# ARGONNE NATIONAL LABORATORY-EAST SITE ENVIRONMENTAL REPORT FOR CALENDAR YEAR 1994

by

N. W. Golchert and R. G. Kolzow



ARGONNE NATIONAL LABORATORY, ARGONNE, ILLINOIS

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## ARGONNE NATIONAL LABORATORY-EAST SITE ENVIRONMENTAL REPORT FOR CALENDAR YEAR 1994

by

N. W. Golchert and R. G. Kolzow Environmental Management Operations

May 1995



9700 South Cass Avenue
Argonne, Illinois 60439

Preceding Report in This Series: ANL-94/10

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**1,2-DCE** cis-1,2-Dichloroethene

ACM Asbestos-Containing Materials

ADS Activity Data Sheets

ALARA As Low As Reasonably Achievable ANL Argonne National Laboratory-East

AOC Area of Concern

APS Advanced Photon Source

ASTM American Society for Testing and Materials
ATLAS Argonne Tandem Linac Accelerating System

BAT Best Available Technology
BOD Biochemical Oxygen Demand

CAA Clean Air Act

CAAPP Clean Air Act Permit Program

CEDE Committed Effective Dose Equivalent

**CERCLA** Comprehensive Environmental Response, Compensation

and Liability Act

CFR Code of Federal Regulations
CLP Contract Laboratory Program
COD Chemical Oxygen Demand

**CP-5** Chicago Pile-Five

**CRADA** Cooperative Research and Development Agreement

CRM Cultural Resource Management

CWA Clean Water Act

CWDD Continuous Wave Deuterium Demonstrator

CY Calendar Year

**D&D** Decontamination and Decommissioning

DCG Derived Concentration Guides
DMR Discharge Monitoring Report
U. S. Department of Energy

DOE-CH U. S. Department of Energy - Chicago Operations Office

DOT Department of Transportation
DSTP Draft Site Treatment Plan
EA Environmental Assessment

**EBWR** Experimental Boiling Water Reactor

**ECR** Environmental Compliance Representative

EIS Environmental Impact Statement

EML Environmental Measurements Laboratory
EMO Environmental Management Operation

EMS Environmental Protection Data Management System

**ENE** East-Northeast

**EPA** U. S. Environmental Protection Agency

**EPCRA** Emergency Planning and Community Right to Know Act

**ESA** Endangered Species Act

ESH Environment, Safety and Health

ESH/DACH Environment, Safety and Health/Dosimetry and Analytical Services,

Chemistry Laboratory

ESH/DACL Environment, Safety and Health/Dosimetry and Analytical Services,

Control Laboratory

ESH/DARC Environment, Safety and Health/Dosimetry and Analytical Services,

Radiochemistry Laboratory

EWM Environment and Waste Management Program

FEUL Fossil Energy Users Laboratory

FFCA Federal Facility Compliance Agreement

FIFRA Federal Insecticide, Fungicide and Rodenticide Act

FWS Fish and Wildlife Service

FY Fiscal Year

GOCO Government-Owned Contractor-Operated

**HEPA** High-Efficiency Particulate Air

HRS Hazard Ranking System

HSWA Hazardous and Solid Waste Amendments
HWMU Hazardous Waste Management Unit

IAC Illinois Administrative Code ICP Inductively Coupled Plasma

ICRP International Commission on Radiological Protection

IDPH Illinois Department of Public Health
IEPA Illinois Environmental Protection Agency
IHPA Illinois Historic Preservation Agency

IPNS Intense Pulsed Neutron Source

LEPC Local Emergency Planning Committee
LWTP Laboratory Wastewater Treatment Plant

MCL Maximum Contaminant Level

MCLG Maximum Contaminant Level Goals

MHD Magneto Hydrodynamics
MSDS Material Safety Data Sheets

MSL Mean Sea Level

NBL New Brunswick Laboratory

NCRP National Commission on Radiation Protection and Measurements

NEPA National Environmental Policy Act

**NESHAP** National Emission Standards for Hazardous Air Pollutants

NHPA National Historic Preservation Act

NIST National Institute of Standards and Technology

NOAA National Oceanic and Atmospheric Administration NPDES National Pollutant Discharge Elimination System

NPL National Priority List

NRHP National Register of Historical Places

OSHA Occupational Safety and Health Administration

PA Preliminary Assessment
PCB Polychlorinated Biphenyls

PCV Polyvinyl Chloride

PFS Plant Facilities and Services
PRP Potentially Responsible Party
PSTP Proposed Site Treatment Plan

QA Quality Assurance

QAP Quality Assurance Program

**RCRA** Resources Conservation and Recovery Act

RFA RCRA Facility Assessment
RFI RCRA Facility Investigation
RMW Radioactive Mixed Waste

SARA Superfund Amendments and Reauthorization Act

SDWA Safe Drinking Water Act

SI Site Investigation

SIP State Implementation Plan SOP Standard Operating Procedure

SPCC Spill Prevention Control and Countermeasures

SRM Standard Reference Material
SSI Site Screening Investigation
SWMU Solid Waste Management Units

TCA 1,1,1-Trichloroethane

TCE Trichloroethene

TDS Total Dissolved Solids

TIE Toxicity Identification Evaluation
TLD Thermoluminescent Dosimeter

TRC Total Residual Chlorine

TRE Toxicity Reduction Evaluation
TSCA Toxic Substances Control Act

TSS Total Suspended Solids

VOC Volatile Organic Compounds WMO Waste Management Operations

WQS Water Quality Standards

ANL Site Environmental Report

This report discusses the results of the environmental protection program at Argonne National Laboratory-East (ANL) for 1994. To evaluate the effects of ANL operations on the environment, samples of environmental media collected on the site, at the site boundary, and off the ANL site were analyzed and compared to applicable guidelines and standards. A variety of radionuclides was measured in air, surface water, groundwater, soil, grass, and bottom sediment samples. In addition, chemical constituents in surface water, groundwater, and ANL effluent water were analyzed. External penetrating radiation doses were measured and the potential for radiation exposure to off-site population groups was estimated. The results of the surveillance program are interpreted in terms of the origin of the radioactive and chemical substances (natural, fallout, ANL, and other) and are compared with applicable environmental quality standards. A U. S. Department of Energy (DOE) dose calculation methodology, based on International Commission on Radiological Protection (ICRP) recommendations and the CAP-88 version of the EPA-AIRDOSE/RADRISK COMPUTER CODE, is used in this report. The status of ANL environmental protection activities with respect to the various laws and regulations which govern waste handling and disposal is discussed. This report also discusses progress being made on environmental corrective actions and restoration projects.

This report is a summary of the ongoing environmental protection program conducted by ANL in 1994. It includes descriptions of the site, the ANL missions and programs, the status of compliance with environmental regulations, environmental protection and restoration activities, and the environmental surveillance program. The surveillance program conducts regular monitoring for radiation, radioactive materials, and nonradiological constituents on the ANL site and in the surrounding region. These activities document compliance with appropriate standards and permit limits, identify trends, provide information to the public, and contribute to a better understanding of ANL's impact on the environment. The surveillance program supports the ANL policy to protect the public, employees, and the environment from harm that could be caused by ANL activities and to reduce environmental impacts to the greatest degree practicable.

#### **Compliance Summary**

Radionuclide emissions, the disposal of asbestos, and conventional air pollutants from ANL facilities are regulated under the Clean Air Act (CAA). A number of airborne radiological emission points at ANL are subject to the National Emission Standards for Hazardous Air Pollutants (NESHAP) regulations for radionuclide releases from DOE facilities (40 CFR 61, Subpart H). All such air emission sources were evaluated to ensure that the requirements are being properly addressed. The ANL individual off-site dose required to be reported by these U. S. Environmental Protection Agency (EPA) regulations was 0.015 mrem/y in 1994. This is 0.15% of the 10 mrem/y standard.

At ANL, asbestos-containing material was frequently encountered during maintenance or renovation of existing facilities and equipment. Asbestos was removed in strict accordance with the NESHAP regulations as well as with the much stricter Occupational Safety and Health Administration (OSHA) worker protection standards. All asbestos waste

material was disposed of at off-site landfills in Illinois. Approximately 153 m<sup>3</sup> (5400 ft<sup>3</sup>) of asbestos-containing materials were removed and disposed of off-site during 1994.

The ANL site contains several sources of conventional air pollutants. The steam plant and fuel dispensing facilities operate continuously and represent the only significant sources of conventional air pollutants. The operating permit for the steam plant requires continuous opacity and sulfur dioxide monitoring of the smoke stack from Boiler No. 5, the only boiler equipped to burn coal. Coal was burned five months during 1994, whereas natural gas was used exclusively as a fuel for seven months of the year. During the period coal was burned, which is in colder weather to supplement the gas-fired boilers, one excursion for opacity was observed. The last of the high sulfur coal was burned in January 1994.

The principal regulatory mechanism designed to achieve the goals of the Clean Water Act (CWA) is the National Pollutant Discharge Elimination System (NPDES). The authority to implement the NPDES program has been delegated to the State of Illinois. Nine surface water discharge points are regulated by the ANL NPDES permit, which identifies the sampling locations, sampling frequency, constituents, and limits. The permit renewal, which became effective October 30, 1994, increased the number of monitored discharge points to 28. Although there was a slight decrease of NPDES exceedances (24) during 1994, overall the number of NPDES exceedances has been declining with 86 in 1990, 44 in 1991, 19 in 1992 and 25 in 1993.

ANL was granted interim status under the Resource Conservation and Recovery Act (RCRA) by submitting a Part A permit application in 1980. In 1990, a Part B permit application was submitted to the Illinois Environmental Protection Agency (IEPA). Fourteen hazardous waste treatment and storage facilities have been identified. The Part B permit application is currently under review.

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ANL has prepared and implemented a site-wide underground storage tank compliance plan. Thirty-three tanks were removed over the past several years and 22 tanks replaced or upgraded in FY 1992 and FY 1993. Three tanks in the 800 Area (Building 827) which are no longer necessary for operation remain to be removed. One additional tank (Tank No. 17) servicing Buildings 813 and 815 will be upgraded or replaced as appropriate in FY 1995. Of the locations from which tanks were removed or replaced, 17 were found to have some degree of exterior contamination from leaks, spills, or overfills. All but one of these contaminated sites were successfully cleaned and filled. One removal completed in 1989 required an IEPA-approved "dirty" closure due to its proximity to a building.

In 1986, ten potential Comprehensive Environmental Response, Compensation and Liability Act (CERCLA) sites were identified. Under the Superfund Amendments and Reauthorization Act (SARA) of 1986, a total of 15 Preliminary Assessments (PA) reports were submitted. In late 1990, Site Screening Investigation (SSI) reports were completed on two individual sites and one composite submittal of three locations (317/319/ENE). Characterization studies are at various stages for a number of the identified sites.

The only Toxic Substances Control Act (TSCA) compounds in significant quantities at ANL are polychlorinated biphenyls (PCBs) contained in electrical capacitors, transformer oil, and PCB-contaminated soil and sludge. All pole-mounted transformers and circuit breakers containing PCBs were replaced or retrofilled with non-PCB oil. All removal and disposal activities were conducted by licensed contractors specializing in such operations. A sludge drying bed, servicing the ANL wastewater treatment plant, is contaminated with PCBs of unknown origin. An extensive characterization study was conducted and the sludge was containerized and stored.

The DOE implementation of the National Environmental Policy Act (NEPA) regulations has been undergoing significant changes since 1992. Most NEPA project reviews sent to

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DOE for review and approval were determined to be categorical exclusions although Environmental Assessments (EA) will be required for several projects. There are currently no active projects at ANL for which an Environmental Impact Statement (EIS) is required.

The 1994 Five-Year Plan requests funds for on-site corrective action projects, environmental restoration projects, and waste management activities. The corrective action projects concentrate on upgrading or replacing existing treatment facilities. Environmental restoration activities are projects which assess and clean up inactive waste sites. These include two inactive landfills, three French drains (dry wells used to dispose of liquid chemicals), two inactive wastewater treatment facilities and a number of areas that may have been contaminated with small amounts of hazardous chemicals. A number of Decontamination and Decommissioning (D&D) projects for on-site nuclear facilities have been identified, including clean up at the Experimental Boiling Water Reactor (EBWR) and Chicago Pile-Five (CP-5) research reactors. The majority of the Waste Management projects involve improvements to existing treatment or storage facilities.

The major compliance issue at ANL in 1994 was compliance with the new NPDES limits for TDS, copper, ammonia, nitrogen, and total residual chlorine. Another significant issue involved the inadvertent shipment, through an independent contractor, of waste oil contaminated with PCBs to an off-site recycling facility. Other compliance issues included exceeding the action levels for lead and copper in drinking water, elevated levels of some routine indicator parameters in the groundwater at the sanitary landfill, and clean-up of environmental contamination caused by previous activities on the ANL site.

#### **Environmental Surveillance Program**

Airborne emissions of gaseous radioactive materials from ANL were monitored and the effective dose equivalents were estimated at the site perimeter and to the maximally-exposed

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member of the public. The CAP-88 version of the EPA/AIRDOSE-RADRISK code was used. The estimated maximum perimeter dose was 0.35 mrem/y in the north direction, while the estimated maximum dose to a member of the public was 0.12 mrem/y. This is 0.12% of the DOE radiation protection standard of 100 mrem/y for all pathways. Approximately 90% of this estimated dose is due to the release of 1750 curies of radon-220 in 1994. If the radon-220 impact is excluded from reporting, as required in 40 CFR 61, Subpart H, the estimated dose to the maximally-exposed individual would be 0.015 mrem/y. The estimated population dose from all releases to the approximately eight million people living within 80 km (50 mi) of the site was 5.8 man-rem compared to 13.0 man-rem in 1993.

Air monitoring was also conducted at ANL for total alpha activity, total beta activity, strontium-90, isotopic thorium, isotopic uranium, and plutonium-239. No statistically significant difference was identified between samples collected at the ANL perimeter and samples collected off the site. Monitoring for hazardous chemical constituents in ambient air was not conducted.

The only source of radionuclides and chemical pollutants in surface water due to ANL releases was in Sawmill Creek below the wastewater discharge point. At various times, measurable levels of hydrogen-3, strontium-90, cesium-137, plutonium-239, and americium-241 were detected. Of these radionuclides, the maximum annual release was 1.14 curies of hydrogen-3. The hydrogen-3 was added to the wastewater as part of normal Laboratory operations. The dose to a hypothetical individual using water from Sawmill Creek as his sole source of drinking water would be 0.0788 mrem/y. However, no one uses this as drinking water and dilution by the Des Plaines River reduces the concentrations of the measured radionuclides to levels below their respective detection limits downstream from ANL at Lemont. Sawmill Creek is also monitored for nonradiological constituents to demonstrate compliance with State of Illinois water quality standards. Silver and copper were occasionally detected above the standard.

Surface soil and grass samples were collected at ten perimeter and ten off-site locations during 1994. The purpose of the sampling was to detect the possible buildup of radionuclides from the deposition of airborne emissions. The results indicate no statistically significant difference between the perimeter and off-site concentrations of potassium-40, cesium-137, radium-226, thorium-228, thorium-232, plutonium-238, plutonium-239, and americium-241.

Sediment samples were collected from Sawmill Creek, above, at, and below the point of wastewater discharge. For comparison purposes, samples were also collected from the beds of ten off-site streams and ponds. The analysis of the off-site samples for selected radionuclides established their current ambient levels. Elevated levels of cesium-137 (up to 0.49 pCi/g), plutonium-238 (up to 0.002 pCi/g), plutonium-239 (up to 0.019 pCi/g), and americium-241 (up to 0.008 pCi/g) were found in the sediment below the outfall and are attributed to past ANL releases.

Dose rates from penetrating radiation (gamma-rays) were measured at 14 perimeter and on-site locations and at five off-site locations in 1994 using thermoluminescent dosimeters. The off-site results averaged  $82 \pm 6$  mrem/y, consistent with the long-term average. Above-background doses occurred at one perimeter location and were due to ANL operations. At the south fence, radiation from a temporary storage facility for radioactive waste resulted in an average dose of 92 mrem/y for 1994. The estimated dose from penetrating radiation to the nearest resident south of the site was < 0.01 mrem/y.

The potential radiation doses to members of the public from ANL operations during 1994 were estimated by combining the exposure from inhalation, ingestion, and direct radiation pathways. The pathway that dominates is the airborne releases. The highest estimated dose was about 0.12 mrem/y to individuals living 500 m north of the site if they were outdoors at that location during the entire year. Doses from other pathways were

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calculated and were small at this location. The magnitude of the doses from ANL operations are well within all applicable standards and are insignificant when compared to doses received by the public from natural radiation (~ 300 mrem/y) or other sources, e.g., medical x-rays and consumer products (~ 60 mrem/y).

Radiological and chemical constituents in the groundwater were monitored in several areas of the ANL site in 1994. The ANL domestic water supply is monitored by collecting quarterly samples from the four wells and a treated water tap. All results were less than the limits established by the Safe Drinking Water Act (SDWA) except for elevated levels of total dissolved solids (TDS). The action levels for copper and lead in drinking water were exceeded during 1994.

Ten monitoring wells screened in the glacial till and two into the dolomite were sampled quarterly at the 317/319 Area and analyzed for radiological and for volatile organic, semivolatile, PCBs, and pesticides/herbicides constituents. The major organic contaminants detected were trichloroethene, 1,1,1-trichloroethane, 1,1-dichloroethane, carbon tetrachloride, 1,2-dichloroethane, tetrachloroethene, and chloroform. Measurable levels of hydrogen-3, strontium-90, and cesium-137 were present in several of the wells. A characterization program to assess the extent of the groundwater contamination was initiated during 1993 and continued in 1994.

Eleven monitoring wells screened in the glacial till and two into the dolomite at the 800 Area sanitary landfill are sampled on a quarterly basis and analyzed for metals, volatile organic compounds, semivolatiles, PCBs, pesticides/herbicides, and hydrogen-3. Levels above Water Quality Standards (WQS) for chloride, iron, manganese, and total dissolved solids were found in some wells. Above background levels of hydrogen-3, 1,4-dioxane, chlorodifluoromethane, and tetrahydrofuran were found in several of the wells. A work

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plan for a groundwater characterization program at this site was completed during 1993 and work began in 1994.

An extensive quality assurance program is maintained to cover all aspects of the environmental surveillance sampling and analysis programs. Approved documents are in place along with the supporting standard operating procedures. Newly collected data were compared both with recent results and historical data to ensure that deviations from previous conditions were identified and promptly evaluated. Samples at all locations were collected using well-established and documented procedures to ensure consistency. Samples were analyzed by documented standard analytical procedures. Data quality was verified by a continuing program of analytical laboratory quality control, participation in inter-laboratory cross-checks, and replicate sampling and analysis. Data were managed and tracked by a dedicated computerized data management system which assigns unique sample numbers, schedules collection and analysis, checks status, and prepares tables and information for the annual report.

## 1. INTRODUCTION



#### 1.1. General

This annual report on the ANL environmental protection program provides the DOE, environmental agencies, and the public with information on the levels of radioactive and chemical pollutants in the vicinity of ANL and on the amounts, if any, added to the environment by ANL operations. It also summarizes compliance of ANL operations with applicable environmental laws and regulations and highlights significant accomplishments and problems related to environmental protection. The report follows the guidelines given in DOE Order 5400.1.1

ANL conducts a continuing program of environmental surveillance on and near the site to determine the identity, magnitude, and origin of radioactive and chemical substances in the environment. The detection of any such materials released to the environment by operations of ANL is of special interest. One important function of the program is to verify the adequacy of ANL's pollution control systems.

ANL is a DOE energy research and development laboratory with several principal objectives. It conducts a broad program of research in the basic energy and related sciences (physical, chemical, material, computer, nuclear, biomedical, and environmental) and serves as an important engineering center for the study of nuclear and nonnuclear energy sources. Energy-related research projects conducted during 1994 included: advanced reactor development; safety studies for light water and breeder reactors; component and material development for fission and fusion reactors; superconductivity advances and applications; improvements in the use of coal for power production (particularly high-sulfur coal); synchrotron radiation accelerator design; development of electrochemical energy sources, including fuel cells and batteries for vehicles and for energy storage; and evaluation of heat exchangers for the recovery of waste heat from engines.

Other areas of research are the use of superconducting magnets for improved nuclear particle accelerators, fundamental coal chemistry studies, the immobilization of radioactive waste products for safe disposal, medical radioisotope technology, carcinogenesis, and the biological effects of small amounts of radiation. Environmental research studies include biological activity of energy-related mutagens and carcinogens; characterization and monitoring of energy-related pollutants; and the effects of acid rain on vegetation, soil, and surface water quality. A significant number of these laboratory studies require the controlled use of radioactive and chemically toxic substances.

The principal nuclear facilities at ANL are: a superconducting heavy ion linear accelerator (Argonne Tandem Linac Accelerating System, ATLAS); a 22 MeV pulsed electron Linac; several other charged particle accelerators (principally of the Van de Graaff and Dynamitron types); a large fast neutron source (Intense Pulsed Neutron Source, IPNS) in which high-energy protons strike a uranium target to produce neutrons; chemical and metallurgical plutonium laboratories; and several hot cells and laboratories designed for work with multi-curie quantities of the actinide elements and with irradiated reactor fuel materials. The DOE New Brunswick Laboratory (NBL), a safeguards plutonium and uranium measurements and analytical chemistry laboratory, is located on the ANL site.

Two activities initiated in 1984 and continued in 1994 have some potential environmental impact: (1) management of radioactive contamination remaining from the proof-of-breeding in light-water reactors project, which involved the dissolution and analysis of irradiated thorium and uranium-233 dioxide fuel elements and (2) recovery of tritium from reactor irradiated ceramic lithium compounds. The shut down 5-MW heavy water enriched uranium research reactor (CP-5) and the EBWR are in various stages of decontamination and decommissioning.

The principal nonnuclear activities at ANL in 1994 that may have measurable impacts on the environment include the use of a coal-fired boiler (No. 5), studies of the closed-loop heat exchanger for waste heat recovery and use of large quantities of chlorine for water treatment. The closed-loop heat exchanger studies involved the use of moderately large quantities of toxic or flammable organic compounds, such as toluene, Freon, biphenyl oxides, methyl pyridine, and trifluoroethanol. Chlorine is used for wastewater treatment. The major potential for environmental impact from these materials would be associated with any accidental releases caused by equipment malfunction. However, no such releases have occurred.

# 1.2. Description of Site

Argonne National Laboratory (Illinois site) occupies the central 688 hectares (1,700 acres) of a 1,514-hectare (3,740-acre) tract in DuPage County. The site is 43 km (27 mi) southwest of downtown Chicago and 39 km (24 mi) west of Lake Michigan. It is north of the Des Plaines River Valley, south of Interstate Highway 55 (I-55) and west of Illinois Highway 83. Figures 1.1 and 1.2 are maps of the site, the surrounding area, and sampling locations of the monitoring program. The 826-hectare (2,040-acre) Waterfall Glen Forest Preserve surrounding the site is mostly former ANL property that was deeded to the DuPage County Forest Preserve District in 1973 for use as a public recreational area, nature preserve, and demonstration forest. Figure 1.1 contains numbers on the abscissa and letters on the ordinate. In this report, facilities are identified by the alpha-numeric designations in Figure 1.1 to facilitate their location.

The terrain of ANL is gently rolling, partially wooded, former prairie and farmland. The grounds contain a number of small ponds and streams. The principal stream is Sawmill Creek, which runs through the site in a southerly direction and enters the Des Plaines River



Figure 1.1 Sampling Locations at Argonne National Laboratory

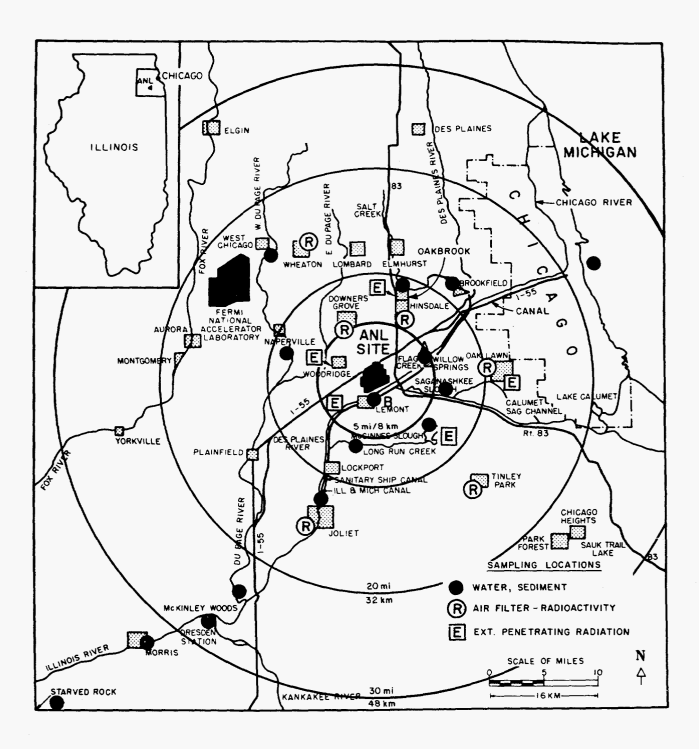


Figure 1.2 Sampling Locations Near Argonne National Laboratory

about 2.1 km (1.3 mi) southeast of the center of the site. The land is drained primarily by Sawmill Creek, although the extreme southern portion drains directly into the Des Plaines River, which flows along the southern boundary of the forest preserve. This river flows southwest until it joins the Kankakee River about 48 km (30 mi) southwest of ANL to form the Illinois River.

The largest topographical feature of the area is the Des Plaines River valley, which is about 1.6 km (1 mi) wide. This valley contains the river, the Chicago Sanitary and Ship Canal, and the Illinois and Michigan Canal. Their presence extends the uninhabited area created by the ANL site and surrounding forest preserve about 1.6 km (1 mi) south of the site. The elevation of the channel surface is 180 m (578 ft) above sea level. The bluffs that form the southern border of the site rise from the river channel at slope angles of 15° to 60°, reaching an average elevation of 200 m (650 ft) above sea level at the top. The land then slopes gradually upward reaching the average site elevation of 220 m (725 ft) above sea level at 915 m (3,000 ft) from the bluffs. Several large ravines oriented in a north-south direction are located in the southern portion of the site. The bluffs and ravines generally are forested with mature deciduous trees. The remaining portion of the site changes in elevation by no more than 7.6 m (25 ft) in a horizontal distance of 150 m (500 ft). The Chicago District Pipe Line Co. and the Atchison, Topeka, and Santa Fe Railroad have rights-of-way in the southern portion of the forest preserve, Additional information about the site is given in the 1982 draft Argonne Environmental Assessment.<sup>2</sup>

# 1.3. Population

The area around ANL has experienced a large population growth in the past 30 years. Large areas of farmland have been converted into housing. Table 1.1 presents directional and annular 80-km (50-mi) population distribution for the area, which is used for the population dose calculations later in this report. The population distribution, centered on

TABLE 1.1

Population Distribution in the Vicinity of ANL, 1991

	Population (individuals) at 0-5 Miles				Population (thousands) at 5-50 Miles					
Direction	0-1	1-2	2-3	3-4	4-5	5-10	10-20	20-30	30-40	40-50
N	0	661	4199	5602	8783	44.7	172.1	336.7	187.5	221.3
NNE	0	22	3684	5925	5287	38.8	302.3	485.8	86.7	0
NE	0	737	2293	2431	1689	40.9	674.4	866.3	0	0
ENE	0	1117	2495	1460	1482	33.5	598.7	178.9	0	0
E	0	16	10	1	42	40.8	467.0	199.8	13.0	25.8
ESE	0	0	55	331	306	22.4	186.1	282.0	245.0	80.9
SE	0	2	219	425	198	19.8	103.2	114.2	28.6	12.2
SSE	0	72	401	221	1800	12.0	22.1	7.7	11.0	16.8
S	0	105	2298	921	860	3.7	23.4	2.0	35.3	35.0
SSW	0	33	3504	1229	759	14.7	89.8	10.8	17.6	7.1
SW	0	80	20	87	79	11.6	36.7	9.4	16.2	9.1
WSW	0	215	86	620	1646	4.8	7.6	3.7	8.0	7.2
W	0	779	1237	8338	9056	26.2	67.2	19.0	14.8	6.7
WNW	0	254	224	5867	4433	44.3	104.6	20.7	6.6	52.9
NW	0	552	2602	6979	6779	41.6	69.1	95.5	18.2	16.7
NNW	0	492	2774	4521	9390	33.4	108.5	225.2	130.6	96.5
Total	0	5137	26101	44958	52589	433.2	3112.8	2857.7	819.1	588.4
Cumulative Total	0	5137	31238	76196	128785	561.9	3674.7	6532.4	7351.5	7939.9

To convert from miles to kilometers, multiply by 1.6.

Cumulative total = total of this sector plus totals of all previous sectors.

the CP-5 reactor (Location 9G in Figure 1.1), was prepared by the Geographic Data Systems Computing and Telecommunications Division at Oak Ridge National Laboratory and represents projections to 1991 based on the 1990 census data.

# 1.4. Climatology

The climate of the area is representative of the upper Mississippi Valley, as moderated by Lake Michigan. Summaries of the meteorological data collected on the site from 1949 to 1964 are available<sup>3</sup> and provide a historical sample of the climatic conditions. The most important meteorological parameters for the purposes of this report are wind direction, wind speed, temperature, and precipitation. The wind data are used to select air sampling locations and distances from sources and to calculate radiation doses from air emissions. Temperature and precipitation data are useful in interpreting some of the monitoring results. The 1994 data were obtained from the on-site ANL meteorological station. The 1994 average monthly and annual wind roses are shown in Figure 1.3. The wind roses are polar coordinate plots in which the lengths of the radii represent the percentage frequency of wind speeds in classes of 2.01-6 m/s (4.5-13.4 mph), 6.01-10 m/s (13.4-22.4 mph), and greater than 10.01 m/s (22.4 mph). The number in the center of each wind rose represents the percentage of observations of wind speed less than 2 m/s (4.5 mph) in all directions. The direction of the radii from the center represents the direction from which the wind blows. Sixteen radii are shown on each plot at 22.5° intervals; each radius represents the average wind speed for the direction covering 11.25° on either side of the radius.

The monthly wind roses indicate that the winds are variable, so that monitoring for airborne releases must be carried out in all directions from the site. For example, the dominant wind direction in January is from the west while in February it is south. The annual average wind rose for 1994 is consistent with the long-term average wind direction, which usually varies from the west to south, but with a significant northeast component.

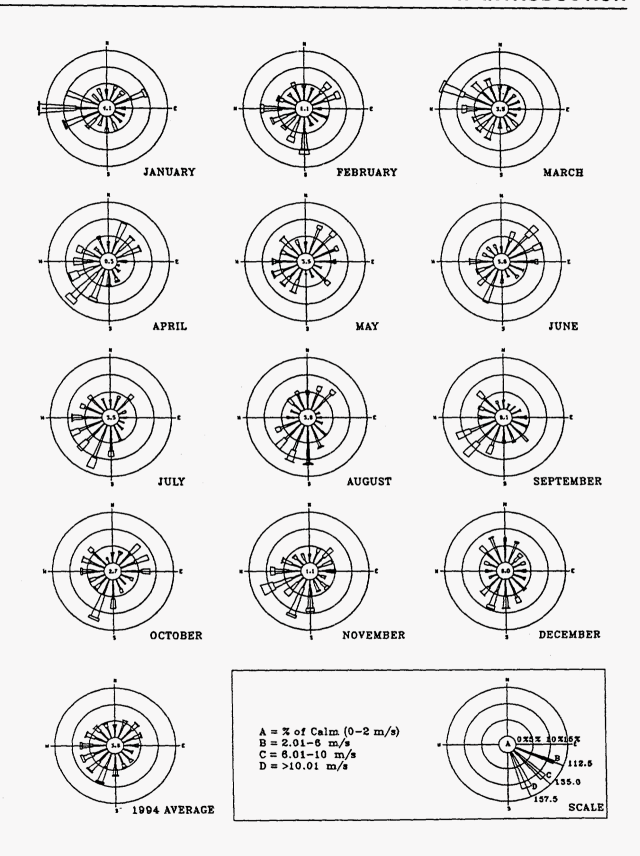


Figure 1.3 Monthly and Annual Wind Roses at Argonne National Laboratory, 1994

Precipitation and temperature data for 1994 are shown in Table 1.2. The monthly precipitation data for 1994 showed some differences from the average. For example, August and November are above the average, while March, May, and July are below the average. The annual total is very similar to the long-term average. Except for a colder January, the temperatures are similar to the long-term averages.

# 1.5. Geology

The geology of the ANL area consists of about 30 m (100 ft) of glacial till on top of bedrock, which is Niagaran and Alexandrian dolomite, underlain by shale and older dolomites and sandstones of Ordovician and Cambrian age. The beds are nearly horizontal. Niagaran and Alexandrian dolomite is about 60 m (200 ft) thick and widely used in DuPage County as a source of groundwater. The shale separating the upper dolomite aquifer from the underlying sandstone and dolomite aquifers retards hydraulic connection between them. The lower aquifer has a much lower piezometric level and does not appear to be affected by pumpage from the overlying bedrock.

The southern boundary of ANL follows the escarpment of a broad valley, now occupied by the Des Plaines River and the Chicago Sanitary and Ship Canal. This valley was carved by waters flowing out of the glacial Lake Michigan about 11,000 to 14,000 years ago. The soils on the site have derived from glacial till over the past 12,000 year, most of which are of the Morley series, which are moderately well-drained upland soils with slope ranging from 2% to 20%. The surface layer is a dark grayish-brown silt loam, the subsoil is a brown silty clay, and the underlying material is a silty clay loam glacial till. Morley soils have a relatively low organic content in the surface layer, moderately slow subsoil permeability, and a large water capacity. These soils are well-suited to growing crops, if good erosion control practices are used. The remaining soils along creeks, intermittent streams, bottomlands, and a few small upland areas are of the Sawmill,

TABLE 1.2

ANL Weather Summary, 1994

Precipitation (cm) ANL			m)	ANL 1994	Temperature (°C) ANL	
Month	ANL 1994	Historical Average**	Historical Average*	Monthly Average	Historical Average**	Historical Average*
January	3.18	3.61	4.06	-9.1	-5.9	-5.9
February	5.69	3.38	3.33	-5.0	-3.7	-3.3
March	1.17	5.56	6.58	3.4	0.6	2.2
April	6.25	9.14	9.30	10.6	8.3	9.3
May	1.60	7.82	8.00	15.0	14.5	15.1
June	11.18	9.47	10.36	20.5	19.7	20.3
July	5.82	10.97	9.22	22.0	21.7	22.8
August	14.86	8.71	8.97	20.1	20.9	22.2
September	4.55	7.14	8.51	18.6	16.8	18.2
October	8.07	6.58	5.79	12.4	11.4	11.9
November	10.92	4.37	5.23	6.6	2.9	4.3
December	5.38	3.20	5.33	1.3	-4.2	-2.4
Total	78.67	80.01	79.95			

<sup>\*</sup>Data obtained from the National Oceanic and Atmospheric Administration (NOAA) for the weather station at O'Hare International Airport. The average is for the years 1951-1980.

<sup>\*\*</sup>ANL data obtained from Reference 3.

Ashkum, Peotone, and Beecher series, which are generally poorly drained. They have a black to dark gray or brown silty clay loam surface layer, high organic-matter content, and a large water capacity.

# 1.6. Seismicity

No tectonic features within 135 km (62 mi) of ANL are known to be seismically active. The longest of these features is the Sandwich Fault. Smaller local features are the Des Plaines disturbance, a few faults in the Chicago area, and a fault of apparently Cambrian age.

Although a few minor earthquakes have occurred in northern Illinois, none has been positively associated with particular tectonic features. Most of the recent local seismic activity is believed to be caused by isostatic adjustments of the earth's crust in response to glacial loading and unloading, rather than by motion along crustal plate boundaries.

There are several areas of considerable seismic activity at moderate distances (hundreds of kilometers) from ANL. These areas include the New Madrid Fault zone (southwestern Missouri), in the St. Louis area, the Wabash Valley Fault zone along the southern Illinois-Indiana border, and the Anna region of western Ohio. Although high-intensity earthquakes have occurred along the New Madrid Fault zone, their relationship to plate motions remains speculative at this time.

According to estimates, ground motions induced by near and distant seismic sources in northern Illinois are expected to be minimal. However, peak accelerations in the ANL area may exceed 10% of gravity (approximate threshold of major damage) once in about 600 years, with an error range of -250 to +450 years.

# 1.7. Hydrology

Most groundwater supplies in the ANL area are derived from the Niagaran, and to some extent, the Alexandrian dolomite bedrock. Dolomite well yields are variable, but many are near 800 gallons per minute. In DuPage County, groundwater pumpage over the past 100 years has led to severe overdraft; in northeastern Illinois, the piezometric surface has been lowered in areas of heavy pumping. Delivery of Lake Michigan water to the major suburban areas is expected to relieve this problem. Because the cones of depression of ANL wells do not extend beyond the site and adjacent forest preserve, ANL water use does not affect neighboring communities.

Two principal aquifers are used as water supplies in the vicinity of ANL. The upper aquifer is the Niagaran and Alexandrian dolomite, which is about 60 m (200 ft) thick in the ANL area and has a piezometric surface between 15 and 30 m (50 and 100 ft) below the ground surface for much of the site. The lower aquifer is Galesville sandstone, which lies between 150 and 450 m (500 and 1,500 ft) below the surface. Maquoketa shale separates the upper dolomite aquifer from the underlying sandstone aquifer. This shale retards hydraulic connection between the two aquifers.

The four domestic water supply wells now in use on the ANL site (see Figure 1.1) are drilled about 90 m (300 ft) deep, terminating in the Niagaran dolomite. A well drilled in the Galesville sandstone 490 m (1,600 ft) deep has been taken out of service. The water level in the Niagaran dolomite has remained reasonably stable under ANL pumping, dropping about 6 m (20 ft) between 1960 and 1990. The aquifer appears to be adequate for future ANL use, but this ground water source is used throughout the area. Several small capacity water wells used for laboratory experiments, fire protection, and sanitary facilities also exist on the site, primarily in the 800 Area and meteorology complex.

#### 1.8. Water and Land Use

Sawmill Creek flows through the eastern portion of the site. This stream originates north of the site, flows through the property in a southerly direction, and discharges into the Des Plaines River. Two small streams originate on-site and combine to form Freund Brook, which discharges into Sawmill Creek. Along the southern margin of the property, the terrain slopes abruptly downward forming forested bluffs. These bluffs are dissected by ravines containing intermittent streams that discharge some site drainage into the Des Plaines River. In addition to the streams, various ponds and cattail marshes are present on the site. There is also a network of ditches and culverts that transport surface runoff toward the smaller streams.

The greater portion of the ANL site is drained by Freund Brook. Two intermittent branches of Freund Brook flow from west to east, draining the interior portion of the site and ultimately discharging into Sawmill Creek. The larger, south branch originates in a marsh adjacent to the western boundary line of the site. It traverses wooded terrain for a distance of about 2 km (1.5 mi) before discharging into the south branch at Lower Freund Pond.

Sawmill Creek carried effluent water continuously from a sewage treatment plant (Marion Brook Treatment Plant) located a few kilometers north of the site until October 27, 1986, when the plant was closed. Residential and commercial development in the area has resulted in the collection and channeling of runoff water into Sawmill Creek. Treated sanitary and laboratory wastewater from ANL are combined and discharged into Sawmill Creek at location 7M in Figure 1.1. This effluent averaged 2.5 million liters (0.66 million gallons) per day. This reduction from the 3.9 million liters discharged in 1993 is attributed to the completion of a sewer replacement project which eliminated infiltration. The combined ANL effluent consisted of 46% laboratory wastewater and 54% sanitary wastewater. The

water flow in Sawmill Creek upstream of the wastewater outfall averaged about 24 million liters (6.4 million gallons) per day during 1994.

Sawmill Creek and the Des Plaines River above Joliet, about 21 km (13 mi) southwest of ANL, receive very little recreational or industrial use. A few people fish in these waters downstream of ANL and some duck hunting takes place on the Des Plaines River. Water from the Chicago Sanitary and Ship Canal is used by ANL for cooling towers and by others for industrial purposes, such as hydroelectric generators and condensers, and for irrigation at the state prison near Joliet. The ANL usage is about 1.1 million liters (290,000 gallons) per day. The canal, which receives Chicago Metropolitan Sanitary District effluent water, is used for industrial transportation and some recreational boating. Near Joliet, the river and canal combine into one waterway, which continues until it joins the Kankakee River to form the Illinois River about 48 km (30 mi) southwest of ANL. The Dresden Nuclear Power Station complex is located at the confluence of the Kankakee, Des Plaines, and Illinois rivers. This station uses water from the Kankakee River for cooling and discharges the water into the Illinois River. The first downstream location where water is used for drinking is at Alton, on the Mississippi River about 710 km (370 mi) downstream from ANL. At that location, water is used indirectly to replenish groundwater supplies by infiltration. In the vicinity of ANL, only subsurface water (from both shallow and deep aquifers) and Lake Michigan water are used for drinking purposes.

The principal recreational area near ANL is Waterfall Glen Forest Preserve, which surrounds the site (see Section 1.2 and Figure 1.1). The area is used for hiking, skiing, and equestrian sports. Sawmill Creek flows south through the eastern portion of the preserve on its way to the Des Plaines River. Several large forest preserves of the Forest Preserve District of Cook County are located east and southeast of ANL and the Des Plaines River. The preserves include the McGinnis and Saganashkee Sloughs (shown in Figure 1.2), as well as other, smaller lakes. These areas are used for picnicking, boating, fishing, and

hiking. A small park located in the eastern portion of the ANL site (Location 12-0 in Figure 1.1) is for the use of ANL and DOE employees. Recently, use of this park has also been provided to DuPage County.

# 1.9. Vegetation

ANL lies within the Prairie Peninsula of the Oak-Hickory Forest Region. The Prairie Peninsula is a mosaic of oak forest, oak openings, and tall-grass prairie occurring in glaciated portions of Illinois, northwest Indiana, southern Wisconsin, and sections of other states. Much of the natural vegetation of this area has been modified by clearing and tillage. Forests in the ANL region, which are predominantly oak-hickory forests, are somewhat limited to slopes of shallow, ill-defined ravines or of low morainal ridges. Gently rolling to flat intervening areas between ridges and ravines were predominantly occupied by prairie before their use for agriculture. The prevailing successional trend on these areas, in the absence of cultivation, is toward oak-hickory forest. Forest dominated by sugar maple, red oak, and basswood may occupy more pronounced slopes. Poorly drained areas, streamside communities, and floodplains may support forests dominated by silver maple, elm, and cottonwood.

From early photographs of the site, it appears that most of the land that ANL now occupies was actively farmed. About 75% was plowed field and 25% was pasture, open oak woodlots, and oak forests. Starting in 1953 and continuing for three seasons, some of the formerly cultivated fields were planted with jack, white, and red pine trees. Other fields are dominated by bluegrass.

Crown vetch has been planted on much of the developed area since 1954, to help control soil erosion and provide low-maintenance ground cover. Other open space in developed areas has been sown to grass, which is mowed regularly.

The deciduous forests on the remainder of the site are dominated by various species of oak, generally as large, old, widely spaced trees, often not forming a complete canopy. Their large low branches indicate that they probably matured in the open, rather than in a dense forest. Other upland tree species include hickory, hawthorn, cheery, and ash.

#### 1.10. Fauna

Terrestrial vertebrates that are commonly observed or likely to occur on the site include about five species of amphibians, seven of reptiles, and about 40 species of summer resident birds, and 25 of mammals. More than a hundred other bird species occur in the area during migration or winter but do not nest on the site or in the surrounding region. An unusual species on the ANL site is the fallow deer, a European species that was introduced to the area by a private landowner prior to government acquisition of the property in 1947 and which subsequently increased to about 400 individuals. In November 1988, about 200 of the deer were removed for population control. A significant population of native white-tailed deer also occur on the ANL reservation. Invertebrate species, as well as plants and other animals, were also observed on the ANL site.

Freund Brook crosses the center of the site, but is impounded by a beaver dam in this area. The gradient of the stream is relatively steep, and riffle habitat predominates. The substrate is coarse rock and gravel on a firm mud base. Primary production in the stream is limited by shading, but diatoms and some filamentous algae are common. Aquatic macrophytes include common arrowhead, pondweed, duckweed, and bulrush. Invertebrate fauna consist primarily of dipteran larvae, crayfish, caddisfly larvae, and midge larvae. Few fish are present because of low summer flows and high temperatures. Other aquatic habitats on the ANL site include additional beaver ponds, artificial ponds, ditches, and Sawmill Creek.

The biotic community of Sawmill Creek is relatively depauperate, reflecting creek's high silt load, steep gradient, and historic release of sewage effluent from the Marion Brook sewage-treatment plant north of the site. The fauna consists primarily of blackflies, midges, isopods, flatworms, segmented worms, and creek chubs. A few other species of minnows, sunfishes, and catfish are also present. Clean water invertebrates, such as mayflies and stoneflies, are rare or absent. The fish species that have been recorded in ANL aquatic habitats include black bullhead, bluegill, creek shub, golden shiner, goldfish, green sunfish, largemouth bass, stoneroller, and orange-spotted sunfish.

The Des Plaines River system, including ANL streams, has been rated as "poor" in terms of the fish species present, as determined by the U. S. Fish and Wildlife Service, a result of domestic and industrial pollution and stream modification.

# 1.11. Archaeology

ANL, located in the Illinois and Michigan Canal National Heritage Corridor, is situated in an area known to have a long and complex cultural history. All periods listed in the cultural chronology of Illinois, with the exception of the earliest period (Paleo-Indian), have been documented in the ANL area by either professional cultural resource investigation or by interviews of ANL staff with local collectors. A variety of site types, including mounds, quarries, lithis workshops, and habitation sites have been reported by amateurs within a 25-km (16-mi) radius of ANL.

There are 26 recorded sites including prehistoric chert quarries, special purpose camps, base camps, and historical farmsteads. The range of human occupation spans several time periods (Early Archaic through Mississipian Prehistoric to Historical). To date, one site may be eligible and 19 of the sites are not eligible for the National Register of

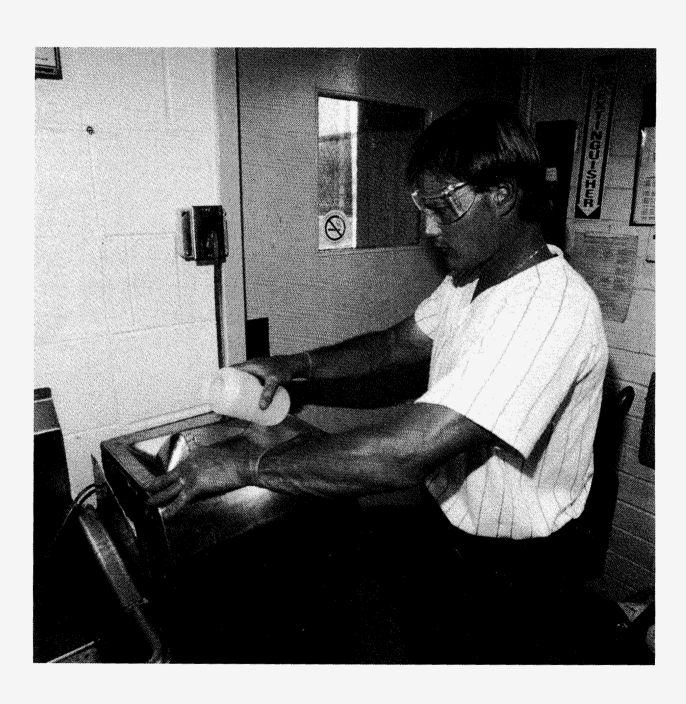
Historic Places (NRHP); the remainder have not been formally evaluated for NRHP eligibility.

# 1.12. Endangered Species

Although the geographic ranges of several federally listed animal species include the northern Illinois region, no suitable habitat for these species is present on the site, with the possible exception of the Indiana bat (Myotis sodalis). An unconfirmed capture of an Indiana bat in nearby Waterfall Glen Forest Preserve indicates that the bat may occur in the ANL region. Consultation with the U. S. Fish and Wildlife Service determined that suitable habitat for this species does not exist in the area that would be affected by APS construction. The bald eagle, peregine falcon, piping plover, interior least tern, and Kirtland's warbler could occur in the ANL area as extremely rare nonbreeders during migration or winter.

Neither federal nor state-listed threatened or endangered species are known to reside on the ANL site. The federally-listed endangered Indiana Bat, Myotis sodalis, and the federally-threatened Hine's Emerald Dragonfly, Somatochlora hineana, reside in the area and may possibly reside on the ANL site. Two state-endangered species, River Otter, Lutra canadensis, and White Lady's Slipper, Cypripededium candidum; and two state-threatened species Kirtland's Snake, Clonophis kirtland; and sedge, Carex crawei, reside in the area and may possibly reside on the ANL site.

# 2. COMPLIANCE SUMMARY



# 2. COMPLIANCE SUMMARY

ANL is a government-owned, contractor-operated (GOCO) non-production facility which is subject to environmental statutes and regulations administered by the U. S. Environmental Protection Agency (EPA), the Illinois Environmental Protection Agency (IEPA), the Illinois Department of Public Health (IDPH), and the State Fire Marshal, as well as numerous DOE Orders and Executive Orders. A detailed listing of applicable regulations is contained in DOE Order 5400.1, which establishes DOE's policy concerning environmental compliance. The status of ANL during 1994 with regard to these authorities is discussed in this Chapter.

To insure compliance with both the letter and spirit of these requirements, ANL has made a commitment to comply with all applicable environmental requirements, as described in the following policy statement revised during 1990:

It is the policy of Argonne National Laboratory that its activities will be conducted in such a manner that worker and public safety, including protection of the environment, is given the highest priority. The Laboratory will comply with all applicable federal and state environmental laws, regulations, and orders.

#### 2.1. Clean Air Act

The Clean Air Act (CAA) is a federal statute that specifies National Ambient Air Quality Standards, sets emission limits for air pollutants and determines emission limits and operating criteria for a number of hazardous air pollutants. The program is implemented by individual states through a State Implementation Plan (SIP), which describes how that state will ensure compliance with the air quality standards for stationary sources.

A number of major changes to the CAA were made with the passage of the Clean Air Act Amendments of 1990. Under Title V of the 1990 Amendments to the CAA, ANL is required to apply for a sitewide, federally enforceable operating permit to supersede the 35 state air permits currently in effect. All facilities that are designated as major emission sources for regulated air pollutants are subject to this requirement. ANL meets the definition of a major source due to potential emissions of oxides of nitrogen (NO<sub>x</sub>) in excess of 25 tons per year and sulfur dioxide (SO<sub>2</sub>) in excess of 100 tons per year at the Building 108 Central Heating Plant.

Facilities subject to Title V must characterize emissions of all regulated air pollutants, not only those that make them qualify as major sources. For ANL, in addition to  $NO_x$  and  $SO_2$ , emissions of carbon monoxide, particulates, volatile organic compounds, hazardous air pollutants (a list of 189 chemicals, including radionuclides), and ozone-depleting substances must also be evaluated. The permit program requires that facilities pay annuals fees based upon the total amount of regulated air pollutants (except carbon monoxide) they will be allowed to emit.

Although it is a federal program, the permit program will be administered by the IEPA under EPA oversight. In 1994, a consultant was contracted to provide support for preparation of the ANL CAA Title V permit application. In February 1994, a six-week stack test was conducted at the ANL Central Heating Plant to obtain data for the Title V inventory on the site's major emission source. During the summer of 1994, all divisions were visited to review their 1992 sitewide emission inventory and obtain baseline information for the application. In November 1994, a meeting at IEPA with ANL, DOE and the consultant was held to discuss the approach for preparing the application and clarify technical issues. IEPA has officially notified ANL that the permit application would be due by December 7, 1995. The final forms for Illinois' Clean Air Act Permit Program (CAAPP), which is the state designation for the Title V permit, were received in January 1995.

Plans for 1995 are completion of the CAAPP application and submittal to IEPA in late summer. Issuance of a CAAPP permit would be expected sometime in 1997.

The ANL site contains a large number of air emission point sources. The vast majority are laboratory ventilation systems which are exempt from state permitting requirements, except for those systems emitting radionuclides. By the end of 1994, a total of 35 air permits were in place covering all known emission points. Section 2.15 contains a listing (Table 2.9) of the permits in effect at ANL.

#### 2.1.1. National Emission Standards for Hazardous Air Pollutants

The National Emission Standards for Hazardous Air Pollutants (NESHAP) are a body of federal regulations that set forth emission limits and other requirements, such as monitoring, record keeping, and operational requirements, for activities generating emissions of certain hazardous air pollutants. The standards for asbestos, radionuclides, and halogenated solvent cleaning are the only standards affecting ANL operations. A total of 16 radiological air permits have been issued by IEPA to ANL.

#### 2.1.1.1. Asbestos Emissions

Many buildings on the ANL site contain large amounts of asbestos-containing materials (ACM), such as thermal system insulation around pipes and tanks, spray applied surfacing material for fireproofing, floor tile, and asbestos-cement panels. This material is removed as necessary during renovations or maintenance of equipment and facilities. The removal and disposal of this material is governed by the asbestos NESHAP.

The standards for asbestos specify detailed requirements for removal and disposal of certain types of ACM. Until the November 1990 revisions, only friable (easily crushed) ACM was regulated. Now, however, many other types of ACM are regulated, including non-friable materials which have been, or could be reduced to a crumbly, pulverized or powder state through the process of removal or disposal. This change greatly increases the amount of material regulated by the NESHAP.

The standard describes accepted procedures for removal of ACM, including notification of the IEPA prior to removal of greater than certain amounts, work practices and procedures to be used and emission control procedures to be used. The use of specially trained individuals for removal of ACM is mandated.

ANL maintains an asbestos abatement program designed to assure compliance with these and other regulatory requirements. The removal of ACM at the Laboratory is done either by a specially trained Waste Management Operations (WMO) crew (for "small-scale" short-duration projects as defined by the OSHA asbestos standard for the Construction Industry - 29 CFR 1926.58) or by outside contractors specializing in ACM removal work (for larger-scale insulation removal projects lasting a day or longer). All removal work is done in strict compliance with both the NESHAP requirements as well as the OSHA requirements governing worker safety at ACM removal sites. When ACM is encountered during a renovation or demolition project, it is carefully wetted or otherwise encapsulated and completely removed. The work area is sealed off using disposable glove bags or temporary plastic sheeting barriers, and high-efficiency particulate air (HEPA) filtration equipment is used to control emissions. Air is monitored in the vicinity of such work by ANL Industrial Hygiene personnel both during the removal work and after the work is completed, in order to verify that adequate precautions have been taken to prevent the release of significant amounts of asbestos. Personal

exposure air samples are collected. Asbestos fiber counts are analyzed using Phase Contrast Microscopy and selected samples are analyzed by Transmission Electron Microscopy.

Approximately 153 m<sup>3</sup> (5400 ft<sup>3</sup>) of ACM were removed from ANL buildings during 1994. These materials included various structural or facility components such as surfacing materials, thermal system insulation, floor tile and mastic, and transite wall-board. Also included are items which were part of the removal activity and became contaminated with asbestos, such as Tyvek coveralls, gloves, and polyethylene sheets. Asbestos-containing laboratory equipment such as an oven, gloves, and asbestos-covered wire were also removed. The primary types of asbestos identified in these materials were chrysotile and amosite.

Approximately 1818 m<sup>3</sup> (64,150 ft<sup>3</sup>) of nonfriable asbestos-containing transite sewer pipe and excavated soil, and 2.7 m<sup>3</sup> (95 ft<sup>3</sup>) of friable pipe insulation and related removal items were disposed of as part of the laboratory and sanitary sewer rehabilitation project.

Most of the ANL asbestos removal activities are "small-scale short-duration" projects as defined by the OSHA Asbestos Standard for the Construction Industry (29 CFR 1926.58). The duration of these projects is usually not more than four hours and glovebag removal techniques are normally used. A total of 142 small removal projects were completed which generated 119 m³ (4183 ft³) of ACM waste. Projects performed by outside contractors accounted for 71 m³ (2500 ft³) of the ACM waste from small projects.

The asbestos NESHAP standards require that the IEPA be notified before beginning large asbestos removal projects involving more than 80 m (260 ft) of pipe insulation or 15 m<sup>2</sup> (160 ft<sup>2</sup>) of other materials or 1 m<sup>3</sup> (35 ft<sup>3</sup>) of ACM where the length or area cannot be measured. The Notification of Demolition and Renovation Form must be forwarded to the IEPA within a prescribed time limit. Six Notification of Demolition and Renovation forms were provided to the IEPA during 1994. The total waste from building projects was 37 m<sup>3</sup> (1309 ft<sup>3</sup>). Project information is provided in Table 2.1.

One project was cancelled. For two projects to (1) excavate nonfriable sewer pipe, and (2) remove floor tile from Building 205, courtesy notifications were provided because these nonfriable materials were not subject to NESHAP reporting requirements.

The NESHAP requires estimation of the total amount of ACM to be removed during renovation or demolition activities during each upcoming calendar year. If this amount exceeds the regulatory levels given above, the IEPA must be notified. ANL made such a notification during December 1994 and February 1995 (revised) for activities planned for 1995. It is estimated that no more than 128 m³ (4500 ft³) of ACM waste will be generated during 1995.

A separate portion of the standard contains requirements for waste disposal sites used for disposal of ACM. The acceptable disposal practice involves placing wetted waste materials into labeled, leakproof plastic bags for disposal in landfills. Off-site shipments are to be accompanied by completed shipping manifests. The principal requirements applicable to landfill disposal of ACM relate to covering the ACM daily with at least six inches of non-asbestos-containing materials and maintenance of disposal records. All shipments except one, which went to Pekin Metro Landfill, Pekin, Illinois, went to Community Landfill Company, Morris, Illinois.

TABLE 2.1

# Asbestos Abatement Projects IEPA Notification, 1994

Completion Date	Notification Quantity (ft)	Notification Quantity (ft²)	Material	Building	Disposal Quantity (ft³)
February 26, 1994	-	540	Floor Tile	600	60
March 15, 1994	70	145	Pipe and Tank Insulation	362	219
None	80	245	Pipe and Tank Insulation	108	None
August 30, 1994	3800	-	Transite Sewer Pipe and Soil	NE Site	64,152
September 25, 1994	-	4,000	Floor Tile	205	540
October 31, 1994	160	2000	Pipe Insultation, Transite Panels	813A, 821, 827 827B	490

Until closure of the ANL landfill in September 1992, asbestos from small-scale projects was disposed on-site in a designated area of the landfill. IEPA conducted an asbestos disposal inspection on April 28, 1994. Noncompliance with 40 CFR 61.154 (g) - failure to modify deed to reflect the presence of asbestos material - was cited. DOE is conducting a survey of the site and the landfill. The deed will be modified after the survey is completed.

