8.0 Groundwater Monitoring

MJ Hartman

The Hanford Site, part of DOE’s nuclear weapons complex, encompasses 586 square miles (~1,500 square kilometers) along the Columbia River in southeastern Washington State. During World War II and the Cold War period (1945 to 1991), the government built a total of nine reactors for the production of plutonium and other nuclear materials.

During reactor operations, chemical and radioactive waste was released into the environment and contaminated the soil and groundwater beneath portions of the Hanford Site, mostly in the 200 East Area, 200 West Area, 300 Area, 1100 Area, and the 100 Area reactor areas along the river (e.g., 100-BC, 100-K) (Figure 8.1). Since 1989, DOE has worked to remediate this contamination. DOE developed a plan to address groundwater and vadose zone contamination in consultation with EPA and Ecology. Key elements associated with managing the Hanford Site’s groundwater and vadose zone contamination are to: (1) protect the Columbia River and groundwater, (2) develop a cleanup decision process, and (3) achieve final cleanup restoring groundwater to its highest beneficial use.

Groundwater occurs in an unconfined aquifer within unconsolidated gravel and sand units. Groundwater in the unconfined aquifer generally flows from upland areas in the west toward the regional discharge areas along the Columbia River (Figure 8.2). The flow of water divides beneath the 200 East Area, with some water flowing toward the north and some flowing southeast. Maximum concentrations of key groundwater contaminants are presented in Tables 8.1 and 8.2; time series graphs of plume area over time for the largest plumes and the combined plume footprint are shown in Figure 8.3.
DOE has taken the following actions to protect the Columbia River from contaminated groundwater:

- Ceasing discharge of all unpermitted liquids in the central Hanford Site.
- Remediating waste sites in the 100 and 300 Areas to reduce the potential for future groundwater contamination.
- Containing groundwater plumes and reducing the mass of primary contaminants through remedial actions such as pump-and-treat (P&T).
Table 8.1. Overview of the River Corridor Groundwater Interest Areas Contaminant Concentrations

<table>
<thead>
<tr>
<th>Area</th>
<th>Primary Operations</th>
<th>Status of Waste Site Remediation</th>
<th>Status of Ground-water ROD</th>
<th>Groundwater Contamination: Maximum Concentration and Plume Area</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Carbon-14</td>
</tr>
<tr>
<td>100-BC</td>
<td>Reactor operations -- B Reactor 1944-69; C Reactor 1952-69</td>
<td>93% complete</td>
<td>None to date</td>
<td>N</td>
</tr>
<tr>
<td>100-K</td>
<td>Reactor operations -- KE Reactor 1955-71; KW Reactor 1955-70</td>
<td>59% complete</td>
<td>Interim action P&amp;T</td>
<td>14,300 pCi/L</td>
</tr>
<tr>
<td>100-N</td>
<td>Reactor operations -- N Reactor 1963-67</td>
<td>92% complete</td>
<td>Interim action permeable reactive barrier</td>
<td>52 pCi/L</td>
</tr>
<tr>
<td>100-D &amp; 100-H</td>
<td>Reactor operations -- D Reactor 1944-67; DR Reactor 1950-64; H Reactor 1949-65</td>
<td>87% complete</td>
<td>Interim action P&amp;T</td>
<td>N</td>
</tr>
<tr>
<td>100-F</td>
<td>Reactor operations -- F Reactor 1945-65; Biological experiments until 1976</td>
<td>98% complete</td>
<td>Final action MNA</td>
<td>N</td>
</tr>
<tr>
<td>300</td>
<td>Nuclear fuel fabrication and research -- 1940s-1960s</td>
<td>91% complete</td>
<td>Final action enhanced attenuation, MNA</td>
<td>N</td>
</tr>
<tr>
<td>1100 and Offsite: Vehicle maintenance, 1954-55; solid waste landfill --1950s-1970</td>
<td>100% complete (final action ROD)</td>
<td>Final action MNA; goals met</td>
<td>N</td>
<td>N</td>
</tr>
</tbody>
</table>

Standards:
- 2,000 pCi/L | 10 μg/L | 45 mg/L | 8 pCi/L | 5 μg/L | 20,000 pCi/L | 30 μg/L

Half-life (years):
- 5.730 | N/A | N/A | 28.8 | N/A | 12 | >159,000

Mobility in subsurface:
- High | High to Moderate | High | Slight | Moderate | High | Moderate

### Legend
- Colors indicate maximum concentration in 2014
- Height of bar indicates plume area above standard (km²)

### Notes
- (a) Approximate percentage by number of waste sites classified as closed, interim closed, no action, rejected, or not accepted (end of 2014).
- (b) Nitrates in 300-FF-5, and nitrates and uranium in 1100-EM-1, originate from offsite sources, so plume areas and maximum concentrations are not shown.
- (c) Drinking water standards for all but hexavalent chromium (aquatic standard).

