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U.S. Nuclear Regulatory Commission Attn: Document Control Desk Washington, DC 20555

Gentlemen,

Subject: Saxton Nuclear Experimental Corporation (SNEC) Operating License No. DPR-4 Docket No. 50-146 2000 Radiological Environmental Monitoring Report

In accordance with SNEC Technical Specification Section 3.8.2.3 and the SNEC Off-Site Dose Calculation Manual Part 3, Section 1.0, the 2000 SNEC Radiological Environmental Monitoring Report is enclosed.

Please contact Art Paynter (Radiation Safety Officer) at (814) 635-4384 if you have any questions concerning this submittal.

Sincerely,

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AFP Enclosure

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IE25



2000 Radiological Environmental Monitoring Report

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ABBREVIATIONS

actinium	Ac	nitrogen	N
antimony	Sb	north	N
argon	Ar	northeast	NE
barium	Ва	north-northeast	NNE
becquerel	Bq	north-northwest	NNW
beryllium	Be	northwest	NW
carbon	С	percent	%
cesium	Cs	picocurie(s) per cubic meter	pCi/m ³
chromium	Cr	picocurie(s) per gram	pCi/g
cobalt	Со	picocurie(s) per liter	pCi/L
cubic meter(s)	m ³	picocurie(s)	pCi
curie(s) per year	Ci/yr	plutonium	Pu
curie(s)	Ci	potassium	К
curium	Cm	radium	Ra
east	E	radon	Rn
east-northeast	ENE	rem per year	rem/yr
east-southeast	ESE	Roentgen equivalent man	rem
gram(s)	g	Roentgen	R
hour(s)	h	ruthenium	Ru
hydrogen (tritium)	H-3	silver	Ag
iodine	I	south	S
iron	Fe	southeast	SE
krypton	Kr	south-southwest	SSW
lanthanum	La	southwest	SW
liter(s)	L	standard deviation	std dev
manganese	Mn	standard month	std month
mean sea level	msl	strontium	Sr
meter(s)	m	thorium	Th
microroentgen per hour	μR/h	uranium	U
millirem per hour	mrem/h	west	W
millirem per standard month	mrem/std	west-northwest	WNW
	month	west-southwest	WSW
millirem per year	mrem/yr	year(s)	yr
millirem	mrem	zinc	Zn
milliroentgen per hour	mR/h	zirconium	Zr
milliroentgen per standard month			
	month		
milliroentgen	mR		
niobium	Nb		

SUMMARY AND CONCLUSIONS

This report reviews the radiological environmental monitoring performed in 2000 for the Saxton Nuclear Experimental Corporation (SNEC) Facility. The environmental sample results indicated that SNEC operations in 2000 had no adverse effect on either the environment or the health and safety of the public.

Many of the radioactive materials discussed in this report are usually present in the environment, either from natural processes or as a result of non-SNEC activities such as prior atmospheric nuclear weapon tests and medical industry activities. To determine the impact of SNEC operations on the environment and the public, results from samples collected close to the SNEC Facility (indicator stations) were compared to results from samples obtained at distant sites (control or background stations).

The results of environmental measurements were used to assess the impact of SNEC decommissioning operations and to demonstrate compliance with the SNEC Facility Offsite Dose Calculation Manual (ODCM) (Reference 1), applicable Federal, and State regulations.

During 2000, samples of air, surface water, sediment, vegetation, and groundwater were collected. Direct radiation exposures were also measured in the vicinity of SNEC. Samples were analyzed for gross alpha and gross beta radioactivity, tritium (H-3), strontium-90 (Sr-90), and/or gamma emitting radionuclides. The results are discussed in the various sections of this report and are summarized in the following highlights:

- 262 samples were collected in 2000 from the aquatic, atmospheric and terrestrial environments around the SNEC Facility. There were approximately 500 analyses performed on these samples. In addition, 112 direct radiation exposure measurements were taken using thermoluminescent dosimeters (TLDs). The monitoring performed in 2000 met or exceeded the sample collection and analysis requirements of the SNEC Facility ODCM.
- The surface water collected downstream of the SNEC liquid discharge outfall resulted in less then detectable activities for radionuclides attributed to SNEC, including tritium (H-3).
- River sediments collected just downstream of the SNEC liquid discharge outfall and at the control station upstream of the site detected low concentrations of Cesium-137 (Cs-137). These concentrations are believed to be attributed to fallout from prior nuclear weapon tests. Cs-137 was also detected in aquatic sediments collected from storm drains that are located on site. Cesium is readily adsorbed by suspended particles in sediment.
- Three groundwater samples collected from the onsite monitoring and supply wells contained H-3 above ambient concentrations. The activities of these samples ranged from 130 to 730 picocuries per liter (pCi/L). Although humans do not consume this water, it was well below the United States Environmental Protection Agency's (USEPA's) Primary Drinking Water standard (Reference 2) of 20,000 pCi/L.

Gamma radiation exposure rates recorded at the offsite indicator TLD stations averaged 70 milliroentgens per year (mR/yr). These exposure rates were consistent with those presented by the National Council on Radiation Protection and Measurements (Reference 3). No increase in ambient gamma radiation levels was detected.

In conclusion, radioactive materials related to SNEC operations were detected in certain onsite environmental samples, but the measured concentrations were very low. During 2000, no measurable radioactive liquid effluents were released and no SNEC related radioactivity was detected in the gaseous effluent releases that occurred from SNEC. Hence, no dose to the public should be attributed to SNEC activities. The environmental sample results indicated that there was no permanent buildup of radioactive materials in the environment and no increase in background radiation levels.

Therefore, based on the results of the radiological environmental monitoring program (REMP) SNEC operations in 2000 did not have any adverse effects on the health and safety of the public or on the environment.

INTRODUCTION

Characteristics of Radiation

Instability within the nucleus of radioactive atoms results in the release of energy in the form of radiation. Radiation is classified according to its nature - particulate and electromagnetic. Particulate radiation consists of energetic subatomic particles such as electrons (beta particles), protons, neutrons, and alpha particles. Because of its limited ability to penetrate the human body, particulate radiation in the environment contributes primarily to internal radiation exposure resulting from inhalation and ingestion of radioactivity.

Electromagnetic radiation in the form of x-rays and gamma rays has characteristics similar to visible light but is more energetic and, hence, more penetrating. Although x-rays and gamma rays are penetrating and can pass through varying thickness' of materials, once they are absorbed they produce energetic electrons which release their energy in a manner that is identical to beta particles. The principal concern for gamma radiation from radionuclides in the environment is their contribution to external radiation exposure.

The rate with which atoms undergo disintegration (radioactive decay) varies among radioactive elements, but is uniquely constant for each specific radionuclide. The term "half-life" defines the time it takes for half of any amount of an element to decay and can vary from a fraction of a second for some radionuclides to millions of years for others. In fact, the natural background radiation to which all mankind has been exposed is largely due to the radionuclides of uranium (U), thorium (Th), and potassium (K). These radioactive elements were formed with the creation of the universe and, owing to their long half-lives, will continue to be present for millions of years to come. For example, potassium-40 (K-40) has a half-life of 1.3 billion years and exists naturally within our bodies. As a result, approximately 4000 atoms of potassium emit radiation within each of us every second of our lives.

In assessing the impact of radioactivity on the environment, it is important to know the quantity of radioactivity released and the resultant radiation doses. The common unit of radioactivity is the curie (Ci). The curie represents the radioactivity in one-gram (g) of natural radium (Ra), which is equivalent to a decay rate of 37 billion radiation emissions every second. Because of the extremely small amounts of radioactive material in the environment, it is more convenient to use fractions of a curie. Subunits like picocurie, pCi, (one trillionth of a curie) are frequently used to express the radioactivity present in environmental and biological samples.

The biological effects of a whole body equivalent dose of radiation are the same whether the radiation source is external or internal to the body. The important factor is how much radiation energy or dose was deposited. The unit of radiation dose is the Roentgen equivalent man (rem), which also incorporates the variable effectiveness of different forms of radiation to produce biological change. For environmental radiation exposures, it is convenient to use the smaller unit of millirem (mrem) to express dose (1000 mrem equals 1 rem). When radiation exposure occurs over periods of time, it is appropriate to refer to the dose rate. Dose rates, therefore, define the total dose for a fixed interval of time, and environmental exposures are usually expressed with reference to one year (mrem/yr).

Sources of Radiation

Life on earth has evolved amid the constant exposure to natural radiation. In fact, the single major source of radiation to which the general population is exposed comes from natural sources. Although everyone on the planet is exposed to natural radiation, some people receive more than others. Radiation exposure from natural background has three components (i.e., cosmic, terrestrial, and internal) and varies with altitude and geographic location, as well as with living habits. For example, cosmic radiation originating from deep interstellar space and the sun increases with altitude, since there is less air which acts as a shield. Similarly, terrestrial radiation resulting from the presence of naturally occurring radionuclides in the soil and rocks varies and may be significantly higher in some areas of the country than in others. Even the use of particular building materials for houses, cooking with natural gas, and home insulation affect exposure to natural radiation. The presence of radioactivity in the human body results from the inhalation and ingestion of air, food, and water containing naturally occurring radionuclides. For example, drinking water contains trace amounts of uranium and radium while milk contains radioactive potassium. Table 1 summarizes the common sources of radiation and their average annual doses.

The average person in the United States receives about 300 mrem/yr (0.3 rem/yr) from natural background radiation sources (Reference 4). This estimate was revised from about 100 to 300 mrem because of the inclusion of radon gas which has always been present but was not previously included in the calculations. In some regions of the country, the amount of natural radiation is significantly higher. Residents of Colorado, for example, receive an additional 60 mrem/yr due to the increase in cosmic and terrestrial radiation levels. In fact, for every 100 feet above sea level, a person will receive an additional 1 mrem/yr from cosmic radiation. In several regions of the world, naturally high concentrations of uranium and radium deposits result in doses of several thousand mrem/yr to their residents (Reference 5).

TABLE 1

Sources and Doses of Radiation

Natural ((32%)
Source	Radiation Dose in mrem/yr
Radon	200 (55%)
Cosmic Rays	27 (8%)
Terrestrial	28 (8%)
Internal	40 (11%)
Approximate Total mrem/yr	300
Manmade	(18%)
Source	Radiation Dose in mrem/yr
Medical X-rays	39 (11%)
Nuclear Medicine	14 (4%)
Consumer Products	10 (3%)
Other (Releases from natural gas, phosphate mining, burning of coal, weapons fallout, & nuclear fuel cycle	<1 (<1%)
Approximate Total mrem/yr	60
* Percentage contribution of the total dose is shown Reference 4.	n in parentheses. This data was obtained from

Recently, public attention has focused on radon (Rn), a naturally occurring radioactive gas produced from uranium and radium decay. These elements are widely distributed in trace amounts in the earth's crust. Unusually high concentrations have been found in certain parts of eastern Pennsylvania and northern New Jersey. Radon levels in some homes in these areas are hundreds of times greater than levels found elsewhere in the United States. Additional surveys, however, are needed to determine the full extent of the problem nationwide.

Radon is the largest component of natural background radiation and may be responsible for a substantial number of lung cancer deaths annually. The National Council on Radiation Protection and Measurements (NCRP) estimates that the average individual in the United States receives an annual dose of about 2,400 mrem to the lung from natural radon gas (Reference 4). This lung dose is considered to be equivalent to a whole body dose of 200 mrem. The NCRP has recommended actions to control indoor radon sources and reduce exposures.

When radioactive substances are inhaled or swallowed, they are not uniformly distributed within the body. For example, radioactive iodine selectively concentrates in the thyroid gland, radioactive cesium is distributed throughout the body water and muscles, and radioactive strontium concentrates in the bones. The total dose to organs by a given radionuclide is influenced by the quantity and the duration of time that the radionuclide remains in the body, including its physical, biological and chemical characteristics. Depending on their rate of radioactive decay and biological elimination from the body, some radionuclides stay in the body for very short times while others remain for years.

In addition to natural radiation, we are exposed to radiation from a number of manmade sources. The single largest of these sources comes from diagnostic medical x-rays and nuclear medicine procedures. Some 180 million Americans receive medical x-rays and nuclear medicine treatment each year. The annual dose to an individual from such radiation averages about 53 mrem. Much smaller doses come from nuclear weapon fallout and consumer products such as televisions, smoke detectors, and fertilizers. Production of commercial nuclear power and its associated fuel cycle contributes less than 1 mrem to the annual dose of about 360 mrem for the average individual living in the United States.

