

RADIOCHEMICAL AND CHEMICAL CONSTITUENTS IN WATER FROM SELECTED WELLS AND SPRINGS FROM THE SOUTHERN BOUNDARY OF THE IDAHO NATIONAL ENGINEERING AND ENVIRONMENTAL LABORATORY TO HAGERMAN AREA, IDAHO, 1997

U.S. GEOLOGICAL SURVEY Open-File Report 98-646



Prepared in cooperation with the

U.S. DEPARTMENT OF ENERGY and IDAHO DEPARTMENT OF WATER RESOURCES



Cover: Center-Pivot Sprinkler System on Snake River Plain.

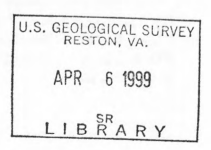
Radiochemical and Chemical Constituents in Water from Selected Wells and Springs from the Southern Boundary of the Idaho National Engineering and Environmental Laboratory to the Hagerman Area, Idaho, 1997

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CONVERSION FACTORS AND ABBREVIATED UNITS

Multiply	Ву	To Obtain
acre-foot per year (acre-ft/yr)	1,233	cubic meter per year
foot (ft)	0.3048	meter
gallon (gal)	3.785	liter
mile (mi)	1.609	kilometer
millirem per year (mrem/yr)	0.010	millisievert per year
picocurie per liter (pCi/L)	0.037	becquerel per liter
square mile (mi ²)	2.590	square kilometer

Temperature can be converted from degrees Celsius (°C) to degrees Fahrenheit (°F) by the equation: ${}^{\circ}F = ({}^{\circ}C \times 1.8) + 32$.

Abbreviated units used in report: mL (milliliter); L (liter); μ g/L (microgram per liter); mg/L (milligram per liter); μ m (micrometer); and μ S/cm (microsiemens per centimeter) at 25 degrees Celsius.

Radiochemical and Chemical Constituents in Water from Selected Wells and Springs from the Southern Boundary of the Idaho National Engineering and Environmental Laboratory to the Hagerman Area, Idaho, 1997

By Roy C. Bartholomay and Linda M. Williams, U.S. Geological Survey, and Linford J. Campbell, Idaho Department of Water Resources

Abstract

The U.S. Geological Survey and the Idaho Department of Water Resources, in cooperation with the U.S. Department of Energy, sampled 18 sites as part of the fourth round of a long-term project to monitor water quality of the Snake River Plain aquifer from the southern boundary of the Idaho National Engineering and Environmental Laboratory to the Hagerman area. Water samples were collected and analyzed for selected radiochemical and chemical constituents. The samples were collected from seven domestic wells, six irrigation wells, two springs, one dairy well, one observation well, and one stock well. Two quality-assurance samples also were collected and analyzed.

None of the radiochemical or chemical constituents exceeded the established maximum contaminant levels for drinking water. Many of the radionuclide- and inorganic-constituent concentrations were greater than their respective reporting levels.

INTRODUCTION

Recently, the public has expressed concern about waste-disposal practices at the Idaho National Engineering and Environmental Laboratory (INEEL) and the effect these practices might have had on the water quality of the Snake River Plain aquifer. The U.S. Department of Energy (DOE) requested that the U.S. Geological Survey (USGS) conduct two studies to respond to the public's concern and to gain a greater understanding of the radiochemical and chemical quality of water in the aquifer. The first study described a one-time

sampling effort in the eastern part of the A & B Irrigation District in Minidoka County (Mann and Knobel, 1990). The second study, an ongoing annual sampling effort in the area between the southern boundary of the INEEL and Hagerman (fig. 1), is being conducted with the Idaho Department of Water Resources (IDWR) in cooperation with the DOE. The first round of sampling for the second study involved analyzing water samples collected from 55 sites during August and September 1989 (Wegner and Campbell, 1991). The second round of sampling involved analyzing water samples collected from 19 of the initial 55 sites in 1990 (Bartholomay and others, 1992), another 18 of the initial 55 sites in 1991 (Bartholomay and others, 1993), and the remaining 18 sites in 1992 (Bartholomay and others, 1994a). An evaluation of data collected during the first four years of the study is found in Bartholomay, Williams, and Campbell (1997a). The third round of sampling involved analyzing water samples collected from 19 of the initial 55 sites in 1993 (Bartholomay and others, 1994b), another 18 of the initial 55 sites during 1994 (Bartholomay and others, 1995). another 17 of the initial 55 sites during 1995 (Bartholomay and others, 1996), and one final site during 1996 (Bartholomay, Williams, and Campbell, 1997b). The first part of the fourth round of sampling involved analyzing water samples collected from 19 of the initial 55 sites in 1996 (Bartholomay, Williams, and Campbell, 1997b). This report summarizes the results of analyses of water samples collected from another 18 sites as part of the fourth round in August 1997.

The INEEL includes about 890 mi² of the northeastern part of the eastern Snake River Plain

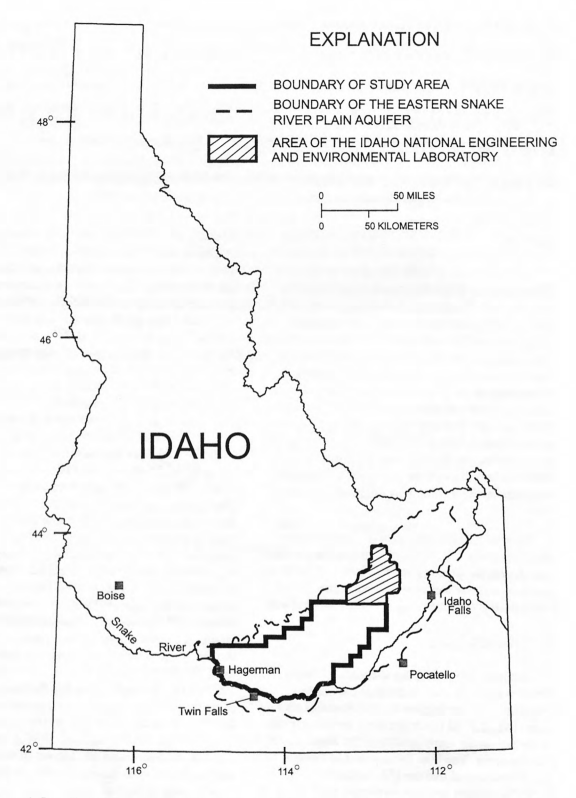


Figure 1. Location of the study area, between the Idaho National Engineering and Environmental Laboratory and Hagerman, Idaho.

and is about 110 mi northeast of the Hagerman area (fig. 1). In the past, wastewater containing radiochemical and chemical wastes generated at the INEEL was discharged mostly to ponds and wells. Since 1983, most aqueous wastes have been discharged to infiltration ponds. Many of the constituents in the wastewater enter the aquifer indirectly by percolation through the unsaturated zone (Pittman and others, 1988).

Chemical and radioactive wastes have migrated from less than 1 to about 9 mi southwest of the disposal areas at the INEEL (Pittman and others, 1988). Tritium was detected intermittently at concentrations of 3,400±200 pCi/L or less in water from three wells along the southern boundary of the INEEL between 1983 and 1985. Since April 1985, tritium concentrations in water from wells near the southern boundary of the INEEL have been less than the reporting level (Bartholomay, Tucker, and others, 1997, p. 27).

Water samples collected from the 18 sites (fig. 2) were analyzed for selected radionuclides, trace elements, common ions, nutrients, purgeable organic compounds, carbamate insecticides, organophosphorus insecticides, gross polychlorinated biphenyls (PCBs), gross polychlorinated naphthalenes (PCNs), chlorophenoxy-acid herbicides, and other herbicides by the USGS National Water Quality Laboratory (NWQL) at Arvada, Colo. Samples also were collected at the same sites for selected radionuclide analyses by the Idaho State University (ISU) Environmental Monitoring Laboratory at Pocatello, Idaho. Two replicate water samples also were collected and analyzed as a measure of quality assurance.

Geohydrologic Setting

The eastern Snake River Plain is a northeast-trending structural basin about 200 mi long and 50 to 70 mi wide. The basin, bounded by faults on the northwest and downwarping and faulting on the southeast, has been filled with basaltic lava flows interbedded with terrestrial sediments (Whitehead, 1986). Individual basalt flows average 20 to 25 ft in thickness with an aggregate thickness of several thousand feet in places. Alluvial fan deposits are composed primarily of sand and gravel, whereas in areas where streams were dammed by

basalt flows, the sediments are predominantly silt and clay (Garabedian, 1986). Rhyolitic lava flows and tuffs are exposed locally at the surface and may exist at depth under most of the eastern plain. A 10,365-ft-deep test hole at the INEEL penetrated about 2,160 ft of basalt and sediment and 8,205 ft of tuffaceous and rhyolitic volcanic rocks (Mann, 1986).