#### 2.1.1.2. Radionuclide Emissions

The NESHAP standard for radionuclide emissions from DOE facilities (40 CFR 61, Subpart H) establishes the emission limits for release of radionuclides to the air and requirements for monitoring, reporting, and record keeping. This regulation was revised in late 1989, resulting in increased monitoring and reporting requirements. A number of emission points at ANL are subject to these requirements. These points include ventilation systems for hot cell facilities for storage and handling of radioactive materials (Buildings 200, 205, and 212), ventilation systems for inactive reactors (Building 330, inactive reactor CP-5), ventilation systems for particle accelerators (Building 375, IPNS facility and the Building 411 APS Linac), and several ventilation systems associated with the New Brunswick Laboratory (Building 350). In addition, many small ventilation systems and fume hoods are occasionally used for processing of small quantities of radioactive materials. The radionuclide NESHAP requires that all air emission sources of radionuclides be evaluated to determine whether the magnitude of these emissions is above a threshold amount which would result in an effective dose equivalent to the maximally exposed individual of greater than 1% of the standard of 10 mrem/yr. Those sources with greater than this amount of emissions must be monitored in accordance with 40 CFR 61.93(b) and a report issued annually summarizing the emissions measured. Any emission point below this threshold must be measured periodically to verify the low rate. At ANL, the significant emission sources are continuously monitored to comply with this requirement. However, to satisfy the determination for monitoring requirements for the large number of smaller sources, all radionuclide air emission sources have been evaluated. The emissions from NBL are included with ANL emissions when calculating dose rates under NESHAP.

Routine continuous monitoring of the larger emission sources has indicated that the amount of radioactive material released to the atmosphere from these sources is extremely small, resulting in a very small incremental radiation dosage to the neighboring population. The calculated potential maximum individual off-site dose to a member of the general public for 1994 was 0.015 mrem which, excluding radon-220, is 0.15% of the 10 mrem per year EPA standard. Section 4.6.1. contains a more detailed discussion of these emission points and compliance with the standard.

## 2.1.1.3. Halogenated Solvent Cleaner Emissions

The NESHAP standard for halogenated solvent cleaner emissions (40 CFR 63, Subpart T) became effective on December 2, 1994, and establishes emission limits on a variety of cleaning and degreasing activities utilizing a number of halogenated solvents. The Central Shops (Building 363) vapor degreaser is subject to this regulation since a listed solvent (perchloroethylene) is used in cleaning small metal parts on a batch basis. The NESHAP requires notification to IEPA by August 29, 1995, of the presence of the vapor degreaser and the applicability of the rule to its operations. The major effect of the regulation will be to maintain recordkeeping on solvent use to verify that the specified emission limits are not exceeded. Compliance with the NESHAP is required no later than December 2, 1997.

#### 2.1.2. Conventional Air Pollutants

The ANL site contains a number of sources of conventional air pollutants, including a steam plant, oil-fired boilers, gasoline and methanol fuel dispensing facilities, two alkali metal reaction booths, a number of bulk chemical tanks, a dust collection system, a medical equipment sterilization unit, and fire training activities. The emission sources that have been granted operating permits by the IEPA are as shown in Section 2.15. During 1994, four new air permits were issued by the IEPA, three air permits were modified, and nine air permits were renewed.

The operating permit for the steam plant requires continuous opacity and sulfur dioxide monitoring of the smoke stack from Boiler No. 5, the only one of the five boilers equipped to burn coal. The permit requires submission of a quarterly report listing any excursions beyond emission limits for this boiler [30% opacity averaged over six minutes and 1.8 lb sulfur dioxide (SO<sub>2</sub>) per million Btu averaged over a one-hour period]. The hours of operation during 1994 of Boiler No. 5 on high and low sulfur coal is presented in Table 2.2. During the first quarter of 1994 due to equipment failure, one excursion for opacity was noted. The last of the high sulfur coal was burned in January 1994. EPA and IEPA conducted an air emission compliance inspection of the steam plant on March 4, 1994, and June 14, 1994, respectively.

Fuel dispensing facilities include a commercial service station and Building 46 Grounds and Transforation facility. Except for methanol vapors, theses facilities have VOC emissions typical of any commercial gasoline service station. Stage II vapor recovery systems were installed at both facilities by November 1, 1994.

TABLE 2.2

Boiler No. 5 - Hours of Operation, 1994

Month	w/High Sulfur Coal	w/Low Sulfur Coal	Total	
January	504	240	744	
February	0	548	548	
March	0	147.5	147.5	
April	0	180	180	
May	0	0	0	
June	0	0	0	
July	0	0	0	
August	0	0	0	
September	0	0	0	
October	0	0	0	
November	0	0	0	
December	0	48	48	
Total	504	1163.5	1667.5	

#### 2.2. Clean Water Act

The Clean Water Act (CWA) was established in 1977 as a major amendment to the Federal Water Pollution Control Act of 1972 and was substantially modified by the Water Quality Act of 1987. The CWA provides for the restoration and maintenance of water quality in all waters throughout the country, with the ultimate goal of "fishable and swimmable" water quality. The act established the NPDES permitting system, which is the regulatory mechanism designed to achieve this goal. The authority to implement the NPDES program has been delegated to those states, including Illinois, that have developed a program substantially the same and at least as stringent as the federal NPDES program.

The 1987 amendments to the CWA significantly changed the thrust of enforcement activities. Greater emphasis is now placed on monitoring and control of toxic constituents in wastewater, the permitting of outfalls composed entirely of stormwater, and the imposition of regulations governing sewage sludge disposal. These changes in the NPDES program resulted in much stricter discharge limits and greatly expanded the number of chemical constituents monitored in the effluent. The wastewater treatment facilities on the ANL site will be upgraded to improve treatment capabilities.

# 2.2.1. Liquid Effluent Discharge Permit

The primary tool for enforcing the requirements of the NPDES program is through the NPDES permitting process administered by the IEPA. Before wastewater can be discharged to any receiving stream, each wastewater discharge point (outfall) must be characterized and described in a permit application. The IEPA then issues a permit that contains numeric limits on certain pollutants likely to be present and sets forth a number of specific and general requirements, including sampling and analysis schedules and reporting and recordkeeping requirements. Wastewater generation activities at ANL are covered by NPDES Permit IL 0034592. This permit was renewed during 1994 and became effective October 30, 1994. The new permit has a major impact on ANL. Although outfalls 003, 005, and 009 were removed from sampling requirements, more locations (20) with increased sampling frequencies and sampling parameters were added. Additionally, more restrictive discharge limits were placed on ANL discharges.

Wastewater at ANL is generated by a number of activities and consists of sanitary wastewater (from restrooms, cafeteria sinks and sinks in certain buildings and laboratories, steam boiler blowdown, and drinking water filter backwash), laboratory wastewater (from laboratory sinks and floor drains in most buildings), and stormwater. Water softener regenerant is discharged to the DuPage County sewer system. Cooling water and cooling tower blowdown are currently discharged into stormwater ditches which are monitored as part of the NPDES permit. The current permit authorizes the release of wastewater from 26 separate outfalls (compared to nine on the previous permit), most of which discharge directly or indirectly into Sawmill Creek. In addition, the permit requires monitoring of the wastewater at two internal sampling points that combine to form the main wastewater outfall, outfall 001. Table 2.3 describes these outfalls, and the locations are shown in Figure 2.1. Two of these outfalls, 009 and 010, are used for emergency overflow discharge from the lime sludge pond and coal pile, respectively. Outfall 009 was removed from the permit by the IEPA during 1994.

# 2.2.1.1. Effluent Monitoring Results and Compliance Issues

Results of the routine monitoring required by the NPDES permit are submitted monthly to the IEPA in a Discharge Monitoring Report (DMR). As required by the permit, any exceedance of permit limits or conditions is reported by telephone to the

### 2. COMPLIANCE SUMMARY

**TABLE 2.3** 

Description of NPDES Outfalls at ANL, 1994

Outfall	Description	Flow	
001A	Sanitary Treatment Plant	0.6	MGD
001B	Laboratory Treatment Plant	0.4	MGD
001	Combined Outfall	1.0	MGD
003A	Swimming Pool	500	GPD
003B	300 Area (Condensate)	1000	GPD
003C	Building 205 Footing Tile Drainage	1000	GPD
003D&E	Steam Trench Drainage (Condensate)	1000	GPD
003F	Building 201 Fire Pond	0	GPD
003G	North Building 201 Storm Sewer (Condensate)	500	GPD
003H	Building 212 Cooling Tower Blowdown	1000	GPD
003I	Buildings 200 and 211 Cooling Tower Blowdown	1000	GPD
003J	Building 213 and Building 213 Parking Lot Stormwater	0	GPD
004	Building 203 Cooling Tower and Building 221 Footing Drainage	0.05	MGD
005A	Westgate Road Stormwater	Stormwater	Only
005B	800 Area East Stormwater	Stormwater	Only
005C	Building 200 West	1000	GPD
005D	Stormwater	Stormwater	Only

# 2. COMPLIANCE SUMMARY

TABLE 2.3 (Contd.)

Outfall	utfall Description		
005E	Building 203 West Footing Drainage and Condensate	500 GPD	
006	Cooling Tower Blowdown	0.13 MGD	
007	Domestic Cooling Water for Compressor	0.02 MGD	
008	Transportation and Grounds Stormwater	Stormwater Only	
010	Coal Pile Runoff Emergency Overflow	Stormwater Only	
101	North Fenceline Marsh Storm Discharge	Stormwater Only	
102	100 Area Stormwater Discharge	Stormwater Only	
103	Southeast 100 Area Stormwater	Stormwater Only	
104	Northern East Area Stormwater Discharge	Stormwater Only	
105A&B	Building 40 Stormwater Discharge	Stormwater Only	
106 A&B	Southern East Area Stormwater Discharge	Stormwater Only	
108	Eastern 300 Area Stormwater and Cooling Water	0.02 MGD	
110	Shooting Range Stormwater Discharge	Stormwater Only	
111	319 Landfill and Northeast 317 Area	Stormwater Only	
112A&B	Southern and Western 317 Area	Stormwater Only	
113	Southern and Eastern 800 Area Landfill	Stormwater Only	
114	Northern and Western 800 Area Landfill	Stormwater Only	
115	314, 315, and 316 Cooling Water, Eastern and Southern APS Construction Area	0.03 MGD	
116	Water Treatment Plant and Stormwater	0.03 MGD	

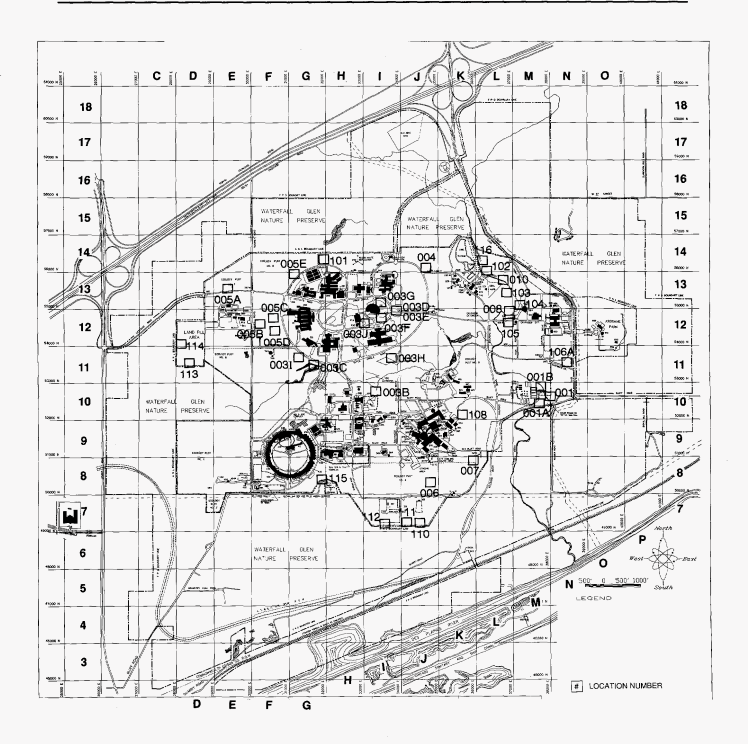
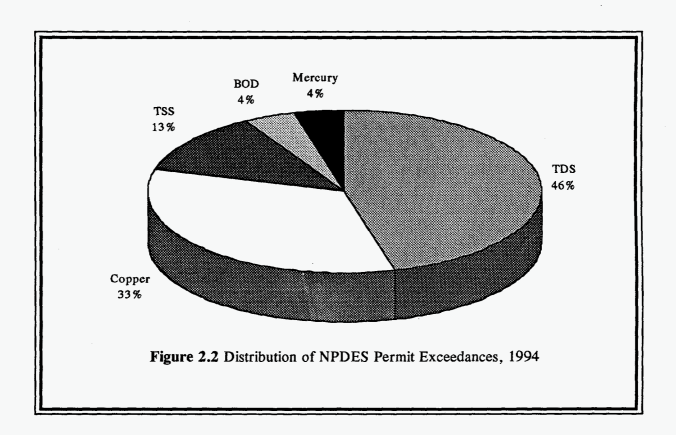


Figure 2.1 NPDES Permit Locations, 1994

IEPA within 24 hours, and a written explanation of the exceedance is submitted with each DMR. During 1994, there were 24 exceedances of NPDES permit limits out of approximately 1000 measurements. This represents a 97.5% compliance rate, similar to the 1993 compliance rate (25 exceedances), 98% compliance rate in 1992 (19 exceedances), a 96% rate in 1991 (44 exceedances), and a 91% rate in 1990 (86 exceedances).

Many of the exceedances experienced were new in comparison to recent years due to new requirements. A statistical breakdown appears in Figure 2.2.

The largest category (11) is total dissolved solids (TDS) exceedances at outfall 001. The source of the TDS underwent an intensive investigation during May 1994. By



monitoring manholes upstream and downstream from potential sources of TDS, the source of regular, high-TDS discharges into the laboratory wastewater sewer system from the area of the domestic water treatment facility was identified. This discharge appears to have originated from the recycle-brine tank overflow piping which connected to the laboratory wastewater sewer system. The recycle-brine overflow was redirected to the DuPage County sewer system for disposal along with other brine wastes from the facility. During late 1994, more exceedances of TDS appeared. Discharge of effluents from the boiler operations may be an active source. With supply-water TDS concentrations in the range of 750 mg/L (well head) to 950 mg/L (treated water) and the new NPDES permit limit at 1000 mg/L, the capacity for dissolved solids addition to the site wastewater is quite limited. Lake Michigan water, which has a much lower TDS concentration, will be incorporated as the ANL source water by late 1997.

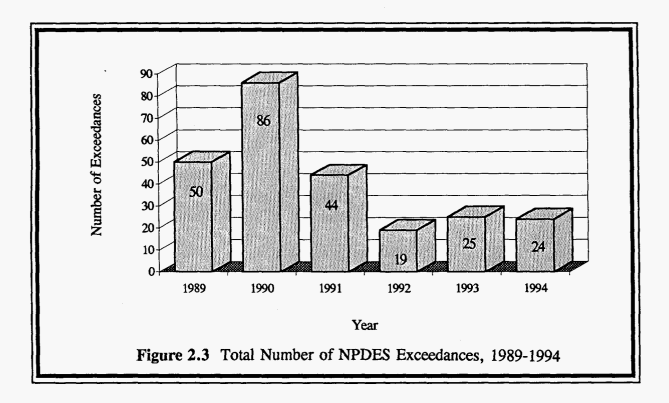
Eight exceedances of copper at outfall 001 were noted. The well water obtained by ANL from the Niagaran dolomite has very low copper concentrations. However, after treatment and distribution through copper pipe, which is used for the domestic water distribution within ANL buildings, a copper concentration range of 0.5 mg/L to 1.0 mg/L is typical at drinking fountains. This range has been determined by the copper/lead monitoring program required by EPA (Section 6.1.1.). The acceptable copper level for human consumption is significantly above the NPDES Permit limit at outfall 001 of 0.051 mg/L. There is no process in the wastewater treatment plant to remove copper. Past samples collected from the wastewater treatment plant effluent have been below the state effluent limit of 0.5 mg/L, but concentrations in Sawmill Creek, below the point where the treated wastewater has been discharged, have consistently exceeded the state stream standard for copper (0.02 mg/L) for the past ten years. This may be indicative of the current ambient levels of copper in surface water as a result of the increased usage of copper pipe. The corrective action currently being taken by ANL to control copper levels in the domestic water supply includes the injection of

polyphosphate and zinc to coat the copper piping. Addition of zinc has a synergistic effect for the coating of the pipes.

Total suspended solids (TSS) exceedances accounted for only 13% of the total. TSS was related to excessive surface runoff from heavy precipitation at outfalls 003A and 004. Sludge carryover from laboratory wastewater holding tanks was responsible for the TSS excursion at outfall 001B.

Discharges from outfalls 003A, 007, and 116 are monitored for total residual chlorine (TRC). Special Condition 8 of the NPDES Permit states that ANL shall achieve compliance with a 0.05 mg/L TRC limit as soon as possible, but no later than two years after October 30, 1994. The discharges from these outfalls consistently exceeded the 0.05 mg/L limit during November and December 1994. The cause of these elevated levels is the infiltration of chlorinated potable water due to water main leakage (003A), water valve malfunction (116), and cross connection of a potable drain to the stormwater collection system, most likely in either a building maintenance area or a programmatic research area (007). Corrective actions to eliminate chlorinated potable water have so far included repair of a swimming pool discharge (003A) and water valve (116). Investigations are continuing to discover the potable water cross connections near outfall 007.

Data regarding the total number of each type of exceedance over the past six years is presented in Figure 2.3. From 1989 to 1994, the total number of exceedances has been reduced; 50 exceedances in 1989, 86 exceedances in 1990, 44 exceedances in 1991, 19 exceedances during 1992, 25 exceedances during 1993, and 24 exceedances in 1994.



The NPDES Permit, effective October 30, 1994, placed more restrictive limits on ANL discharges. Specifically, for reasons stated above, copper limits have resulted in a substantial increase of exceedances. Future exceedances of the TDS limit remain possible since only moderate additions of soluble materials to the wastewater stream could result in exceedances. Water conservation efforts currently being implemented at ANL may be expected to decrease the dissolved solids capacity of the wastewater system and thereby make TDS compliance even more difficult in the future. Prior to November 1994, ANL experienced 12 NPDES exceedances. During November and December 1994 alone, 12 exceedances were noted, 67% due to exceedance of the new copper limits.

To improve the level of compliance with permit limits, ANL is in the fourth year of an intensive effort to build additional wastewater treatment facilities or upgrade existing facilities. Projects to upgrade and refurbish the laboratory and sanitary

wastewater treatment plant are scheduled for 1995 through 1996. Upgrade and repair of a large portion of ANL's wastewater sewer system took place during 1994. These and other corrective action projects are described in the Environmental Management Five Year Plan for ANL and identified in Chapter 3.

#### 2.2.1.2. Additional NPDES Monitoring

The current permit requires semiannual testing of outfall 001B, the laboratory wastewater treatment plant outfall, for all the priority pollutants (a list of 126 metals and organic compounds identified by the IEPA as being of particular concern). During 1994, this sampling was conducted in June and December. Chloroform (4  $\mu$ g/L and 33.6  $\mu g/L$ ), dibromochloromethane (2.1  $\mu g/L$  and 1.4  $\mu g/L$ ), and bromodichloromethane (2.5 μg/L and 1.7 μg/L) were detected in both the June and December samples at low concentrations which resulted from normal ANL operations. Low concentrations of bromoform (2.1  $\mu$ g/L) were noted in the June sample and methylene chloride (123  $\mu$ g/L) was found in the December sample. The source of most of these materials is suspected to be from the contact of chlorinated water with organic chemicals in the laboratory, as well as the discharge of small amounts of chemicals from various research and support operations. All semivolatiles concentrations were below the detection limits. Low concentrations of zinc (0.076 mg/L), copper (0.049 mg/L), and cyanide (0.028 mg/L) were detected. Lead was detected in the December sample at low concentrations (0.006) mg/L). No chrysotile asbestos greater than 10 millimicrons in length was detected. Dioxins were not detected. These findings are discussed further in Chapter 5.

In addition to the priority pollutant analysis, the permit requires annual biological toxicity testing of the combined effluent stream, outfall 001. This was done between June 21 and June 29, 1994. The data indicate that no adverse effects were noted in any of the toxicity tests at any concentration. Comparing 1994 results with previous years'

data, it appears that the cessation of chlorination of ANL's effluent correlated with a beneficial effect on aquatic life in the receiving streams.

The new requirements of the revised NPDES Permit will require acute toxicity testing at outfalls 003H, 003I, 003J, 004, 006, and 115 twice a year during the months of July and August, as well as the annual acute toxicity testing during June at outfall 001.

As a result of the outfall 001 effluent exhibiting acute toxicity to the fathead minnow and Ceriodaphnia for two consecutive years, on September 10, 1992, the IEPA requested that ANL continue the biomonitoring plan and initiate a Toxicity Reduction Evaluation (TRE). The purpose of the TRE is to identify the substance or substances causing whole effluent toxicity and to propose solutions to the problem. A Toxicity Identification Evaluation (TIE) as an initial phase to the project was completed during September 1993. Testing performed in 1993 at the sanitary, laboratory, and combined wastewater outfalls showed that toxicity was present in each system, but it was not consistent and was usually at low concentrations. Two TRE toxicity screening tests were performed to determine the group of chemicals responsible for the toxicity, however, the toxicity was at low levels and the responsible group of chemicals could not be identified. All results were submitted to the IEPA during December 1993.

TRE testing for acute toxicity at the laboratory wastewater (outfall 001B), sanitary wastewater (outfall 001A), and stream (NPDES outfall 001) continued during 1994. Results were submitted to the IEPA during December 1994 and showed that although some toxicity occurred at outfalls 001A and 001B, the effects were minor at the combined outfall 001.

A biosurvey of Sawmill Creek was also performed during the summer of 1994. The results showed that ANL discharges had no impact on the biota of Sawmill Creek. IEPA reviewed the biosurvey report and concurred with ANL that the ANL wastewater does not affect the biota of Sawmill Creek.

#### 2.2.2. Stormwater Regulations

In November 1990, the EPA promulgated new regulations governing the permitting and discharge of stormwater from industrial sites. The ANL site contains a large number of small scale operations which are considered industrial activities by the regulation, and thus, is subject to these requirements. To satisfy the stormwater permit application information needs, an extensive stormwater characterization program began in 1991 resulting in a stormwater permit application sent to the IEPA during 1992.

The NPDES Permit Special Conditions address ANL's stormwater control needs. ANL is required to develop a stormwater pollution prevention plan for stormwater associated with industrial activities from outfalls 003B, 003C, 003G, 003H, 003I, 003J, 004, 005A, 005B, 005C, 005D, 006, 007, 008, 101, 102, 103, 104, 105A, 105B, 106A, 106B, 108, 110, 111, 112A, 112B, 113, 114, 115, and 116. Potential non-stormwater discharges from outfalls 101, 102, 103, and 104 are to be addressed as well as existing sources of pollution which may be expected to affect the quality of stormwater, such as janitorial drains, from outfalls 003C, 003H, 003J, and 005C.

In addition to the above requirement, ANL is to develop a stormwater prevention plan for stormwater runoff associated with construction site activity from outfalls 008, 104, and 115.

#### 2.2.3. NPDES Inspections and Audits

On October 12-13, 1994, the IEPA conducted a Compliance Sampling Inspection of NPDES outfalls and related facilities, as well as associated sampling and analysis and record keeping requirements. No significant issues were identified.

#### 2.2.4. General Effluent and Stream Quality Standards

In addition to specific permit conditions, ANL discharges are required to comply with general effluent limits contained in 35 Illinois Administrative Code, Subtitle C, Chapter I, Part 304. Also, wastewater discharges must be of sufficient quality to insure that Sawmill Creek complies with the IEPA's General Use Water Quality Standards found in 35 Illinois Administrative Code, Subtitle C, Chapter I, Part 302, Subpart B. Chapter 5 of this report, which presents the results of the routine environmental monitoring program, also describes the general effluent limits and water quality standards applicable to the outfalls and discusses compliance with these standards.

#### 2.2.5. NPDES Analytical Quality Assurance

ANL conducts the majority of the analyses required for inclusion in the Discharge Monitoring Report. These analyses are conducted using EPA approved methods in 40 CFR 136. To demonstrate the capabilities of the ANL laboratory for these analyses, the IEPA requires the laboratory to participate in the DMR Quality Assurance program. The IEPA sends a series of control samples to ANL annually and the results of analyses of these samples are submitted to the IEPA and EPA for review. The proficiency of the laboratory is determined by how close the analytical results for the submitted samples come to the actual values. The ANL laboratory has consistently performed very well on these tests.

#### 2.2.6. Spill Prevention Control and Countermeasures Plan

ANL maintains a Spill Prevention Control and Countermeasures (SPCC) plan as required by the Clean Water Act and EPA implementing regulations set forth in 40 CFR 112. This plan describes the actions to be taken in case of oil or oil product releases to waterways in the environment. Persons with specific duties and responsibilities in such situations are identified, as are reporting and recordkeeping requirements mandated by the regulations. Effective use of this plan is ensured by regular training, including both classroom instruction and field exercises. This plan is due to be updated in 1995. There were no reportable spills during 1994.

#### 2.3. Resource Conservation and Recovery Act

The Resource Conservation and Recovery Act (RCRA) and its implementing regulations are intended to insure that hazardous waste are disposed of in an environmentally safe manner and that facilities that treat, store, or dispose of hazardous waste do so in a way that protects human health and the environment. The Hazardous and Solid Waste Amendments of 1984 (HSWA) created a set of restrictions on land disposal of hazardous waste. In addition, HSWA also requires that releases of hazardous waste or hazardous constituents from any solid waste management unit located on the site of a RCRA-permitted facility be cleaned up, regardless of when the waste was placed in the unit or if the unit was originally intended as a waste disposal unit. As discussed below, these RCRA corrective action provisions will have a far-reaching impact on ANL. The RCRA program includes regulations governing management of underground storage tanks containing hazardous materials or petroleum products. The IEPA has been authorized to administer most aspects of the RCRA program in Illinois, except for toxicity characteristic waste and organic solvent emissions.

#### 2.3.1. Hazardous Waste Treatment and Disposal

Because of the nature of the research activities conducted at ANL, small quantities of a large number of waste chemicals are generated. Many of these materials are classified as hazardous waste under RCRA. A number of these wastes also exhibit radioactivity, making them "mixed waste." The hazardous components of mixed waste are subject to RCRA regulations by IEPA, while the radioactive component is subject to DOE regulation under the Atomic Energy Act of 1954. Hazardous waste is collected by the ANL Waste Management Operations (WMO) Department from individual on-site generators and shipped off-site for disposal at an approved hazardous waste disposal facility. Small quantities of reactive hazardous waste are treated on-site. To provide for on-site management of hazardous and mixed waste before off-site shipment or on-site treatment, ANL operates several RCRA-permitted storage and treatment facilities. These facilities, designed and operated in compliance with RCRA requirements, allow for accumulation and storage of waste pending identification of a disposal site. Mixed waste generated on-site does not have any approved disposal mechanism. An aqueous mixed waste treatment system is currently operating under an IEPA Treatability Study. This system uses pH adjustment and sulfide addition followed by precipitation and further volume reduction by concentration. Some mixed waste (scintillation liquids, contaminated lead) is being sent to Hanford for storage and future disposal and some mixed waste will be treated in accordance with a Federal Facilities Compliance Act -Site Treatment Plan.

Twelve Hazardous Waste Management Units (HWMUs) are used for storage of hazardous or mixed waste. All HWMUs, with the exception of the Building 306 - Waste Storage Tank, are used for container storage. Two HWMUs are used for treatment of nonradioactively-contaminated alkali metals and radioactively-contaminated alkali metals. All units are described in Table 2.4.

Hazardous Waste Treatment and Storage Facilities - 1994

TABLE 2.4

Description	Location	Purpose		
Current Interim Status Facilities				
Waste Treatment and Storage	Building 306 - Storage Room A-142	Lab packing and storage of bulk and lab- packed liquid radioactive mixed waste (RMW)		
	Building 306 - Storage Room A-150	Storage of solid and liquid RMW		
	Building 306 - Storage Room C-131	Drum storage and lab packing solid and liquid hazardous waste		
	Building 306 - Storage Room C-157	Drum storage and lab packing of hazardous waste		
	Building 306 - Storage Room D-001	Storage for solid RMW containing toxic metal constituents		
Tank Storage	Building 306	Storage of mixed waste (4000 gal)		
Portable Storage Units	Building 306	Storage of mixed waste (3) Bulking nonradioactive hazardous, flammable, and corrosive liquids (1)		

# TABLE 2.4 (Contd.)

Description	Location	Purpose
Container Storage Area	Building 325C. East	Storage of liquid and solid bulk or lab-packed reactive hazardous waste and solid and liquid bulk PCBs and miscellaneous PCB units
	Building 325C, West	Storage of bulk and lab-packed liquid flamma- ble hazardous waste
Mixed Waste Container Storage	Building 329	Storage of containers of mixed liquid waste
Dry Mixed Waste Storage Area	Building 374A	Storage of solid RMW
Concrete Storage Pad	317 Area	Storage of solid radioactive waste and solid RMW in the form of steel-encased lead shielding containers and containerized solid mixed waste
Alkali Metal Reaction Booth	Building 308	Destruction of water reactive alkali metals
Alkali Metal Reaction Booth	Building 206	Destruction of water reactive alkali metals, possibly contaminated with radionuclides
Interim Status Facilities Closed D	uring 1994	
Water Reaction Tank	317 Area	Destruction of water reactive alkali metals and other reactive chemicals
Shock Sensitive Treatment Area	317 Area	Treatment (detonation) of extremely reactive, or shock-sensitive waste

Two HWMUs will be used for the treatment and storage of mixed waste. The Building 306 Metal Precipitation/Filtration Unit will be used for the treatment of liquid radioactive mixed waste (RMW) containing hazardous metals and/or acids and bases. The metal precipitation tank system, the filtration system, and the mixed waste concentrator will be permitted under the revised Part B application to be submitted to the IEPA in 1995. The Mixed Waste Storage Facility (Building 303) will consist of a building to store mixed waste. The facility has a design capacity large enough to hold approximately five times the current ANL inventory of mixed waste.

The current Part A (interim status) permit lists two units which were formally closed during 1994. These units, also shown in Table 2.4, are the water reaction tank, used in the past for treatment of alkali metals and other water reactive materials, and the shock-sensitive treatment area, used for treatment of highly unstable or explosive materials. Both units are located in the 317 Area. The locations of the major hazardous and non-hazardous waste treatment, storage, and disposal areas at ANL are presented in Figure 2.4.

#### 2.3.2. Permit Status

ANL was granted interim status under RCRA after submitting a notification of Waste Handling Activities and a Part A application in 1980. In 1990, a new Part B permit application (one had previously been sent to the EPA but not acted upon) was prepared for submittal to the IEPA, which had been granted authority to administer most of the RCRA program. The application was submitted to the IEPA and EPA on December 21, 1990. Revisions to the permit application were submitted on June 17, 1991, and September 24, 1991, in response to IEPA and EPA comments.

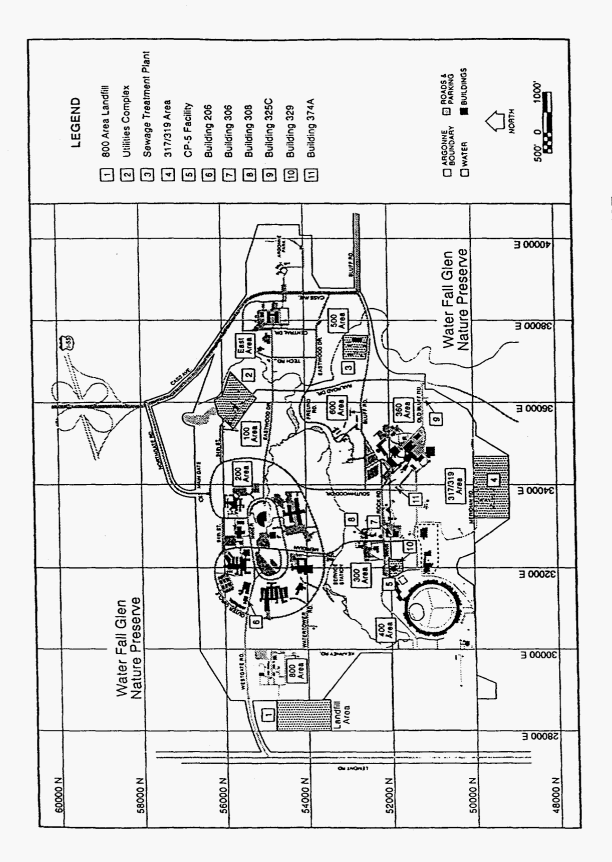


Figure 2.4 Major Treatment, Storage, and/or Disposal Areas at ANL

The RCRA Part B Permit application was revised and updated in 1993. Revision I was submitted to the IEPA during November 1993, which includes information on four new portable hazardous waste storage units and a mixed waste storage tank. ANL responded to EPA notice of deficiency comments regarding the alkali metal passivation booths in Buildings 308 and 206 and incorporated the response into the revised application. Revision II of the Part B application is being prepared to include a new hazardous waste storage facility, a new mixed waste storage facility, and a transuranic mixed waste storage facility. Revision II will be submitted in 1995. IEPA will conduct a technical review of the Part B application and may issue ANL a RCRA draft permit by the end of 1995.

A RCRA Facility Assessment (RFA) was completed by the IEPA during summer 1991. The RFA report from IEPA was received during late 1993 and identified 740 units [735 solid waste management units (SWMUs) and five Areas of Concern (AOC)]. The report identified 432 units (427 SWMUs and five AOCs) for further work. ANL prepared a report entitled "Proposed Revisions to the RCRA Facility Assessment Report for Argonne National Laboratory-East." This report included recommendations to reduce the number of units that IEPA identified in the RFA Report for further work from 432 units to 71 units (69 SWMUs and two AOCs). When the Part B permit is issued, it will most likely contain requirements to characterize and assess the SWMUs. ANL is working proactively to characterize and investigate these SWMUs with emphasis in the 800 and 317 Areas. ANL continues to abide by its Part A permit and the interim status standards found in 40 CFR 265 and 35 IAC Part 725.

#### 2.3.3. Hazardous Waste Generation

ANL typically generates a wide variety of hazardous waste and mixed waste each year. The quantity of mixed waste generated during 1994 was 42,016 liters (11,099)

gallons). Of this amount, 84% of the mixed waste was sent to the DOE Hanford site for storage. In 1994, 47,879 kg (105,556 lbs) and 21,838 liters (5,769 gals) of hazardous waste were shipped to a disposal site by an IEPA-permitted hazardous waste disposal company. About 3.6 kg (8 lbs) of reactive hazardous waste (sodium potassium alloy) were treated on the site in the Building 308 alkali metal reaction booth. This unit renders the waste nonhazardous. During 1994, corrosive liquid mixed waste was treated in the Building 306 metal precipitation/filtration unit's treatability study.

#### 2.3.4. Facility Modifications

New radioactive and hazardous waste storage facilities are being planned. Various facilities are scheduled for completion between 1995 and 1998. The mixed waste storage building is currently entering the final design phase. The hazardous waste, radioactive waste, and Building 306 rehabilitation completed preliminary design during 1993. The Part B permit application will be revised to incorporate these facilities when the final design details are known.

In 1994, four prefabricated portable hazardous waste storage buildings became operational. These units are used to store containers of flammable and corrosive waste and other hazardous, radiological, or mixed wastes. The units, each 5 m (16 ft) long by 3 m (10 ft) wide by 3 m (9 ft) high, store the above hazardous or mixed waste until ANL-E ships the waste to an approved treatment, storage, and disposal facility. The waste stored in the waste storage units includes flammable (D001) and corrosive (D002) hazardous waste and radioactive mixed waste. Bulk liquid and lab-packed waste will be stored in containers of various sizes (e.g., 55-gallon and 30-gallon drums, 5-gallon containers, and other container types) in these storage units. Based on the maximum retention sump capacity and configuration of containers, a maximum of 4600 L (1,200 gal) of waste can be stored in each waste storage unit. Bulking operations are conducted

to consolidate and reduce the volume of lab-packed waste in containers in a designated storage unit.

#### 2.3.5. Mixed Waste Handling

The hazardous component of mixed waste is governed by RCRA regulations, while the radioactive component is subject to regulation under the Atomic Energy Act as implemented by DOE Orders. Accordingly, facilities storing or disposing of mixed waste must comply with both DOE requirements and RCRA permitting and facility standards. ANL generates several types of mixed waste, including acids, solvents, and sludges contaminated with radionuclides. Corrosive mixed waste is undergoing a treatability study to remove the hazardous characteristics. Some mixed waste that meets a Hanford Waste Acceptance Criteria is sent to Hanford for storage and eventual disposal. The Part B permit application addresses mixed waste management procedures. A number of mixed waste treatment processes will be operational during 1995 and 1996. All of these processes will be incorporated into a Part B Permit revision to the IEPA late in 1995.

## 2.3.6. Federal Facility Compliance Agreement (FFCA) Activities

The Draft Site Treatment Plan (DSTP) was submitted to the DOE for review in August 1994. The DSTP outlined the mixed waste on-site and identified several potential treatment options for each waste stream. The Proposed Site Treatment Plan (PSTP) was submitted to the DOE which in turn submitted it to the state in March 1995. The PSTP identifies six on-site treatment systems and one off-site treatment system that ANL proposes to use for its existing inventory of mixed waste. Once the PSTP is approved by the state, treatment schedules for the various mixed waste streams in the ANL

inventory will be developed and the hazardous constituent treated in compliance with RCRA regulations.

#### 2.3.7. RCRA Inspections

Two RCRA inspections were conducted by IEPA during 1994. A RCRA compliance inspection was conducted on April 28, 1994. No significant issues were identified. A RCRA closure inspection of the 317 Area Shoot and Burn Unit was conducted on June 24, 1994. No significant issues were identified. IEPA certified closure for the above unit on September 29, 1994.

#### 2.3.8. Underground Storage Tanks

In response to underground storage tank regulations, ANL prepared a Site-Wide Underground Tank Compliance Plan. The ANL site currently contains 25 existing, upgraded, replaced or new underground storage tanks; 33 tanks have been removed over the last several years. The majority of these tanks are being used, or were used in the past, for storage of fuel oil for emergency generators or space heaters. The on-site vehicle maintenance facilities use underground gasoline and methanol tanks. The Compliance Plan sets out a program for the replacement or upgrading of tanks that must remain in use. Currently, all tanks remaining in use are being monitored under a new recordkeeping program initiated by ANL.

During 1993, nine regulated underground tanks were upgraded to current technical requirements (secondary containment, corrosion protection, leak detection, double-walled piping, spill and overfill protection) and 10 in-use underground storage tanks were replaced with new double-wall fiberglass tanks and required monitoring equipment. In 1994, three new tanks to store vehicle fuel were installed at the new Grounds and

Transportation Center. The tanks they replaced, located in the 800 Area, will be removed and closed in 1995.

#### 2.3.9. Corrective Action for Solid Waste Management Units

As mentioned previously, the HSWA amendments to RCRA require that any Part B permit issued must include provisions for corrective actions for all releases of hazardous materials from any solid waste management unit (SWMU) at the site, regardless of when the waste was placed in the unit. When issued, the Part B permit will contain a compliance schedule which will govern the characterization and any required remediation of such units. The Part B permit application submitted to the IEPA identified and provided information on 56 SWMUs, both active and inactive. The recently issued RFA report from the IEPA identified more than 700 SWMUs (see Section 2.3.2.). The great majority of these sites are believed to contain little or no residual contamination; however, a number may be required to undergo some type of corrective action. The process of conducting detailed characterization studies to determine if hazardous materials have been released from a number of these units was begun in 1989. A summary of the preliminary results of these investigations can be found in Chapter 6. More extensive characterization is currently underway at a number of the SWMUs in accordance with the IEPA-approved corrective action work plans for the 800 Area and 317/319 Areas. Interim removal actions for two SWMUs were also being implemented.

# 2.4. Solid Waste Disposal

During September 1992, ANL closed the operation of its sanitary landfill. This facility began operation in 1969. The original operating permit was issued by the IEPA in 1981. A supplemental permit addressing final elevations, a groundwater monitoring

program, and closure/post closure costs was issued by the IEPA on April 24, 1992, and revised on September 15, 1992, and October 22, 1992. The IEPA conducted a closure inspection of this landfill on April 28, 1994, and no significant issues were identified.

The April 24, 1992, supplemental landfill permit required ANL to implement a specific groundwater monitoring program at the sanitary landfill. The program is designed to identify any releases from the landfill and demonstrate compliance with the applicable groundwater quality standards. Quarterly monitoring of 11 locations began during July 1992. Exceedances of the groundwater quality standards for chloride, iron, total dissolved solids, and manganese were noted at monitoring locations where these levels have been historically reported. One exceedance of the groundwater quality standard for phenols was noted at one location only during the first quarter of 1993. On January 11, 1995, IEPA issued a supplemental permit which, among other things, revised the monitoring program to add 13 new well locations, delete the previous 11 well locations, and increase the number and types of contaminants monitored.

The IEPA began requiring annual non-hazardous special waste reporting in 1991. The report is submitted by February 1 of each year and describes the activity of the previous year. It is a summation of all manifested nonhazardous and PCB wastes. Nonhazardous special waste generated and disposed during 1994 are described in Table 2.5.

# 2.5. National Environmental Policy Act

The National Environmental Policy Act of 1969 (NEPA) established a national environmental policy that promotes consideration of environmental factors in federal or federally-sponsored projects. NEPA requires the review of the environmental impacts of

**TABLE 2.5** 

Generation of Nonhazardous Special Waste, 1994

Waste	Quantity		
PCB contaminated oil	5640 L (1490 gals)		
PCB (miscellaneous contaminated oils, ballasts)	5734 L (1515 gals)		
PCB (transformers and transformer oil)	52,605 kg (115,974 lbs)		
Oil	22,599 L (5970 gals)		
Mineral spirits	2328 L (615 gals)		
Sanitary sludge	84 m <sup>3</sup> (110 yds <sup>3</sup> )		
Sandblast media (water tower project)	126 m <sup>3</sup> (165 yds <sup>3</sup> )		
Soil contaminated with paint	255 m <sup>3</sup> (333 yds <sup>3</sup> )		
Soil contaminated with diesel fuel	23 m <sup>3</sup> (30 yds <sup>3</sup> )		
Bag House waste	1799 m³ (2353 yds³)		
Medical waste	249 kg (550 lbs)		
Miscellaneous nonhazardous solid chemicals	11,205 gals (56,532 lbs)		
Miscellaneous nonhazardous liquid chemicals	6505 gals (34,513 lbs)		

a project. To ensure compliance with this policy, NEPA requires that projects with potentially significant impacts be reviewed carefully through the generation of either an Environmental Assessment (EA) or Environmental Impact Statement (EIS). This review process is designed to insure that all potential impacts are identified, all available options are considered, and all affected parties are informed and given opportunity to comment on a project.

The DOE implementation of NEPA has undergone significant change during recent years. The threshold at which projects are subject to NEPA review has been reduced to such an extent that virtually all activities are now required to undergo some sort of NEPA review and documentation. On the other hand, the list of Categorical Exclusions, which is a list of project types that normally do not require an EA or EIS, has been expanded to help streamline the process. The DOE final rule on NEPA implementing procedures and guidelines revocation was published on April 24, 1992.

The ANL NEPA compliance program is designed to ensure that all activities under consideration are reviewed to determine any significant environmental impacts. This program subjects each proposed project to a careful consideration of potential impacts to air (dust, gaseous emissions), water (liquid effluents, wetland impacts), and soil (solid waste generation, construction activity), as well as impacts involving critical wildlife habitats, historic and cultural resources, radiation, noise, workers and other considerations. A questionnaire is completed for the project and is used as documentation of the review of potential impacts. This form (DOE/CH Form 560) is submitted to DOE for review and determination of the proper level of NEPA documentation. Projects that exhibit potentially adverse impacts in any area are subject to further review, including, if necessary, preparation of an EA or EIS. Any EA or EIS prepared by ANL is reviewed by DOE according to the procedures specified in DOE Order 5440.1E and DOE/CH Order 5440.1C.

During 1994, approximately 200 projects underwent NEPA review. There was one EA determination for projects reviewed in 1994. Preparation of that EA began in 1994 as well as preparation of a second EA from a determination made prior to 1994. No EIS determinations were made in 1994. There are no known major actions planned for 1995.

#### 2.6. Safe Drinking Water Act

The Safe Drinking Water Act of 1974 (SDWA) established a program to ensure that public drinking water supplies are free of potentially harmful materials. This mandate is carried out through the institution of national drinking water quality standards, such as Maximum Contaminant Levels (MCL) and Maximum Contaminant Level Goals (MCLG), as well as through imposition of wellhead protection requirements, monitoring requirements, treatment standards, and regulation of underground injection activities. The SDWA established Primary and Secondary National Drinking Water Regulations, which set forth requirements to protect human health (primary standards) and provide aesthetically acceptable water (secondary standards).

#### 2.6.1. Applicability to ANL

The drinking water supply at ANL consists of four on-site wells that supply raw water to the water treatment plant. The treatment plant removes iron, softens the water by ion-exchange, and adds chlorine before pumping it to the site-wide distribution system. Because of the nature of the ANL drinking water system and the persons served by it, the system is classified as a non-transient, non-community water supply, and as such is subject to the regulations applicable to such systems. The Laboratory is subject to regulations under the State of Illinois program administered by the Illinois Department of

Public Health (IDPH). IDPH adopted the Illinois Pollution Control Board regulations at 35 IAC Parts 605, 607, and 611. These regulations are incorporated by reference into 77 IAC Part 900. These regulatory programs establish a monitoring program, design, operation and maintenance requirements, and secondary water quality standards.

#### 2.6.2. Monitoring Requirements

The primary drinking water standards establish certain monitoring and analytical requirements. Both federal and state regulations apply to the ANL drinking water monitoring program. ANL routinely samples each of the four wells and the treated water quarterly for compliance with applicable regulations. Treated water is also sampled annually for radiological analyses. Chapter 6 of this report presents a detailed discussion of the results of the drinking water program. During 1994, samples continued to be collected and all state and federally-required analyses were conducted. EPA-approved procedures were employed by a certified laboratory. The treated water sample results were not required to be sent to IDPH during 1994 because of monitoring waivers. As a result of meeting current federal and state drinking water standards, waivers allowing reduction in future sampling were granted to ANL by the IDPH during March and August of 1993.

Since 1993, results indicated that the "action levels" for lead and copper were not exceeded. A reduced frequency in sampling (annually) and number of locations (20) was authorized for 1994. The action levels for lead and copper were exceeded during 1994 and ANL is required to return to semi-annual sampling of 40 locations during 1995. When action levels are exceeded, a user education program must be established within 60 days of the end of the monitoring period. Notice was provided to system users and posted on all ANL bulletin boards during February 1995.

#### 2.7. Federal Insecticide, Fungicide and Rodenticide Act

The Federal Insecticide, Fungicide and Rodenticide Act (FIFRA) establishes a program to register pesticides, regulate their transportation and disposal, and determine standards for their use. Within ANL, all applications of pesticides are by licensed contractors who provide any pesticides used and remove any unused portions. Herbicides are rarely used, but when they are needed, a licensed contractor is brought in to apply them. In these situations, ANL ensures that the herbicide is EPA-approved, that it is used properly and any residue is disposed of in accordance with applicable regulations. This is carried out by oversight inspections and maintenance of records.

# 2.8. Comprehensive Environmental Response, Compensation and Liability Act

The Comprehensive Environmental Response, Compensation and Liability Act (CERCLA) addresses the cleanup of hazardous waste disposal sites and the response to hazardous substance spills. Under CERCLA, the EPA collects data regarding sites subject to CERCLA action through generation of a Preliminary Assessment (PA) report, followed up by a Site Investigation (SI). Based on the data collected, the sites are ranked according to their potential to cause human health impacts or environmental damage. The sites with the highest ranking are placed on the National Priority List (NPL) and are subject to mandatory cleanup actions, funded either by Potentially Responsible Parties (PRPs) or by the allocation of Superfund money to the project. Federal agencies are responsible for their own cleanup costs.

#### 2.8.1. CERCLA Program at ANL

In the past, federal facilities were allowed to develop and manage their own independent CERCLA program subject to EPA oversight. The DOE's CERCLA program was detailed in DOE Order 5480.14. This DOE Order has since been superseded by DOE Order 5400.4. Under the provisions of DOE Order 5480.14, in July 1986, ANL submitted preliminary assessment (PA) reports to DOE for the seven inactive units on the current ANL site and one inactive unit located on land deeded to the DuPage County Forest Preserve District in 1973 as shown in Table 2.6. Because of changes in the EPA CERCLA program brought about by the Superfund Amendments and Reauthorization Act of 1986 (SARA), the EPA is now required to publish a comprehensive inventory of federal facility sites known as the Federal Agency Hazardous Waste Compliance Docket. These sites are ranked, using the Hazardous Ranking System (HRS), and placed on the NPL list if they score high enough. However, since they are federal facilities, Superfund money is not available to support cleanup operations. In support of this effort, the EPA required submittal of PA reports for sites at ANL (as listed in Table 2.6). These reports were submitted in April 1988. Four sites not included in the original DOE submittal were included in the subsequent submission. In late 1990, ANL prepared and submitted one additional PA for a solvent disposal site used for a number of years by the ANL 810 Area paint shop for disposal of waste paint solvents. The site in Waterfall Glen Forest Preserve is currently owned by the DuPage County Forest Preserve Preserve District and thus is no longer part of a federal facility subject to SARA. The PA for this site was submitted in an effort to inform the EPA of past ANL activities.

During early 1990, the EPA requested that ANL submit Site Screening Investigation (SSI) reports for six of the 13 sites. Upon further discussions between the EPA and DOE, one of the six sites was eliminated from consideration and three units

TABLE 2.6

List of Inactive Waste Disposal Sites at ANL
Described in Various CERCLA Reports

Site Name	DOE/CERCLA	EPA/SARA	EPA/SSI
Waste Sites on Current ANL Property			
800 Area Landfill and French Drain	X	х	X
319 Area Landfill and French Drain	X	X	X (1)
Landfill East-Northeast of the 319 Area	x	x	X (1)
Compressed Gas Cylinder Disposal Area, 318 Area	X	X	X (1)
French Drain, 317 Area	x	X	X (1)
Mixed Waste Storage Vaults, 317 Area		X	X (1)
Shock Treatment Facility, 317 Area	x	X	X (1)
Wastewater Holding Basin, Sewage Treatment Plant		X	
Liquid Waste Treatment Facility, Building 34	X	X	
Decommissioned Reactor CP-5, Building 330		x	X
Gasoline Spill, Gasoline Station		X	
810 Area Paint Shop		X	
Waste Sites on Old ANL Property, Currently Waterfall Glen Forest Preserve			
Reactive Waste Disposal, Underwriters Pond	X	X	

<sup>(1)</sup> All units located in the 317/319/ENE Area were described in a single Site Screening Investigation (SSI) report.

(317/319/ENE) were treated as a single site due to their physical proximity. As a result, three SSI reports were completed by ANL and submitted to DOE in December 1990. They were subsequently transmitted to EPA in January 1991. Table 2.6 lists those sites for which an SSI was submitted.

Recent inquiries into waste disposal practices during the 1950s and 1960s have identified a number of smaller waste disposal sites, some of which could contain hazard-ous materials. These sites are under investigation; however, their potential to impact groundwater is thought to be minimal.

#### 2.8.2. CERCLA Remedial Actions

Remedial actions to clean up any release of hazardous materials from these sites could occur in a number of different ways. Since all but one of the CERCLA sites are on the ANL site and are included as SWMUs in the RCRA Part B permit application, they may be subject to RCRA corrective action and come under the authority of the IEPA. However, since several of the sites contain radiological contamination, over which RCRA has no authority, the sites may be subject to a combined RCRA/CERCLA action.

Regardless of which regulatory vehicle is ultimately used to facilitate the cleanup of these sites, the DOE, through various initiatives put forth by the Secretary of Energy, has made the commitment to clean up all such sites voluntarily within the next 30 years, wherever possible returning them to unrestricted use. As a response to these commitments, ANL has requested funding for the characterization and remediation of all but two of these sites. The two remaining sites are the one off-site unit, which is no longer under the control of ANL or DOE, and a small gasoline spill which was completely

cleaned up immediately after the spill occurred. Several of the characterization projects have already begun and will continue over the next few years.

# 2.8.3. Emergency Planning and Community Right to Know Act (EPCRA), Superfund Amendments and Reauthorization Act (SARA) Title III

#### Sections 311 and 312

Title III of the 1986 SARA amendments to CERCLA created EPCRA as a freestanding provision for response to emergency situations involving hazardous materials and for making known to federal, state, and local emergency planning authorities information regarding the presence and storage of hazardous substances and their planned and unplanned environmental releases. Under EPCRA, ANL is required to provide to applicable emergency response agencies an inventory of hazardous substances stored on the site, Material Safety Data Sheets (MSDS), and completed SARA data sheets (Tier I or II forms) for each hazardous substance stored in quantities above a certain threshold planning quantity (typically 10,000 lbs; but as low as one pound for certain compounds). However, chemicals used in research laboratories under the direct supervision of a technically qualified individual, are exempt from reporting. In November 1987, an inventory and MSDS forms for nine chemicals were submitted to the Local Emergency Planning Committee (LEPC); in March 1988, Tier I reports providing additional information on these chemicals were submitted. Updated Tier II forms were submitted to the LEPC by the required March 1 deadline for the years 1989 through 1994. The Tier II forms updated the previous listings and provided more information regarding the amount of material stored and the location of the material. Table 2.7 lists hazardous compounds reported under SARA Title III for 1994.

# 2. COMPLIANCE SUMMARY

**TABLE 2.7** Compounds Reported Under SARA Title III - 1994

	Hazard Class				
Compound	Sudden Release Fire of Pressure Reactive		Acute Health Hazard	Chronic Health Hazard	
Diesel Fuel/ Heating Oil	X				
Gasoline	X				
Methanol/ Gasoline	X				
Chlorine		X		X	
Chlorofluoro- carbon 11		X			
Sulfuric Acid				X	
Calcium Oxide				X	
Oils containing PCBs					X
Lubricating Oils	X				
NALCO 356 Amine Corrosion Inhibitor	X			X	

Section 304 of SARA Title III requires that the LEPC and state emergency planning agencies be notified of accidental or unplanned releases of certain hazardous substances to the environment. The procedures for notification are described in the Argonne Comprehensive Emergency Management Plan. There were no incidents during 1994 that required notification of the LEPC and Illinois Emergency Management Agency.

#### Section 313

Section 313 of the EPCRA was enacted as a free-standing provision of the SARA in 1986. It requires facilities to prepare an annual report titled "Toxic Chemical Release Inventory, Form R" if annual usage quantities of listed toxic chemicals exceed certain thresholds. In 1994, threshold chemical usage quantities were 10,000 lbs for listed chemicals either manufactured or processed.

Seven Form R reports for 1993 were transmitted to DOE on June 28, 1994, for chemicals where use exceeded the 10,000 lb/yr reporting threshold. Table 2.8 lists chemicals reported under SARA Title III, Section 313 for 1993. SARA 313 reports are due to EPA each July 1. Seven Toxic Release Inventory reports will be filed for 1994. The principal chemical uses for ANL are sulfuric acid at the boiler house, methanol for alternate fueled vehicles, and additives to gasoline. The vehicle fuels are all used on-site and consumption cannot be delineated between mobile sources and other uses.

On July 15, 1994, the Laboratory Activity Toxic Chemical Usage Report for ANL was transmitted to DOE showing usage of acetone of 1000-3000 lbs for 1993. This report is due to DOE on August 1 to evaluate the significance of emissions from SARA 313 chemicals used in laboratory activities. (This use is currently exempt from reporting.) ANL releases originate principally from research and development activities.

TABLE 2.8

SARA Title III, Section 313, Chemicals, 1994

		Pounds Emitted				
Chemicals	Air	Water	Off-site Transfer	Other*		
Sulfuric Acid		A**	4000	7		
Methanol	В		1400¹	12		
Benzene	В		$\mathbf{B}^2$	23		
Toluene	В		$\mathbb{C}^3$	144		
Xylene (mixed isomers)	В		$\mathbf{B}^4$	144		
Methyl tert-Butyl ether	В			86		
1,2,4-Trimethylbenzene	В			29		

<sup>\*</sup>Quantity released pursuant to EPCRA Section 329(8); reported in Section 8: "Source Reduction and Recycling Activities."

B 11-449 lbs

C 500-999 lbs

<sup>\*\*</sup>Range Codes A 1-10 lb

<sup>&</sup>lt;sup>1</sup>Reported for use in off-site energy recovery.

<sup>&</sup>lt;sup>2</sup>450 lbs listed for off-site energy recovery.

<sup>&</sup>lt;sup>3</sup>620 lbs listed for off-site energy recovery.

<sup>&</sup>lt;sup>4</sup>100 lbs listed for off-site energy recover.

On November 30, 1994, EPA included an additional 286 chemicals and chemical categories to the Section 313 toxic chemical list. Reporting on these additional chemicals will be required beginning in 1995 calendar year (due July 1, 1996). ANL is currently reviewing material use to identify and implement source reduction techniques.

#### 2.9. Toxic Substances Control Act

The Toxic Substances Control Act of 1976 (TSCA) provides for testing of manufactured substances to determine toxic or otherwise harmful characteristics and regulation of the manufacture, distribution, use, and disposal of regulated substances. The only TSCA-regulated compounds in significant quantities at ANL are polychlorinated biphenyls (PCBs) contained in electrical capacitors and transformer oil and PCB-contaminated sludge. Regulations governing PCB management, such as use and disposal and remediation of spills, are set forth in 40 CFR 761. These regulations provide detailed requirements for use and disposal of materials containing concentrations of PCBs above 50 ppm. Most of these regulations relate to PCBs contained in dielectric fluids within electrical equipment, such as transformers and capacitors.

#### 2.9.1. PCBs in Use at ANL

The majority of PCBs at ANL were contained in a number of transformers, capacitors, and switches throughout the site. Starting in 1987, ANL began removing and disposing of all PCB and PCB-contaminated electrical equipment. All indoor and outdoor transformers, with the exception of one unit in Building 211, have been removed and transported off the site for proper disposal. During 1990, all pole mounted transformers and circuit breakers containing PCBs were replaced or retrofilled with non-PCB oil. All removal and disposal activities were conducted by licensed contractors specializing in

such activities. Accordingly, the equipment has been labelled to reflect their non-PCB status. Operation, removal, storage, and disposal of PCB-containing articles were conducted in compliance with applicable TSCA regulations. The PCB Annual Report for Calendar Year 1993 was prepared during June 1994. A computerized database for tracking PCB-containing articles has been developed.

Large capacitors (466) with PCBs are still present on-site. All are either in use or standby units. There are no plans to dispose of any of these units. The total weight of PCBs in these capacitors is estimated to be 2920 kg (6437 lbs).

# 2.9.2. Disposal of PCBs

Disposal of PCBs from the ANL site includes material lab-packed and bulked through Waste Management Operations (WMO), bulk solids sent off-site through WMO, and bulk solids sent off-site directly from the transformer removal/retrofill projects.

#### 2.9.3. PCB Spills

In 1989, ANL characterized sludge from the Laboratory Wastewater Treatment Plant (LWTP) for organic constituents. The characterization was for acceptance of the sludge at the Hanford site. The analysis revealed that the sludge contained PCBs in concentrations greater than 50 ppm; therefore, defining it as a TSCA regulated waste. In 1992, sludge contained in the LWTP holding tanks, and in the LWTP drying beds was characterized for TSCA, RCRA, and radioactive constituents. Results of that study showed the sludge to be a TSCA waste (total PCBs ≥ 50 ppm), nonhazardous under RCRA and a Class A low-level waste. In 1993, a process to evaluate and recommend a viable treatment alternative to separate the PCBs from the sludge began. The process first evaluated several methods to treat the waste by assessing their relative cost and

implementability. ANL also is assessing the effectiveness of potential technologies using treatability studies. These treatability studies will be completed in FY 1995 and a comparative analysis document recommending the most viable treatment alternative submitted to DOE for approval in FY 1995.

