

APPENDIX E SAMPLE COLLECTION, TRACKING, AND QA/QC RESULTS

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1.0 GROUNDWATER SAMPLING

This section describes the tracking system, quality assurance, and quality control (QA/QC) for samples collected for the Environmental Restoration (ER) and Environmental Surveillance (ES) Groundwater Monitoring Programs, and contains the procedures used to collect groundwater samples from monitoring wells. QA/QC issues and the procedures for collecting groundwater samples were formalized during 1999 into BNL Standard Operating Procedures (SOPs) and revised in 2002. These SOPs will be discussed in the appropriate sections below.

1.1 Sample Collection

Groundwater samples were collected during CY 2003 by the BNL Field Sampling Team for the ES samples and by Dvirka and Bartilucci, Consulting Engineers for the ER samples. A low flow purge technique was used to collect all the groundwater samples. The USEPA approved low flow purge technique allows a groundwater sample to be collected using a very low purge rate, approximately 100 to 500 milliliters per minute. This low flow rate does not draw any standing water from the well casing, allowing all water to be drawn directly from the aquifer being sampled. Since the groundwater is drawn directly from the aquifer, very little is purged out prior to sample collection. A typical low flow sampling produces 1.5 to 5 gallons of purge water waste per sampling event as compared to 25 to 200 gallons of purge water waste using the three purge volume technique.

BNL EM-SOP-302, Low Purge Sampling of Monitoring Wells Using Dedicated Pumps, was followed by field personnel collecting groundwater samples from wells with dedicated pumps installed. Most of the wells in the monitoring program were equipped with dedicated pumps designed to collect water samples using the low flow technique. When a well was designated to be sampled but a dedicated pump was not installed, the procedures outlined in BNL EM-SOP-307, Low Purge Sampling of Monitoring Wells using Non-dedicated Pumps, was used.

Purge water from groundwater samples is disposed of in one of three methods as described in BNL EM-SOP-802, Well Development, Purge and Decontamination Water Handling Procedure. If the groundwater does not contain any analytes above action levels, NYSDEC and Federal drinking water standards and DOE groundwater screening levels, the purge water is discharged approximately 20 feet down gradient from the wellhead. If the groundwater contains chemical analytes above action levels but no radiological parameters above action levels, the purge water is

run through activated carbon and discharge at least 20 feet downgradient of the wellhead. Purge water which contains radiological parameters above action levels is containerized and disposed of off-site in accordance with all local, state, and federal regulations.

1.1.1 Decontamination

Most groundwater sampling equipment is dedicated to each well, and was decontaminated by the manufacturer. The only non-dedicated equipment requiring field decontamination was the submersible pump used for sampling some OU III (AOC 29 HFBR) wells. Decontamination was performed as per BNL EM-SOP-801.

1.2 Sample Tracking System

Samples are tracked by the Environmental Information Management System (EIMS). Tracking is started when a sample is recorded on a chain-of-custody form. Sampling personnel submit these forms to the sampling coordinator, and the information is entered into the EIMS.

1.2.1 Sample Identification

Samples were identified using a code consisting of the chain-of-custody (COC) number and the unique ID number. This ID is written on the sampling logs along with the BNL well ID. BNL well IDs also were placed on the COC forms in the Site ID column. QA/QC samples are identified in the same manner as environmental samples. The blind duplicate samples were recorded as BD in the sample ID column of the COC. Supplemental forms are used by field sampling personnel to distinguish information about the QC samples, such as blind duplicate IDs and associated field and trip blanks. BNL EM-SOP-102 details BNL's Chain of Custody Procedure. COC records are filed at BNL and are available for review.

1.2.2 Sample Tracking

Copies of the COC forms are provided weekly to the sampling coordinator to enter into the EIMS. The status of each sample is updated when

1. The sample is assigned to a Sample Delivery Group (SDG) and the analytical laboratory communicates this information to the Sampling Coordinator;
2. The Project Manager receives and approves the hard copy with the results of the sample analysis;

3. The Electronic Data Deliverable (EDD) analytical results are received and entered into the EIMS;
4. The Project Manager receives and approves of the hard copy of the data validation package (when applicable);
5. The results of the EDD data-validation results are received and entered (when applicable) into the EIMS.