Movement of water in the aquifer generally is from the northeast to the southwest. Water moves horizontally through basalt interflow zones and vertically through joints and interfingering edges of the interflow zones. Infiltration of surface water, heavy pumpage, geologic conditions, and seasonal fluxes of recharge and discharge locally affect the movement of ground water (Garabedian, 1986).

The Snake River Plain aquifer is recharged by seepage from the upper reaches of the Snake River, tributaries and canals, infiltration from irrigation and precipitation, and underflow from tributary valleys on the perimeter of the plain. Discharge from the aquifer primarily is by pumpage for irrigation and spring flow to the Snake River (Mann and Knobel, 1990). Discharge from all of the springs in the eastern Snake River Plain has fluctuated over the years as a result of changes in water use, irrigation practices, and precipitation (Kjelstrom, 1992, p. 2).

Acknowledgments

The authors gratefully acknowledge the well owners for granting permission to collect the water samples and extend special thanks to Downy Strode of the Tikura Cattlemen's Association for his help in locating and pumping one of the wells for sample collection. The authors are grateful for technical review of the manuscript by Scott Anderson of the IDWR and Mary Pace of the USGS.

METHODS AND QUALITY ASSURANCE

The methods used in sampling and analyzing for selected chemicals generally followed the guidelines established by the USGS (Goerlitz and Brown, 1972; Stevens and others, 1975; Wood, 1981; Claassen, 1982; W.L. Bradford, USGS, written commun., 1985; Wershaw and others,

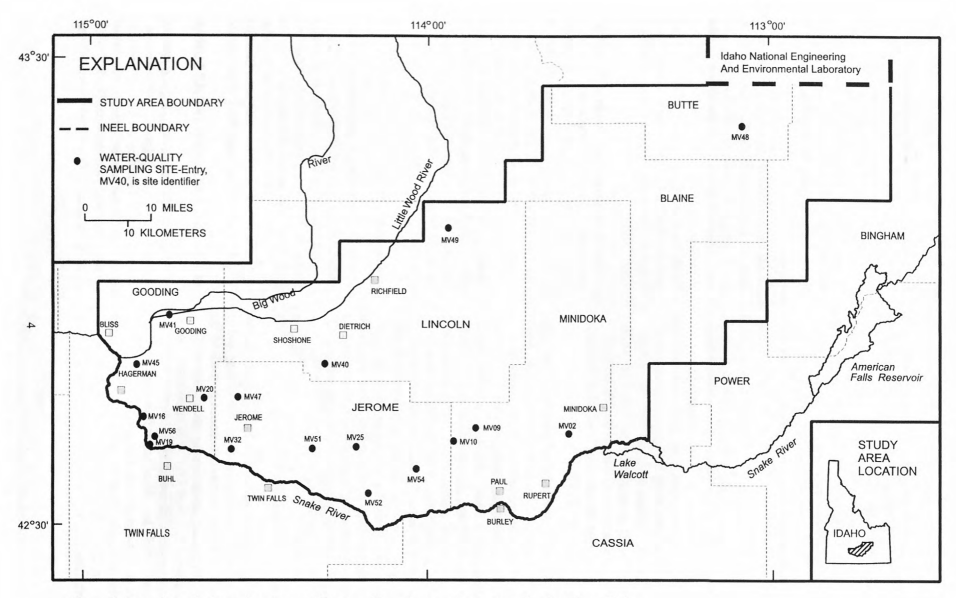


Figure 2. Location of selected water-quality sampling sites on the eastern Snake River Plain.

1987; Fishman and Friedman, 1989; Faires, 1992; and Fishman, 1993). The methods used in the field and quality-assurance practices are described in the following sections.

Site Selection

Water samples were collected at 18 sites (fig. 2), including 7 domestic wells, 6 irrigation wells, 2 springs, 1 dairy well, 1 observation well, and 1 stock well. Two replicate water samples also were collected. The irrigation wells were equipped with turbine pumps. The domestic, dairy, observation, and stock wells were equipped with submersible pumps. Criteria for site selection were geographic location, ease of sample collection, and long-term access.

Sample Containers and Preservatives

Sample containers and preservatives differ depending on the constituent(s) for which analyses are requested. Samples analyzed by the NWQL were collected and preserved in accordance with laboratory requirements specified by Timme (1995). Water samples analyzed by ISU were collected in accordance with laboratory requirements specified by the Director of the Environmental Monitoring Program at ISU. Containers and preservatives were supplied by the respective laboratories. Containers and preservatives used for this study are listed on table 1.

Sample Collection

One of the irrigation wells discharged into a stilling pond and was sampled from the discharge pipe. The remaining irrigation wells were sampled from spigots in discharge lines or from open ports near pumps; domestic, dairy, and stock wells were sampled from spigots closest to pumps; the observation well was sampled at the well head. All the wells either were pumping on arrival of the sampling team or were started on arrival and pumped long enough to ensure that pressure tanks and pumping systems had been thoroughly flushed as evidenced by stable pH, specific conductance, and water-temperature measurements. The two springs were sampled as near the sources as possible by

collecting a grab sample from an area of moving water.

Chemical and physical characteristics monitored at the water-sampling sites included pH, specific conductance, and water temperature. These characteristics were monitored during pumping using methods described by Wood (1981) and Hardy and others (1989). A water sample was collected when measurements of these characteristics indicated probable hydraulic and chemical stability. After collection, sample containers were sealed with laboratory film, labeled, and packed into ice chests for shipment to the NWQL. The samples collected for ISU were stored in coolers until they were hand-delivered to the laboratory.

Field measurements of pH, specific conductance, and water temperature are shown on table 2. Ranges for these measurements were from 7.5 to 8.1 for pH, which is within the U.S. Environmental Protection Agency's (1997) recommended range of 6.5 to 8.5 for community water systems; from 355 to 996 μ S/cm for specific conductance; and from 10.5 to 16.5°C for water temperature.

Conditions at the sampling site during sample collection were recorded in a field logbook; a chain-of-custody record was used to track the samples from the time of collection until delivery to the analyzing laboratory. These records are available for inspection at the USGS Project Office at the INEEL.

Quality Assurance

Detailed descriptions of internal quality control and overall quality-assurance practices used by the NWQL are provided in reports by Friedman and Erdmann (1982), Jones (1987), and Pritt and Raese (1995). Water samples analyzed by the NWQL were collected in accordance with a quality-assurance plan for quality-of-water activities conducted by personnel at the INEEL Project Office. The plan was finalized in June 1989, revised in March 1992 and in 1996 (Mann, 1996), and is available for inspection at the USGS Project Office at the INEEL. Water samples analyzed by ISU were collected in accordance with procedures described by B. Graham (ISU, written commun., 1991). About 10 percent of the water samples were

quality-assurance samples. Sample MV-28 is a replicate of sample MV-32. Sample MV-34 is a replicate of MV-41.

RADIOCHEMICAL CONSTITUENTS

Water samples were analyzed for strontium-90, tritium, gross alpha- and gross beta-particle radioactivity, and gamma-emitting radio-nuclides. The samples were analyzed using methods described by Thatcher and others (1977). Maximum contaminant levels for the types of radioactivity and for selected radionuclides are listed on table 3.

An analytical uncertainty, s, is calculated for each radionuclide concentration or radioactivity measurement. This report presents the analytical uncertainty as 2s. Guidelines for interpreting analytical results are based on an extension of the method described by Currie (1984). In radiochemical analyses, laboratory measurements are made on a target sample and a prepared blank. Instrument signals for the sample and blank vary randomly. Therefore, it is essential to distinguish between two key aspects of the problem of detection: (1) The instrument signal for the sample must be greater than the signal for the blank to make the decision that there was detection; and (2) an estimation must be made of the minimum concentration that will yield a sufficiently large signal to make the correct decision for detection or nondetection most of the time. The first aspect of the problem is a qualitative decision based on signals and a definite criterion for detection. The second aspect of the problem is an estimation of the detection capabilities of a complete measurement process that includes hypothesis testing.