**Abbreviations**
- MNA: Monitored natural attenuation
- N/A: Not applicable
- ROD: Record of decision

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Section 8: Groundwater Monitoring

Hanford Site Environmental Report for CY 2014

8.3
## Table 8.2. Overview of Central Plateau Groundwater Interest Areas Contaminant Concentrations

<table>
<thead>
<tr>
<th>Area</th>
<th>Primary Operations</th>
<th>Status of Groundwater ROD</th>
<th>Groundwater Remedial Action</th>
<th>Groundwater Contamination: Maximum Concentration and Plume Area</th>
</tr>
</thead>
</table>

**Standards**
- Carbon Tetrachloride: 5 µg/L
- Cyanide: 48 µg/L
- Iodide-129: 200 µg/L
- Nitrate: 45 mg/L
- Strontium-90: 8 µg/L
- Technetium-99: 5 µg/L
- Plutonium: 900 µg/L
- Tritium: 20,000 µg/L
- Uranium: 30 µg/L

**Half-life (years)**
- Carbon Tetrachloride: N/A
- Cyanide: N/A
- Iodide-129: 16,000 years
- Nitrate: N/A
- Strontium-90: 28.8 years
- Technetium-99: N/A
- Plutonium: 212,000 years
- Tritium: 12 years
- Uranium: >559,000 years

**Mobility in subsurface**
- Carbon Tetrachloride: Multi-phase
- Cyanide: High to Moderate
- Iodide-129: Moderate
- Nitrate: High
- Strontium-90: High
- Technetium-99: High
- Plutonium: High
- Tritium: Moderate
- Uranium: Moderate

**Legend**
- Colors indicate maximum concentration in 2014
- Height of bar indicates plume area above standard (km²)
- >100 x standard and <1,000 x standard
- >10 x standard and <100 x standard
- >10 x standard and <10 x standard
- Not detected or not analyzed

**Notes**
- Drinking water standards for all but hexavalent chromium (MTCA standard)

**Abbreviations**
- N/A = Not applicable
- P&T = pump-and-treat
- ROD = record of decision
- MTCA = Model Toxics Control Act

*CHS282014G20605*
DOE operates an extensive groundwater monitoring program on the Hanford Site, collecting thousands of samples from hundreds of wells each year. In addition to monitoring wells, DOE monitors hundreds of sampling points near the Columbia River, known as aquifer tubes, for general information about groundwater approaching the river. DOE sampled 977 monitoring and extraction wells, and 324 aquifer tubes in 2014. Many of them were sampled multiple times, for a total of 4,654 sampling events (Figure 8.4).
Figure 8.5 compares maximum concentrations of the major groundwater contaminants in various parts of the Site in 2014. These contaminants are discussed further in the following sections. This chapter is organized by geographic regions known as “groundwater interest areas” (Figure 8.5) within the River Corridor (100 and 300 Areas) and the Central Plateau (200 Areas).

![Figure 8.5. Exceedance Ratios of Groundwater Contaminants in 2014](image)

Bar height shows contaminant concentrations in multiples of the applicable water quality standard.
8.1 River Corridor

Since the 1990s, DOE has been remediating waste sites and groundwater in the River Corridor under interim action RODs. Removal of contaminated soil has reduced the potential for exposure to contaminants, including future groundwater impacts. As of the end of 2014, 89 percent of the waste sites in the River Corridor had been remediated or were classified as not needing remediation under interim action RODs, as compared to 85 percent in 2013 and 74 percent in 2012. Cleanup of the remaining sites is underway. Based on remedy performance monitoring and the reduction in length of shoreline impacted by contaminant plumes, groundwater remediation systems in 100-HR, 100-KR, and 100-NR are reducing the amount of contamination entering the Columbia River.

Table 8.1 provides a summary of the River Corridor groundwater interest areas and associated contamination plumes. In the 100 Area, groundwater contamination is related to past disposal of waste associated with water cooled nuclear reactors. The primary groundwater contaminants of concern (COC) in the 100 Area are chromium (hexavalent and total), strontium-90, nitrate, trichloroethene, and tritium (Figure 8.6). Sources of hexavalent chromium contamination included the routine disposal of reactor cooling water, which contained the corrosion inhibitor sodium dichromate, and unplanned spills and leaks of the high-concentration sodium dichromate stock solution. In the 300 Area, the groundwater COCs are uranium, tritium, nitrate, gross alpha, trichloroethene and cis-1,2-dichloroethene.

