer.

During 2017, DEQ analyzed 45 milk samples. Radioiodine ($^{131}$I) was not detected in any milk sample. The DEQ action level of 4.4 pCi/L is based upon the radioiodine concentration in milk necessary for an infant to receive an annual thyroid radiation dose of 5 millirem. The Food and
Drug Administration (FDA) recommended maximum concentration of $^{131}$I for food, including milk, is 4600 pCi/kg.

**Terrestrial Monitoring Verification Results**

Naturally occurring Potassium-40 ($^{40}$K) is present in milk and soil and is ideal as a quality control measurement and indicator of measurement sensitivity. Therefore, many of the comparisons conducted between DEQ and DOE sample results include this isotope, especially since the target radionuclide (such as Iodine-131) is seldom detected in milk samples.

Gamma spectroscopic analysis results of the 24 milk split samples collected by the DOE contractor and submitted to DEQ for analysis were compared with DOE results. $^{40}$K results obtained by DEQ showed 100% agreement with DOE contractor results, which is considered satisfactory. All $^{131}$I results were below the minimum detectable activity for both agencies.

The DOE contractor did not conduct any *in-situ* soil sampling in 2017.

**Terrestrial Monitoring Impacts and Conclusions**

Based upon terrestrial radiological measurements of soil and milk, there were no discernable impacts to the environment from INL operations. Long-term accumulation of radionuclides observed by soil monitoring was consistent with historical measurements and was in the range of concentrations expected as a result of historic above-ground testing of nuclear weapons.

**Quality Assurance for the ESP**

**Data Assessment Summary**

This section summarizes the results of the quality assurance (QA) assessment of the data collected during calendar year 2017 by the DEQ’s Environmental Surveillance Program. All analyses and quality control (QC) measures at the analytical laboratories used by the DEQ were performed in accordance with approved written procedures maintained by each laboratory. Sample collection and those analyses performed by DEQ were in accordance with written procedures maintained by the DEQ.

During calendar year 2017 the DEQ submitted QC samples for 293 radiological and non-radiological analyses, representing 12 percent of the 2446 field sample analyses completed. Analytical results for these QC samples (170 blank results, 81 duplicate results, and 42 spike results) were used to assess the precision, accuracy, and representativeness of results from analyzing laboratories. All analytical results for QC samples and field samples are found in the DEQ quarterly reports for 2017.

During 2017, one QC blank result, five QC duplicate results, and one QC spiked sample result failed DEQ acceptance criteria for groundwater. These failures resulted in fourteen associated groundwater field sample results being qualified as estimates (three enriched tritium, one chromium, one gross beta, eight total phosphorus, and one zinc). Also during 2017, 26 gross alpha and 26 gross beta results for weekly TSP air samples were qualified as rejected, all due to insufficient air sample volume.
The 2017 data usability (non-rejected results divided by total field sample results) was acceptable at 97.9%. In addition to the 2446 field sample analysis results completed, another 96 expected results were not obtained for 2017, due primarily to TSP sampler issues and environmental radiation detector electronic problems. The 2017 data completeness (non-qualified results divided by total field sample results expected) was acceptable at 93.6%. The field data were validated, assigned qualifiers to designate restrictions on their use, and deemed usable and complete, meeting the program’s data quality objectives.

**Issues and Problems**

No major issues or problems affecting data quality were identified during 2017.

**Comparing Data**

DEQ compares its data with DOE’s to determine whether the programs’ data sets are statistically equivalent, or if each program’s data support the same conclusions relative to environmental impacts and public health. To evaluate statistically the degree of agreement between organizations’ split sampling and co-sampling measurements, DEQ evaluates the Relative Percent Difference (RPD) between paired results using the following equation:

$$ \text{RPD} = \left( \frac{\text{DOE result} - \text{DEQ result}}{\frac{\text{DEQ result} + \text{DOE result}}{2}} \right) \times 100 $$

An RPD in the range of ±20% is considered to indicate acceptable agreement between measurements. For non-radiological analysis, the RPD is used to compare paired samples in which both of the results exceed five times the detection level. If one or both of the sample results are less than five times the detection level, the absolute difference between the two results is acceptable if it is less than or equal to the larger method detection limit.