In the laboratory, instrument signals must exceed a critical level of 1.6s to make the qualitative decision whether the radionuclide or radioactivity was detected. At 1.6s, there is about a 95-percent probability that the correct decision—not detected—will be made. Given a large number of samples, as many as 5 percent of the samples with measured concentrations greater than or equal to 1.6s, which were concluded as being detected, might not contain the radioactive constituent. These measurements are referred to as false

positives and are errors of the first kind in hypothesis testing.

Once the critical level of 1.6s has been defined, the minimum detectable concentration can be established. Concentrations that equal 3s represent a measurement at the minimum detectable concentration. For true concentrations of 3s or greater, there is a 95-percent-or-more probability of correctly concluding that the radioactive constituent was detected in a sample. Given a large number of samples, as many as 5 percent of the samples with true concentrations greater than or equal to 3s, which were concluded as being not detected, could contain the radioactive constituent at the minimum detectable concentration. These measurements are referred to as false negatives and are errors of the second kind in hypothesis testing.

True radionuclide concentrations between 1.6s and 3s have larger errors of the second kind. That is, there is a greater-than-5-percent probability of false negative results for samples with true concentrations between 1.6s and 3s, and although the radionuclide or radioactivity might have been detected, such detection may not be considered reliable; at 1.6s, the probability of a false negative is about 50 percent.

These guidelines are based on counting statistics alone and do not include systematic or random errors inherent in laboratory procedures. The values 1.6s and 3s vary slightly with background or blank counts and with the number of gross counts for individual analyses. The use of the critical level and minimum detectable concentration aids the reader in the interpretation of analytical results and does not represent absolute concentrations of radioactivity that may or may not have been detected. In this report, if the concentration of a selected radionuclide was equal to or greater than 3s, the concentration is considered to be above a "reporting level." The reporting level should not be confused with the analytical method detection limit, which is based on laboratory procedures. At small concentrations, the reporting level approaches the analytical method detection limit; however, at larger concentrations, they may be significantly different.

Many analytical results of environmental radioactivity measurements are at or near zero. If the true concentration for a given radionuclide is zero, a given set of analytical results for that radionuclide should be distributed about zero, with an equal number of negative and positive measurements. Negative analytical results occur if the radioactivity of a water sample is less than the background radioactivity or the radioactivity of the prepared blank sample in the laboratory (American Society for Testing and Materials, 1992, p. 126; Knobel and others, 1992, p. 51).

Strontium-90

Strontium-90 is a fission product that was widely distributed in the environment during atmospheric weapons tests. Strontium-90 generally is present in ground water as a result of these tests and from nuclear industry waste-disposal practices. No concentration of dissolved strontium-90 exceeded the reporting level for the water samples analyzed (table 4).

Tritium

Tritium, a radioactive isotope of hydrogen, is formed in nature by interactions of cosmic rays with gases in the upper atmosphere. Tritium also is produced in thermonuclear detonations and is a waste product of the nuclear power industry. Samples were submitted to the ISU laboratory and the NWQL. The ISU laboratory used two techniques: a standard liquid-scintillation technique and an enrichment and liquid-scintillation technique. The NWOL used an enrichment and gas-counting technique. The analytical method detection limit for the laboratories differed. The analytical method detection limits for the ISU laboratory were 150 and less than 25 pCi/L using ten 20-minute counting periods, and that for the NWQL was 1 pCi/L using a 1,000-minute counting period.

The concentrations of tritium in the water samples are shown on table 4. Concentrations of tritium in all of the water samples analyzed by the NWQL were greater than the reporting level and ranged from 7.7 ± 1.0 to 68.8 ± 4.5 pCi/L (table 4). Concentrations of tritium in 18 of the 20 water samples analyzed by the ISU laboratory using the enrichment technique were equal to or greater than

the reporting level and ranged from 12±8 to 76±8 pCi/L. Two of the samples (MV-9, MV-28) had split samples analyzed. For the purpose of comparison, background concentrations of tritium in ground water in Idaho generally range from 0 to 40 pCi/L (Knobel and others, 1992). The maximum contaminant level for tritium in public drinking-water supplies is 20,000 pCi/L (table 3).

Gross Alpha-Particle Radioactivity

Gross alpha-particle radioactivity is a measure of the total radioactivity given off as alpha particles during the radioactive decay process; however, laboratories normally report the radioactivity as if it were all given off by one radionuclide. In this report, concentrations are reported as dissolved thorium-230 in picocuries per liter by the NWQL, and as total americium-241 in picocuries per liter by the ISU laboratory.

The concentration of gross alpha-particle radioactivity reported as dissolved thorium-230 in one of the water samples analyzed by the NWQL (MV-20) was greater than the reporting level (table 5) and was 5.5±3.0 pCi/L. Total concentrations of gross alpha-particle radioactivity reported as americium-241 in two water-sample splits analyzed by ISU (MV-47, MV-48) were greater than the reporting level (table 5).

Gross Beta-Particle Radioactivity

Gross beta-particle radioactivity is a measure of the total radioactivity given off as beta particles during the radioactive decay process; however, laboratories normally report the radioactivity as if it were all given off by one radionuclide. In this report, concentrations are reported as dissolved cesium-137 in picocuries per liter by the NWQL, and as total cesium-137 in picocuries per liter by the ISU laboratory. The average annual concentrations of strontium-90 and cesium-137 in public drinking-water supplies that produce a 4-mrem/yr dose are 8 pCi/L and 120 pCi/L, respectively. Gross beta-particle radioactivity measurements should not be compared directly with these concentrations.

Concentrations of gross beta-particle radioactivity reported as dissolved cesium-137 in 13 of

the water samples analyzed by the NWQL were greater than the reporting level (table 5) and ranged from 7.0 ± 4.2 to 11.5 ± 4.4 pCi/L. Concentrations of gross beta-particle radioactivity reported as total cesium-137 in 18 of the water samples analyzed by ISU were equal to or greater than the reporting level (table 5) and ranged from 1.2 ± 0.8 to 6.5 ± 1.9 pCi/L.

Cesium-137 and Potassium-40

Gamma spectrometry involves using a series of detectors to simultaneously determine the concentrations of a variety of radionuclides by the identification of their characteristic gamma emissions. When no specific gamma-emitting radionuclides are identified, the concentration is reported by ISU as total cesium-137. In addition, as part of a special request, concentrations also were reported as potassium-40. Cesium-137 is a fission product of uranium-235, uranium-233, or plutonium-239. Concentrations of total cesium-137 in all water samples were less than the reporting level (table 6). The concentration of potassium-40 in one sample (MV-32) was greater than the reporting level (table 6).

CHEMICAL CONSTITUENTS

Water samples were analyzed for selected chemical constituents. These constituents included trace elements, common ions, nutrients, purgeable organic compounds, insecticides, polychlorinated compounds, and herbicides. In this report, minimum reporting levels and method detection limits established for these constituents are not to be confused with reporting levels and analytical method detection limits for selected radionuclides. The minimum reporting level for inorganic and organic constituents is the smallest measured concentration of a constituent that may be reliably reported using a given analytical method (Timme, 1995, p. 92). The method detection limit is defined as the minimum concentration of a substance that can be identified, measured, and reported with 99-percent confidence that the analyte concentration is greater than zero and is determined from analysis of a sample in a given matrix containing the analyte (Timme, 1995, p. 92).

Trace Elements

Water samples were analyzed for selected dissolved trace elements including arsenic, barium, beryllium, cadmium, chromium, cobalt, copper, iron, lead, lithium, manganese, mercury, molybdenum, nickel, selenium, silver, strontium, vanadium, and zinc. The maximum or secondary maximum contaminant levels and minimum reporting levels of selected trace elements are shown on table 7. The concentrations of dissolved trace elements are shown on table 8.

<u>Arsenic.</u>—Concentrations of arsenic in 18 samples were greater than the minimum reporting level and ranged from 1 to 6 μ g/L. The maximum contaminant level is 50 μ g/L.

Barium.—Concentrations of barium in all samples were greater than the minimum reporting level and ranged from 18 to 139 μ g/L. The maximum contaminant level is 2,000 μ g/L.

Beryllium.—Concentrations of beryllium in all the samples were less than the minimum reporting level of 0.5 $\mu g/L$. The maximum contaminant level is 4 $\mu g/L$.

<u>Cadmium.</u>—The concentration of cadmium in one sample was 1 μ g/L; concentrations in the other samples were less than the minimum reporting level. The maximum contaminant level is 5 μ g/L.