Under interim action RODs, groundwater remediation systems in the 100-HR-3 and 100-KR-4 Operable Units are limiting the amount of contamination reaching the Columbia River and reducing the mass of contaminants. The primary contaminant addressed is hexavalent chromium. The comparison concentration for inland groundwater wells is 20 \( \mu g/L \). Similar to other river corridor decisions (e.g., 100-FR-3), the cleanup level for groundwater discharges to the river for the final action ROD for 100-HR-3 and 100-KR-4 is expected to be 10 \( \mu g/L \), when they are completed. EPA and DOE signed a final action ROD for 100-FR waste sites and groundwater in 2014. Final action RODs previously were signed for the 300-FF-5 and 1100-EM-1 OUs. Final action RODs for the other portions of the River Corridor are expected to be developed in the next few years.

8.1.1 100-BC

Groundwater contaminants in 100-BC include hexavalent chromium and strontium-90. Tritium concentrations remained below the DWS in 2014. Waste sites in 100-BC have been remediated under an interim action ROD, so contaminant levels in groundwater are expected to continue to decline.

Remedial investigation (RI) studies continued in 100-BC in 2014, with additional sampling of wells and river shoreline sampling points installed in 2013. The studies, which are expected to conclude in 2016, will provide data to support remedy decisions for groundwater cleanup. This includes ongoing, intensive sampling of water in the shallow river bed to evaluate variable concentrations of hexavalent chromium, at the groundwater/surface water interface.

DOE and EPA have agreed that additional RI studies are needed to reduce uncertainties relating to (1) the completion of waste site remediation; (2) short term changes in groundwater contaminants related to waste site remediation; (3) modeling results predicting that the hexavalent chromium plume could persist for over 100 years; and (4) the level of risk associated with variable hexavalent chromium concentrations in Columbia River pore water.
8.1.2 100-KR

Hexavalent chromium is the primary contaminant in 100-KR groundwater. Smaller plumes of carbon 14, tritium, strontium 90, nitrate, and trichloroethene also are present. Cleanup actions for these other contaminants will be defined in an upcoming ROD. DOE has proposed additional P&T for hexavalent chromium as part of a preferred alternative for groundwater remediation. The draft RI/Feasibility Study (FS) and Proposed Plan underwent review in 2012, and DOE will incorporate the results of supplemental source characterization activities that includes drilling boreholes near the KE Fuel Storage Basin and 116-KE-3 Crib and Reverse Well. These boreholes are expected to be drilled in 2015.

Approximately 59 percent of 100-KR waste sites have been remediated or were determined not to require remediation under an interim action ROD. Three P&T systems continued to operate in 100-KR to remove hexavalent chromium from groundwater. In 2014, over 607.6 million gallons (2.3 billion liters) of groundwater was pumped from 41 extraction wells and a total of 1,757 pounds (797 kilograms) of...
hexavalent chromium have been removed to date. The hexavalent chromium plume area (greater than 20 µg/L) was estimated to be 0.29 square miles (0.76 square kilometers) in 2014, a decrease from 2013. Since 2007, the plume area above 20 µg/L has decreased by approximately 70 percent, and the length of shoreline that the plume is interpreted to intersect (based on data from wells and aquifer tubes) has decreased from 7,200 feet (2,200 meters) to 660 feet (200 meters) (Figure 8.7).

For AEA purposes, DOE monitors groundwater near the KW and former KE Fuel Storage Basins, which were integral parts of each reactor building. Groundwater monitoring in 2014 did not show new groundwater impacts from the basins.

**Figure 8.7. 100-KR Hexavalent Chromium Plume in 1996 (Before Interim Action) and 2014 (During Interim Action)**

8.1.3 **100-NR**

DOE submitted a draft RI/FS report and Proposed Plan to Ecology for review in 2013. When finalized, these documents will be used to develop a ROD documenting remediation of waste sites and groundwater. Work continued in 2014 on revisions to the RI/FS report in response to Ecology comments. The major liquid waste disposal sites have been remediated, and excavation is continuing at remaining waste sites. Work is expected to be complete in 2015. Principal groundwater activities for 100-NR include RCRA monitoring and remediation of strontium-90 and total petroleum hydrocarbons. Other groundwater contaminants include nitrate and tritium. Hexavalent chromium from 100-KR has affected 100-NR groundwater in some locations.
Strontium-90, which originated at the 116-N-1 and 116-N-3 waste sites, is the primary contaminant. Strontium (including the strontium-90 isotope) substitutes for calcium in the sediment, reducing the mobility of this contaminant in the vadose zone and groundwater. As a result, the shape and size of the plume (Figure 8.8) has not changed significantly since 1996.