Fallout commonly refers to the radioactive debris that settles to the surface of the earth following the detonation of a nuclear weapon. It is dispersed throughout the environment either by dry deposition or washed down to the earth's surface by precipitation. There are approximately 200 radionuclides produced in the nuclear weapon detonation process; a number of these are detected in fallout. The radionuclides found in fallout, which produce most of the fallout radiation exposures to humans are I-131, Cs-137, Sr-89, and Sr-90.

There has been no atmospheric nuclear weapon testing since 1980 and many of the radionuclides, still present in our environment, have decayed significantly. Consequently, doses to the public from fallout have been decreasing. As a result of the nuclear accident at Chernobyl, Ukraine, on April 26, 1986, radioactive materials were dispersed throughout the environment and detected in various media such as air, milk, and soil. Cesium-134, Cs-137, I-131 and other radionuclides were detected in the weeks following the Chernobyl accident.

Description of the SNEC Site

The site is located about 100 miles east of Pittsburgh and 90 miles west of Harrisburg, Pennsylvania in the Allegheny Mountains, three-fourths of a mile north of the Borough of Saxton in Liberty Township, Bedford County, Pennsylvania. The site is on the north side of Pennsylvania Route 913, 17 miles south of U.S. Route 22, and about 15 miles north of the Breezewood Interchange of the Pennsylvania Turnpike.

The SNEC Facility was built adjacent to the Saxton Steam Electric Generating Station of Pennsylvania Electric Company (Penelec), a subsidiary of GPU. This coal fired station operated from 1923 to 1974 and was demolished between 1975 and 1977. The SNEC site consists of 1.148 fenced acres of the approximate 150 acres owned by Penelec. An additional 9.6-acre fenced area contains an electrical switchyard and buildings under Penelec control. A general property layout is shown in Figure 1. The SNEC site, as well as a portion of the Penelec area and the surrounding uncontrolled lands, is in the 100-year floodplain of the

Raystown Branch of the Juniata River which borders the north and west portion of the property. A small stream known as Shoup Run crosses the central portion of the property and joins the Juniata River. Normal elevation of the river near the SNEC site is approximately 794 feet above mean sea level (msl).

The SNEC site and adjacent property lie about 17 feet above river level. Much of the property consists of gently sloping open grassland, a result of the restoration activities following the demolition of the Saxton Steam Electric Generating Station.

SNEC Decommissioning Operations

The Saxton Nuclear Experimental Facility was a pioneer in the development of the nuclear energy program for the United States. It operated for ten years, from 1962 to 1972, and provided valuable information on operations and training. The fuel was removed from the Containment Vessel (CV) in 1972 and shipped to the Atomic Energy Commission (AEC) Facility at Savannah River, South Carolina. Following fuel removal, equipment, tanks, and piping located outside the CV were removed. Final decontamination and dismantlement of reactor support structures and buildings were completed in 1992.

On April 20, 1998, the U.S. Nuclear Regulatory Commission (NRC) gave its approval for the final stage of decommissioning. The following is a list of work activities that had potential to impact sample results required by the SNEC Facility REMP since April 20, 1998:

- November 1998 The large component structures, pressurizer, steam generator and reactor vessel, were removed and shipped to Chem-Nuclear's low-level waste facility in Barnwell, South Carolina.
- Early 2000 The remaining miscellaneous components were removed from the CV and concrete remediation operations commenced.
- May 2000 Soil excavation and remediation activities began in the CV North Yard.
- October 2000 Soil excavation and remediation activities began in the Saxton Steam Generating Station (SSGS) footprint.

The only remaining SNEC Facility structures include the CV, the two remaining concrete shield walls, and tunnel sections that are immediately adjacent to the outer circumference of the CV. Concrete barrier walls have been installed to isolate the open ends of the tunnel that were connected to the Control & Auxiliary Buildings, the Radioactive Waste Disposal Facility and the Steam Plant. Portions of the Steam Plant Tunnel still exist beyond the point where the tunnel was blocked-off. Additional information can be obtained from the 2000 SNEC Annual Operations Report (Reference 6).

Containment Vessel

The SNEC CV is a circular steel structure approximately 100 ft. tall by 50 ft. in diameter with approximately 50 percent of the structure below grade. The CV is subdivided into a Reactor Compartment/Storage Well, Primary Compartment, Auxiliary Compartment and an Operating Floor. Concrete walls, floors, and ceilings separate these areas from each other. The below grade portion of the CV is lined with concrete, as well.

Demography - Human Activities in the Environs

The area surrounding the SNEC site is generally rural forested and mountainous terrain. The population density of the area is low with small concentrations in the valleys and along main highways. The site lies about three-fourths of a mile north of the Borough of Saxton in Liberty Township, Bedford County, Pennsylvania. The population and population trends for the Borough of Saxton in Bedford County have decreased by approximately 4.2% between 1990 and 2000 (Reference 7). At the time the SNEC Facility was constructed, the estimated population of the Borough of Saxton was 975 as recorded during the 1960 census. Forty years later, the population as recorded during the 2000 census was 803, a decline of 17.6%.

The nearest population center (as defined by 10 CFR 100) of 25,000 or more is the city of Altoona in Blair County which lies approximately 20 miles north-northwest of the SNEC site. The 2000 population of Altoona was 49,523. The closest incorporated towns other than the Borough of Saxton are Coalmont Borough about 2.5 miles to the east (2000 population of 128), Dudley Borough about 3.4 miles to the east (2000 population of 192) and Broad Top about 5.3 miles also to the east (2000 population of 384).

Current uses of adjoining properties include undeveloped wooded and residential areas. A cemetery lies along the eastern property boundary while undeveloped wooded and residential areas border the northern, southern and western property boundaries.

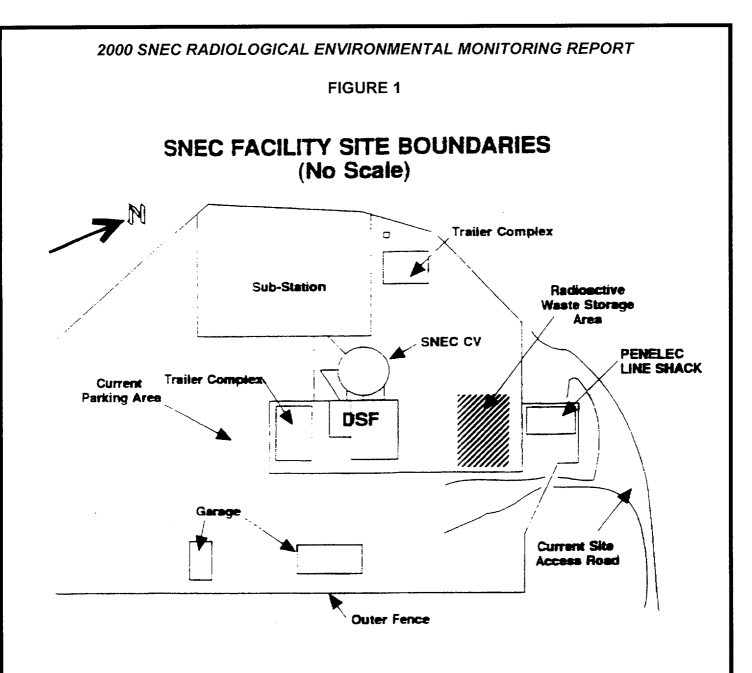
The Raystown Branch of the Juniata River in the vicinity of the site is primarily used for recreational boating and fishing by local residences. The vast majority of recreational activities along the river, however, are located downstream of the site on Raystown Lake.

Approximately 34 miles downstream of the site, the Raystown Branch of the Juniata River is dammed, impounding the river to form Raystown Lake. The dam built by the US Army Corps of Engineers (COE) for flood control, recreation and water quality purposes was constructed from 1968 to 1973. At normal pool level, the lake is 27 miles long and has an area of 8,300 acres. Raystown Lake provides one of the most popular recreational areas for boating, fishing, camping, hunting, and picnicking in this part of Pennsylvania.

<u>Geology</u>

The SNEC site lies in the Appalachian Highlands in the Ridge and Valley Physiographic Province. This province comprises alternate successions of narrow ridges and broad or narrow valleys trending generally northeast. This region contains alternating hard and soft sedimentary rocks that have been severely folded by lateral compression into a series of anticlines and synclines. The ridge consists of Tuscarora quartzite and a small amount of

Pleistocene gravel. Most of the area is underlain by strata of Upper Devonian age. Although coal was mined in the general area of the site, no coal has been reported to lie beneath the site, nor has the site been undermined. The ridges immediately to the northwest of the site rise to 1300 feet and to the southeast rise to 1500 feet with site elevation being approximately 811 feet above msl.



RADIOLOGICAL ENVIRONMENTAL MONITORING

A comprehensive Radiological Environmental Monitoring Program (REMP) is performed at the SNEC Facility to measure levels of radiation and radioactive materials in the environment. The information obtained from the REMP is then used to determine the effect of SNEC operations, if any, on the environment and the public.

The USNRC has established regulatory guides that contain acceptable monitoring practices. The SNEC REMP was designed on the basis of these regulatory guides along with the guidance provided by the USNRC Radiological Assessment Branch Technical Position for an acceptable radiological environmental monitoring program (Reference 8). The SNEC REMP meets or exceeds the monitoring requirements set forth by the USNRC.

The important objectives of the REMP are:

- To assess dose impacts to the public from the SNEC Facility.
- To verify decommissioning controls for the containment of radioactive materials.
- To determine buildup of long-lived radionuclides in the environment and changes in background radiation levels.
- To provide reassurance to the public that the program is capable of adequately assessing impacts and identifying noteworthy changes in the radiological status of the environment.
- To fulfill the requirements of the SNEC Technical Specifications.

<u>Sampling</u>

The program consists of taking radiation measurements and collecting samples from the environment, analyzing them for radioactivity content, and then interpreting the results. These samples include, but are not limited to; air, water, sediment, vegetation, groundwater and thermoluminescent dosimeters (TLDs) to measure gamma radiation levels.

The SNEC Facility ODCM (Reference 1) defines the sample types to be collected and the analyses to be performed. However, the minimum sampling and analysis requirements specified in the ODCM are maintained or exceeded. As appropriate, changes to the REMP are initiated by recommendations from SNEC Radiological Controls Management.

Sampling locations were established by considering topography, meteorology, population distribution, hydrology, and areas of public interest. The sampling locations are divided into two classes, indicator and control. Indicator locations are those that are expected to show effects from SNEC activities, if any exist. These locations were selected primarily on the basis of where the highest predicted environmental concentrations would occur. The indicator locations are typically within the site boundary, along the perimeter fence or a few miles from the SNEC Facility.

Control stations are located generally at distances greater than 10 miles from the SNEC Facility. The samples collected at these sites are expected to be unaffected by SNEC operations. Data from control locations provide a basis for evaluating indicator data relative to natural background radioactivity and fallout from prior nuclear weapon tests. Figures 2 and 3 show the current sampling locations around the SNEC Facility. Table 11 in Appendix A describes the sampling locations along with the type(s) of samples collected at each sampling location.

<u>Analysis</u>

In addition to specifying the media to be collected and the number of sampling locations, the ODCM also specifies the frequency of sample collection and the types and frequency of analyses to be performed. Also specified are analytical sensitivities (detection limits) and reporting levels. Table 12 in Appendix A provides a synopsis of the sample types, number of sampling locations, collection frequencies, number of samples collected, types and frequencies of analyses, and number of samples analyzed. Table 13 in Appendix "A" lists samples, which were not collected or analyzed as per the requirements of the ODCM. Changes to the REMP are described in Appendix C. All samples analyzed in 2000 met the required analytical sensitivities.

Measurement of low radionuclide concentrations in environmental media requires special analysis techniques. Analytical laboratories use state-of-the-art laboratory equipment designed to detect all three types of radiation emitted (alpha, beta, and gamma). This equipment must meet the analytical sensitivities required by the ODCM. Examples of the specialized laboratory equipment used are germanium detectors with multichannel analyzers for determining specific gamma-emitting radionuclides, liquid scintillation counters for detecting H-3, low level proportional counters for detecting gross alpha and beta radioactivity and alpha spectroscopy for determining specific transuranic isotopes.

Counting equipment calibrations are performed by using standards traceable to the National Institute of Standards and Technology (NIST). Computer hardware and software are used in conjunction with the counting equipment to perform calculations and provide data management. Analysis methods are described in Appendix H.