<u>Chromium.</u>—Concentrations of chromium in all samples were less than the minimum reporting level of 5 μ g/L. The maximum contaminant level is 100 μ g/L.

<u>Cobalt.</u>—Concentrations of cobalt in all samples were less than the minimum reporting level of 3 μ g/L.

<u>Copper.</u>—Concentrations of copper in all samples were less than the minimum reporting level of $10 \mu g/L$. The secondary maximum contaminant level is $1,000 \mu g/L$.

<u>Iron.</u>—Concentrations of iron in five samples were greater than the minimum reporting level and ranged from 5 to 72 μ g/L. The secondary maximum contaminant level is 300 μ g/L.

<u>Lead.</u>—Concentrations of lead in 11 samples were greater than the minimum reporting level and

ranged from 12 to 34 μ g/L. Lead has an action level of 15 μ g/L.

<u>Lithium.</u>—Concentrations of lithium in 19 samples were greater than the minimum reporting level and ranged from 5 to 50 µg/L.

<u>Manganese.</u>—Concentrations of manganese in three samples were greater than the minimum reporting level and ranged from 1 to 9 μ g/L. The secondary maximum contaminant level is 50 μ g/L.

Mercury.—The concentration of mercury in one sample (MV-02) was equal to the minimum reporting level of $0.1 \mu g/L$. The maximum contaminant level is $2 \mu g/L$.

Molýbdenum.—Concentrations of molybdenum in two samples (MV-20, MV-47) were 11 and 14 μ g/L, respectively; concentrations in the other samples were less than the minimum reporting level of 10 μ g/L.

Nickel.—Concentrations of nickel in all samples were less than the minimum reporting level of $10 \mu g/L$. The maximum contaminant level is $100 \mu g/L$.

<u>Selenium.</u>—Concentrations of selenium in four samples were equal to the minimum reporting level of 1 μ g/L; concentrations in the other samples were less than the minimum reporting level. The maximum contaminant level is 50 μ g/L.

<u>Silver.</u>—Concentrations of silver in all samples were less than the minimum reporting level of 1 μ g/L. The secondary maximum contaminant level is 100 μ g/L.

Strontium.—Concentrations of strontium in all samples were greater than the minimum reporting level and ranged from 168 to 443 μ g/L.

<u>Vanadium.</u>—Concentrations of vanadium in 14 samples were equal to or greater than the minimum reporting level and ranged from 6 to $17~\mu g/L$.

Zinc.—Concentrations of zinc in 13 samples were equal to or greater than the minimum reporting level and ranged from 3 to 355 μ g/L. The secondary maximum contaminant level is 5,000 μ g/L.

Common lons

Water samples were analyzed for dissolved common ions including calcium, magnesium, silica, and sodium. The minimum reporting levels of these ions are shown on table 9. Maximum contaminant levels have not been established for any of these common ions. The concentrations of dissolved common ions are shown on table 10.

<u>Calcium</u>.—Concentrations of calcium in all samples were greater than the minimum reporting level and ranged from 31 to 72 mg/L.

Magnesium.—Concentrations of magnesium in all samples were greater than the minimum reporting level and ranged from 13 to 33 mg/L.

<u>Silica.</u>—Concentrations of silica in all samples were greater than the minimum reporting level and ranged from 23 to 43 mg/L.

Sodium.—Concentrations of sodium in all samples were greater than the minimum reporting level and ranged from 7.4 to 77 mg/L. The Idaho Department of Health and Welfare (1989) recommends an optimum concentration of 20 mg/L of sodium for public drinking-water supplies.

Nutrients

Water samples were analyzed for dissolved ammonia (as nitrogen), nitrite (as nitrogen), nitrite plus nitrate (as nitrogen), and orthophosphate (as phosphorus). The maximum contaminant levels and the minimum reporting levels are shown on table 9. A maximum contaminant level has not been established or proposed for ammonia or orthophosphate. Concentrations of nutrients are shown on table 10.

Ammonia (as nitrogen).—Concentrations of ammonia (as nitrogen) in 2 of the samples (MV-49, MV-51) were 0.018 and 0.015 mg/L, respectively; concentrations in the other samples were less than the minimum reporting level.

Nitrite (as nitrogen).—Concentrations of nitrite (as nitrogen) in two of the samples were equal to the minimum reporting level of 0.01 mg/L; concentrations in the other samples were less than the minimum reporting level. The maximum contaminant level is 1 mg/L.

Nitrite plus nitrate (as nitrogen).— Concentrations of nitrite plus nitrate (as nitrogen) in all samples were greater than the minimum reporting level and ranged from 0.64 to 6.7 mg/L. The maximum contaminant level is 10 mg/L.

Orthophosphate (as phosphorus).—Concentrations of orthophosphate (as phosphorus) in 19 of the samples were equal to or greater than the minimum reporting level and ranged from 0.01 to 0.10 mg/L.

Purgeable Organic Compounds

Water samples were analyzed for 60 purgeable organic compounds. The maximum contaminant levels and minimum reporting levels of these compounds are shown on table 11. Two of the compounds (1,2-Dibromo-3-chloropropane and 1,2,-Dibromoethane) had maximum contaminant levels less than their minimum reporting levels. Concentrations of purgeable organic compounds in all the samples were less than their respective minimum reporting levels.

Insecticides and Gross Polychlorinated Compounds

Water samples were analyzed for concentrations of 11 carbamate insecticides, 11 organophosphate insecticides, 15 organochlorine insecticides, gross PCBs, and gross PCNs (table 12). The minimum reporting levels ranged from 0.007 to 1.0 µg/L. Water samples also were analyzed for an additional group of 24 insecticides that included 11 of the carbamate, organophosphate, and organochlorine insecticides mentioned above; therefore, 11 insecticides are listed twice and minimum reporting levels may be different. The maximum contaminant levels and minimum reporting levels for these compounds are shown on table 12. None of the samples contained concentrations of insecticides or polychlorinated compounds greater than their respective minimum reporting levels.

Herbicides

Water samples were analyzed for concentrations of 5 chlorophenoxy-acid herbicides and 47 other herbicides. Maximum contaminant levels

and minimum reporting levels for these compounds are shown on table 13. Because new laboratory schedules with lower method detection limits and minimum reporting levels than past schedules were used, concentrations of some herbicides in several samples exceeded the method detection limits or minimum reporting levels (table 14). Concentrations in some of the samples listed in table 14 exceeded the method detection limits but were less than the minimum reporting levels. One sample (MV-19) contained an estimated concentration of alachlor of 0.002 µg/L. The estimated concentrations in table 14 need to be evaluated carefully because of variable performance (Zaugg and others, 1995). Estimated and actual concentrations of atrazine in 16 samples ranged from 0.001 to 0.018 µg/L. One sample (MV-19) contained an estimated concentration of bentazon of 0.009 µg/L. Estimated concentrations of desethyl atrazine in 19 samples ranged from 0.001 to 0.026 µg/L. The analyses for desethyl atrazine demonstrate low recovery because of poor retention on the solid-phase extraction column (Zaugg and others, 1995). Actual and estimated concentrations of metribuzin in two samples (MV-20 and MV-45) were 0.006 and 0.002 µg/L. respectively. The estimated concentration of prometon in two samples (MV-34 and MV-41) were 0.003 and 0.004 µg/L, respectively. Estimated and actual concentrations of simazine in three samples ranged from 0.002 to 0.006 µg/L. Concentrations of herbicides not listed in table 14 were less than the minimum reporting levels and method detection limits in all the samples.

SUMMARY

The USGS and the IDWR, in cooperation with the DOE, sampled 18 sites as part of the fourth round of a long-term project to monitor water quality of the Snake River Plain aquifer from the southern boundary of the Idaho National Engineering and Environmental Laboratory to the Hagerman area. Water samples were collected and analyzed for selected radiochemical and chemical constituents. The samples were collected from seven domestic wells, six irrigation wells, two springs, one dairy well, one observation well and one stock well. Two quality-assurance samples also were collected and analyzed.

Concentrations of strontium-90 in all samples were less than the reporting level. Concentrations of tritium in all of the samples analyzed by the NWOL and 18 of the samples analyzed by ISU using an enrichment technique were equal to or greater than the reporting level, but none exceeded the maximum contaminant level for drinking water. The concentration of gross alpha-particle radioactivity reported as dissolved thorium-230 in one sample analyzed by the NWOL was greater than the reporting level; it did not exceed the maximum contaminant level. Concentrations of gross beta-particle radioactivity reported as dissolved cesium-137 in 13 samples analyzed by the NWOL were greater than the reporting level. Concentrations of gross beta-particle radioactivity reported as total cesium-137 in 18 of the samples analyzed by ISU were equal to or greater than the reporting level. Concentrations of total cesium-137 were analyzed using gamma spectrometry and concentrations in all the samples were less than the reporting level. Concentrations of total potassium-40, also analyzed using gamma spectrometry, were less than the reporting level except in one sample.