P&T technology, which operated from 1995 to 2006, was found to be ineffective in cleaning up strontium-90, so DOE is now applying an in situ technology called strontium-90 sequestration, using an apatite chemical solution. Under an interim action ROD, a 900 foot (170 meter) section of a permeable reactive barrier was placed along the shoreline, reducing the amount of strontium-90 migrating from groundwater into the river. Expansion of the barrier to its full 2,500 foot (760 meter) length is pending.

In 2014, RCRA monitoring continued under final status detection programs at the 1301-N, 1324-N/NA, and 1325-N facilities (waste sites 116-N-1, 120-N-1, 120-N-2, and 116-N-3). Results indicated no releases of dangerous waste constituents from the RCRA units.

8.1.4 100-HR

The 100-HR-3 Groundwater OU in the northern Hanford Site includes the 100-HR-D and 100-HR-H groundwater interest areas. Hexavalent chromium is the primary COC. Additional groundwater contaminants in 100-HR include strontium-90 and nitrate (Figure 8.6). Investigation of groundwater conditions at 100 HR greatly changed the understanding of the extent of chromium contamination since P&T began, primarily because many more wells were installed. In 1997, 110 wells and aquifer tubes were sampled, and in 2014, over 330 wells and aquifer tubes were sampled in 100-HR. The added wells and aquifer tubes identified areas of higher chromium concentrations at 100 D, in the Horn, and in the Ringold Formation upper mud unit (RUM). Even with areas of high levels of contamination being identified, the overall areal extent of the plume has decreased as a result of remediation (Figure 8.9).

The CERCLA process is underway to make final cleanup decisions for 100-HR. DOE submitted the Draft A RI/FS and Proposed Plan in 2012. In 2013 and 2014, DOE and Ecology worked through the comment resolution process, and Ecology accepted the 100-HR-3 RI/FS Report in October 2014. The Proposed Plan is expected to be available for public comment in 2015 or 2016. A ROD will then be issued that identifies the final remedial alternatives. DOE has proposed ongoing P&T as the preferred alternative for remediating hexavalent chromium in groundwater.

Approximately 87 percent of the former waste sites have been remediated or were determined not to require remediation under an interim action ROD. Remediation of waste sites continued in 2014, including 100-D-100, a major source of hexavalent chromium. Contaminated sediment was excavated down to the water table, and excavation into the top of the aquifer continues in 2015.

Two P&T systems continued to operate under an interim action ROD, removing hexavalent chromium. In 2014, 634 million gallons (2.4 billion liters) of groundwater were pumped from 74 extraction wells. A total of 4,952 pounds (2,246 kilograms) of hexavalent chromium have been removed to date. The plume area (greater than 20 µg/L) was estimated to be 1.4 square miles (3.5 square kilometers in 2014, a decline from 2013. Since 2005 the plume has decreased in area by over 60 percent, and the length of shoreline intersected by the plume (above 20 µg/L) has been reduced from 8,200 feet (2,550 meters) to zero (Figure 8.9). The changes are a result of contaminant removal, remediation of sources, hydraulic control, and natural processes.
The former 183-H Solar Evaporation Basins (waste site 116-H-6) constitute the only RCRA site in 100-HR. The site is monitored in accordance with RCRA corrective action requirements during the post-closure period to track contaminant trends during operation of the CERCLA interim action for hexavalent chromium.

*Figure 8.8.* 100-NR 2014 Strontium-90 Plume and Apatite Barrier
Figure 8.9. 100-HR Hexavalent Chromium Plume in 1999 (Early in Interim Action Period) and 2014 (During Interim Action)
8.1.5 100-FR

Groundwater contamination in 100-FR originated from disposal of solid and liquid waste associated with operation of the water-cooled F Reactor and biological experiments. Nitrate concentrations in groundwater exceed the DWS beneath much of the 100-F Area and the plume extends southward approximately 3.1 miles (5 kilometers). Smaller plumes of hexavalent chromium, strontium-90, and trichloroethene are present (Figure 8.6). Contaminant concentrations are below cleanup standards near the river and are declining.