Data Review

The REMP Coordinator and Quality Assurance Officer routinely review REMP analytical results to assure that sensitivities have been achieved and that the proper analyses have been performed. Investigations are conducted when action levels or USNRC reporting levels are reached or when anomalous values are discovered. This review process is discussed in more detail in Appendix D.

Tables 2 and 3 summarize gamma radiation exposure rates near the SNEC Facility from thermoluminescent dosimeter (TLD) measurements. Tables 4 and 5 depict airborne particulate gross beta and gross alpha results, respectively. Table 6 lists the tritium concentrations from station GEO-5 and Table 7 lists tritium results from all 2000 monitoring wells. Table 10 provides a summary of radionuclide concentrations detected in the aquatic

sediment samples for 2000. Statistical methods used to derive these tables along with other statistical conclusions are detailed in Appendix G. Quality control (QC) sample results were used mainly to verify the primary sample result or the first result in the case of a duplicate analysis. Therefore, the QC results were excluded from these tables and the main text of this report to avoid biasing the results.

Quality Assurance Program

A quality assurance (QA) program is conducted in accordance with guidelines provided in Regulatory Guide 4.15, "Quality Assurance for Radiological Monitoring Programs" (Reference 9) and as required by the ODCM. SNEC maintains written policies, procedures, and records that encompass all aspects of the REMP including sample collection, equipment calibration, laboratory analysis, and data review.

The QA program is designed to identify possible deficiencies so that immediate corrective action can be taken. The program also provides assurance to the regulatory agencies and the public that the results are valid. The QA program for the measurement of radioactivity in environmental samples is implemented by:

- Auditing all REMP-related activities including analytical laboratories.
- Requiring analytical laboratories to participate in a crosscheck program(s).
- Requiring analytical laboratories to split samples for separate analysis (recounts are performed when samples cannot be split).
- Splitting samples, having the samples analyzed by independent laboratories, and then comparing the results for agreement.
- Reviewing QC results of the analytical laboratories including spike and blank sample results and duplicate analysis results.

The QA program and the results of the crosscheck programs are outlined in Appendix E and F, respectively.

The TLD readers are calibrated monthly against standard TLDs to within five percent of the standard TLD values. Also, each group of TLDs processed by a reader contains control TLDs that are used to correct for minor variations in the reader. The accuracy and variability of the results for the control TLDs are examined for each group of TLDs to assure the reader is functioning properly. In addition, each element (TLD) has an individual correction factor based on its response to a known exposure.

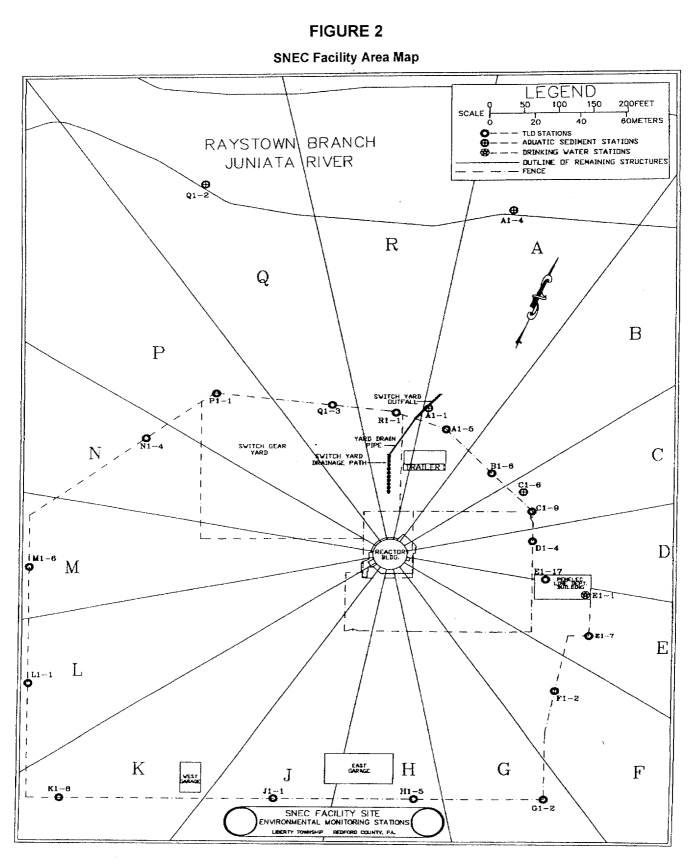
Other crosschecks, calibrations, and certifications used to assure the accuracy of the TLD program include:

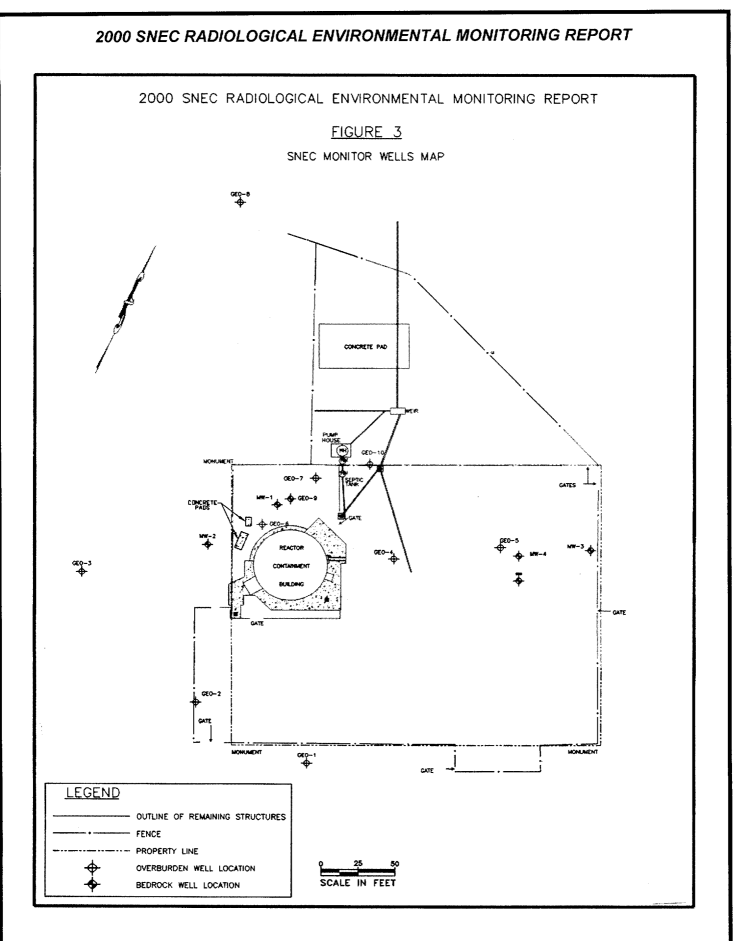
• Semiannually, randomly selected TLDs are sent to an independent laboratory where they are irradiated to set doses unknown to SNEC. TLDs, which meet the criteria specified by the National Voluntary Laboratory Accreditation Program (NVLAP), are used for this test.

The dosimetry vendor (AmerGen) processes the TLDs and the results are compared against established limits.

- Every two years, each TLD is checked to ensure an appropriate correction factor is assigned to each element of the TLD.
- Every two years, the dosimetry program is examined and NVLAP recertified by the NIST.

The environmental dosimeters were tested and qualified to the American National Standard Institutes (ANSI) publication N545-1975 and the USNRC Regulatory Guide 4.13 (References 10 and 11).





DIRECT RADIATION MONITORING

Radiation is a normal component of the environment resulting primarily from natural sources, such as cosmic radiation and naturally occurring radionuclides, and to a lesser extent from manmade sources, such as fallout from prior nuclear weapon tests. The cessation of atmospheric nuclear weapon tests and the decay of fallout products have resulted in a gradual decrease in environmental radiation levels. Direct radiation monitoring is used to measure ionizing radiation primarily from cosmic and terrestrial sources.

Gamma radiation exposure rates near the SNEC Facility were measured using thermoluminescent dosimeters (TLDs). There are 28 TLDs that surround the SNEC Facility. Sixteen Indicator Stations, one per compass sector, are located at the outer perimeter fence. One station is located in the Penelec Line Department garage. There are nine Offsite Indicator Stations in various sectors within two miles of the site. There are also two Control Stations, each about 10 miles from the site. The indicator stations are used to detect any potential effect of SNEC Facility activities on environmental radiation levels. No relationship between the SNEC Facility activities and offsite exposure rates were detected at any station. The 2000 quarterly exposure rates for the individual TLD stations are contained in Appendix I.

Sample Collection and Analysis

A thermoluminescent dosimeter (TLD) is composed of a crystal (phosphor) which absorbs and stores energy in traps when exposed to ionizing radiation. These traps are so stable that they do not decay appreciably over time. When heated, the crystal emits light proportional to the amount of radiation received, and the light is measured to determine the integrated exposure. This process is referred to as thermoluminescence. The reading process 'rezeros' (anneals) the TLD and prepares it for reuse. The TLDs in use for environmental monitoring at the SNEC Facility are capable of accurately measuring exposures from a minimum of 1 mR (well below normal environmental exposures for the quarterly monitoring periods) to a maximum of 200,000 mR.

Each TLD station consists of 2 TLD badges, each of which has 4 phosphors or elements. Since each TLD responds to radiation independently, this provides 8 independent detectors at each station. The elements within the TLDs are composed of calcium sulfate and lithium borate. The calcium sulfate elements are shielded with a thin layer of lead making the response to different energies of gamma radiation more linear. The lead also shields the elements from beta radiation, making them sensitive to gamma radiation only. The lithium borate element is shielded differently to permit the detection of beta radiation as well as gamma. The combination of different phosphor materials, shielding, and multiple phosphors per badge permit quantification of both gamma and beta radiation. Only the calcium sulfate elements can be used to evaluate beta exposures or as a backup to the calcium sulfate elements should more data be required.

Data from the TLDs were evaluated by obtaining the average of the usable element results at each station. All TLD exposure rate data presented in this report were normalized to a

standard month (std month) to adjust for variable field exposure periods. A std month is 30.4 days. Several control TLDs were used to quantify transit exposure during TLD storage and handling. Transit exposures were subtracted from gross field exposures to produce net field exposures.

<u>Results</u>

In 2000, the average annual exposure rate for offsite indicator stations was 5.81 mR/std month. Quarterly exposure rates ranged from 4.5 to 8.6 mR/std month. This equates to an annual exposure rate of 69.7 mR/yr. Exposure of this magnitude is comparable with the annual average dose a person receives from cosmic and terrestrial sources (Table 1, "Sources and Doses of Radiation").

Offsite indicator station E2-1, located 0.25 mile from the CV, displayed the highest elevated exposures. An investigation revealed that the elevated results were due to residual waste from the coal-fired station formerly located adjacent to the SNEC Facility. The soil at station E2-1 consists of a mixture of coal slag and cinders, which emit a slightly elevated gamma from naturally occurring Th-232 and Ra-226, and thus adding to the TLD results.

The average annual exposure rate for the two control stations, those stations farther than 10 miles from SNEC, was 5.5 mR/std month. Quarterly exposure rates at the control stations ranged from 4.8 to 5.9 mR/std month. Table 2 depicts the average offsite indicator results with the average control results. The TLD along the perimeter fence from sector D showed the highest results due to the storage of radioactive materials proximal to the subject TLD station.

No elevated exposure rates attributed to the SNEC Facility activities were observed at any offsite station. TLDs are sensitive and accurate mechanisms for measuring the low exposure rates characteristic of environmental levels. Effects of normal SNEC Facility activities, however, are too small to be discernible outside the normal range of background radiation levels. Table 3 compares the highest site boundary exposure result to the allowable maximum exposure rate based on 40 CFR 190 (Reference 12).

TABLE 2

2000 SNEC TLD Summary

Field Cycle: January 6, 2000 to January 15 and 16, 2001

	MEAN (mR/std month)	MINIMUM (mR/std month)	MAXIMUM (mR/std month)
Average Offsite Indicator Stations	5.8	4.5 @ K1-5	8.6 @ E2-1
Average Control Stations	5.5	4.8 @ H10-1	5.9 @ G10-2

TABLE 3

Highest Site Boundary Exposure Comparison

7.2	mR/std month		Compared to an allowable maximum exposure rate of 0.37 mR/hr. This is equivalent to the 25 mR annual limit specified by 40 CFR 190
0.010	mR/hr	@ Station F1-2	adjusted by the 67-hour recreational factor specified in Reg. Guide 1.109 (shoreline exposure for maximum exposed teenager) (Reference 12).