All the samples contained one or more dissolved trace elements and common ions in concentrations greater than the minimum reporting levels. No concentration exceeded an established maximum contaminant level.

Concentrations of ammonia (as nitrogen) in two of the water samples were equal to or greater than the minimum reporting level. Concentrations of nitrite (as nitrogen) in two samples were equal to the minimum reporting level. Concentrations of nitrite plus nitrate (as nitrogen) in all the water samples were greater than the minimum reporting level. Concentrations of orthophosphate (as phosphorus) in 19 of the water samples were equal to or greater than the minimum reporting level. No nutrient concentration exceeded an established maximum contaminant level.

Concentrations of purgeable organic compounds, carbamate insecticides, organophosphate insecticides, organochlorine insecticides, gross PCBs, and gross PCNs in all samples were less than their respective minimum reporting levels. Concentrations of some herbicides were greater than their method detection limits and minimum

reporting levels, but none exceeded established maximum contaminant levels.

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Table 1. Containers and preservatives used for water-sample collection [Abbreviations: L, liter; mL, milliliter; μm, micrometer; gal, gallon; °C, degrees Celsius. Chemical formulas: HNO₃, nitric acid; K₂Cr₂O₇, potassium dichromate. Chilled samples were shipped by overnight-delivery mail. Analyzing laboratory: NWQL, U.S. Geological Survey National Water Quality Laboratory; ISU, Idaho State University Environmental Monitoring Laboratory]

C	Container		Preservati	ive			
Constituent or type of constituent	Туре	Size	Туре	Size	Other treatment	Analyzing laborator	
Strontium-90	Polyethylene, acid rinsed	1 L	HNO ₃	4 mL	0.45-μm filter	NWQL	
Tritium	Polyethylene	1 L	None	None	None	NWQL	
	Polyethylene	1 L	None	None	None	ISU	
Other radionuclides	Polyethylene, acid rinsed	1 L	HNO ₃	4 mL	.45-μm filter	NWQL	
	Polyethylene	1 gal	None	None	None	ISU	
	Polyethylene	1 L	HNO ₃	5 mL	.45-μm filter	ISU	
Trace elements	Polyethylene, acid rinsed	250 mL	HNO ₃	1 mL	.45-μm filter	NWQL	
	Polyethylene	125 mL	None	None	None	NWQL	
Mercury	Glass, acid rinsed	250 mL	HNO ₃ /K ₂ Cr ₂ O ₇	10 mL	.45-μm filter	NWQL	
Common ions	Polyethylene, acid rinsed	250 mL	HNO ₃	1 mL	.45-μm filter	NWQL	
Nutrients	Polyethylene, brown	125 mL	None	None	.45-μm filter, chill 4°C	NWQL	
Purgeable organic compounds	Glass, baked	40 mL	None	None	Chill 4°C	NWQL	
Pesticides	Glass, baked	1 L	None	None	Chill 4°C	NWQL	

Table 2. Results of field measurements of water for pH, specific conductance, and temperature from selected wells and springs, eastern Snake River Plain

[See figure 2 for location of sites. Site type: H, domestic; I, irrigation; Sp, spring; QA, quality assurance (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41); D, dairy; O, observation; S, stock. Date sampled: month/day/year. Units: pH, negative base-10 logarithm of hydrogen-ion activity in moles per liter; specific conductance, microsiemens per centimeter at 25°C (degrees Celsius); temperature, °C]

Site identifier	Site type	Date sampled	pН	Specific conductance	Temperature
MV-02	Н	8/26/97	7.8	630	14.0
MV-09	I	8/26/97	7.7	996	13.5
MV-10	I	8/26/97	7.8	760	13.5
MV-16	Sp	8/27/97	7.9	413	14.5
MV-19	Sp	8/27/97	7.9	524	14.5
MV-20	I	8/25/97	7.8	434	15.0
MV-25	H	8/26/97	7.7	668	15.0
MV-28	QA	8/27/97	7.8	732	15.0
MV-32	Н	8/27/97	7.8	732	15.0
MV-34	QA	8/25/97	7.5	675	14.0
MV-40	I	8/25/97	8.0	374	14.5
MV-41	I	8/25/97	7.5	675	14.0
MV-45	I	8/25/97	8.0	398	16.0
MV-47	D	8/27/97	8.0	376	15.0
MV-48	0	8/28/87	8.1	355	12.0
MV-49	S	8/28/97	7.7	361	10.5
MV-51	Н	8/25/97	7.8	687	14.5
MV-52	Н	8/26/97	7.8	626	16.5
MV-54	Н	8/26/97	7.8	833	14.5
MV-56	Н	8/27/97	7.9	425	14.5

Table 3. Maximum contaminant levels for types of radioactivity and selected radionuclides in drinking water [The maximum contaminant levels were established pursuant to the recommendations of the U.S. Environmental Protection Agency (1997, p. 296) for community water systems and are included for comparison purposes only. Maximum contaminant levels given for strontium-90 and tritium are average annual concentrations assumed to produce a total body or organ dose of 4 mrem/yr (milliremper year) of beta-particle radiation. The maximum contaminant level given for gross alpha-particle radioactivity includes radioactivity excludes radioactivity from natural sources and is included for comparison purposes only. Abbreviation: pCi/L, picocurie per liter]

Type of radioactivity or radionuclide	Maximum contaminant level
Strontium-90	8 pCi/L
Tritium	20,000 pCi/L
Gross alpha-particle radioactivity	15 pCi/L
Gross beta-particle and gamma radioactivity	4 mrem/yr

Table 4. Concentrations of strontium-90 and tritium in water from selected wells and springs, eastern Snake River Plain

[See figure 2 for location of sites. Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory (NWQL) and Idaho State University (ISU) Environmental Monitoring Laboratory using the following methods: strontium-90 by chemical separation and precipitation; tritium by enrichment and gas counting at the NWQL and by liquid scintillation at ISU. Analytical results and uncertainties—for example 0.15±0.22—in picocuries per liter. Analytical uncertainties are reported as 2s. Concentrations that exceed the reporting level of 3 times 1s are shown in boldface type. Symbol: *, quality-assurance sample (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41). ISU tritium values were determined using two different detection methods. Two values for ISU tritium and ISU tritium (enriched) represent laboratory split samples]

Site identifier	Strontium-90, NWQL	Tritium, NWQL	Tritium, ISU	Tritium, ISU (Enriched)
MV-02	0.15±0.22	39.0±2.6	-30±90	47±8
MV-09	.25±0.26	68.8±4.5	10±90; 60±90	71±9; 76±8
MV-10	.27±0.26	38.7±2.6	-20±90	42±7
MV-16	.06±0.24	9.6±1.0	-30±90	8±8
MV-19	.27±0.26	12.5±1.3	20±90	13±7
MV-20	.32±0.26	16.6±1.3	-10±90	14±7
MV-25	.31±0.27	53.1±3.2	-30±90	53±8
MV-28*	.26±0.26	54.1±3.2	70±90	52±9; 52±9
MV-32	.45±0.35	51.8±3.2	50±90	56±8
MV-34*	.31±0.30	47.4±3.2	10±90	53±9
MV-40	.24±0.27	13.1±1.3	20±90	12±8
MV-41	.21±0.27	45.1±2.6	0±90	55±9
MV-45	.32±0.29	18.9±1.6	-80±90	16±7
MV-47	.35±0.27	7.7±1.0	30±90	8±7
MV-48	.26±0.24	31.0±1.9	60±90	31±8
MV-49	.27±0.26	58.6±3.8	40±90	57±8
MV-51	.27±0.26	46.4±3.2	30±90	49±8
MV-52	.26±0.24	57.6±3.8	-30±90	61±8
MV-54	.24±0.26	58.6±3.8	-50±90	65±8
MV-56	.37±0.28	7.7±1.0	-20±90	13±8

Table 5. Concentrations of gross alpha-particle and gross beta-particle radioactivity in water from selected wells and springs, eastern Snake River Plain