For radiological analysis, the RPD is calculated (using the above equation) to compare paired samples if both results are greater than the sample-specific minimum detectable concentration (MDC). DEQ-INL OP also considers paired sample results with an absolute difference of no more than three times the pooled error (or “3 sigma”) to be in acceptable agreement. This is accomplished using the following equation:

$$ |R_1 - R_2| \leq 3(S_1^2 + S_2^2)^{1/2} $$

Where:
- $R_1$ = First sample value.
- $R_2$ = Second sample value.
- $S_1$ = Uncertainty (one standard deviation) associated with the laboratory measurement of the first sample.
- $S_2$ = Uncertainty (one standard deviation) associated with the laboratory measurement of the second sample.

Individual pairs of measurements having an absolute difference of no more than three times their pooled uncertainty, or with an RPD in the range of ±20%, are considered to be statistically in agreement. Paired data sets are considered to be in satisfactory statistical agreement if at least 80% of the individual paired results are in agreement.
Radiological Emergency Response Planning and Preparedness

DEQ’s role in emergency response planning and preparedness is defined in detail in the Environmental Oversight and Monitoring Agreement (EOMA) with the DOE. DEQ works with DOE and INL contractors to evaluate and participate in response planning, and to respond to incidents. DEQ works with state, federal and local agencies to respond to incidents, as described in the Idaho Hazardous Materials Response Plan. The Idaho Bureau of Homeland Security (IBHS) coordinates state emergency response actions in Idaho. Most of DEQ’s emergency response activities are directed towards planning and response to INL incidents. DEQ also responds to non-INL radiological incidents to help maintain lines of communication with the State’s emergency response organization, and as opportunities to test organizational readiness under real-world conditions. As a part of public outreach, DEQ can provide technical information, assistance, and training to local and state authorities for incidents involving radioactive materials at the INL or elsewhere in Idaho.

By agreement with DOE, INL radiological incident response planning is based on hazard assessment documents (HADs) developed by DOE contractors. These documents describe potential incidents at INL facilities that could release radionuclides to the environment. Review of current INL HADs is a key element of preparing for INL radiological emergencies. This information allows DEQ to identify scenarios that could potentially result in off-site radiological impacts, and plan appropriate responses. DEQ uses the source inventory and accident scenarios from the HADs to develop input for atmospheric dispersion and dose modeling using the Radiological Assessment System for Consequence Analysis (RASCAL) code. RASCAL uses real time National Oceanic and Atmospheric Administration (NOAA) weather data for regional-scale dispersion modeling. This allows DEQ to make independent radiological dose assessments for planning purposes, and would support development of timely technical and protective action recommendations for state authorities during actual emergencies. DEQ also receive text messages from the INL Warning Communication Center anytime their emergency resources are deployed; primarily the INL Fire Department.

Non-INL Radiological Activities

1. DEQ manager participated in 19 regional and county emergency planning meetings.
2. DEQ manager attended the National Transportation Stakeholders Forum meeting in June 2017.
3. DEQ manager was designated the State of Idaho Nuclear Regulatory Commission interface and attended a NRC State Liaison’s meeting.
4. DEQ manager attended Western Interstate Energy Board High Level Waste Committee meeting November 24-25, 2017. The group continued work on a set of policy papers.
Drills and Exercises

1. DEQ observed an emergency drill conducted to test the mutual aid agreement between Butte County and the INL on April 11th.
2. INL held a communications exercise on April 13th that included Idaho State Communications and DEQ was notified per procedure.
3. DEQ attended an INL emergency planning exercise on August 3rd.
4. DEQ participated in the INL annual exercise on September 20th.
5. DEQ participated in an INL emergency drill on November 9, 2017.