The PCB-contaminated sludge was removed, containerized, and stored on-site during 1994. Removal of the drying beds and closure of the area will also be recommended to DOE. The area has been identified as a solid waste management unit (SWMU) subject the corrective action provisions of RCRA. The schedule for closure of the drying beds will be decided as part of the RCRA corrective action process.

A secondary issue, finding the source of the PCBs contaminating the LWTP, has been a subject of concern. To address it, ANL initiated a project in FY 1994 to characterize the contents of several retention tanks across the Laboratory. The retention tanks serve as collection vessels for buildings at ANL which handle radioactive materials. The purpose of the tanks is to ensure that laboratory wastewater containing unacceptable quantities of radioactive materials does not have a direct link to the LWTP sewer. When full, ANL analyzes the tanks' contents for radioactivity, and if results are below the release limits, the water is released to the LWTP. Sludge from a set of tanks was sampled and analyzed in FY 1994 and showed that one tank, in Building 203, contained PCB concentrations near 4100 ppm. ANL intends to sample a second set of tanks in FY 1995. Cleanout of the sludge in the tanks in Building 203 is scheduled for FY 1995. The tank cleaning will be performed according to regulatory requirements. The cleaned tank will no longer be classified as a "PCB container." Cleanout of any additional tanks containing PCBs in concentrations ≥ 50 ppm during sampling of the second set, is scheduled for FY 1995. The purpose of the sampling and cleanout is to isolate the sources of PCB contamination across the ANL site and cleanout the system from the "headwaters" to the treatment plant.

EPA conducted a TSCA (PCB) Compliance Inspection on July 20, 1994. Several minor issues were identified: the PCB annual report for 1993 was inaccurate, several manifests for off-site shipments of PCBs in 1993 were not readily available and PCB Quarterly Inspection Reports were not readily available. These issues were addressed by ANL in a follow-up letter to EPA dated August 11, 1994.

# 2.10. Endangered Species Act

The Endangered Species Act of 1973 (ESA) is designed to protect plant and animal resources from the adverse effects of development. Under the Act, the Secretaries of the Interior and Commerce are directed to establish programs to insure the conservation of endangered or threatened species or critical habitat of such species. For ANL, the Fish and Wildlife Service (FWS) has been delegated authority to conduct these consultations and enforce the ESA.

To comply with the ESA, federal agencies are required to make an assessment of the proposed project area to determine if any threatened or endangered species or critical habitat of these species exist. If no such species or habitat are present this fact is to be documented in a letter to the FWS. If such species or habitat are found to exist, the FWS is to be notified and a series of consultations and studies are then carried out to determine the extent of impact and any special actions which must be taken to minimize this impact.

At ANL, the provisions of the ESA are implemented through the NEPA project review process. All proposed projects must provide a statement describing the potential impact to threatened or endangered species and critical habitat. This statement is included in the general Project Environmental Evaluation Form. If there is potential adverse

impact, this impact will be further assessed and evaluated through the preparation of a more detailed NEPA document, such as an EA or EIS.

Neither federal nor state-listed threatened or endangered species are known to reside on the ANL site. The federally-listed endangered Indiana Bat, Myotis sodalis, and the federally-threatened Hine's Emerald Dragonfly, Somatochlora hineana, reside in the area and may possibly reside on the ANL site. Two state-endangered species, (River Otter, Lutra canadensis; and White Lady's Slipper, Cypripedium candidum; and two state-threatened species Kirtland's Snake, Clonophis kirtlandi; and sedge, Carex crawei, reside in the area and may possibly reside on the ANL site. Impacts to these species are also assessed during the NEPA process. No project at ANL has ever had to be stopped, delayed or modified as a result of potential impact to endangered species.

# 2.11. National Historic Preservation Act

The National Historic Preservation Act (NHPA) requires federal agencies to assess the impact of proposed projects on historic or culturally important sites, structures or objects within the site of the proposed projects. It further requires federal agencies to assess all sites, buildings, and objects on the site to determine if any qualify for inclusion in the National Registry of Historic Places. The Act also establishes a procedure for archaeological investigation activities and a system of civil and criminal penalties for unlawfully damaging or removing such artifacts.

The NHPA is implemented at ANL through the NEPA review process, as well as through the internal digging permit process. All proposed actions must consider the potential impact to historic or culturally important artifacts and document this consideration in the Project Environmental Evaluation Form. If the proposed site has not been

surveyed for the presence of such artifacts, a cultural resources survey is conducted and any artifacts found are carefully documented and removed. Prior to disturbing the soil, an ANL digging permit must be obtained from the PFS division. This permit must be signed by the Cultural Resources Officer at ANL prior to digging to document the fact that no significant cultural resources will be impacted.

During 1993, fieldwork was completed on the archaeological survey of the ANL site. Phase I archaeological surveys were conducted to identify the location of potential historic and prehistoric sites at ANL. The results of the Phase I surveys were documented in reports which were subsequently submitted to the Illinois Historic Preservation Agency (IHPA) for their review. Upon completion of the review, the IHPA makes a determination whether or not the sites are eligible for nomination to the National Registry of Historic Places. If the sites are deemed not eligible, the area is considered "cleared" and no further archaeological review is required. If however, the IHPA believes that a site is significant and potentially eligible for nomination to the National Registry, a Phase II survey is recommended. The Phase II survey intensively characterizes the site and the reported findings are again reviewed by the IHPA to determine the site's eligibility to the National Register.

During 1994, two Phase II archaeological surveys were conducted. From August 30 through September 1, 1994, a Phase II survey was conducted on the North Meadow Site. IHPA determined that the site was not significant and no further action required. From August 10 through August 26, 1994, a Phase II survey was conducted on five prehistoric sites. The Phase II report submitted to IHPA states that these sites appear to have sufficient integrity and cultural materials to warrant possible National Register status. As of February 15, 1995, ANL has not received a determination from IHPA.

The text of the Cultural Resource Management (CRM) plan and the final site map have been completed. A final CRM plan will be issued after the documents for the appendices have been prepared. The CRM plan and final site map will be distributed to appropriate ANL and DOE personnel in 1995.

ANL currently does not contain any sites, buildings or structures included in the National Register of Historic Places. It does, however, contain several facilities which represent historically important scientific or technical achievements, such as the first experimental boiling water reactor. If it is determined that such sites are suitable for listing, they will be investigated and submitted to the Department of the Interior for possible listing.

# 2.12. Floodplain Management

Federal policy on managing flood plains is contained in Executive Order 11988 (May 24, 1977). This Executive Order requires federal facilities to avoid to the extent possible adverse impacts associated with the occupancy and modifications of floodplains. A project proposed for construction in a floodplain must demonstrate that there is no reasonable alternative to the floodplain location.

The ANL site is located approximately 150 feet above the nearest large body of water (Des Plaines River) and thus is not subject to major flooding. A number of small areas, associated with Sawmill Creek and other small streams or low-lying areas, are subject to local flood conditions following extremely heavy precipitation. To insure that these areas are not adversely impacted, ANL has maintained a practice of not permitting new facility construction within these areas, unless there is no practical alternative. Any

impact to floodplains are fully assessed in a floodplain assessment, and, as appropriate, documented in the NEPA documents prepared for a proposed project.

# 2.13. Protection of Wetlands

Federal policy on wetland protection is contained in Executive Order 11990. In addition, 10 CFR Part 1022 describes DOE's implementation of this Executive Order. This Order requires federal agencies to identify potential impacts to wetlands resulting from proposed activities and to minimize these impacts. Where impacts cannot be avoided, action must be taken to mitigate the damage by repairing the damage or replacing the wetlands with an equal or greater amount of a man-made wetland as much like the original wetland as possible. The current DOE policy is for no net decrease in the amount of wetland as a result of DOE activities.

Due to the topography and nature of the soil at ANL, the site contains a significant number of natural and man-made wetlands. These range from small stormwater ditches which are overgrown with cattails to natural depressions, beaver ponds and man-made ponds. The potential impact to these areas caused by a proposed action is described in the NEPA Project Environmental Evaluation Form for the project. The impacts are assessed in a wetlands assessment, as appropriate. If the involvement of wetland negates use of a categorical exclusion, the DOE will require preparation of an EA or EIS. The APS project, currently under construction, required a U. S. Army Corps of Engineers Section 404 permit and extensive wetland mitigation activities, since several small natural wetlands occupied the construction site and had to be replaced elsewhere. These actions were documented in the EA which was approved in early 1990.

During 1993, a site-wide wetlands delineation was completed of the ANL site. A survey was conducted to identify and delineate all jurisdictional wetlands present on-site in accordance with the 1987 U. S. Army Corps of Engineers Wetlands Delineation Manual. The results of the survey were delineated on a site map indicating the areal extent of all wetlands present at ANL down to 500 m<sup>2</sup> (1/8th acre). The findings are documented in an accompanying report which describes in detail the soil, vegetation, and hydrology of each wetlands area delineated on the map. Thirty-five (35) individual wetland areas were identified totalling approximately 45 acres. The wetlands areas were also digitized onto a computer-aided design file in order that scale maps are available to ANL engineers for planning and designing projects. The delineation will also be useful for determining project impacts under NEPA review.

Activities in 1994 were limited to the dissemination of the sitewide wetlands map to ANL and DOE personnel requiring wetlands information. Delineations were also performed for several specific projects to be conducted in close proximity to wetlands identified on the sitewide map. The purpose of these delineations was to specifically identify wetlands boundaries in order to assess project impacts.

In 1995, wetlands maps will continue to be disseminated as necessary and specific delineations will continue to be performed for projects occurring near wetlands identified on the sitewide map.

#### 2.14. Current Issues and Actions

The purpose of this section is to summarize the most important issues related to environmental protection encountered during 1994. Since preceding sections of this chapter contain detailed discussions of specific issues related to each major piece of

environmental regulation, discussions of specific issues will not be repeated in this section. Please refer to the appropriate section of this chapter for these details.

## 2.14.1. Major Compliance Issues

The most significant ongoing issue encountered at ANL during 1994 involves wastewater discharges: compliance with existing NPDES wastewater discharge permit requirements. ANL was issued a new NPDES permit, effective October 30, 1994. The new permit included a number of changes to sampling parameters and frequency for various outfalls. Since the permit became effective, ANL has not been able to consistently meet the permit limits for TDS, copper, and total residual chlorine. It is also anticipated that ANL will have difficulty consistently meeting the new permit limits for ammonia nitrogen. A significant compliance issue for 1995 is identifying and implementing actions needed to ensure compliance with ANL's NPDES permit. Plans to upgrade or construct the necessary wastewater treatment facilities are underway.

Another significant issue at ANL during 1994 involves the inadvertent shipment, through an independent contractor, of waste oil contaminated with PCBs to an off-site recycling facility. EPA and IEPA were made aware of the incident and both agencies conducted investigations at ANL relating to the incident. IEPA issued a Compliance Inquiry Letter on November 14, 1994, stating that ANL was in apparent violation of certain special waste regulations in 35 IAC Parts 808 and 809. ANL formally responded through DOE on December 14, 1994. On January 10, 1995, IEPA formally notified ANL that ANL had returned to compliance for the apparent violations. There has been no correspondence from EPA following that agency's investigations.

During 1994, ANL exceeded action levels for copper and lead in drinking water samples. As a result, ANL must perform a number of follow-up actions. These actions

include additional sampling for copper, lead, and water quality parameters during 1995, distribution of educational materials to users of the water supply, and development of a corrosion control plan.

The IEPA-approved sanitary landfill groundwater monitoring program continues to indicate the Ground Water Quality Standards of some routine indicator parameters are consistently being exceeded. Continued monitoring of this site may provide data showing that additional extensive characterization and remediation are required.

Identification and clean-up of environmental contamination caused by previous activities on the ANL site remains an issue. These activities will primarily come under the purview of the RCRA programs administered by the EPA and IEPA. The ANL site has a significant number of such sites which will probably require extensive remediation to remove residual contamination resulting from past activities. The Five Year Plan contains a number of projects, termed Environmental Restoration projects, to provide for characterization and remediation of the sites. Several characterization projects are ongoing, while others are planned for the next few years. Remedial actions are scheduled to begin within three years, depending on the results of the characterization studies.

# 2.14.2. Regulatory Agency Interactions

The regulatory agency interactions with ANL during 1994 related primarily to site inspections regarding permit requirements and a PCB-contaminated oil issue. One outstanding compliance issue that has been formally identified by regulatory agencies relates to the closed ANL solid waste landfill. ANL's failure to modify its property deed to reflect the fact that the landfill had been used for the disposal of asbestos-containing materials, thus making said disposal part of the public record and subject to

discovery during a title search of the property, resulted in a formal notice of noncompliance from the IEPA. DOE is responsible for any required modifications to the property deed and provided a response to IEPA on July 8, 1994, indicating that steps would be taken to modify the deed. This issue remains open.

#### 2.14.3. Tiger Team Assessment

To resolve the deficiencies identified by the Tiger Team and the ANL self assessment, an Action Plan was prepared in December 1990. This plan lists specific actions to be taken to resolve each of the 256 Tiger Team findings and many of the self assessment findings. This document was approved by DOE Headquarters in early 1991. A number of the activities listed in the Action Plan were either ongoing actions or previously planned actions, many of which appear in the Five Year Plan. In addition, a series of new activities, not previously anticipated, were identified. These activities were started in 1991, contingent on additional funding provided by the DOE. A total of 153 action plans have been completed since 1991. An internal tracking system was developed to insure that the various commitments contained in the Action Plan are satisfied and the milestones are met.

#### 2.14.4. Progress Assessment Team

The DOE conducted a Tiger Team Progress Assessment of the ANL site from October 24, 1993, through November 9, 1993. The Team consisted of 13 professionals from various DOE offices and their support contractors, with expertise in the areas of management, quality assurance, environment, safety, and health. The purpose of the assessment was to provide the Secretary of Energy and senior DOE managers with concise independent information on the following: 1) change in culture and attitude related to ES&H activities; 2) progress and effectiveness of the ES&H corrective actions

resulting from previous Tiger Team Assessment; 3) adequacy and effectiveness of the ES&H self-assessment process; and 4) effectiveness of DOE and contractor management structures, resources, and systems to address ES&H problems and new ES&H initiatives effectively.

The results of the Assessment were: 12 concerns; four weaknesses; and six strengths. These were distilled into two key accomplishments (ANL and DOE-CH have effected ES&H cultural changes and there are organizational enhancements and have mobilized resources); two key concerns (inconsistent sitewide implementation of ES&H activities and DOE responsibility and accountability for integrated planning, budgeting, and resource allocation is not clearly understood or accepted); and one probable root cause (ANL and DOE have not developed and implemented integrated planning processes that define, guide, and set priorities for accomplishing the ES&H aspects of the ANL mission on the basis of sitewide risks and vulnerabilities). The conclusion of the Progress Assessment Team was that three years of intense activity on the part of ANL and DOE has resulted in substantial progress in correcting the deficiencies identified by the Tiger Team. Among the more important accomplishments are a positive culture change, an effective construction safety program, a graded approach to conduct of operations, enhanced ES&H training program, and improved ES&H related program plans.

#### 2.15. Environmental Permits

Table 2.9 lists all environmental permits in effect at the end of 1994. Other portions of this Chapter discuss special requirements of these permits and compliance with those requirements. The results of monitoring required by these permits are discussed in those sections, as well as in Chapters 5 and 6.

TABLE 2.9

ANL Environmental Permits in Effect on December 31, 1994

Permit Requirement	Source Name	Building	Date Issued	Expiration Date
Air	ALEX Alkali Metal Scrubber	370	12/5/91	12/3/96
Air	Alkali Booth	308	2/15/89	11/18/98
NESHAP	Alkali Booth	206	6/19/89	6/9/97
Air	Argonne Service Station	300	1/9/91	1/7/96
Air	Central Shops Rotoclone Dust Collection System	363	3/12/91	3/7/96
Air	Coal/Oil Fired MHD (FEUL Facility) - Withdrawn	146	3/30/90	3/27/95
Air	Gasoline Dispensing Facility (Modified) <sup>1</sup>	46	2/1/93	9/17/95
Air	Medical Department Steri-Vac Sterilizer	201	3/27/91	3/22/96
Air	Methanol/Gasoline Storage Tank	46	2/27/86	9/23/96
Air	Oil Fired Boilers	800 Area	11/1/91	10/29/96
Air	Open-Burning - Fire Training	Site-Wide	2/4/94	4/16/95
Air	Proton Decay Project Grieve Oven	366	8/8/91	8/6/96
Air	Boiler House	108	12/28/93	12/28/98
Air	Sulfuric Acid Storage Tank	108	1/17/91	12/1/99
Air	Vapor Degreaser	363	3/13/90	12/1/99
Air	Wastewater Air Stripper <sup>2</sup>	306	4/27/93	4/27/98
Air	Wood Shop Rotoclone Dust Collection System - Operate	809	10/22/91	10/17/96
Air	Dust Collector, Wood Shop - Construct	809	12/16/93	
NESHAP	Advanced Photon Source - Operating Permit - IEPA	400	12/21/93	7/26/98
NESHAP	Alpha-Gamma Hot Cell Facility	212	3/25/91	11/30/95
NESHAP	Building 212 Exhausts <sup>3</sup>	212	7/30/91	7/23/96
NESHAP	Building 306 Vents and 317 Area	306	8/6/91	7/25/96

TABLE 2.9 (Contd.)

Permit Requirement	Source Name	Building	Date Issued	Expiration Date
NESHAP	Continuous Wave Deuterium Detector (CWDD)	369	5/9/91	12/28/99
NESHAP	CP-5	330	5/10/91	12/28/99
NESHAP	Cyclotron	211	5/10/91	
NESHAP	EBWR HEPA Filtration System	331	3/25/91	12/1/99
NESHAP	Intense Pulsed Neutron Source	375		12/5/99
NESHAP	JANUS Reactor	202	3/25/91	11/30/95
NESHAP			5/10/91	11/30/95
NESHAP	M-Wing Hot Cells NBL Plutonium & Uranium Hoods	200	3/25/91	11/30/95
		350	4/25/91	4/19/96
NESHAP	Rad Hoods	Site-Wide	7/9/92	7/9/97
NESHAP	Map Tube Facility	317	5/10/94	5/10/99
Air	APS Emergency Generators	400	5/16/94	3/15/99
Air	Waste Bulking Sheds <sup>4</sup>	306	6/14/94	7/25/96
NESHAP	WMO HEPA Filter System	Site-Wide	9/28/94	9/28/99
Hazardous Waste	RCRA Part A Permit	Site-Wide	4/30/82	-
Hazardous Waste	RCRA Part A Modification - Storage Units	Site-Wide	2/18/93	-
Hazardous Waste	RCRA Part A Modification - Scintillation Vials	Site-Wide	9/22/93	-
Miscellaneous	Nuisance Wildlife Control	Site-Wide	1/31/95	12/31/95
Solid Waste	Landfill	800 Area	3/31/82	_
Solid Waste	Landfill	800 Area	3/30/89	-
Solid Waste	Landfill	800 Area	4/12/89	
Solid Waste	Landfill Leachate Test Wells	800 Area	8/31/90	

TABLE 2.9 (Contd.)

Permit	Source Name	Building	Date Issued	Expiration Date
Requirement	Source Traine	Dunuing	1550000	Date
Solid Waste	Landfill Groundwater Assessment	800 Area	9/30/91	-
Solid Waste	Landfill Leachate Characterization	800 Area	9/30/91	-
Solid Waste	Landfill Revised Closure Plan	800 Area	4/24/92 <sup>5</sup>	-
Solid Waste	Landfill Supplemental Closure Plan	800 Area	9/15/92	-
Water	APS Wetland	Site-Wide	11/22/88	-
Water	Landfill Wetlands	800 Area	5/20/81	· -
Water	Lime Sludge Application - Land Application	Site-Wide	1/12/94	12/31/98
Water	NPDES Permitted Outfalls	Site-Wide	10/31/94	7/1/99
Water	NPDES Stormwater Outfalls	Site-Wide	10/31/94	7/1/99
Water	Canal Water Treatment Plant Overflow	Site-Wide	3/5/93	-
Water	317 Area Pump Station and Force Main	Site-Wide	3/9/93	-
Water	Outfall #008 Rehabilitation	Site-Wide	6/1/93	-
Water	Solar Pond Dischage to DuPage County <sup>6</sup>	Site-Wide	6/3/93	₹.

<sup>&</sup>lt;sup>1</sup>Includes EtOH/Gasoline Tank.

<sup>&</sup>lt;sup>2</sup>Issued in conjunction with emergency RCRA Permit (expired).

<sup>&</sup>lt;sup>3</sup>Plasma Spray Booth added to permit May 27, 1994. <sup>4</sup>Construction Permit issued; operated under Building 306 permit. <sup>5</sup>Revised September 15, 1992, and October 22, 1992.

<sup>&</sup>lt;sup>6</sup>Solar Pond Discharge completed August 1993.



3.	ENVIRONMENTAL	PROGRAM INFO	RMATION	
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It is the policy of the DOE and ANL to conduct all operations in compliance with applicable environmental statutes, regulations, and standards and to ensure that environmental obligations are carried out consistently across all operations and organizations. Protection of the environment and human health and safety are given the highest priority. At ANL, a number of programs and organizations exist to ensure compliance with these regulations and to monitor and minimize the impact ANL operations have on the environment.

# 3.1. Environment and Waste Management Program

ANL management has designated the Environment and Waste Management Program (EWM) as the lead environmental support organization. The mission of EWM is to proactively support the ANL operations by conducting those activities that ensure compliance with applicable environmental statutes, regulations, DOE Orders, and ANL policies and procedures. These activities include: the technical support in the preparation of permits and compliance documents, consideration of applicable regulatory requirements, and liaison with oversight and compliance organizations; proper collection, treatment, and disposal of radioactive, hazardous, and non-hazardous waste materials; the characterization, remediation, decontamination and decommissioning of facilities, operations, and areas; and the conduct of the ANL environmental monitoring and surveillance program. These activities are carried out to minimize the potential adverse effects to the health and safety of persons at the ANL site and the general public, to property and to the environment.

EWM is divided into five major operational departments: Environmental Projects; D&D Projects (transferred to the Technology Development Division on March 22, 1994); Waste Management Operations; Waste Reduction; and Monitoring, Surveillance, and Environmental Compliance. The principal function of EWM is to serve as the ANL focal

point for the execution of the DOE Environmental Restoration and Waste Management Program (now Environmental Management Operations).

In 1989, the DOE established the goal of achieving compliance with applicable regulations and assessing and cleaning up releases of hazardous materials from inactive waste sites, returning all such sites to unrestricted use within 30 years. As a management tool to improve the achievement of this goal, the DOE established the Environmental Restoration and Waste Management Program. This program identifies specific needs and established a system for allocating funds to resolve the various deficiencies. Each of the DOE facilities has prepared a set of planning documents (Activity Data Sheets, or ADSs) describing the activities necessary to bring that specific site into compliance and to identify and clean up inactive waste sites. These planning documents are contained in a report which is updated and published annually, the Environmental Restoration and Waste Management Five Year Plan. Five Year Plan projects and activities are subdivided into three categories, namely, corrective activities (those actions necessary in the short term to bring a facility into compliance with environmental regulations), environmental restoration activities (those activities necessary to identify and clean up inactive waste sites and other sites potentially contaminated as a result of DOE activities) and waste management activities( designed to ensure that hazardous and radioactive wastes are stored and disposed of safely and the volume of waste is minimized). The 1994 Five Year Plan contained information on 19 separate projects. The on-site activities are listed in Table 3.1. The Five Year Plan is a public document available upon request from the DOE.

### 3.1.1. Environmental Projects

The role of the Environmental Projects Department is to support ANL operations, organizations, and DOE environmental missions by managing environmental projects in

**TABLE 3.1** 

Environmental Restoration and Waste Management Projects

ADS Number	Title
CH 1300	Facility Operations and Maintenance - Defense Programs
CH 1301A	Waste Management Projects - Non-Defense Programs
CH 1301B	Facility Operations and Maintenance - Non-Defense Programs
CH 1303	Rehabilitation of Waste Management Building
CH 1304	Hazardous, Radioactive, and Mixed Waste Storage Facility
CH 1306	Sanitary Wastewater Treatment Plant Improvements
CH 1308	Laboratory and Sanitary Sewage Collection System Rehabilitation
CH 1309	Laboratory Wastewater Treatment Plant Improvements
CH 1310	Treatment of Boiler House Area Wastewater
CH 1311	Canal Water Treatment Plant Rehabilitation
CH 1313	Cooling Tower Blowdown Water Diversion
CH 1432	Remedial Support Activities
CH 1433	Solid Waste Storage/Disposal
CH 1434	Mixed Waste Storage/Disposal Facilities
CH 1435	Treatment Sites
CH 1436	Facility Conversion D&D Projects
CH 1437	Reactor Facilities D&D Projects
CH 1438	Support Facilities D&D Projects
CH 1439	Program Management

accordance with applicable DOE Orders and environmental statutes. The Department provides project management and engineering support for environmental remediation projects. Project management functions include: the development of work scopes; project budgets; and schedules. The projects implemented by this Department are designed to minimize any current or future impact to the environment or human health.

The corrective activity projects at ANL involve the construction of new or upgraded wastewater treatment facilities used for disposal of wastewater from the ANL. As discussed in Chapters 2 and 5, the site has experienced a number of violations of its NPDES wastewater discharge permit in recent years. The reason for many of these violations is the lack of appropriate treatment technology to comply with current effluent limits. These deficiencies will be resolved as these corrective action projects are completed. During 1994, design work on several facilities was started.

Environmental Restoration activities represent the projects designed to carry out the objective of assessing and cleaning up inactive waste sites. The ANL site contains a number of inactive waste sites used for disposal of waste during the early years of Laboratory operations. These sites include two inactive landfills, three French drains (which consisted of shallow pits used for disposal of liquid wastes), two inactive wastewater treatment facilities and a number of areas which may have been contaminated through the discharge of small amounts of hazardous chemicals. Several sites used from the 1940s through the 1970s for open burning of combustible waste and construction debris also exist. A series of ongoing and planned activities has been designed to foster the clean up of these sites.

The Environmental Restoration projects at ANL are typically broken down into two phases, the characterization phase and the remediation phase. Several of the characterization projects were started in 1989 and 1990. Additional characterization is

required before significant remediation can be undertaken. The results of some of this early characterization work is presented in Chapter 6. Following the characterization phase, projects designed to clean up and dispose of residual contamination found during characterization will commence.

#### 3.1.2. D&D Projects

The mission of the D&D Projects is to proactively support the ANL and DOE mission in the D&D area by conducting those activities that promote compliance with applicable DOE, ANL, federal, and state regulations and procedures. This includes: directing and planning all D&D Program activities at the ANL site, interfacing with DOE on the ANL D&D Program, performing planning and scheduling of D&D at the ANL site and investigating new or innovative approaches to accomplish D&D project work in a more timely and cost effective manner. Conduct of the D&D Program at ANL shall be done in a manner with due regard for the environment and public and worker safety and health.

In addition to the inactive waste site clean up projects, the Five Year Plan also contains a number of Decontamination and Decommissioning (D&D) projects for on-site nuclear facilities. The ANL site contains several inactive nuclear reactors and hot cells used in the past for processing of radioactive materials. These facilities are either currently undergoing D&D or are scheduled for D&D in the next few years. The D&D operations will remove residual radiological contamination, will dispose of radiologically contaminated materials, and will return the facilities to unrestricted use status. The largest such activities are the D&D of the Experimental Boiling Water Reactor (EBWR) and the CP-5 research reactor.

Current technology is not adequate to process and dispose properly many of the waste materials that may be generated by these activities. Much of the waste is a mixture of

radioactive and chemically hazardous materials. This waste will be treated in accordance with the Federal Facility Compliance Act Site Treatment Plan.

## 3.1.3. Waste Management Department

The mission of the Waste Management Department is to provide for treatment, storage, and disposal of all regulated waste generated at ANL in compliance with state and federal regulations at minimal cost; and to supply skilled craftspeople, uniquely trained to safely provide decontamination and operational support activities for facilities which generate radioactive, hazardous, and other special wastes.

The projects included in this section of the Five Year Plan represent activities necessary to ensure that waste materials currently being generated are properly stored, treated and disposed. A primary motivation for the improvement in waste handling and disposal operation is the need to upgrade such facilities to comply with increasingly stringent RCRA requirements as well as other state and federal regulations and DOE orders. The majority of the Waste Management projects involve improvements to existing treatment or storage facilities.

# 3.1.4. Waste Reduction Department

The role of the Department is to develop, promote, and implement waste reduction technologies, practices, policies, and environmental quality through training, review, culture change, and operational activities at ANL in support of Laboratory and DOE missions.

## 3.1.5. Monitoring, Surveillance and Environmental Compliance Department

The Department is composed of the Environmental Compliance Section and the Monitoring and Surveillance Section. Environmental protection activities are those sets of actions conducted at ANL which are needed to ensure the safety of the public, protection of the environment, and compliance with applicable federal, state, and local environmental regulations and with the DOE Orders.

The mission of the Department is to define the applicable compliance requirements with assistance from the Legal Department of ANL. The Department helps to ensure that ANL is in compliance with these standards. The activities of the Department include: defining applicable federal, state, and local regulations; define applicable DOE Orders; develop site-wide regulatory programs; provide technical support in preparing permits and NEPA documents; provide technical support and guidance to the ANL programs through the ECR and ESH representatives; conduct reviews of construction projects and experiments; and act as liaison with external regulatory agencies and to coordinate with internal research and support groups. This information and services for the Department are transmitted to the programmatic and operations groups at the Laboratory.

Monitoring and surveillance are conducted to determine the effects, if any, of ANL activities on the public and on the on-site and off-site environment. Effluent monitoring is the collection and analyses of samples, or measurements of liquid and gaseous effluents for the purpose of characterizing and quantifying contaminants, assessing radiation exposures to members of the public, providing a means to control effluents at or near the point of discharge, and to demonstrate compliance with applicable standards and permit requirements. Environmental surveillance is the collection and analysis of samples or direct measurements of air, water, soil, foodstuffs, biota, and other media from the ANL site and its environs for the purpose of determining compliance with applicable standards and permit require-

ments, assessing radiation exposure of the public and assessing the effects, if any, of ANL operations on the local environment. The information generated by this program is compiled each year in the ANL Site Environmental Report which is distributed to ANL and DOE personnel and to the federal, state, and local regulators.

# 3.2. Environmental Support Programs

#### 3.2.1. Self-Assessment

In preparation for the 1990 Tiger Team visit, ANL conducted a formal top-down environmental self-assessment. This was followed in the next two years with self-assessments on Tiger Team findings, root causes, and action plans. For 1994, the self-assessment was expanded and formalized to document accomplishments, clarify and evaluate the organization's status, and to develop short- and long-term goals and plans for improvement. All ANL organizational units were required to participate in the 1994 assessment.

A guidance document was prepared to standardize the self-assessment process and organizational assignments were made to coordinate the effort. Nine topics were selected for the 1994 self-assessment and these are listed in Table 3.2. The product of the assessment was a report that briefly described the mission of the organization, the assessment process, provided a summary of the overall results of the assessment against the nine topics, and established goals, timelines, and milestones. Organizational unit reports were rolled-up into higher level reports.

## **TABLE 3.2**

# Topics for the 1994 ANL Self-Assessment

A.	Job or Facility Specific Training
В.	Documentation and Records
C.	Implementation of Building Emergency Plans
D.	Risk Management and Occupational Injuries
E.	Egress Management (Life Safety)
F.	Implementation of Chemical Hygiene Plans
G.	Implementation of ALARA Process for Radiological Exposure
H.	Environmental Protection
I.	Waste Minimization

## 3.2.2. Environmental Training Programs

ANL has a comprehensive environmental protection training program which includes mechanisms to identify, track, and document requirements for every employee. Environmental protection training for ANL personnel is primarily provided by the ESH Training Section, although ancillary training may be delivered by subject matter experts from other organizations. Personnel training requirements are mandated by DOE Orders, DOT, EPA, and OSHA regulations are identified by a Job Hazards Checklist form which is completed by every employee and reviewed by the employee's supervisor. A positive answer to any one of a battery of specific questions triggers the training requirements specific to that question. There are also options for divisionally-required training, recommended training, and elective training.

The logistics for ensuring that the training 1) meets compliance, 2) is completed, and 3) is documented is managed through the Training Management System, an "on-line" main frame computer-based system. Environmental protection training courses and course descriptions are listed in the Training Course Catalog available from divisional representatives, the ESH Training System, or from Human Resources.

#### 3.2.3. Pollution Prevention - Waste Minimization

Waste minimization is a policy specifically mandated by the Congress in the 1984 Hazardous and Solid Wastes Amendments to the Resource Conservation and Recovery Act (RCRA). RCRA requires hazardous waste generators to establish a program to reduce the volume or toxicity of waste to the degree determined by the generator to be "economically practicable." Hazardous waste generators, such as ANL, must certify in their waste manifest that this requirement has been fulfilled. Generators must also identify in their

biennial reports to the IEPA the efforts undertaken during the year to reduce the volume and toxicity of waste generated and the changes in volume and toxicity actually achieved.

Pollution is to be prevented at the source wherever and whenever possible. Those potential waste materials that cannot be eliminated or minimized by source reduction are to be recycled, i.e., used, reused, or reclaimed. All waste that is nevertheless generated is to be treated to reduce volume, toxicity, or mobility before storage or disposal. Reducing or eliminating the generation of waste should be given prime consideration in research, process design, and plant operations.

DOE Orders 5400.1, 5400.3, and 5820.2A mandate that the management of radioactive wastes and other pollutants shall be accomplished in a manner that minimizes the generation of such wastes.

ANL has a long-standing history of pollution prevention and waste minimization dating back to the Manhattan Engineering District. Early activities included recycling of metals and rare materials. Although no formal pollution prevention and waste minimization plan is in place, through the efforts of the Waste Reduction Department, ANL continues to focus on pollution prevention and waste minimization activities. A few of these activities include:

- Paper, metal, and toner cartridge recycling
- · Actinide recovery for re-use and recycle
- Refuse recycling through contractor service
- Use of alternate cleaning fluids for the Advanced Photon Source beam lines
- Use of alternate methods of cleaning and painting water towers
- Filtering of Central Shops aqueous lubricants
- Development of a process to concentrate human excrement samples for radiological analysis resulting in source reduction

- Adaptation of micro-experimentation techniques in chemical synthesis
- Adaptation of bacteriological testing to replace animal testing applications
- Substitution of non-hazardous scintillation flours for hazardous flours
- Specifications requiring use of retread tires and fly ash in concrete are in place
- ANL research programs order minimal quantities of material on an as needed basis
- White paper recycling campaign for reducing waste generation
- Development of a Chemical Management System for control of inventory

The level of achievement for the above-mentioned activities will be determined by the amount of funding provided to the Waste Reduction Department during Fiscal Year 1995. The coordination and development of several pollution prevention and waste minimization concepts will be provided by a Pollution Prevention Task Force that was formed during 1993.

# 3.3. Environmental Monitoring Program Description

As required by DOE Order 5400.1, ANL conducts a routine environmental monitoring program. This program is designed to determine the effect the operation of ANL is having on the environment surrounding the site. This section describes this monitoring program. A general description of the techniques used to sample each environmental medium is provided. This is followed by the collection procedures, the sampling schedule and analytical techniques used. Greater detail is provided in the Environmental Monitoring Plan.

## 3.3.1. Air Sampling

Continuously operating air samplers are used at ANL to measure the concentrations of airborne particulate radioactivity. There is currently no monitoring of non-radiological air contaminants in ambient air. Particulate samplers are placed at 15 locations around the ANL perimeter and at six off-site locations, approximately five miles from ANL to determine the ambient or background concentrations.

Airborne particulate samples for direct radiation measurement are collected continuously at 13 perimeter locations and at five off-site locations on glass fiber filter media. Average flow rates on the air samplers are about 70 m<sup>3</sup>/hr. Filters are changed weekly. The filters on perimeter samplers are changed by ANL staff and the filters on off-site samplers are changed and mailed to ANL by cooperating local agencies. The sampling units are serviced every six months and the flow meters are recalibrated annually.

Additional air samples, used for radiochemical analysis of plutonium and other radionuclides, are collected at two perimeter locations and one off-site location. These samples are collected on special filter media which are changed every ten days by ANL staff. The flow rate calibration and servicing schedule is the same as discussed above.

At the time of sample collection, the date and time when sample collection began, the initial flow rate, the date and time when the sample was collected and final flow rate are recorded on a label attached to the sample container. The samples are then transported to ANL where this information is then transferred to the environmental protection data management system.

Each air filter sample collected for direct measurement is cut in half. Half of each sample for any calendar week is combined with all the other perimeter samples from that

week and packaged for gamma-ray spectrometry. A similar package is prepared for the offsite filters for each week. A two-inch circle is cut from the other half of the filter, mounted in a two-inch low-lip stainless steel planchet, and counted for alpha and for beta activity. The balance of the filter is saved.

The air filter samples collected for radiochemical analysis are composited by location for each month. After addition of the appropriate tracers, the samples are ashed, and then sequentially analyzed for plutonium, thorium, uranium, and strontium.

Stack monitoring is conducted continuously at those emission points that have a probability of releasing measurable radioactive effluents. The results of these measurements are used for calculating the theoretical annual off-site dose using the required CAP-88 version of the EPA-AIRDOSE atmospheric dispersion computer code and dose conversion.

#### 3.3.2. Water Sampling

Water samples are collected to determine what, if any, radioactive materials or selected hazardous chemicals used or generated at ANL enter the environment by the water pathway. The samples are collected from Sawmill Creek below the point at which ANL discharges its treated wastewater and stormwater. The results of radiological analysis of water collecting at this location are compared to upstream and off-site results to determine the ANL contribution. The results of the chemical analysis are compared to the applicable IEPA stream quality standards to determine if the site is degrading the quality of the creek. These results are discussed in more detail in Chapters 4 and 5.

In addition to surface water, subsurface water samples are also collected at approximately 34 locations. These samples are collected from monitoring wells located near

sites which have the potential for adversely impacting groundwater. These sites are the 800 Area landfill, the 317/319 waste management area, and the site of the inactive CP-5 reactor. Samples of the domestic water, which comes from four on-site wells, are also collected and analyzed for hazardous or radioactive constituents.

Surface water samples are collected from Sawmill Creek and composited into a single weekly composite sample. A continuous sampling device has been installed at this location to improve sample collection efficiency. To provide control samples, Sawmill Creek is sampled upstream of ANL once a month. The Des Plaines River is sampled twice a month below, and monthly above, the mouth of Sawmill Creek to determine if the radioactivity in the Creek had any effect on the activity in the River. Water samples are collected from remote locations in the spring and fall to serve as additional control samples.

Subsurface water samples are collected quarterly from the monitoring wells located in the 317/319 Area, 330 (CP-5), and the 800 Area Sanitary Landfill. The monitoring wells are purged and samples collected from the recharged well water. These samples are analyzed for both chemical and radiological constituents, as discussed in Chapter 6. Samples are collected quarterly from the well-heads of the four ANL wells used to provide the Laboratory domestic water supply. The water is pumped to the surface and collected in appropriate containers depending on the required analysis.

At the time of sample collection for radiological analysis, the sampling location, time, date and collector identification number are recorded on a label attached to the sample container. Upon return to the laboratory, the information is transferred to the EMS system. Each sample is assigned a unique number, which accompanies it through all analyses.

After the sample has been logged in, an aliquot is removed for tritium analysis, 20 mL of conc. HNO<sub>3</sub> is added per gallon of water as a preservative, and the sample is filtered

through Whatman #2 filter paper to remove sediment present in the sample. Appropriate aliquots are then taken depending on the analysis.

For nonradiological analysis, samples are collected and preserved using EPA prescribed procedures. Cooling is used for organic analysis and nitric acid is used to preserve samples to be analyzed for metals. Specific collection procedures are used for other components and EPA methods are used. All samples are analyzed within the required holding period or noncompliance is documented. The quality control requirements of either SW-846 and/or CLP are met or deviations are documented. All samples are assigned a unique number which serves as a reference source for each sample. When duplicate samples are obtained, unique numbers are assigned and the indication that duplicates exist is noted in the data management system.

#### 3.3.3. Bottom Sediment

Bottom sediment accumulates small amounts of radioactive materials which may be present from time to time in the stream and, as a result, acts as an integrator of radioactive material that was present in the water. It provides a historical record of radioactive materials in that surface water system. These samples are not routinely analyzed for chemical constituents.

Bottom sediment samples are collected annually from Sawmill Creek above, at, and several locations below the point at which ANL discharges its treated wastewater. Periodically, sediment samples are collected from several on-site ponds and lagoons. Ten off-site bottom sediment samples are collected each year, five in the spring and five in the fall, from remote locations to serve as controls. One gallon of sediment is collected from each location with a stainless steel scoop and transferred to a glass bottle.

At the time of sample collection, the date, time, and sample collector identification are recorded on sample labels affixed to the sample container. Upon return to the laboratory, the information is transferred to the EMS system. Each sample is assigned a unique number which accompanies it through the process.

Each sample is dried for several days at 110°C, ball milled, and sieved through a No. 70 mesh screen. The material that does not pass the No. 70 screen is discarded. A 100 g portion is taken for gamma-ray spectrometric measurement and other appropriate aliquots are used for specific radiochemical analyses.

#### 3.3.4. Soil

Soil accumulates small amounts of particulate matter and serves as an integrator of the deposition of airborne releases of radioactive materials. Although it should not be used as the primary measurement system for air monitoring, in many cases, it may be the only available avenue if insufficient air sampling occurred at the time of an incident. The ANL program is designed to provide samples for analysis to determine if any changes in concentrations have occurred over the year. No analysis for chemical constituents is carried out on these samples.

Each year soil from ten locations is collected at the site perimeter (five spring and five fall) and ten at remote locations (five spring and five fall). Sampling sites are selected in reasonably level areas that represent undisturbed soil. Two one-meter squares are marked off and soil samples are collected from the corners and center of each square. Samples are collected with a 10.4 cm-diameter coring tool to a depth of 5 cm. All ten cores are composited as a single sample. This procedure follows the ASTM Standard Method for Sampling Surface Soil for Radionuclides, C-998.

At the time of sample collection, the date, time, and sample collector identification number are recorded on a preprinted sample label affixed to the sample container. Upon return to the laboratory, the information is transferred to the EMS system. Each sample is assigned a unique number which accompanies it through the process.

The entire sample is dried at 110°C for several days, ball milled, and sieved through a No. 70 mesh screen. The material that does not pass the No. 70 mesh screen is discarded. A 100 g portion is taken for gamma-ray spectrometric measurement and appropriate aliquots taken for radiochemical analysis. Because a known area of surface soil was collected, results are calculated in terms of concentration and deposition.

#### 3.3.5. Vegetation

Grass samples are collected to determine the uptake of radionuclides from the soil by vegetation. This is done to monitor that part of the food chain pathway.

Grass samples are collected each year from ten perimeter and ten off-site locations at the same places as the soil samples. All the grass within one of the one-meter plots used for soil sampling is cut just above the soil surface and collected.

At the time of sample collection, the date, time, and sample collector identification number are recorded on a preprinted sample label affixed to the sample container. Upon return to the laboratory, the information is transferred to the EMS system. Each sample is assigned a unique number which accompanies it through the process.

Grass samples are washed in water to remove surface dirt, dried at 110°C for several days, and ground. A 100 g aliquot is measured by gamma-ray spectrometry and appropriate aliquots taken for radiochemical analysis.

#### 3.3.6. External Penetrating Radiation

Measurements of direct penetrating radiation emanating from several sources within ANL are made using calcium fluoride thermoluminescent dosimeter (TLD) chips. Each measurement is the average of four chips exposed in the same packet. All calcium fluoride packets are shielded with 1/16 inch copper foil to reduce or eliminate the beta and low-energy x-ray components. The response of the chips is determined with a U. S. National Institute of Standards and Technology (NIST) standard radium-226 source.

Dosimeters are exposed at 14 locations at the site perimeter and on the site and at five locations off the site. All dosimeters are changed quarterly.

At the time of dosimeter collection, the date, time, and collector identification number are recorded on a preprinted label affixed to the container. Upon return to the laboratory, the information is transferred to the EMS system. Each sample is assigned a unique number which accompanies it through the process.

The individual chips are read on an Eberline Model TLR-6 TLD reader. Control chips are read and their contribution subtracted from the values of the field chips. A set of chips irradiated with a radium-226 standard source is also read and these values are used to convert the individual field readings to dose.

## 3.3.7. Data Management

The management of the large amount of data assembled in the environmental monitoring program is handled by ANL in a very structured manner that allows a number of reports to be generated. Basic radiological data management, including sample recordkeeping, is done with the EMS computerized recordkeeping program. All sample and analytical data are kept in the EMS for eventual output in formats required for either regulatory compliance reports or for the annual reports. In addition, reports are provided for trend analysis, statistical analysis, and tracking.

The ANL-developed EMS program is the basic data management tool; it generates sampling schedules, all other tracking and calculation routines, and the final analytical result tabulations. The EMS program is set up for the radiological portion of the monitoring program and nonradiological monitoring for groundwater and NPDES surface water effluents. For purposes of this plan, the procedures for nonradiological sampling follow the same basic protocol as shown below.

The starting point for effluent monitoring and environmental surveillance is establishing a set of sampling locations and a sample schedule. Based on either regulatory parameters, pathway analysis, or professional judgement, sample locations for the various media are identified and entered into the EMS. For each sample location, nine categories of data are entered into the EMS. They are: geographic code; location description; sampling frequency; sample type (water, soil, plant, etc.); exact sampling position; last date sampled; sampling priority (same location with multiple samples); size of sample to collect; and analytes.

Once the data are entered, the EMS program is used to generate a sampling schedule. Every week a schedule for the next week is printed out, along with uniquely numbered, preprinted labels for the sample containers. These items are provided to the staff who are doing the sampling in the field. Field data is entered into the EMS system. At the time when the samples are submitted to the analytical laboratory, chain-of-custody documents are generated. In addition, the same information is provided electronically to assist the laboratory in data entry. All necessary information required by the laboratory is provided.

As the laboratory results are compiled, the data are entered into the EMS program. This permits up-to-date tracking of all samples currently in process. When the analysis is completed on each sample and the results electronically entered into the EMS, the completed final results sample card is retained in a file as an additional quality assurance measure.

Complete data sets for all samples are maintained by the EMS program. When all radiochemical results are complete and entered into the EMS, a final result card is generated listing all data related to each sample. The electronic files are backed up by the EWM computer network server. The printed final result card is filed after review, then ultimately put in DOE's archives in Chicago. Annually, EWM staff print and bind for reference the complete results, by sample type for the past calendar year. Final results are thus available both on-line via the network and in hard copy.

## 3.4. Compliance with DOE Order 5820.2A

DOE Order 5820.2A "Radioactive Waste Management" requires that an environmental monitoring and surveillance program be conducted, Section III-3 (k), to determine any releases or migration from low-level treatment, storage, or disposal sites. Compliance with these requirements is an integral part of the ANL site-wide monitoring and

#### 3. ENVIRONMENTAL PROGRAM INFORMATION

surveillance program. To cover the waste management operations in general, reliance is made on the perimeter air monitoring network and monitoring of the liquid effluent streams and the Sawmill Creek. The analytical results are collected in Chapter 4 of this report.

Of particular interest is monitoring of the waste management activities conducted in the 317 Area. This includes: air monitoring for total alpha, total beta, gamma-ray emitters, and radiochemical determinations of plutonium, uranium, thorium, and strontium-90; direct radiation measurements with TLDs; surface water discharges for tritium and gamma-ray emitters; perimeter soil and plant samples analyzed for gamma-ray emitters, plutonium, and americium; and subsurface water samples at 15 monitoring wells with analyses for tritium, strontium-90, and gamma-ray emitters, plus selected monitoring for volatile organic compounds. The results are collected in Chapter 4 and Chapter 6 of this report.



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## 4.1. Description of Monitoring Program

The radioactivity of the environment around ANL was determined by measuring the concentrations of radioactive nuclides in naturally occurring materials and by measuring the external penetrating radiation dose. Sample collections and measurements were made at the site perimeter and off the site for comparative purposes. Some on-site results are also reported when they are useful in interpreting perimeter and off-site results.

Since radioactivity is primarily transported by air and water, the sample collection program concentrated on these media. In addition, samples of soil, plants, and materials from the beds of lakes and streams also were analyzed. The program followed the guidance provided in the DOE Environmental Regulatory Guide.<sup>5</sup> About 1,274 samples were collected and approximately 3,900 analyses were performed. The results of radioactivity measurements are expressed in terms of picocuries per liter (pCi/L) for water; femtocuries per cubic meter (fCi/m³) and attocuries per cubic meter (aCi/m³) for air; and picocuries per gram (pCi/g), femtocuries per gram (fCi/g), and/or nanocuries per square meter (nCi/m²) for soil, bottom sediment, and vegetation. Penetrating radiation measurements are reported in units of millirem per year (mrem/y) and population dose in man-rem. Other units are defined in the text.

The DOE has provided guidance<sup>6</sup> for effective dose equivalent calculations for members of the public, based on ICRP-26 and ICRP-30.<sup>7</sup> Those procedures have been used in this report. The methodology requires three components to be calculated: (1) the committed effective dose equivalent from all sources of ingestion, (2) the committed effective dose equivalent from inhalation, and (3) direct effective dose equivalent from external radiation. These three components are summed for comparison with the DOE effective dose equivalent limits for environmental exposure. The guidance requires that sufficient data on exposure

to radionuclide sources be available to assure that at least 90% of the total committed effective dose equivalent is accounted for. The primary radiation dose limit for members of the public is 100 mrem/y. The effective dose equivalents for members of the public from all routine DOE operations, natural background and medical exposures excluded, shall not exceed the values and shall be as low as reasonably achievable (ALARA), or as far below the limits as is practical taking into account social, economic, technical, practical, and public policy considerations. Routine DOE operations are normally planned operations, which exclude actual or potential accidental or unplanned releases.

The measured or calculated environmental radionuclide concentrations are converted to a 50-year committed effective dose equivalent with the use of the Committed Effective Dose Equivalent (CEDE) Factors<sup>8</sup> and compared to the annual dose limits for uncontrolled areas. The CEDE are calculated from the DOE Derived Concentration Guides (DCG)<sup>6</sup> for members of the public from ingested water and inhalation resulting in a radiation dose of 100 mrem/y. The numerical values of the CEDE factors used in this report are given in Table 4.30. Although the CEDE factors apply only to concentrations above natural levels, the calculated dose is sometimes given in this report for radioactivities that are primarily of natural origin for comparison purposes. Such values are enclosed in parentheses to indicate this. Occasionally, other standards are used, and their sources are identified in the text.

#### 4.2. Air

The radioactive content of particulate matter in the air was determined by collecting and analyzing air-filter samples. The sampling locations are shown in Figures 1.1 and 1.2. Separate collections were made for specific radiochemical analyses and for gross alpha, gross beta, and gamma-ray spectrometry. The latter measurements were made on samples

collected continuously on laminated glass fiber filters (changed weekly) at 13 locations at the ANL site perimeter using PM-10 units and at five off-site locations.

Samples were collected at the site perimeter to determine if a statistically significant difference exists between perimeter measurements and measurements made on samples collected at various off-site locations. The off-site samples establish the local background concentrations of naturally-occurring or ubiquitous man-made radionuclides, such as from nuclear weapons testing fallout. Higher levels of radioactivity in the air measured at the site perimeter may indicate radioactivity releases from ANL, providing the perimeter samples are greater than the background samples by an amount greater than the relative error of the measurement. The relative error is a result of natural variation in background concentrations as well as sampling and measurement error. This relative error is typically 5% to 20% of the measurement value for most of the analyses, but approaches 100% at values near the detection limit of the instrument.

The total alpha and beta activities in the individual weekly samples are summarized in Table 4.1. These measurements were made in low-background gas-flow proportional counters, and the counting efficiencies used to convert counting rates to disintegration rates were those measured for a 0.05 MeV beta and a 5.5 MeV alpha on filter paper. The average concentrations of gamma-ray emitters, as determined by gamma-ray spectrometry performed on composite weekly samples, are given in Table 4.2. The gamma-ray detector is a shielded germanium diode calibrated for each gamma-ray emitting nuclide measured.

The alpha activity, principally due to naturally-occurring nuclides, averaged the same as in the past several years and was in its normal range. The perimeter beta activity averaged 30 fCi/m³, which is slightly higher than the average value for the past five years. The gamma-ray emitters listed in Table 4.2 are those that have been present in the air for the past few years and are of natural origin. The beryllium-7 exhibits an increase in

TABLE 4.1

Total Alpha and Beta Activities in Air-Filter Samples, 1994\*

(concentrations in fCi/m³)

		No. of	Alph	a Activ	ity	Bet	a Activit	y
Month	Location	Samples	Avg.	Min.	Max.	Avg.	Min.	Max.
January	Perimeter	34	2.2	1.3	2.9	42.1	33.3	58.2
	Off-Site	18	2.6	1.0	4.6	37.7	23.9	63.8
February	Perimeter	47	1.7	1.3	2.1	32.4	21.1	36.7
	Off-Site	17	1.6	0.3	3.2	26.7	9.3	36.5
March	Perimeter	50	1.6	1.2	2.0	27.1	19.7	34.5
	Off-Site	23	2.0	1.2	4.2	28.1	17.9	47.5
April	Perimeter	42	1.5	0.7	2.2	20.4	9.0	28.5
	Off-Site	18	1.7	0.9	2.3	20.6	10.8	29.9
May	Perimeter	51	1.3	0.5	2.0	18.1	9.9	25.8
	Off-Site	18	1.6	0.8	4.4	18.3	10.9	28.6
June	Perimeter	52	1.6	0.9	2.3	24.2	15.3	29.6
	Off-Site	23	1.8	0.9	3.3	23.7	14.6	30.9
July	Perimeter	40	1.2	0.1	2.3	20.5	1.6	28.5
	Off-Site	19	1.4	0.5	2.2	21.0	11.2	35.0
August	Perimeter	60	1.9	0.7	2.8	28.4	14.7	38.3
	Off-Site	21	2.0	0.7	2.9	28.1	13.2	39.6
September	Perimeter	49	2.0	1.3	3.1	34.7	17.9	61.5
	Off-Site	17	1.9	0.6	4.1	32.4	12.6	66.4
October	Perimeter	45	1.8	1.3	2.6	29.0	23.0	35.8
	Off-Site	17	1.4	0.3	2.8	23.9	13.7	39.3
November	Perimeter	53	1.6	1.0	2.4	30.7	24.3	45.2
	Off-Site	22	1.6	0.5	2.7	25.5	11.3	44.3
December	Perimeter	32	2.4	1.3	3.3	54.6	26.9	73.9
	Off-Site	15	2.8	0.8	5.7	46.3	21.4	81.1
Annual	Perimeter	555	$1.7 \pm 0.2$		3.3		1.6	73.9
Summary	Off-Site	228	$1.9 \pm 0.3$	0.3	5.7	$27.7 \pm 5.0$	9.3	81.1

<sup>\*</sup> These results were obtained by measuring the samples four days after they were collected to avoid counting the natural activity due to short-lived radon and thoron decay products. This activity is normally present in the air and disappears within four days by radioactive decay.

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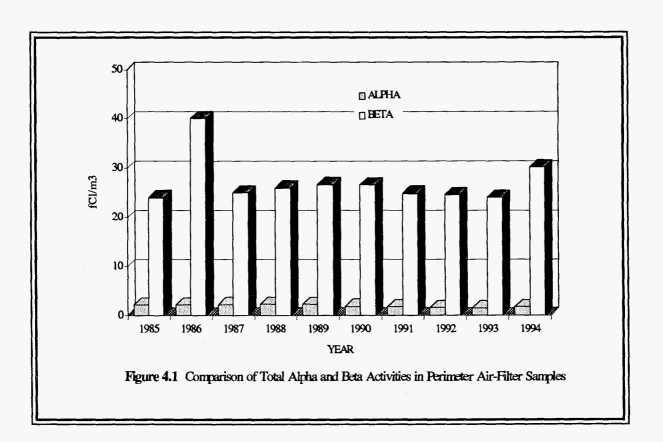
TABLE 4.2

Gamma-Ray Activity in Air-Filter Samples, 1994
(concentrations in fCi/m³)

Month	Location	Beryllium-7	Lead-210
January	Perimeter	94	38
,	Off-Site	65	26
February	Perimeter	126	23
•	Off-Site	77	27
March	Perimeter	132	19 .
	Off-Site	117	19
April	Perimeter	129	12
	Off-Site	97	10
May	Perimeter	128	10
	Off-Site	104	9
June	Perimeter	155	14
	Off-Site	93	29
July	Perimeter	108	14
	Off-Site	75	11
August	Perimeter	128	21
	Off-Site	98	19
September	Perimeter	151	30
	Off-Site	102	23
October	Perimeter	99	19
	Off-Site	88	19
November	Perimeter	103	23
	Off-Site	72	18
December	Perimeter	114	42
	Off-Site	92	40
Annual	Perimeter	122 ± 12	$22 \pm 6$
Summary	Off-Site	90 ± 9	21 ± 5
Dose(mrem)	Perimeter	(0.00031)	(2.52)
	Off-Site	(0.00023)	(2.36)

concentration in the spring, indicating its stratospheric origin. The lead-210 in air is due to the radioactive decay of gaseous radon-222 and is similar to last year.

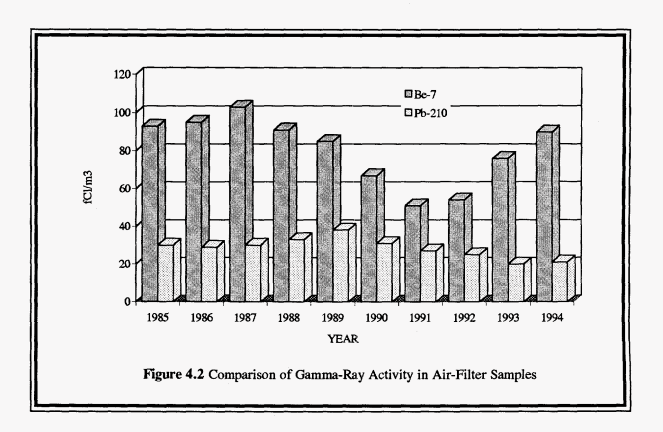
The annual average alpha and beta activities since 1985 are displayed in Figure 4.1. The elevated beta activity in 1986 was due to fallout from the Chernobyl incident. If the radionuclides attributed to the Chernobyl incident are subtracted from the annual beta average of 40 fCi/m³, the net would be 27 fCi/m³, very similar to the averages of the other years. Figure 4.2 presents the annual average concentrations of the two major gamma-ray-emitting radionuclides in air. The annual average beryllium-7 concentrations have decreased regularly since 1987, reached a minimum in 1991, and are now increasing. The downward trend in the beryllium-7 air concentrations has been observed worldwide by the DOE



Environmental Laboratory's Surface Air Sampling Program and is attributed to an increase in solar activity.<sup>9</sup>

Samples for radiochemical analyses were collected at perimeter locations 12N and 7I (Figure 1.1) and off the site in Downers Grove (Figure 1.2). Collections were made on polystyrene filters. The total air volume filtered for the monthly samples was about 20,000 m<sup>3</sup> (700,000 ft<sup>3</sup>). Samples were ignited at 600°C (1080°F) to remove organic matter and were prepared for analysis by vigorous treatment with hot hydrochloric, hydrofluoric, and nitric acids.