[See figure 2 for location of sites. Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory (NWQL) and the Idaho State University (ISU) Environmental Monitoring Laboratory using a residue procedure. Analytical results and uncertainties—for example 3.4±2.9—in piccouries per liter. Analytical uncertainties are reported as 2 s. Concentrations that equal or exceed the reporting level of 3 times 1s are shown in boldface type. Symbol: *, quality-assurance sample (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41). Two or three values for ISU gross alpha and gross beta analyses represent laboratory split samples]

Site identifier	Gross alpha, as dissolved thorium-230,		alpha, as m-241, ISU	Gross beta, as dissolved cesium-137,	Gross beta, as cesium-137, ISU		
identifier	NWQL	Filtered	Raw	NWQL	Filtered	Raw	
MV-02	3.4±2.9	-2.4±3.6	08±3.9	11.1±4.3	2.4±1.6	5.1±1.7	
MV-09	3.9±3.6	3.4±4.2; .4±5.5	-4.9±4.7; 1.3±5.2	10.2±4.4	6.2±1.8; 5.4±2.7	6.4±1.8 4.5±2.6	
MV-10	2.3±2.7	-1.1±3.9	9±4.2	7.1±4.2	4.8±1.7	6.5±1.9	
MV-16	1.1±2.1	4±3.0	4±2.4	7.6±4.1	2.5±1.5	2.4±1.4	
MV-19	1.4±2.3	-1.2±3.4	-1.3±2.6	3.2±3.9	2.6±1.6	2.2±1.5	
MV-20	5.5±3.0	2.0±2.1; .9±3.4	-1.6±3.2	7.4±4.1	2.3±1.2; 2.9±1.3	3.4±1.4	
MV-25	3.9±3.1	6.7±4.0	8±3.2	11.5±4.4	1.6±1.4	6.0±1.7	
MV-28*	4.1±3.1	3.0±3.2	3.9±4.5; -2.6±5.0	7.1±4.2	2.0±1.4	4.6±1.8 ; 2.4±2.7	
MV-32	3.9±3.1	9±3.6	1.4±4.0	7.5±4.2	4.7±1.8	3.5±1.7	
MV-34*	1.6±2.5	-3.2±4.2	-1.6±3.7	7.1±4.2	3.6±1.5	5.0±1.5	
MV-40	2.6±2.3	2.3±2.5; 1.0±2.2	1.6±2.7	5.4±4.0	3.4±1.3; 3.1±1.3	3.4±1.4	
MV-41	4.3±3.2	2.7±3.4	3.8±3.6	7.0 ± 4.2	4.0±1.5	4.0±1.5	
MV-45	2.1±2.2	-3.2±2.1	9±2.8	4.0±3.9	2.6±1.2	3.3±1.3	
MV-47	.80±1.9	-1.6±2.7	5.1±2.8 ; 3.8±2.8	3.6±3.9	1.5±1.4	2.6±1.4 2.4±1.4	
MV-48	3.3±2.5	.3±1.8	9.1±3.9; 1±1.6	3.6±3.9	1.9±1.3	2.3±1.5 1.2±0.8	
MV-49	2.8±2.4	6.6±3.3	9±1.6	4.2±3.9	1.3±1.4	1.8±1.3	
MV-51	3.2±3.0	-4.0±3.8	.5±4.5; 2.6±3.8	11.2±4.4	6.4±1.6	5.8±1.6; 5.9±1.7	
MV-52	2.8±2.8	4.0±3.8; 9.7±5.0; 21.1±3.4	.3±4.4	8.4±4.2	4.0±1.6; 4.5±1.7	4.7±1.0; 6.1±1.6	
MV-54	4.4±3.5	1.5±3.2	-1.2±4.2	8.4±4.3	1.6±1.4	5.2±1.8	
MV-56	2.1±2.3	2.8±2.7	2±2.4	3.8±3.9	4.2±1.4	2.5±1.4	

Table 6. Concentrations of cesium-137 and potassium-40 in water from selected wells and springs, eastern Snake River Plain

[See figure 2 for location of sites. Analyses were performed by the Idaho State University Environmental Monitoring Laboratory using gamma spectrometry. Analytical results and uncertainties—for example -2.5±2.8—are for total concentrations in picocuries per liter. Analytical uncertainties are reported as 2s.Concentration that exceeds the reporting level of 3 times 1s is shown in boldface type. Symbol: *, quality-assurance sample (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41). Two values represent laboratory split samples]

Site identifier	Cesium-137	Potassium-40
MV-02	-2.5±2.8	-51±47
MV-09	-3.3±2.7	-10±49
MV-10	.3±3.2	-16±60
MV-16	.1±3.3	74±64
MV-19	-2.5±3.2	-5±54
MV-20	9±3.0	-30±50
MV-25	1.0±3.2	29±59
MV-28*	-3.5±2.8	-33±50
MV-32	.2±3.2	54±27
MV-34*	.1±3.5	58±56
MV-40	-1.6±2.2	-21±46
MV-41	.3±3.2	48±65
MV-45	.8±3.3	-16±63
MV-47	-1.4±2.7; -2.1±2.6	-21±50; -13±51
MV-48	-1.2±2.6	-15±49
MV-49	1.6±2.7	23±65
MV-51	-2.0±3.3; -1.3±2.6	31±66; 11±60
MV-52	1.1±3.4	-43±58
MV-54	1.9±3.1	17±60
MV-56	2±3.5	2±60

Table 7. Maximum or secondary maximum contaminant levels and minimum reporting levels of selected trace elements in drinking water

[The maximum contaminant levels are for total measurements and were established pursuant to the recommendations of the U.S. Environmental Protection Agency (1994; 1997, p. 295, 360) for community water systems and are for comparison purposes only. Secondary maximum contaminant levels—in brackets—are from U.S. Environmental Protection Agency (1997, p. 474). Minimum reporting levels are from Timme (1995). Units are in micrograms per liter (µg/L). Symbols: ••, maximum contaminant level has not been established; *, lead has an action level of 15 µg/L]

Trace element	Maximum or secondary maximum contaminant level	Minimum reporting level
Arsenic	50	1
Barium	2,000	1
Beryllium	4	.5
Cadmium	5	1
Chromium	100	5
Cobalt		3
Copper	[1,000]	10
Iron	[300]	3
Lead	*	10
Lithium	••	4
Manganese	[50]	10
Mercury	2	.1
Molybdenum		10
Nickel	100	10
Selenium	50	1
Silver	[100]	1
Strontium		.5
Vanadium		6
Zinc	[5,000]	3

Table 8. Concentrations of dissolved trace elements in water from selected wells and springs, eastern Snake River Plain
[See figure 2 for location of sites. Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory. Analytical results in micrograms per liter. Symbols: <, concentration is less than the respective minimum reporting level; *, quality-assurance sample (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41)]

Site identifier	Arsenic	Barium	Beryllium	Cadmium	Chromium	Cobalt	Copper	Iron	Lead	Lithiun
MV-02	3	67	<0.5	<1	<5	<3	<10	6	28	41
MV-09	4	139	<.5	<1	<5	<3	<10	<3	12	50
MV-10	3	79	<.5	<1	<5	<3	<10	<3	<10	46
MV-16	2	22	<.5	<1	<5	<3	<10	<3	15	24
MV-19	2	34	<.5	<1	<5	<3	<10	<3	34	32
MV-20	2	21	<.5	<1	<5	<3	<10	<3	12	20
MV-25	4	60	<.5	<1	<5	<3	<10	<3	<10	23
MV-28*	1	87	<.5	<1	<5	<3	<10	<3	<10	39
MV-32	2	87	<.5	<1	<5	<3	<10	<3	30	38
MV-34*	3	63	<.5	<1	<5	<3	<10	<3	<10	9
MV-40	3	19	<.5	<1	<5	<3	<10	7	33	13
MV-41	3	64	<.5	<1	<5	<3	<10	<3	14	9
MV-45	2	23	<.5	<1	<5	<3	<10	<3	<10	16
MV-47	2	18	<.5	<1	<5	<3	<10	<3	31	23
MV-48	<1	51	<.5	<1	<5	<3	<10	6	18	<4
MV-49	<1	49	<.5	<1	<5	<3	<10	72	<10	5
MV-51	3	61	<.5	<1	<5	<3	<10	5	33	37
MV-52	6	76	<.5	1	<5	<3	<10	<3	<10	27
MV-54	2	99	<.5	<1	<5	<3	<10	<3	<10	48
MV-56	2	24	<.5	<1	<5	<3	<10	<3	<10	27