ATMOSPHERIC MONITORING

A potential exposure pathway to humans is the inhalation of airborne radioactive materials. To monitor this exposure pathway, ambient air was sampled by a network of continuously operating samplers and then analyzed for radioactivity content. Based on the analytical results, no contribution to the general levels of airborne radioactivity was attributed to the SNEC Facility during 2000.

The indicator air sampling stations are located in the three predominant wind sectors around the Containment Vessel (CV), the north sector (A1-2), the east sector (D1-1), and the south sector (J1-3). The control air sampling station (G10-1), which is 10 miles from the site, provided background airborne radioactivity data for comparison.

Sample Collection and Analysis

Mechanical air samplers were used to continuously draw air through glass fiber filters. To maintain a constant flow rate throughout the collection period, each sampler was equipped with a mass flow probe. This electronic device maintains a constant airflow across the filter paper. All air samplers were calibrated semiannually and maintained by Radiological Instrument Technicians.

The glass fiber filters were used to collect airborne particulate matter. The filters were collected weekly or bi-weekly and analyzed for gross alpha and gross beta radioactivity. The filters were then combined (composited) quarterly by individual station locations and analyzed for gamma-emitting radionuclides.

Air Results

During 2000, 181 air particulate samples (filters) were collected and analyzed for gross alpha and gross beta radioactivity. The particulate matter (dust particles) collected on all indicator and control filters contained gross beta radioactivity above the minimum detectable concentration (MDC). The gross beta concentrations measured on the filters collected from indicator sites ranged from 0.0065 \pm 0.0025 pCi/m³ to 0.041 \pm 0.004 pCi/m³ and averaged 0.017 \pm 0.0024 pCi/m³. The air particulate samples collected from the control location had gross beta concentrations, which ranged from 0.0046 \pm 0.0023 pCi/m³ to 0.037 \pm 0.001 pCi/m³ and averaged 0.0166 \pm 0.0026 pCi/m³. The average results are listed in Table 4.

As depicted in Figure 4, average weekly gross beta concentrations at indicator and control air monitoring locations were analogous and trended similarly throughout the monitoring period. The weekly gross beta concentrations and trends at individual air sampling sites also were similar. The 2000 data indicated that gross beta radioactivity levels did not change as a result of SNEC operations. Additionally, the gross beta radioactivity associated with airborne particulate was due to naturally occurring radionuclides.

Air particulate gross alpha concentrations (detected above the MDC) at indicator stations ranged from 0.00079 \pm 0.00048 pCi/m³ to 0.0045 \pm 0.0009 pCi/m³ and averaged 0.0017 \pm 0.0006 pCi/m³. Control samples averaged 0.0017 \pm 0.0006 pCi/m³ and ranged from 0.00083 \pm 0.00054 pCi/m³ to 0.0041 \pm 0.0008 pCi/m³.

Average weekly gross alpha concentrations are depicted in Figure 5. Due to a naturally occurring alpha ingrowth that occurs in air particulate samples, variations in concentrations were observed. As the time between sample collection and sample analysis increased, so did the ingrowth of alpha resulting in higher sample activity. Inconsistent or late sample deliveries to the analytical laboratory promoted these variations of concentrations. Actual concentrations (whether the count rates were above, below, or equal to the blank count rates) were used to calculate weekly averages because approximately 22% (40 of 181) of the weekly results were below the MDC. Using actual concentrations eliminates biases in the data and missing data points on graphs. As depicted in Figure 5, average weekly gross alpha concentrations at indicator and control stations remained relatively constant throughout the monitoring period. Generally, the trends of average gross alpha concentrations at indicator and control sites are listed in Table 5.

The data obtained in 2000 indicated that gross alpha radioactivity levels did not change as a result of SNEC Facility operations. In addition, the gross alpha radioactivity measured on the particulate filters was caused by naturally occurring radionuclides.

Gamma-emitting radionuclides related to the SNEC Facility were not detected on any of the quarterly composites that were analyzed in 2000. As expected, all of the quarterly composite samples contained naturally occurring beryllium-7 (Be-7). Concentrations detected on indicator samples were similar to those detected on control filters.

TABLE 4

2000 Average Gross Beta Concentrations in Airborne Particulates (pCi/m^3)

Station	Description	Average ± 2 std deviations*
A1-2 (I)	North Sector	0.0147 ± 0.002
D1-1 (I)	East Sector	0.02 ± 0.0027
J1-3 (I)	South Sector	0.015 ± 0.0025
G10-1 (C)	New Granada	0.0166 ± 0.0026

TABLE 5

2000 Average Gross Alpha Concentrations in Airborne Particulates (pCi/m³)

Station	Description	Average ± 2 std deviations*
A1-2 (I)	North Sector	0.0016 ± 0.006
D1-1 (I)	East Sector	0.0018 ± 0.006
J1-3 (I)	South Sector	0.0017 ± 0.0006
G10-1 (C)	New Granada	0.0017 ± 0.0006

FIGURE 4

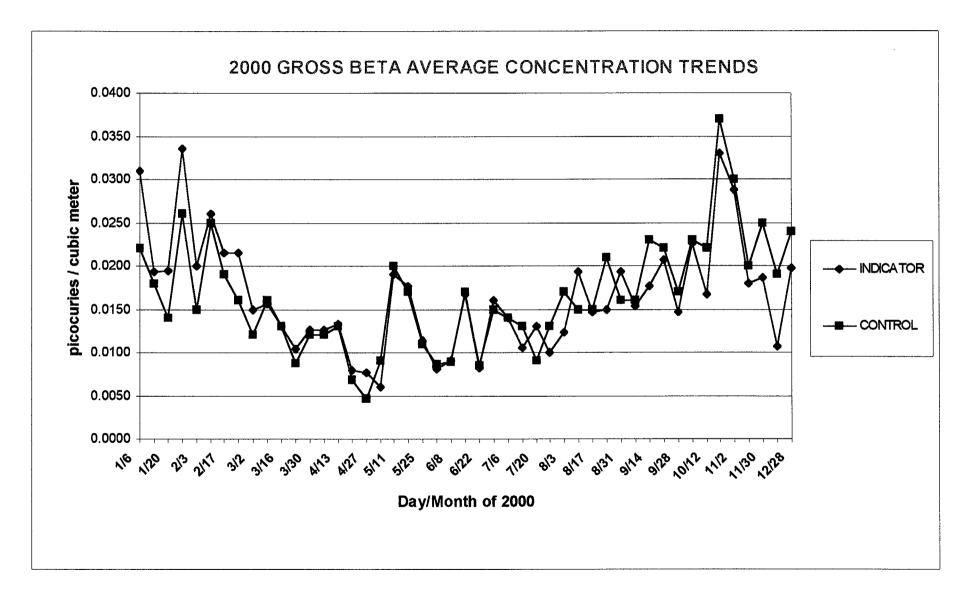
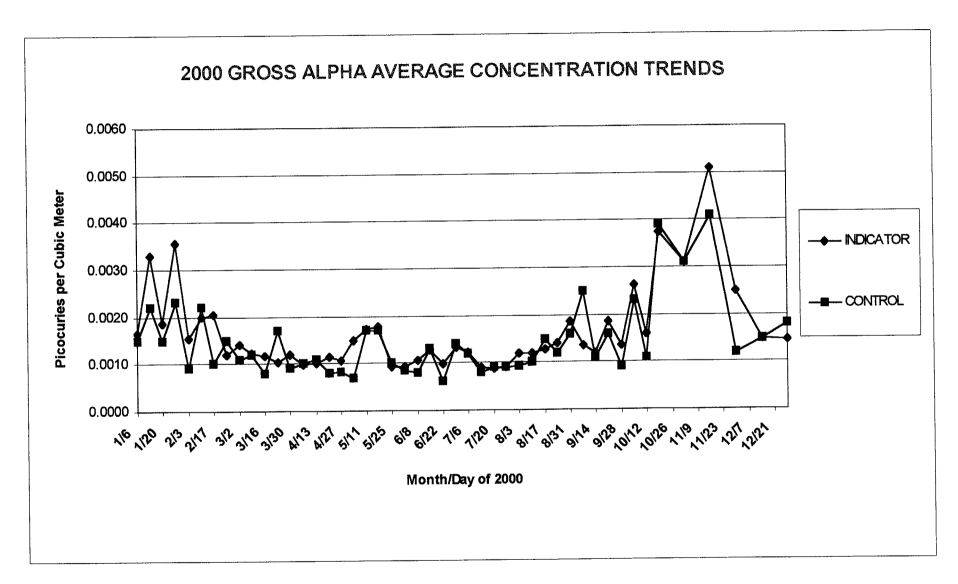


FIGURE 5



GROUNDWATER MONITORING

Groundwater monitoring is conducted to check for water leakage, if any, from the SNEC Containment Vessel. An investigation was performed to define the depth of the bedrock surface and the orientation of the bedrock groundwater flow pathways (Reference 14). The site is immediately underlain by a fill-layer composed of fly ash, cinders and/or silt and sand-size sediment. A layer of boulders in a silty clay matrix underlies this fill-layer. The surface of the bedrock lies beneath this boulder layer at a depth of approximately 7.5 to 18 feet.

The results of this investigation indicate that the overburden groundwater occurs at a depth ranging from approximately 4 to 16 feet. Groundwater elevation contour maps of this data indicate that the groundwater within the overburden soil flows west toward the Raystown Branch of the Juniata River.

Groundwater movement within the bedrock beneath the site is predominately controlled by fractures in the bedrock. There are two major fracture patterns; one trends northeast to southwest, and dips moderately toward the northwest. The second fracture pattern trends northwest to southeast, and dips steeply toward the southwest (Reference 14). Groundwater also moves within the spaces (bedding planes) between the individual layers of the siltstone bedrock.

In 1994, eight overburden groundwater wells were restored. Four of the wells are hydraulically downgradient of the Containment Vessel (GEO-3, GEO-6, GEO-7, and GEO-8). The other four wells (GEO-1, GEO-2, GEO-4, and GEO-5) serve as background monitoring points, since these wells are located hydraulically upgradient of the Containment Vessel.

Two bedrock wells (MW-1 and MW-2) were also monitored. As part of the analysis performed by the contracted hydrogeologic consultants (GEO Engineering), it was determined that bedrock monitoring wells should be installed at an angle in order to maximize the interception of fractures and bedding planes. The boreholes were drilled into bedrock at an angle of approximately 25 degrees from vertical to accomplish this. Filling the annular space with a sand filter pack, a bentonite pellet seal and cement grout allows these wells to monitor only the significant fractures and bedding planes of the bedrock ground water.

In May of 1998, three additional monitoring wells were drilled. Two bedrock wells (MW-3 and MW-4) were installed to determine if there was contamination in the vicinity of the former Radwaste Facility Building. This area was monitored by GEO-5, which in the past was the only well to show positive tritium levels. An additional overburden well (GEO-10) was installed to supplement the existing monitoring wells to monitor for the possible migration of trace amounts of tritium or other contaminants.

In addition, two off-site (potable water) samples were collected. One site monitored the well water from the Pennelec Line Department garage located adjacent to the site. The other sample collected was from a resident in the borough of Saxton. The resident water sample was initially believed to be well water, but it was later ascertained that this water was actually from the township water supply. All Saxton township residents get their domestic water from one of two sources. Putts Hollow reservoir is the primary source, but during low water levels,

the township switches to the Seton Plant water supply, which draws from the Juniata River upstream of the SNEC Facility. Neither of these samples detected any radioactive contaminants.

In May of 2000, three monitoring wells (MW-1, GEO-6, and GEO-7) and piezometer (GEO-9) were removed from the SNEC Facility Yard, North of the CV. These wells were removed to support soil remediation activities.

In October of 2000, one additional monitoring well (GEO-2) was removed from the area west of the SNEC property line near the Saxton Steam Generating Station (SSGS) footprint. This well was removed to support remediation of the SSGS footprint. It was initially believed to be sampling groundwater outside of the SSGS footprint. But when excavation began, the well was actually found installed within the SSGS footprint.

Groundwater Results

Locations of the onsite groundwater stations sampled in 2000 are shown in Figure 3. The results from the analyses performed on these samples indicated no radioactive contamination from plant-related radionuclides other than tritium. Of the 41 groundwater samples collected in 2000, three samples contained H-3 ranging from 130 to 730 pCi/L, which is considered slightly above ambient concentrations, but are well below the USEPA's Primary Drinking Water Standard of 20,000 pCi/L. The required sensitivities for SNEC are contained in Table 16.