Plutonium and thorium were separated on an anion exchange column, and the uranium was extracted from the column effluent. Following the extraction, the aqueous phase was analyzed for radiostrontium by a standard radiochemical procedure. The separated plutonium, thorium, and uranium fractions were electrodeposited and measured



by alpha spectrometry. The chemical recoveries were monitored by adding known amounts of plutonium-242, thorium-229, and uranium-236 tracers prior to ignition. Since alpha spectrometry cannot distinguish between plutonium-239 and plutonium-240, it should be understood that when plutonium-239 is mentioned in this report, the alpha activity due to the plutonium-240 isotope is also included. The results are given in Table 4.3.

The strontium-90 concentrations have decreased over the past several years so that during 1994 most of the results were less than the detection limit of 10 aCi/m<sup>3</sup>. Strontium-89 was not observed above the detection limit of 100 aCi/m<sup>3</sup>. The plutonium-239 concentrations are similar to last year at Location 7I, Location 12N, and at the off-site sampling location.

The thorium and uranium concentrations are in the same range found in the past and are considered to be of natural origin. The amounts of thorium and uranium in a sample were proportional to the mass of inorganic material collected on the filter paper. The bulk of these elements in the air was due to resuspension of soil.

The major airborne effluents released at ANL during 1994 are listed by location in Table 4.4 and the annual releases of the major sources since 1985 are illustrated in Figure 4.3. The radon-220 released from Building 200 is due to radioactive contamination from the "proof-of-breeding" program and from nuclear medicine studies. Even though the CP-5 reactor ceased operations in 1979, hydrogen-3 continues to be emitted from Building 330. The hydrogen-3 emitted from Building 212 is from tritium recovery studies. In addition to the nuclides listed in Table 4.4, several other fission products also were released in millicurie or smaller amounts. The quantities listed in Table 4.4 were measured by on-line stack monitors in the exhaust systems of the buildings, except for Building 350.

TABLE 4.3

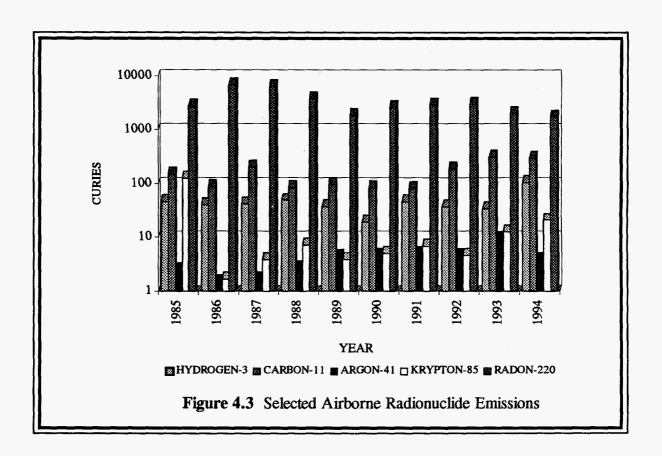
Strontium, Thorium, Uranium, and Plutonium Concentrations in Air-Filter Samples, 1994

(Concentrations in aCi/m³)

January   71							
12N   Off-Site	<10	10 ± 1	15 ± 1	7 ± 1	16 ± 2	15 ± 2	0.2 ± 0.4
Off-Site  February 7I 12N Off-Site  March 7I 12N Off-Site  April 7I 12N Off-Site  May 7I 12N Off-Site  June 7I 12N Off-Site  June 7I 12N Off-Site  July 7I 12N Off-Site  August 7I 12N Off-Site  September 7I 12N Off-Site  October 7I 12N Off-Site  November 7I 12N Off-Site  December 7I 12N Off-Site	<10	8 ± 3	7 ± 1	4 ± 1	10 ± 1	9 ± 1	$2.0 \pm 1.6$
12N   Off-Site	11 ± 15	7 ± 2	9 ± 1	5 ± 1	10 ± 1	10 ± 1	0.5 ± 0.3
12N   Off-Site	<10	11 ± 2	17 ± 1	11 ± 1	15 ± 2	16 ± 2	0.9 ± 0.3
Off-Site  March 71	<10	4 ± 1	9 ± 1	5 ± 1	11 ± 1	8 ± 1	0.7 ± 0.4
12N   Off-Site	<10	17 ± 5	23 ± 4	9 ± 2	20 ± 1	20 ± 1	0.5 ± 0.6
12N   Off-Site	<10	11 ± 6	15 ± 1	8 ± 1	15 ± 1	13 ± 1	0.4 ± 0.3
Off-Site  April 71	<10	11 ± 5	11 ± 1	7 ± 1	15 ± 1	12 ± 1	1.9 ± 0.4
12N   Off-Site	<10	4 ± 1	4 ± 1	3 ± 1	6 ± 1	4 ± 1	$0.4 \pm 0.3$
12N   Off-Site	<b>~10</b>	16 · A	17 . 2	12 . 1	19 . 2	17 ± 2	3.9 ± 0.7
Off-Site  May 71 12N Off-Site  June 71 12N Off-Site  July 71 12N Off-Site  August 71 12N Off-Site  September 71 12N Off-Site  October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N Off-Site	<10	16 ± 4	17 ± 2	12 ± 1	18 ± 2		
May 71 12N Off-Site  June 71 12N Off-Site  July 71 12N Off-Site  August 71 12N Off-Site  September 71 12N Off-Site  October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N Off-Site	<10	17 ± 7	20 ± 2	15 ± 1	18 ± 1	19 ± 1	1.5 • 0.4
12N   Off-Site	21 ± 2	4 ± 3	5 ± 1	3 ± 1	7 ± 1	5 ± 1	$0.5 \pm 0.2$
Off-Site  June 71 12N Off-Site  July 71 12N Off-Site  August 71 12N Off-Site  September 71 12N Off-Site  October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N Off-Site	<10	11 ± 3	14 ± 1	9 ± 1	17 ± 2	14 ± 2	$0.9 \pm 0.6$
June 71 12N Off-Site  July 71 12N Off-Site  August 71 12N Off-Site  September 71 12N Off-Site  October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N Off-Site	$15 \pm 2$	$12 \pm 5$	$15 \pm 1$	$9 \pm 1$	17 ± 2	16 ± 1	$1.1 \pm 0.7$
12N   Off-Site	<10	4 ± 3	5 ± 1	3 ± 1	8 ± 1	6 ± 1	$0.4 \pm 0.2$
Off-Site  July 71 12N Off-Site  August 71 12N Off-Site  September 71 12N Off-Site  October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N Off-Site	<10	12 ± 4	11 ± 2	8 ± 2	13 ± 2	11 ± 2	$0.3 \pm 0.2$
July 71 12N Off-Site  August 71 12N Off-Site  September 71 12N Off-Site  October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N Off-Site	18 ± 2	10 🏚 4	$16 \pm 2$	7 ± 1	$15 \pm 2$	15 ± 2	$1.7 \pm 0.4$
12N	<10	9 ± 6	8 ± 1	6 ± 1	10 ± 2	10 ± 2	$1.2 \pm 0.4$
12N	<10	8 ± 5	10 ± 1	6 ± 1	10 ± 1	10 ± 1	0.4 ± 0.2
Off-Site  August 71	<10	7 ± 2	8 ± 1	$4 \pm 1$	8 ± 1	7 ± 1	$0.6 \pm 0.3$
12N Off-Site  September 71	<10	8 ± 5	9 ± 1	4 ± 1	9 ± 2	8 ± 2	$0.5 \pm 0.3$
12N   Off-Site	<10	5 ± 11	6 ± 1	3 ± 1	6 ± 1	5 ± 1	0.2 ± 0.2
Off-Site   September   71   12N   Off-Site	<10	7 ± 7	6 ± 1	4 ± 1	7 ± 1	7 ± 1	$0.4 \pm 0.3$
12N   Off-Site	<10	6 ± 5	6 ± 1	3 ± 1	6 ± 1	8 ± 1	0.7 ± 0.3
12N   Off-Site	<10	8 ± 2	8 ± 1	5 ± 1	10 ± 1	10 ± 1	0.2 ± 0.2
October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N		5 ± 2				6 ± 1	0.3 ± 0.2
October 71 12N Off-Site  November 71 12N Off-Site  December 71 12N	<10			_	_		$0.3 \pm 0.2$ $0.4 \pm 0.2$
12N	<10	4 ± 2	3 ± 1.	2 ± 1	5 ± 1	4 ± 1	0.4 ± 0.2
Off-Site  November 7I 12N Off-Site  December 7I 12N	-	-	-	-	•	-	-
November 7I 12N Off-Site  December 7I 12N	-	-	. •	-	-	•	-
12N Off-Site  December π 12N	•	•	•	•	. •	•	•
Off-Site  December 71 12N	<10	7 ± 3	6 ± 1	4 ± 1	9 ± 2	11 ± 2	0.2 ± 0.4
December 71 12N	< 10	6 ± 4	$5 \pm 1$	$4 \pm 1$	7 ± 1	8 ± 1	<0.1
12N	<10	3 ± 5	2 ± 1	<1	4 ± 1	3 ± 1	$0.2 \pm 0.3$
12N			-	•	•	-	•
	-	-	-	-	•	-	-
	<10	3 ± 3	3 ± 1	1 ± 1	4 ± 1	4 ± 1	$0.2 \pm 0.3$
Annual 7I	<10	10 ± 7	12 ± 10	7 ± 7	13 ± 9	12 ± 8	0.8 ± 2.5
Summary 12N	<10	9 ± 9	10 ± 11	6 ± 8	$12 \pm 10$	11 ± 10	1.0 ± 1.6
Off-Site	<10	6 ± 9	7 ± 13	4 ± 5	8 ± 10	8 ± 11	$0.5 \pm 0.6$
Dose 7I	<(0.00004)	(0.0248)	(0.0238)	(0.073)	(0.00065)	(0.00060)	(0.0019)
(mrem) 12N	<(0.00007)	(0.0216)	(0.0207)	(0.061)	(0.00058)	(0.00054)	(0.0026)
Off-Site	<(0.00007)	(0.0156)	(0.0141)	(0.036)	(0.00041)	(0.00038)	(0.0013)

<sup>•</sup> Perimeter locations are given in terms of the grid coordinates in Figure 1.1

			Amount Released	Amount Released
Building	Nuclide	Half-Life	(Ci)	(Bq)
200	Radon-220	56 s	1750.1	$6.5 \times 10^{13}$
205	Hydrogen-3 (HTO)	12.3 y	0.39	1.4 x 10 <sup>10</sup>
.12	Hydrogen-3 (HTO)	12.3 y	17.85	6.6 x 10 <sup>11</sup>
	Hydrogen-3 (HT)	12.3 y	40.15	$1.5 \times 10^{12}$
	Krypton-85	10.7 y	20.96	$7.8 \times 10^{11}$
	Antimony-125	2.71 y	0.000022	$8.1 \times 10^5$
	Radon-220	56 s	2.6	$9.6 \times 10^{10}$
330 (CP-5)	Hydrogen-3 (HTO)	12.3 y	49.17	$1.8 \times 10^{12}$
350 (NBL)	Uranium-234	$2.4 \times 10^5 \text{ y}$	1.6 x 10 <sup>-8</sup>	$5.9 \times 10^2$
,	Uranium-238	4.5 x 10 <sup>9</sup> y	1.6 x 10 <sup>-8</sup>	$5.9 \times 10^2$
	Plutonium-238	87.7 y	2.5 x 10 <sup>-13</sup>	9.2 x 10 <sup>-3</sup>
	Plutonium-239	$2.4 \times 10^4 \text{ y}$	$7.7 \times 10^{-8}$	$2.8 \times 10^3$
	Plutonium-240	$6.6 \times 10^4 \text{ y}$	2.1 x 10 <sup>-9</sup>	$7.8 \times 10^2$
	Plutonium-241	14.4 y	$8.7 \times 10^{-10}$	$3.2 \times 10^{1}$
	Plutonium-242	$3.76 \times 10^5 \text{ y}$	$6.4 \times 10^{-15}$	$2.4 \times 10^{-4}$
375 (IPNS)	Carbon-11	20 m	305.9	1.1 x 10 <sup>13</sup>
<b>,</b> , , ,	Argon-41	1.8 h	4.4	$1.6 \times 10^{11}$
11/415 (APS)	Carbon-11	20 m	0.94	$2.4 \times 10^{10}$
( ()	Nitrogen-13	10 m	40.6	$1.5 \times 10^{12}$
	Oxygen-15	122 s	4.3	$1.6 \times 10^{11}$



#### 4.3. Surface Water

All surface water samples collected in the monitoring program were acidified to 0.1N with HNO<sub>3</sub> and filtered immediately after collection. Total nonvolatile alpha and beta activities were determined by counting the residue remaining after evaporation of the water and then applying counting efficiency corrections determined for plutonium-239 (for alpha activity) and thallium-204 (for beta activity) to obtain disintegration rates. Hydrogen-3 was measured from a separate aliquot, and this activity does not appear in the results for total nonvolatile beta activity. Analyses for the radionuclides were performed by specific radio-chemical separations followed by appropriate counting. One-liter aliquots were used for all

analyses except for hydrogen-3 and the transuranium nuclides. Hydrogen-3 analyses were performed by liquid scintillation counting of 9 mL of a distilled sample in a non-hazardous cocktail. Analyses for transuranium nuclides were performed on 10-liter samples with chemical separation methods followed by alpha spectrometry. Plutonium-236 was used to determine the yields of plutonium and neptunium, which were separated from the sample together. A group separation of a fraction containing the transplutonium elements was monitored for recovery with americium-243 tracer. Isotopic uranium concentrations were determined by alpha spectrometry using uranium-236 as an isotopic tracer.

Argonne wastewater is discharged into Sawmill Creek, which runs through the ANL grounds, drains surface water from much of the site, and flows into the Des Plaines River about 500 m (0.3 mi) downstream from the ANL wastewater outfall. Sawmill Creek was sampled upstream from the ANL site and downstream from the wastewater outfall to determine if radioactivity was added to the stream by ANL wastewater or surface drainage. The sampling locations are shown in Figure 1.1. Below the wastewater outfall, daily samples were collected. Equal portions of the daily samples collected each week were combined and analyzed to obtain an average weekly concentration. Upstream of the site, samples were collected once a month and were analyzed for the same radionuclides measured in the below-outfall samples.

Annual summaries of the results obtained for Sawmill Creek are given in Table 4.5. Comparison of the results and 95% confidence levels of the averages for the two sampling locations shows that the nuclides found in the creek water that can be attributed to ANL operations were hydrogen-3, strontium-90, cesium-137, plutonium-239, americium-241, and occasionally neptunium-237, plutonium-238, curium-242 and/or californium-252, and curium-244 and/or californium-249. The percentage of individual samples containing activity attributable to ANL was 32% for hydrogen-3, 78% for strontium-90, 26% for cesium-137, 66% for plutonium-239, and 78% for americium-241. The concentrations of all these

TABLE 4.5

Radionuclides in Sawmill Creek Water, 1994

Activity	Location*	No. of Samples	Avg.	Concentrations i Min.	n pCi/L Max.	Avg.	Dose (mrem) Min.	Max.
Alpha (Nonvolatile)	16K 7M	12 50	1.3 ± 1.3 1.7 ± 1.4	0.6 0.6	2.4 4.4	-	-	<u>-</u>
Beta	16K	12	8 ± 8	6	16	-	-	-
(Nonvolatile)	7M	50	16 ± 14	7	37	-	•	
Hydrogen-3	16K	12	< 100	< 100	114	< 0.0046	< 0.0046	0.0052
	7M	50	137 ± 606	< 100	1931	0.0063	< 0.0046	0.0886
Strontium-90	16K	12	0.26 ± 0.20	< 0.25	0.40	0.025	< 0.024	0.038
	7M	50	0.59 ± 0.99	< 0.25	2.34	0.056	< 0.024	0.222
Cesium-137	16K	12	< 1.0	< 1.0	< 1.0	< 0.04	< 0.04	< 0.04
	7M	50	1.0 ± 3.0	< 1.0	6.4	0.04	< 0.04	0.23
Uranium-234	16K	12	0.772 ± 0.767	0.441	1.291	0.147	0.084	0.245
	7M	50	0.601 ± 0.425	0.293	1.030	0.114	0.056	0.196
Uranium-238	16K	12	0.726 ± 0.679	0.380	1.139	0.122	0.064	0.191
	7M	50	0.518 ± 0.434	0.232	0.932	0.087	0.039	0.157
Neptunium-237	16K	12	< 0.0010	< 0.0010	0.0011	< 0.0028	< 0.0028	0.0031
	7M	50	< 0.0010	< 0.0010	0.0013	< 0.0028	< 0.0028	0.0038
Plutonium-238	16K	12	< 0.0010	< 0.0010	< 0.0010	< 0.0028	< 0.0028	< 0.0028
	7M	50	< 0.0010	< 0.0010	0.0015	< 0.0028	< 0.0028	0.0043
Plutonium-239	16K	12	< 0.0010	< 0.0010	0.0010	< 0.0031	< 0.0031	0.0031
	7M	50	0.0023 ± 0.004	7 < 0.0010	0.0106	0.0071	< 0.0031	0.0331
Americium-241	16K 7M	12 50	< 0.0010 0.0034 ± 0.0074	< 0.0010	0.0010 0.0155	< 0.0033 0.0110	< 0.0033 < 0.0033	0.0034 0.0508
Curium-242 and/or	· 16K	12	< 0.0010	< 0.0010	0.0011	< 0.0007	< 0.0007	0.0007
Californium-252	7M	50	< 0.0010	< 0.0010	0.0010	< 0.0007	< 0.0007	0.0007
Curium-244 and/or	16K	12	< 0.0010	< 0.0010	0.0015	< 0.0034	< 0.0034	0.0050
Californium-249	7M	50	< 0.0010	< 0.0010	0.0060	< 0.0034	< 0.0034	0.0200

<sup>\*</sup> Location 16K is upstream from the Argonne site and location 7M is downstream from the Argonne wastewater outfall.

nuclides were low and a small fraction of the allowed DOE limits. If the concentrations of the radionuclides listed in Table 4.5 were increased by a factor of five, which approximates the effect of the dilution by Sawmill Creek on the ANL effluent water, the concentrations would still be below the DOE limits. This demonstrates compliance with DOE Order 5400.5 for use of Best Available Technology (BAT) for release of liquid effluents.

Liquid wastewater from buildings or facilities that use or process radioactive materials are collected in retention tanks. When a tank is full, it is sampled and analyzed for alpha and beta radioactivity. If the radioactivity exceeds the release limits, the tank is processed by evaporation and the residue disposed of as solid low-level radioactive waste. If the radioactivity is below the release limits, the wastewater is conveyed to the Laboratory wastewater treatment plant in dedicated pipes to waste storage tanks. These tanks are again sampled and analyzed for radioactivity and if below the release limits, discharged to the environment. The release limits are based on the DCGs of plutonium-239 (0.03 pCi/mL) for alpha activity and for strontium-90 (1.0 pCi/mL) for beta activity. These radionuclides were selected because of their potential for release and their conservative allowable limits in the environment. This effluent monitoring program documents that no liquid releases above the DCGs have occurred and reinforces the demonstration of compliance with the use of BAT as required by DOE Order 5400.5.

At location 7M, below the ANL outfall, the annual average concentrations of most measured radionuclides were similar to recent annual averages. All the annual averages were well below the applicable standards. The annual total radioactive effluent discharged to the creek in ANL wastewater can be estimated from the average net concentrations and the volume of water carried by the creek. These totals are collected in Table 4.6.

TABLE 4.6

Total Radioactivity Released to Sawmill Creek, 1994

Released (Ci)	Percent	
1.14	99.0	
0.0033	0.3	
0.0083	0.7	
0.000013	< 0.1	
0.000024	< 0.1	
1.15		
	1.14 0.0033 0.0083 0.000013 0.000024	

Based on the results of the Stormwater Characterization Study (see Section 2.2.2), two perimeter surface water locations were identified which contained measurable levels of radionuclides. There were south of the 319 Area, Location 7J, and from the 800 Area Landfill, Location 11D in Figure 1.1. Samples were scheduled to be collected quarterly and analyzed for hydrogen-3, strontium-90, and by gamma-ray spectrometry. The locations were dry during the third quarter. The results are collected below:

## (Concentrations in pCi/L)

Date Collected	Location 7J Hydrogen-3	Location 7J Strontium-90	Location 7J Cesium-137	Location 11D Hydrogen-3
February 18, 1994	842	0.6	< 1	< 100
April 13, 1994	7632	1.4	< 1	167
October 31, 1994	< 100	1.7	< 1	< 100

The source of the radionuclides at Location 7J appears to be leachate from the 319 Area Landfill. Interim actions are being planned to collect and divert or treat the leachate. The hydrogen-3 at Location 11D is probably also from the leachate and the decrease in the concentration from earlier years is due to the completion of the clay cap on the Landfill in the fall of 1993.

Because Sawmill Creek empties into the Des Plaines River, which in turn flows into the Illinois River, data on the radioactivity in the two rivers are important in assessing the contribution of ANL wastewater to environmental radioactivity. The Des Plaines River was sampled twice a month below, and once a month above, the mouth of Sawmill Creek to determine if the radioactivity in the creek had any effect on the radioactivity in the river.

Table 4.7 presents annual summaries of the results obtained for these two locations. The average nonvolatile alpha, beta, and uranium concentrations in the river were very similar to past averages and remained in the normal range. Results were quite similar above and below the creek for all radionuclides, because the activity in Sawmill Creek was reduced by dilution to the point that it was not detectable in the Des Plaines River. The average nonvolatile alpha and beta activities, 1.5 pCi/L and 12.1 pCi/L, respectively, of 26 off-site surface water samples collected in 1994 were similar to the levels found in previous years. The hydrogen-3 concentration in these surface water samples averaged 56 pCi/L.

The radioactivity levels in samples of Illinois River water, shown in Table 4.8, were similar to those found previously at these same locations. No radioactivity originating at ANL could be detected in the Des Plaines or Illinois rivers. The elevated hydrogen-3 levels appear to be due to discharges from the Dresden nuclear power station complex.

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TABLE 4.7

Radionuclides in Des Plaines River Water, 1994

	No. of Concentrations in pCi/L Dose (mrem)  Location* Samples Avg. Min. Max. Avg. Min. M							
Activity	Location*	Samples	Avg.	Min.	Max.	Avg.	Min.	Max.
Alpha	A	12	1.3 ± 0.9	0.6	2.1	-	•	· •
(Nonvolatile)	В	24	$1.3 \pm 1.4$	0.1	3.0	•	-	-
Beta	A	12	16 ± 11	9	25	•	-	•
(Nonvolatile)	В	24	17 ± 12	7	30	•	•	•
Hydrogen-3	A	12	< 100	< 100	< 100	< 0.0046	< 0.0046	< 0.0046
	В	24	< 100	< 100	< 100	< 0.0046	< 0.0046	< 0.0046
Strontium-90	A	12	$0.32 \pm 0.13$	5 < 0.25	0.46	0.031	< 0.024	0.044
	В	24	$0.36 \pm 0.22$	2 < 0.25	0.59	0.034	< 0.024	0.056
Uranium-234	Ä	12	0.490 ± 0.4	457 0.216	0.896	0.093	0.041	0.170
	В	24	0,498 ± 0.2	374 0.138	0.816	0.095	0.026	0.155
Uranium-238	A	12	0.404 ± 0.2		0.750	0.068	0.031	0.126
	В	24	$0.412 \pm 0.3$	310 0.115	0.710	0.069	0.019	0.119
Neptunium-237	A	12	< 0.0010	< 0.0010	0.0023	< 0.0028	< 0.0028	0.0064
	В	12	< 0.0010	< 0.0010	< 0.0010	< 0.0028	< 0.0028	< 0.0028
Plutonium-238	A	12	< 0.0010	< 0.0010	< 0.0010	< 0.0028	< 0.0028	< 0.0028
•	В	12	< 0.0010	< 0.0010	< 0.0010	< 0.0028	< 0.0028	< 0.0028
Plutonium-239	A	12	< 0.0010	< 0.0010	< 0.0010	< 0.0031	< 0.0031	< 0.0031
	В	12	< 0.0010	< 0.0010	< 0.0010	< 0.0031	< 0.0031	< 0.0031
Americium-241	A	12	< 0.0010	< 0.0010	0.0036	< 0.0033	< 0.0033	0.0119
	В	12	< 0.0010	< 0.0010	< 0.0010	< 0.0033	< 0.0033	< 0.0033
Curium-242 and/o	or A	12	< 0.0010	< 0.0010	< 0.0010	< 0.0007	< 0.0007	< 0.0007
Californium-252	В	12	< 0.0010	< 0.0010	< 0.0010	< 0.0007	< 0.0007	< 0.0007
Curium-244 and/o	or A	12	< 0.0010	< 0.0010	< 0.0010	< 0.0034	< 0.0034	< 0.0034
Californium-249	В	12	< 0.0010	< 0.0010	0.0013	< 0.0034	< 0.0034	0.0043

<sup>\*</sup> Location A, near Willow Springs, is upstream and location B, near Lemont, is downstream from the mouth of Sawmill Creek. See Figure 1.2

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TABLE 4.8

Radionuclides in Illinois River Water, 1994

Concentrations in pCi/L

Date Collected	Location	Alpha*	Beta*	Hydrogen-3	Uranium-234	Uranium-238	Plutonium-239
April 21	McKinley Woods Park, IL	1.6 ± 0.4	12.6 ± 0.4	< 100	0.52 ± 0.04	0.39 ± 0.04	0.006 ± 0.001
April 21	Dresden Lock & Dam, IL	$3.2 \pm 0.4$	$8.1 \pm 0.3$	186 ± 48	0.78 ± 0.09	0.61 ± 0.07	< 0.001
April 21	Morris, IL	$1.5 \pm 0.3$	$7.0 \pm 0.3$	129 ± 47	$0.55~\pm~0.04$	$0.47 \pm 0.04$	-
April 21	Starved Rock State Park, IL	1.0 ± 0.2	$6.5 \pm 0.3$	< 100	$0.60 \pm 0.07$	0.49 ± 0.06	-
October 26	McKinley Woods Park, IL	$0.6 \pm 0.3$	11.7 ± 0.3	< 100	$0.15 \pm 0.05$	0.11 ± 0.04	< 0.001
October 26	Dresden Lock & Dam, IL	$0.4 \pm 0.2$	$10.5 \pm 0.3$	113 ± 49	$0.23 \pm 0.07$	$0.16 \pm 0.05$	< 0.001
October 26	Dresden Lock & Dam, IL	$0.5 \pm 0.2$	$10.0 \pm 0.3$	105 ± 45	$0.20 \pm 0.04$	$0.18 \pm 0.03$	< 0.001
October 26	Morris, IL	$0.7 \pm 0.3$	$10.4 \pm 0.3$	163 ± 46	$0.31 \pm 0.07$	$0.20 \pm 0.05$	-
October 26	Starved Rock State Park, IL	0.7 ± 0.3	10.6 ± 0.4	231 ± 47	0.30 ± 0.07	0.26 ± 0.06	-

<sup>\*</sup>Nonvolatile activity.

## 4.4. Soil, Grass, and Bottom Sediment

The radioactive content of soil, grass, and bottom sediment was measured at the site perimeter and off the site. The purpose of the off-site sampling was to measure deposition for comparison with perimeter samples and with results obtained by other organizations for samples collected at large distances from nuclear installations. Such comparisons are useful in determining if the radioactivity of soil near ANL is normal. For this purpose, site-selection criteria and sample collection and sample preparation techniques recommended by the American Society for Testing and Materials (ASTM) were used. Sites were selected in several directions and at various distances from ANL. Each site was selected on the basis that the soil appeared, or was known to have been, undisturbed for a number of years. Attempts were made to select open, level, grassy areas that were mowed at reasonable intervals. Public parks were selected when available.

As part of the quality assurance program, replicate samples are taken from ten percent of the locations. The EMS data management system has been programmed to schedule the replicate samples on a rotating basis. The following tables will show paired results from the same location. Comparison of the analytical data in these tables of pairs of samples collected at the same location will provide a measure of the heterogeneity of the media, i.e., soil, grass, or bottom sediment.

Each soil sample consisted of ten cores, totaling 864 cm<sup>2</sup> (134 in<sup>2</sup>) in area by 5 cm (2 in) deep. Through 1976, samples had been collected down to 30 cm (12 in) to measure total deposition. The results of five years of sample collection at this depth has established the total deposition in the ANL environment. Reducing the sampling depth to 5 cm (2 in) will make the analysis more sensitive to changes in current deposition. The grass samples were obtained by collecting the grass from a 1 m<sup>2</sup> (10 ft<sup>2</sup>) area in the immediate vicinity of

a soil sample. A grab sample technique was used to obtain bottom sediments from water bodies. After drying, grinding, and mixing, 100 g portions of each soil, bottom sediment, and grass samples were analyzed by the same methods described in Section 4.2 for air-filter residues. The plutonium and americium were separated from the same 10 g aliquot of soil. Results are given in terms of the oven-dried (110°C) weight.

The results for the gamma-ray emitting nuclides in soil are presented in Table 4.9. Intermediate half-life fission products reported in 1986 have decayed to below their detection limits and no evidence of Chernobyl fallout is apparent. The cesium-137 levels are similar to those found over the past several years and represent an accumulation from nuclear tests over a period of many years. The annual average concentrations for the perimeter and off-site samples were similar. The plutonium and americium concentrations are given in Table 4.10. The range and average concentrations of plutonium and americium in soil were similar at both perimeter and off-site sampling points. For fallout americium-241 in soil, about 10% is due to direct deposition, while about 90% is from the decay of the previously deposited plutonium-241. The americium-241/plutonium-239 ratio is consistent with the current estimated value for this ratio of 0.43 in fallout derived material.<sup>11</sup>

The radionuclide concentrations measured in grass are listed in Table 4.11. The annual averages and concentration ranges were similar at the perimeter and off-site locations and were similar to those of previous years, indicating no contribution from ANL operations. In terms of deposition, the plutonium-239 concentration was a factor of about 10<sup>4</sup> less in the grass than in the soil from the same location.

Results of analyses of bottom sediment samples for gamma-ray emitters and transuranics are given in Table 4.12. The annual off-site averages were in the same range found in off-site samples collected in previous years. Plutonium results varied widely among locations and were strongly dependent on the retentiveness of the bottom material.

TABLE 4.9

Gamma-Ray Emitting Radionuclides in Soil, 1994
(Concentrations in pCi/g)

Collected	Location	Potassium-40	Cesium-137	Radium-226	Thorium-228	Thorium-232
	Perimeter*					
April 20	12C	19.45 ± 0.70	$0.65 \pm 0.04$	$1.36 \pm 0.07$	$1.37 \pm 0.05$	$0.96 \pm 0.10$
April 20	12D	$18.58 \pm 0.70$	$0.28 \pm 0.03$	$1.15 \pm 0.06$	$1.03 \pm 0.04$	$0.84 \pm 0.09$
April 20	14L	$18.82 \pm 0.68$	$0.38 \pm 0.03$	$1.17 \pm 0.06$	$1.18 \pm 0.04$	$0.87 \pm 0.05$
April 20	5D	$20.19 \pm 0.70$	$0.55 \pm 0.03$	$1.12 \pm 0.06$	$1.17 \pm 0.04$	$0.97 \pm 0.09$
April 20	8N	$16.80 \pm 0.57$	$0.72 \pm 0.03$	$1.03 \pm 0.06$	$1.01 \pm 0.04$	$0.72 \pm 0.08$
April 20	12 <b>D</b>	$18.41 \pm 0.66$	$0.23 \pm 0.03$	$1.22 \pm 0.07$	$1.16 \pm 0.05$	$0.80 \pm 0.09$
October 24	10 <b>E</b>	$18.79 \pm 0.68$	$0.64 \pm 0.03$	$1.02 \pm 0.06$	$1.15 \pm 0.04$	$0.87 \pm 0.09$
October 24	10N	$16.59 \pm 0.64$	$0.38 \pm 0.03$	$0.83 \pm 0.05$	$0.83 \pm 0.04$	$0.75 \pm 0.09$
October 24	14 <b>I</b>	$18.91 \pm 0.68$	$0.66 \pm 0.03$	$1.03 \pm 0.06$	$1.19 \pm 0.04$	$0.94 \pm 0.09$
October 24	6J	19.71 ± 0.69	$0.63 \pm 0.03$	$0.95 \pm 0.06$	$1.20 \pm 0.04$	$0.90 \pm 0.09$
October 24	9 <b>N</b>	$17.86 \pm 0.66$	$0.51 \pm 0.03$	$0.75 \pm 0.05$	$0.89 \pm 0.04$	$0.84 \pm 0.09$
October 24	10 <b>N</b>	$19.30 \pm 0.69$	$0.20 \pm 0.02$	$1.24 \pm 0.06$	$1.11 \pm 0.04$	$0.86 \pm 0.09$
	Average	$18.62 \pm 2.39$	$0.49 \pm 0.40$	$1.07 \pm 0.39$	$1.11 \pm 0.32$	$0.86 \pm 0.17$
	Off-site					
April 21	Dresden Lock & Dam, IL	$20.74 \pm 0.72$	$0.19 \pm 0.02$	$1.49 \pm 0.07$	$1.25 \pm 0.04$	$0.98 \pm 0.10$
April 21	McKinley Woods Park, IL	$20.15 \pm 0.65$	$0.49 \pm 0.03$	$0.93 \pm 0.06$	$1.13 \pm 0.05$	$0.79 \pm 0.09$
April 21	Morris, IL	$14.63 \pm 0.64$	$0.20 \pm 0.02$	$0.73 \pm 0.06$	$0.84 \pm 0.04$	$0.72 \pm 0.09$
April 21	Morris, IL	$13.16 \pm 0.58$	$0.48 \pm 0.03$	$0.87 \pm 0.06$	$0.83 \pm 0.04$	$0.57 \pm 0.08$
April 25	McCormick Woods, Brookfield, IL	20.70 ± 0.70	$0.23 \pm 0.03$	$1.26 \pm 0.07$	1.12 ± 0.04	$0.89 \pm 0.09$
April 25	Bemis Woods, Western Springs, IL	20.56 ± 0.62	$0.42 \pm 0.03$	$1.55 \pm 0.07$	$1.30 \pm 0.07$	$0.81 \pm 0.09$
April 25	McCormick Woods, Brookfield, IL	$19.83 \pm 0.70$	$0.26 \pm 0.03$	$1.39~\pm~0.07$	$1.05 \pm 0.04$	$0.88 \pm 0.09$
October 25	Orland Park, IL	$23.23 \pm 0.66$	$0.57 \pm 0.03$	$1.82 \pm 0.07$	$1.26 \pm 0.04$	$0.91 \pm 0.09$
October 25	Palos Hills, IL	$19.50 \pm 0.70$	$0.21 \pm 0.02$	1.52 ± 0.07	$1.23 \pm 0.05$	$0.93 \pm 0.09$
October 25	Orland Park, IL	$23.92 \pm 0.76$	$0.51 \pm 0.03$	$1.67 \pm 0.07$	$1.18 \pm 0.05$	$0.87 \pm 0.10$
October 26	Channahon, IL	$17.84 \pm 0.66$	$0.23~\pm~0.03$	$1.27 \pm 0.06$	$1.42 \pm 0.05$	$0.93 \pm 0.09$
October 26	Starved Rock State Park, IL	14.67 ± 0.55	$0.29 \pm 0.02$	$0.82 \pm 0.05$	$0.80 \pm 0.03$	$0.64 \pm 0.07$
October 26	Channahon, IL	$19.06 \pm 0.69$	$0.25~\pm~0.03$	$0.53~\pm~0.04$	$0.90 \pm 0.04$	$0.63 \pm 0.07$
	Average	19.03 ± 7.01	0.33 ± 0.31	1.22 ± 0.88	1.10 ± 0.44	0.81 ± 0.29

<sup>\*</sup> The perimeter locations are given in terms of the grid coordinates in Figure 1.1

TABLE 4.10

Transuranics in Soil, 1994

Date Collected	Location	Plutonium-238 (fCi/g)	Plutonium-238 (nCi/m²)	Plutonium-239 (fCi/g)	Plutonium-239 (nCi/m²)	Pu-238/Pu-239	Americium-241 (fCi/g)	Americium-241 (nCi/m²)	Am-241/Pu-239
	Perimeter*			- "					
April 20	12C	$0.6 \pm 0.4$	$0.020 \pm 0.012$	$15.8 \pm 1.8$	$0.538 \pm 0.063$	0.038	$5.6 \pm 1.5$	$0.190 \pm 0.052$	0.354
April 20	12D	$0.3 \pm 0.3$	$0.011 \pm 0.014$	$7.0 \pm 1.4$	$0.304 \pm 0.061$	0.038	$2.9 \pm 0.8$	$0.125 \pm 0.037$	0.410
April 20	14L	< 0.1	< 0.001	$9.8 \pm 1.5$	$0.281 \pm 0.044$	0.001	$4.2 \pm 1.2$	$0.119 \pm 0.036$	0.424
April 20	5D ·	$1.5 \pm 1.4$	$0.058 \pm 0.056$	$17.4 \pm 4.7$	$0.666 \pm 0.179$	0.087	$5.5 \pm 1.2$	$0.213 \pm 0.046$	0.320
April 20	8N	$0.7 \pm 0.5$	$0.023 \pm 0.016$	$15.6 \pm 2.4$	$0.504 \pm 0.076$	0.046	$6.4 \pm 0.1$	$0.207 \pm 0.004$	0.411
April 20	12D	$0.3 \pm 0.2$	$0.010 \pm 0.008$	$7.3 \pm 1.0$	$0.242 \pm 0.033$	0.042	$2.9 \pm 0.9$	$0.098 \pm 0.030$	0.406
October 24	10E	$0.6 \pm 0.4$	$0.029 \pm 0.017$	$17.6 \pm 2.0$	$0.836 \pm 0.096$	0.034	$5.4 \pm 1.1$	$0.259 \pm 0.055$	0.310
October 24	10N	$0.4 \pm 0.3$	$0.012 \pm 0.010$	$13.5 \pm 1.9$	$0.398 \pm 0.057$	0.031	$3.8 \pm 0.8$	$0.112 \pm 0.023$	0.282
October 24	141	$0.5 \pm 0.3$	$0.021 \pm 0.013$	$14.5 \pm 1.8$	$0.610 \pm 0.075$	0.034	$4.8 \pm 1.1$	$0.200 \pm 0.048$	0.328
October 24	63	$0.4 \pm 0.3$	$0.016 \pm 0.012$	$18.1 \pm 2.3$	$0.720 \pm 0.091$	0.022	$6.1 \pm 1.0$	$0.244 \pm 0.039$	0.339
October 24	9N	$0.5 \pm 0.4$	$0.013 \pm 0.010$	$14.9 \pm 2.1$	$0.392 \pm 0.054$	0.033	$5.0 \pm 0.9$	$0.131 \pm 0.023$	0.334
October 24	10N	$0.3 \pm 0.3$	$0.008 \pm 0.007$	$6.7 \pm 1.2$	$0.186 \pm 0.033$	0.045	$1.8 \pm 0.7$	$0.050 \pm 0.020$	0.267
	Average	$0.5 \pm 0.2$	$0.018 \pm 0.009$	$13.2 \pm 2.7$	0.473 ± 0.131	0.037	$4.5\pm0.9$	$0.162 \pm 0.041$	0.349
	Off-nite			•					
April 21	Dresden Lock & Dam, IL	$0.3 \pm 0.3$	$0.011 \pm 0.012$	$6.6 \pm 1.4$	$0.290 \pm 0.061$	0.039	$1.6 \pm 0.6$	0.072 ± 0.028	0.247
April 21	McKinley Woods Park, IL	$0.5 \pm 0.5$	0.019 ± 0.017	$12.7 \pm 2.3$	$0.473 \pm 0.087$	0.040	5.1 ± 0.9	$0.189 \pm 0.035$	0.400
April 21	Morris IL	$0.4 \pm 0.4$	$0.023 \pm 0.022$	$4.0 \pm 1.1$	$0.251 \pm 0.068$	0.090	$3.3 \pm 1.0$	$0.206 \pm 0.060$	0.821
April 21	Morris, IL	$0.1 \pm 0.9$	$0.007 \pm 0.048$	$12.1 \pm 1.9$	$0.658 \pm 0.102$	0.010	$6.0 \pm 1.6$	$0.327 \pm 0.089$	0.497
April 25	McCormick Woods, Brookfield, IL	< 0.1	$0.002 \pm 0.004$	$6.1 \pm 1.2$	$0.181 \pm 0.036$	0.010	$2.4 \pm 0.8$	0.071 ± 0.025	0.396
April 25	Bemis Woods, Western Springs, IL	$0.3\pm0.2$	$0.009 \pm 0.006$	$9.3 \pm 1.1$	$0.279 \pm 0.034$	0.032	$4.0 \pm 0.9$	$0.120 \pm 0.028$	0.429
April 25	McCormick Woods, Brookfield, IL	$0.2\pm0.3$	$0.006 \pm 0.009$	8.0 ± 1.2	$0.221 \pm 0.032$	0.027	$2.6\pm0.7$	0.073 ± 0.019	0.329
October 25	Orland Park, IL	$0.6 \pm 0.3$	$0.021 \pm 0.012$	15.4 ± 1.9	$0.516 \pm 0.062$	0.040	$6.5 \pm 1.7$	$0.217 \pm 0.056$	0.421
October 25	Palos Hills, IL	$0.3 \pm 0.2$	$0.008 \pm 0.007$	$5.5 \pm 1.0$	$0.169 \pm 0.031$	0.050	$2.0 \pm 0.7$	$0.061 \pm 0.021$	0.358
October 25	Orland Park, IL	$0.9 \pm 0.4$	$0.031 \pm 0.015$	14.6 ± 1.8	$0.515 \pm 0.063$	0.059	4.5 ± 1.2	$0.158 \pm 0.042$	0.307
October 26	Channahon, IL	$0.3 \pm 0.3$	$0.009 \pm 0.007$	$6.0 \pm 1.1$	$0.153 \pm 0.028$	0.056	1.8 ± 0.7	$0.045 \pm 0.018$	0.294
October 26	Starved Rock State Park, IL	$0.3 \pm 0.3$	$0.013 \pm 0.011$	$6.3 \pm 1.2$	0.277 ± 0.051	0.048	$2.3 \pm 0.7$	$0.100 \pm 0.033$	0.362
October 26	Channahon, IL	$0.2\pm0.2$	$0.006 \pm 0.006$	$6.1\pm1.1$	$0.213 \pm 0.037$	0.027	$1.9 \pm 0.6$	$0.065 \pm 0.021$	0.306
	Average	0.3 ± 0.1	0.013 ± 0.005	8.7 ± 2.3	0.323 ± 0.098	0.041	$3.4 \pm 1.0$	0.131 ± 0.051	0.398

<sup>\*</sup> The perimeter locations are given in terms of grid coordinates in Figure 1.1

TABLE 4.11

Radionuclides in Grass, 1994

Date Collected	Location	Potassium-40 (pCi/g)	Cesium-137 (fCi/g)	Plutonium-239 (fCi/g)	Deposited Plutonium-239 (pCi/m2)
-	Perimeter*				
April 20	12C	15.57 ± 0.54	29 ± 16	$0.4 \pm 0.1$	0.04 ± 0.01
April 20	12D	$10.38 \pm 0.40$	$15 \pm 13$	$0.4 \pm 0.1$	$0.05 \pm 0.02$
April 20	14L	$14.81 \pm 0.54$	$23 \pm 21$	$0.2 \pm 0.1$	$0.02 \pm 0.01$
April 20	5D	$18.59 \pm 0.61$	< 10	$0.2 \pm 0.1$	$0.02 \pm 0.01$
April 20	8N	10.81 ± 0.44	< 10	< 0.1	< 0.01
April 20	12D	19.88 ± 0.56	< 10	$0.2 \pm 0.1$	$0.01 \pm 0.01$
October 24	10E	$25.16 \pm 0.60$	< 10	< 0.1	< 0.01
October 24	141	$26.95 \pm 0.56$	$23 \pm 16$	$0.2 \pm 0.1$	$0.02 \pm 0.01$
October 24	6J	$17.77 \pm 0.51$	< 10	$0.4 \pm 0.1$	$0.08 \pm 0.02$
October 25	10N	$21.02 \pm 0.51$	< 10	< 0.1	< 0.01
October 25	9N	$17.48 \pm 0.51$	< 10	< 0.1	$0.01 \pm 0.01$
October 25	10N	$23.56 \pm 0.52$	< 10	< 0.1	< 0.01
	Average	18.50 ± 1.00	< 10	0.2 ± 0.1	0.02 ± 0.0
	Off-site				
April 21	Dresden Lock & Dam, IL	35.69 ± 0.84	< 10	$0.1 \pm 0.1$	< 0.01
April 21	McKinley Woods Park, IL	$11.57 \pm 0.36$	$14 \pm 10$	$0.6 \pm 0.1$	$0.09 \pm 0.03$
April 21	Morris, IL	$7.50 \pm 0.39$	< 10	$0.1 \pm 0.1$	$0.02 \pm 0.0$
April 21	Morris, IL	$13.41 \pm 0.45$	$10 \pm 13$	$0.4 \pm 0.1$	$0.04 \pm 0.0$
April 25	McCormick Woods, Brookfield, IL	$40.56 \pm 0.87$	16 ± 19	$0.3 \pm 0.1$	$0.02 \pm 0.0$
April 25	Bemis Woods, Western Springs, IL	23.28 ± 0.55	$26 \pm 14$	$0.7\pm0.1$	$0.08 \pm 0.0$
April 25	McCormick Woods, Brookfield, IL	64.44 ± 1.16	< 10	$0.2\pm0.1$	$0.02 \pm 0.0$
October 25	Orland Park, IL	21.31 ± 0.54	< 10	$0.1 \pm 0.1$	$0.02 \pm 0.0$
October 25	Palos Hills, IL	$12.75 \pm 0.45$	< 10	$0.1 \pm 0.1$	$0.01 \pm 0.0$
October 25	Orland Park, IL	$25.51 \pm 0.55$	< 10	< 0.1	< 0.01
October 26	Channahon, IL	$21.00 \pm 0.56$	< 10	< 0.1	< 0.01
October 26	Starved Rock State	25.60 ± 0.55	18 ± 16	< 0.1	$0.01 \pm 0.0$
O000001 20	Park, IL	_	-		
October 26	Channahon, IL	$22.37 \pm 0.57$	< 10	< 0.1	< 0.01
	Average	25.00 ± 2.63	< 10	$0.2\pm0.1$	$0.03 \pm 0.0$

<sup>\*</sup> The perimeter locations are given in terms of the grid coordinates in Figure 1.1

Radionuclides in Bottom Sediment, 1994

**TABLE 4.12** 

Date			Concentrations in pCi/g						Concentrations in fCi/g				
Collected	Location	Potassium-40	C	sium-137	Radio	um-226	Thori	um-228	Thorium-232	Plutonium-238	Plutonium-239	Ameri	icium-241
	Perimeter*								,				
July 21	Sawmill Creek 25 m Above Outfall	10.17 ± 0.	52 0.0	3 ± 0.02	0.59	± 0.05	0.45	± 0.04	0.33 ± 0.07	$0.2 \pm 0.2$	1.9 ± 0.8	1.1	± 0.5
July 21	Sawmill Creek At Outfall	2.93 ± 0.3	23 0.49	9 ± 0.03	0.22	± 0.03	0.15	± 0.02	0.15 ± 0.04	$2.0 \pm 0.7$	18.7 ± 2.3	8.1	± 1.3
July 21	Sawmill Creek 50 m Below Outfall	7.82 ± 0.4	7 0.4	4 ± 0.03	0.47	± 0.05	0.36	± 0.03	0.30 ± 0.07	$0.6 \pm 0.5$	$7.6 \pm 1.4$	3.3	± 0.8
July 21	Sawmill Creek 100 m Below Outfall	10.22 ± 0.4	32 0.4	7 ± 0.03	0.50	± 0.04	0.38	± 0.03	0.31 ± 0.06	0.3 ± 0.4	$7.4 \pm 1.5$	2.5	± 0.7
July 21	Sawmill Creek At Des Plaines River	10.24 ± 0.5	3 0,3	7 ± 0.03	0.62	± 0.05	0.55	± 0.04	0.40 ± 0.08	0.5 ± 0.5	15.7 ± 2.3	4.9	± 1.1
	Off-Site		•										
April 21	Illinois River Dresden Lock & Dam, IL	16.86 ± 0.6	5 0.19	£ 0.03	0.80	± 0.06	0.72	± 0.04	0.56 ± 0.09	< 0.1	4.4 ± 1.1	2.0	± 0.7
April 21	Illinois River McKinley Woods Park, IL	9.03 ± 0.4	9 0.02	£ 0.02	0.39	± 0.04	0.32	± 0.03	0.25 ± 0.07	0.1 ± 0.2	$0.5 \pm 0.4$		± 0.5
April 21	Illinois River Morris, IL	10.08 ± 0.4	0 0.04	± 0.02	0.29	± 0.03	0.34	± 0.03	0.31 ± 0.05	0.1 ± 0.2	$0.5 \pm 0.4$	3.1	± 1.0
April 21	Illinois River Morris, IL	11.08 ± 0.4	3 0.03	± 0.02	0.33	± 0.04	0.33	± 0.03	0.27 ± 0.05	0.2 ± 0.2	1.6 ± 0.5	0.8	± 0.4
April 25	Des Plaines River McCormick Woods, Brookfield, IL	18.59 ± 0.6	8 0.32	£ ± 0.03	1.26	± 0.07	1.16	± 0.05	0.85 ± 0.10	0.4 ± 0.4	8.9 ± 1.3	3.4	± 1.0
April 25	Salt Creek  Bemis Woods,  Western Springs, IL	12.81 ± 0.6	1 0.04	± 0.02	1.39	± 0.07	0.67	± 0.04	0.64 ± 0.09	< 0.1	1.4 ± 0.6	1.5	± 0.8
October 25	McGinnis Slough Orland Park, IL	2.20 ± 0.7	0 0.41	± 0.04	1.45	± 0.07	1.17	± 0.05	0.91 ± 0.10	0.4 ± 0.3	9.1 ± 1.3	3.3	± 0.7
October 25	Saganashkee Slough Palos Hills, IL	12.34 ± 0.5	2 0.09	± 0.02	0.64	± 0.05	0.52	± 0.03	0.42 ± 0.07	< 0.1	3.5 ± 1.0	1.6	± 0.6
October 25	Des Plaines River Romeoville, IL	18.63 ± 0.6	9 0.40	± 0.03	1.49	± 0.07	1.26	± 0.05	0.85 ± 0.10	0.4 ± 0.3	12.2 ± 1.5	3.7	± 1.0
October 26	DuPage River Channahon, IL	10.79 ± 0.5	3 0.01	± 0.02	0.55	± 0.05	0.80	± 0.04	0.72 ± 0.08	$0.1 \pm 0.1$	$0.8 \pm 0.4$	0.7	± 0.4
October 26	Illinois River Starved Rock State Park, IL	5.27 ± 0.3	0.02	± 0.02	0.46	± 0.04	0.26	± 0.03	0.18 ± 0.05	< 0.1	0.6 ± 0.3	0.4	± 0.3
October 26	Illinois River Starved Rock State Park, IL	5.41 ± 0.3	2 0.02	± 0.02	0.55	± 0.04	0.27	± 0.03	0.25 ± 0.05	< 0.1	0.7 ± 0.4	0.6	± 0.4
	Average	11.09 ± 11.5	3 0.13	± 0.35	0.80	± 1.03	0.65	± 0.83	0.52 ± 0.60	0.1 ± 0.4	3.7 ± 9.0	1.9	± 2.7

<sup>•</sup> The perimeter locations are given in terms of the grid coordinates in Figure 1.1

A set of sediment samples was collected on July 21, 1994, from the Sawmill Creek bed, above, at the outfall, and at several locations below the point at which ANL discharges its treated wastewater (location 7M in Figure 1.1). The results, as listed in Table 4.12, show that the concentrations in the sample above the 7M outfall are similar to those of the off-site samples. The plutonium, americium, and cesium-137 concentrations are elevated below the outfall, indicating that their origin is in ANL wastewater. The changes in concentrations of these nuclides with time and location indicate the dynamic nature of the sediment material in this area.

## 4.5. External Penetrating Radiation

Levels of external penetrating radiation at and in the vicinity of the ANL site were measured with calcium fluoride thermoluminescent dosimeter (TLD) chips. Each measurement reported represents the average of four chips exposed in the same packet. All calcium fluoride packets were shielded with 1.6 mm (1/16 in) copper foil to reduce or eliminate the beta and low-energy X-ray components. The response of the chips was determined with a U. S. National Institute of Standards and Technology (NIST) standard radium-226 source, and the results were calculated in terms of the air dose. Dosimeters were exposed at several locations at the site boundary and on the site. Readings were also taken at five off-site locations for comparison purposes. These locations are shown in Figure 1.2.

The results are summarized in Tables 4.13 and 4.14, and the site boundary and on-site readings are also shown in Figure 4.4. Measurements were made for the four successive exposure periods shown in the tables, and the results were calculated in terms of annual dose for ease in comparing measurements made for different elapsed times. The uncertainty given in the tables for an average is the 95% confidence limit calculated from the standard deviation of the average.

The off-site results averaged  $82 \pm 10$  mrem/y and were similar to last year's off-site average of  $76 \pm 6$  mrem/y. To compare boundary results for individual sampling periods, the standard deviation of the 20 individual off-site results is useful. This value is 9 mrem/y, so individual results in the range of  $82 \pm 19$  mrem/y may be considered to be the average natural background with a 95% probability.

TABLE 4.13

Environmental Penetrating Radiation at Off-Site Locations, 1994

		D	ose Rate (mren	n/year)	
Location	1/7-4/7	4/7-7/7	7/7-10/6	10/6-1/5	Average
Lemont	81	125	77	73	89 ± 24
Oak Brook	91	89	63	133	$94 \pm 28$
Orland Park	81	73	98	83	$84 \pm 10$
Woodridge	74	79	111	65	$82 \pm 20$
Willow Springs	62	70	61	43	$59 \pm 11$
Average	78 ± 9	87 ± 20	82 ± 19	79 ± 29	82 ± 10

In the past, two site boundary locations, 7I (south) and 14I (north), the dose rates were consistently above the average background. At 7I this was due to radiation from ANL's Radioactive Waste Storage Facility (317 Area) in the northern half of grid 7I. Waste is packaged and temporarily kept in this area before removal for permanent disposal off-site. The dose at this perimeter fence location was about  $95 \pm 7$  mrem/y, one of the lowest values since these measurements were conducted. In previous years, this value has ranged up to 941 mrem/y which was in 1985. About 300 m (0.2 mi) south of the fence in grid 6I,

Environmental Penetrating Radiation at ANL, 1994

**TABLE 4.14** 

	D	ose Rate (m	rem/year)					
		Period	of Measuren	nent				
Location*	1/7-4/7	4/7-7/7 7/7-10/6		10/6-1/5	Avera	Average		
14G - Boundary	81	83	87	70	80 ±	7		
•		80	53	68		12		
14I - Boundary	77							
14L - Boundary	66	77	50	73	67 ±	12		
6I - 200 m N of Quarry Road	87	85	70	65	77 ±	11		
7I - Center, Waste Storage Area Facility 317	645	1621	1596	1958	1455 ±	553		
7I - Boundary	86	101	100	92	95 ±	7		
8H - Boundary	77	71	76	62	72 ±	7		
8H - 65 m S of Building 316	74	70	79	77	73 ±	4		
8H - 200 m NW of Waste Storage Area (Heliport)	70	84	89	72	79 ±	9		
8H - Boundary, Center, St. Patrick Cemetary	70	75	81	54	70 ±	11		
9H - 50 m SE of CP-5	113	91	116	74	99 ±	19		
9I - 65 m NE of Building 350, 230 m NE of Building 316	64	65	69	50	62 ±	8		
9/10EF - Boundary	84	121	78	70	88 ±	22		
10/11K - Lodging Facilities	65	75	58	63	65 ±	7		

<sup>\*</sup>See Figure 4.4

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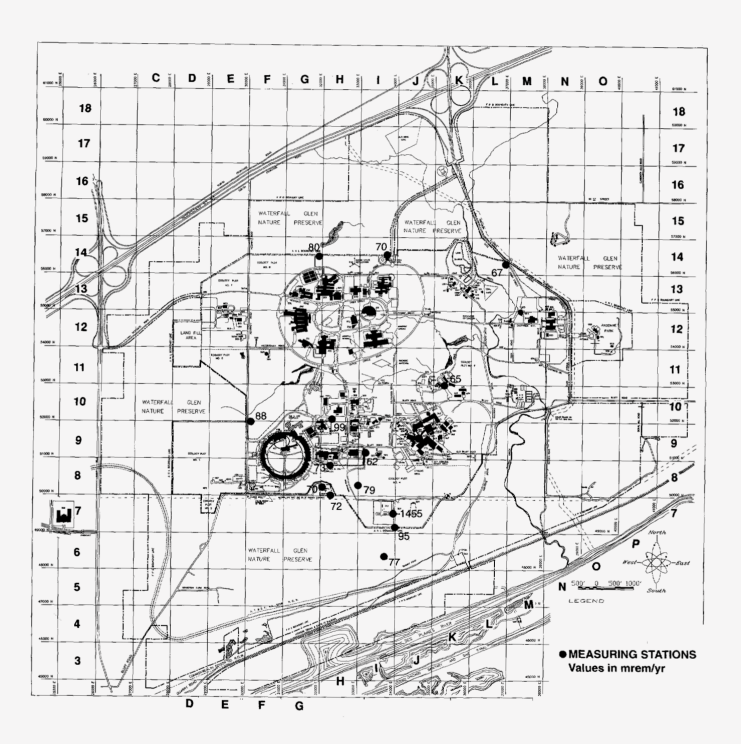


Figure 4.4 Penetrating Radiation Measurements at the ANL Site, 1994

the measured dose dropped to  $77 \pm 11$  mrem/y, within the normal background range. The recent decrease in dose in the 317 Area has been due to a concerted effort to transport radioactive historic waste off the site.

In the past, an elevated perimeter area was at Location 14I, at the north boundary. This dose was attributed to the use of cobalt-60 irradiation sources in Building 202. However, the irradiation program using the cobalt-60 source was terminated at the end of FY 1990 and not used at all since then. The perimeter dose at Location 14I,  $70 \pm 12$  mrem/y, was within the normal background range.

An elevated on-site dose was measured in the past at Location 9H, next to the CP-5 facility, where irradiated hardware from CP-5 was stored. During the past few years, considerable cleanup of the CP-5 yard occurred as part of the CP-5 D&D project. The dose at Location 9H decreased from about 1200 mrem/y in 1989 to 99 mrem/y in 1994. By the end of 1994, the clean-up was completed and the residual dose was from sources in the building.