Table 8. Concentrations of dissolved trace elements in water from selected wells and springs, eastern Snake River Plain—Continued

Site identifier	Manganese	Mercury	Molybdenum	Nickel	Selenium	Silver	Strontium	Vanadium	Zinc
MV-02	<1	0.1	<10	<10	<1	<1	328	<6	50
MV-09	<1	<.1	<10	<10	1	<1	434	6	<3
MV-10	<1	<.1	<10	<10	1	<1	382	<6	<3
MV-16	<1	<.1	<10	<10	<1	<1	206	9	10
MV-19	<1	<.1	<10	<10	<1	<1	262	7	<3
MV-20	1	<.1	11	<10	<1	<1	194	9	3
MV-25	<1	<.1	<10	<10	<1	<1	238	17	125
MV-28*	<1	<.1	<10	<10	<1	<1	381	7	27
MV-32	<1	<.1	<10	<10	<1	<1	380	<6	27
MV-34*	<1	<.1	<10	<10	<1	<1	334	12	<3
MV-40	<1	<.1	<10	<10	<1	<1	172	7	7
MV-41	<1	<.1	<10	<10	<1	<1	333	11	<3
MV-45	2	<.1	<10	<10	<1	<1	205	9	<3
MV-47	<1	<.1	14	<10	<1	<1	183	9	7
MV-48	<1	<.1	<10	<10	1	<1	234	<6	95
MV-49	9	<.1	<10	<10	<1	<1	168	<6	355
MV-51	<1	<.1	<10	<10	<1	<1	325	9	23
MV-52	<1	<.1	<10	<10	<1	<1	259	15	40
MV-54	<1	<.1	<10	<10	1	<1	443	<6	21
MV-56	<1	<.1	<10	<10	<1	<1	207	7	<3

Table 9. Maximum contaminant levels and minimum reporting levels of selected common ions and nutrients in drinking water

[The maximum contaminant levels are for total measurements and were established pursuant to the recommendations of the U.S. Environmental Protection Agency (1997, p. 360) for community water systems and are for comparison purposes only. Minimum reporting levels are from Timme (1995). Units are in milligrams per liter (mg/L). Symbol: ••, maximum contaminant level has not been established]

Constituent	Maximum contaminant level	Minimum reporting level
Calcium	••	0.02
Magnesium	••	.01
Silica	••	.01
Sodium ¹	••	.2
Ammonia (as nitrogen)	••	.015
Nitrite (as nitrogen)	1	.01
Nitrite plus nitrate (as nitrogen)	10	.05
Orthophosphate (as phosphorus)	••	.01

¹Idaho Department of Health and Welfare (1989) recommends an optimum concentration of 20 mg/L for public drinking-water supplies.

Table 10. Concentrations of dissolved common ions and nutrients in water from selected wells and springs, eastern Snake River Plain

[See figure 2 for location of sites. Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory. Analytical results in milligrams per liter. Symbols: <, concentration is less than the respective minimum reporting level; *, quality-assurance sample (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41)]

Site identifier	Calcium	Mag- nesium	Silica	Sodium	Ammonia (as nitrogen)	Nitrite (as nitrogen)	Nitrite plus nitrate (as nitrogen)	Orthophosphate (as phosphorus)
MV-02	52	20	36	40	< 0.015	< 0.01	1.5	0.02
MV-09	72	33	35	77	<.015	<.01	6.7	.03
MV-10	60	24	34	51	<.015	.01	2.7	.01
MV-16	35	17	33	20	<.015	<.01	1.6	.02
MV-19	45	20	33	24	<.015	<.01	3.3	.02
MV-20	34	18	32	23	<.015	<.01	2.0	.02
MV-25	54	25	36	48	<.015	<.01	3.2	.10
MV-28*	65	26	37	41	<.015	<.01	3.7	.01
MV-32	64	26	37	41	<.015	<.01	3.7	.01
MV-34*	61	26	38	37	<.015	<.01	3.1	.08
MV-40	32	16	32	18	<.015	<.01	1.2	.02
MV-41	62	26	39	36	<.015	.01	3.3	.07
MV-45	34	17	32	18	<.015	<.01	1.3	.02
MV-47	31	15	32	18	<.015	<.01	.89	.01
MV-48	41	14	23	8.3	<.015	<.01	.64	<.01
MV-49	45	13	25	7.4	.018	<.01	1.8	.02
MV-51	59	23	33	42	.015	<.01	2.4	.05
MV-52	47	21	43	49	<.015	<.01	2.0	.02
MV-54	72	27	36	50	<.015	<.01	4.6	.03
MV-56	37	17	32	21	<.015	<.01	1.2	.02

Table 11. Maximum contaminant levels and minimum reporting levels of selected purgeable organic compounds in drinking water

[Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory using an analytical method equivalent to U.S. Environmental Protection Agency method 524.2. Maximum contaminant levels were established pursuant to the recommendations of the U.S. Environmental Protection Agency (1994; 1997, p. 358) for community water systems and are included for comparison purposes only. Minimum reporting levels are from Timme (1995). Units are in micrograms per liter (µg/L). Symbols: ••, maximum contaminant level has not been established or proposed; *, total trihalomethanes—which include bromoform, chlorodibromomethane, chloroform, and dichlorobromomethane—in community water systems serving 10,000 or more persons cannot exceed 100 µg/L (U.S. Environmental Protection Agency, 1997, p. 295). Abbreviations: MCL, maximum contaminant level; MRL, minimum reporting level]

Compound	MCL	MRL	Compound	MCL	MRL	
Benzene	5	0.2	1,3-Dichloropropane	••	0.2	
Bromobenzene	••	.2	2,2-Dichloropropane	••	.2	
Bromochloromethane		.2	cis-1,3-Dichloropropene	••	.2	
Bromoform	*	.2	trans-1,3-Dichloropropene		.2	
Bromomethane		.2	1,1-Dichloropropene	••	.2	
n-Butylbenzene		.2	Ethylbenzene	700	.2	
sec-Butylbenzene	••	.2	Hexachlorobutadiene	••	.2	
tert-Butylbenzene		.2	Isopropylbenzene		.2	
Carbon tetrachloride	5	.2	p-Isopropyltoluene		.2	
Chlorobenzene	100	.2	Methylene chloride	5	.2	
Chlorodibromomethane	*	.2	Methyl tert-butylether	••	1	
Chloroethane	••	.2	Naphthalene	••	.2	
Chloroform	*	.2	n-Propylbenzene		.2	
Chloromethane	••	.2	Styrene	100	.2	
2-Chlorotoluene		.2	1,1,1,2-Tetrachloroethane		.2	
4-Chlorotoluene		.2	1,1,2,2-Tetrachloroethane		.2	
1,2-Dibromo-3-chloropropane	.2	1	Tetrachloroethylene	5	.2	
1,2-Dibromoethane	.05	.2	Toluene	1,000	.2	
Dibromomethane		.2	1,2,3-Trichlorobenzene	••	.2	
1,2-Dichlorobenzene	600	.2	1,2,4-Trichlorobenzene	70	.2	
1,3-Dichlorobenzene	600	.2	1,1,1-Trichloroethane	200	.2	
1,4-Dichlorobenzene	75	.2	1,1,2-Trichloroethane	5	.2	
Dichlorobromomethane	*	.2	Trichloroethene	5	.2	
Dichlorodifluoromethane	••	.2	Trichlorofluoromethane		.2	
1,1-Dichloroethane		.2	1,2,3-Trichloropropane	••	.2	
1,2-Dichloroethane	5	.2	1,1,2-Trichloro 1,2,2-trifluoroethane	••	.2	
cis-1,2-Dichloroethene	70	.2	1,2,4-Trimethylbenzene	••	.2	
1,1-Dichloroethene	7	.2	1,3,5-Trimethylbenzene		.2	
trans-1,2-dichloroethene	100	.2	Vinyl chloride	2	.2	
1,2-Dichloropropane	5	.2	Xylenes, total ortho, meta, and para	10,000	.2	

Table 12. Maximum contaminant levels and minimum reporting levels of selected insecticides and gross polychlorinated compounds in drinking water

[Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory. The maximum contaminant levels were established pursuant to the recommendations of the U.S. Environmental Protection Agency (1997, p. 359) for community water systems and are included for comparison purposes only. Minimum reporting levels are from Timme (1995). Method detection limits are from Zaugg and others (1995). Units are in micrograms per liter. Symbols: ••, maximum contaminant level has not been established or proposed; *, samples analyzed using two different laboratory schedules and different minimum reporting levels. **, chlorthalonil is a fungicide, DNOC is listed as an insecticide and herbicide. Abbreviations: MCL, maximum contaminant level; MRL, minimum reporting level; MDL, method detection limit]