Former 100-FR waste sites have been excavated and backfilled under an interim action ROD. In 2014, EPA and DOE signed a ROD that includes monitored natural attenuation (MNA) as the preferred alternative for 100-FR-3 OU groundwater remediation. Preparation of a work plan and sampling and analysis plan is underway.

8.1.6 300-FF

Three geographic regions comprise 300-FF: the 300 Area Industrial Complex; the 618-11 Burial Ground region; and a region including the 618-10 Burial Ground and 316-4 Cribs (Figure 8.6). Approximately 91 percent of the waste sites have been remediated or classified as not requiring remediation. Remediation is continuing at the remaining sites. DOE is conducting field and laboratory studies to understand and model the processes that control contaminant flux between groundwater and the Columbia River. In 2014, studies focused on seasonal water quality dynamics (including uranium and nitrate).

EPA and DOE signed a ROD in 2013 and the remedial design report/remedial action work plan that implements the 2013 final action ROD is anticipated to be issued in 2015. The remedial action for groundwater includes enhanced attenuation of uranium using sequestration by phosphate application. MNA is the selected remedy for other COCs: trichloroethene and cis-1,2-dichloroethene at the 300 Area Industrial Complex and tritium and nitrate at the 618-11 Burial Ground. Uranium concentrations remain above the cleanup level (30 µg/L) in groundwater in the 300 Area Industrial Complex and downgradient from the 618-7 Burial Ground (Figure 8.10). Contamination from 618-7 was mobilized by waste site remediation activities in recent years.

Trichloroethene concentrations exceeded the cleanup level (4 µg/L) in one 300-FF monitoring well and several aquifer tubes in 2014. Concentrations of nitrate above 45 mg/L are also present in groundwater beneath part of the 300 Area Industrial Complex, but these originated from sources off the Hanford Site; nitrate in the 300 Area Industrial Complex is not a COC for 300-FF.

Groundwater associated with the 618-11 Burial Ground, north of the 300 Area Industrial Complex, contains a high-concentration tritium plume originating from irradiated material in the burial ground. The waste site has not been remediated. Nitrate concentrations near the 618-11 Burial Ground also continued to exceed the cleanup level (45 mg/L).

RCRA groundwater monitoring continued at the 300 Area Process Trenches (waste site 316-5). The unit is monitored in accordance with post-closure corrective action requirements (WAC 173-303-645[11]). Uranium and cis-1,2-dichloroethene continued to exceed permit concentration limits in 2014.

In accordance with the closure plan, groundwater corrective action will be addressed as part of the remediation for the CERCLA 300-FF-5 Groundwater OU.
8.1.7 1100-EM and Richland North

Remediation of the former 1100-EM-1 OU is complete and the 1100-EM-1 Groundwater OU was removed from the “National Priorities List” (40 CFR 300, Appendix B) in 1996. The selected remedy was MNA of volatile organic compounds, with institutional controls (IC) on drilling of new water supply wells. Trichloroethene is the primary COC, but concentrations have remained below the cleanup level since 2001.

Uranium concentrations in Hanford Site wells in the vicinity of DOE’s inactive Horn Rapids Landfill have increased gradually since 1996, exceeding the DWS in 2012 and dropping slightly below the standard in 2014. The presence of uranium at these locations is attributed to a plume moving northeast from an active offsite facility, AREVA NP, Inc. a nuclear fuel production facility.

DOE monitors wells in and near the north Richland well field, which is part of the municipal water supply system. Tritium concentrations are at background levels.

8.2 Central Plateau

When the Hanford Site was operating as a plutonium-production facility, irradiated fuel reprocessing, isotope recovery, and associated waste management activities occurred in the 200 East and 200 West Areas in the central portion of the Site. Ponds, cribs, and ditches used for disposal of liquid waste were primary sources of groundwater contamination. There are also seven single-shell tank WMAs in the 200 Area. Some of these tanks have leaked, contaminating the vadose zone and groundwater beneath the tanks. Contamination is still present in many parts of the thick vadose zone, and may continue to drain into the
groundwater. Remediation of the Central Plateau waste sites and vadose zone will accelerate after River Corridor remediation is complete. Meanwhile, DOE has been remediating groundwater and testing methods to remediate the deep vadose zone.