1.2.3 Sample Packaging and Shipping

Samples that are shipped to external contract laboratories are packaged by placing each sample bottle inside a plastic bag and sealing it. The bottles with VOC samples are placed in protective cans with foam inserts. Glass bottles are wrapped with protective packaging to protect against breakage during shipment. Plastic bags are filled with ice and sealed, or blue-ice containers are placed inside each cooler with the samples to ensure that they arrive at the analytical laboratory at four degrees Celsius (plus or minus two degrees). A BNL chain-of-custody form completed by the sampling team accompanies the samples to the laboratory. The form is placed in a plastic bag, sealed, and put inside the cooler with the samples. The analytical samples are shipped to the analytical laboratory via an overnight mail carrier. Samples that are transported to the BNL ASL for analyses are treated in much the same way, except that the packaging requirements to protect against breakage during shipment are less critical.

1.2.4 Sample Documentation

The sample teams maintain field notebooks, bound weatherproof logbooks, that are filled out at the location where the sample is collected. It contains the sample's designation, collection time, description, collection method, and the weather conditions, field measurements, and other site-specific observations.

The sample teams also complete collection logs for every sample that is collected. The completed sample-collection logs are submitted to the sample coordinator each week.

1.3 Analytical Methods

The following sections describe the analytical methods used for the BNL Groundwater Monitoring Program.

1.3.1 Chemical Analytical Methods

The EM and ES samples collected during CY 2000 were analyzed by organic, inorganic, and various wet-chemical methods. EM Chemical analyses were performed by GEL Laboratories, Severn Trent Laboratories and the BNL ASL. ES chemical analyses were performed by the ASL and H2M Labs. Table 1-6 summarizes the analytes and/or methods used for specific EM monitoring programs and samples. Table 1-7 summarizes this information for the ES program.

The following inorganic, organic, and wet chemical methods were used: United States Environmental Protection Agency (USEPA) methods including 200 and 500 Series methods (40 Code of Federal Regulations (CFR) 141); 600 Series methods (40 CFR 136); and SW-846 methods (40 CFR 261. Other standard methods include those listed in *Standard Methods for the Analysis of Wastewater* (latest edition) and those in the American Society for Testing and Materials (ASTM) publications (latest revision).

1.3.2 Radiological Analytical Methods

Unlike organic and inorganic chemical analytical methods, few standard methods are available for the radiological analysis of environmental samples. There are no standard established QA/QC requirements and acceptance criteria for environmental radiological methods; therefore, different USEPA, U.S. Department of Energy (DOE), and commercial laboratories may have different methods of preparing samples preparing and analytical techniques for specific radiological analytes. Hence, laboratory-reported detection limits may vary. Nonetheless, multi laboratory validation studies and interlaboratory comparisons have demonstrated that accurate, comparable, radiological data are obtainable even when different procedures are used.

Tables 1-6 and 1-7 provide the analytical parameters and/or methods used for specific monitoring programs and samples. Radiological analyses were undertaken for onsite, offsite, and site-boundary locations. The radiological analyses enable BNL to monitor radiological water-quality status throughout the site.

1.4 Quality Assurance and Quality Control

This section describes the QA/QC requirements for field work conducted under the ER and ES Groundwater Monitoring Programs. The *BNL Groundwater Monitoring Program Quality Assurance Project Plan, August 1999*, provides detailed information regarding QA/QC requirements. In general, the quality of the analytical results from groundwater samples collected during CY2001 met data quality objectives.

1.4.1 Calibration and Preventive Maintenance of Field Instruments

Sampling team personnel are responsible for assuring that a master calibration/maintenance log is maintained for each field-measuring device (i.e., pH, conductivity, turbidity meters, etc.). The sample coordinator provides a calibration/maintenance logbook for equipment supplied to contracted sampling teams.