As stated earlier, GEO-5 originally was the only well to show positive tritium levels. The first sample obtained from GEO-5 was collected and analyzed July of 1994. A "Less Than" result for tritium was reported. Gamma analysis performed on this sample yielded "Less Than" activities. The October 1994 sample reported 560 pCi/L tritium. A special collection was performed two weeks later to confirm the positive tritium and a result of 310 pCi/L was obtained. Gamma analysis continued to show no reportable activity. The highest activity of tritium (760 pCi/L) was observed in October 1995. Since that time, no concentrations above 200 pCi/L were observed. Upon review of these results, it appears that the activity in the GEO-5 area can be attributed to pockets of tritiated water trapped in fractures leading to the overburden groundwater. Table 6 is a list of all tritium results that have been performed since the start of GEO-5 monitoring.

In order to assess the possibility of other contaminants in the vicinity of GEO-5, SNEC contracted Haley & Aldrich, Inc. (formally GEO Engineering) to add supplemental monitoring wells in this location (Reference 15). The new wells showed infrequent tritium activity slightly above the MDC except for one sample that was obtained on 7/13/00 from monitoring well MW-4, which is located immediately adjacent to GEO-5. This sample yielded a positive tritium result of 730 pCi/L \pm 110 pCi/L; well below the reporting level of 2.0 E4 pCi/L. Subsequent samples obtained after 7/13/00 yielded "Less Than" activities. An ongoing evaluation is being performed by the REMP Coordinator to determine the cause of the increase in activity.

All of the new monitoring wells, like the former wells, yielded "less than" activities for gamma analysis. Table 7 is a list of tritium results from all the monitoring wells sampled in 2000. The results indicate that no other contaminants are present in the groundwater.

TABLE 6

SX-GW-GEO-5 Tritium Results in pCi/L Activity $\pm 2\sigma$

DATE	RESULTS
7/13/94	< 170
10/06/94	560 ± 130
10/27/94	310 ±120
1/12/95	< 190
4/05/95	< 180
5/30/95	270 ± 120
6/13/95	370 ± 130
7/13/95	370 ± 110
8/17/95	390 ± 130
9/15/95	410 ± 130
10/18/95	760 ± 140
11/17/95	< 200
1/25/96	< 190
4/03/96	< 150
7/10/96	< 140
10/03/96	< 140
1/08/97	< 140
4/16/97	< 150
7/09/97	< 150
10/01/97	180 ± 100
1/08/98	< 150
4/15/98	140 ± 80
7/09/98	< 120
10/08/98	< 130
1/19/99	200 ± 90
4/15/99	< 160
7/22/99	200 ± 90
10/14/99	< 130
1/06/00	< 130
4/06/00	< 120
7/13/00	190 ± 80
10/11/00	< 644

TABLE 7

2000 Tritium Results of Ground Water Analysis in pCi/I

Monitoring Well	First Qtr 01/06/00 ②	Second Qtr 04/06/00 ②	Third Qtr 07/13/00 ②	Fourth Qtr 10/11/00 ③
GEO-1	< 130	0	< 120	< 644
GEO-2	< 130	< 120	< 120	< 644
GEO-3	< 130	< 120	< 120	< 644
GEO-4	< 130	130 ± 80	< 120	< 644
GEO-5	< 130	< 120	190 ± 80	< 644
GEO-6	0	0	0	0
GEO-7	< 130	< 120	0	0
GEO-8	< 130	< 120	< 120	0
GEO-10	< 130	< 120	< 120	< 644
SX-GW-MW1	0	< 120	Ð	0
SX-GW-MW2	< 130	< 120	< 120	< 644
SX-GW-MW3	< 130	< 120	< 120	< 644
SX-GW-MW4	< 130	< 120	730 ± 110	< 644
SX-GW-E1-1	< 130	< 120	< 120	0
SX-DW-G1-1	0	< 120	< 120	< 644

Activity $\pm 2 \sigma$

① No sample collected. See sampling and analysis exceptions in Table 13.

② Samples were analyzed by AmerGen Environmental Radioactivity Laboratory.

③ Samples were analyzed by BWX Technologies Inc.

BROAD LEAF VEGETATION MONITORING

Radionuclides released into the atmosphere may deposit on vegetation. To assess the deposition, yearly broad leaf vegetation samples are collected and analyzed for gammaemitting isotopes. Collection occurred during the growing season from two different sectors on site where the prevailing wind direction has been determined. Naturally-occurring Be-7 and K-40 were measured in all samples. No radionuclides attributable to SNEC operations were detected above the MDC.

TABLE 8

2000 Gamma Spectrometry Results from Broad Leaf Vegetation (pCi/g Wet)

Sample Designation	Date	Co-60	Cs-134	Cs-137
SX-BR-A1-6 (Sector A)	7/13/00	< 0.018	< 0.016	< 0.018
SX-BR-B1-7 (Sector B)	7/13/00	< 0.02	< 0.019	< 0.02

SURFACE WATER MONITORING

The Juniata River surface water was monitored for radionuclides of potential SNEC origin. Two grab samples, one control and one indicator, were collected on a quarterly basis and analyzed for gamma emitting radionuclides and tritium. The indicator sample was collected at the discharge bulkhead leading into the river, while the control sample was collected upstream of the discharge. No tritium or radionuclides attributed to SNEC operations were detected above the MDC.

TABLE 9

Quarterly Results of Surface Water in pCi/I

Sample Designation	First Qtr 01/06/00 ①	Second Qtr 04/06/00 ①	Third Qtr 07/13/00 ①	Fourth Qtr 10/11/00 ②
	< 1.5 Cs-137	< 3 Cs-137	< 1.5 Cs-137	< 10.2 Cs-137
	< 1.5 Cs-134	< 3 Cs-134	< 1.6 Cs-134	< 10.1 Cs-134
A1-4 (I)	< 1.6 Co-60	< 3 Co-60	< 1.9 Co-60	< 12.5 Co-60
	< 130 H-3	< 130 H-3	< 120 H-3	< 130 H-3
	< 3 Cs-137	< 1.6 Cs-137	< 1.6 Cs-137	< 14.4 Cs-137
	< 2 Cs-134	< 1.6 Cs-134	< 1.5 Cs-134	< 12.6 Cs-134
Q1-2 (C)	< 3 Co-60	< 1.9 Co-60	< 1.7 Co-60	< 7.5 Co-60
	< 130 H-3	< 130 H-3	< 120 H-3	< 130 H-3

① Samples were analyzed by AmerGen Environmental Radioactivity Laboratory.

② Samples were analyzed by BWX Technologies Inc.

(I) = Indicator Station

(C) = Control Station

AQUATIC SEDIMENT MONITORING

Sediment samples were collected from on-site storm drains on a quarterly basis (Stations A1-1 and C1-6). In addition, a sediment sample taken directly from the Juniata River at the discharge bulkhead (A1-4), as well as a control sediment sample (Q1-2) taken up river from the discharge, was also collected. The sediment is dried and then analyzed for gamma emitting radioisotopes.

Low concentrations of cesium-137 (Cs-137) were detected in both river sediments (A1-4 and Q1-2). These concentrations are believed to be attributed to fallout from prior nuclear weapon tests.

SNEC-related cesium-137 (Cs-137) was detected in all the sediments collected on site. The average activity was 0.82 pCi/g dried. Cs-137 is readily adsorbed by suspended particles and is concentrated in the storm drains.

TABLE 10

Quarterly Results of Sediment Analysis in pCi/g (Dry) Activity ± 2 σ

Sample	First Qtr	Second Qtr	Third Qtr	Fourth Qtr
Designation	01/06/00	04/06/00	07/13/00	10/11/00
A1-1 (I)	0.92 ± 0.09 Cs-137	1.9 ± 0.2 Cs-137	1.3 ± 0.1 Cs-137	1.18 ± 0.07 Cs-137
	< 0.03 Cs-134	< 0.02 Cs-134	< 0.02 Cs-134	< 0.04 Cs-134
	< 0.03 Co-60	< 0.03 Co-60	< 0.03 Co-60	< 0.03 Co-60
C1-6 (I)	0.81 ± 0.08 Cs-137	1.5 ± 0.1 Cs-137	0.44 ± 0.04 Cs-137	1.0 ± 0.06 Cs-137
	< 0.05 Cs-134	< 0.04 Cs-134	< 0.03 Cs-134	< 0.07 Cs-134
	< 0.06 Co-60	< 0.05 Co-60	< 0.04 Co-60	< 0.04 Co-60
A1-4 (I)	0.034 ± 0.018 Cs-137	0.03 ± 0.014 Cs-137	0.25 ± 0.04 Cs-137	0.43 ± 0.03 Cs-137
	< 0.02 Cs-134	< 0.013 Cs-134	< 0.02 Cs-134	< 0.02 Cs-134
	< 0.03 Co-60	< 0.015 Co-60	< 0.04 Co-60	< 0.03 Co-60
Q1-2 (C)	0.043 ± 0.019 Cs-137	0.05 ± 0.01 Cs-137	< 0.017 Cs-137	< 0.03 Cs-137
	< 0.017 Cs-134	< 0.011 Cs-134	< 0.011 Cs-134	< 0.03 Cs-134
	< 0.019 Co-60	< 0.014 Co-60	< 0.013 Co-60	< 0.03 Co-60

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- 13. GEO Engineering "Phase I Report of Findings Groundwater Investigation", November 18, 1992
- 14. GEO Engineering "Summary of Field Work", June 7, 1994
- 15. Haley and Aldrich "Summary of Field Work", July 24, 1998

APPENDIX A

REMP Sampling Locations and Descriptions, Synopsis of REMP, and Exceptions in Sampling and Analysis

TABLE 11

Radiological Environmental Monitoring Program Description

Station Code	Sample Medium	Description	Comments
A1-1	Sediment	Drain Outfall Outside Perimeter Fence	Water rarely present
A1-2	Air Particulate	Westinghouse Yard Area	
A1-4	Surface Water Sediment	Juniata River at the Westinghouse Weir Bulkhead	
A1-5	TLD	N Sector, Perimeter Fence	
A1-6	Broadleaf Vegetation	N Sector, Outside Perimeter Fence	
B1-4	Surface Water Sediment	Drop Weir In The Westinghouse Yard Area	Weir was removed from the ground
B1-6	TLD	NNE Sector, Perimeter Fence	
B1-7	Broadleaf Vegetation	NE Sector, Outside Perimeter Fence	
C1-6	Sediment	Drain Outfall, NE Corner Of Perimeter Fence	Water rarely present
C1-9	TLD	NE Sector, Perimeter Fence	
C2-1	TLD	Weaver Ridge, 0.8 Mile from CV	
D1-1	Air Particulate	Open Field ENE Sector	
D1-4	TLD	ENE Sector, Perimeter Fence	
D2-1	TLD	Weaver Bridge, 1.3 Miles from CV	
E1-1	Potable Water	Penelec Line Shack	
E1-7	TLD	E Sector, Perimeter Fence	
E1-17	TLD	Penelec Line Shack	
E2-1	TLD	E Sector, 0.25 Mile from CV	
E3-1	TLD	3 Miles East of CV in State Game Land #67	
F1-2	TLD	ESE Sector, Perimeter Fence	
G1-1	TLD and Potable Water	SE Sector, Private Residence in Saxton (Putts Hollow Reservoir or Seton Water Supply Plant	

TABLE 11 (Continued)

Radiological Environmental Monitoring Program Description

Station Code	Sample Medium	Description	Comments
G1-2	TLD	SE Sector, Perimeter Fence	
G2-1	TLD	SE Sector, Closest Private Residence	
G10-1	Air Particulate	Reichley Microwave Tower	
G10-2	TLD	New Granada	
H1-5	TLD	SSE Sector, Perimeter Fence	
H2-1	TLD	Tussey Mountain High School	
H10-1	TLD	Wells Tannery	Offsite Control Station
J1-1	TLD	Penelec Fence, 100 Feet from SE Corner of West Garage	
J1-3	Air Particulate	Penelec Area S Sector	
K1-5	TLD	Saxton Borough Hall	
K1-8	TLD	SSW Sector, Perimeter Fence	
L1-1	TLD	SW Sector, Perimeter Fence	
L2-1	TLD	SW Sector, Stonerstown, 1 Mile From CV	
M1-6	TLD	WSW Sector, Perimeter Fence	
N1-4	TLD	W Sector, Perimeter Fence	
P1-1	TLD	WNW Sector, Perimeter Fence	
Q1-2	Surface Water Sediment	Old Station Discharge	Upstream (Control)
Q1-3	TLD	NW Sector, Perimeter Fence	
R1-1	TLD	NNW Sector, Perimeter Fence	
GEO 1	Groundwater	Monitoring Well, South of SNEC Property Fence	