Groundwater contaminant plumes of tritium, nitrate, and iodine-129 formed when the waste discharged to ponds and cribs reached the aquifer. These contaminants form regional plumes originating on the Central Plateau. The plumes have decreased in area over the years as a result of dispersion and, in the case of tritium, radioactive decay. A large carbon tetrachloride plume originated in the PFP area of 200 West. Other groundwater contaminants in the Central Plateau include technetium-99, uranium, strontium-90, trichloroethene, hexavalent chromium, cyanide, and other dangerous waste constituents (Table 8.2 and Figure 8.11).

*Figure 8.11. Groundwater Contaminants in the Central Plateau*
An estimated 1,256,635 to 2,028,253 pounds (570,000 to 920,000 kilograms) of liquid wastes with carbon tetrachloride was discharged to waste sites in 200-ZP-1. Remediation has reduced the size of the high-concentration core and the overall footprint of the carbon tetrachloride plume. Combined, the final action system, the interim action system, and a soil vapor extraction system have removed a total of 221,556 pounds (100,496 kilograms) of carbon tetrachloride from the subsurface.

A P&T system at WMA S-SX in the 200-UP-1 OU, which began operating in July 2012, has removed 1.66 Ci of technetium-99, 35,891 pounds (16,280 kilograms) of nitrate, 62 pounds (28.1 kilograms) of chromium, and 551 pounds (250 kilograms) of carbon tetrachloride from groundwater.

The size of the regional tritium plume associated with 200-PO-1 has decreased from 71 to 49 square miles (185 to 79 square kilometers) since 1980, primarily as a result of radioactive decay and dispersion.

### 8.2.1 200-ZP

Contaminant sources in 200-ZP, located in the 200 West Area, included cribs, ponds, and single-shell storage tanks. A final action ROD for 200-ZP-1 OU groundwater identified carbon tetrachloride as the primary COC. Other COCs are trichloroethene, iodine-129, technetium-99, nitrate, chromium, and tritium. Two Low-Level Waste Management Areas (LLWMA-3 and LLWMA-4) in 200-ZP are monitored under RCRA interim status, contaminant indicator parameter programs. Monitoring results showed no indication that either of these is contaminating groundwater.

RCRA assessment monitoring continued at WMA T and WMA TX-TY. Due to CERCLA remediation activities (operation of the 200 West P&T system) near WMA T, chromium concentrations are declining and the plume extents at both WMAs are shrinking.

The State-Approved Land Disposal Site (SALDS) receives treated water from the Hanford Site’s Effluent Treatment Facility. It is regulated under a state waste discharge permit and has created a local tritium plume. All groundwater sampling results from the SALDS proximal wells were within permit compliance limits during 2014.

A P&T system has reduced the high-concentration core area of the carbon tetrachloride plume (Figure 8.12). The plume area above 5 µg/L was 5.1 square miles (13.1 square kilometers) in 2014, compared to 5.9 square miles (15.4 square kilometers) in 2013. In 2014, 20 extraction wells and 20 injection wells were in use and the treatment plant operated at a flow rate of 1,562 gallons per minutes (5,913 liters per minute) (71 percent of its design capacity). Additional wells were installed in 2014 that will be used to expand the extraction network. In 2014, the system processed 819 million gallons (3.1 billion liters) of groundwater and removed 6,164 pounds (2,796 kilograms) of carbon tetrachloride, 517,240 pounds (234,616 kilograms) of nitrate, and other contaminants from groundwater. Combined, the final action system, the interim action system, and a soil vapor extraction system have removed a total of 221,556 pounds (100,496 kilograms) of carbon tetrachloride from the subsurface (Figure 8.13).
Figure 8.12. 200-West Carbon Tetrachloride Plume in 1996 (Upper Part of Unconfined Aquifer) and 2014 (Including Available Vertical Interval Data)

Figure 8-13. 200-ZP Carbon Tetrachloride Mass Removed by Final Pump-and-Treat, Interim Pump-and-Treat, and Soil Vapor Extraction
8.2.2 200-UP

The southern portion of the 200 West Area and adjacent areas to the east and south comprise 200-UP. Contaminant sources included cribs, ponds, and single-shell tanks. Carbon tetrachloride, technetium-99, uranium, tritium, iodine-129, nitrate, and chromium plumes are present. Carbon tetrachloride in this region originated from sources in 200-ZP. Wells near WMA S-SX monitor the highest technetium-99 concentrations on the Hanford Site, and the plume has grown in recent years (Figure 8.14). An interim action ROD addressing all of the major contaminant plumes within the 200-UP-1 OU was published in 2012. The selected remedy in the ROD consists of a combination of P&T, MNA, hydraulic containment, and institutional controls.