Carbamate insecticides								
Insecticide	MCL	MRL	Insecticide	MCL	MRL			
Aldicarb	3	0.016	Methomyl	••	.017			
Aldicarb sulfone	2	.016	1-Naphthol	••	.007			
Aldicarb sulfoxide	4	.021	Oxamyl	200	.050			
*Carbaryl (Sevin)	••	.008	Propham	••	.035			
*Carbofuran	40	.028	Propoxur		.035			
Methiocarb		.026						

Organophosphate insecticides

Insecticide	MCL	MRL	Insecticide	MCL	MRL
*Chlorpyrifos; Dursban	•	0.01	*Malathion	••.	0.01
DEF		.01	Methylparathion	••	.01
*Diazinon	••	.01	Parathion	••	.01
*Di-Syston (Disulfoton)		.01	*Phorate	••	.01
Ethion		.01	Trithion		.01
*Fonofos	••	.01			

Organochlorine insecticides

Insecticide	MCL	MRL	Insecticide	MCL	MRL
Aldrin	••	0.01	Heptachlor	0.4	0.01
Chlordane, technical	2	.1	Heptachlor epoxide	.2	.01
DDD, p,p'-	••	.01	*Lindane	.2	.01
*DDE, p,p'-		.01	Methoxychlor, p,p'-	40	.01
DDT, p,p'-	•	.01	Mirex	••	.01
*Dieldrin	••	.01	Perthane	••	.1
Endosulfan I		.01	Toxaphene	3	1.0
Endrin	2	.01			

Gross polychlorinated compounds

Compound	MCL	MRL
Gross polychlorinated biphenyls (PCBs)	0.5	0.1
Gross polychlorinated naphthalenes (PCNs)	••	0.1

Table 12. Maximum contaminant levels and minimum reporting levels of selected insecticides and gross polychlorinated compounds in drinking water—Continued

Additional insecticides

Insecticide	MCL	MRL	MDL	Insecticide	MCL	MRL	MDL
Azinphos methyl-	••	0.038	0.001	Ethoprop	••	.012	.003
*Carbaryl (Sevin)		.046	.003	*Fonofos	••	.008	.003
*Carbofuran	40	.013	.003	HCH, alpha-	••	.007	.002
*Chlorpyrifos		.005	.004	*HCH, gamma- (Lindane)	.2	.011	.004
**Chlorthalonil	••	.035	.035	Hydroxycarbofuran, 3-	••	.014	.014
*DDE, p,p'-		.010	.006	*Malathion	••	.010	.005
*Diazinon		.008	.002	Parathion, ethyl-		.022	.004
*Dieldrin	••	.008	.001	Parathion, methyl-	••	.035	.006
Dinoseb		.035	.035	Permethrine, cis-	••	.019	.005
*Disulfoton		.028	.017	*Phorate		.011	.002
**DNOC		.035	.035	Propargite I & II	••	.006	.013
Esfenvalerate		.019	.019	Terbufos		.012	.013

Table 13. Maximum contaminant levels and minimum reporting levels of chlorophenoxy-acid herbicides and other herbicides in drinking water

[Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory. The maximum contaminant levels were established pursuant to the recommendations of the U.S. Environmental Protection Agency (1997, p. 359) for community water systems and are included for comparison purposes only. Minimum reporting levels are from Timme (1995). Method detection limits are from Zaugg and others (1995). Units are in micrograms per liter. Symbols: ••, maximum contaminant level has not been established or proposed. *, samples analyzed using two different laboratory schedules with different minimum reporting levels. Abbreviations: MCL, maximum contaminant level; MRL, minimum reporting level; MDL, method detection limit]

Chlorophenoxy-acid herbicides									
Herbicide	MCL	MRL	Herbicide	MCL	MRL				
*2,4-D	70	0.01	*Silvex	50	0.01				
(dissolved)	70	.035	(dissolved)		.021				
2,4-DB	••	.035	*2,4,5-T	••	.01				
2,4-DP	••	.01	(dissolved)		.035				

Other herbicides

Herbicide	MCL	MRL	MDL	Herbicide	MCL	MRL	MDL
Acetochlor	••	0.009	0.002	*Linuron	••	.039	.002
Acifluorfen		.035	.035	(dissolved)	••	.018	.018
Alachlor	2	.009	.002	MCPA	••	.050	.050
Atrazine	3	.017	.001	MCPB	••	.035	.035
Atrazine, desethyl-	••	.007	.002	Metolachlor	••	.009	.002
Benfluralin	••	.013	.002	Metribuzin	••	.012	.004
Bentazon		.014	.014	Molinate		.007	.004
Bromacil		.035	.035	Napropamide	••	.010	.003
Bromoxynil	••	.035	.035	Neburon	••	.015	.015
Butylate		.008	.002	Norflurazon		.024	.024
Chloramben	••	.011	.011	Oryzalin		.019	.019
Clopyralid	••	.050	.050	Pebulate	••	.009	.004
Cyanazine	••	.013	.004	Pendimethalin	••	.018	.004
*DCPA (Dacthal)	••	.004	.002	Picloram	500	.050	.050
(dissolved)	••	.017	.017	Prometon	••	.008	.018
Dicamba	••	.035	.035	Pronamide	••	.009	.003
Dichlobenil	••	.020	.020	Propachlor	••	.015	.007
Dichlorprop		.032	.032	Propanil	••	.016	.004
Diethylaniline	••	.006	.003	Simazine	4	.008	.005
Diuron	••	.020	.020	Tebuthiuron	••	.015	.010
EPTC (Eptam)	••	.005	.002	Terbacil		.030	.007
Ethalfluralin	••	.013	.004	Thiobencarb		.008	.002
Fenuron	••	.013	.013	Triallate	••	.008	.001
Fluometuron	••	.035	.035	Triclopyr		.050	.050
				Trifluralin	••	.012	.002

Table 14. Concentrations of herbicides in water from selected wells and springs, eastern Snake River Plain [See figure 2 for location of sites. Analyses were performed by the U.S. Geological Survey National Water Quality Laboratory. Units are in micrograms per liter. Symbols: <, concentration is less than the respective minimum reporting level; *, quality-assurance sample (MV-28 is a replicate of MV-32; MV-34 is a replicate of MV-41); E, concentration was estimated and needs to be evaluated carefully because of variable performance. Concentrations of herbicides not listed were less than the minimum reporting levels and method detection limits in all samples]

Site identifier	Alachlor	Atrazone	Bentazon	Desethyl atrazine	Metrbuzin	Prometon	Simazine
MV-02	< 0.002	E, 0.002	< 0.014	E, 0.001	< 0.004	< 0.018	< 0.005
MV-09	<.002	.012	<.014	E, .014	<.004	<.018	.000
MV-10	<.002	.006	<.014	E, .005	<.004	<.018	<.00
MV-16	<.002	E, .002	<.014	E, .004	<.004	<.018	<.005
MV-19	E, .002	.012	E, .009	E, .014	<.004	<.018	<.005
MV-20	<.002	.004	<.014	E, .007	.006	<.018	<.005
MV-25	<.002	.011	<.014	E, .006	<.004	<.018	<.003
MV-28*	<.002	<.001	<.014	E, .003	<.004	<.018	<.005
MV-32	<.002	<.001	<.014	E, .002	<.004	<.018	<.00:
MV-34*	<.002	.018	<.014	E, .026	<.004	E, .003	<.005
MV-40	<.002	E, .002	<.014	E, .002	<.004	<.018	<.005
MV-41	<.002	.017	<.014	E, .026	<.004	E, .004	E, .002
MV-45	<.002	.004	<.014	E, .003	E, .002	<.018	<.00:
MV-47	<.002	E, .001	<.014	E, .001	<.004	<.018	<.005
MV-48	<.002	<.001	<.014	E, .024	<.004	<.018	<.00.5
MV-49	<.002	<.001	<.014	<.002	<.004	<.018	<.00:
MV-51	<.002	E, .002	<.014	E, .004	<.004	<.018	<.005
MV-52	<.002	E, .001	<.014	E, .002	<.004	<.018	<.00.
MV-54	<.002	.011	<.014	E, .012	<.004	<.018	.003
MV-56	<.002	E, .002	<.014	E, .002	<.004	<.018	<.003