TABLE 11 (Continued)

Radiological Environmental Monitoring Program Description

Station Code	Sample Medium	Description	Comments
GEO 2	Groundwater	Monitoring Well South of CV Fenced Area	2
GEO 3	Groundwater	Monitoring Well West of CV Fenced Area	
GEO 4	Groundwater	Monitoring Well West of CV Fenced Area	
GEO 5	Groundwater	Monitoring Well East of CV Fenced Area	
GEO 6	Groundwater	Monitoring Well North of CV Fenced Area	0
GEO 7	Groundwater	Monitoring Well East of CV Fenced Area	1
GEO 8	Groundwater	Monitoring Well North of GPU Energy Fence	
GEO 9	Groundwater	Piezometer Inside of CV Fenced Area	1
GEO 10	Groundwater	Monitoring Well NE of CV Fenced Area	
MW-1	Groundwater	NE to NW Diagonal Well	0
MW-2	Groundwater	NW to SW Diagonal Well	
MW-3	Groundwater	Monitoring Well East of CV Fenced Area	
MW-4	Groundwater	Monitoring Well East of CV Fenced Area	

TABLE 12

Synopsis of the 2000 Radiological Environmental Monitoring Program

Sample Type	Number of Sampling Locations	Collection Frequency	Number of Samples Collected	Type of Analysis	Analysis Frequency	Number of Samples Analyzed ⁽²⁾	
		<u>r i in denere</u>		Gross Beta	Weekly/Biweekly	404	
Air Particulate	4	Weekly or Biweekly	181	Gross Alpha	Weekly/Biweekly	181	
				Gamma	Quarterly	16	
Aquatic Sediment	4	Quarterly	16	Gamma	Quarterly	16	
Broad Leaf Vegetation	2	Annually	2	Gamma	Annually	2	
				H-3	Quarterly	41	
Groundwater	13	Quarterly	41	Gamma	Quarterly	41	
				Sr-90	Quarterly	5	
				H-3	Quarterly		
Potable Water	2	Quarterly	6	Gamma	Quarterly	6	
				Gross Beta	Quarterly		
Dosimeters (TLD) ⁽³⁾	28	Quarterly	896	Immersion Dose	Quarterly	896	
				Gamma	Quarterly	0	
Surface Water	2	Quarterly	8	H-3	Quarterly	8	

Notes:

(1) This table represents results from the primary (base) program. It does not include quality control (QC) results.

(2) The total number of analyses does not include duplicate analyses, recounts or reanalysis.

(3) For the purposes of this table, a dosimeter is considered to be a phosphor element.

(4) The total number of samples or elements (TLDs) used for data analysis.

(5) Biweekly means once every two weeks.

TABLE 13

Sampling and Analysis Exceptions 2000*

Period of Deviation	Description of Deviation and Corrective Action (as applicable)
4/27/00 to 5/04/00	Sample not obtained at GMW particulate air sample monitoring station D1-1 due to sampler malfunction.
1/13/00 to 2/24/00	Sample not obtained at GMW particulate air sample monitoring station J1-3 due to sampler malfunction.
4/06/00	Water sample was not obtained at well GEO-1 due to utility pole installation.
1/06/00 and 4/06/00	Water sample was not obtained at well GEO-6 because well was dry.
1/06/00	Water sample was not obtained at well MW-1 because well was dry.
1/06/00	Water sample was not obtained at G1-1 because Mrs. Weaver was not home.
7/13/00 and 10/11/00	Water samples were not obtained at wells MW-1, GEO-6, and GEO-7. These well were removed to support soil remediation of the CV Yard.
10/11/00	Water sample was not obtained at well GEO-8 because well was dry.
4 th Quarter	TLD element 3 of location G1-2 determined to be outlier. It was excluded from the data set. Investigation performed by vendor.
10/11/00	Water sample was not obtained at E1-1 because GPU Energy Line Departmer door was locked.

APPENDIX B

LOWER LIMIT OF DETECTION (LLD) EXCEPTIONS

TABLE 14

Analytical Results that Failed to Meet the SNEC REMP Required LLD During 2000

Sample Media	Analysis	Required LLD	Number of Samples that Failed to Meet the LLD	Comments
0	Ū	0	O	٥

 There were no analytical results that failed to meet the required Lower Limit of Detection (LLD). A list of the detection limits is found in Table 16.

APPENDIX C

REMP CHANGES

TABLE 15

2000 REMP Changes

Date of Change	Description of Changes to SNEC Procedure E900-ADM-4500.22 (Environmental Monitoring)
5/03/01	1. Added a definition of Lower Limit of Detection (LLD) to the procedure.
	2. Changed the responsibility for tracking samples offsite from "Environmental Affairs" to "Radiological or Environmental Controls Personnel".
	 Changed the responsibility for replicate analysis review from "Environmental Controls" to "SNEC Quality Assurance Officer or Environmental Controls Personnel".
	4. Added personnel instructions for deficiencies, which involve samples counted that do not meet the required LLD and limits that have been exceeded.
	 Added a note to Exhibit 1 explaining that monitoring wells Geo-6, Geo- 7, Geo-9 (piezometer), and MW-1 were removed from the ground in the month of May 2000. This change was necessary to support soi remediation of the CV Yard North.
	6. Updated the Site Monitoring Wells Map to reflect the changes.
	Revised the sensitivity and reporting levels to comply with limits stated in the ODCM.

APPENDIX D

ACTION LEVELS

Analytical results of environmental samples were routinely reviewed and evaluated by the REMP Coordinator and Quality Assurance Officer (QAO). The results were checked for LLD violations, anomalous values, USNRC reporting levels, main sample and quality control (QC) sample agreement (Appendix E), and action levels.

Established by SNEC, the action level is defined as that level of reactor-related radioactivity which when detected in environmental samples initiates an investigation and subsequent actions, as necessary. An action level is reached if either of the following two criteria is met:

- The radioactivity concentration at an indicator station reaches or exceeds those concentrations listed in Table 16.
- The radioactivity concentration at the indicator station reaches or exceeds 10 times the mean concentration for the control locations. (This criteria applies only to those media and analyses which are not listed in Table 16.)

Action levels for gamma exposure rates measured by TLDs have also been established. For TLDs, an action level is reached if any of the following three criteria is met:

- The exposure rate at an indicator station not on the owner controlled area fence exceeds three times the mean of the control stations.
- The exposure rate at an indicator station on the owner controlled area fence exceeds 0.185mR/Hr (50% of the 40 CFR 190 limit of 25 mR/yr adjusted by a 67 hour recreational factor).
- The exposure rate at an indicator station not on the owner controlled area fence exceeds either two times the previous quarterly result or two times the historical average for the station.

If an action level is reached, an investigation is initiated which consists of some or all of the following actions:

- Examine the collection sheets for an indication of any equipment malfunctions, collection or delivery errors.
- Examine the running tables (prior data) for trends.
- Review control station data.
- Review QC or duplicate sample data (if available).
- Recount and/or reanalyze the sample.
- Collect and analyze an additional sample.

The results of the investigation are then documented. As appropriate, site personnel are apprised of plant-related radioactivity that exceeds the SNEC action level. If it is concluded that the detected activity is related to SNEC operations and also exceeds the USNRC reporting limits as defined in the ODCM, a detailed report will be issued to the USNRC.

There were no sample results that equaled or exceeded action level violations in 2000.

TABLE 16

SNEC REMP Analytical Required Sensitivities (LLD) and Reporting Levels

Exposure/Pathways and/or Sample	Units	Analysis	Required LLD	Reporting Level
Air Particulate (AP)	pCi/m3	Gross Alpha	1.5 E-3	1.0 E-1
		Gross Beta	1.0 E-2	1.0
		Cs-134	5.0 E-2	1.0 E1
		Cs-137	6.0 E-2	2.0 E1
		Sr-90	1.0 E-2	1.0 E-1
Sediment/Soil (SD/S)	pCi/g (Dry)	Cs-134	1.5E-1	1.0
		Cs-137	1.8E-1	5.0
		Sr-90	5.0 E-2	5.0 E-1
Water (SW/GW)	pCi/L	Gross Alpha	5.0	1.0 E2
		Gross Beta	4.0	5.0 E1
		Tritium	2.0 E3	2.0 E4
		Co-60	1.5 E1	3.0 E2
		Cs-134	1.5 E1	3.0 E1
		Cs-137	1.8 E1	5.0 E1
		Sr-90	2.0	8.0
Vegetation	pCi/g (Wet)	Cs-134	6.0 E-2	1.0
		Cs-137	8.0 E-2	2.0
		Sr-90	1.0 E-2	1.0 E-1

APPENDIX E

QUALITY CONTROL PROGRAM

Basis for a Quality Assurance Program

A quality assurance (QA) program is an essential part of any radiological environmental monitoring program (REMP). It provides reasonable assurance that the results of radiation measurements are valid. To be effective, elements of quality assurance must be evident in all phases of the monitoring program. These include, but are not limited to, sample collection, preservation and shipment, receipt of samples by the analysis laboratory, preparation and analysis of samples and data review and reporting. An effective QA program will allow for the identification of deficiencies in all monitoring processes so that appropriate investigative and corrective actions can be implemented.

The USNRC published Regulatory Guide 4.15, "Quality Assurance for Radiological Monitoring Programs (Normal Operations) - Effluent Streams and the Environment", which defines an acceptable QA program (Reference 9). SNEC has adopted the guidance contained in Regulatory Guide 4.15. To meet the objectives of this position document, procedures and plans have been written and implemented.

In the laboratory, samples are typically analyzed one time. Therefore, laboratory personnel must be reasonably confident with the analytical results which are generated. One means of achieving confidence in the results is through the analysis of quality control (QC) samples.

Three types of QC samples are routinely analyzed as part of SNEC's QA Program. They include intralaboratory-split samples, crosscheck program samples, and interlaboratory split samples. A discussion of each QC sample type is provided below.

Intralaboratory Split Samples

Each laboratory is required to split at a minimum every twentieth sample (at least 5%) and perform an analysis (or analyses) on each portion. The samples which can not be split (e.g., air particulate filters) are counted twice. The results of the two analyses are then checked by staff scientists and/or the Quality Assurance Officer for agreement using the criteria defined in procedure E900-ADM-4500.22, "Environmental Monitoring". Agreement is considered acceptable if the value of the ratio fall within certain limits similar to those listed in USNRC Inspection Procedure 84750 "Radioactive Waste Treatment, Effluent and Environmental Monitoring" with minor adjustments to account for activity concentrations with large uncertainties. Non-agreement of the sample concentrations may result in recounting or reanalyzing the sample(s) in question.

There was one intralaboratory non-agreement during 2000. REMP sample SX-SD-Q1-2 was analyzed using a Gamma Spectroscopy System and the initial Cs-137 result (less than 0.02 pCi/g) did not agree with the duplicate result (0.043 ± 0.019 pCi/g). It was determined that if the net or actual concentration of the original result (0.024 ± 0.021 pCi/g) were used, the results were not statistically different (i.e. the results with their counting uncertainties overlapped). No follow up action was required. The positive result of 0.043 ± 0.019 pCi/g was used as the record data.

Cross-check Program Samples

Each laboratory analyzing environmental samples for the SNEC Facility participates in at least two separate crosscheck programs. USDOE, USEPA, and Analytics supply either water, air particulates, vegetation and soil samples. All samples are sent to the laboratories as unknowns. Participation in these programs provides an independent check on the ability of each laboratory to perform analyses on various kinds of samples containing detectable concentrations of radioactivity. The results submitted by the laboratories are compared to: 1) limits established by the USEPA, or 2) agreement criteria used by the NRC in their Configuratory Measurement Inspection Program. If the results are outside the established limits or agreement criteria, the laboratories are requested to perform an investigation and take corrective action as necessary.

The 2000 crosscheck program results from each laboratory are listed in Appendix F. Explanations are provided for those results which were not submitted and/or which were not within the established limits.