1.4.2 QA/QC Sample Collection

Guidance on collecting the QA/QC samples is given in the *BNL Groundwater Monitoring Program Quality Assurance Project Plan, August 1999* and in *BNL EM-SOP-200 "Collection and Frequency of Field Quality Control Samples."* Sample-specific requirements are listed separately, below.

The collection of QA/QC samples is dependent on the data quality objectives of each project. The following is a general breakdown of the QA/QC samples collected by project type:

ER Groundwater Monitoring: trip blanks, field blanks, equipment blanks, matrix spike/matrix spike duplicates (MS/MSDs), and blind duplicates.

Remediation Treatment System Sampling: trip blanks.

ES Groundwater Monitoring: trip blanks, field blanks, equipment blanks, and field duplicates.

1.4.2.1 Equipment Blanks

Equipment blanks are collected to evaluate any potential cross-contamination of samples due to the sampling equipment. They are collected by pouring laboratory grade water over the sampling

equipment which comes in contact to the groundwater sample. Equipment blanks are only collected for projects that require the use on non-dedicated sampling equipment. The frequency of collecting equipment blanks is one for every 20 groundwater samples shipped to the analytical laboratory. The sampling equipment for all projects except for the HFBR is dedicated, therefore no equipment blanks are needed. In the HFBR project, all but 20 monitoring wells have dedicated equipment, therefore only one equipment blank is needed per sampling round. Equipment blank detections are summarized on Table E-1. Tetrachloroethene and toluene were each detected once in equipment blanks collected during 2003. The only constituent detected was toluene. Toluene is a typical contaminant introduced during field collection activities. In addition to the three volatile organic equipment blanks listed on Table E-1, four equipment blanks were also collected and analyzed for tritium. No tritium was detected.

1.4.2.2 Field Blanks

Field blanks are obtained by pouring laboratory grade water into clean sample bottles containing preservatives. The field blanks are collected in the field and accompany field personnel to the sampling location. They are analyzed for the same parameters that the groundwater is being analyzed for on that day. The frequency of collecting equipment blanks is one for every 20 groundwater samples shipped to the analytical laboratory. For projects with less than 20 monitoring wells, a minimum of one field blank is collected per project for each sampling event. Field blank results are summarized on Table E-2. The most common constituent detected was methylene chloride which was detected in 12 of the 66 field blanks collected for VOCs. Methylene chloride is defined by the USEPA data validation guidelines as a common laboratory contaminant. Strontium-90, gross beta, and tritium were each detected once in the 72 field blanks collected for radiological analyses. The gross beta detection was in the same field blank as the strontium-90 detection, which confirmed both detections as actual contamination rather than analytical error.

1.4.2.3 Trip Blanks

A trip blank is an aliquot of deionized water that is sealed in a sample bottle (glass vials (40 ml) with Teflon septa). It is used to determine if there is any cross-contamination between aqueous samples during shipment. Trip blanks are analyzed for aqueous VOCs only. A trip blank is shipped to the analytical laboratory with each set of samples submitted for VOC analyses. Upon

arrival at BNL, the sealed trip blank bottles are placed in a cooler and brought to the field by the sampling team. If several coolers are required, each cooler must contain an individual trip blank. Trip blank detections are summarized on Table E-3. 169 trip blanks were collected in 2003. The constituents detected the most were methylene chloride and toluene which were detected in 30 and 10 trip blanks, respectively. Methylene chloride is defined by the USEPA data validation guidelines as a common laboratory contaminant. Toluene is a typical contaminant introduced during field collection activities.