#### 4.6. Estimates of Potential Radiation Doses

The radiation doses at the site boundary and off the site that could have been received by the public from radioactive materials and radiation leaving the site were calculated. These calculations were made for three exposure pathways, airborne, water, and direct radiation from external sources.

#### 4.6.1. Airborne Pathway

Guidance issued by the DOE<sup>6</sup> stipulates that DOE facilities with airborne releases of radioactive materials are subject to 40 CFR Part 61, Subpart H, <sup>13</sup> which requires the use

of the CAP-88 version of the EPA-AIRDOSE/RADRISK code to calculate the dose for radionuclides released to the air and to demonstrate compliance with the regulation. The dose limit applicable for CY 1994 for the air pathway is 10 mrem/y effective dose equiva lent. The EPA-AIRDOSE/RADRISK computer code uses a modified Gaussian plume equation to estimate both horizontal and vertical dispersion of radionuclides released to the air from stacks or area sources. For 1994, doses were calculated for hydrogen-3, carbon-11, nitrogen-13, oxygen-15, argon-41, krypton-85, radon-220 plus daughters and a number of actinide radionuclides. The annual release rates are those listed in Table 4.4, and separate calculations were performed for each of the seven release points. The wind speed and direction data shown in Figure 1.3 were used for these calculations. Doses were calculated for an area extending out to 80 km (50 mi) from ANL. The upgraded population distribution of the 16 compass segments and ten distance increments given in Table 1.1 was used. The dose rate was calculated at the midpoint of each interval and integrated over the entire area to give the annual population cumulative dose.

Distances from the specific facilities that exhaust radiological airborne emissions (see Table 4.4) to the fenceline (perimeter) and nearest resident were determined in the 16 compass segments. Calculations also were performed to evaluate the major airborne pathways; ingestion, inhalation, and immersion, both at the point of maximum perimeter exposure and to the maximally exposed resident. The perimeter and resident doses and the maximum doses are listed, respectively, for releases from Buildings 200 (Tables 4.15 and 4.16), Building 205 (Tables 4.17 and 4.18), Building 212 (Tables 4.19 and 4.20), Building 330 (Tables 4.21 and 4.22), Building 350 (Tables 4.23 and 4.24), Building 375 (Tables 4.25 and 4.26) and Building 411 (Tables 4.27 and 4.28). The doses given in these tables are the committed whole body effective dose equivalents.

The dominant contributor to the calculated doses was the radon-220 and daughters released from Building 200. This accounted for 90% of the off-site dose in 1994. The

TABLE 4.15

Radiological Airborne Releases from Building 200, 1994

Source Term: Radon-220 = 1750.1 Ci (plus daughters)

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N	500	0.280	1000	0.070
NNE	600	0.310	1100	0.088
NE	750	0.170	2600	0.017
ENE	1700	0.029	3100	0.011
E	2400	0.016	3500	0.009
ESE	2200	0.015	3600	0.007
SE	2100	0.018	4000	0.007
SSE	2000	0.020	4000	0.007
S	1500	0.013	4000	0.003
SSW	1000	0.055	2500	0.012
SW	800	0.019	2200	0.031
wsw	1100	0.049	1500	0.027
W	750	0.100	1500	0.026
WNW	800	0.069	1300	0.028
NW	600	0.150	1100	0.048
NNW	600	0.180	800	0.100

**TABLE 4.16** 

# Maximum Perimeter and Individual Doses from Building 200 Air Emissions, 1994

Dose (mrem/y)

Perimeter (600 m NNE)	Individual (800 m NNW)
3.0 x 10 <sup>-13</sup>	1.0 x 10 <sup>-13</sup>
$3.1 \times 10^{-1}$	1.0 x 10 <sup>-1</sup>
$1.7 \times 10^{-3}$	$5.6 \times 10^{-4}$
1.1 x 10 <sup>-4</sup>	4.8 x 10 <sup>-5</sup>
0.31	0.10
$1.4 \times 10^{-3}$	4.6 x 10 <sup>-4</sup>
2.8 x 10 <sup>-2</sup>	1.2 x 10 <sup>-2</sup>
1.4 x 10 <sup>-1</sup>	$6.4 \times 10^{-2}$
1.4 x 10 <sup>-1</sup>	2.8 x 10 <sup>-2</sup>
0.31	0.10
	3.0 x 10 <sup>-13</sup> 3.1 x 10 <sup>-1</sup> 1.7 x 10 <sup>-3</sup> 1.1 x 10 <sup>-4</sup> 0.31  1.4 x 10 <sup>-3</sup> 2.8 x 10 <sup>-2</sup> 1.4 x 10 <sup>-1</sup> 1.4 x 10 <sup>-1</sup>

TABLE 4.17

Radiological Airborne Releases from Building 205, 1994

Source Term: Hydrogen-3 = 0.39 Ci

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N	850	2 x 10 <sup>-5</sup>	1300	1 x 10 <sup>-5</sup>
NNE	1000	2 x 10 <sup>-5</sup>	2100	8 x 10 <sup>-6</sup>
NE	1200	2 x 10 <sup>-5</sup>	2700	5 x 10 <sup>-6</sup>
ENE	2400	5 x 10 <sup>-6</sup>	3000	3 x 10 <sup>-6</sup>
E	2200	5 x 10 <sup>-6</sup>	2400	3 x 10 <sup>-6</sup>
ESE	2000	5 x 10 <sup>-6</sup>	3500	2 x 10 <sup>-6</sup>
SE	1800	6 x 10 <sup>-6</sup>	3900	2 x 10 <sup>-6</sup>
SSE	1500	9 x 10 <sup>-6</sup>	4000	2 x 10 <sup>-6</sup>
S	1300	4 x 10 <sup>-6</sup>	3900	8 x 10 <sup>-7</sup>
SSW	1100	1 x 10 <sup>-5</sup>	2400	4 x 10 <sup>-6</sup>
SW	900	3 x 10 <sup>-5</sup>	2100	1 x 10 <sup>-5</sup>
WSW	1100	1 x 10 <sup>-5</sup>	1800	5 x 10 <sup>-6</sup>
W	1300	8 x 10 <sup>-6</sup>	1800	5 x 10 <sup>-6</sup>
WNW	1100	9 x 10 <sup>-6</sup>	1700	5 x 10 <sup>-6</sup>
NW	1100	1 x 10 <sup>-5</sup>	1500	8 x 10 <sup>-6</sup>
NNW	900	2 x 10 <sup>-5</sup>	1500	9 x 10 <sup>-6</sup>

**TABLE 4.18** 

# Maximum Perimeter and Individual Doses from Building 205 Air Emissions, 1994

Dose (mrem/y)

Pathway	Perimeter (850 m N)	Individual (1300 m N)	
Ingestion	8 x 10 <sup>-6</sup>	3 x 10 <sup>-6</sup>	
Inhalation	3 x 10 <sup>-5</sup>	8 x 10 <sup>-6</sup>	
Air Immersion	-	-	
Ground Surface	- -	-	
Total	3 x 10 <sup>-5</sup>	1 x 10 <sup>-5</sup>	
Radionuclide			
Hydrogen-3	3 x 10 <sup>-5</sup>	1 x 10 <sup>-5</sup>	

**TABLE 4.19** 

Radiological Airborne Releases from Building 212, 1994

Source Term: Hydrogen-3 (HT) = 40.15 Ci Hydrogen-3 (HTO) = 17.85 Ci Krypton-85 = 20.96 Ci Antimony-125 = 2.2 x 10<sup>-5</sup> Ci Radon-220 = 2.6 Ci

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N.	800	3.6 x 10 <sup>-3</sup>	2000	9.2 x 10 <sup>-4</sup>
NNE	1000	3.6 x 10 <sup>-3</sup>	2500	9.2 x 10 <sup>-4</sup>
NE	1300	2.2 x 10 <sup>-3</sup>	2000	1.2 x 10 <sup>-4</sup>
ENE	1500	1.5 x 10 <sup>-3</sup>	2500	7.0 x 10 <sup>-4</sup>
Е	1600	1.3 x 10 <sup>-3</sup>	2800	5.8 x 10 <sup>-4</sup>
ESE	1200	1.7 x 10 <sup>-3</sup>	2500	5.8 x 10 <sup>-4</sup>
SE	1400	1.5 x 10 <sup>-3</sup>	3500	3.9 x 10 <sup>-4</sup>
SSE	1400	1.5 x 10 <sup>-3</sup>	4500	2.8 x 10 <sup>-4</sup>
S	1500	5.7 x 10 <sup>-4</sup>	5000	9.5 x 10 <sup>-5</sup>
SSW	1600	1.1 x 10 <sup>-3</sup>	5000	2.0 x 10 <sup>-4</sup>
sw	1400	2.9 x 10 <sup>-3</sup>	2400	1.3 x 10 <sup>-3</sup>
wsw	1300	1.4 x 10 <sup>-3</sup>	2300	6.0 x 10 <sup>-4</sup>
W	1700	9.1 x 10 <sup>-4</sup>	2200	6.2 x 10 <sup>-4</sup>
WNW	1500	9.6 x 10 <sup>-4</sup>	2000	6.4 x 10 <sup>-4</sup>
NW	1300	1.5 x 10 <sup>-3</sup>	2000	8.4 x 10 <sup>-4</sup>
NNW	1000	2.6 x 10 <sup>-3</sup>	2000	9.6 x 10 <sup>-4</sup>

**TABLE 4.20** 

# Maximum Perimeter and Individual Doses from Building 212 Air Emissions, 1994

Dose (mrem/y)

Pathway	Perimeter (1000 m NNE)	Individual (2400 m SW)
Ingestion	8.0 x 10 <sup>-4</sup>	3.0 x 10 <sup>-4</sup>
Inhalation	$2.8 \times 10^{-3}$	$1.0 \times 10^{-3}$
Air Immersion	$1.0 \times 10^{-5}$	4.0 x 10 <sup>-6</sup>
Ground Surface	5.0 x 10 <sup>-6</sup>	2.0 x 10 <sup>-6</sup>
Total	3.6 x 10 <sup>-3</sup>	1.3 x 10 <sup>-3</sup>
Radionuclide		
Hydrogen-3	$3.4 \times 10^{-3}$	1.2 x 10 <sup>-3</sup>
Krypton-85	2.0 x 10 <sup>-5</sup>	6.0 x 10 <sup>-6</sup>
Antimony-125	5.0 x 10 <sup>-6</sup>	2.0 x 10 <sup>-6</sup>
Radon-220	4.0 x 10 <sup>-5</sup>	5.0 x 10 <sup>-7</sup>
Total	$3.6 \times 10^{-3}$	1.3 x 10 <sup>-3</sup>

TABLE 4.21

Radiological Airborne Releases from Building 330 (CP-5), 1994

Source Term: Hydrogen-3 (HTO) = 49.17 Ci

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N	1500	1.1 x 10 <sup>-3</sup>	2000	7.2 x 10 <sup>-4</sup>
NNE	1800	1.2 x 10 <sup>-3</sup>	3300	4.7 x 10 <sup>-4</sup>
NE	2100	8.7 x 10 <sup>-4</sup>	2800	5.6 x 10 <sup>-4</sup>
ENE	2200	6.7 x 10 <sup>-4</sup>	3300	3.6 x 10 <sup>-4</sup>
E	1500	1.2 x 10 <sup>-3</sup>	3100	3.9 x 10 <sup>-4</sup>
ESE	1300	1.2 x 10 <sup>-3</sup>	3500	2.7 x 10 <sup>-4</sup>
SE	1200	1.5 x 10 <sup>-3</sup>	3500	3.1 x 10 <sup>-4</sup>
SSE	1000	2.0 x 10 <sup>-3</sup>	3500	3.1 x 10 <sup>-4</sup>
S	500	$2.3 \times 10^{-3}$	3000	1.5 x 10 <sup>-4</sup>
SSW	700	2.8 x 10 <sup>-3</sup>	3500	2.6 x 10 <sup>-4</sup>
SW	900	4.2 x 10 <sup>-3</sup>	2400	1.0 x 10 <sup>-3</sup>
WSW	1400	1.0 x 10 <sup>-3</sup>	2000	5.8 x 10 <sup>-4</sup>
W	700	2.7 x 10 <sup>-3</sup>	2000	5.6 x 10 <sup>-4</sup>
WNW	700	2.3 x 10 <sup>-3</sup>	1900	5.4 x 10 <sup>-4</sup>
NW	1500	9.8 x 10 <sup>-4</sup>	2000	6.6 x 10 <sup>-4</sup>
NNW	1600	1.0 x 10 <sup>-3</sup>	1900	8.1 x 10 <sup>-4</sup>

**TABLE 4.22** 

# Maximum Perimeter and Individual Doses from Building 330 (CP-5) Air Emissions, 1994

# Dose (mrem/y)

Pathway	Perimeter (900 m SW)	Individual (2400 m SW)
Ingestion	1.0 x 10 <sup>-3</sup>	2.4 x 10 <sup>-4</sup>
Inhalation	$3.2 \times 10^{-3}$	7.8 x 10 <sup>-4</sup>
Air Immersion	-	-
Ground Surface	. <del>-</del>	-
Total	4.2 x 10 <sup>-3</sup>	$1.0 \times 10^{-3}$
Radionuclide		
Hydrogen-3	4.2 x 10 <sup>-3</sup>	1.0 x 10 <sup>-3</sup>

**TABLE 4.23** 

Radiological Airborne Releases from Building 350, 1994

Source Term: Uranium-234 =  $1.6 \times 10^{-8}$  Ci Uranium-238 =  $1.6 \times 10^{-8}$  Ci Plutonium-238 =  $2.5 \times 10^{-13}$  Ci Plutonium-239 =  $7.7 \times 10^{-8}$  Ci Plutonium-240 =  $2.1 \times 10^{-9}$  Ci Plutonium-241 =  $8.7 \times 10^{-10}$  Ci Plutonium-242 =  $6.9 \times 10^{-15}$  Ci

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N	1700	2 x 10 <sup>-6</sup>	2200	2 x 10 <sup>-6</sup>
NNE	1800	3 x 10 <sup>-6</sup>	3200	2 x 10 <sup>-6</sup>
NE	2200	2 x 10 <sup>-6</sup>	3100	1 x 10 <sup>-6</sup>
ENE	2000	2 x 10 <sup>-6</sup>	3100	$1 \times 10^{-6}$
E	1700	3 x 10 <sup>-6</sup>	2500	$2 \times 10^{-6}$
ESE	900	4 x 10 <sup>-6</sup>	3000	1 x 10 <sup>-6</sup>
SE	900	4 x 10 <sup>-6</sup>	3000	1 x 10 <sup>-6</sup>
SSE	700	6 x 10 <sup>-6</sup>	2700	1 x 10 <sup>-6</sup>
S	600	2 x 10 <sup>-6</sup>	2700	6 x 10 <sup>-7</sup>
SSW	400	3 x 10 <sup>-6</sup>	2500	1 x 10 <sup>-6</sup>
sw	600	8 x 10 <sup>-6</sup>	2700	2 x 10 <sup>-6</sup>
wsw	800	4 x 10 <sup>-6</sup>	2100	2 x 10 <sup>-6</sup>
W	900	$3 \times 10^{-6}$	2200	1 x 10 <sup>-6</sup>
WNW	1000	2 x 10 <sup>-6</sup>	2100	1 x 10 <sup>-6</sup>
NW	1900	2 x 10 <sup>-6</sup>	2400	1 x 10 <sup>-6</sup>
NNW	1900	2 x 10 <sup>-6</sup>	2200	2 x 10 <sup>-6</sup>

Maximum Perimeter and Individual Doses from

Dose (mrem/y)

Building 350 Air Emissions, 1994

**TABLE 4.24** 

Pathway	Perimeter (600 m SW)	Individual (2700 m SW
Ingestion	8 x 10 <sup>-8</sup>	2 x 10 <sup>-8</sup>
Inhalation	7 x 10 <sup>-6</sup>	2 x 10 <sup>-6</sup>
Air Immersion	1 x 10 <sup>-15</sup>	4 x 10 <sup>-16</sup>
Ground Surface	2 x 10 <sup>-10</sup>	5 x 10 <sup>-11</sup>
Total	8 x 10 <sup>-6</sup>	2 x 10 <sup>-6</sup>
<u>Radionuclide</u>		
Uranium-234	5 x 10 <sup>-7</sup>	2 x 10 <sup>-7</sup>
Uranium-238	4 x 10 <sup>-7</sup>	1 x 10 <sup>-7</sup>
Plutonium-238	2 x 10 <sup>-11</sup>	6 x 10 <sup>-12</sup>
Plutonium-239	6 x 10 <sup>-6</sup>	2 x 10 <sup>-6</sup>
Plutonium-240	2 x 10 <sup>-7</sup>	6 x 10 <sup>-8</sup>
Plutonium-241	1 x 10 <sup>-9</sup>	$4 \times 10^{-10}$
Plutonium-242	5 x 10 <sup>-11</sup>	2 x 10 <sup>-13</sup>
Total	8 x 10 <sup>-6</sup>	2 x 10 <sup>-6</sup>

TABLE 4.25

Radiological Airborne Releases from Building 375 (IPNS), 1994

Source Term Carbon-11 =305.9 Ci Argon-41 = 4.4 Ci

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N	1600	2.2 x 10 <sup>-2</sup>	3200	6.3 x 10 <sup>-3</sup>
NNE	1700	2.9 x 10 <sup>-2</sup>	3100	9.9 x 10 <sup>-3</sup>
NE	1700	2.6 x 10 <sup>-2</sup>	2700	1.2 x 10 <sup>-2</sup>
ENE	1500	2.7 x 10 <sup>-2</sup>	2500	1.1 x 10 <sup>-2</sup>
E	600	1.1 x 10 <sup>-1</sup>	2500	1.2 x 10 <sup>-2</sup>
ESE	600	9.6 x 10 <sup>-2</sup>	2500	9.1 x 10 <sup>-3</sup>
SE	600	9.9 x 10 <sup>-2</sup>	2500	1.0 x 10 <sup>-2</sup>
SSE	600	1.1 x 10 <sup>-1</sup>	3000	7.2 x 10 <sup>-3</sup>
S	800	2.7 x 10 <sup>-2</sup>	3000	2.8 x 10 <sup>-3</sup>
SSW	800	5.4 x 10 <sup>-2</sup>	3500	$4.3 \times 10^{-3}$
sw	800	1.1 x 10 <sup>-1</sup>	4000	$6.2 \times 10^{-3}$
WSW	1500	$2.0 \times 10^{-2}$	2700	7.1 x 10 <sup>-3</sup>
W	2200	1.1 x 10 <sup>-2</sup>	2700	$6.7 \times 10^{-3}$
WNW	1500	1.6 x 10 <sup>-2</sup>	2600	6.2 x 10 <sup>-3</sup>
NW	2200	1.1 x 10 <sup>-6</sup>	2500	8.8 x 10 <sup>-3</sup>
NNW	1800	1.8 x 10 <sup>-2</sup>	2200	1.3 x 10 <sup>-2</sup>

Maximum Perimeter and Individual Doses from Building 375 (IPNS) Air Emissions, 1994 Dose (mrem/y)

**TABLE 4.26** 

Pathway	Perimeter (600 m E)	Individual (2200 m NNW)
Ingestion	-	-
Inhalation	$5.0 \times 10^{-3}$	5.4 x 10 <sup>-4</sup>
Air Immersion	1.0 x 10 <sup>-1</sup>	$1.2 \times 10^{-2}$
Ground Surface	3.8 x 10 <sup>-3</sup>	4.5 x 10 <sup>-4</sup>
Total	1.1 x 10 <sup>-1</sup>	1.3 x 10 <sup>-2</sup>
Radionuclide		
Carbon-11	1.1 x 10 <sup>-1</sup>	1.2 x 10 <sup>-2</sup>
Argon-41	$2.0 \times 10^{-3}$	2.8 x 10 <sup>-4</sup>
Total	1.1 x 10 <sup>-1</sup>	1.3 x 10 <sup>-2</sup>

TABLE 4.27

Radiological Airborne Releases from Building 411/415 (APS), 1994

Source Term: Carbon-11 = 0.94 Ci

Nitrogen-1 = 40.6 Ci Oxygen-15 = 4.3 Ci

Direction	Distance to Perimeter (m)	Dose (mrem/y)	Distance to Nearest Resident (m)	Dose (mrem/y)
N	1500	2.7 x 10 <sup>-3</sup>	2000	4.0 x 10 <sup>-7</sup>
NNE	1600	$3.6 \times 10^{-3}$	2100	4.0 x 10 <sup>-7</sup>
NE	2200	1.8 x 10 <sup>-3</sup>	3100	1.6 x 10 <sup>-7</sup>
ENE	2500	1.1 x 10 <sup>-3</sup>	3300	1.3 x 10 <sup>-7</sup>
Е	1600	2.7 x 10 <sup>-3</sup>	3400	1.2 x 10 <sup>-7</sup>
ESE	1500	2.5 x 10 <sup>-3</sup>	3500	9.5 x 10 <sup>-8</sup>
SE	400	2.3 x 10 <sup>-3</sup>	3000	1.2 x 10 <sup>-7</sup>
SSE	400	$2.5 \times 10^{-3}$	3000	1.2 x 10 <sup>-7</sup>
S	350	1.2 x 10 <sup>-2</sup>	2500	7.2 x 10 <sup>-8</sup>
SSW	400	1.9 x 10 <sup>-2</sup>	2800	1.3 x 10 <sup>-7</sup>
sw	550	2.4 x 10 <sup>-2</sup>	3000	1.9 x 10 <sup>-7</sup>
wsw	800	7.0 x 10 <sup>-3</sup>	1400	6.6 x 10 <sup>-7</sup>
W	800	6.4 x 10 <sup>-3</sup>	1500	5.3 x 10 <sup>-7</sup>
WNW	500	1.1 x 10 <sup>-2</sup>	1400	5.0 x 10 <sup>-7</sup>
NW	350	2.1 x 10 <sup>-2</sup>	1600	4.4 x 10 <sup>-7</sup>
NNW	1500	2.6 x 10 <sup>-3</sup>	2000	2.9 x 10 <sup>-7</sup>

**TABLE 4.28** 

## Maximum Perimeter and Individual Doses from Building 411/415 (APS) Air Emissions, 1994

Dose (mrem/y)

Pathway	Perimeter (400 m SSE)	Individual (1400 m SW)
Ingestion	-	-
Inhalation	7.2 x 10 <sup>-4</sup>	1.8 x 10 <sup>-8</sup>
Air Immersion	2.4 x 10 <sup>-2</sup>	6.3 x 10 <sup>-7</sup>
Ground Surface	4.2 x 10 <sup>-4</sup>	1.1 x 10 <sup>-8</sup>
Total	2.5 x 10 <sup>-2</sup>	6.6 x 10 <sup>-7</sup>
Radionuclide		
Carbon-11	5.8 x 10 <sup>-4</sup>	8.3 x 10 <sup>-8</sup>
Nitrogen-13	$2.3 \times 10^{-2}$	$4.2 \times 10^{-7}$
Oxygen-15	$1.6 \times 10^{-3}$	1.5 x 10 <sup>-7</sup>
Total	2.5 x 10 <sup>-2</sup>	6.6 x 10 <sup>-7</sup>

highest perimeter dose rates were in the north-northwest sector with a maximum dose of 0.35 mrem/y at a fenceline location north of Building 203 (location 14H in Figure 1.1). The major contributor to this dose was inhalation of lead-212 (0.14 mrem/y) and the organs receiving the greatest dose were the lung and the bone. The releases from the other facilities are very minor contributors to the total dose.

During 1994, a significant program progressed to D&D the M-wing hot cells in Building 200, the source of the radon-220 emissions. Much of the year was spent in planning and preparation activities, although by the end of 1994, a considerable amount of waste was packaged and/or removed from the hot cells. This has resulted in a decrease of radon-220 emissions, 2023 Ci in 1993 to 1750 Ci in 1994. Work on the major source of the radon-220, cell M-1, is planned for 1995.

In August of 1992, the JANUS reactor (Building 202) terminated operation because of a lack of programmatic support. In early 1993, the fuel was removed and shipped to the Savannah River Plant for reprocessing. Likewise, the cyclotron in Building 211 ceased operation at the end of 1992 because of the lack of use. The facility was placed in standby status awaiting future D&D. Neither facility will produce radiological airborne emissions in the future.

The full-time resident who would receive the largest annual dose (0.12 mrem/y) is located approximately 0.8 km (0.5 mi) NNW of the site boundary. The major contributor to the whole body dose is the inhalation dose from lead-212 (0.06 mrem/y). If radon-220 and daughters were excluded from the calculation, as required by NESHAP, <sup>13</sup> the maximally exposed resident would receive a dose of 0.015 mrem/y, primarily carbon-11 from the IPNS facility (Building 375).

The individual doses to the maximally exposed member of the public and the maximum fenceline dose is shown in Figure 4.5. The decrease in individual and population doses since 1988 are due in part to the decrease of the radon-220 emissions from the Proof-of-Breeding Program.

The population data in Table 1.1 were used to calculate the cumulative population dose from gaseous radioactive effluents from ANL operations. The results are given in Table 4.29, together with the natural external radiation dose. The natural radiation dose listed is the product of the 80-km (50-mi) population and the natural radiation dose of 300 mrem/y.<sup>14</sup> It is assumed that this dose is representative of the entire area within an 80-km

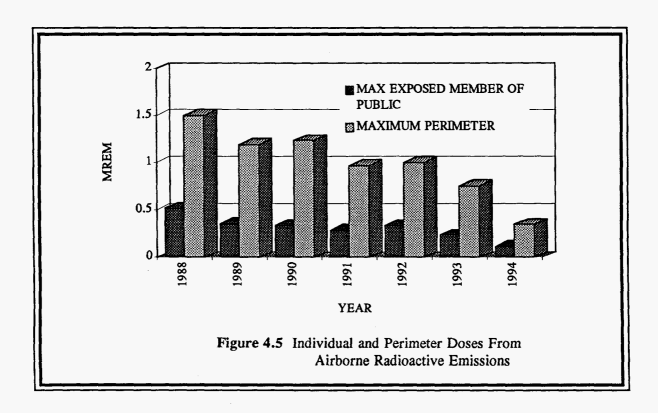


TABLE 4.29

80 km Population Dose, 1994

Radionuclide	man-rems
Hydrogen-3	0.40
Carbon-11	0.92
Nitrogen-13	0.04
Oxygen-15	< 0.01
Argon-41	0.06
Krypton-85	< 0.01
Antimony-125	< 0.01
Thallium-208	< 0.01
Lead-212	3.97
Bismuth-212	0.43
Radon-220	< 0.01
Uranium-234	< 0.01
Uranium-238	< 0.01
Plutonium-238	< 0.01
Plutonium-239	< 0.01
Plutonium-240	< 0.01
Plutonium-241	< 0.01
Plutonium-242	< 0.01
Total	5.8
Natural	$2.4 \times 10^6$

(50-mi) radius. The population dose since 1987, due to ANL operations, is shown in Figure 4.6.

The potential radiation exposures by the inhalation pathways also were calculated by the methodology specified in DOE Order 5400.5.<sup>6</sup> The total quantity for each radionuclide inhaled, in microcuries ( $\mu$ Ci), is calculated by multiplying the annual average air concentrations by the general public breathing rate of 8,400 m<sup>3</sup>/y.<sup>15</sup> This annual intake is then multiplied by the CEDE for the appropriate lung retention class.<sup>8</sup> Because the CEDE factors are in units of Rem per microcurie (Rem/ $\mu$ Ci), this calculation gives the 50-year committed effective dose equivalent. The applicable CEDE factors are listed in Table 4.30.

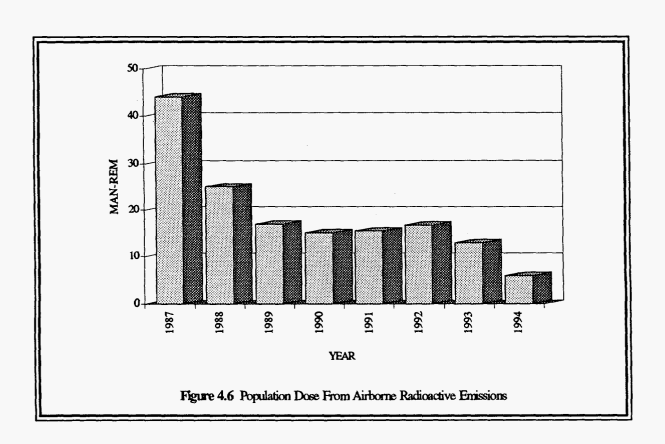


TABLE 4.30 
50-Year Committed Effective Dose Equivalent (CEDE) Factors  $(\text{Rem}/\mu\text{Ci})$ 

Nuclide	Ingestion	Inhalation	
Hydrogen-3	6.3 x 10 <sup>-5</sup>	6.3 x 10 <sup>-5</sup>	
Beryllium-7	-	2.7 x 10 <sup>-4</sup>	
Carbon-11	-	8.0 x 10 <sup>-6</sup>	
Strontium-90	0.13	1.32	
Cesium-137	0.05	0.032	
Lead-210	-	13.2	
Radium-226	1.1	-	
Thorium-228	-	310	
Thorium-230	-	260	
Thorium-232	-	1100	
Uranium-234	0.26	130	
Uranium-235	0.25	120	
Uranium-238	0.23	120	
Neptunium-237	3.90	-	
Plutonium-238	3.80	-	
Plutonium-239	4.30	330	
Americium-241	4.50	-	
Curium-242	0.11	-	
Curium-244	2.30	-	
Californium-249	4.60	-	
Californium-252	0.94	-	

The calculated doses in Tables 4.2 and 4.3 were obtained using this procedure. Because they are all essentially at perimeter locations, these doses represent the fenceline values for those radionuclides measured. In most cases, these doses also are the same as the off-site measurements and represent the ambient dose for the area from these nuclides. No doses are calculated for the total alpha and total beta measurements since the guidance does not provide CEDE factors for such measurements.

#### 4.6.2. Water Pathway

Following the methodology outlined in DOE Order 5400.5, the annual intake of radionuclides (in  $\mu$ Ci) ingested with water is obtained by multiplying the concentration of radionuclides in microcuries per milliliter ( $\mu$ Ci/mL) by the average annual water consumption of a member of the general public (7.3 x 10<sup>5</sup> mL). This annual intake is then multiplied by the CEDE factor for ingestion (Table 4.30) to obtain the dose received in that year. This procedure is carried out for all radionuclides and the individual results are summed to obtain the total ingestion dose.

The only location where radionuclides attributable to ANL operations could be found in off-site water was Sawmill Creek below the waste-water outfall, see Table 4.5. Although this water is not used for drinking purposes, the 50-year effective dose equivalent was calculated for a hypothetical individual ingesting water at the radionuclide concentrations measured at that location. Those radionuclides added to Sawmill Creek by ANL wastewater, their net concentrations in the creek and the corresponding dose rates (if water at these concentrations were used as the sole water supply by an individual) are given in Table 4.31. The dose rates were all well below the standards for the general population. It should be emphasized that Sawmill Creek is not used for drinking, swimming, or boating. Inspection of the area shows there are fish in the stream, but they do not constitute a

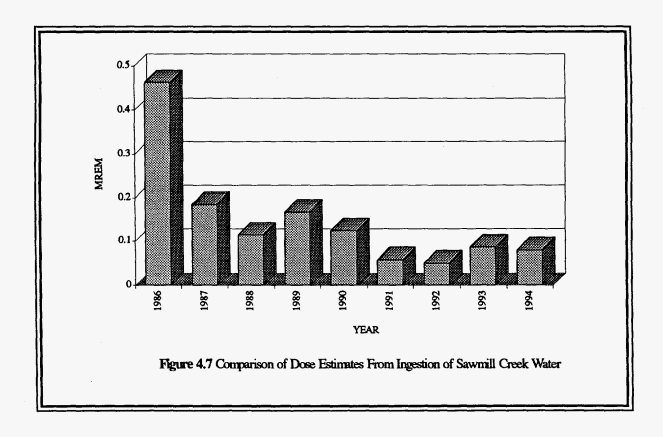
significant source of food for any individual. Figure 4.7 is a plot of the estimated dose an individual would receive if ingesting Sawmill Creek water.

Radionuclide Concentrations and Dose Estimates for Sawmill Creek Water, 1994

**TABLE 4.31** 

Radionuclide	Total Released (curies)	Net Avg Conc (pCi/L)	Dose (mrem)
Hydrogen-3	1.14	114	0.0052
Strontium-90	0.0033	0.33	0.0313
Cesium-137	0.0083	0.83	0.0303
Plutonium-239	0.000013	0.0013	0.0041
Americium-241	0.000024	0.0024	0.0079
Sum	1.15		0.0788

As indicated in Table 4.5, occasional Sawmill Creek samples (fewer than ten percent) contained traces of neptunium-237, plutonium-238, curium-242,244, or californium-249,252, but the averages were only slightly greater than the detection limit. The annual dose to an individual consuming water at these concentrations can be calculated with the same method used for those radionuclides more commonly found in creek water, but the method of averaging probably overestimates the true concentration. Annual doses range from  $3 \times 10^{-3}$  to  $6 \times 10^{-5}$  mrem/y for these radionuclides.



DOE Order 5400.5<sup>6</sup> requires that an evaluation be made of the dose to aquatic organisms from liquid effluents. The dose limit is one rad/day or 365 rad/y. The location that could result in the highest dose to aquatic organisms is in Sawmill Creek downstream of the point where ANL discharges its treated wastewater. Based on inspection of the creek at this location, small bluegill and carp (about 100 g each) have been observed. Using the annual average concentrations of the radionuclides listed in Table 4.5, a dose can be estimated. The sum of the exposure from these radionuclides is estimated to be about 5 x 10<sup>-6</sup> rad/y, well within the DOE standard, and therefore demonstrating compliance with that portion of the Order.

The EPA has established drinking water standards based on a maximum dose of 4 mrem/y for man-made beta particle and photon-emitting radionuclides. <sup>16</sup> The EPA standard is 2 x 10<sup>4</sup> pCi/L for hydrogen-3 and 8 pCi/L for strontium-90. The net concentrations in Table 4.31 correspond to 0.57% (hydrogen-3) and 4.1% (strontium-90) of the EPA standards. No specific EPA standards exist for the transuranic nuclides.

Sawmill Creek flows into the Des Plaines River. The flow rate of Sawmill Creek (see Section 1.6) is about 10 cfs, while the flow rate of the Des Plaines River in the vicinity of ANL is about 900 cfs. Applying this ratio to the concentration of radionuclides in Sawmill Creek listed in Table 4.31, the dose to a hypothetical individual ingesting water from the Des Plaines River at Lemont would be about 0.0010 mrem/y. Significant additional dilution occurs further downstream. Very few people, either directly or indirectly, use the Des Plaines River as a source of drinking water. If 100 people used Des Plaines River water at the hypothetical concentration at Lemont, the estimated population dose would be about  $10^{-4}$  man-rem.

## 4.6.3. External Direct Radiation Pathway

The TLD measurements given in Section 4.5 were used to calculate the radiation dose from external sources. Above-normal fenceline doses attributable to ANL operations were found at the southern boundary near the Waste Storage Facility (Location 7I).

At Location 7I, the fenceline dose from ANL was about 95  $\pm$  7 mrem/y. Approximately 300 m (0.3 mi) south of the fenceline (grid 6I), the measured dose was 77  $\pm$  11 mrem/y, the same as the normal range of the off-site average (82  $\pm$  10 mrem/y). No

individuals live in this area. The closest residents are about 1.6 km (1 mi) south of the fenceline. At this distance, the calculated dose rate from the Waste Storage Facility was 0.002 mrem/y, if the energy of the radiation were that of 0.66 MeV cesium-137 gammaray, and about 0.006 mrem/y if the energy were that of 1.33 MeV cobalt-60 gamma-ray.

At the fenceline, where higher doses were measured, the land is wooded and unoccupied. All of these dose calculations are based on full-time, outdoor exposure. Actual exposures to individuals would be substantially less, since some of the individuals are indoors (which provides shielding) or away from their dwellings for some of the time.

In addition to the permanent residences in the area, occasionally visitors may conduct activities around ANL that could result in exposure to radiation from this site. Examples of these activities could be cross country skiing, horseback riding, or running in the fire lane next to the perimeter fence. If the individual spent ten minutes per week adjacent to the 317 Area, the dose would be 0.01 mrem/y at the 317 Area fence (location 7I).

#### 4.6.4. Dose Summary

The total effective dose equivalent received by off-site residents during 1994 was a combination of the individual doses received through the separate pathways that contributed to exposure: hydrogen-3, carbon-11, nitrogen-13, oxygen-15, argon-41, krypton-85, radon-220 (plus daughters), and actinides through the airborne pathway. The highest dose was about 0.12 mrem/y to individuals living north of the site if they were outdoors at that location during the entire year. The total annual population dose to the entire area within

an 80-km (50-mi) radius is 5.8 man-rem. The dose pathways are collected in Table 4.32 and compared to the applicable standards.

TABLE 4.32

Summary of the Estimated Dose to the Public, 1994

(mrem/y)

Pathway	ANL Estimate	Applicable Standard
Air (Less radon)	0.015	10 (EPA)
Air Total	0.12	100 (DOE)
Water	0.079	100 (DOE)
Direct Radiation	0.01	100 (DOE)
Maximum Public	0.12	

To put the maximum individual dose of 0.12 mrem/y attributable to ANL operations into perspective, comparisons can be made to annual average doses received by the public from natural or accepted sources of radiation. These values are listed in Table 4.33. It is obvious that the magnitude of the doses received from ANL operations is insignificant compared with these sources. Therefore, the monitoring program results establish that the radioactive emissions from ANL are very low and do not endanger the health or safety of those living in the vicinity of the site.

**TABLE 4.33** 

# Annual Average Dose Equivalent in the U. S. Population\*

Source		Dose (mrem)
Natural Sources		
Radon		200
Internal ( <sup>40</sup> K and <sup>226</sup> Ra)		39
Cosmic		28
Terrestrial		28
Medical		
Diagnostic X-rays		39
Nuclear Medicine		14
Consumer Products		
Domestic Water Supplies,		10
Building Materials, etc.		
Occupational (medical		1
radiology, industrial		
radiography, research, etc.)		
Nuclear Fuel Cycle		< 1
Fallout		< 1
Other Miscellaneous Sources		< 1
	Total	360

<sup>\*</sup>NCRP Report No. 93.14



5.	ENVIRONMENTAL	. NONRADIOLOGICAL	PROGRAM INFORMATION
			•
5	-2		ANL Site Environmental Repor

The nonradiological monitoring program involves the collection and analysis of surface water and groundwater samples from numerous locations throughout the site. The release of nonradiological pollutants to the air from ANL is extremely small, except for the boiler house, which is equipped with dedicated monitoring equipment for sulfur dioxide (SO<sub>2</sub>) and opacity. One excursion for opacity was noted during 1994 over a period of 4100 hours of operation of Boiler No. 5, the coal-burning boiler. No SO<sub>2</sub> excursions were noted for Boiler No. 5 during 1994. As a result, the ambient air is not routinely monitored. Chapter 3 discusses the entire environmental monitoring program in more detail.

Surface water samples for nonradiological chemical analyses are collected from NPDES permitted outfalls and Sawmill Creek. Analyses conducted on the samples from the NPDES outfalls vary depending on the permit-mandated monitoring requirements for each outfall. The results of the analyses are compared with the permit limits for each outfall to determine whether they comply with the permit. Besides being published in this report, the NPDES monitoring results are transmitted monthly to the IEPA in an official Discharge Monitoring Report (DMR). A summary of exceedances of permit limits during 1994 appears in Table 5.1.

In addition to the permit-required monitoring, other analyses are conducted on samples collected from the combined wastewater outfall (NPDES outfall 001) to provide a more complete evaluation of the impact of the wastewater on the environment. Samples of water from Sawmill Creek are also collected and analyzed for a number of inorganic constituents. The results of these additional analyses of the main outfall and receiving streams are then compared with IEPA General Effluent Standards and Stream Quality Standards listed in the IAC, Title 35, Subtitle C, Chapter I.<sup>17</sup>

TABLE 5.1

NPDES Permit Limit Exceedances, 1994

Outfall	Parameter	Number of Exceedances
001	Total Dissolved Solids Copper	11 8
001A	Biochemical Oxygen Demand	1
001B	Total Suspended Solids Mercury	1 1
003A	Total Suspended Solids	1
004	Total Suspended Solids	1

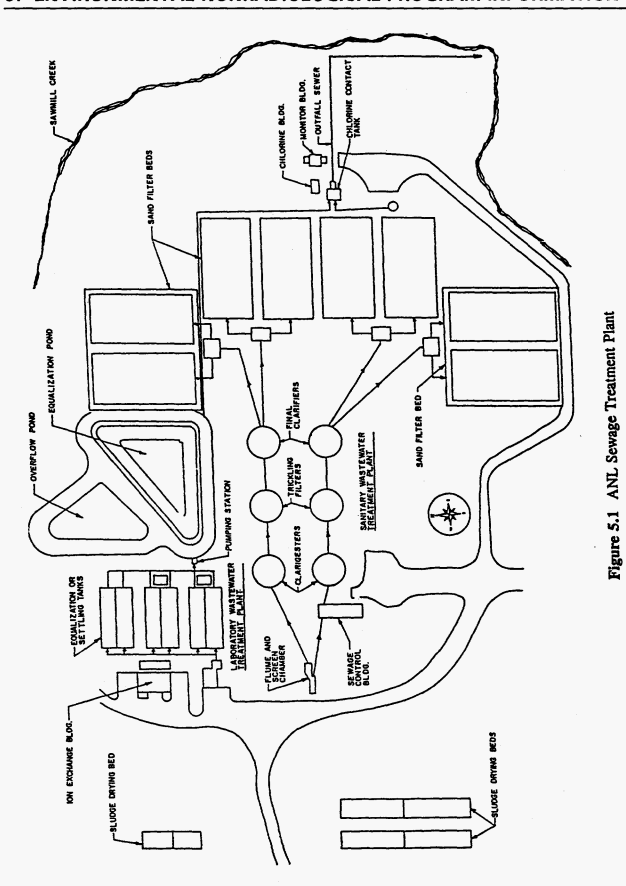
# 5.1. National Pollutant Discharge Elimination System Monitoring Results

Wastewater is processed at ANL in two independent treatment systems, the sanitary system and the laboratory system. The sanitary wastewater collection and treatment system collects wastewater from lavatories, the cafeteria, office buildings, and other portions of the site which do not contain radioactive or hazardous materials. This wastewater is treated in a biological wastewater treatment system consisting of primary clarifiers, trickling filters, final clarifiers, and slow sand filters. Wastewater generated by research-related activities, such as laboratories and experimental equipment, flows to a series of retention tanks located in each building. When a retention tank is full, a sample is collected and analyzed for radioactivity. If the wastewater is found to be below the release limits for discharge, it is pumped to the laboratory wastewater collection system, which directs the flow to the laboratory wastewater treatment system. This system consists of a series of concrete

holding tanks which collect the wastewater prior to discharge. As with the retention tanks, once a holding tank is full, it is sampled and analyzed for radioactivity. If the level of radioactivity is below ANL discharge criteria, which were selected to ensure compliance with DOE Orders, it is pumped to a lined equalization basin, slowly combined with the sanitary waste stream, chlorinated, and discharged to Sawmill Creek. If either a retention tank or holding tank is found to contain unacceptable levels of radioactivity, the wastewater is pumped into portable tanks, treated by evaporation in Building 306 and the residue is disposed of as radioactive waste. Figure 5.1 shows the two wastewater treatment systems that are located adjacent to each other. The volume of wastewater discharged from these facilities averaged 1.37 million liters per day (0.36 million gallons per day) sanitary wastewater and 1.15 millions liters per day (0.30 million gallon) laboratory process wastewater.

# 5.1.1. Sewage Treatment Plant Rehabilitation

Two projects to rehabilitate the existing Laboratory and Sanitary Wastewater Treatment Plant facilities are currently in design. The existing laboratory treatment facilities will be rehabilitated and additional treatment units will be provided. New facilities will provide treatment capability for heavy metals, suspended solids, and volatile and semivolatile organics. The hydraulic capacity of the plant will be expanded to enable treatment of the existing flow rate and anticipated future loadings. Existing equipment to be replaced includes flow meters, bar screens, sludge pumps and piping, sludge scrapers, and flow regulating valves and chambers for holding tanks. Major treatment process equipment areas to be provided in the rehabilitation include: oil and grease removal, grit removal, metals precipitation, suspended solids filtering, air stripping, carbon adsorption, sludge handling, flow monitoring, and process control instrumentation. This project is currently in final design.



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The Sanitary Wastewater Treatment Plant will be upgraded to replace equipment that has reached its design life and to add equipment where a more efficient or environmentally-sound process applies. The equipment to be rehabilitated includes the headworks, the clarigesters, the existing trickling filters, the final clarifiers, and the intermittent sand filters. The disinfection system will not be upgraded or replaced as it is no longer used or required. An additional 1000 m<sup>2</sup> (10,000 sq<sup>2</sup>) of sludge drying area will be provided with two additional drying beds utilizing sand with an underdrain system. The preliminary design has been completed and the final design was initiated during FY 1994.

#### 5.1.2. Effluent Monitoring

The two treatment plant systems process the vast majority of wastewater generated by ANL. However, a small amount of process wastewater, primarily cooling tower blowdown and cooling water, is discharged directly to a number of small streams and ditches throughout the site. This wastewater does not contain significant amounts of contaminants and does not require treatment before discharge. However, these discharge points are included in the site NPDES permit as separate regulated outfalls.

ANL-processed wastewater discharges are regulated by NPDES Permit No. IL 0034592 (renewed effective October 30, 1994). <sup>18</sup> Twenty-six surface water discharge points (outfalls) and two internal monitoring points are included in this permit. The analyses required and the frequency of analysis for each point are specified in the permit. The analytical methods required for NPDES monitoring are listed in Table 1B of 40 CFR Part 136. <sup>19</sup> Sample collection, preservation, and holding times are also mandated by requirements stipulated in Table 2 of 40 CFR Part 136. <sup>19</sup>

The NPDES outfall locations are shown in Figure 5.2. Outfalls 001A and 001B, the two internal monitoring points representing the effluent from the sanitary system and

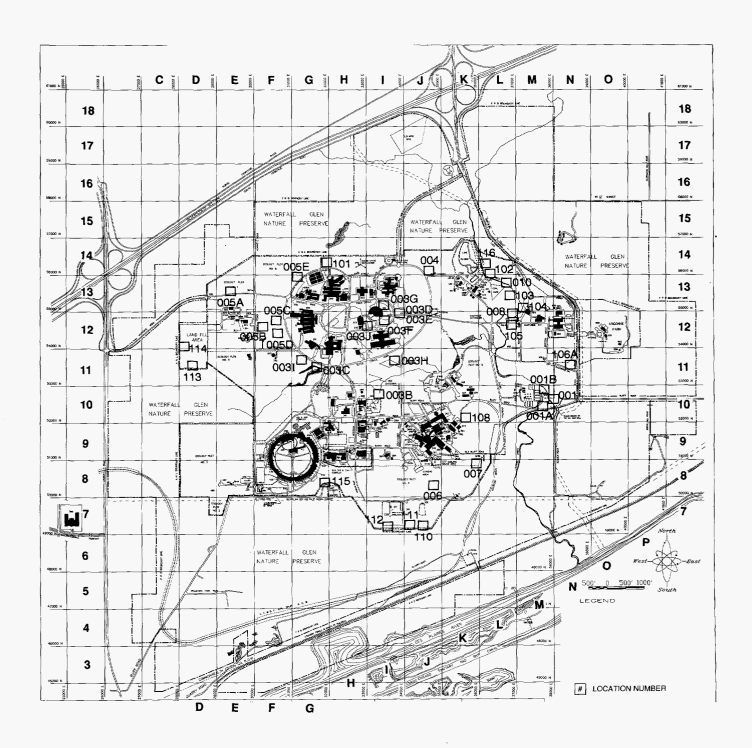


Figure 5.2 NPDES Outfall Locations

laboratory system, respectively, are both located at the wastewater treatment facility. Their flows combine to form outfall 001 which is also located at the treatment facility. The combined stream flows through an outfall pipe which discharges into Sawmill Creek approximately 1100 m (3500 ft) south of the treatment plant.

#### 5.1.2.1. Sample Collection

NPDES samples are collected by ANL's Environment and Waste Management Program (EWM) personnel, with the exception of samples from locations 001, 001A, and 001B, which are collected by Plant Facilities and Services Division (PFS) personnel. All samples are collected using specially cleaned and labelled bottles with appropriate preservatives added. Custody seals and chain-of-custody sheets are also used. All samples are analyzed within the required holding time. Samples are collected at locations 001A, 001B, and 001 a weekly basis. Samples are collected at the other locations in accordance with the NPDES permit.

#### 5.1.2.2. Sample Analysis - NPDES

NPDES sample analyses were performed using standard operating procedures (SOPs) written, reviewed, and issued as controlled documents by members of the Environment, Safety, and Health Division, Dosimetry and Analytical Services Section, Radiochemistry Laboratory (ESH-DACH), and Control Laboratory (ESH-DACL). These SOPs reference protocols found in 40 CFR Part 136, "Test Procedures for the Analysis of Pollutants Under the Clean Water Act." Six metal analyses were performed using flame atomic absorption spectroscopy. Mercury was determined by cold vapor atomic absorption spectroscopy. Hexavalent chromium determination was performed using a colorimetric technique. Five-day biochemical oxygen demand (BOD<sub>5</sub>) was determined using a dissolved oxygen probe. Total suspended solids, total dissolved solids,

and fats, oils, and grease were determined gravimetrically. Sulfate determination was performed using a turbidimetric technique while chloride was determined by titrimetry. Ammonia was determined by distillation followed by an ion selective electrode finish. Five volatile organic compounds were determined by using a purge and trap sample pretreatment followed by gas chromatography-mass spectroscopy detection. Beta radioactivity was performed using a gas flow proportional counting technique. Hydrogen-3 was determined by distillation followed by a beta liquid scintillation counting technique.

Semiannually, NPDES outfall 001B is sampled and analyzed for priority pollutant compounds. An off-site contracted laboratory performed these determinations using methods specified in 40 CFR Part 136. Volatile organic compounds were determined using a purge and trap sample pretreatment followed by gas chromatography-mass spectroscopy detection. Semivolatile organic compounds and 2,3,7,8-tetrachlorodibenzo-p-dioxin were determined by solvent extraction followed by gas chromatography-mass spectroscopy detection. PCB/pesticides were determined by solvent extraction followed by gas chromatography-electron capture detection. Thirteen metals were determined by graphite furnace atomic absorption, flame atomic absorption, and inductively coupled plasma-atomic emission spectrometry. Asbestos analysis was performed using transmission electron microscopy. Cyanide and phenol were determined by distillation followed by a spectrophotometric finish.

Annually, NPDES outfall 001 is sampled and analyzed for acute and chronic aquatic toxicity parameters. An off-site contracted laboratory performed both the sample collection and analyses. The methods used are described in "Methods for Measuring the Acute Toxicity of Effluents and Receiving Water to Freshwater and Marine Organisms," (EPA/600/4-90/027F, 4th Edition, "Environmental Effects Tests Guidelines," USEPA/560-6-82-002), and IEPA specifications given in "Effluent Biomonitoring and Toxicity Assessment-Aquatic Life Concerns." The testing is performed by using ANL effluent with Sawmill Creek receiving water, introducing species of fish, invertebrates, and aquatic plants

and measuring survival, growth and/or reproduction over two to seven days. Statistically, significant mortality, inhibition of growth and/or reproduction is reported as a function of effluent concentration.

#### 5.1.2.3. Results

During 1994, approximately 97.5% of all NPDES analyses were in compliance with their applicable permit limits as compared to 1993, 1992, and 1991 rates of 97.5%, 98%, and 96%, respectively. Specific limit exceedances are discussed later in this section as well as in Chapter 2. A discussion of the analytical results for each outfall follows.

#### 5.1.2.4. Outfalls

#### Outfall 001A

This outfall is composed of treated sanitary wastewater and various wastewater streams from the boiler house area, including coal pile stormwater runoff. The effectiveness of the sanitary wastewater treatment systems is evaluated by weekly monitoring for biochemical oxygen demand (BOD), pH, and total suspended solids (TSS). The limits for five-day BOD are a monthly average of 10 mg/L with a maximum value of 20 mg/L. The permit limits for TSS are a maximum concentration of 24 mg/L and a monthly average of 12 mg/L. The pH must range between values of 6 and 9. There was one exceedance of the BOD limit at outfall 001A during June 1994. Heavy rains resulted in higher than normal coal pile runoff. The higher than normal iron content for this runoff results in depletion of oxygen in the effluent, resulting in increased BOD.

The permit requires weekly monitoring for total chromium, copper, iron, lead, manganese, zinc, and oil and grease. The effluent limits for these parameters and results are

shown in Table 5.2. There are two limits listed, one is a maximum limit for any single sample and the other is for the average of all samples collected during the month. The constituents presented in Table 5.2 are present in the coal pile runoff which discharges to the sanitary sewage system. All samples collected and analyzed for these parameters were within the permit limits during 1994. The average shown in the table is the annual average for each constituent.

Outfall 001A Effluent Limits and Monitoring Results, 1994
(Concentrations in mg/L)

Constituent	Minimum	Average	Average Limit	Maximum	Maximum Limit
Chromium	_	< 0.02	1.00	< 0.02	2.00
Copper	0.03	0.08	0.50	0.32	1.00
Iron	0.08	0.22	2.00	0.85	4.00
Lead	-	< 0.10	0.20	< 0.10	0.40
Manganese	0.01	0.04	1.00	0.14	2.00
Zinc	0.02	0.08	1.00	0.15	2.00
Oil & Grease	-	< 5.0	15.0	< 5.0	30.0

### Outfall 001B

This outfall consists of processed wastewater from the laboratory wastewater system. The permit requires that weekly samples be collected and analyzed for BOD, TSS, mercury, pH, and chemical oxygen demand (COD).

The limits established for BOD are a daily maximum of 20 mg/L with a 30-day average of 10 mg/L. The permit also contains BOD mass loading limits of 114 lbs/day as a daily maximum and 57 lbs/day as a 30-day average. The mass loading represents the weight of material discharged per day and is a function of concentration and flow. The daily maximum limit for TSS is 24 mg/L with a 30-day average of 12 mg/L. The TSS mass loading limits are 136 and 68 lbs/day, respectively. There was one exceedance of the concentration limit for TSS.

The daily maximum concentration limit for mercury is 6  $\mu$ g/L and the 30-day average is 3  $\mu$ g/L. The corresponding loading values are 0.034 lbs/day and 0.017 lbs/day. In 1994, there was one exceedance of the concentration limit.

There are no concentration limits established for COD. The once-per-week grab samples give a rough indication of the organic content of this stream. The values obtained in 1994 ranged from less than 10 mg/L to 38 mg/L.

There is a special condition for location 001B that requires the monitoring for the 126 priority pollutants, listed in the permit, during the months of June and December. The June sampling is to be conducted at the same time that aquatic toxicity testing of outfall 001 is conducted. In addition to the typical list of priority pollutants, fibrous asbestos and 2,3,7,8-tetrachlorodibenzo-p-dioxin (commonly called dioxin) are to be determined. Samples were collected on June 7, 1994, and December 8, 1994, and analyzed within the required holding times.

Analysis of these samples indicated that very small amounts of a few chemicals were present. The results for semivolatile organic compounds, PCBs, and pesticides were all less than the detection limits. The results for metals were similar to concentrations found in ANL treated drinking water. The samples contained several volatile organic compounds at

very low levels. The majority of compounds found are halomethanes. The concentrations of volatile organics identified in these samples are contained in Table 5.3. While there are currently no permit limits or effluent standards for these compounds with which to compare these results, the concentrations found are believed to be of little concern because they are below acceptable standards for drinking water supplies, where such standards exist.

TABLE 5.3

Outfall 001B Priority Pollutant Monitoring Results, 1994

(Concentrations in μg/L)

Compound	Concentration in June Sample	Concentration in December Sample
Bromodichloromethane	2.5	1.7
Bromoform	2.1	< 1.0
Chloroform	4.0	33.6
Dibromochloromethane	2.1	1.4
Methylene Chloride	< 5.0	123

No chrysotile asbestos fibers greater than 10 micrometers ( $\mu$ m) in length were detected. The December sample had 123  $\mu$ g/L of methylene chloride. Neither of the samples had detectable levels of dioxin.

The laboratory wastewater treatment system consists of six 69,000 gallon equalization or settling (holding) tanks (see Figure 5.1) which are pumped to a lined equalization pond before being discharged to Sawmill Creek. During 1989, a study was performed to determine the levels of volatile organic compounds in the influent to these tanks and to determine

the variability of this concentration. A number of different volatile organics were found to be present from time to time, with the concentration varying greatly throughout the day. Maximum levels were found to occur in the late afternoon. As a follow-up to this study, each month one influent sample is obtained at about 1300 hours and analyzed for volatile organic compounds. During August 1993, the discharge of water from Manhole 2E (refer to Section 6.2.2.3.) in the 317 Area began on a regular basis. This water is known to contain volatile organics at consistent levels. A modified NPDES permit was issued by the IEPA to reflect this discharge.

The 1994 results for the most common compounds found are shown in Table 5.4. In addition to these compounds, most samples contained very low concentrations of bromodichloromethane, dichlorobromomethane, and bromoform. These halomethanes, at the levels found, including some of the chloroform results, are thought to be due to the contact of the chlorinated supply water with organic chemicals. Chloroform levels above  $10 \mu g/L$  are probably due to other causes.

Similar to 1992 and 1993, acetone was found in only 42% of the samples and at lower concentrations (see Figure 5.3). The levels found ranged to 5969  $\mu$ g/L. Methylene chloride was found in 58% of the samples and ranged to 75  $\mu$ g/L, significantly lower concentrations than reported during 1992 and 1993 (see Figure 5.4). A sample obtained in October contained 1-propanol. Infrequent trace levels of other chemicals, i.e., carbon tetrachloride, trichloroethene, toluene, xylene, cis-1,2-dichloroethane, 2-butanone, tetrahydrofuran, chloromethane, butyl ether, ethylbenzene, 1,1,2-trichloromethane, 1,2-dichloroethane, 4-methyl-2-pentanone, d-limonene, and tetrachloroethane were also noted.