Waste Isolation Pilot Plant Shipment Safety

DOE contracts with the Western Governors Association (WGA) to coordinate activities related to the safe shipment of transuranic waste to the Waste Isolation Pilot Plant (WIPP) through western states. DEQ works with the Idaho State Police (ISP) and the Idaho Office of Emergency Management to manage WIPP shipment safety activities on the US Route 20/26, Interstate 15, and Interstate 84 / 86 corridors in Idaho.

During 2017, DEQ:

- Oversaw radiological equipment repairs and calibrations for ISP, all seven Idaho regional response teams, the Shoshone-Bannock Tribes, and three area hospitals.
- Staff members attended the National Transportation Stakeholders Forum and two meetings of the WIPP Technical Advisory Group and Western Governors Association Waste Isolation Pilot Plant Technical Advisory Group. DEQ also participated in monthly conference calls with the WIPP Technical Advisory Group.

Emergency Response

- DEQ attended fifteen Local Emergency Planning Committee (LEPC) meetings, and the seven regional emergency planning meetings. DEQ manager attended a Northwest Emergency Managers Workshop on December 14, 2017.
- DEQ responded to a facility in Boise to evaluate scrap metal for radionuclide content.

Classes and Presentations

1. DEQ reviewed and commented on updates to the Modular Emergency Response Radiological Transportation Training (MERRT) modules and videos.
2. DEQ provided training to Twin Falls Fire Department on March 20, 2017.
3. DEQ received training in INL Web Emergency Operations Center access and database.
4. One DEQ staff member attended the National Radiological Emergency Preparedness meeting April 10-14, 2017.
5. DEQ conducted radiological training for the 101st Civil Support Team at Gowen Field on November 20-21, 2017.
6. DEQ manager attended WIPP Transportation Tracking and Communication System (TRANSCOM) training.
Public Outreach

A fundamental aspect of DEQ’s work is sharing our findings with the public and factoring public input into our activities and policy recommendations. DEQ uses several tools to provide Idahoans with independent, accurate, and timely information about activities relating to the INL and other DOE activities in Idaho – publications, events, our Web site, and our community monitoring network.

Publications

DEQ regularly issues technical and non-technical publications to communicate the findings and activities of our program. In 2017, we issued:

- Four quarterly environmental surveillance data reports.


Presentations and Events

DEQ also communicates with the public about INL-related issues through schools, fairs, special interest groups, and public events. In 2017, we gave public presentations on the aquifer, and INL Site issues to a range of schools, civic groups, and special interest groups.

The Water Festival begins with a distribution of water education materials to approximately 3100 eastern Idaho students from 44 schools. Each year, some of the students from the Water Festival participate in the Poetry contest. The poems and winners are displayed in the Idaho Falls Library three weeks prior to the event (Figure 25). The event has now grown so large that we have extended it to two days attended by over 1,600 students. The Edible Aquifer, Rain Stick and Macro Invertebrates activities are presented to students (Figures 26, 27, 28).

Idaho Falls Earth Day continues to offer several activities for the youth and adults to enjoy. DEQ provides an Edible Aquifer activity to teach about the importance of water in our aquifer (Figure 29). DEQ-INL OP provides carry-all bags with Earth Day giveaways at the booth (Figure 30).
Figure 25. Water Awareness Poetry Contest 2017 on display at the Idaho Falls Library.

Figure 26. Children enjoying “Edible Aquifer” food activity at Water Festival 2017.
Figure 27. Children preparing their rain sticks at the Water Festival event 2017.

Figure 28. Children learning about Macro Invertebrates at the Water Festival event 2017.
Figure 29. Children participating in the Edible Aquifer activity at the 2017 Earth Day event.

Figure 30. DEQ staff handing out give-away items at the 2017 Earth Day event.
Community Monitoring Network

DEQ also participates in a community monitoring network in Eastern Idaho in cooperation with the Shoshone-Bannock Tribes, the U.S. Department of Energy, and NOAA. Strategically located community monitoring stations provide real-time atmospheric and radiological data to the public at each station location and also transmit data to the World Wide Web at http://www.idahoop.org/. Figure 31 shows the community monitoring station at Fort Hall. The monitoring boards have been updated at each location in 2017.

Figure 31. Community monitoring station at the Fort Hall location.