1.4.2.4 Duplicate Samples

Field duplicate samples are analyzed to check the reproducibility of the laboratory's analytical results. Duplicates are either blind (the laboratory doesn't know the identity of the sample location) or field (the laboratory is told the identity of the sampling location). The specific type of duplicate used on a project is dependent on the project data quality objectives. At least 5 percent (one out of every 20 samples) of the total number of collected groundwater samples are duplicated to evaluate the precision of the methods. For projects with less than 20 monitoring wells, a minimum of one blind duplicate sample is collected per project for each sampling event. USEPA Region II USEPA Region II data validation criteria were used for field duplicate interpretation. For detects above 5 times the contract required detection limit (CRDL), a relative percent difference (RPD) was calculated. An acceptable RPD was 50% or below. For detects below 5 times the CRDL, the QC requirements is that the difference between the duplicate results must be less than or equal to the CRDL. A total of 83 duplicate samples were collected for non-radiological analyses and 85 duplicates were collected for radiologic analyses. Not all parameters were analyzed in every duplicate. The parameters in each duplicate were consistent with those required for the specific program the duplicate was monitoring. Of the 5310 parameters analyzed, only 66 (1.2%) of the non-radiologic analyses failed to meet QA criteria. For the radiologic parameters only 7 of the 353 parameters (2%) failed to meet QA criteria. The results are indicative of consistency with the laboratory and sampling team that is resulting in valid, reproducible data.

1.4.2.5 Requirements for Matrix Spike/Matrix Spike Duplicate Volumes

MS/MSDs for organic analysis are performed at a frequency of one MS/MSD for every 20 groundwater samples in an SDG. Reanalysis may be necessary in certain situations. To ensure

that the laboratory has sufficient volume for MS/MSD analysis, triple the sample volume must be collected.

1.4.3 Data Verification

There are two stages of data verification. One stage consists of reviewing the latest data in comparison to historical data generated at the sampling site. All groundwater data collected at BNL undergoes this type of verification. The other stage is a formal, documented, data verification. The procedures for the formal data verification are given in BNL EM-SOP-203, Chemical Data Verification and BNL EM-SOP-204, Radiochemical Data Verification. This is BNL's internal process to verify the accuracy and/or completeness of analytical data.

The decision to perform a formal verification is based on the data quality objectives of the specific projects. Data generated under of the EM Groundwater Monitoring Program that were not validated underwent data verification. ES data and treatment system data do not undergo formal verification, but are compared to known baseline data. Therefore, the ES and treatment system analytical data need only to undergo the historical review. If the comparison of historical data to new data indicates an inconsistency with the expected results, a further review is conducted which may include, formal data verification, data validation, and/or a data usability review.

The formal data verification process is designed to detect the most common analytical problems that affect the quality of the results. To accomplish this task, QA/QC items such as the following are checked: holding times; matrix spikes; laboratory and field blanks; and, field logs. If items are detected that can affect the use of the data, they are either corrected, as in the case of unintelligible information on the field logs, or the data is qualified, as in the case of blank contamination or holding time violations.

1.4.4 Data Usability

Data usability is the process by which data that does not meet the expected results, but which has been deemed acceptable by a data validation or verification, is reviewed.

Determining the usability of chemical data is relatively straightforward. Laboratory analytical data are validated or verified, and validation qualifiers are assigned to them. Table E-4 defines the qualifiers placed on the data by the analytical laboratory and Table E-5 defines the data validation, verification, and usability qualifiers.

The usability of radiological data for the 2001 data was determined through a two-step process. The project manager initially reviewed all groundwater monitoring data. Data were considered acceptable for use if they were not significantly different than expected for a particular well, based on historical trends and were not qualified as unusable during the validation and/or verification procedures. Results for a particular well that were not expected, based on historical trends, were referred to the ER radiochemist. The data then were assessed according to BNL's Procedures for Radiochemical Data Validation (BNL EM-SOP-209) and Radiochemical Data Usability (BNL EM-SOP-210). The data then were assigned (if applicable) a revised qualifier and a data-usability code. A usability code of "N3" was assigned if the data were not usable based on the lack of expected daughter products. A usability code of "N2" was assigned if the data were not usable because the results and the propagated error are indistinguishable from background (i.e., the result minus the 2 sigma error is less than the detection limit). Data identified as being "not usable" were not considered in characterizing the presence or extent of contamination. Data usability report summaries are included as Appendix G.