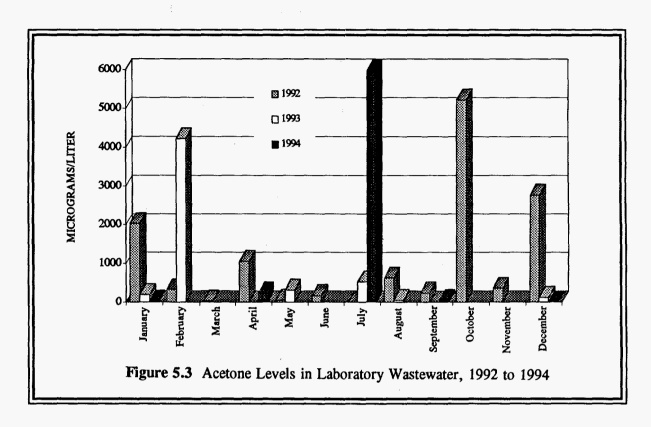
Figures 5.3, 5.4, and 5.5 present comparisons of the 1992, 1993, and 1994 laboratory wastewater results for the more persistent volatile organic compounds and show an overall reduction in the frequency and concentrations of the compounds. ANL's waste generator

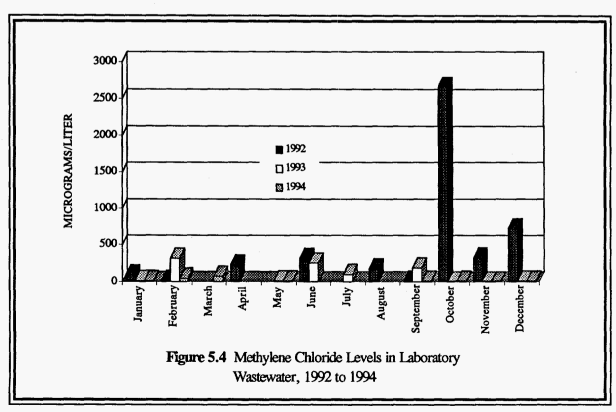
TABLE 5.4

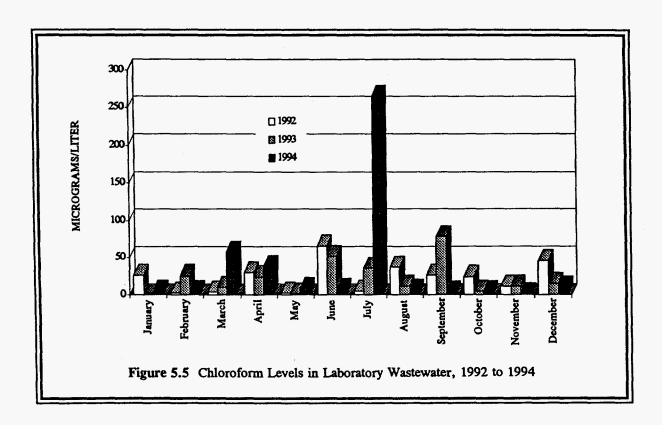
Volatile Organic Compounds in Laboratory Wastewater, 1994

(Concentrations in  $\mu$ g/L)

Month	Acetone	Chloroform	Methylene Chloride	Bromodi- chloroethane	Dibromo- chloromethane	Bromoform
January	27	5	3	6	7	1
February	< 1	5	< 1	6	, 6	< 1
March	< 1	57	75	4	2	< 1
April	236	38	< 1	4	2	< 1
May	< 1	9	9	7	5	< 1
June	< 1	7	< 1	10	13	6
July	5969	265	< 1	9	18	15
August	< 1	6	< 1	6	7	3
September	41	4	7	4	<b>5</b> .	, 7
October	< 1	4	6	4	6	6
November	< 1	2	1	4	5	4
December	11	10	12	9	18	13







education program regarding disposal of chemicals down laboratory drains continues to be a contributing factor to the reduction of volatile organics in the laboratory wastewater.

### Outfall 001

The treated wastewater streams from the two treatment systems are combined, chlorinated, and samples for analysis of most of the permit parameters are collected from a manhole downstream of the chlorine contact chamber. This combined effluent then flows through the outfall sewer to Sawmill Creek. The effluent travels through this sewer for approximately 0.8 mile before being discharged.

The permit requires analysis of the combined effluent once a week for TDS, chloride, and sulfate. The results, limits, and number of exceedances are presented in Table 5.5.

TABLE 5.5

Outfall 001 Monitoring Results and Effluent Limits, 1994

(Concentrations in mg/L)

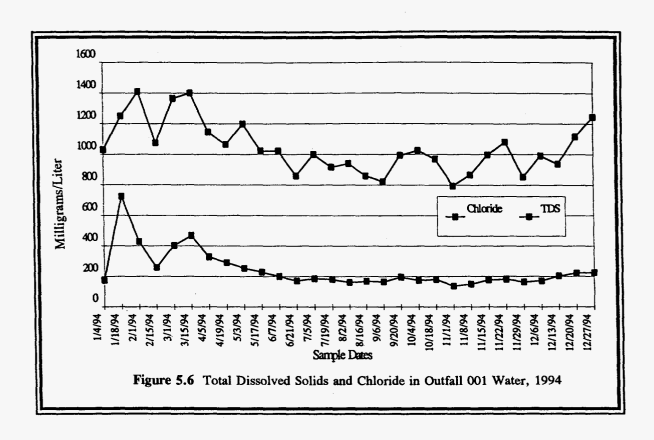
Constituent	Minimum	Average	Maximum	Limit	Exceedances
Copper	0.044	0.098	0.193	0.051*	8
Total Dissolved Solids	795	1057	1409	1045 1000*	8 3
Sulfates	171	236	287	575 500*	0 0
Chlorides	137	235	467	550 500*	0 0

<sup>\*</sup>Limit effective October 30, 1994.

The limit for TDS was exceeded 11 times in 1994. Frequent TDS exceedances (1-January, 2-February, 2-March, 2-April, 1-May) occurred early in 1994. A Task Force was formed to expeditiously determine the cause of the problem and recommend corrective actions. Through a strategic sampling and analysis program and review of operations at the facility where excess TDS appeared to originate, one major TDS source was identified and corrected in May 1994. This source involved a connection of the overflow piping from the recycle-brine tankage at the domestic water treatment plant, which permitted the brine overflow to enter the laboratory sewer system. After the recycle-brine overflow was redirected to the DuPage County sewer line, TDS peaks in the laboratory sewer wastewater disappeared and TDS values at Outfall 001 met the NPDES requirement (1045 ppm). TDS exceedances (3) that occurred after that time were believed to be related to discharges from boiler operations, i.e., boiler blowdown, which are known to contain high levels of TDS, and domestic water treatment. Chemical analysis for chloride shows a close relationship

between TDS levels and chloride levels. Figure 5.6 shows the results of TDS and chloride analyses for 1994. The groundwater at ANL is characterized by high TDS levels, i.e., approximately 800 ppm. This elevated concentration allows a narrow margin of added TDS (about 200-250 ppm) to the wastewater in meeting the new NPDES effluent standard of 1000 ppm (effective October 30, 1994). Levels for sulfate and chloride were not exceeded during 1994.

During the months of November and December, ANL exceeded the new (October 30, 1994) copper limit (0.051 mg/L Daily Maximum) in eight of nine samples. The well water obtained by ANL from the Niagaran dolomite has very low copper concentrations. However, after treatment and distribution through domestic water distribution copper piping, a copper concentration range of 0.5 mg/L to 1.0 mg/L is typical at drinking fountains. The



range has been determined by the lead/copper monitoring program (refer to Chapter 6) required by the EPA. The action level for copper in drinking water has been established at 1.3 mg/L. Essentially all the Laboratory monitoring locations are below this action level. These acceptable levels for human consumption are significantly above the NPDES permit limit at outfall 001. There is no process in the wastewater treatment plant to remove copper. Past samples collected from the wastewater treatment plant effluent have been below the IEPA effluent limit of 0.5 mg/L, but concentrations measured in Sawmill Creek, below the point where the treated wastewater has been discharged, have consistently exceeded the IEPA stream standard for copper (0.02 mg/L) for several years. This is indicative of the current ambient levels of copper in surface water as a result of the increased usage of copper pipe for domestic water distribution.

The permit requires that a biological toxicity screening test be performed on wastewater from outfall 001 in June of each year. The toxicity testing is run on at least three trophic levels of aquatic species for both chronic and acute toxicity. The 1994 testing was conducted on samples collected during the period June 2 to June 29, 1994, using the water flea, Ceriodaphnia dubia, the fathead minnow, Pimephales promelas, and the green alga, Selenastrum capricornutum. Testing was done in accordance with EPA protocols (USEPA, 1982, 1985, 1989) and the IEPA specifications given in "Effluent Biomonitoring and Toxicity Assessment - Aquatic Life Concerns."

No adverse effects were observed in any of the toxicity tests performed in any test concentration, including 100% effluent. Adverse effects which were monitored included lethality, decreased growth and decreased reproduction. It appears the cessation of chlorination of ANL's wastewater effluent has had a beneficial effect on aquatic life. Table 5.6 summarizes the results from the various toxicity tests for 1994. Table 5.7 summarizes the test results from 1991 to 1994.

TABLE 5.6

Outfall 001 Aquatic Toxicity Test Results, 1994

Test	Endpoint	96/48-Hour LC <sub>50</sub> *	7-Day NOEC**	7-Day LOEC***
96-Hour Fathead Minnow Acute Toxicity	Survival	100.0%	N/A	N/A
48-Hour Ceriodaphnia Acute Toxicity	Survival	100.0%	N/A	N/A
7-Day Fathead Minnow Chronic Toxicity	Survival Growth	N/A N/A	100.0% 100.0%	100.0% N/A
7-Day Ceriodaphnia Chronic Toxicity	Survival Reproduction	N/A N/A	100.0% 100.0%	100.0% N/A
96-Hour Algal Growth	Cell-Growth	N/A	100.0%	100.0%

<sup>\*</sup>LC<sub>50</sub> - Concentration of wastewater that produces 50% mortality in the test population.

<sup>\*\*</sup>NOEC - No Observable Effect Concentration is the highest concentration of the effluent at which no adverse effect is observed.

<sup>\*\*\*</sup>LOEC - Lowest Observable Effect Concentration is the lowest concentration of the effluent at which an adverse effect is observed.

TABLE 5.7

Aquatic Toxicity Test Results - 1991 to 1994

Test	1991	1992	1993	1994
Minnow, Acute, LC50	61.6%	< 6.2%	100.0%	100.0%
Ceriodaphnia, Acute, LC50	17.1%	35.4%	100.0%	100.0%
Minnow, Chronic, Survival, NOEC	50.0%	100.0%	50.0%	100.0%
Minnow, Chronic, Survival, LOEC	100.0%	100.0%	100.0%	100.0%
Minnow, Chronic, Growth, NOEC	50.0%	100.0%	50.0%	100.0%
Ceriodaphnia, Chronic, Survival, NOEC	50.0%	50.0%	50.0%	100.0%
Ceriodaphnia, Chronic, Survival, LOEC	100.0%	100.0%	100.0%	100.0%
Ceriodaphnia, Chronic, Reproduction, NOEC	50.0%	50.0%	25.0%	100.0%
Algal Growth, LOEC	6.2%	6.2%	100.0%	100.0%
Algal Growth, NOEC	3.1%	< 6.25%	100.0%	100.00%

The permit also requires that weekly pH, ammonia nitrogen, dissolved iron, manganese, and zinc measurements be made. There were no results outside the NPDES limits during 1994. Monthly monitoring for lead, hexavalent and trivalent chromium, and beta radioactivity is required.

### Outfall 003

This outfall is the discharge point from a series of small man-made ponds and is composed primarily of stormwater, with small amounts of process wastewater, such as cooling tower blowdown. It was sampled monthly through October 1994 and analyzed for pH, TSS, and temperature. Permit limits exist for TSS (15 mg/L average and 30 mg/L maximum), pH (between 6 and 9 pH units) and temperature (less than 5°F temp. rise). During 1994, there were no exceedances of the NPDES limits. This outfall is now inactive. For the outfalls 003 through 116, the number of samples collected, permit constituents, limits, and exceedances are shown in Table 5.8.

#### Outfall 003A

This discharge is located approximately 25 m (75 ft) north of the swimming pool, and is a vitrified clay pipe which was originally used as the discharge point for all the swimming pool activities (filter backwash, draining and overflow). The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There was one exceedance of the TSS limit in 1994. The effluent discharging directly from the pipe contains noticeable solids. The solids appear to be pipe material that is sloughing off the outfall pipe.

Total residual chlorine (TRC) levels exceeded the 0.05 mg/L TRC NPDES permit limit in eight of nine samples. Special Condition 8 of the NPDES permit allows ANL two years to achieve compliance with the 0.05 mg/L TRC limit. Therefore, IEPA does not consider

TABLE 5.8

NPDES Effluent Summary, 1994

Discharge ocation	Number of Samples Collected	Permit Constituent	Limit 30-Day Daily Average Maximum	Number Exceeding Limit
0031	10	Flow	None	0
		pН	6-9	0
		TSS	15 30	0
		Temperature	< 2.8°C Rise	0
003A <sup>2</sup>	5	Flow	None	0
		pН	6-9	0
		TSS	15 30	1
		TRC <sup>3</sup>	0.05	NA
$003B^2$	2	Flow	None	0
		pН	6-9	0
,		Temperature	< 2.8°C Rise	0
$003C^{2}$	2	Flow	None	. 0
		pН	6-9	0
$003D^{2}$	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0
$003E^{2}$	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0

# TABLE 5.8 (Contd.)

	Number		Limit	Number
Discharge Location	of Samples Collected	Permit Constituent	30-Day Daily Average Maximum	Exceeding Limit
$003F^{2}$	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0
		TDS	Monitor only	NA
$003G^{2}$	2	Flow	None	0
		рH	6-9	0
		Temperature	< 2.8°C Rise	0
003H <sup>2</sup>	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0
		TDS	Monitor only	NA
$003I^{2}$	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0
		TDS	Monitor only	NA
		Oil & Grease	Monitor only	NA
$003J^{2}$	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0
		TDS	Monitor only	NA

TABLE 5.8 (Contd.)

1	1					
	Number Exceeding Limit	0 0 1 0	0000	0 0 0 <b>V</b>	0 0	0 0 0 V
	20-Day Daily Average Maximum	None 6-9 15 30 < 2.8°C Rise	None 6-9 < 2.8°C Rise 15	None 6-9 < 2.8°C Rise Monitor only	None 6-9	None 6-9 30 Monitor only < 2.8°C Rise
	Permit Constituent	Flow pH TSS Temperature⁴	Flow pH Temperature Oil & Grease	Flow pH Temperature Oil & Grease	Flow pH	Flow pH TSS TDS <sup>2</sup> Temperature <sup>2</sup>
	Number of Samples Collected	12	10	2	2	12
	Discharge Location	900	0051	005C <sup>2</sup>	$005\mathrm{E}^2$	900

TABLE 5.8 (Contd.)

	Number		Lim	nit	Number
Discharge Location	of Samples Collected	Permit Constituent	30-Day Average	Daily Maximum	Exceeding Limit
007	7	Flow	No	one	0
		pН	6-9	)	0
		Temperature	< 2	.8°C Rise	0
		$TRC^{2}$	0.0	)5	NA
		Oil & Grease <sup>2</sup>	Mon	itor only	NA
008	10	Flow	No	one	0
		pН	6-9	)	0
		VOC <sup>2</sup>	Mon	itor only	NA
$009^{1}$	0	Flow	None		0
		рH	6-9	)	0
		TSS	15	30	. 0
010	0	Flow	No	ne	NA
		рН	6-9	)	NA
*		TSS	15	30	NA
		Total Iron	2	4	NA
		Dissolved Iron <sup>2</sup>		1.0	NA
		Lead		0.1	NA
		Zinc		1.0	NA
		Manganese		1.0	NA
		Hexavalent Chromium <sup>2</sup>	0.011	0.016	NA
		Trivalent Chromium <sup>2</sup>	0.519	2.0	NA
		Copper	0.031	0.051	NA
		Oil & Grease	15	30	NA

TABLE 5.8 (Contd.)

Discharge Location	Number of Samples Collected	Permit Constituent	Limit 30-Day Daily Average Maximum	Number Exceeding Limit
_			·.	
$108^2$	2	Flow	None	0
		pН	6-9	0
		Temperature	< 2.8°C Rise	0
$111^{2}$	0	Flow	None	NA
		Tritium	Monitor Only	NA
112A <sup>2</sup>	0	Flow	None	NA
		Tritium	Monitor Only	NA
112B <sup>2</sup>	0	Flow	None	NA.
	_	Tritium	Monitor Only	NA
113 <sup>2</sup>	2	Flow	None	0
		Tritium	Monitor Only	NA
		PCB 1260	Monitor Only	NA
		Lead, Copper, Nickel, Zinc	Monitor Only	NA
114 <sup>2</sup>	2	Flow	None	0
		Tritium	Monitor Only	NA
		PCB 1260	Monitor Only	NA
		Lead, Copper, Nickel, Zinc	Monitor Only	NA

Number			Limit		Number
Discharge Location	of Samples Collected	Permit Constituent	30-Day Average	Daily Maximum	Exceeding Limit
115 <sup>2</sup>	2	Flow	No	ne	0
113	-	pH Temperature	6-9		0
		TDS		itor Only	NA
$116^{2}$	2	Flow	No	ne	0
		рH	6-9	)	0
		pH TRC³	0.0	)5	NA

<sup>&</sup>lt;sup>1</sup>Inactive effective October 30, 1994.

<sup>&</sup>lt;sup>2</sup>Added October 30, 1994.

<sup>3</sup>NPDES Special Condition allows ANL two years to comply with TRC limit.

<sup>4</sup>Removed October 30, 1994.

an exceedance of the TRC limit to be "reportable" until October 30, 1996. The present flow at 003A appears to be coming from a manhole which is probably receiving water from a leaking domestic water main. Corrective actions are underway to stop the incursion of chlorinated water to outfall 003A. Also, a "scupper" drain system from the swimming pool drains to this outfall. This will be rerouted during Spring 1995.

#### Outfall 003B

The outfall is located approximately 150 m (500 ft) northeast of Building 308, and is composed of storm water run-off and condensate from the buildings in the watershed of the outfall. The discharge point is a 1 m (36 in) concrete pipe to a tributary brook flowing north to the Freund Brook. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 003C

The discharge from this outfall is made up of footing tile drainage and storm water runoff. The discharge point is a 0.65 m (24 in) concrete pipe discharging into Freund Brook approximately 50 meters upstream of the gas station, south of Building 205. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfalls 003D and 003E

These two discharge points are from the steam trench around Inner Circle Drive, and discharge into the north fork of Freund Brook approximately 150 m (500 ft) east of the intersection of Inner Circle Drive and Eastwood Extension. The sampling requirements and

effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 003F

This outfall is intended to discharge excess water from the fire pond during storm events. The building discharges cooling tower water to the fire pond, and the rate is low enough to generally not discharge without rain water to attain flow. The discharge is through a cement raceway to the south fork of the north branch of Freund Brook. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 003G

Footing tile drainage from the Inner Circle steam trench is pumped to the storm sewer passing around the northeast portion of Building 201 and discharging into the northern fork of the southern branch of Freund Brook. Condensate leaks in the steam trench produce discharge on a regular basis to the storm sewer. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 003H

This discharge originates from the footing tile drainage around Building 212 and storm water collected from around Building 212 and 214 and their associated parking lots. The cooling tower located on the south roof of Building 212 discharges into the tile drainage system, and is the source of the industrial discharge. The sampling requirements and

effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 0031

This outfall collects stormwater from the Building 200, 211, and western portion of Building 205 areas and also gets cooling tower discharge from the cooling tower located behind Building 200. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 003J

This outfall collects stormwater from the Building 213 area and parking lot, which passes through a storm sewer around the Building 201. Cooling tower blowdown is the industrial discharge to this system. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 004

This outfall discharges stormwater from the Building 203 and 221 areas and cooling water from Building 221. The discharge is to a drainage ditch and sewer system, passing around the northeastern portion of Outer Circle Drive and to a ditch leading north to the fenceline, east of the Visitor's Center. The sampling requirements and effluent limits are in Table 5.8. There was one exceedance of TSS limits in 1994, most likely due to surface runoff from the surrounding area during periods of heavy precipitation.

#### Outfall 005

This outfall consisted of stormwater and process wastewater from the Building 206 cooling system and the 800 Area, which includes vehicle and other maintenance areas. The permit requirements include monthly sampling and analyses for oil and grease, pH, and temperature. Limits of 15 mg/L average and 30 mg/L maximum exist for oil and grease. The pH and TSS limits are the same as for outfall 003. There were no exceedances in 1994. Effective October 30, 1994, this outfall is inactive.

#### Outfall 005A

This outfall discharges runoff from the northwestern portion of the 800 Area. The flow passes under Westgate Road, east of the west gate and flows toward the northwestern fenceline. This is a stormwater only outfall.

#### Outfall 005B

The outfall for this watershed discharges runoff collected from the majority portion of the 800 Area. The flow is collected from the parking lots and roadways and flows by storm sewers to the east, where it is discharged to the marsh located on the eastern side of Kearney Road. This is a stormwater only outfall.

#### Outfall 005C

This outfall collects stormwater from the northern side and the loading dock area of Building 200. The Building 200 clear water system discharges to this outfall, which passes through sewers to the west of the loading dock, and to the beaver pond west of Building

200. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 005D

The Building 200 M-Wing loading dock area stormwater runoff is collected in a storm sewer and passes west to the Beaver pond located west of Building 200. The discharge is through a 1 m (36 in) corrugated pipe into the pond. This is a stormwater only discharge.

### Outfall 005E

This outfall discharges footing tile drainage from Building 203 west and Building 208. The outfall also discharges stormwater collected from the same area. The industrial discharge arises from cup drains and compressors discharging into the footing tile sumps. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 006

Cooling towers at Building 350 and the 377 Area discharge into the drainage ditch flowing south of the Canal Water Treatment Plant, bending south and flowing to the south fenceline. The permit requires monthly sampling for pH, TSS, and temperature. The limits are in Table 5.8. In 1994, there were no exceedances of the NPDES limits.

### Outfall 007

The watershed for outfall 007 includes the southeastern section of the 300 Area, extending from Building 370 east to Building 366, and north to Building 367. Water is

collected in catchment basins and conveyed toward the southeast to a point approximately 30 m (100 ft) southeast of Building 366, where it is discharged into a ditch on the south side of Old Bluff Road. This ditch runs along the roadside for 15 m (50 ft), at which point it turns south and runs to the fenceline where it is discharge to the forest preserve. Compressors' once-through cooling water is the industrial component to this outfall.

The sampling requirements and effluent limits are in Table 5.8. TRC levels exceeded the NPDES permit limit of 0.05 mg/L in 6 out of 9 samples collected during November and December 1994. For reasons explained above for outfall 003A, these are not "reportable" exceedances to the IEPA. Apparently some discharge of potable water from the 300 Area flows through this outfall. The source of this flow has not yet been determined but it is apparent that there is a cross-connect of a potable drain to the stormwater collection system.

#### Outfall 008

The watershed for this outfall includes the area around the new Vehicle Maintenance and Grounds Building 46. Runoff is collected in stormwater grates and catchments and conveyed through sewers to the discharge point in Sawmill Creek, located directly west of Building 24. Industrial activity in this small watershed involves the activities associated with maintenance of all facility vehicles, grounds, maintenance, the storage of the equipment associated with these activities, and fueling for the vehicles.

Five volatile organic compounds are monitored once a month, effective October 30, 1994. pH is the only NPDES limit that applies at this point. During 1994, no exceedances were noted.

#### Outfall 009

This outfall was an emergency overflow for an inactive lime sludge lagoon near the domestic water treatment plant. This lagoon has not been used since 1986. In the event that an extremely heavy storms occurs, rainwater could flow out of this outlet. The permit contains limits for pH and TSS, as shown in Table 5.8. The permit required monitoring monthly, when discharge was occurring. There is no discharge during 1994. This outfall is inactive effective October 30, 1994.

#### Outfall 010

This outfall is the outfall for the coal pile storage area runoff collection system over-flow line. The collection system consists of a trench on the north and west sides of the coal pile, with a sump located at the extreme southern end of the western trenchline. The overflow line comes into use only when the runoff reaches the level at which the trench system would overflow, and was put into place to insure against overflow conditions. During normal operations, the water is pumped to the equalization basin located in the western area of the 200 Area. The industrial activity associated with this outfall is solely the coal pile operation. The berm and trench system in place to collect runoff has been improved to eliminate discharge from the outfall.

This outfall is sampled once per day when flow occurs. Analyses are performed for pH, TSS, TDS, iron, lead, zinc, manganese, trivalent and hexavalent chromium, copper, and oil and grease. No flow occurred at this site during 1994.

#### Outfall 101

The drainage to the outfall is through ditches along the streets, and sewer conduits from the parking lot to a marsh located between Outer Circle Drive and the fenceline, to the outfall, which consists of a 0.65 m (24 in) corrugated metal pipe with a Palmer-Bowlus flume. This then discharges on the other side of the fenceline into the forest preserve. The sources of stormwater runoff to the outfall are the Building 203 parking lot with loading dock and the excess equipment storage area on the north side of Outer Circle Drive. This is a stormwater only discharge.

#### Outfall 102

This watershed includes portions of the 100 Area. There are large amounts of paved area involved with the industrial activities for the production of steam such as those areas associated with the water treatment plant, the lime sludge pond, and the tarmac around the boiler house. The contributing runoff flows are collected from stormwater inlet grates and catch basins, through storm sewers to a discharge point of a 0.30 m (12 in) corrugated metal pipe extending out of the bank of Sawmill Creek. This is a stormwater only discharge.

### Outfall 103

The watershed for outfall 103 includes the southern and southeastern extreme of the 100 Area and the area south of the coal pile. These areas drain into a storm sewer which runs due east of the coal pile towards Sawmill Creek. The outfall is located at the outlet of a 0.35 m (15 in) corrugated metal pipe culvert located approximately 50 m (150 ft) from the creek. Activities which are industrial in nature take place in and around the Utilities area, and consist of boiler house steam generation, storage of plastic and metal, loading dock activities, a flue gas scrubber and cooling pond (no longer in use), steam condensate

return storage (two tanks), and the southern access road to the coal pile storage area. This is a stormwater only discharge.

#### Outfall 104

This outfall includes the buildings and parking areas remaining in the East Area, excluding Buildings 40 and 46. This includes Buildings 4, 5, 6, and their smaller attendant buildings, and Buildings 24 and 25. The area is served by a number of roadways leading to and from these buildings, with contributing storm grate inlets on the roadways and parking areas. This is a stormwater only discharge.

#### Outfalls 105A and 105B

There are two discharge points within this watershed. The contributing sources of storm water with this watershed receive runoff from areas around Building 40, and elevated water tower tanks for potable water distribution, and scrub vegetation areas on the west side of Tech Road. Industrial activity within this watershed includes receiving, loading, parking and storage areas, and oil-containing transformers. These are stormwater only discharges.

### Outfalls 106A and 106B

The watershed for these outfalls encompasses the largest portion of the East Area, most of which is now demolished, with the buildings razed. There is a portion of the eastern end of the Shipping and Receiving Area which is part of this watershed, Building 33, which has electrical transformers located outside of it, and a portion of Argonne Park is within this watershed. As with outfall 105 above, this watershed is served by two distinct outfalls. The industrial activity within this watershed are the receiving and shipping areas with loading docks, and the transformer area. These are stormwater only discharges.

#### Outfall 108

This watershed encompasses a portion of the 300 Area. The drainage area includes the parking areas north of Building 360, the buildings in and around Building 360, excluding Buildings 370 and 390 and the southern and western end of the 300 Area, and the paved parking and loading dock areas in and around the eastern portions of the 300 Area (surrounding Building 363). The industrial activities ongoing in this watershed are shipping and receiving, a metals reclaim dumpster (Building 363), loading dock activities and numerous outdoor equipment storage areas. The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 110

The watershed for this outfall includes the shooting range (inactive since March 1993) and the area just south of the range. There are no other industrial activities taking place within this watershed at present. Past industrial activity involved use of the shooting range for practice by the security force. This is a stormwater only discharge.

#### Outfall 111

This outfall is located on the south fenceline of the site due south of the old, closed 319 Area Landfill, between the watershed for outfall 110 and the watershed for outfalls 112A and 112B. This watershed encompasses the 319 Landfill, the 318 Area (landfill area for compressed gases), and portions of the 317 Area, primarily the paved area. As well, the roadways for access to these areas drain to this outfall through a small ditch running along the southern extreme of the 319 Landfill, turning south to the fenceline, and then to the outfall location, which is a 0.65 m (24 in) corrugated metal pipe culvert which passes under the fence and discharges into the forest preserve. Industrial activities taking place within

this watershed are the 317 Area radioactive waste storage and remediation activities, the 319 Landfill Area, and the associated roadways for access. This outfall is sampled semiannually and has no permit limits. Sampling from this outfall will commence during 1995.

#### Outfalls 112A and 112B

The contributing sources of stormwater within this watershed receive runoff from the southern and western sections of the 317 Area radioactive waste storage. Runoff flow is generally towards the south in sheet flow from the source areas, with the eastern portions consolidating at the fenceline at the southeastern corner of the 317 Area to pass under the fence through rough concrete fill. The western and central portions of the drainage area sheet flow consolidate in the same manner and pass under the fence through the same material approximately 50 m to the west. Both flows discharge into large gullies in the forest preserve, to conjoin into one flow approximately 100 m south of the ANL fenceline. Industrial activity within this watershed are the 317 Area radioactive waste storage and remediation activities, loading activities at Building 350, and the associated roadways for access. These outfalls are sampled semiannually and have no permit limits. Sampling from these outfalls will commence during 1995.

# Outfall 113

This outfall is the discharge point for runoff from the eastern, southern, and southwestern sections of the closed 800 Area Landfill. The outfall is located in a ditch on the extreme southern end of the landfill, approximately 50 m from the southwestern corner of the landfill fenceline. This discharge flows under the fence in the ditch, and empties into the creek which flows south from the wetland marsh west of the site. The marsh is the headwaters of one leg of the Freund Brood system which runs through the middle of the ANL site and discharges into Sawmill Creek. Industrial activity within this watershed is

limited to the landfill. Beginning October 30, 1994, this outfall was sampled monthly and has no permit limits.

#### **Outfall 114**

This outfall is the discharge point for runoff coming from the northern and northwestern sections of the closed 800 Area Landfill. The outfall is located in a ditch on the extreme western side of the landfill, approximately halfway between the northern and southern boundaries of the landfill. The flow proceeds along the western edge of the landfill discharging into a small marsh located on the western side of landfill. The water from the marsh flows south through another ditch, which eventually combines with the ditch from the outfall 113 flow, and then into the creek which flows south from the wetland marsh west of the ANL site. Industrial activity within this watershed is limited to the landfill. Beginning October 30, 1994, this outfall was sampled monthly and has no permit limits.

### Outfall 115

The watershed encompasses the Advanced Photon Source (APS) construction site and the southern areas around the Building 314, 315, and 316 complex. The APS flow drains into ditches that discharge through a cement culvert into a collection pond located on the southeastern portion of the APS site. The 0.65 m (24 in) sewer conduit from the Building 314, 315, and 316 complex discharges into the same collection pond approximately 10 m (30 ft) east of the ditch culvert. The flow from this pond discharges south through a culvert into another pond, flowing through this pond and discharging through a 1 m (36 in) corrugated metal pipe culvert under the south fenceline into the forest preserve. Industrial activities within the watershed are the APS construction, all roadways associated with APS construction, loading docks in the APS buildings, and the Building 314, 315, and 316 complex storage, loading areas, and cooling water discharges. The sampling requirements

and effluent limits (effective October 30, 1994) are in Table 5.8. There were no exceedances during 1994.

#### Outfall 116

This outfall was originally intended as a stormwater discharge point only. It has been determined to contain non-storm water discharge as well. The source of the discharge was traced back and found to be potable water from the water treatment plant located uphill from the main Utilities area. This source will be investigated for corrective action and the flow stopped. The watershed for this outfall contains sections of the water treatment plant, including the garage and storage area, the area around well No. 5, and the associated access roads for the water treatment plant. Flow is conducted through stormwater sewers and discharged at the outfall, which is a 0.25 m (10 in) vitrified clay pipe with a cement raceway, into Sawmill Creek. Industrial activities for this watershed include parking, loading and materials storage around the water treatment plant, the water treatment plant operation including bulk chemical storage (brine tank), and transformers (Building 129), outdoor equipment storage area and four flammable materials storage cabinets (Building 130), outdoor materials storage (Buildings 107 and 163), well operation and maintenance (Building 160), and the associated roadways for these activities.

The sampling requirements and effluent limits (effective October 30, 1994) are in Table 5.8. TRC levels exceeded the NPDES permit limit of 0.05 mg/L in two samples collected in November and December 1994. For reasons explained above under outfall 003A, no reportable exceedance occurred. The source of chlorinated potable water was identified as the drain system from the vertical ground tank which contains the clear water from the domestic water treatment plant. The leakage was related to the rehabilitation of the tank. In order to supply water to the system, water was pumped under very high pressure into the line in which the valve is located. The high pressure reduced the valve's ability to hold the

water and the water leaked through to the drain system, discharged to the closest stormwater collection basin, discharging at outfall 116.

# 5.2. Additional Effluent Monitoring

To characterize the wastewater from the ANL site more fully, composite samples of the combined effluent are collected each week and analyzed for the constituents shown in Table 5.9. The results are then compared to the IEPA General Effluent Limits found in 35 IAC, Subtitle C, Part 304.<sup>20</sup>

### 5.2.1. Sample Collection

Samples for analysis of inorganic constituents are collected daily from outfall 001 located at the Wastewater Treatment Plant using a refrigerated time proportional sampler. A portion of the sample is transferred to a specially cleaned bottle, a security seal is affixed and chain-of-custody is maintained. Five daily samples are composited on an equal volume basis to produce a weekly sample, which is then analyzed.

#### 5.2.2. Results

Fifteen metals were determined by inductively coupled plasma emission spectroscopy, flame atomic absorption spectroscopy, and graphite furnace atomic absorption spectroscopy. Mercury was analyzed using cold vapor atomic absorption spectroscopy. Fluoride was determined by a specific ion electrode. The results for 1994 appear in Table 5.9. None of the annual average results or maximum results exceeded General Effluent Limits. Higher maximum concentrations were noted for barium, chromium, cobalt, and nickel than in previous years, but within the General Effluent Limits.<sup>20</sup>

TABLE 5.9

Chemical Constituents in Effluents from ANL Wastewater Treatment Plant, 1994

(Concentrations in mg/L)

Constituent	No. of	on			
	Samples	Avg.	Min.	Max.	Limit
Arsenic	52	0.0026	< 0.0020	0.0043	0.25
Barium	52	0.0673	0.0500	0.5000	2.0
Beryllium	52	-	-	< 0.0002	-
Cadmium	52	0.0002	< 0.0001	0.0005	0.15
Chromium	52	0.0926	< 0.0200	0.2000	1.0
Cobalt	52	0.0553	< 0.0200	0.1000	-
Copper	52	0.0647	0.0320	0.1270	0.5
Fluoride	29	0.4597	0.2940	0.7640	15.0
Iron	52	0.2249	< 0.0400	0.4680	2.0
Lead	52	0.0031	< 0.0010	0.0054	0.2
Manganese	52	0.0399	< 0.0100	0.0840	1.0
Mercury	52	0.0008	< 0.0001	0.3150	0.5
Nickel	52	0.0996	< 0.0200	0.2000	1.0
Silver	52	0.0015	< 0.0010	0.0072	0.1
Thallium	52	0.0030	< 0.0030	0.0033	-
Vanadium	52	-	-	< 0.2000	-
Zinc	52	0.0576	0.0250	0.0880	1.0
pH (units)	52	-	7.23	8.50	6.0-9.0

### 5.3. Sawmill Creek

Sawmill Creek is a small natural stream that is fed primarily by stormwater runoff. During periods of low precipitation, the creek above ANL has a very low flow. At these times, a major portion of the water in Sawmill Creek south of the site consists of ANL wastewater and discharges to assorted storm drains. To determine the impact ANL wastewaters have on Sawmill Creek, samples of the creek downstream of all ANL discharge points are collected and analyzed. The results are then compared to the IEPA General Use Water Quality Standards found in 35 IAC, Subtitle C, Part 302.<sup>21</sup>

# 5.3.1. Sample Collection

A proportional sampler is used to collect a daily sample at a point well downstream of the combined wastewater discharge point where thorough mixing of the ANL effluent and Sawmill Creek water is assured. Samples are collected in precleaned, labelled bottles and security seals are used. After pH measurement, the daily samples are acidified and then combined into equal volume weekly composites and analyzed for the same set of inorganic constituents analyzed in the wastewater described in Table 5.10.

Fifteen metals were determined by inductively coupled plasma emission spectroscopy, flame atomic absorption spectroscopy, and graphite furnace atomic absorption spectroscopy. Mercury was analyzed using cold water atomic absorption spectroscopy. Fluoride was determined by a specific ion electrode.

TABLE 5.10

Chemical Constituents in Sawmill Creek, Location 7M,\* 1994

(Concentrations in mg/L)

	No. of Concentrations						
Constituent	Samples	Avg.	Min.	Max.	Limit		
Arsenic	50	0.0025	< 0.0020	0.0048	1.0		
Barium	50	0.0531	0.0500	0.0830	5.0		
Beryllium	50	-	-	< 0.0002	-		
Cadmium	50	0.0004	< 0.0001	0.0039	0.05		
Chromium	50	0.0884	< 0.0200	0.2000	1.0		
Cobalt	50	0.0536	< 0.0200	0.1000	-		
Copper	50	0.0416	0.0110	0.1800	0.02		
Fluoride	27	0.3537	0.1760	0.7800	1.4		
Iron	50	0.4108	0.1650	1.5830	1.0		
Lead	50	0.0064	0.0013	0.1047	0.1		
Manganese	50	0.0911	0.0160	0.3580	1.0		
Mercury	50	0.2001	< 0.0001	10.0000	0.5		
Nickel	50	0.0957	< 0.0200	0.2000	1.0		
Silver	50	0.0010	< 0.0010	0.0023	0.005		
Thallium	50	0.0035	< 0.0030	0.0300	-		
Vanadium	50	-	-	< 0.2000	-		
Zinc	50	0.1134	0.0300	1.8000	1.0		
pH (Units)	50	-	7.51	8.47	6.5-9.0		

<sup>\*</sup> Location 7M is 15 m (50 ft) downstream from the ANL wastewater outfall

### 5. ENVIRONMENTAL NONRADIOLOGICAL PROGRAM INFORMATION

### 5.3.2. Results

The results obtained are shown in Table 5.10. As in previous years, the annual average concentration for copper was above the Water Quality Standards (WQS). The maximum concentrations for copper, iron, mercury, and zinc exceeded the WQS. A possible explanation for these elevated levels may be due to the release of constituents during the sewer rehabilitation project. However, since these elevated levels were not noted at Outfall 001, further investigation would be needed.

### 5.4. Des Plaines River

Based on previous sampling results, it was determined that mercury would be the only element likely to have a measurable impact on the Des Plaines River. During previous years, the effect of Sawmill Creek on the levels of mercury in the Des Plaines River was evaluated by collecting samples in the river at Willow Springs (upstream of ANL) and at Lemont (downstream of ANL). All of the samples analyzed showed that the total mercury concentration was less than the detection limit of  $0.1~\mu g/L$ . Based on these consistently low results, the decision was made to remove this specific monitoring from the ANL program during 1993.



The groundwater below the ANL site is monitored through the collection and analysis of samples obtained from the on-site water supply wells and from a series of groundwater monitoring wells located near several sites which have the potential for causing groundwater impact. Federal and state drinking water regulations are used to evaluate the quality of groundwater used for human consumption at ANL. Regulations establishing comprehensive water quality standards for the protection of groundwater have been enacted, IEPA Groundwater Quality Standards, 35 IAC, Subtitle F, Part 620.<sup>22</sup> In addition, compliance with the groundwater protection requirements in DOE Order 5400.1, as related to sitewide characterization studies and monitoring well requirements, are demonstrated in this Chapter. The permit for the 800 Area landfill requires a groundwater monitoring program and this was initiated during July 1992.

# 6.1. Potable Water System

The ANL domestic water is supplied by four wells. The wells are described in Section 1.7 and Table 6.1. Their locations are shown in Figure 1.1. According to the National Primary Drinking Water Regulations, <sup>16</sup> ANL's system is classified as a non-transient, non-community public water system, since it regularly serves at least 25 of the same persons over six months of the year. This designation determines the parameters to be monitored and the frequency of monitoring. Monitoring of the ANL domestic water supply is conducted to demonstrate compliance with applicable regulations and to obtain information on the concentrations of other constituents.

# 6.1.1. Regulatory Required Monitoring

The primary regulations that apply to ANL are the Illinois Department of Public Health, *Drinking Water System Code* 77 IAC Part 900.<sup>23</sup> These regulations identify the

inorganic (900.50) and organic (900.65) constituents that require monitoring and set the State limits. In addition, ANL must also demonstrate compliance with 40 CFR 141.40 of the National Primary Drinking Water Regulations<sup>16</sup> by conducting the Special Monitoring for Organic Chemicals.

TABLE 6.1

ANL Water Supply Wells

Well No.	Location	Well Elevation <sup>1</sup>	Pumping Level	Bedrock Elevation	Well Depth <sup>2</sup>	Diameter <sup>3</sup>	Year <sup>4</sup>
1	Building 31	671	~ 613	605	284	12	1948
2	Building 32	664	~ 600	601	300	12	1948
3	Building 163	689	~ 597	600	318	12	1955
4	Building 264	716	~ 608	595	340	14	1959

<sup>&</sup>lt;sup>1</sup>Feet mean sea level.

All chemical analyses were performed by a commercial laboratory which is certified by the State of Illinois to conduct Safe Drinking Water Act analyses. Lead was determined by graphite furnace atomic absorption spectroscopy. Copper was determined by inductively coupled plasma atomic emission spectroscopy. Volatile organic compounds were determined by using a purge and trap sample pretreatment followed by gas chromatography-mass spectroscopy detection. Nitrate/nitrite was analyzed using a colorimetric technique. Samples were collected quarterly from each of the four ANL domestic wells and a treated tap water sample in Building 128. The samples were analyzed for nitrate/nitrite, metals,

<sup>&</sup>lt;sup>2</sup>Feet below ground.

<sup>&</sup>lt;sup>3</sup>Inner diameter (inches).

<sup>&</sup>lt;sup>4</sup>Year drilled.

volatile organic compounds, pesticides, and herbicides by a commercial laboratory which is certified to conduct Safe Drinking Water Act analyses. The samples were analyzed for the constituents specified in the regulations by approved methods which allowed the minimum detectable limit of 0.0005 mg/L to be met for the organic chemicals. The results were provided to the DuPage County Health Department and the Illinois Department of Public Health (IDPH).

On March 17, 1993, the IDPH provided ANL with a permanent waiver for sampling for asbestos and an extension to November 24, 1995, for the testing of inorganics/metals. On July 22, 1993, ANL submitted seven quarters of organic data and petitioned the IDPH for sampling/analysis waivers from the requirements. On August 6, 1993, the IDPH approved a waiver for the 18 Phase II Volatile Organic Compounds to November 24, 1998, a waiver for Synthetic Organic Chemicals/Herbicides-Pesticides to November 24, 1995, and a permanent waiver for all future sampling for unregulated chemicals. The federal organic list in 40 CFR 141.40 is covered under the Phase II organics and the unregulated list and, therefore, future sampling and analysis of these compounds is covered. In addition, future sampling is only required at a representative tap, however, well head sampling will be continued for information monitoring of radionuclides and volatile organic compounds. Because of the waivers, no analyses were required in 1994 for metals, volatile organics, synthetic organics, or the federal organic list.

On June 7, 1991, the EPA promulgated final rules establishing National Primary Drinking Water Regulations for lead and copper. The regulations require collection of finished water samples for lead and copper analyses at selected sites and to determine if the concentrations are below the action level of 0.015 mg/L for lead and 1.3 mg/L for copper. The required sampling protocols maximizes the opportunity for having lead and copper present. Sampling locations are determined after a water piping materials survey. Priority sampling locations are those that have lead pipes, are served by lead service lines, or have

copper pipes with lead solder joints. Samples must be first draw water where the water has stood motionless in the piping for at least six hours. Because of compliance with the lead and copper limits in 1993, only annual sampling was required in 1994 at 20 sites.

Samples were collected, following the above protocols, on June 15, 1994, analyzed by a laboratory certified to conduct Safe Drinking Water Act analyses, and transmitted to the Illinois Department of Public Health, through DOE, on January 3, 1995. The results indicated three of the 20 lead results exceeded the action level and three of the 20 copper samples exceeded the action level. The results are collected in Table 6.2. Since both action levels were exceeded expanded sampling will be required in 1995.

### 6.1.2. Informational Monitoring

Samples were collected quarterly at the wellhead. These samples were analyzed for several types of radioactive constituents and volatile organic compounds to determine their presence in the ANL drinking water. Samples from each well were tested for total alpha, total beta, hydrogen-3 and strontium-90. Annually, samples were also analyzed for radium-226, radium-228, and isotopic uranium. Alpha and beta radioactivity were determined by a gas flow proportional counting technique. Hydrogen-3 was determined by distillation followed by a beta liquid scintillation counting technique. Strontium-90 and isotopic uranium were performed by ion exchange separations followed by proportional counting and alpha spectrometry techniques, respectively. The results are presented in Table 6.3. Since ANL is a "non-transient, non-community" water system, the following EPA limits are established for the nuclides measured in Table 6.3:

Gross Alpha Particle Activity = 15 pCi/L

Gross Beta Particle Activity = 50 pCi/L

Hydrogen-3 = 2 x 10<sup>4</sup> pCi/L

Radium-226 = 5 pCi/L

Strontium-90 = 8 pCi/L

TABLE 6.2

Lead/Copper Samples Collected June 15, 1994

(Concentrations in mg/L)

	Lead	Copper
1.	< 0.005 (lowest)	0.02 (lowest)
2.	< 0.005	0.1
3.	< 0.005	0.2
4.	< 0.005	0.2
5.	< 0.005	0.3
6.	< 0.005	0.5
7.	< 0.005	0.5
8.	< 0.005	0.6
9.	< 0.005	0.6
10.	0.005	0.7
11.	0.006	0.8
12.	0.007	0.8
13.	0.008	0.9
14.	0.008	0.9
15.	0.009	1.1
16.	0.009	1.1
17.	0.014	1.2
18.	0.016 (90th Percentile)	1.4 (90th Percentile)
19.	0.017	2.0
20.	0.020	3.3
Action Level	0.015	1.3

TABLE 6.3

Radioactivity in ANL Domestic Wells, 1994

(Concentrations in pCi/L)

Type of Activity	Location	No. of Samples	Avg.	Min.	Max.
Alpha (nonvolatile)	Well #1 Well #2 Well #3 Well #4 Tap	4 4 4 4	10.6 5.8 4.5 3.7 1.4	7.1 4.7 3.6 2.6 0.7	14.7 7.2 5.1 4.8 2.2
Beta (nonvolatile)	Well #1 Well #2 Well #3 Well #4 Tap	4 4 4 4	16.7 10.0 11.6 10.4 5.1	14.2 8.8 11.1 9.0 4.6	22.4 10.7 12.4 11.5 5.5
Hydrogen-3	Well #1 Well #2 Well #3 Well #4 Tap	4 4 4 4	< 100 < 100 < 100 < 100 < 100	< 100 < 100 < 100 < 100 < 100	140 < 100 < 100 < 100 < 100
Strontium-90	Well #1 Well #2 Well #3 Well #4 Tap	4 4 4 4	< 0.25 < 0.25 < 0.25 < 0.25 < 0.25	< 0.25 < 0.25 < 0.25 < 0.25 < 0.25	0.28 < 0.25 < 0.25 < 0.25 < 0.25
Radium-226	Well #1 Well #2 Well #3 Well #4 Tap	1 1 1 1	- - - -	: : :	0.80 1.10 1.03 1.18 0.40
Radium-228	Well #1 Well #2 Well #3 Well #4 Tap	1 1 1 1	- - - -	- - - - -	0.56 0.86 0.92 0.68 0.27
Uranium-234	Well #1 Well #2 Well #3 Well #4 Tap	1 1 1 1	: : :	- - - - -	3.31 0.68 0.17 0.25 0.25
Uranium-235	Well #1 Well #2 Well #3 Well #4 Tap	1 1 1 1	-	-	0.12 0.02 < 0.01 < 0.01 < 0.01
Uranium-238	Well #1 Well #2 Well #3 Well #4 Tap	1 1 1 1	- - - -	- - -	2.29 0.45 0.13 0.16 0.15

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Well #1 was removed from service in 1990 and the system was not operated during 1994; however, samples were collected for monitoring. All the radiological results are in the normal range of concentrations for the various constituents and wells below the EPA drinking water standards except for strontium-90 and isotopic uranium in Well #1. Since hydrogen-3 had been identified in Well #1 in the past, these constitutes may be indicators of continued migration from the same source.

Although volatile organic compounds were not required to demonstrate compliance with the Drinking Water Regulations for 1994, the results are included in the informational monitoring section to determine if any past disposal practices have resulted in groundwater contamination and to support the environmental restoration program. Samples were collected on quarterly and the results are collected in Table 6.4 through Table 6.7, respectively. Samples were analyzed for the SDWA volatile compounds and quantified by EPA Method 524.2. The limit of detection reported in the tables is the practical quantification limit which is defined as ten times the method detection limit.

No volatile organic compounds were detected in any of the wells. The tap water samples showed four volatile organic compounds (dichlorobromomethane, bromoform, chlorodibromomethane, and chloroform) present. These compounds are known to be associated with chlorination of drinking water, i.e., trihalomethanes.

# 6.2. Groundwater Monitoring at Waste Management Sites

ANL has occupied its current site since 1948. Since that time, waste generated by the Laboratory had been placed in a number of on-site disposal units ranging from ditches filled with construction and demolition debris during the 1950s to a modern sanitary landfill used for nonhazardous solid waste disposal until September 1992. Several of these units

TABLE 6.4

Volatile Organic Compounds in Drinking Water Collected February 10, 1994

(Concentrations in mg/L)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
Benzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Vinyl Chloride	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Carbon Tetrachloride	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Trichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1-Trichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
p-Dichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Bromobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorobromomethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0096
Bromoform	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0019
Bromomethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Chlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Chlorodibromomethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0102
Chloroethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Chloroform	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0050
Chloromethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
o-Chlorotoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Chlorotoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dibromomethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m-Dichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
o-Dichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1-Dichloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
cis-1,2-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
trans-1,2-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Dichloromethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dichloropropane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005

TABLE 6.4 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Tap
1,3-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
2,2-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Ethylbenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Styrene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1,2,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Tetrachloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Toluene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,2-Trichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,3-Trichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m&p-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
o-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dibromo-3-Chloropropar	ne < 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Ethylenedibromide (EDB)	< 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Bromochloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
sec-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
tert-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorodifluoromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Fluorotrichloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Hexachlorobutadiene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Isopropylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Isopropyltoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Naphthalene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Propylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,2,3-Trichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.4 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Tap
1,2,4-Trichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,4-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< .0.001
1,3,5-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.5

Volatile Organic Compounds in Drinking Water Collected May 26, 1994

(Concentrations in mg/L)

Vinyl Chloride       < 0.0005       < 0.0005       < 0.         Carbon Tetrachloride       < 0.0005       < 0.0005       < 0.         1,2-Dichloroethane       < 0.0005       < 0.0005       < 0.         Trichloroethylene       < 0.0005       < 0.0005       < 0.	.0005 < .0005 < .0005 <	0.0005 < 0.0005 <	0.0005 0.0005 0.0005
Carbon Tetrachloride       < 0.0005       < 0.0005       < 0.         1,2-Dichloroethane       < 0.0005       < 0.0005       < 0.         Trichloroethylene       < 0.0005       < 0.0005       < 0.	.0005 <	0.0005	0.0005
1,2-Dichloroethane       < 0.0005	.0005 <		
Trichloroethylene < 0.0005 < 0.0005 < 0.		0.0005 <	
·	.0005 <		0.0005
1.1-Dichloroethylene < 0.0005 < 0.0005 < 0.0005		0.0005 <	0.0005
1,1 Dichloroctayione \ \ 0.0003 \ \ \ 0.0005	.0005 <	0.0005 <	0.0005
1,1,1-Trichloroethylene < 0.0005 < 0.0005 < 0.	.0005 <	0.0005	0.0005
p-Dichlorobenzene < 0.0005 < 0.0005 < 0.	.0005 <	0.0005 <	0.0005
Bromobenzene < 0.001 < 0.001 < 0.	.001 <	0.001 <	0.001
Dichlorobromomethane $< 0.0005$ $< 0.0005$ $< 0.$	.0005 <	0.0005	0.0111
Bromoform $< 0.0005 < 0.0005 < 0.$	.0005 <	0.0005	0.0033
Bromomethane $< 0.002 < 0.002 < 0.002$	.002 <	0.002 <	0.002
Chlorobenzene < 0.0005 < 0.0005 < 0.	.0005 <	0.0005 <	0.0005
Chlorodibromomethane $< 0.0005$ $< 0.0005$ $< 0.$	.0005 <	0.0005	0.0138
Chloroethane < 0.002 < 0.002 < 0.	.002 <	0.002 <	0.002
Chloroform < 0.0005 < 0.0005 < 0.	.0005 <	0.0005	0.0056
Chloromethane $< 0.002 < 0.002 < 0.002$	.002 <	0.002 <	0.002
o-Chlorotoluene < 0.001 < 0.001 < 0.	.001 <	0.001 <	0.001
p-Chlorotoluene < 0.001 < 0.001 < 0.	.001 <	0.001 <	0.001
Dibromomethane < 0.001 < 0.001 < 0.	.001 <	0.001 <	< 0.001
m-Dichlorobenzene < 0.001 < 0.001 < 0.	.001 <	0.001 <	< 0.001
o-Dichlorobenzene < 0.0005 < 0.0005 < 0.	.0005 <	0.0005 <	0.0005
1,1-Dichloroethane < 0.001 < 0.001 < 0.	.001 <	0.001 <	0.001
cis-1,2-Dichloroethylene $< 0.0005$ $< 0.0005$ $< 0.$	.0005 <	0.0005 <	0.0005
trans-1,2-Dichloroethylene $< 0.0005$ $< 0.0005$ $< 0.$	.0005 <	0.0005 <	0.0005
Dichloromethane $< 0.0005$ $< 0.0005$ $< 0.$	.0005 <	0.0005 <	0.0005
1,2-Dichloropropane < 0.0005 < 0.0005 < 0.	.0005 <	0.0005 <	0.0005

TABLE 6.5 (Contd.)

			•		
Parameter	Well #1	Well #2	Well #3	Well #4	Tap
1,3-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
2,2-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Ethylbenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Styrene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1,2,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Tetrachloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Toluene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,2-Trichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,3-Trichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m&p-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
o-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dibromo-3-Chloropropane < 0.010	e < 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Ethylenedibromide (EDB)	< 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Bromochloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
sec-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
tert-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorodifluoromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Fluorotrichloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Hexachlorobutadiene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Isopropylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Isopropyltoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Naphthalene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Propylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,2,3-Trichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.5 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
1,2,4-Trichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,4-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3,5-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.6

Volatile Organic Compounds in Drinking Water Collected August 18, 1994

(Concentrations in mg/L)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
Benzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Vinyl Chloride	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Carbon Tetrachloride	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Trichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1-Trichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
p-Dichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Bromobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorobromomethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0126
Bromoform	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0030
Bromomethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Chlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Chlorodibromomethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0137
Chloroethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Chloroform	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0073
Chloromethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
o-Chlorotoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Chlorotoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dibromomethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m-Dichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
o-Dichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1-Dichloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
cis-1,2-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
trans-1,2-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Dichloromethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0006
1,2-Dichloropropane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005

TABLE 6.6 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
1,3-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
2,2-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Ethylbenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Styrene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1,2,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Tetrachloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Toluene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,2-Trichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,3-Trichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m&p-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
o-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dibromo-3-Chloropropar	ne < 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Ethylenedibromide (EDB)	< 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Bromochloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
sec-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
tert-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorodifluoromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Fluorotrichloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Hexachlorobutadiene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Isopropylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Isopropyltoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Naphthalene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Propylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,2,3-Trichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.6 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
1,2,4-Trichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,4-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3,5-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.7

Volatile Organic Compounds in Drinking Water Collected November 10, 1994

(Concentrations in mg/L)

Parameter	Well #1	Well #2	Well #3	Well #4	Tap
Benzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Vinyl Chloride	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Carbon Tetrachloride	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Trichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1-Trichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
p-Dichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Bromobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorobromomethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0152
Bromoform	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0006
Bromomethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Chlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Chlorodibromomethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0071
Chloroethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Chloroform	< 0.0005	< 0.0005	< 0.0005	< 0.0005	0.0155
Chloromethane	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
o-Chlorotoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Chlorotoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dibromomethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m-Dichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
o-Dichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1-Dichloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
cis-1,2-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
trans-1,2-Dichloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Dichloromethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2-Dichloropropane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005

TABLE 6.7 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
1,3-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
2,2-Dichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3-Dichloropropene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Ethylbenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Styrene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,1,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,1,2,2-Tetrachloroethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Tetrachloroethylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
Toluene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,1,2-Trichloroethane	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,3-Trichloropropane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
m&p-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.000
o-Xylene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.000
1,2-Dibromo-3-Chloropropa	ne < 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Ethylenedibromide (EDB)	< 0.010	< 0.010	< 0.010	< 0.010	< 0.010
Bromochloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
sec-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
tert-Butylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Dichlorodifluoromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Fluorotrichloromethane	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Hexachlorobutadiene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Isopropylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-Isopropyltoluene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Naphthalene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
n-Propylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,2,3-Trichlorobenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

TABLE 6.7 (Contd.)

Parameter	Well #1	Well #2	Well #3	Well #4	Тар
1,2,4-Trichlorobenzene	< 0.0005	< 0.0005	< 0.0005	< 0.0005	< 0.0005
1,2,4-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
1,3,5-Trimethylbenzene	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001

contain significant amounts of hazardous materials and therefore represent a potential threat to the environment. Groundwater below these sites is monitored routinely to assess the amount and nature of hazardous chemical releases from these units. The sites which are routinely monitored are the sanitary landfill in the 800 Area and the 317/319 Area, which consists of eight separate waste management units located within a small geographical area. The site of an inactive experimental reactor, CP-5, is also monitored periodically to determine if any releases of radionuclides occurred from this unit.

### 6.2.1, 317/319 Area

Management of waste has been conducted in eight separate units within the 317 and 319 Areas. The 317 Area is currently used as a temporary storage area for radioactive waste before it is shipped off-site for disposal. The area also contained two RCRA-permitted units which were formally closed during 1994. The 319 Area is an inactive landfill adjacent to the 317 Area. In addition to these units, a second landfill site, the ENE landfill, is located to the east-northeast of the 319 Area. This unit was used in the late 1940s and early 1950s primarily for the disposal of construction debris from several sites, including the University of Chicago's Manhattan Project. A sketch of the 317/319 Area is shown in Figure 6.1.

The most significant units in this area in terms of groundwater impact are an inactive French drain (dry well) in the 317 area and the landfill and French drain in the 319 Area. The 317 Area French drain operated until the mid 1950s and was used for disposal of unknown amounts of liquid chemical wastes. The landfill at 319 was operated from the mid-1950s until 1968 when the sanitary landfill in the 800 Area was put into use. The French drain, similar to the one in the 317 Area, was operated until 1968. Quantities of a wide variety of liquid wastes, including heavy metals, solvents and waste oil, some containing PCBs, were poured into this drain.