The P&T system at WMA S-SX began operating in July 2012. From 2012 to 2014 the system removed 1.66 Ci of technetium-99, 35,891 pounds (16,280 kilograms) of nitrate, 62 pounds (28.1 kilograms) of chromium, and 551 pounds (250 kilograms) of carbon tetrachloride from groundwater. Another part of groundwater remediation under the interim action ROD is a groundwater extraction system to remediate the uranium and technetium-99 plumes in the U Plant area. The system is currently being designed and will be constructed in 2015.

![Figure 8.14. WMA S-SX Technetium-99 Plume in 2007 and 2014](image)

RCRA monitoring in 200-UP includes interim status groundwater quality assessment monitoring at WMA S-SX and WMA U, and interim status indicator parameter evaluation monitoring at the 216-S-10 Pond and Ditch. WMA S-SX has contaminated groundwater with chromium, nitrate, and the non-RCRA
constituent technetium-99. Water levels have declined at WMA S-SX due to groundwater extraction, causing some monitoring wells to go dry sooner than they would have otherwise. One new well was installed in 2014 and four additional replacement wells are planned in 2015. Sources within WMA U have contaminated groundwater with nitrate and chromium. The groundwater beneath this tank farm is within the capture zone of a nearby extraction well. Indicator parameters did not exceed statistical comparison values at the 216-S-10 Pond and Ditch during 2014.

The Environmental Restoration Disposal Facility (ERDF) is a CERCLA disposal facility used for disposal of low-level radioactive mixed waste generated by remedial actions. The results of groundwater monitoring in 2014 continued to indicate that the facility has not impacted groundwater.

8.2.3 200-BP

The 200-BP groundwater interest area includes the northern 200 East Area and the region to the northwest where mobile contaminants have migrated between Gable Mountain and Gable Butte. Most of the groundwater contamination is concentrated beneath WMA B-BX-BY and adjacent waste sites in the northwestern portion of the 200 East Area. Nitrate, iodine-129, and technetium-99 form the largest contaminant plumes. The high-concentration cores of these plumes have grown in size since 2007 due to continued drainage of contaminated water from the vadose zone into the aquifer (Figure 8.15). Smaller plumes of uranium, cyanide, strontium-90, and tritium also exceed their respective DWSs. Cesium-137 and plutonium-239/240 contamination is limited to one or two wells.

Wells in the northwestern part of 200-BP detect the highest concentrations of uranium in Hanford Site groundwater. Concentrations are even higher in a zone of perched water that lies above the water table. DOE, EPA, and Ecology signed an Action Memorandum in 2014 that directs continuing the extraction of contaminated perched water as a non-time-critical removal action under CERCLA. Approximately 117 pounds (53 kilograms) of uranium were removed from the perched zone through the end of 2014. A draft RI report for the 200-BP-5 OU was prepared in 2014, describing the nature and extent of contamination and identifying contaminants of potential concern to support a future FS. In addition, work began on the FS in late 2014.

Six RCRA sites with groundwater monitoring requirements are located in 200-BP. RCRA groundwater quality assessment monitoring at WMA B-BX-BY and WMA C indicates that the dangerous waste constituent cyanide in groundwater originated in the WMAs. Because of the continued migration of this dangerous waste constituent an additional well was installed at WMA B-BX-BY in 2014. RCRA contamination indicator parameter monitoring continued at the 216-B-63 Trench, LLWMA-1, and LLWMA-2 in 2014. Results continued to show that these units have not impacted groundwater. DOE monitors the LERF under a RCRA final status detection program. Results showed no indication that the site has affected groundwater. A new groundwater monitoring plan for LERF was implemented in 2014.
8.2.4 200-PO

The southern portion of the 200 East Area and a large region of the Hanford Site to the east and southeast comprise 200-PO. Disposal of large volumes of liquid waste created regional groundwater plumes of tritium, iodine-129, and nitrate. An RI addendum report for the 200-PO-1 OU is currently being prepared to update the risk assessment based on additional groundwater data collected since the RI was completed in 2008 and 2009. In addition, work began on the FS in late 2014.

The tritium plume had an estimated area of 30.6 square miles (79.2 square kilometers) in 2014, a 5 percent decrease from 2013. The size of the regional tritium plume (Figure 8.16) from 200-PO has decreased from 71 to 31 square miles (185 to 79 square kilometers) since 1980, primarily as a result of radioactive decay and dispersion. Concentrations of tritium are declining as the groundwater plume attenuates naturally as a result of radioactive decay and dispersion. The maximum concentration has declined from over 6 million pCi/L in the 1980s to 510,000 pCi/L in 2014.