1.4.5 Data Qualification

During the data validation, verification and/or usability processes, the data may be qualified to alert the user to limitations in the use of the data based on QA/QC violations. Table E-2 defines the qualifiers placed on the data by the analytical laboratory and Table E-3 defines the data validation, verification, and usability qualifiers. For organic and inorganic analytes, three primary qualifiers may be applied to laboratory data: "U," "J," and "R." In addition, there may be no qualifier if QA/QC issues are not identified. For radiological data, in addition to the "U," "J," and "R" qualifiers, qualifiers such as "DL," "N2," and "N3" were also applied to the results.

A "U" qualifier, which is a laboratory qualifier, indicates that the analyte was a target of the method but was not detected. The "U" qualifier also may be used in conjunction with the "J" qualifier, which indicates that the reported concentration is an estimated value because the reported value is lower than the required reporting limit, or because one or more analytical

deficiencies were noted during the data validation review. Thus the designation “UJ” indicates that the analyte was not detected and the reported quantitation or detection limit is an estimate due to QA/QC deficiencies. The “R” qualifier indicates that the datum is rejected. An “R” qualifier can be reported for analytes that either were, or were not, detected. In other words, an “R” qualifier may be assessed upon a reported concentration or a result reported with a “U” qualifier.

Data reported as either unqualified, or with “U” or “J” qualifiers are typically usable, in assessments of the extent of contamination or effectiveness of remedial actions. Data qualified by “R” are considered unusable.

Table E-1 Concentrations of Constituents Detected in Equipment Blank Samples Associated with the EM and ES Grounwater Sampling Programs

| Constituent | Number of Analyses | Number of Detects | Minimum | Maximum | Average Detection Limit | Units |
|---------------------|--------------------|-------------------|---------|---------|-------------------------|-------|
| Tetrachloroethylene | 3 | 1 | 1 | 1 | 1.5 | ug/L |
| Toluene | 3 | 1 | 0.26 | 0.26 | 1.5 | ug/L |

ug/L Micrograms per liter.

Table E-2 Concentrations of Constituents Detected in Field Blank Samples Associated with the EM and ES Grounwater Sampling Programs