Interlaboratory Split Samples

The third type of QC sample is the interlaboratory split sample. These samples are routinely collected for the REMP. After or during the collection process, the sample is thoroughly mixed (as necessary) to ensure that, as much as possible, the distribution of radioactivity in the sample is homogeneous. The sample is then split into two portions. One portion is sent to the primary (main) laboratory and the other portion is sent to the QC laboratory.

Analysis results from the QC laboratory are then compared to those from the primary laboratory. The agreement criteria are the same as that used for the intralaboratory-split samples. Corrective action for disagreements may include recounting or reanalyzing the sample(s). There were no interlaboratory non-agreements during the year 2000.

APPENDIX F

CROSS-CHECK PROGRAM RESULTS

TABLE 17

			EI	RL .	DC	E EML				
Collection Date	Media	Nuclide	Value (a) & (d)	Uncertainty (a)	Value (b) & (d)	Uncertainty (c)	Ratio	Minimum Ratio	Maximum Ratio	Agreement
		Am-241	0.15	0.02	0.088	0.005	1.705	0.73	2.58	Yes
		Co-57	5.9	0.6	5.31	0.22	1.111	0.65	1.39	Yes
		Co-60	6.4	0.6	5.32	0.26	1.203	0.75	1.32	Yes
		Cs-137	7.1	0.7	6.1	0.3	1.164	0.73	1.37	Yes
		Mn-54	31	3	27.2	0.8	1.140	0.73	2.58	Yes
3/1/2000	Air Filter	Pu-238	0.079	0.013	0.08	0.001	0.988	0.74	1.4	Yes
		Pu-239	0.1	0.01	0.089	0.003	1.124	0.76	1.44	Yes
		Ru-106	3.3	1.7	2.01	1.94	1.642	0.59	1.3	No
		U-234	0.055	0.009	0.062	0.001	0.887	0.83	1.92	Yes
		U-238	0.051	0.009	0.062	0.001	0.823	0.84	2.61	No
		U-NAT	0.11		0.126	0.001	0.873	0.8	2.61	Yes
	A	Alpha	2.7	0.3	3.02	0.3	0.894	0.5	1.55	Yes
3/1/2000	Air Filter	Beta	2.8	0.3	2.42	0.2	1.157	0.72	1.67	Yes
	1	Am-241	13	3	3.36	0.51	3.869	0.63	2.31	No
		Cs-137	393	40	339	9.3	1.159	0.83	1.32	Yes
		K-40	943	97	811	29	1.163	0.78	1.53	Yes
		Pu-238	20	2	18.6	0.5	1.075	0.52	2.84	Yes
3/1/2000	Soil	Pu-239	7.7	1.3	7	0.34	1.100	0.69	1.74	Yes
		Sr-90	18	5	20.2	0.2	0.891	0.6	3.66	Yes
		U-234	105	10	111	11	0.946	0.47	1.3	Yes
		U-238	105	10	114	12	0.921	0.44	1.42	Yes
		U-NAT	214		229	23	0.934	0.42	1.3	Yes
		Am-241	16	2	10.4	1.4	1.538	0.68	2.7	Yes
		Cm-244	10	1	5	1.8	2.000	0.47	1.74	No
		Co-60	55	6	52.8	1	1.042	0.69	1.46	Yes
		Cs-137	1400	100	1380	20	1.014	0.8	1.4	Yes
3/1/2000	Vegetation	K-40	550	60	521	20	1.056	0.79	1.42	Yes
		Pu-238	1.7	0.2	1.09	0.1	1.560	0.66	7.94	Yes
		Pu-239	16	2	15.5	2.1	1.032	0.68	1.59	Yes
		Sr-90	2200	200	1780	17.8	1.236	0.5	1.33	Yes

AmerGen ERL & DOE EML Cross Check Program Results for 2000

TABLE 17 (Continued)

AmerGen ERL & DOE EML Cross Check Program Results for 2000

			ER	L	DOE	EML			A days	
Collection Date	Media	Nuclide	Value (a) & (d)	Uncertainty (a)	Value (b) & (d)	Uncertainty (c)	Ratio	Min. Ratio	Max Ratio	Agreement
		Am-241	3.8	0.4	1.95	0.18	1.949	0.75	1.49	No
		Co-60	53	5	48.9	1.8	1.084	0.8	1.2	Yes
		Cs-137	110	10	103	4	1.068	0.8	1.26	Yes
		Fe-55	31	5	33.1	0.7	0.937	0.44	1.53	Yes
		H-3	81	7	79.4	2.5	1.020	0.71	1.79	Yes
3/1/2000	Water	Pu-238	1.2	0.1	0.944	0.04	1.271	0.78	1.25	No
		Pu-239	1.2	0.1	0.918	0.03	1.307	0.8	1.39	Yes
		Sr-90	3.1	0.8	3.39	0.12	0.914	0.75	1.5	Yes
		U-234	0.53	0.06	0.482	0.04	1.100	0.8	1.4	Yes
		U-238	0.51	0.05	0.492	0.04	1.037	0.8	1.26	Yes
		U-NAT	1.1		0.995	0.087	1.106	0.67	1.42	Yes
		Alpha	1600	100	1700	170	0.941	0.61	1.32	Yes
3/1/2000	Water	Beta	940	100	690	70	1.362	0.55	1.54	Yes

Table 17 - Notes:

(a) The AmerGen ERL Value is an average of 1 to 4 determinations.

(b) The DOE EML value is the mean of replicate determinations for each nuclide.

(c) The DOE EML uncertainty is the standard error of the mean.

(d) Units are Bg/L for Water, Bg/kg (dry) for Soil, Bg/kg (wet) for Vegetation and total Bg for Air Filter.

(e) This sample was analyzed three times for Ru-106 and the average (3.3 +/- 1.7 Bq/un) was reported. The individual results were as follows:

3.5 +/- 2.0 Bq/un

3.0 +/- 1.4

3.4 +/- 1.7

The EML value was 2.01 +/- 1.94. The ERL/EML ratio was 1.642 and was not acceptable. The acceptance range was between 0.59 and 1.3.No follow-up actions were requested because the concentrations were not statistically different (i.e. the results with their counting uncertainties overlapped.).

(f) An investigation was conducted to determine why five out of twenty-two TRU radionuclides failed to achieve acceptable results from various media submitted by the EML crosscheck program. When processing EML crosscheck samples, separate glassware is used in order to avoid cross contaminating other client's samples. This glassware has become etched throughout the years due to acid digestions and flouric acid precipitations. What used to be very good test results from the various crosscheck programs has progressively depreciated. It has been determined that the etching causes fluctuations in radionuclide recoveries. This is true not only with TRU radionuclides but also other fission and activation product nuclides. The ERL is now disposing glassware that shows deterioration to prevent this problem from reoccurring. Also, when it is appropriate, plastic beakers are being substituted for glass. Both corrective actions should eliminate the problems that result from etched glassware.

The control limit concept was established from percentiles of historic data distributions (1982 - 1992). The evaluation of this historic data and the development of the control limits are presented in DOE report EML-564. The control limits for QAP-XLVIII were developed from percentiles of data distributions for the years 1993 - 1999.

TABLE 18

AmerGen ERL & Analytics Cross Check Program Results for 2000

				· · · · · · · · · · · ·	Analytics						
Collection Date	Media	Nuclide	ERL Value	Value	Uncei	rtainty	Resolution	Ratio	Min. Ratio	Max. Ratio	Agreement
Date			(b)	(a)	(3 Sigma)	(1 Sigma)					
		I-131	92	84	4	1.3	63.0	1.10	0.8	1.25	Yes
6/22/2000	Milk	I-131	89	84	4	1.3	63.0	1.06	0.8	1.25	Yes
6/22/2000	Cartridge	I-131	66	72	4	1.3	54.0	0.92	0.8	1.25	Yes

Table 18 - Notes:

a. The Analytics Value is the known concentration. Units are pCi/L for Milk, pCi/g (dry) for Soil and total pCi for Filter and Cartridge.

b. The AmerGen ERL Value is an average of three or more determinations. Units are pCi/L for Milk, pCi/g (dry) for Soil and total pCi for Filter and Cartridge.

To determine agreement or possible agreement:

- 1. Divide each Analytics value by its associated one sigma uncertainty to obtain the resolution.
- 2. Divide each ERL value by the corresponding Analytics value to obtain the ratio.
- 3. The ERL measurement is in agreement if the value of the ratio falls within the limits shown in the following table for the corresponding resolution.

Resolution	<u>Agreement</u>
< 4	0.4 - 2.5
4 - 7	0.5 - 2.0
8 - 15	0.6 - 1.66
16 - 50	0.75 - 1.33
51 - 200	0.80 - 1.25
> 200	0.85 - 1.18

Criteria are similar to those listed in USNRC Inspection Procedure 84750 "Radioactive Waste Treatment, and Effluent and Environmental Monitoring" with minor adjustments to account for activity concentrations with large uncertainties.

TABLE 19

Teledyne Brown Engineering & Analytics Cross Check Program Results for 2000

Collection Date	Media	Nuclide	TBE Value	Analytics Value (a)	2 Sigma Uncertainty	Resolution	Ratio (b)	Minimum Ratio	Maximum Ratio	Agreement
		I-131	18	20	1	36	0.9	0.75	1.33	Yes
		Cr-51	381	387	38	20	0.98	0.75	1.33	Yes
		Cs-134	132	143	13	20	0.92	0.75	1.33	Yes
		Cs-137	128	114	13	20	1.12	0.75	1.33	Yes
6/20/00	Milk	Co-58	89	79	9	20	1.13	0.75	1.33	Yes
		Mn-54	195	176	20	20	1.11	0.75	1.33	Yes
		Fe-59	161	144	16	20	1.12	0.75	1.33	Yes
		Zn-65	171	165	17	20	1.04	0.75	1.33	Yes
		Co-60	179	176	18	20	1.02	0.75	1.33	Yes
		Sr-89	13	25	3	9	0.52	0.6	1.66	No
3/20/00	Milk	Sr-90	16	19	1	32	0.84	0.75	1.33	Yes
		Ce-141	143	132	8	36	1.08	0.75	1.33	Yes
		Cr-51	229	198	17	27	1.16	0.75	1.33	Yes
6/19/00 Air Filter		Cs-134	74	81	4	37	0.91	0.75	1.33	Yes
		Cs-137	143	115	8	36	1.24	0.75	1.33	Yes
	Air Filter	Co-58	89	77	5	36	1.16	0.75	1.33	Yes
0,10,00		Mn-54	102	84	6	34	1.21	0.75	1.33	Yes
		Fe-59	98	75	6	33	1.31	0.75	1.33	Yes
		Zn-65	188	139	11	34	1.35	0.75	1.33	No
		Co-60	113	104	7	32	1.09	0.75	1.33	Yes
6/19/00	Cartridge	I-131	106	88	6	35	1.2	0.75	1.33	Yes
6/19/00	Air Filter	Sr-90	88	96	5	35	0.92	0.75	1.33	Yes
0/10/00	7 11 7 1100	Gr-A	103	93	6	34	1.11	0.75	1.33	Yes
6/19/00	Air Filter	Gr-B	210	193	6	70	1.09	0.8	1.25	Yes
		I-131	97	87	10	19	1.11	0.75	1.33	Yes
		Ce-141	83	77	8	21	1.08	0.75	1.33	Yes
		Cr-51	323	304	40	16	1.06	0.75	1.33	Yes
		Cs-134	98	102	10	20	0.96	0.75	1.33	Yes
		Cs-137	117	107	12	20	1.09	0.75	1.33	Yes
9/18/00	Milk	Co-58	64	60	6	21	1.07	0.75	1.33	Yes
		Mn-54	99	88	10	20	1.13	0.75	1.33	Yes
		Fe-59	132	119	13	20	1.11	0.75	1.33	Yes
		Zn-65	218	196	22	20	1.11	0.75	1.33	Yes
		Co-60	209	198	22	20	1.06	0.75	1.33	Yes
		Sr-89	14	15	1	28	0.93	0.75	1.33	Yes
9/18/00	Milk	Sr-89 Sr-90	14	15	1	36	1.29	0.75	1.33	Yes

Table 19 - Notes:

- (a) The Analytics Value is the known concentration. Units are in pCi/L for Milk, pCi/g (dry) for Soil and total pCi for Filter and Cartridge.
- (b) Ratio of Teledyne Brown Engineering to Analytics results.
- (c) Under investigation.
- (d) Caused by incorrect rinsing of the strontium extraction column. Additional training was conducted and was documented in the analyst's training file. Subsequent tests on two milk samples spiked with Sr-89 produced correct results.