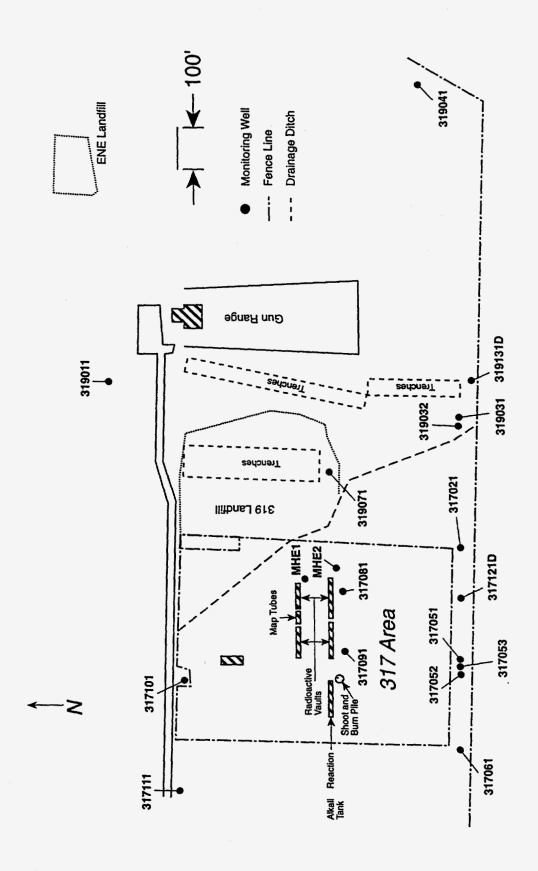


Figure 6.1 Locations of Components Within the 317/319/ENE Area

The 317 Area contains six vaults used for temporary storage of solid radioactive waste. Water from footing drains and/or sumps is collected and discharged into a sewer system. This sewer system, which was designed to drain off-site, was permanently closed in 1986 after it was discovered that the water contained very small amounts of several radionuclides. Water collecting in the sewer system was periodically pumped out from manholes into portable tanks, transported to the Waste Management Building and analyzed for radioactivity before release to the laboratory sewage collection system. During August 1993, the discharge of water from these manholes to the laboratory sewer system began. Monthly samples from two manholes associated with this system are analyzed for volatile organic compounds. The results are presented in Section 6.2.2.3.

The 319 Area currently consists of a mound created by waste fill activities. The waste consisted of noncombustible refuse, demolition and construction debris. In addition, suspect waste (material which was not known to be contaminated but which had the potential for hidden radioactive contamination which could not be confirmed by direct measurement, such as the inside of long pipes or ductwork) was also placed in this unit. The landfill consisted of a number of trenches, 3 to 5 m (10 to 15 ft) deep, which were filled with waste material. When the trenches were filled with waste, they were covered with soil. A recent geophysical survey has identified at least three of these trenches. Recently, five test pits were dug into the waste; the waste was inspected, photographed, and surveyed for the presence of radioactive or volatile organic compound contamination; then placed back into the evacuation. Field measurements of radioactivity levels and organic vapor concentrations indicated that this material is not significantly contaminated.

The French drain in the 319 Area was constructed in the late 1950s in an area of the fill material by placing a corrugated steel pipe vertically into a gravel-filled excavation and backfilling around the pipe. Waste liquids were poured into the pit and flowed into the pipe.

The ENE landfill is believed to consist primarily of construction debris, and other noncombustible rubbish, such as metal turnings and empty steel drums. The waste was placed in a natural ravine and covered with soil.

### 6.2.2. Groundwater Monitoring at the 317/319 Area

Twelve active monitoring wells (some of which are clustered or nested) are installed at the locations shown in Figure 6.2. Well data are listed in Table 6.8. The wide range in water level elevations shown in Table 6.8 is not unusual and is due to the fact that some of the wells are screened at different depths in different saturated zones. This variation in water level also may be indicative of "perched" (i.e., discontinuous) groundwater conditions within the glacial till. Samples are collected quarterly following EPA sampling protocols listed in the RCRA Ground-Water Monitoring Technical Enforcement Guidance Document (September 1986).<sup>24</sup>

Groundwater monitoring in the 317/319 Area has been conducted since 1986. Wells 319011, 317021, 319031, and 319041 were installed in September 1986; 317051 and 317061 in August 1987; 319071, 317081, and 317091 in July 1988; 317101 and 317111 in September 1988; and wells 319032, 317053, and 317052 were installed in June 1989. These wells were all completed in the glacial till. In addition, wells 317121D and 319131D were installed in November 1989 and reach the dolomite aquifer at about 25 m (80 ft) below the surface.

Wells 317101 and 317111 are upgradient of the 317 storage area and well 319011 is upgradient of the 319 landfill area. A sand lens present at 5 to 8 m (15 to 25 ft) was recently discovered and wells 317053, 317052 and 319032 were placed at this depth. This layer is also intercepted by wells 317081, 317091, and 317101.

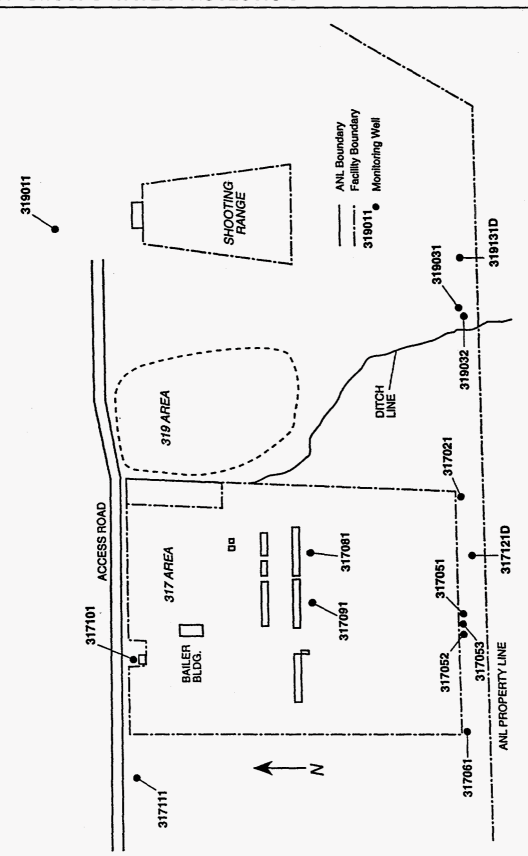


Figure 6.2 Active Monitoring Wells in the 317 and 319 Areas

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Groundwater Monitoring Wells - 317/319 Area

TABLE 6.8

ID Number	Well Depth <sup>1</sup>	Ground Elevation <sup>2</sup>	Monitoring Zone <sup>3</sup>	Well Type <sup>4</sup>	Date Drilled	Water Elevation
319011	40	688.3	35-40/653-648	2/PVC	9/86	655.8
317021	40	686.2	35-40/651-646	2/PVC	9/86	664.2
319031	41	670.2	36-41/634-629	2/PVC	9/86	633.0
319032	25	670.2	20-25/650-645	2/PVC	8/89	650.5
317051	20	683.4	15-20/668-663	2/PVC	7/87	not measured
317053	22	683.4	17-22/666-661	2/PVC	8/89	dry
317052	14	683.4	9-14/674-669	2/PVC	8/89	676.0
317061	40	680.9	35-40/646-641	2/PVC	7/87	659.4
317081	30	682.8	20-30/662-652	2/PVC	7/88	675.4
317091	30	684.0	20-30/663-653	2/PVC	7/88	674.4
317101	39	692.3	29-39/663/653	2/PVC	8/89	670.2
317111	39	698.8	29-39/670-660	2/PVC	8/89	682.8
317121D	79	681.0	69-79/612-602	6/CS	9/88	612.4
319131D	69	667.8	59-69/609-599	6/CS	9/88	607.1

<sup>&</sup>lt;sup>1</sup>feet below ground

Note: Wells identified by a "D" are deeper wells monitoring the dolomite bedrock aquifer.

<sup>&</sup>lt;sup>2</sup>feet mean sea level

<sup>&</sup>lt;sup>3</sup>depth/elevation

<sup>4</sup>inner diameter (inches)/well material (PVC=polyvinyl chloride, CS=carbon steel))

In addition to wells in this area, two manholes associated with the vault sewer system were monitored on a monthly basis. The location of the manholes is shown in Figure 6.1.

## 6.2.2.1. Sample Collection

The monitoring wells are sampled using the protocol listed in the RCRA Ground-Water Monitoring Technical Enforcement Guidance Document.<sup>23</sup> The volume of the water in the casing is determined by measuring the water depth from the surface and the depth to the bottom of the well. This latter measurement also determines whether siltation has occurred that might restrict water movement in the screen area. For those wells in the glacial till that do not recharge rapidly, the well is emptied and the volume removed compared to the calculated volume. In most cases these volumes are nearly identical. The well is then sampled by bailing with a dedicated Teflon bailer. The field parameters for these samples (pH, specific conductance, redox potential, and temperature) are measured statically. For those samples in the porous, saturated zone which recharge rapidly, three well volumes are purged using dedicated submersible pumps while the field parameters are measured continuously. These parameters stabilize quickly in these wells. In the case of the dolomite wells, samples are collected as soon as these readings stabilize. Samples for volatile organics, semivolatile organics, PCB/pesticides, metals, and radioactivity are collected in that order. The samples are placed in precleaned bottles, labelled, and preserved.

During each sampling event, one well is selected for replicate sampling. An effort is made to vary this selection so that replicates are obtained at every well over the course of time. In addition, a field blank is also obtained.

# 6.2.2.2. Sample Analysis - 317/319 Area

The 317/319 groundwater chemical analyses were performed using standard operating procedures (SOPs) written, reviewed, and issued as controlled documents by members of the ESH-DACH. These SOPs reference protocols found in SW-846, 3rd edition, "Test Methods for Evaluating Solid Waste." Sixteen metals were routinely determined. Ten were done using flame atomic absorption spectroscopy and inductively coupled plasma atomic emission spectroscopy and five by graphite furnace atomic absorption spectroscopy. Mercury was determined by cold vapor atomic absorption spectroscopy. Chloride was determined by titrimetry. Volatile organic compounds were determined by using a purge and trap sample pretreatment followed by gas chromatography-mass spectroscopy detection. Semivolatile organic compounds were determined by solvent extraction followed by gas chromatography-mass spectroscopy detection. In the case of organic compound analyses, efforts were made to identify compounds which were present, but not included on the method list. This was accomplished and standard solutions of these compounds were prepared and analyzed.

PCB/pesticides were determined by solvent extraction followed by gas chromatography-electron capture detection at an off-site-contracted laboratory. SW-846 Procedure 8080 was specified and used.

The 317/319 groundwater radiological analyses were performed using SOPs written, reviewed, and issued as controlled documents by members of the ESH-DARC. Cesium-137 was determined by gamma-ray spectrometry. Hydrogen-3 was determined by distillation followed by a beta liquid scintillation counting technique. Strontium-90 was determined by an ion-exchange separation followed by a proportional counting technique.

### 6.2.2.3 Results of Analyses

The description of each well, a listing of field parameters measured during sample collection, and the results of chemical and radiological analyses of samples from the wells in the 317/319 Area are contained in Tables 6.9 through 6.20. All radiological and inorganic analyses results are shown in these tables. The analysis methods used for organic compounds will identify and quantify all the compounds contained in the CLP Target Compound List. However, the vast majority of these compounds were not detected in the samples. To simplify the format of these tables, those results less than the detection limit are not included. Only those constituents which were present in amounts great enough to quantify are shown. The detection limits for the organic compounds listed were typically 1 to 5  $\mu$ g/L.

### Field Results

The purging of wells to produce water representative of the groundwater being studied is followed by measuring the field parameters. For the wells reported in this study, temperature, pH, and specific conductance remain fairly constant after two well volumes are removed. The redox potential stabilizes after two well volumes are removed. On the basis of this information, sampling is conducted after the removal of three well volumes. The field parameters listed in the tables are the final readings obtained at the time of sampling. Wells 319011, 317021, 317061, 317111, and 319031 usually dry up after one well volume is removed. Therefore, field parameters are measured on one well volume. Well 319031 was dry during the third and fourth quarters.

TABLE 6.9 Groundwater Monitoring Results, 300 Area Well #319011, 1994

m(MSL) Well Point Elevation 196.95 Ground Surface Elevation 209.81 **PVC** 

Casing Material:

Constituent	Units	03/14/94	06/20/94	09/13/94	11/28/94
Water Elevation	m	198.94	198.82	198.21	198.00
Temperature	°C	10.0	11.1	12.1	10.2
pН	pН	7.09	6.99	7.08	6.66
Redox	mV	276	13	323	-162
Conductivity	µmhos/cm	829	809	839	808
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.08	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.015	0.031	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	1.1	1.6
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	26	23	24	26

TABLE 6.10

Groundwater Monitoring Results, 300 Area Well #317021, 1994

Well Point Elevation 196.90
Ground Surface Elevation 209.17
Casing Material: PVC

Constituent	Units	03/14/94	06/20/94	09/13/94	11/28/94
Water Elevation	m	200.97	200.87	199.80	198.94
Temperature	$^{\circ}\mathrm{C}$	10.3	10.9	10.9	10.2
pН	pН	7.24	7.19	7.26	6.80
Redox	mV	265	28	332	-110
Conductivity	μmhos/em	746	554	587	551
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	< 0.100	0.102	< 0.100	0.104
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	11	17	19	11
1,1,1-Trichloroethane	μg/L	128	215	191	84
1,1-Dichloroethane	μg/L	63	96	163	79
1,1-Dichloroethene	μg/L	2	2	4	1
1,2-Dichloroethane	μg/L	10	26	17	6
Carbon tetrachloride	μg/L	6	10	10	4
Chloroform	μg/L	4	5	6	3
Methylene chloride	μg/L	-	-	1	-
Acetone	μg/L	-	-	-	4
Tetrachloroethene	μg/L	1	2	2	1
Trichloroethene	μg/L	3	5	5	3
cis-1,2-Dichloroethene		=	0.6	-	-

TABLE 6.11

Groundwater Monitoring Results, 300 Area Well #319031, 1994

Well Point Elevation 19
Ground Surface Elevation 20
Casing Material: P

m(MSL) 192.08 204.28 PVC

Constituent	Units	03/14/94	03/14/94	06/27/94
Water Elevation	m	193.17	193.17	192.80
Temperature	°C	10.2	10.2	10.9
рН	рH	7.05	7.05	7.03
Redox	mV	260	260	304
Conductivity	μmhos/cm	975	975	702
Arsenic	mg/L	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	0.053	0.052
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020
Copper	mg/L	< 0.010	< 0.010	< 0.010
Iron	mg/L	< 0.04	< 0.04	< 0.04
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.010	< 0.010
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010
Cesium-137	pCi/L	< 1.0	< 1.0	<b>-</b>
Hydrogen-3	nCi/L	1.221	1.148	0.994
Strontium-90	pCi/L	0.26	0.32	0.27
Chloride	mg/L	30	30	27
1,1,1-Trichloroethane	$\mu \mathrm{g}/\mathrm{L}$	1	0.6	1
1,1-Dichloroethane	$\mu \mathrm{g}/\mathrm{L}$	5	4	5
Tetrachlorethene	$\mu \mathrm{g/L}$	-	-	0.7
Trichloroethene	$\mu$ g/L	5	5	7

TABLE 6.12

Groundwater Monitoring Results, 300 Area Well #319032, 1994

	m(MSL)
Well Point Elevation	195.82
Ground Surface Elevation	204.28
Casing Material:	PVC

Constituent	Units	03/14/94	06/27/94	09/13/94	11/28/94
Water Elevation	m	197.77	197.43	197.15	196.83
Temperature	°C	10.1	10.9	10.9	10.7
pH	pН	6.98	7.02	7.07	6.58
Redox	mV	296	233	330	-141
Conductivity	μmhos/cm	772	756	806	840
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.052	0.058	0.057	0.051
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	0.028
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.017	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	1.4	< 1.0
Hydrogen-3	nCi/L	0.660	0.523	0.664	0.680
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	27	24	25	172
Methylene chloride	$\mu \mathrm{g}/\mathrm{L}$	-	-	1	-

TABLE 6.13

Groundwater Monitoring Results, 300 Area Well #317052, 1994

Well Point Elevation 203.70
Ground Surface Elevation 208.32
Casing Material: PVC

Constituent	Units	03/14/94	06/27/94	09/13/94	11/29/94
Water Elevation	m	205.56	204.54	204.46	205.20
Temperature	°C	6.9	10.4	13.6	12.1
pH	pН	7.30	7.20	7.33	7.05
Redox	mV	176	304	323	267
Conductivity	µmhos/cm	450	501	559	532
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.010
Cesium-137	pCi/L	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	2	2	2	1
Methylene Chloride	μg/L		<b>-</b>	<del>-</del>	1

TABLE 6.14

Groundwater Monitoring Results, 300 Area Well #317061, 1994

Well Point Elevation 194.93
Ground Surface Elevation 207.54
Casing Material: PVC

Constituent	Units	03/14/94	06/20/94	06/20/94	09/14/94	11/28/94
Water Elevation	m	199.52	199.09	199.09	198.59	198.50
Temperature	°C	10.5	10.9	10.9	10.6	9.9
pH	pН	7.07	7.04	7.04	7.12	7.06
Redox	mV	129	25	25	348	-96
Conductivity	μmhos/cm	766	769	769	794	776
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	0.051	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.024	0.016	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.026
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	< 0.100	0.116	< 0.100	< 0.100	0.114
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	57	56	54	67	57
Acetone	μg/L	-	-	-	_	5
Methylene chloride	μg/L	-	-	-	-	1

TABLE 6.15

Groundwater Monitoring Results, 300 Area Well #317081, 1994

Well Point Elevation 192.08
Ground Surface Elevation 208.14
Casing Material: PVC

Constituent	Units	03/14/94	06/27/94	09/16/94	09/16/94
Water Elevation	m	204.16	204.09	204.36	204.36
Temperature	°C	8.6	10.4	13.5	13.5
рН	pН	7.04	6.94	7.16	7.16
Redox	mV	194	289	86	86
Conductivity	μmhos/cm	544	574	646	646
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pČi/L	4.2	11.5	9.0	6.6
Hydrogen-3	nCi/L	0.253	0.172	0.547	0.673
Strontium-90	pCi/L	3.42	4.79	7.22	7.68
Chloride	mg/L	9	15	44	47
Carbon tetrachloride	μg/L	2	18	8	8
Chloroform	μg/L	1	9	3	3
Methylene Chloride	μg/L	-	-	0.6	1
Tetrachloroethene	μg/L	0.7	4	5	6
Trichloroethene	μg/L	91	102	77	83
cis-1,2-Dichloroethene	μg/L	3	5	9	11
trans-1,2-Dichloroethene	μg/L	0.5	-	-	0.5

TABLE 6.16

Groundwater Monitoring Results, 300 Area Well #317091, 1994

Well Point Elevation 199.16
Ground Surface Elevation 208.14
Casing Material: PVC

Constituent	Units	03/14/94	06/27/94	09/16/94	11/29/94	11/29/94
Water Elevation	m	202.71	201.68	201.42	201.75	201.75
Temperature	°C	7.7	10.5	13.5	13.7	13.7
pH	pН	7.20	7.11	7.24	6.89	6.89
Redox	mV	174	272	-94	303	303
Conductivity	μmhos/cm	490	521	609	665	665
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	0.0020	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.001	< 0.036	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050	< 0.050
Cesium-137	pČi/L	< 1.0	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	6	7	18	42	44
Chloroform	$\mu \mathrm{g}/\mathrm{L}$	-	-	0.7	-	-
Methylene chloride	μg/L	-	-	1	-	49
Trichloroethene	μg/L	-	-	-	0.4	-
1,1 Dichloroethane	μg/L	-	2	15	-	<b>-</b>
1,1,1 Trichloroethane	μg/L	-	-	1	-	· <u>-</u>
1,2 Dichloroethane	μg/L	-	-	0.5	-	-
Acetone	μg/L	-	-	-	-	6
Bromochloromethane	μg/L	-	-	-	-	3

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TABLE 6.17

Groundwater Monitoring Results, 300 Area Well #317101, 1994

Well Point Elevation 198.66
Ground SurfaceElevation 211.04
Casing Material: PVC

Constituent	Units	03/14/94	06/27/94	09/14/94	11/19/94
Water Elevation	m	203.01	202.73	202.19	201.90
Temperature	$^{\circ}\mathrm{C}$	11.6	11.4	11.4	11.4
pН	pН	7.05	7.01	7.16	6.74
Redox	mV	1 <b>70</b>	272	352	284
Conductivity	μmhos/cm	1394	1272	1201	1479
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	0.056	0.064
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.061	0.069	0.076	0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	0.022	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	328	275	247	369
Methylene chloride	$\mu g/L$	-	-	2	_

TABLE 6.18

Groundwater Monitoring Results, 300 Area Well #317111, 1994

Welll Point Elevation 200.72
Ground Surface Elevation 213.02
Casing Material: PVC

Constituent	Units	03/14/94	06/20/94	09/14/94	11/28/94
Water Elevation	m	207.52	206.66	202.87	202.26
Temperature	.C	10.4	10.6	11.5	11.7
pН	pН	7.28	7.06	7.09	6.51
Redox	mV	145	-6	346	-198
Conductivity	μmhos/cm	752	758	844	1097
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.061	0.057	0.067	< 0.085
Beryllium	mg/L	< 0.0001	0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	0.0002	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	0.024
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	0.17	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.028	0.024	0.092	0.069
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	94	94	105	25

TABLE 6.19

Groundwater Monitoring Results, 300 Area Well #317121D, 1994

Well Point Elevation 183.17
Ground Surface Elevation 207.57
Casing Material: Steel

Constituent	Units	03/14/94	06/21/94	09/14/94	11/28/94
Water Elevation	m	186.47	186.38	186.37	186.34
Temperature	$^{\circ}\mathrm{C}$	11.0	12.1	11.5	10.7
pH	рH	11.33	9.15	11.40	10.65
Redox	mV	92	88	264	-178
Conductivity	μmhos/cm	587	583	<b>5</b> 11	<i>5</i> 66
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.011	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	0.021
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pCi/L	< 1.0	< 1.0	< 1.0	< 1.0
Hydrogen-3	nCi/L	0.112	0.107	0.135	0.171
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	47	46	50	47
Methylene chloride	μg/L	-	-	-	0.6

TABLE 6.20

Groundwater Monitoring Results, 300 Area Well #319131D, 1994

Well Point Elevation 182.06
Ground Surface Elevation 203.56
Casing Material: Steel

Constituent	Units	03/14/94	06/21/94	09/14/94	11/28/94
Water Elevation	m	184.69	184.42	184.24	184.23
Temperature	°C	11.1	16.0	11.5	10.7
pH	pН	7.06	7.47	7.34	6.57
Redox	mV	302	87	309	-157
Conductivity	μmhos/cm	<i>7</i> 71	773	788	810
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.076	0.071	0.072	0.066
Beryllium	mg/L	0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	< 0.010	< 0.036	< 0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	0.020	< 0.020	0.026	0.028
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.2000	< 0.2000	< 0.2000	< 0.2000
Zinc	mg/L	< 0.0100	< 0.0100	< 0.0500	< 0.0500
Cesium-137	pCi/L	< 1.0	< 1.0	1.0	< 1.0
Hydrogen-3	nCi/L	0.730	1.148	1.349	1.325
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	46	44	51	47

#### Inorganic Results

The Illinois Groundwater Quality Standards for Class I: Potable Resource Groundwater, 35 IAC Section 620.410, were used as the standard for evaluation of the inorganic results based on evaluation of the regulations. The standards are presented in Tables 6.21 and 6.22. In 1994, all samples for metals analyses were field filtered prior to preservation with acid (IEPA requirement for the IEPA-approved groundwater monitoring program at the 800 Area Landfill, Section 6.3.2.3). In previous years, ESH-DACH performed the majority of the metals analyses using flame atomic absorption. Inductively Coupled Argon Plasma (ICP) Spectrometry was used during late 1994. The ICP method is not as sensitive as the flame method and, therefore, the reporting levels for metals are inconsistently higher during the second half of 1994. No elevated levels, with respect to the WQS for the inorganics were noted with the exception of pH at dolomite well 317121D and chloride at well 317101. The pH changes drastically between the purging of 2 to 5 volumes of water. In each case, the last value obtained was recorded. Several wells had elevated levels of barium and manganese, but well below the WQS. Barium concentrations in these wells ranged from 0.05 mg/L to 0.085 mg/L and manganese levels ranged from 0.02 mg/L to 0.09 mg/L. The source of the elevated barium and manganese levels is unknown. Elevated levels of manganese have been reported in previous annual reports. 12

### Organic Results

Each well was sampled quarterly and analyzed for volatile organic compounds. The results for 1994 are similar to those reported for 1993. Volatile organic compounds were detected in wells 317021, 319031, 319032, 317052, 317061, 317081, 317091, 317101, and 317121D. The levels of volatile organics are persistent and appear to be indicative of different sources of contamination. Once during the year the wells were sampled and analyzed for semivolatile organic compounds, polychlorinated biphenyls (PCBs) and

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TABLE 6.21

Illinois Class I Groundwater Quality Standards - Inorganics (Concentrations in mg/L, except radionuclides and pH)

Constituent	Standard
Antimony	0.006
Arsenic	0.05
Barium	2
Beryllium	0.004
Boron	2
Cadmium	0.005
Chloride	200
Chromium	0.1
Cobalt	1
Copper	0.65
Cyanide	0.2
Fluoride	4
Iron	5
Lead	0.0075
Manganese	0.15
Mercury	0.002
Nickel	0.1
Nitrate, as N	10
Radium-226	20 pCi/
Radium-228	20 pCi/
Selenium	0.05
Silver	0.05
Sulfate	400
Thallium	0.002
Total Dissolved Solids	1,200
Zinc	5
pH	6.5-9.0 un

TABLE 6.22

Illinois Class I Groundwater Quality Standards - Organics
(Concentrations in mg/L)

Constituent	Standard
Alachlor	0.002
Aldicarb	0.003
Atrazine	0.003
Benzene	0.005
Benzo(a)pyrene	0.0002
Carbofuran	0.04
Carbon Tetrachloride	0.005
Chlordane	0.002
Dalapon	0.2
Dichloromethane	0.005
Di(2-ethyhexyl)phthalate	0.006
Dinoseb	0.007
Endothall	0.1
Endrin	0.002
Ethylene Dibromide	0.00005
Heptachlor	. 0.0004
Heptachlor Epoxide	0.0002
Hexachlorocyclopentadiene	0.05
Lindane	0.0002
2,4-D	0.07
o-Dichlorobenzene	0.6
p-Dichlorobenzene	0.075

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TABLE 6.22 (Contd.)

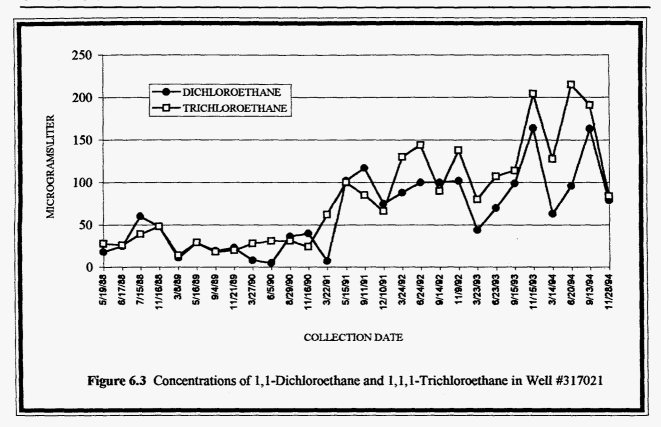
Constituent	Standard
1,2-Dibromo-3-Chloropropane	0.0002
1,2-Dichloroethane	0.005
1,1-Dichloroethane	0.007
cis-1,2-Dichloroethylene	0.07
trans-1,2-Dichloroethylene	0.1
1,2-Dichloropropane	0.005
Ethylbenzene	0.7
Methoxychlor	0.04
Monochlorobenzene	0.1
Pentachlorophenol	0.001
Phenols	0.1
Picloram	0.5
PCBs(decachlorobiphenyl)	0.0005
Simazine	0.004
Styrene	0.1
2,4-5-TP (Silvex)	0.05
Tetrachloroethylene	0.005
Toluene	1
Toxaphene	0.003
1,1,1-Trichloroethane	0.2
1,1,2-Trichloroethane	0.005
1,2,4-Trichlorobenzene	0.07
Trichloroethylene	0.005
Vinyl Chloride	0.002
Xylenes	10

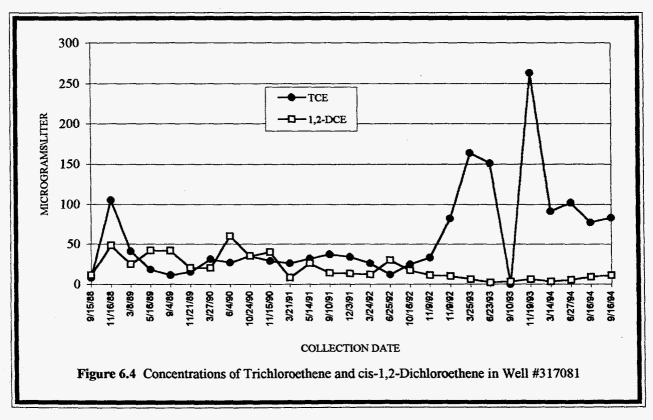
pesticides and herbicides. In 1994, no semivolatiles, PCBs, pesticides, or herbicides were found.

The organic WQSs were exceeded for at least one parameter in wells 317021, 317081, 317091, and 319031. The results for well 317021 are shown in Figure 6.3. The major components are 1,1,1-trichloroethane (TCA, exceeding the WQS) and 1,1-dichloroethane, which can be a decomposition product of TCA. As can be seen, the concentrations roughly parallel each other and the levels found are remarkably constant until 1991 at which time a substantial increase is seen. The previous consistency would indicate that this well is sampling a large area of contaminated water which is unaffected by seasonal water level changes. The large increase in the summer and fall of 1991 clearly is related to a period of intense drought and could be related to restricted flow of normal dilution water. Levels exceeding the WQS for carbon tetrachloride, 1,2,-dichloroethane, and tetrachloroethene were also found. Trace levels of acetone, chloroform, trichloroethene (TCE), methylene chloride, and cis-1,2-dichloroethene (1,2-DCE) were also found in this well. The well is immediately below the plugged sewer line previously discussed and this sewer line is known to be contaminated with these four compounds. 1,1-dichloroethene was found in trace amounts, which can be a decomposition product of TCE.

Wells 317081 and 317091 are adjacent to the storage vaults and are close to one another. The chemical characteristics are quite dissimilar. The principal volatile organic compounds found in well 317081 in 1994 are TCE, 1,2-DCE, and carbon tetrachloride. The results obtained from the beginning of sampling until the end of 1994 are shown in Figure 6.4 for TCE and 1,2-DCE. When TCE breaks down in the presence of soil bacteria, the cis isomer of 1,2-DCE is produced almost exclusively. The fact that they are both present in these samples at relatively stable concentrations indicates that there may be ongoing release of TCE into the groundwater, such as from highly contaminated soil. The half life for the conversion indicated is about 30 days. The end product of this conversion

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is vinyl chloride which has a half life of 26,000 days. Vinyl chloride has never been detected in these samples. Carbon tetrachloride appeared each quarter in 1994 and at higher levels than 1993 (two quarters). Chloroform, tetrachloroethene, trans-1,2-dichloroethene, and methylene chloride are occasionally found in trace amounts in this well. In contrast, the levels and variety of volatile organics found in well 317091 are quite variable. In the initial samples obtained in 1988, very high amounts of 1,1,1,-trichloroethane and 1,1-dichloroethane (170 and 160  $\mu$ g/L, respectively) were found. In subsequent samples, values for 1,1-dichloroethane have ranged from 0.5  $\mu$ g/L to 186  $\mu$ g/L and values for 1,1,1-trichloroethane have ranged from 1  $\mu$ g/L to 31  $\mu$ g/L. Trichloroethene, tetrachloroethene and 1,2-dichloroethane have also been detected on occasion. During 1992 and 1993, trace levels of chloroform, methylene chloride, cis-1,2-dichloroethene, and trichloroethane were found. Methylene chloride was found often in most samples and blanks. During 1994, trace levels of acetone, chloroform, bromochloromethane, 1,2-dichloroethane, and 1,1,1-trichloroethane were found during one quarter and trace levels of methylene chloride were found during two quarters.

Dolomite well 317121D had only trace levels of toluene during one quarter. Low levels of trichloroethene, tetrachloroethene, 1,1-dichloroethane, and 1,1,1-trichloroethane were detected in well 319031. This is consistent with results noted in previous reports. This well is frequently dry but it contains organic constituents when water is present.

Polychlorinated biphenyl compounds were reported in several of the wells in 1990. These wells were resampled in 1991, 1992, 1993, and 1994 and no PCBs were indicated. This confirms previous sampling results. Semivolatile organics and pesticides/herbicides were not detected in any of the wells during 1994.

Manholes E1 and E2 described in Sections 6.2.1 and 6.2.2, in the 317 Area are sampled monthly and analyzed for volatile organic compounds. The results are presented

in Table 6.23. Existing foundation drains around storage vaults convey groundwater away from the structures and into manholes E1 and E2. Volatile organics are detected at fairly consistent levels in all samples as shown in Figure 6.5. As previously discussed for well 317081, the consistency would indicate that these manholes are collecting an area of contaminated water. The fact that levels are constant and the TCE and 1,2-DCE are present in most of the samples, indicates an ongoing release of these compounds into the groundwater, such as from highly contaminated soils. Trace levels of acetone, benzene, trans-1,2-dichloroethene,1,2-dichloroethane,1,1,1-trichloroethane,trichlorofluoromethane, 4-methyl-2-pentanone, ethyl ether, methylene chloride, and toluene have been found but not on a consistent basis. Carbon tetrachloride was detected in 75% of the samples in manhole E2 and only the February sample in manhole E1. The source of these compounds is believed to be the French drains previously described in Section 6.2.1 but additional characterization activities described in Section 6.5.2 will better define the nature, rate, and extent of contamination at this location.

#### Radioactive Constituents

Samples collected quarterly from the monitoring wells in the 317 and 319 Areas were analyzed for hydrogen-3, strontium-90, and for gamma-ray emitters. The results are presented in Tables 6.9 to 6.20 Evidence of possible migration of radionuclides off the site is noted by the low concentrations of hydrogen-3 in wells 317021, 317061, 317121D, 319031, 319032, and 319131D, which are located near the south perimeter fence. During two quarters, a small amount of cesium-137 was detected in well 319011. A small amount of strontium-90 was also detected in well 319031. These monitoring wells are directly below a small drainage swale from the 319 Area that has contained water intermittently with measurable concentrations of hydrogen-3 and strontium-90. Well 317081 contains measurable levels of hydrogen-3, strontium-90, cesium-137. This well is next to facilities that have

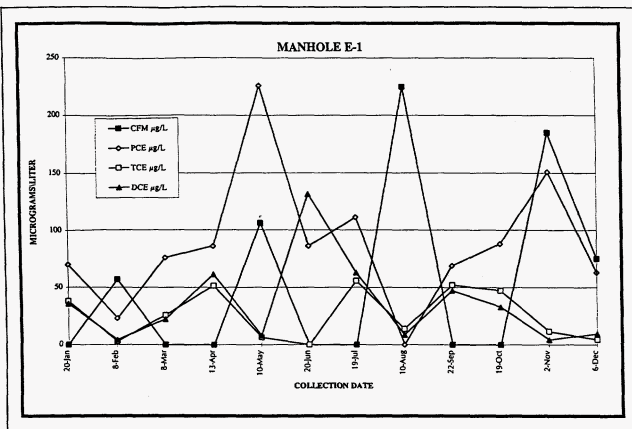
TABLE 6.23

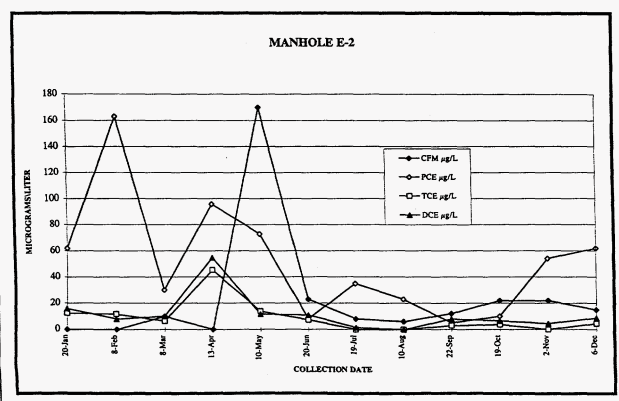
Volatile Organic Compounds in 317 Area Manholes E-1 and E-2, 1994

(Concentrations in  $\mu g/L$ )

Date	Chlor	roform	Tetra- chloro	ethene	Tri- chloro	ethene	cis-1,2 Dichlo	- roethene
	E-1	E-2	E-1	E-2	E-1	E-2	E-1	E-2
January 20	< 1	< 1	70	62	53	17	50	22
February 8	57	< 1	23	163	4	16	6	11
March 8	< 1	10	76	30	36	9	31	14
April 13	< 1	< 1	86	96	72	63	86	76
<b>M</b> ay 10	106	170	226	73	9	19	11	16
June 20	< 1	23	86	8	< 1	10	184	15
July 19	< 1	8	111	35	78	< 1	88	2
August 10	225	6	< 1	23	19	< 1	12	< 1
September 22	< 1	12	69	5	73	4	66	11
October 19	< 1	22	88	10	66	5	46	9
November 2	185	22	151	54	16	< 1	6	6
December 6	75	15	63	62	6	6	13	12

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stored radioactive materials in the past. All concentrations are well below any applicable standards.

### 6.3. Sanitary Landfill

The 800 Area is the site of ANL's sanitary landfill. The 21.8-acre landfill is located on the western edge of ANL property (Figure 1.1). The landfill has received waste since 1966 and operated under IEPA permit No. 1981-29-OP which was issued on September 18, 1981. The landfill received general refuse, construction debris, boiler house ash, and other nonradioactive solid waste until September 1992. The landfill is now being closed pursuant to permit number 1992-002-SP and supplemental permit number 1994-506-SP.

#### 6.3.1. French Drain

The landfill area was used for the disposal of certain types of liquid wastes from 1969 to 1978. The wastes were poured into a French drain which consisted of a corrugated steel pipe placed in a gravel-filled pit dug into an area previously filled with waste. The liquid waste was poured into the drain and allowed to permeate into the gravel and thence into the soil and fill material. There is documentation available that indicates that 29,000 gallons of liquid waste were placed in this drain. Many of the wastes disposed of in this manner are now defined as hazardous wastes. The presence of volatile and other toxic organic compounds has been confirmed by soil gas surveys conducted at the landfill. Measurable amounts of these materials were identified in soil vapors and in shallow groundwater of the landfill.

#### 6.3.2. Monitoring Studies

In 1979, an investigation was conducted to determine the subsurface characteristics of the site and to place monitoring wells around the landfill (see Figure 6.6 and Table 6.24). The topography and initial studies indicated that water flow was primarily southerly. Wells 800011R and 800051 were located outside the landfill and were meant to measure water entering and leaving the landfill. Wells 800020, 800031, and 800040 were placed at the perimeter of the landfill. In April 1980, a more comprehensive study was initiated to develop information required for the State of Illinois operating permit.<sup>24</sup> Three additional wells were placed at the perimeter to improve coverage as well as to measure vertical movement. Well 800061 was placed in the eastern section to sample any water flowing out of the landfill in a southeasterly direction. Wells 800070 and 800071 were located along the southern boundary and were nested. In September 1986, six new wells were installed. Wells 800010, 800020, and 800040 were suspected of being poorly sealed and were removed and replaced by 800011R, 800021R and 800041R. The replacement wells were located within 2 m (6 ft) of the original wells. In addition, wells 800081, 800091, and 800101 were constructed to improve peripheral coverage. In November 1987, additional wells were added to provide sampling at a deeper level. Well 800121 is 10 m (32 ft). Well 800131, which is next to 800091, was installed to a depth of 24 m (78 ft). Finally, in September 1989, two wells (800141D and 800151D) were placed into the dolomite at depths of 45 m (148 ft) and 46 m (151 ft), respectively. These wells will no longer be sampled, effective January 1995.

During October 1992, 15 additional stainless steel wells, Wells Nos. 800161 through 800203D, were installed around the landfill as part of the IEPA-approved closure plan. These wells are required to be monitored as part of the IEPA-approved groundwater monitoring program beginning in 1995.

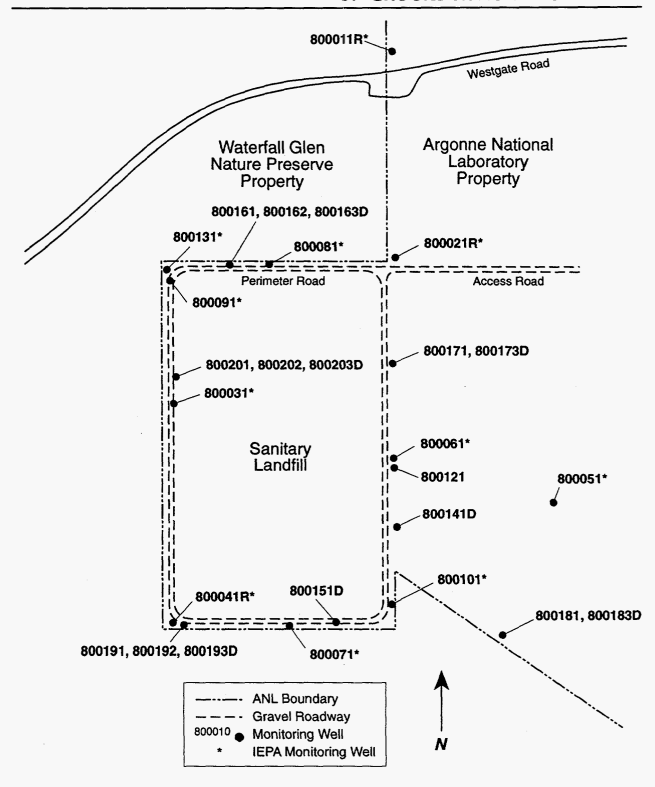


Figure 6.6 Active Monitoring Wells in the 800 Area Landfill (not to scale)

TABLE 6.24

Groundwater Monitoring Wells - 800 Area Landfill

ID Number	Well Depth <sup>1</sup>	Ground Elevation <sup>2</sup>	Monitoring Zone <sup>3</sup>	Well Type <sup>4</sup>	Date Drilled	Water Elevation
800011R	25	747.0	20-25/727-722	2/PVC	9/86	744.88
800021R	55	757.3	50-55/707-702	2/PVC	9/86	741.26
800031	27	744.0	17-27/727-717	2/PVC	10/79	737.02
800041R	25	745.5	20-25/725-720	2/PVC	9/86	740.24
800051	44	746.5	32-44/714-702	2/PVC	10/79	dry
800061	44	748.5	41-44/725-722	2/PVC	4/80	717.72
800071	25	747.4	22-25/725-722	2/PVC	4/80	733.22
800081	30	759.6	25-30/735-730	2/PVC	9/86	753.31
800091	20	754.5	15-20/739-734	2/PVC	9/86	748.06
800101	22	751.8	17-22/735-730	2/PVC	9/86	751.63
800121	32	748.5	27-32/721-716	4/SS	11/87	not measured
800131	78	754.5	68-78/686-676	4/SS	12/87	716.18
800141D	148	753.0	139-148/614-605	6/CS	9/88	not measured
800151D	151	747.4	147-151/600-596	6/CS	9/88	not measured
800161	25	757.1	20-25/737-732	2/SS	10/92	754.53
800162	70	757.0	65-70/692-687	2/SS	10/92	718.35
800163D	154	757.1	144-154/613-603	2/SS	9/92	642.78

TABLE 6.24 (Contd.)

ID Number	Well Depth <sup>1</sup>	Ground Elevation <sup>2</sup>	Monitoring Zone <sup>3</sup>	Well Type⁴	Date Drilled	Water Elevation
800171	25	749.4	20-25/729-724	2/SS	10/92	739.73
800173D	129	749.4	119-129/630-620	2/SS	10/92	632.84
800181	35	756.3	30-35/726-721	2/SS	10/92	734.65
800183D	164	755.8	154-164/602-592	<b>2/SS</b>	10/92	632.71
800191	15	746.0	10-15/736-731	2/SS	10/92	743.46
800192	60	746.0	55-60/691-686	2/SS	10/92	733.31
800193D	151	746.0	141-151/605-595	2/SS	10/92	632.72
800201	35	747.8	30-35/718-713	2/SS	10/92	734.42
800202	60	747.8	55-60/693-688	2/\$\$	10/92	718.35
800203D	126	747.8	116-126/632-622	2/SS	9/92	632.64

<sup>&</sup>lt;sup>1</sup>feet below ground

Note: Wells identified by a "D" are deeper wells monitoring the dolomite bedrock aquifer. Wells identified by an "R" are replacement wells.

<sup>&</sup>lt;sup>2</sup>feet mean sea level

<sup>&</sup>lt;sup>3</sup>depth/elevation

<sup>&</sup>lt;sup>4</sup>inner diameter (inches)/well material (PVC=polyvinyl chloride, SS=stainless steel, CS=carbon steel)

#### 6.3.2.1. Sample Collection

The same procedure for well water sample collection previously described for the 300 Area was used for this area. Previous water level measurements have indicated that a perched water layer exists at a depth of about 6 m (20 ft) on the north to about 7.6 m (25 ft) on the south. Wells 800011R through 800101 sample this layer. Well 800051 was dry during 1994. Wells 800121 and 800131, which are at depths of 10 m (32 ft) and 24 m (80 ft), respectively, exhibit very different characteristics. Well 800131 has an abundant supply of water [casing volume of about 100 L (27 gal)] while well 800131 is usually dry. It is not known if there is a water layer at this depth or if well 800131 is in a local body of water. The dolomite wells are at a depth of about 45 m (148 ft), and both have an abundant supply of water.

### 6.3.2.2. Sample Analysis - 800 Area

The 800 Area sample analyses were performed using SOPs written, reviewed, and issued as controlled documents by members of the ESH-DACH and ESH-DACE. These SOPs reference protocols found in SW-846, 3rd edition, "Test Methods for Evaluating Solid Waste." Sixteen metals were routinely determined. Ten metals were done using flame atomic absorption spectroscopy and inductively coupled plasma atomic emission spectroscopy and five metals were analyzed by graphite furnace atomic absorption spectroscopy. Mercury was determined by cold vapor atomic absorption spectroscopy. Volatile organic compounds were determined by using a purge and trap sample pretreatment followed by gas chromatography-mass spectroscopy detection. Semivolatile organic compounds were determined by solvent extraction followed by gas chromatography-mass spectroscopy detection. In the case of organic compound analyses, efforts were made to identify compounds which were present, but not included on the method list. This was accomplished and standard solutions of these compounds were prepared and analyzed. Total

dissolved solids (TDS) were determined gravimetrically. Sulfate determination was performed using a turbidimetric technique while chloride was determined by titrimetry. Ammonia was determined by using distillation followed by an ion selective specific electrode finish.

Some analyses were performed at an off-site contracted laboratory. SW-846 procedures were specified and used. PCB/pesticides were determined by solvent extraction followed by gas chromatography-electron capture detection. Cyanide and phenol were determined by distillation followed by a spectrophotometric finish. Total organic carbon and total organic halide were determined by combustion techniques followed by infrared detection and coulometric titration, respectively.

The 800 Area groundwater radiological analyses were performed using SOPs written, reviewed, and issued as controlled documents by members of the ESH-DARC Section. Hydrogen-3 was determined by distillation followed by a beta liquid scintillation counting technique. Strontium-90 was determined by an ion-exchange separation followed by a proportional counting technique.

#### 6.3.2.3. Results of Analyses

A description of each well, a listing of field parameters measured during sample collection, and the results of chemical and radiological analysis of samples from the wells in the 800 Area are contained in Tables 6.25 to 6.37. All radiological and inorganic analysis results are shown in these tables. The analysis methods used for organic compounds will identify and quantify all the compounds contained in the CLP Target Compound List. However, the vast majority of these compounds were not detected in the samples. Only those constituents which were present in amounts great enough to quantify are shown. The detection limits for the organic compounds listed were typically 1 to 5  $\mu$ g/L. The trend

TABLE 6.25

Groundwater Monitoring Results, Sanitary Landfill Well #800011R, 1994

Well Point Elevation 219.91
Ground Surface Elevation 227.69
Casing Material: PVC

Constituent	Units	01/24/94	04/04/94	07/11/94	10/04/94
Water Elevation	m	226.26	226.95	225.83	225.30
Temperature	°C	11.2	9.8	11.0	12.6
pH	рH	7.30	7.11	7.05	6.69
Redox	mV	-47	120	128	59
Conductivity	μmhos/cm	2310	2280	2450	2270
Cyanide (Total)	mg/L	< 0.002	0.003	< 0.002	< 0.002
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.135	0.124	0.131	0.112
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	0.04	0.08	0.27	< 0.04
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.235	0.214	0.194	0.180
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	0.034
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.050
Ammonia nitrogen	mg/L	0.3	0.2	0.3	0.3
Phenols	mg/L	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	800	756	837	687
Sulfate	mg/L	121	116	134	131
Total dissolved solids	mg/L	1663	1617	1845	1506
Total organic carbons	mg/L	1.7	1.9	1.1	1.3
Total organic carbons	mg/L	1.6	<b>1.6</b> .	1.8	1.5
Total organic carbons	mg/L	1.4	1.5	1.7	1.8
Total organic carbons	mg/L	1.5	1.5	1.7	1.3
Total organic halogens	mg/L	0.031	0.018	0.022	< 0.005
Total organic halogens	mg/L	0.020	0.021	0.023	< 0.005

6-60

TABLE 6.26

Groundwater Monitoring Results, Sanitary Landfill Well #800021R, 1994

Well Point Elevation 214.70
Ground Surface Elevation 230.83
Casing Material: PVC

Constituent	Units	01/24/94	01/24/94	04/04/94	07/11/94	10/04/94
Water Elevation	m	225.17	225.17	225.60	224.79	224.38
Temperature	°C	10.9	10.9	11.5	11.3	11.0
pH	рH	7.23	7.23	7.13	7.21	6.92
Redox	mV	47	47	104	23	98
Conductivity	μmhos/cm	638	638	657	652	659
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.124	0.125	0.106	0.112	0.110
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	0.0002	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.010
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	0.09	0.07	< 0.04	< 0.04	< 0.04
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.356	0.354	0.296	0.266	0.245
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	0.025	0.019	< 0.050
Ammonia nitrogen	mg/L	0.2	0.4	0.2	0.2	0.3
Phenols	mg/L	< 0.02	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nČi/L	< 0.100	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	16	17	17	19	18
Sulfate	mg/L	116	96	84	104	99
Total dissolved solids	mg/L	518	526	495	493	548
Total organic carbons	mg/L	1.1	2.5	1.1	1.2	1.2
Total organic carbons	mg/L	1.1	1.1	1.1	1.1	1.1
Total organic carbons	mg/L	1.4	1.2	1.1	1.0	1.0
Total organic carbons	mg/L	1.5	1.1	< 1.0	1.2	1.1
Total organic halogens	~	< 0.010	< 0.010	< 0.010	< 0.010	< 0.005
Total organic halogens	•	< 0.010	< 0.010	< 0.010	< 0.010	< 0.005
Total organic halogens		-	-	2	-	-

TABLE 6.27

Groundwater Monitoring Results, Sanitary Landfill Well #800031, 1994

Well Point Elevation 217.51
Ground Surface Elevation 226.77
Casing Material: PVC

Constituent	Units	01/24/94	04/05/94	07/14/94	10/05/94
Water Elevation	m	223.87	224.19	223.19	223.44
Temperature	°C	10.7	10.0	10.5	10.8
pH	pН	6.97	6.91	6.92	6.74
Redox	mV	- 69	- 46	7	21
Conductivity	μmhos/cm	1243	1244	1317	1521
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	< 0.003	0.002	< 0.006	0.002
Barium	mg/L	0.233	0.246	0.285	0.234
Beryllium	mg/L	< 0.0001	0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	7.614	2.517	8.224	0.099
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.192	0.134	0.254	0.174
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	0.020	0.025	0.034	0.037
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	0.011	< 0.010	< 0.050
Ammonia nitrogen	mg/L	0.4	0.9	1.0	1.0
Phenols	mg/L	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	18	8	25	12
Sulfate	mg/L	203	223	257	244
Total dissolved solids	mg/L	1156	971	1331	1212
Total organic carbons	mg/Ľ	26.9	26.6	25.8	31.9
Total organic carbons	mg/L	28.2	26.7	27.8	31.7
Total organic carbons	mg/L	27.8	27.4	27.7	32.0
Total organic carbons	mg/L	28.0	26.7	27.5	35.2
Total organic halogens	mg/L	< 0.010	0.014	< 0.010	< 0.005
Total organic halogens	mg/L	< 0.010	0.016	< 0.010	< 0.005

\_ ANL Site Environmental Report

TABLE 6.28

Groundwater Monitoring Results, Sanitary Landfill Well #800041R, 1994

Well Point Elevation 219.48
Ground Surface Elevation 227.23
Casing Material: PVC

Constituent	Units	01/24/94	04/04/94	04/04/94	07/11/94	10/04/94
Water Elevation	m	224.61	224.85	224.85	224.59	224.49
Temperature	°C	10.8	9.4	9.4	10.0	11.1
pH	рH	7.02	6.76	6.76	6.90	6.72
Redox	mV	-44	108	108	255	132
Conductivity	μmhos/cm	1022	988	988	997	1063
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.210	0.199	0.193	0.209	0.204
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	0.0002	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.04	< 0.04	< 0.04
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.015	0.013	0.027	0.033	0.010
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	0.021	0.021	< 0.020	< 0.020	0.024
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.050
Ammonia nitrogen	mg/L	0.1	0.1	0.5	0.1	0.8
Phenols	mg/L	< 0.02	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	< 0.100	< 0.100	-	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	-	< 0.25	< 0.25
Chloride	mg/L	106	102	106	96	90
Sulfate	mg/L	82	184	182	221	217
Total dissolved solids	mg/L	815	837	839	884	898
Total organic carbons	mg/L	4.4	4.2	5.3	4.9	4.8
Total organic carbons	mg/L	4.4	4.5	4.5	4.5	3.8
Total organic carbons	mg/L	4.4	4.4	4.5	4.5	4.0
Total organic carbons	mg/L	4.5	4.2	4.5	4.8	4.1
Total organic halogens	mg/L	< 0.010	0.018	0.018	< 0.010	< 0.005
Total organic halogens	mg/L	< 0.010	< 0.010	0.010	< 0.010	< 0.005

TABLE 6.29

Groundwater Monitoring Results, Sanitary Landfill Well #800061, 1994

Well Point Elevation
Ground Surface Elevation
Casing Material:

m(MSL) 206.89 229.91 PVC

Constituent	Units	01/24/94	04/04/94	07/11/94	10/05/94
Water Elevation	m	217.17	217.20	216.78	216.02
Temperature	°C	10.2	11.2	12.3	11.1
pН	pН	6.72	6.70	6.78	6.51
Redox	mV	- 43	-13	-14	- 10
Conductivity	μmhos/cm	1343	1369	1420	1426
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.161	0.150	0.175	0.168
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	6.59	0.08	1.10	0.57
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.6020	0.6250	0.5450	0.0672
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	0.026	0.028	0.034
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zine	mg/L	< 0.010	< 0.010	0.015	< 0.050
Ammonia nitrogen	mg/L	0.3	0.5	0.6	0.7
Phenois	mg/L	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	0.541	0.570	0.585	0.623
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	225	234	231	234
Sulfate	mg/L	16	21	14	14
Total dissolved solids	mg/L	985	1003	1007	1064
Total organic carbons	mg/L	10.6	9.9	9.1	9.1
Total organic carbons	mg/L	9.4	8.2	9.4	9.0
Total organic carbons	mg/L	9.3	9.0	9.2	9.2
Total organic carbons	mg/L	9.3	8.9	9.4	8.8
Total organic halogens	mg/L	0.130	0.250	0.166	0.056
Total organic halogens	mg/L	0.142	0.250	0.104	0.059
1,2 Dichloroethane	μg/L	•	-	_	0.6
1,4-Dioxane	μg/L	201	173	220	185
Acetone	μg/L	-	5	-	5
Chlorodifluoromethane	μg/L	82	258	67	199
Ethyl Ether	μg/L	7	9	5	4
Methylene chloride	μg/L	-	-	-	1
Tetrahydrofuran	μg/L	31	23	33	32

TABLE 6.30

Groundwater Monitoring Results, Sanitary Landfill Well #800071, 1994

Well Point Elevation
Ground Surface Elevation
Casing Material:

m(MSL) 220.05 227.80 PVC

Water Elevation         m         221.8           Temperature         °C         10.7           pH         pH         7.0           Redox         mV         -50           Conductivity         μmhos/cm         800           Cyanide (Total)         mg/L         < 0.0           Arsenic         mg/L         < 0.0           Barium         mg/L         < 0.0           Beryllium         mg/L         < 0.0           Cadmium         mg/L         < 0.0           Chromium         mg/L         < 0.0           Cobalt         mg/L         < 0.0           Copper         mg/L         < 0.0           Iron         mg/L         0.1	4/94 04/04/94 07/11/94 10/04/94
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	89 222.24 221.45 221.20
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	7 10.0 10.7 11.2
$\begin{array}{llllllllllllllllllllllllllllllllllll$	02 6.97 7.24 7.07
$\begin{array}{llllllllllllllllllllllllllllllllllll$	98 158 125
$\begin{array}{llllllllllllllllllllllllllllllllllll$	810 729 771
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	002 < 0.002 < 0.002 < 0.002
$\begin{array}{llllllllllllllllllllllllllllllllllll$	002 < 0.002 < 0.002 < 0.002
$\begin{array}{llllllllllllllllllllllllllllllllllll$	0.080 0.071 0.096
$\begin{array}{llllllllllllllllllllllllllllllllllll$	0001 < 0.0001 < 0.0001 < 0.0001
$\begin{array}{lll} \mbox{Cobalt} & \mbox{mg/L} & < 0.0 \\ \mbox{Copper} & \mbox{mg/L} & < 0.0 \end{array}$	0001 < 0.0001 < 0.0001 < 0.0001
Copper mg/L < 0.0	020 < 0.020 < 0.020 < 0.020
••	020 < 0.020 < 0.020 < 0.100
	010 < 0.010 < 0.010 < 0.065
11011 111g/L 0.1	12 < 0.04 < 0.04 0.34
Lead mg/L < 0.0	0.0010 < 0.0010 < 0.0010 < 0.0010
Manganese mg/L 0.0	058 0.209 0.168 0.079
Mercury $mg/L$ < 0.0	0001 < 0.0001 < 0.0001 < 0.0001
Nickel mg/L < 0.0	020 < 0.020 0.020 0.028
Silver $mg/L$ < 0.0	001 < 0.001 < 0.001 < 0.001
Thallium $mg/L$ < 0.0	0.0030 < 0.0030 < 0.0030 < 0.0030
Vanadium mg/L < 0.2	200 < 0.200 < 0.200 < 0.200
Zinc $mg/L$ < 0.0	010 < 0.010 0.022 < 0.050
Ammonia nitrogen mg/L 0.6	6 0.2 0.1 0.7
Phenols $mg/L$ < 0.0	02 < 0.02 < 0.02 < 0.02
Hydrogen-3 nCi/L 0.4	491 0.300 0.207 0.715
Strontium-90 pCi/L < 0.2	25 < 0.25 < 0.25 < 0.25
Chloride mg/L 40	23 25 46
Sulfate mg/L 160	177 161 178
Total dissolved solids mg/L 746	667 550 716
Total organic carbons mg/L 2.2	2 2.1 2.0 3.7
Total organic carbons mg/L 2.2	2 1.8 2.1 4.3
Total organic carbons mg/L 2.2	2 2.0 2.0 5.3
Total organic carbons mg/L 2.4	4 2.2 1.9 3.9
Total organic halogens mg/L 0.0	
Total organic halogens mg/L < 0.0	
1,2 Dichloroethane $\mu g/L$ -	0.5
Acetone $\mu g/L$ -	- 4
Methylene Chloride $\mu g/L$ -	
Tetrahydrofuran μg/L -	2