| Constituent | Number of Analyses | Number of Detects | Minimum | Maximum | Average Detection Limit | Units |
|-------------------------------------|--------------------|-------------------|---------|---------|-------------------------|-------|
| Organic Compounds | | | | | | |
| Acetone | 2 | 2 | 2.1 | 20.4 | 3.5 | ug/L |
| Benzene | 66 | 1 | 0.9 | 0.9 | 0.66 | ug/L |
| Decachlorobiphenyl | 1 | 1 | 0.023 | 0.023 | 0.05 | ug/L |
| Methyl bromide | 66 | 1 | 0.62 | 0.62 | 0.66 | ug/L |
| Methyl chloride | 66 | 1 | 0.28 | 0.28 | 0.66 | ug/L |
| Methylene chloride | 66 | 12 | 0.3 | 2 | 0.66 | ug/L |
| Tetrachloro-m-xylene | 1 | 1 | 0.017 | 0.017 | 0.05 | ug/L |
| Toluene | 66 | 2 | 0.26 | 0.27 | 0.66 | ug/L |
| Trichloroethylene | 66 | 1 | 1.6 | 1.6 | 0.66 | ug/L |
| Metals | | | | | | |
| Aluminum | 11 | 2 | 2.2 | 35.1 | 35.7 | ug/L |
| Antimony | 11 | 1 | 0.9 | 0.9 | 4.69 | ug/L |
| Arsenic | 11 | 2 | 1.6 | 3 | 3.61 | ug/L |
| Barium | 11 | 2 | 0.2 | 1.8 | 5.87 | ug/L |
| Beryllium | 11 | 3 | 0.24 | 0.7 | 0.728 | ug/L |
| Cadmium | 11 | 1 | 1.1 | 1.1 | 1.04 | ug/L |
| Calcium | 10 | 2 | 10.2 | 12.5 | 81.9 | ug/L |
| Chromium hexavalent ion | 4 | 1 | 0.01 | 0.01 | 0.0054 | mg/L |
| Cobalt | 11 | 1 | 0.01 | 0.01 | 2.23 | ug/L |
| Copper | 11 | 1 | 2.5 | 2.5 | 4.63 | ug/L |
| Iron | 11 | 3 | 0.1 | 11.9 | 20.8 | ug/L |
| Lead | 11 | 1 | 1.3 | 1.3 | 2.39 | ug/L |
| Magnesium | 10 | 1 | 12.3 | 12.3 | 85.0 | ug/L |
| Manganese | 11 | 3 | 0.351 | 4.2 | 1.9 | ug/L |
| Mercury | 11 | 1 | 0.061 | 0.061 | 0.093 | ug/L |
| Nickel | 11 | 2 | 1.03 | 1.1 | 4.28 | ug/L |
| Potassium | 10 | 3 | 23.7 | 1830 | 621 | ug/L |
| Selenium | 11 | 1 | 5 | 5 | 4.4 | ug/L |
| Sodium | 11 | 4 | 34 | 138 | 86 | ug/L |
| Thallium | 11 | 4 | 0.7 | 0.7 | 4.58 | ug/L |
| Vanadium | 11 | 1 | 5.5 | 5.5 | 3.32 | ug/L |
| Zinc | 11 | 4 | 0.973 | 22.1 | 3.50 | ug/L |
| General Chemistry Parameters | | | | | | |
| Alkalinity (as CaCO3) | 8 | 3 | 2.06 | 7.2 | 3.23 | mg/L |
| Cyanide | 12 | 3 | 4.34 | 6.65 | 2.81 | ug/L |
| Bromoform | 66 | 1 | 0.64 | 0.64 | 0.66 | ug/L |
| Chloride | 9 | 3 | 0.048 | 4.1 | 0.55 | mg/L |
| Nitrate (as N) | 13 | 1 | 0.012 | 0.012 | 0.10 | mg/L |
| Nitrogen | 12 | 8 | 0.05 | 0.258 | 0.07 | mg/L |
| Sulfate | 9 | 1 | 4 | 4 | 0.75 | ug/L |
| TDS | 8 | 3 | 5 | 227 | 4.0 | ug/L |
| Total Kjeldahl Nitrogen | 12 | 8 | 0.03 | 0.258 | 0.058 | mg/L |

Table E-2 Concentrations of Constituents Detected in Field Blank Samples Associated with the EM and ES Grounwater Sampling Programs

| Constituent | Number of Analyses | Number of Detects | Minimum | Maximum | Average Detection Limit | Units |
|--------------|--------------------|-------------------|---------|---------|-------------------------|-------|
| Radiolucides | | | | | | |
| Gross Beta | 16 | 1 | 53.8 | 53.8 | 1.22 | pCi/L |
| Strontium-90 | 21 | 1 | 29.1 | 29.1 | 0.490 | pCi/L |
| Tritium | 72 | 1 | 569 | 569 | 380 | pCi/L |

ug/L Micrograms per liter.

mg/L Milligrams per liter.

pCi/L Picocuries per liter.

Table E-3 Concentrations of Constituents Detected in Trip Blank Samples Associated with the EM and ES Grounwater Sampling Programs