The area of the iodine-129 plume above the 1 pCi/L contour has decreased slightly over the past decade, and maximum concentrations have declined as a result of dispersion. Radioactive decay has not decreased the level of iodine-129 contamination noticeably because this isotope has a half-life of 15.7 million years. The nitrate plume covers a large area, with concentrations above background, but mostly below the DWS. Other contaminants in 200-PO include strontium-90, technetium-99, and uranium in smaller areas near their discharge sources (Figure 8.11).
In 2014, RCRA assessment monitoring continued at WMA A-AX and interim status indicator parameter programs continued at the 216-A-36B Crib, 216-A-37-1 Crib, 216-A-29 Ditch, 216-B-3 Pond, and NRDWL. One monitoring well with casing corrosion associated with WMA A-AX was decommissioned in 2013. Drilling of a replacement well started in November 2014 and will be completed in early 2015. Monitoring results from the interim status sites provided no indication of releases from these facilities to groundwater.

The IDF is an expandable, double-lined landfill that is regulated under RCRA and the AEA. It is not yet in use, and current groundwater monitoring is directed at obtaining baseline data.

The Solid Waste Landfill is regulated under Washington State solid waste handling regulations. As in previous years, some of the monitoring wells showed higher concentrations of regulated constituents than the statistically calculated background threshold values. Background threshold values exceeded during 2014 included specific conductance, nitrite, sulfate, and total organic carbon.

Three onsite water supply wells provide drinking water and serve as an emergency water supply for the 400 Area, which is in the footprint of 200-PO. Because the 400 Area is in the path of the Hanford Site-wide tritium plume, DOE routinely monitors the wells for tritium.
8.3 Confined Aquifers

Although most Hanford Site groundwater contamination is found in the unconfined aquifer, DOE monitors wells in deeper aquifers because of potential downward movement of contamination.

One confined aquifer occurs within sand and gravel at the base of the Ringold Formation. Carbon tetrachloride, nitrate, and technetium-99 have contaminated this unit in a portion of the 200 West Area where the upper confining unit is absent. Newer wells have been installed to monitor and remediate this contamination. The Ringold confined aquifer is the uppermost aquifer in a region east of 200 East (within portions of 200-BP and 200-PO). Iodine-129 and tritium are detected in wells at this location, but the contamination has not migrated farther to the east or southeast.

In the northern Hanford Site, fine-grained sedimentary units, informally called the RUM, confine deeper sediments in the Ringold Formation. In some parts of 100-HR this unit is contaminated with hexavalent chromium at concentrations over 100 µg/L and is being remediated by a P&T system.

Groundwater within basalt fractures and joints, interflow contacts, and sedimentary interbeds make up the upper basalt-confined aquifer system.

8.4 Wells

Over the lifetime of the Hanford Site, DOE has installed thousands of wells to monitor and remediate groundwater and provide geologic data. During 2014, DOE installed 30 new wells and 6 new aquifer tubes. Table 8.3 lists the new wells and aquifer tubes installed in 2014 and Figure 8.17 illustrates the number of wells installed during the past 10 years.

DOE identifies wells, boreholes, or other subsurface installations for decommissioning when they are no longer needed. This involves sealing the wells in compliance with Washington State standards for construction and maintenance of wells (WAC 173-160). Four temporary wells in 100-D Area (199-D5-155, 199-D5-156, 199-D5-157, and 199-D5-158), were decommissioned in 2014.

<table>
<thead>
<tr>
<th>Groundwater Interest Area</th>
<th>Wells</th>
<th>Aquifer Tubes</th>
</tr>
</thead>
<tbody>
<tr>
<td>100-BC</td>
<td>8</td>
<td>6</td>
</tr>
<tr>
<td>100-HR-D</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>100-KR</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>200-BP</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>200-PO</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>200-UP</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>200-ZP</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>30</td>
<td>6</td>
</tr>
</tbody>
</table>
8.5 Additional Information

The monitoring data presented in this chapter—and information on monitoring well locations, construction details, and screened intervals—can be found through the DOE Environmental Dashboard Application at [https://ehs.hanford.gov/eda/](https://ehs.hanford.gov/eda/) or on the PHOENIX website at [http://phoenix.pnnl.gov](http://phoenix.pnnl.gov). The data and additional groundwater monitoring details are available in the Hanford Site Groundwater Monitoring Report for 2014 ([DOE/RL-2015-07](https://ehs.hanford.gov/eda/)).
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