TABLE 6.31

Groundwater Monitoring Results, Sanitary Landfill Well #800081, 1994

Well Point Elevation 218.71
Ground Surface Elevation 231.53
Casing Material: PVC

Constituent	Units	01/24/94	04/04/94	07/11/94	10/05/94
Water Elevation	m	227.49	231.53	226.40	225.88
Temperature	°C	10.6	9.6	10.6	10.9
pH	pН	6.99	6.92	6.95	6.96
Redox	mV	-43	114	207	147
Conductivity	μmhos/cm	1190	1180	1196	1083
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.052	0.052	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	0.0010
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.04	< 0.04
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.758	0.815	0.549	0.397
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	0.026	0.028	0.025	0.026
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	0.025	< 0.050
Ammonia nitrogen	mg/L	0.4	0.2	0.1	0.2
PhenoIs	mg/L	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	< 0.100	0.137	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	101	101	89	70
Sulfate	mg/L	314	286	285	288
Total dissolved solids	mg/L	1074	1105	1088	966
Total organic carbons	mg/L	2.8	2.5	2.9	2.2
Total organic carbons	mg/L	2.7	2.8	3.1	2.2
Total organic carbons	mg/L	2.7	2.8	2.8	2.2
Total organic carbons	mg/L	3.0	2.6	3.0	2.1
Total organic halogens	mg/L	0.036	0.025	0.013	0.009
Total organic halogens	mg/L	0.032	0.023	0.013	0.006
Acetone	$\mu g/L$	-	2	-	-
Chlorodifluoromethane	$\mu$ g/L	5	9	-	-

TABLE 6.32

Groundwater Monitoring Results, Sanitary Landfill Well #800091, 1994

Well Point Elevation
Ground Surface Elevation
Casing Material:

m(MSL) 223.79 230.00 PVC

Constituent	Units	01/28/94	04/08/94	07/14/94	07/14/94	10/10/94
Water Elevation	m	227.30	227.44	226.51	226.51	226.14
Temperature	°C	10.8	9.7	10.7	10.7	11.3
pH	pН	6.85	6.88	6.90	6.90	6.75
Redox	mV	-18	-5	<b>-4</b> 1	-41	79
Conductivity	μmhos/cm	1133	1054	1214	1214	1295
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	0.005	0.004	0.006	0.005	0.007
Barium	mg/L	0.259	0.235	0.282	0.277	0.253
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	4.00	4.78	4.78	7.63	18.11
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.589	0.554	0.832	0.832	1.397
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	0.020	< 0.020	0.020	0.021	0.040
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.050
Ammonia nitrogen	mg/L	0.2	0.3	1.0	0.4	0.5
Phenois	mg/L	< 0.02	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	0.370	1.859	0.497	0.446	0.511
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	99	76	111	116	126
Sulfate	mg/L	8	35	41	26	34
Total dissolved solids	mg/L	852	854	927	961	1049
Total organic carbons	mg/L	10.2	11.2	14.6	13.8	23.8
Total organic carbons	mg/L	11.5	11.2	13.8	13.6	26.3
Total organic carbons	mg/L	11.2	11.1	13.4	15.0	13.5
Total organic carbons	mg/L	11.5	13.1	14.6	15.0	20.3
Total organic halogens	mg/L	0.070	0.223	0.059	0.057	0.052
Total organic halogens	mg/L	0.071	0.112	0.070	0.076	0.049
1,4-Dioxane	μg/L	158	141	141	61	188
Acetone	μg/L	-	-	-	-	4
Chlorodifluoromethane	μg/L	67	118	52	<i>5</i> 0	188
Ethyl Ether	μg/L	6	7	4	4	5
Methylene Chloride	μg/L	-	1	-	-	3
Nonanal	μg/L	-	-	34	-	-
Octanal	μg/L	-	-	26	-	-
Tetrahydrofuran	μg/L	-	-	-	-	1

# 6. GROUNDWATER PROTECTION

TABLE 6.33

Groundwater Monitoring Results, Sanitary Landfill Well #800101, 1994

Well Point Elevation 222.28
Ground Surface Elevation 229.15
Casing Material: PVC

			Casing iv	iaici iai.	PVC
Constituent	Units	01/28/94	04/05/94	07/14/94	10/10/94
Water Elevation	m	227.80	228,31	227.05	226.39
Temperature	°C	10.1	8.5	11.4	13.0
рН	pН	7.01	7.36	7.24	6.98
Redox	mV	64	25	219	24
Conductivity	μmhos/cm	675	648	681	725
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	0.0037	0.0020	0.0038	0.0043
Barium	mg/L	0.059	0.055	0.061	0.056
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	1.32	0.99	< 0.04	0.88
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.106	0.136	0.136	0.151
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	0.024
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.050
Ammonia nitrogen	mg/L	0.6	0.2	0.2	0.6
Phenols	mg/L	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	5	5	6	8
Sulfate	mg/L	210	175	221	162
Total dissolved solids	mg/L	551	592	568	597
Total organic carbons	mg/L	1.5	1.2	1.8	1.3
Total organic carbons	mg/L	1.3	1.4	1.6	2.1
Total organic carbons	mg/L	1.3	1.3	2.1	1.2
Total organic carbons	mg/L	1.4	1.4	1.7	1.2
Total organic halogens	mg/L	< 0.010	< 0.010	< 0.010	< 0.005
Total organic halogens	mg/L	< 0.010	< 0.010	< 0.010	< 0.005
Acetone	μg/L	-	-	-	4
Methylene Chloride	μg/L	_	-	_	3

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TABLE 6.34

Groundwater Monitoring Results, Sanitary Landfill Well #800121, 1994

	m(MSL)
Well Point Elevation	219.17
Ground Surface Elevation	229.91
Casing Material:	S. Steel

Constituent	Units	01/24/94	04/05/94	07/20/94	10/05/94
Water Elevation	m	221.72	222.09	221.08	220.62
Temperature	°C	10.8	10.5	11.1	10.8
pH	рH	7.39	7.49	7.26	6.95
Redox	mV	-43	43	-165	-12
Conductivity	μmhos/cm	902	720	915	1021
Arsenic	mg/L	0.006	0.006	0.004	0.004
Barium	mg/L	0.059	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	0.019	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.04	< 0.04
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	< 0.010	0.019	< 0.010	0.056
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	0.022	0.022	0.020	0.032
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	0.019	< 0.050
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Chloride	mg/L	27	34	34	33
Sulfate	mg/L	137	223	246	246
Total dissolved solids	mg/L	862	677	827	940
Acetone	μg/L	13	_	-	-
Methylene Chloride	μg/L	-	2	-	3

TABLE 6.35

Groundwater Monitoring Results, Sanitary Landfill Well #800131, 1994

Well Point Elevation 205.66
Ground Surface Elevation 230.00
Casing Material: S. Steel

Constituent	Units	01/28/94	04/08/94	07/14/94	10/12/94	10/12/94
Water Elevation	m	216.77	217.08	214.79	216.90	216.90
Temperature	°C	10.9	10.9	11.3	11.0	11.0
pH	рH	7.05	6.98	6.94	6.82	6.82
Redox	mV	-83	-30	-58	-78	-78
Conductivity	μmhos/cm	704	774	702	719	719
Cyanide (Total)	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Arsenic	mg/L	0.0038	0.0031	0.0033	0.0043	0.0045
Barium	mg/L	< 0.050	0.149	0.146	0.150	0.147
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065	< 0.065
Iron	mg/L	4.23	3.95	3.90	3.56	3.48
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.092	0.189	0.095	0.081	0.079
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	0.022
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.020	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.050	< 0.050
Ammonia nitrogen	mg/L	0.4	0.2	0.7	1.5	1.5
Phenols	mg/L	< 0.02	< 0.02	< 0.02	< 0.02	< 0.02
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100	< 0.100
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	30	41	35	35	32
Sulfate	mg/L	94	95	98	89	93
Total dissolved solids	mg/L	524	579	553	557	542
Total organic carbons	mg/L	4.8	6.1	5.0	6.3	4.4
Total organic carbons	mg/L	4.8	6.1	4.8	4.4	4.6
Total organic carbons	mg/L	4.9	6.3	4.6	5.2	4.3
Total organic carbons	_	4.9	6.2	4.7	4.3	4.2
Total organic halogens	•	< 0.010	0.250	0.027	< 0.005	0.0054
Total organic halogens		< 0.010	0.290	0.011	< 0.005	0.0052
Chlorodifluoromethane	_	-	-	10	-	-
Methylene chloride	μg/L	-	-	-	2	_

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TABLE 6.36

Groundwater Monitoring Results, Sanitary Landfill Well #800141D, 1994

Well Point Elevation 183.13
Ground Surface Elevation 229.53
Casing Material: Steel

Constituent	Units	02/07/94	04/08/94	07/20/94	10/14/94
Water Elevation	m	192.91	193.06	192.98	192.94
Temperature	°C	11.3	11.4	11.6	11.4
pH	pН	7.25	7.15	7.16	6.89
Redox	mV	-89	-52	-257	-307
Conductivity	μmhos/cm	926	928	974	1012
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.058	0.063	0.066	0.062
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.065
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010
Iron	mg/L	1.19	1.29	1.24	1.40
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.018	0.019	0.016	0.022
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	0.021	0.032
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.050
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Chloride	mg/L	139	135	140	126
Sulfate	mg/L	129	173	225	204
Total dissolved solids	mg/L	746	745	768	772
Methylene Chloride	μg/L	-	1	-	-

# 6. GROUNDWATER PROTECTION

TABLE 6.37

Groundwater Monitoring Results, Sanitary Landfill Well #800151D, 1994

Well Point Elevation 182.31
Ground Surface Elevation 227.81
Casing Material: Steel

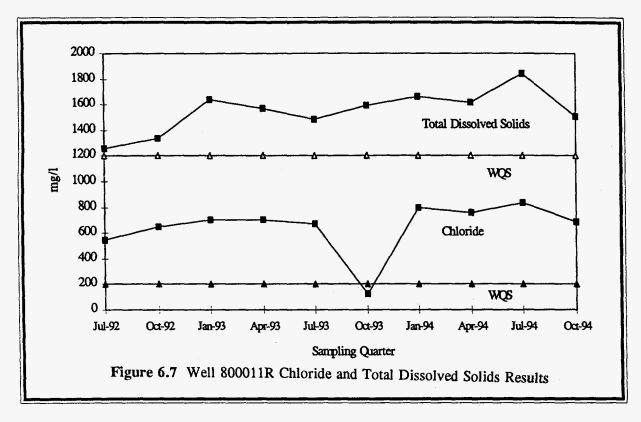
Constituent	Units	02/04/94	04/04/94	07/18/94	10/14/94
Water Elevation	m	191.90	192.19	191.88	191.84
Temperature	°C	11.3	11.3	11.4	11.4
pH	pН	7.28	7.23	7.12	7.00
Redox	mV	-111	-74	-61	-256
Conductivity	μmhos/cm	899.	902	896	931
Arsenic	mg/L	0.003	0.003	0.002	0.003
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	0.924	1.109	1.108	1.108
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.016	0.017	0.013	0.020
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.0030	< 0.0030	< 0.0030	< 0.0030
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.010	< 0.050
Hydrogen-3	nCi/L	< 0.100	< 0.100	< 0.100	< 0.100
Chloride	mg/L	126	117	112	119
Sulfate	mg/L	192	168	242	222
Total dissolved solids	mg/L	690	719	706	684
Acetone	μg/L	-	-	-	3
Methylene chloride	μg/L	-	2	-	1

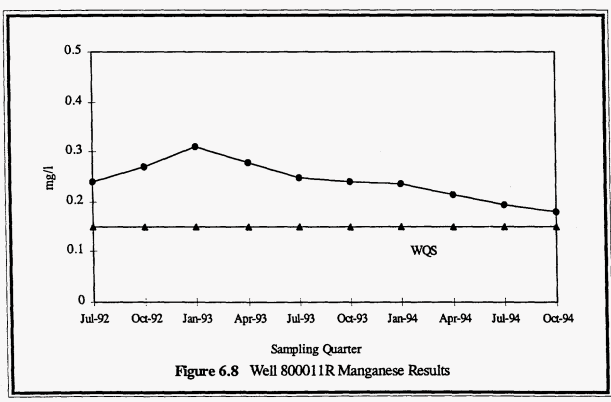
for exceedances of the WQS for wells monitored as part of the IEPA-approved groundwater monitoring program for the sanitary landfill are presented in Figures 6.7 to 6.21.

# Inorganic Constituents

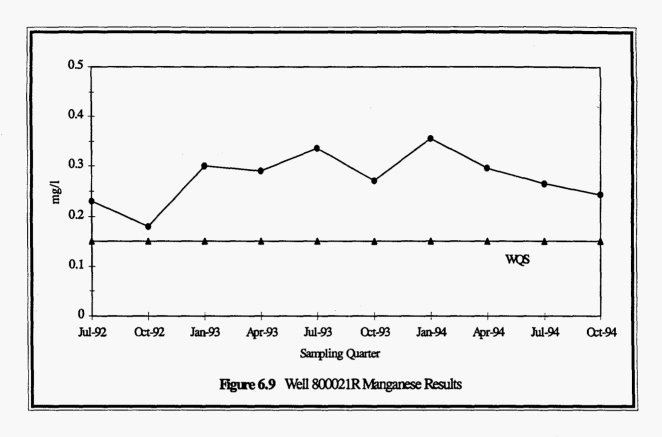
On April 24, 1992, and January 11, 1995, the IEPA issued supplemental permits to ANL which in part approved a groundwater monitoring program for the sanitary landfill. The program is to be capable of identifying any releases from the facility and demonstrate compliance with the applicable groundwater quality standards. Under the April 1992 supplemental permit, IEPA chose 11 groundwater monitoring points (800011R, 800021R, 800031, 800041R, 800051, 800061, 800071, 800081, 800091, 800101, 800131) to be sampled on a quarterly basis commencing July 1992. Parameters to be monitored include field parameters, routine indicator parameters, and volatile organic parameters. Volatile organic parameters are to be monitored only during the second quarter of monitoring.

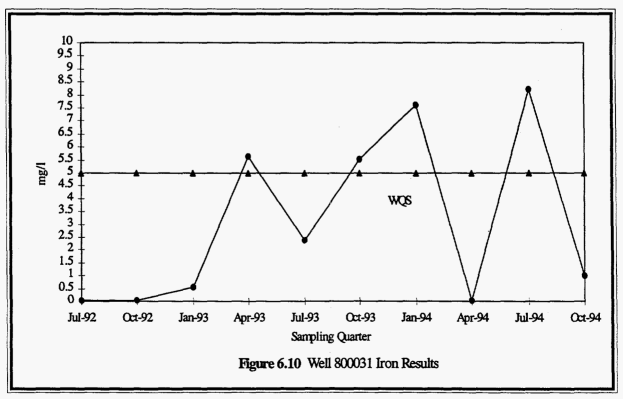
The Illinois Groundwater Quality Standards for Class I: Potable Resource Groundwater, 35 IAC Section 620.410, were used as the standard for evaluation of the inorganic results. Inorganic results are fairly consistent with results reported in previous years. The most common constituents at levels above the WQS (see Table 6.21) are chloride, iron, total dissolved solids (TDS), and manganese. Wells 800031, 800061, and 800091 exceeded the WQS for iron. Iron levels in these wells ranged from 6.6 to 18.1 mg/L. The TDS levels were exceeded in wells 800011R and 800031 and ranged from 1212 to 1845 mg/L. The manganese WQS was exceeded in wells 800011R, 800021R, 800031, 800061, 800071, 800081, 800091, and 800131. Manganese levels in these wells ranged from 0.17 to 1.4 mg/L. The WQS for chloride was consistently exceeded in wells 800011R and 800061 where the levels vary from 225 to 800 mg/L. The inorganic results for dolomite wells 800141D and 800151D were within normal ranges.

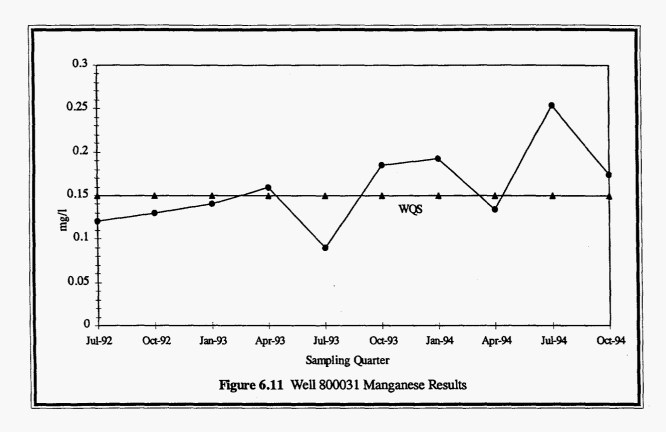


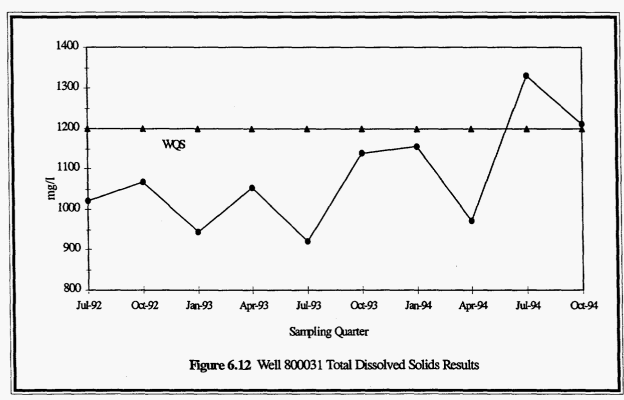


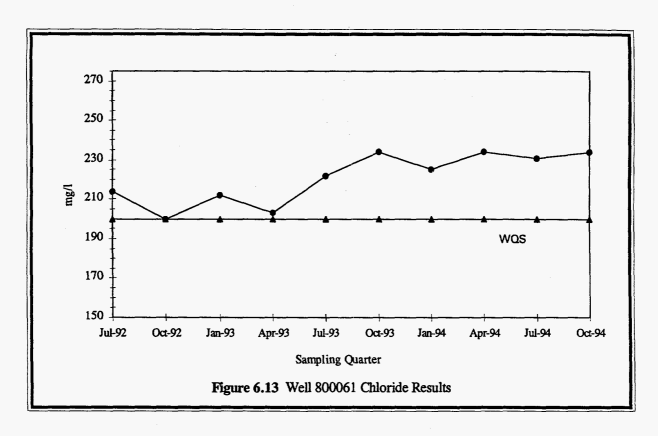
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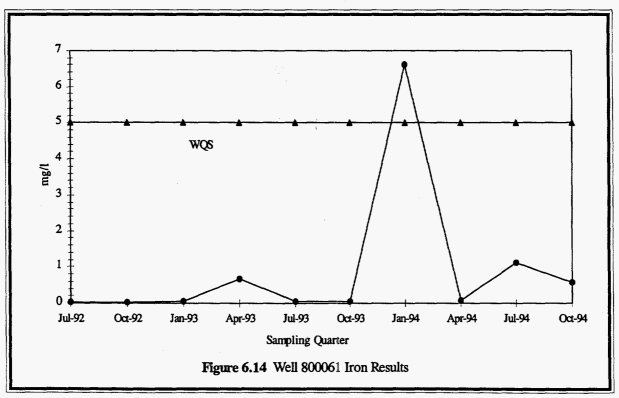


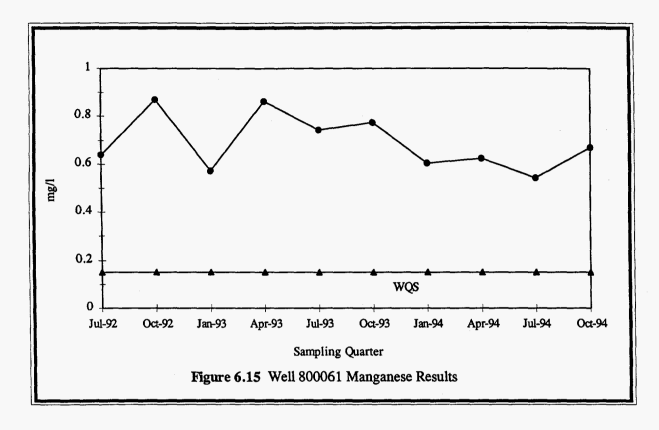


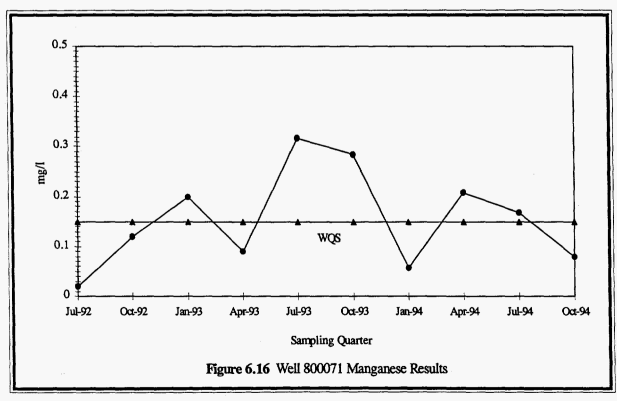


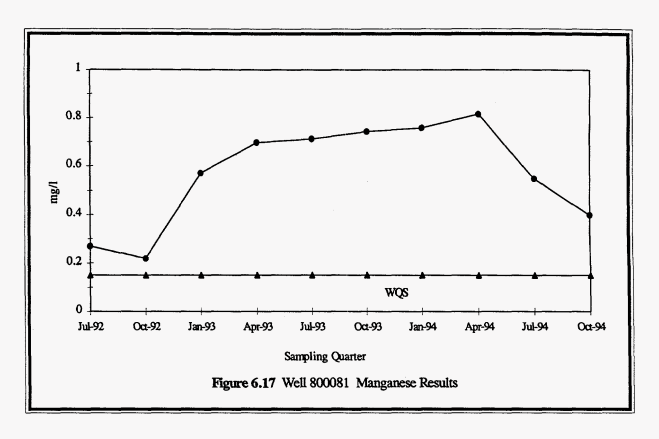


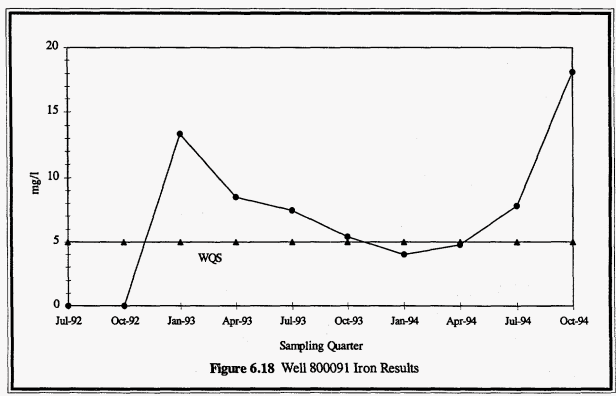


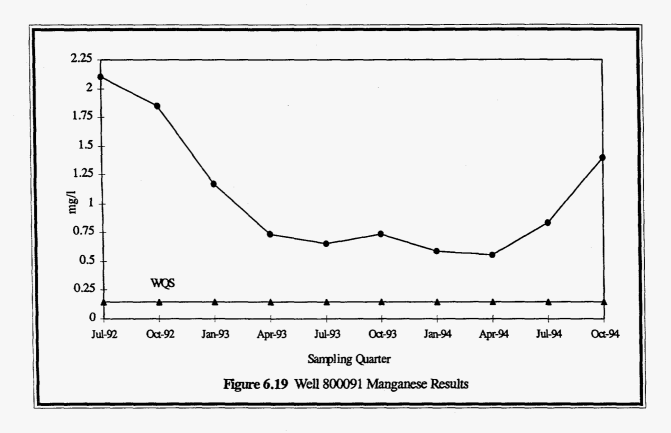


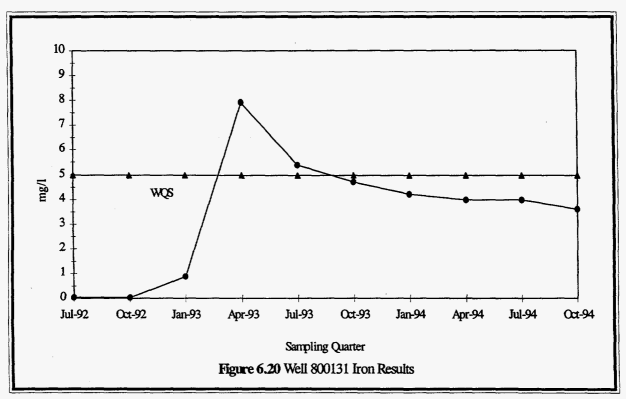


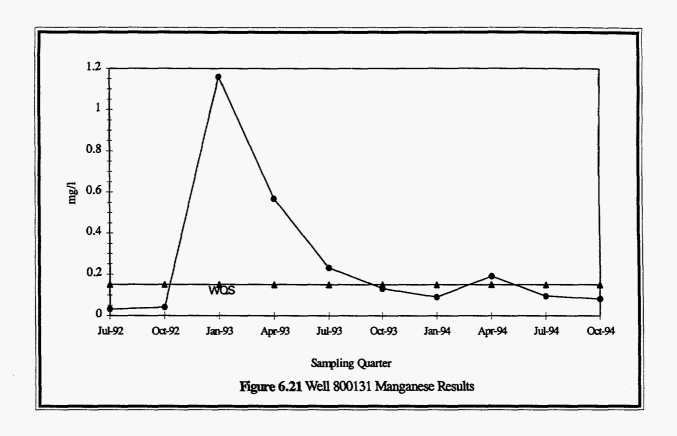












# Organic Constituents

No exceedances of the organic WQS were noted during 1994. Contrary to 1993, trace levels of methylene chloride were identified in more wells but usually for just one quarter. These inconsistent results may be indicative of laboratory contamination. During one quarter, trace levels of methylene chloride were identified in wells 800021R, 800061, 800071, 800101, 800131, and 800141D. During two quarters, trace levels of methylene chloride were identified in wells 800091, 800121, and 800151D.

As noted in 1993, ethyl ether was found in trace amounts in wells 800061 and 800091; chlorodifluoromethane was identified in wells 800061, 800081, 800091, and 800131; and 1-4-dioxane and tetrahydrofuran were identified in well 800061. Tetrahydrofuran was also detected but only during one quarter in wells 800071 and 800091. Trace levels of 1,2-dichloroethane were found in only one quarter in wells 800061 and 800071.

## Radioactive Constituents

Samples collected from the 800 Area sanitary landfill monitoring wells were also analyzed for hydrogen-3 and strontium-90. The results are shown in Tables 6.25 to 6.37. Although the disposal of radioactive materials was prohibited in the sanitary landfill, very low concentrations of hydrogen-3 were detected in wells 800061, 800071, 800081 and 800091, probably due to inadvertent disposal of radioactivity in the ANL trash. However, the presence of hydrogen-3 as tritiated water allows information to be obtained on the subsurface water flow pathway in the sanitary landfill area. The data indicate that the principal direction of subsurface water flow is to the south-southeast, with a small component to the northwest. This is consistent with the estimated subsurface water flow based on

water level measurements and general flow patterns in the area. No strontium-90 was detected.

## 6.4. CP-5 Reactor Area

The CP-5 reactor is an inactive research reactor located in Building 330 (see Figure 1.1). CP-5 was a 5 megawatt research reactor which was used from 1954 until operations were ceased in 1977. In addition to the reactor vessel itself, the CP-5 complex contained several large cooling towers and an outdoor equipment yard used for storage of equipment and supplies. The reactor and associated yard area is in the process of being decommissioned. A single exploratory monitoring well was installed in 1989 in the yard, immediately behind the reactor building, just outside the reactor fuel storage area of the complex. This well (330011) was sampled quarterly in 1994 and analyzed for radionuclides, metals, and volatile organic compounds. A sample collected in January was also analyzed for semivolatiles, pesticides, herbicides and polychlorinated biphenyls. Two new wells were installed as part of a full characterization study of this site which took place during 1993 (Section 6.5.3.) and were sampled quarterly in 1994. The results are shown in Tables 6.38 to 6.40. All wells in this area are described in Table 6.41 (see Figure 6.22 for location).

Well 330011 is installed in a relatively porous, unsaturated region of soil and as a result, recharges quickly. Purging the well by removing several well volumes of water does not lower the water level appreciably. The water has a higher conductivity than similar wells at other locations. The chloride WQS was exceeded in one quarter and levels ranged from 65  $\mu$ g/L to 231  $\mu$ g/L. Manganese concentrations are elevated and the WQS for manganese was exceeded during each quarter. Levels ranged from 0.74 mg/L to 0.95 mg/L. Low levels of barium, nickel, and iron were found, all well below the WQS.

TABLE 6.38

Groundwater Monitoring Results, 300 Area Well #330011, 1994

Well Point Elevation 215.70
Ground Surface Elevation 222.56
Casing Material: Steel

Constituent	Units	03/15/94	06/22/94	09/16/94	11/29/94
Water Elevation	m	219.45	219.13	219.32	219.59
Temperature	°C	12.8	14.7	18.3	17.4
pH	pН	6.90	6.92	7.00	6.61
Redox	mV	148	145	273	276
Conductivity	μmhos/cm	967	1356	1492	1185
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	0.0570	0.078	0.080	0.070
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0012	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	0.11	0.30	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.850	0.908	0.954	0.744
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	0.026	0.024	0.025
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.003	< 0.003	< 0.003	< 0.003
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	0.075	< 0.050	< 0.050
Cesium-137	pCi/L	2.37	< 1.00	< 1.00	< 1.00
Hydrogen-3	nCi/L	5.504	14.280	14.980	7.754
Strontium-90	pCi/L	0.75	2.71	2.53	1.54
Chloride	mg/L	65	231	186	83
Dichlorofluoromethane	μg/L	4	11	15	5
Methylene Chloride	μg/L	-	-	1	-
Trichlorofluoromethane	μg/L	3	-	20	6

TABLE 6.39

Groundwater Monitoring Results, 300 Area Well #330021, 1994

Well Point Elevation 215.88
Ground Surface Elevation 227.52
Casing Material: S. Steel

Constituent	Units	03/14/94	03/14/94	06/22/94	09/19/94	11/29/94
Water Elevation	m	226.70	226.70	225.31	224.96	226.09
Temperature	°C	7.0	7.0	10.1	13.2	12.2
pH	Нq	7.35	7.35	7.12	7.32	6.92
Redox	mV	172	172	126	353	275
Conductivity	μmhos/cm	527	527	568	620	642
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.04	< 0.14	0.87
Lead	mg/L	< 0.0010	< 0.0010	< 0.0020	< 0.0010	< 0.0010
Manganese	mg/L	0.026	0.028	0.142	0.081	0.415
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0 .0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	0.067	< 0.020	0.918
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.003	< 0.003	< 0.003	< 0.003	< 0.003
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	0.032	< 0.050	< 0.050
Cesium-137	pČi/L	< 1.00	1.41	< 1.00	< 1.00	< 1.00
Hydrogen-3	nCi/L	0.201	0.259	0.144	0.482	0.303
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	10	11	6	5	12
Acetone	μg/L	-	_	-	-	8
Methylene Chloride	μg/L	-	-	-	-	2

TABLE 6.40

Groundwater Monitoring Results, 300 Area Well #330031, 1994

Well Point Elevation 220.97
Ground Surface Elevation 226.18
Casing Material: S. Steel

Constituent	Units	03/14/94	06/22/94	09/19/94	11/19/94
Water Elevation	m	226.12	224.98	225.24	225.84
Temperature	°C	8.9	10.7	13.5	12.3
pH	pН	7.27	7.20	7.30	6.87
Redox	mV	173	130	349	279
Conductivity	$\mu$ mhos/cm	744	787	900	884
Arsenic	mg/L	< 0.002	< 0.002	< 0.002	< 0.002
Barium	mg/L	< 0.050	< 0.050	< 0.050	< 0.050
Beryllium	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Cadmium	mg/L	< 0.0001	< 0.0001	< 0.0001	0.0004
Chromium	mg/L	< 0.020	< 0.020	< 0.020	< 0.020
Cobalt	mg/L	< 0.020	< 0.020	< 0.100	< 0.100
Copper	mg/L	< 0.010	< 0.010	< 0.010	< 0.065
Iron	mg/L	< 0.04	< 0.04	< 0.14	< 0.14
Lead	mg/L	< 0.0010	< 0.0010	< 0.0010	< 0.0010
Manganese	mg/L	0.024	0.090	0.061	0.036
Mercury	mg/L	< 0.0001	< 0.0001	< 0.0001	< 0.0001
Nickel	mg/L	< 0.020	< 0.020	< 0.020	< 0.026
Silver	mg/L	< 0.001	< 0.001	< 0.001	< 0.001
Thallium	mg/L	< 0.003	< 0.003	< 0.003	< 0.003
Vanadium	mg/L	< 0.200	< 0.200	< 0.200	< 0.200
Zinc	mg/L	< 0.010	< 0.010	< 0.050	< 0.050
Cesium-137	pČi/L	< 1.00	< 1.00	< 1.00	< 1.00
Hydrogen-3	nCi/L	0.329	0.458	0.220	0.438
Strontium-90	pCi/L	< 0.25	< 0.25	< 0.25	< 0.25
Chloride	mg/L	41	37	39	53
Acetone	μg/L	-	-	-	5
Methylene Chloride	μg/L	-	-	2	-

Manganese was detected each quarter in wells 330021 and 330031 but the WQS was exceeded only during one quarter in well 330021.

Similar to results in 1991, 1992, and 1993 collected from well 330011, three of the samples collected and analyzed in 1994 contained trichlorofluoromethane ranging from 3  $\mu$ g/L to 20  $\mu$ g/L. These levels are significantly lower than those levels reported in 1991 and 1992. Each quarter, dichlorofluoromethane was noted at levels ranging from 4  $\mu$ g/L to 15  $\mu$ g/L.

Methylene chloride was noted at trace levels for only one quarter in each well. Trace levels of acetone were noted for only one quarter in wells 330021 and 33031. Acetone and methylene chloride are found in blanks and their presence may be indicative of laboratory contamination.

TABLE 6.41

Groundwater Monitoring Wells - 330 Area/CP-5

ID Number	Well Depth <sup>1</sup>	Ground Elevation <sup>2</sup>	Monitoring Zone <sup>3</sup>	Well Type⁴	Date Drilled
330011	20	745.5	10-20/736-726	2/PVC	8/89
330021	19	746.5	4-19/743-728	2/SS	9/93
330031	17.1	742.1	2-17/740-725	2/SS	9/93

<sup>&</sup>lt;sup>1</sup>Feet below ground

<sup>&</sup>lt;sup>2</sup>Feet mean sea level

<sup>&</sup>lt;sup>3</sup>Depth/elevation

<sup>&</sup>lt;sup>4</sup>Inner diameter (inches)/well material (PVC = polyvinyl chloride, SS = stainless steel)

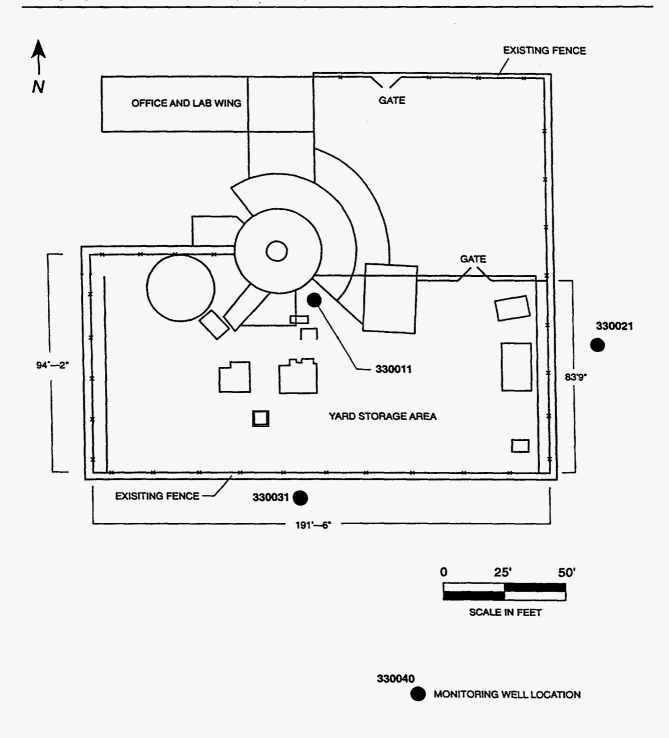


Figure 6.22 Active Monitoring Wells in the CP-5 Reactor Area

The levels of hydrogen-3 ranged from 0.14 to 14.98 nCi/L and the levels of strontium-90 ranged from < 0.25 to 2.7 pCi/L and was noted only in well 330011. Cesium-137 ranged from < 1 to 2.4 pCi/L. CP-5 was a heavy water-moderated reactor. During its operational life, several incidents occurred which released small amounts of this heavy water containing high concentrations of hydrogen-3, to the environment. In addition, the normal operation released significant amounts of water vapor containing hydrogen-3 from the main ventilation system which may have condensed and fallen to the ground in the form of precipitation. These activities are believed to be responsible for the residual amounts of hydrogen-3 now found in the groundwater. The source of the strontium-90 is not known.

## 6.5. Site Characterization Activities

Historical information about waste disposal activities on the ANL site, as well as groundwater monitoring results, indicate that several sites are either currently releasing small amounts of hazardous materials to the environment or have the potential to do so in the future. As a first step to stopping these releases and cleaning up any residual contamination, a series of site characterization projects is underway. To date, these projects have focused on the most significant sites, the 800 Area landfill and the 317/319 Areas. The studies are in the characterization stage, and thus the information available is currently incomplete and may not accurately represent the actual conditions at these sites. Characterization activities are currently scheduled to extend beyond 1995.

## 6.5.1. 317/319 Area Characterization

Preliminary characterization conducted prior to 1994 in the 317/319 Area indicates that two distinct areas of highly contaminated soil exist, one near the site of the French drain in the 317 Area and the other in the 319 Area landfill. A larger number of organic compounds

were identified in the shallow groundwater in the 317 Area, some at very high concentrations (over  $100,000~\mu g/L$ ). A relatively small area of highly contaminated soil was found to exist, just north of the vaults used for storage of radioactive wastes. Significant, but much lower concentrations of volatile organics were found several hundred feet south of the vault area, indicating that movement of the contamination through the soil is occurring. This is consistent with the results of the monitoring well sampling discussed in this chapter. Samples of shallow groundwater [less than 3 m (10 ft) deep] collected several years ago on Forest Preserve property south of the ANL fenceline indicate that low levels of several volatile organic compounds may have moved off-site. The depth and extent of groundwater contamination is not fully defined at this point.

The 319 Area, which contained a similar French drain, was also found to contain a large number of organic compounds, although the concentrations were much lower than in the 317 Area. The French drain in this area was much deeper than the one in the 317 Area. Since the techniques used in this preliminary investigation were limited to a depth of approximately 3 m (10 ft) below the surface, they may not have been able to detect contamination located deep within the 319 waste pile.

One sample recovered from the 319 Area was found to contain low concentrations of two PCBs, Aroclor 1254 and Aroclor 1260 (220  $\mu$ g/L total). A floating oil layer was encountered at this point, indicating the PCBs were the result of disposal of PCB-containing waste oils.

During 1993, additional preliminary characterization activities were conducted in this area to further define the extent of contamination from the French drain, inactive landfills, and other waste units in this area. A series of geophysical investigations were performed. The non-intrusive investigation techniques were able to identify the presence of buried waste material, as well as provide a description of the underlying soils. More detailed

investigations of the soil were carried out using a technique called Cone Penetration Testing. This technique involved pushing an instrumented probe into the soil while recording data on soil composition, groundwater depth, and soil permeability. Samples of groundwater were also collected from porous soil layers containing water. Temporary wells were installed in the shallow surface soils near the 317 Area French drain and 319 Area landfill and additional groundwater samples were collected. The samples were analyzed for volatile organic chemicals, metals, and tritium.

The results of this work indicate that the soil beneath the 317 Area is composed primarily of low permeability clay, interspersed with isolated pockets of porous sand and gravel. Organic contamination of the groundwater present in these layers was present; however, it appears to be limited to the immediate vicinity of the 317 Area French drain. The migration of this contamination beyond the French drain area is being limited by the presence of clay soils.

Very little contamination was found beneath the 319 Area landfill. The existence of what is thought to be previously unknown waste buried east of the 319 Area landfill was confirmed. The nature of this waste materials is unknown but it is likely to be innocuous demolition debris and similar solid wastes since analysis of groundwater samples in the area did not detect significant contamination.

This characterization information was used to develop a detailed work plan for the next phase of characterization of this area. This phase, known as the RCRA Facility Investigation, is a formal, IEPA-approved investigation which will form the basis for eventual remedial action of contamination identified. A draft work plan was written and submitted to IEPA for review. The work plan was approved in August of 1994. In December, RFI field work was begun. A series of soil borings were drilled into the soil south of the 319 landfill to obtain subsurface soil samples. The first monitoring well

cluster, also south of the 319 landfill, was installed. Analytical data from these samples indicates the subsurface soil is free of hazardous chemical contaminants. The groundwater was found to contain low concentrations of several volatile organic compounds, include vinyl chloride (63  $\mu$ g/L), 1,2 dichloroethene (470  $\mu$ g/L) and trichloroethane (911  $\mu$ g/L). Additional characterization field work is scheduled for 1995 and 1996. The final RFI report for this phase of the investigation will be completed in late 1996.

A project to identify and clean up residual radiological contamination within certain facilities within the 317 Area was also begun. A radiological survey of all active and inactive radiological waste management facilities was performed. Based on this data, a plan was developed to begin the decontamination of those facilities which are no longer needed for programmatic purposes. The decontamination operation began early in 1994.

During 1994, a project to characterize and clean up residual radiological contamination within a facility in the 317 Area, known as the Map Tube Facility, was completed. This facility consisted of a monolithic concrete structure with 129 cast iron pipes set into the concrete matrix. The pipes were used to store small quantities of highly radioactive material. Over the years of operation of this unit, some of the containers leaked or broke open, releasing contamination into the inside of the pipes. Because of the presence of two lead-sealed pipe joints in each pipe, the entire pipe, joints and all, were cut from the concrete matrix, disassembled to remove the lead, and shipped to the Hanford site for disposal. The holes in the concrete matrix were then filled with concrete. By late October, all 129 tubes had been removed and shipped off-site. This action prevented further releases of radioactive materials from the unit and reduced the potential for radiation exposure to the workers in the 317 Area.

Two inactive hazardous waste treatment units located in the 317 Area underwent closure activities during 1994 (Section 2.3.1.). Closure activities included sampling and

analysis of residual materials present in the units (sludge and solid waste), and disposal of these waste materials and analysis of the soil underneath and next to these units to verify that all hazardous materials have been removed. Closure was complete in late 1994.

## 6.5.2. 800 Area Landfill Characterization

Characterization activities at the landfill began in 1989 with the collection of a series of soil gas and leachate samples from in and near the fill material. The results of soil gas and leachate analysis showed that volatile organic compounds are present in the fill material and leachate. A large number of the compounds detected are also listed on the log of wastes poured into the old French drain in the north end of the site. It appeared that volatile organics are present throughout most of the fill material. The distribution of these chemicals throughout the fill was found to be highly variable, indicating the possibility of multiple sources within the waste.

In 1992, 14 soil borings were completed along with the installation of 15 monitoring wells around the outside of the landfill. The 15 monitoring wells installed in 1992 enable the measurement of vertical hydraulic gradients and groundwater quality variations at the solid waste management unit boundary and following an action by ANL to modify the solid waste disposal permit, they now comprise the monitoring well network for the now-closed 800 Area Landfill. Since 1992, results from the groundwater monitoring program have revealed little in the way of groundwater contamination migrating from the landfill itself. However, given the type and amounts of hazardous substances disposed of in the landfill, the potential still exists for future groundwater contamination. Capping of the 800 Area Landfill was completed as an interim measure prior to conducting a RCRA Facility Investigation (RFI) and any corrective activities for that area. Future work to be conducted under the RCRA permit will establish the effect the landfill has had to its surrounding environment.

# 6. GROUNDWATER PROTECTION

In 1994, ANL wrote and submitted the RFI work plan for the 800 Area to the IEPA. In November 1994, IEPA approved the RFI work plan. Work on the RFI began in December 1994 with the collection of surface water and sediment samples and shallow soil samples.

Work scheduled for 1995 includes: the installation of nine new groundwater monitoring wells; collection of samples from 12 SWMUs; installation of leachate wells in the landfill; and collection of two rounds of groundwater samples.

# 6.5.3. Building 810 Paint and Solvent Disposal Area Interim Action Project

In 1994, ANL proposed an interim action for a SWMU located in the 800 Area. The SWMU, known as the Building 810 Paint and Solvent Disposal Area, was adjacent to an access door for Building 810. This building served as the site's paint shop for several years, until 1987. During its active period, paint and paint solvent were routinely poured onto the ground surface for disposal.

In 1994, ANL wrote an interim action work plan and submitted the plan to DOE. DOE submitted the plan to IEPA who subsequently approved the work plan. ANL in following the work plan prepared the area and removed approximately 225 m³ of paint and paint solvent contaminated soil from the SWMU. Following removal of the contaminated soil, ANL restored the area to its original condition and produced a final report of the cleanup for submittal to IEPA.

# 6.5.4. Sitewide Hydrogeological Characterization Project

This multiphase project will fully characterize the hydrogeology of the ANL site and support a sitewide RFI. The project is critical to defining the baseline hydrogeological

conditions beneath ANL and the surrounding Forest Preserve including the geological and hydrogeological characteristics of the uppermost aquifer system. This information will define the sitewide ground water quality and flow regime and provide important baseline data for other, smaller-scale, site characterization and remediation projects. It also will be an important resource for new construction project designs and environmental spill responses. The data will reveal whether ANL operations overall have impacted groundwater.

In 1994, the Phase I field investigation was completed. This included geophysical measurements, cone penetration testing, soil borings, rock corings, and the installation of five bedrock monitoring wells in the western portion of the ANL site. Groundwater samples were collected and analyzed to determine baseline groundwater quality. Constituents analyzed for included Illinois Class I groundwater quality parameters, as well as selected radionuclides and stable isotopes.

Also during 1994, 18 old unsealed wells of record were located and evaluated for their potential use in collecting hydrogeological information. Wells requiring proper closure were identified. The unusable wells will be sealed in accordance with State and County health regulations. The usable wells will be incorporated into the site-wide monitoring well network.

The current plan calls for the Phase II field investigation to be conducted in 1996. Phase II will characterize the detailed hydrogeology of the remainder of the ANL site and complete the major field portion of the project.

# 7. QUALITY ASSURANCE



# 7. QUALITY ASSURANCE

Quality Assurance (QA) plans exist for both radiological (ESH-DARC-QAP-001) and non-radiological (ESH-DACH-QAP-001) analyses. Both QA documents were prepared in accordance with DOE Order 5700.6C. The plans discuss responsibilities and auditability. Both documents are supplemented by operating manuals.

# 7.I. Radiochemical Analysis and Radioactivity Measurements

All nuclear instrumentation is calibrated with standard sources obtained from or traceable to the National Institute of Standards and Technology (NIST). The equipment is usually checked daily with secondary counting standards to ensure proper operation. Samples are periodically analyzed in duplicate or with the addition of known amounts of a radionuclide to check precision and accuracy. When a nuclide was not detected, the result is given as "less than" (<) the detection limit by the analytical method used. The detection limits were chosen so that the measurement uncertainty at the 95% confidence level is equal to the measured value. The air and water detection limits for all radionuclides for which measurements were made are given in Table 7.1. The relative error in a result decreases with increasing concentration. At a concentration equal to twice the detection limit, the error is about 50% of the measured value and at ten times the detection limit, the error is about 10%.

Average values are usually accompanied by a plus-or-minus ( $\pm$ ) limit value. Unless otherwise stated, this value is the standard error at the 95% confidence level calculated from the standard deviation of the average. The  $\pm$  limit value is a measure of the range in the concentrations encountered at that location; it does not represent the conventional uncertainty in the average of repeated measurements on the same or identical samples. Since many of the variations observed in environmental radioactivity are not random but occur for specific reasons (e.g., seasonal variations), samples collected from the same location at different

**TABLE 7.1**Detection Limits

Nuclide or Activity	Air (fCi/m³)	Water (pCi/L)
Americium-241	•	0.001
Beryllium-7	5	-
Californium-249	~	0.001
Californium-252	~	0.001
Cesium-137	0.1	1
Curium-242	-	0.001
Curium-244	<del>-</del> '	0.001
Hydrogen-3	100	100
Lead-210	1	-
Neptunium-237	-	0.001
Plutonium-238	0.0003	0.001
Plutonium-239	0.0003	0.001
Radium-226	-	0.1
Radium-228	-	0.1
Strontium-89	0.1	2
Strontium-90	0.01	0.25
Thorium-228	0.001	-
Thorium-230	0.001	-
Thorium-232	0.001	-
Uranium-234	0.0003	0.01
Uranium-235	0.0003	0.01
Uranium-238	0.0003	0.01
Uranium - natural	0.02	0.2
Alpha	0.2	0.2
Beta	0.5	1

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times are not replicates. The more random the variation in activity at a particular location, the closer the confidence limits will represent the actual distribution of values at that location. The averages and confidence limits should be interpreted with this in mind. When a plus-or-minus value accompanies an individual result in this report, it represents the statistical counting error at the 95% confidence level.

Standard and intercomparison samples distributed by the Quality Assurance Branch of the EPA are analyzed regularly. Results of ANL's participation in the EPA program during 1994 are given in Table 7.2. In the table, the comparison is made between the EPA value, which is the quantity added to the sample by that laboratory, and the value obtained in the ANL laboratory. Certain information may assist in judging the quality of the results, including the fact that typical uncertainties for the ANL analyses are 2% to 50%, depending on the concentration and the nuclide, and the uncertainties in the EPA results are 2% to 5% (ANL estimate).

In addition, participation continued in the DOE Environmental Measurements Laboratory Quality Assurance Program (DOE-EML-QAP), a semi-annual distribution of four different sample matrices containing various combinations of radionuclides that are analyzed. Results for 1994 are summarized in Table 7.3. In the table, the EML value, which is the result of duplicate determinations by that laboratory, is compared with the average value obtained in the ANL laboratory. Information that will assist in judging the quality of the results includes the fact that typical uncertainties for ANL's analyses are 2% to 50% and that the uncertainties in the EML results are 1% to 30% (depending on the nuclide and the amount present). For most analyses for which the differences are large (> 20%), the concentrations were quite low and the differences were within the measurement uncertainties.

TABLE 7.2

Summary of EPA Samples, 1994

Type of Sample	Analysis	Number Analyzed	Average Difference from Added (%)
Air Filter	Total Alpha	1	21
	Total Beta	1	33
	Strontium-90	1	2
	Cesium-137	1	2
Water	Total Alpha	4	34
	Total Beta	4	23
	Hydrogen-3	2	3
	Cobalt-60	3	3
	Zinc-65	2	11
	Strontium-89	3	4
	Strontium-90	3	0
	Ruthenium-106	1	9
	Iodine-131	1	3
	Cesium-134	3	7
	Cesium-137	3	10
	Barium-133	2	4
	Radium-226	4	2
	Radium-228	4	2
	Total Uranium	4	6
	Plutonium-239	1	5

TABLE 7.3

Summary of DOE-EML-QAP Samples, 1994

	Percer	nt Difference Fi	rom EML Value	
Radionuclide	Air Filters	Soil	Vegetation	Water
Hydrogen-3	<u>-</u>	· <u>-</u>	-	3 (2)
Potassium-40	-	10 (2)	5 (2)	-
Manganese-54	9 (2)	-	-	6 (2)
Cobalt-57	15 (2)	-	-	-
Cobalt-60	5 (2)	-	8 (2)	13 (2)
Strontium-90	13 (2)	24 (2)	9 (2)	8 (2)
Ruthenium-106	42 (1)	•	-	-
Antimony-125	6 (2)	-	-	-
Cesium-134	14 (1)	-	-	4 (2)
Cesium-137	5 (2)	9 (2)	10 (2)	12 (2)
Cerium-144	20 (2)	-	-	15 (1)
Uranium-234	6 (2)	13 (2)	-	7 (2)
Uranium-238	9 (2)	13 (2)	-	9 (2)
Plutonium-238	9 (2)	18 (2)	41 (1)	9 (2)
Plutonium-239	10 (2)	6 (2)	13 (2)	4 (2)
Americium-241	5 (2)	14 (2)	11 (2)	3 (2)

Note: The value in parentheses is the number of samples.

# 7.2. Chemical Analysis

The documentation for nonradiological analyses is contained in the ESH-DA Chemistry Laboratory Procedure Manual. All samples for NPDES and groundwater are collected and analyzed in accordance with EPA regulations found in 40 CFR Part 136, <sup>19</sup> EPA-600/4-84-017, <sup>26</sup> and SW-846. <sup>27</sup>

Standard Reference Materials (SRM), traceable to the NIST, exist for most inorganic analyses (see Table 7.4). These are replaced annually. Detection limits are determined with techniques listed in Report SW-846<sup>27</sup> and are listed in Table 7.5. In general, the detection limit is the measure of the variability (6) of a standard material measurement at 5-10 times the instrument detection limit as measured over an extended time period. Recovery of inorganic metals, as determined by "spiking" unknown solutions, must be in the range of 75% to 125%. The precision, as determined by analysis of duplicate samples, must be within 20%. These measurements must be made on at least 10% of the samples. Comparison samples for organic constituents were formerly available from the EPA, but are now commercially available under the Cooperative Research and Development Agreement (CRADA) which exists between the EPA and commercial laboratories. In addition, standards are available which are certified by the American Association for Laboratory Accreditation, under a memorandum of understanding with the EPA. Many of these standards are used in this work. At least one standard mixture is analyzed each month and the results for 1994 are shown in Table 7.6 for volatile organic compounds and Table 7.7 for semivolatiles. The recoveries listed are those required by the respective methods.

TABLE 7.4

Reference Materials Used for Inorganic Analysis

	<u> </u>
Reference Material	Constituent
HPS-10003-1	Arsenic
HPS-10004-1	Barium
HPS-10005-1	Beryllium
HPS-10008-1	Cadmium
HPS-100012-1	Chromium
HPS-100013-1	Cobalt
HPS-100014-1	Copper
HPS-100026-1	Iron
HPS-100028-1	Lead
HPS-100032-1	Manganese
HPS-100033-1	Mercury
HPS-100036-1	Nickel
HPS-100049-1	Selenium
HPS-100051-1	Silver
HPS-100065-1	Vanadium
HPS-100068-1	Zinc
NIST-SRM 3181	Sulfate
NIST-SRM 3182	Chloride
NIST-SRM 3183	Fluoride

HPS = High Purity Standards, Inc.

NIST-SRM = National Institute of Standards and Technology - Standard Reference Materials.

TABLE 7.5

Limit of Detection for Metal Analysis

	Limits of Detection Milligrams/Liter		
Constituent	AA*	ICP**	
Arsenic	0.0020	0.110	
Barium	- 1	0.015	
Beryllium	0.0001	0.025	
Cadmium	0.0001	0.040	
Chromium	0.020	0.060	
Cobalt	0.020	0.025	
Copper	0.010	0.045	
Hexavalent Chromium	0.006	-	
Iron	0.040	0.025	
Lead	0.0010	0.110	
Manganese	0.010	0.020	
Mercury	0.0001	-	
Nickel	0.020	0.040	
Silver	0.0010	0.110	
Thallium	0.0030	0.145	
Vanadium	-	0.055	
Zinc	0.010	0.015	

<sup>\*</sup>AA = Atomic Absorption Spectroscopy.

<sup>\*\*</sup>ICP = Inductively Coupled Plasma - Atomic Emission Spectroscopy.

Quality Check Sample Results, Volatile Analyses, 1994

**TABLE 7.6** 

Compound	Percent Recovery	Percent Quality Limits
Benzene	101	73-126
Bromobenzene	101	76-133
Bromodichloromethane	98	101-138
Bromoform	76	57-156
Butylbenzene	108	71-125
sec-Butylbenzene	108	71-145
t-Butylbenzene	112	69-134
Carbon Tetrachloride	101	86-118
Chlorobenzene	108	80-137
Chloroform	103	68-120
o-Chlorotoluene	111	81-146
p-Chlorotoluene	107	73-144
1,2-Dibromo-3-chloropropane	117	36-154
Dibromochloromethane	89	68-130
1,2-Dibromomethane	113	75-149
Dibromomethane	99	65-143
1,2-Dichlorobenzene	121	59-174
1,3-Dichlorobenzene	116	84-143
1,4-Dichlorobenzene	118	58-172
1,1-Dichloroethane	92	71-142
1,2-Dichloroethane	119	70-134
1,1-Dichloroethene	127	18-209
cis-1,2-Dichloroethene	99	85-124
trans-1,2-Dichloroethene	114	67-141
1,2-Dichloropropane	125	19-179
1,3-Dichloropropane	93	73-145
1,1-Dichloropropene	92	71-133
Ethyl Benzene	100	84-130
Isopropylbenzene	101	70-144
4-Isopropyltoluene	105	72-140
Methylene Chloride	89	D-197
n-Propylbenzene	108	78-139
1,1,1,2-Tetrachloroethane	94	88-133
Tetrachloroethene	108	84-132
Toluene	99	81-130
1,1,1-Trichloroethane	90	68-149
1,1,2-Trichloroethane	119	70-133
Trichloroethene	100	91-135
1,2,3-Trichloropropane	120	50-158
1,2,4-Trimethylbenzene	105	80-144
1,3,5-Trimethylbenzene	109	76-142
o-Xylene	100	79-141
p-Xylene	108	74-138

Note: D denotes the compound was detected.

# 7. QUALITY ASSURANCE

Quality Check Sample Results, Semivolatile Analyses, 1994

**TABLE 7.7** 

Compound	Percent Recovery <sup>a</sup>	Percent Quality Limits
2-Fluorophenol <sup>b</sup>	62.1	21-100
Phenol-d5 <sup>b</sup>	42.6	10-94
Phenol	43.1	17-100
2-Chlorophenol	78.7	36-120
1,4-Dichlorobenzene	82.1	37-106
n-Nitroso-n-Propyl Amine	61.6	24-198
Nitrobenzene-d5 <sup>b</sup>	85.9	35-114
1,2,4-Trichlorobenzene	92.0	57-129
4-Chloro-3-Methylphenol	89.3	41-128
2-Fluorobiphenyl <sup>b</sup>	68.0	43-116
Acenaphthene	126.0	47-145
4-Nitrophenol	65,2	13-107
2,4-Dinitrotoluene	126.0	48-127
2,4,6-Tribromophenol <sup>b</sup>	88.6	10-123
Pentachlorophenol	107.0	38-152
Pyrene	90.2	70-100
Terphenyl-d14 <sup>b</sup>	99.9	33-141

<sup>&</sup>lt;sup>a</sup>Average of three determinations.

<sup>&</sup>lt;sup>b</sup>Required surrogates.

# 8. APPENDIX



# 8. APPENDIX

## 8.1. References

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