| Constituent | Number of Analyses | Number of Detects | Minimum | Maximum | Average Detection Limit | Units |
|----------------------------|--------------------|-------------------|---------|---------|-------------------------|-------|
| Volatile Organic Compounds | | | | | | |
| 1,1,2,2-Tetrachloroethane | 169 | 1 | 0.21 | 0.21 | 0.86 | ug/L |
| 1,2,3-Trichlorobenzene | 157 | 1 | 0.36 | 0.36 | 0.74 | ug/L |
| 1,2,4-Trichlorobenzene | 157 | 1 | 0.23 | 0.23 | 0.74 | ug/L |
| 2-Hexanone | 13 | 1 | 1.3 | 1.3 | 3.5 | ug/L |
| Acetone | 13 | 1 | 6.7 | 6.7 | 3.7 | ug/L |
| Benzene | 169 | 1 | 0.49 | 0.49 | 0.86 | ug/L |
| Bromoform | 169 | 1 | 1.8 | 1.8 | 0.86 | ug/L |
| Carbon tetrachloride | 169 | 1 | 2.6 | 2.6 | 0.86 | ug/L |
| Chloroform | 169 | 1 | 0.24 | 0.24 | 0.86 | ug/L |
| Dibromochloromethane | 169 | 1 | 0.73 | 0.73 | 0.86 | ug/L |
| Ethylbenzene | 169 | 1 | 0.49 | 0.49 | 0.86 | ug/L |
| m/p xylene | 166 | 2 | 0.31 | 0.98 | 0.81 | ug/L |
| Methyl chloride | 169 | 1 | 0.24 | 0.24 | 0.92 | ug/L |
| Methyl ethyl ketone | 13 | 1 | 5.9 | 5.9 | 3.7 | ug/L |
| Methyl tert-butyl ether | 133 | 1 | 0.31 | 0.31 | 1.0 | ug/L |
| Methylene chloride | 169 | 30 | 0.1 | 2.6 | 0.86 | ug/L |
| Naphthalene | 157 | 3 | 0.28 | 0.79 | 0.74 | ug/L |
| o-Xylene | 166 | 1 | 0.74 | 0.74 | 0.81 | ug/L |
| Tetrachloroethylene | 169 | 1 | 17 | 17 | 0.86 | ug/L |
| Toluene | 169 | 10 | 0.26 | 3 | 0.86 | ug/L |
| Trichloroethylene | 169 | 1 | 1.9 | 1.9 | 0.86 | ug/L |

ug/L Micrograms per liter.

Table E-4 LABORATORY DATA QUALIFIERS

2003 BNL Groundwater Status Report
Environmental Services Division
Environmental Management

Organic Analytical Data

- U Indicates that the compound was analyzed for but not detected. The sample quantitation limit must be corrected for dilution. For a soil/sediment sample, the value must also be corrected for percent moisture.

- J Indicates an estimated value. This flag is used either when estimating a concentration for tentatively identified compounds (TICs) where a 1:1 response is assumed, or when the mass spectral data indicates the presence of a compound that meets the identification criteria but the result is less than the sample quantitation limit but greater than zero.

- N Indicates presumptive evidence of a compound. This flag is used only for TICs, where the identification is based on a mass spectral library search.

- P Used for pesticide/Aroclor target analytes when there is greater than 25% difference for detected concentrations between the two gas chromatograph (GC) columns.

- C Applies to pesticide results where the identification has been confirmed by gas chromatography/mass spectrometry (GC/MS). If GC/MS confirmation was attempted but was unsuccessful, this flag is not applied; a laboratory-defined flag is used instead.

- B Used when the analyte is found in the associated blank as well as in the sample. It indicates possible/probable blank contamination and warns the data user to take appropriate action. This flag must be used for TICs as well as for positively identified target compounds.

- E Identifies compounds whose concentrations exceed the calibration range of the GC/MS instrument for that specific analysis.

- D Identifies all compounds identified in an analysis at a secondary dilution factor. This flag alerts data users that any discrepancies between the concentrations reported may be due to dilution of the sample or extract.

- A Indicates that a TIC is a suspected aldol-condensation product.

- X Other specific flags may be required to properly define the results. If used, they must be fully described and such description must be attached to the Sample Data Summary Package and the sample delivery group (SDG) narrative.

Table E-4, cont.
LABORATORY DATA QUALIFIERS

2003 BNL Groundwater Status Report
Environmental Services Division
Environmental Management

Inorganic Analytical Data

- B Indicates that the reported value was obtained from a reading that was less than the Contract Required Detection Limit (CRDL), but greater than or equal to the Instrument Detection Limit (IDL).
- U Indicates that the analyte was analyzed for but not detected.
- E Used when the reported value is estimated because of the presence of interference.
- M Indicates that the duplicate injection precision was not met.
- N Indicates that the spiked sample recovery is not within control limits.
- S Indicates that the reported value was determined by the method of standard additions (MSA).
- W Used when the post-digestion spike for furnace atomic absorption analysis is not within control limits (85%-115%), while sample absorbance is less than 50% of spike absorbance.
- * Indicates that the duplicate analysis is not within control limits.
- + Indicates that the correlation coefficient for the MSA is less than 0.995.

Table E-4, cont.
LABORATORY DATA QUALIFIERS

2003 Sitewide Groundwater Monitoring Report
Environmental Restoration Division
Brookhaven National Laboratory

Radiologic Analytical Data

- J The associated numerical value is an estimated quantity.
- JN Presumptive evidence of the presence of the material at an estimated quantity.
- DL Detection limit requirements not met. Data quality objectives may not be met.
- R The data are unusable (radionuclide may or may not be present).
- UI (Uncertain identification for gamma spectroscopy) - Radionuclide peaks that are detected but fail to meet the positive identification criteria.

Table E-5

BNL DATA VALIDATION QUALIFIERS

2003 BNL Groundwater Status Report
Environmental Services Division
Environmental Management

Data Qualifier Codes (Flags) - Code letters affixed to analytical results by the data validator to indicate the reliability and quantitative status of reported data.

- U - The analyte was analyzed for, but not detected above the reported detection limit. The associated numerical value is the sample quantitation limit.
- J - The associated numerical value was an estimated quantity.
- R - The data are unusable/unreliable.
- UJ - The analyte was analyzed for, but was not detected. The sample quantitation limit is an estimated quantity.
- F - The result is faulty due to problems outside the realm of typical validation rules/flags. This qualifier may be affixed to a result when the data validator has reason to consider the result suspect, warranting notification of the end-user of a possible problem.
- N - Tentatively Identified.

Data Subqualifier Codes (Flags) - Subqualifier codes provide additional detail on the type and amount of qualification a given data point has received.

- H Qualified due to holding time violation.
- I Qualified due to interference problems (Inductively Coupled Plasma (ICP) serial dilution or poor analytical spike recovery/RSD/CV by graphite furnace).
- D Qualified due to precision problems (duplicate control limits being exceeded).
- S Qualified due to accuracy problems (matrix spike recoveries outside control limits).
- C Qualified due to instrument calibration problems.
- L Qualified due to accuracy problems (Laboratory Control Standard (LCS) recoveries outside control limits).
- B Qualified due to blank contamination problems.
- K Qualified due to negative blank value problems.
- Q Qualified for other reasons (refer to the text of the report).
- G Qualified due to background problems.
- T Qualified due to chemical tracer or internal standard problem.
- J-X1 Result does not meet USEPA Region II validation criteria of at least 10% solids for a solid sample. However, results meet data quality objectives of the project.

Table E-5
BNL DATA VALIDATION QUALIFIERS

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Data Subqualifier Codes (continued)

- J-X2 Result is estimated and considered field data. Analytical method used was not a Federal or State approved method.
- X3 Sample was not collected by BNL personnel or a BNL representative. Location of sample collection is estimated.

Data Usability Codes (Flags)

- N1 Not usable based on potential false positives. Radionuclide results have reported activity less than or equal to the MDA and/or the uncertainty, or the LOD (2.33 times the one-sigma uncertainty)
- N2 Not usable based on the results that are not distinguishable from background. The reported activity value is less than or equal to the sum of the MDA and the uncertainty or the RDL (4.66 times the one-sigma uncertainty).
- N3 Not usable based on lack of expected daughter products.
- N4 Not usable based on local knowledge of radioactive sources and/or environmental levels.
- N5 Not usable based on level of quality for precision.
- A1 The data was considered usable.