To determine agreement or possible agreement:

- 1. Divide each Analytics value by its associated one sigma uncertainty to obtain the resolution.
- 2. Divide each TBE value by the corresponding Analytics value to obtain the ratio.
- 3. The measurement is in agreement if the value of the ratio falls within the limits shown in the following table for the corresponding resolution.

Resolution	Agreement
< 4	0.4 - 2.5
4 - 7	0.5 - 2.0
8 - 15	0.6 - 1.66
16 - 50	0.75 - 1.33
51 - 200	0.80 - 1.25
> 200	0.85 - 1.18

Criteria are similar to those listed in USNRC Inspection Procedure 84750 "Radioactive Waste Treatment, and Effluent and Environmental Monitoring" with minor adjustments to account for activity concentrations with large uncertainties.

TABLE 20

Teledyne Brown Engineering & Environmental Resource Assessment (ERA) Statistical Summary

2000 Proficiency Testing (Pt) Program

ERA No.	Date	Media	Nuclide	ERA Known Value (pCi/L) (a)	TBE Result (pCi/L) (b)	Expected Dev. Known (pCi/L) (c)	Control Limits (pCi/L) (d)	Performance Evaluation (e)	
			U-Nat	53.0	61.3	5.3	44.0 - 62.0	CE	
			Ra-226	4.05	3.67	0.608	3.0 - 5.1	А	
			Ra-228	2.29	1.33	0.573	1.31 - 3.27	CE	
			Gr-A	71.8	14.0	18.0	40.9 - 103	NA	
			Gr-B	194	34.0	29.1	144 - 244	NA	
Rad 13	2/26/00		Sr-89	16.4	15.7	5.0	7.7 - 25.1	А	
			Sr-90	28.9	29.0	5.0	20.2 - 37.6	А	
			Co-60	64.4	68.3	5.0	55.7 - 73.1	А	
			Cs-134	12.3	12.0	5.0	3.6 - 21.0	А	
			Cs-137	72.2	76.3	5.0	63.5 - 80.9	А	
			Gr-A	25.4	14.0	6.35	14.5 - 36.3	NA	
Rad 14	2/24/00		Gr-B	42.1	34.0	5.0	33.4 - 50.8	CE	
			Ba-133	98.2	91.7	9.82	81.5 - 115	A	
				Co-60	99.6	101	5.0	90.9 - 108	А
Rad 15	2/25/00		Cs-134	49.2	48.0	5.0	40.5 - 57.9	А	
			Cs-137	209	76.3	10.4	191 - 227	NA	
			Zn-65	313	<1.0	31.3	260 - 367	NA	
			Sr-89	22.5	18.3	5.0	13.8 - 31.2	А	
Rad 16	5/18/00		Sr-90	9.6	8.33	5.0	0.9 - 18.3	А	
			Gr-A	58.4	83.6	14.6	33.3 - 83.5	NA	
Rad 17	2/10/00		Gr-B	16.8	15.4	5.0	8.1 - 25.5	А	
Rad 18	5/23/00		I-131	19.9	2.03	3.00	14.7 - 25.1	NA	
			U-Nat	6.07	5.77	3.0	0.87 - 11.3	A	
Rad 19	2/24/00		Ra-226	8.26	7.2	1.24	6.11 - 10.4	А	
		1	Ra-228	2.25	2.37	0.56	1.28 - 3.22	А	
Rad 20	3/01/00		H-3	23800	22300	12380	21100 - 26500	A	

TABLE 20 (Continued)

Teledyne Brown Engineering & Environmental Resource Assessment (ERA) Statistical Summary

2000 Proficiency Testing (Pt) Program

ERA No.	Media	Nuclide	ERA Known Value (pCi/L) (a)	TBE Result (pCi/L) (b)	Expected Dev. Known (pCi/L) (c)	Control Limits (pCi/L) (d)	Performance Evaluation (e)
<u></u>		Ra-226	13.0	9.7	1.15	7.41 - 18.6	A
		U-Nat	63.4	57.0	4.44	52.6 - 74.2	A
Rad 23	Water	Ra-228	2.83	2.99	6.34	2.21 - 3.77	A
		Ra-228	13.0	10.0	3.25	7.41 - 16.8	A
Rad 24	Water	Sr-90	26.2	28.6	1.4	17.5 - 34.9	A
Rad 25	Water	Gr-A	7.17	6.9	1.11	DL - 15.9	A
Rad 25	Water	Gr-B	87.5	88.8	9.76	70.2 - 105	A
Rad 26	Water	H-3	8320	8740	174	6910 - 9730	A

Table 20 - Notes:

- (a) The ERA Known Value is equal to 100% of the parameter present in the standard as determined by gravimetric and/or volumetric measurements made during standard preparation.
- (b) Average result.
- (c) Established per the guidelines contained in the EPA's National Standards for Water Proficiency Testing Criteria Document, December 1998, as applicable.
- (d) Established per the guidelines contained in the EPA's National Standards for Water Proficiency Testing Criteria Document, December 1998, as applicable.
- (e) A = Acceptable. Reported Result falls within the Warning Limits.
 NA = Not Acceptable. Reported Result falls outside of the Control Limits.
 CE = Check for Error. Reported Result falls within the Control Limits and outside of the Warning Limits.
- (f) RAD 23 through RAD 26 were received and analyzed in the Westwood, New Jersey laboratory in September 2000.

TABLE 21

B&W Nuclear Environmental Services & DOE EML Cross Check Program Results for 2000

Sample ID No.	Media	Nuclide	Units	Reported Value	EML Value	Reported/EML Ratio	Performance Evaluation
		Mn-54		47.0 <u>+</u> 3.7	43.2 <u>+</u> 1.3	1.088	А
		Co-57	÷	14.7 <u>+</u> 0.4	14.5 <u>+</u> 0.46	1.014	Α
		Co-60		9.06 <u>+</u> 0.51	8.43 <u>+</u> 0.48	1.075	А
		Sr-90		1.35 <u>+</u> 0.16	1.64 <u>+</u> 0.11	0.823	А
		Ru-106		9.06 <u>+</u> 1.6	(a)	(a)	(a)
2009 AIBC	Air Sample	Cs-137	Bq/Filter	8.07 <u>+</u> 0.38	7.41 <u>+</u> 0.36	1.089	А
		U-234		0.052 <u>+</u> 0.006	0.04 <u>+</u> 0.003	1.3	А
		U-238		0.051 <u>+</u> 0.005	0.04 <u>+</u> 0.002	1.275	W
		Gr-A		1.99 <u>+</u> 0.07	2.35 <u>+</u> 0.23	0.847	А
		Gr-B	-	1.24 <u>+</u> 0.04	1.52 <u>+</u> 0.15	0.816	W
		K-40	873 <u>+</u> 96	713.0 <u>+</u> 38.0	1.224	A	
		Cs-137	Datifica	1220.0 <u>+</u> 120.0	1020 <u>+</u> 51.0	1.196	W
2009 SOBC	Soil Sample	U-234	Bq/Kg	138.0 <u>+</u> 12.0	157.0 <u>+</u> 10.0	0.879	А
		U-238		136.0 <u>+</u> 12.0	163.0 <u>+</u> 10.0	(a) 1.089 1.3 1.275 0.847 0.816 1.224 1.196 0.879 0.834 1.083 1.195 0.911 1.165 0.989 0.781 0.972 1.089 1.171	А
		K-40	Bq/Kg	692.0 <u>+</u> 75	639.0 <u>+</u> 34.0	1.083	А
	Vegetation Sample	Co-60		39.2 <u>+</u> 4.5	32.8 <u>+</u> 1.3	1.195	А
2009 VEBC		Sr-90		1050.0 <u>+</u> 40.0	1152.0 <u>+</u> 94.0	0.911	А
		Cs-137		1010.0 <u>+</u> 100.0	867.0 <u>+</u> 44.0	1.165	А
	· · · · · · · · · · · · · · · · · · ·	Co-60		72.9 <u>+</u> 3.2	73.7 <u>+</u> 2.9	0.989	А
		Sr-90		3.54 <u>+</u> 0.3	4.53 <u>+</u> 0.12	0.781	W
		Cs-137		65.1 <u>+</u> 2.2	67.0 <u>+</u> 3.5	0.972	А
2009 WABC	Water Sample	U-234	Bk/Kg	0.524 <u>+</u> 0.045	0.481 <u>+</u> 0.023	1.089	А
		U-238		0.431 <u>+</u> 0.04	0.368 <u>+</u> 0.012	1.171	А
		Gr-A		1280.0 <u>+</u> 60.0	1070.0 <u>+</u> 100.0	1.196	W
		Gr-B		910.0 <u>+</u> 36.0	950.0 <u>+</u> 90.0	0.958	А
		Mn-54	1	47.7 <u>+</u> 3.7	43.2 <u>+</u> 1.3	1.104	А
2009 AIBX	Air Sample	Co-57	Bq/Filter	15.0 <u>+</u> 0.4	14.5 <u>+</u> 0.46	1.034	А
	•	Co-60		9.25 <u>+</u> 0.53	8.43 <u>+</u> 0.48	1.097	А

TABLE 21 (Continued)

B&W Nuclear Environmental Services & DOE EML Cross Check Program Results for 2000

Sample ID No.	Media	Nuclide	Units	Reported Value	EML Value	Reported/EML Ratio	Performance Evaluation
		Sr-90		1.23 <u>+</u> 0.15	1.64 <u>+</u> 0.11	0.75	W
		Ru-106		9.66 <u>+</u> 1.52	(a)	(a)	(a)
		Cs-137		7.96 <u>+</u> 0.37	7.41 <u>+</u> 0.36	1.074	А
		U-234		0.055 <u>+</u> 0.005	0.04 <u>+</u> 0.003	1.375	W
		Pu-238		0.066 <u>+</u> 0.012	0.045 <u>+</u> 0.0003	1.467	N (b)
2009 AIBX	Air Sample	U-238	Bq/Filter	0.053 <u>+</u> 0.005	0.04 <u>+</u> 0.002	1.325	W
		Pu-239		0.078 <u>+</u> 0.007	0.074 <u>+</u> 0.007	1.054	А
		Am-241		0.058 <u>+</u> 0.015	0.032 <u>+</u> 0.001	1.813	W
		Gr-A		1.83 <u>+</u> 0.07	2.35 <u>+</u> 0.23	0.779	W
		Gr-B		1.21 <u>+</u> 0.04	1.52 <u>+</u> 0.15	0.796	W
		K-40		821.0 <u>+</u> 86.0	713 <u>+</u> 38.0	1.151	A
		Cs-137	Bq/Kg	1220.0 <u>+</u> 120.0	1020.0 <u>+</u> 51.0	1.196	W
		Bi-212		50.3 <u>+</u> 9.9	80.5 <u>+</u> 6.6	0.625	А
		Pb-212		108.0 <u>+</u> 10.0	79.3 <u>+</u> 4.3	1.362	N (b)
	Soil Sample	Bi-214		92.5 <u>+</u> 10.5	83.3 <u>+</u> 4.2	1.110	А
		Pb-214		111.0 <u>+</u> 12.0	86.3 <u>+</u> 4.3	1.286	А
2009 SOBX		Ac-228		83.2 <u>+</u> 11.1	80.2 <u>+</u> 3.6	1.037	А
		Th-234		65.9 <u>+</u> 79.7	148.0 <u>+</u> 10.0	0.445	N (b)
		U-234		128.0 <u>+</u> 11.0	157.0 <u>+</u> 10.0	0.815	W
		U-238		141.0 <u>+</u> 12.0	163.0 <u>+</u> 10.0	0.865	А
		Pu-239		16.1 <u>+</u> 1.9	16.8 <u>+</u> 0.3	0.958	А
		Am-241		2.48 <u>+</u> 0.96	8.27 <u>+</u> 0.69	0.3	N (b)
		K-40		755.0 <u>+</u> 77.0	639.0 <u>+</u> 34.0	1.182	Α
		Co-60		36.5 <u>+</u> 4.0	32.8 <u>+</u> 1.3	1.113	А
		Sr-90		1000.0 <u>+</u> 40.0	1152.0 <u>+</u> 94.0	0.868	А
2009 VEBX	Vegetation Sample	Cs-137	Bq/Kg	1010.0 <u>+</u> 100.0	867.0 <u>+</u> 44.0	1.165	А
		Pu-239	_	8.44 <u>+</u> 1.11	9.6 <u>+</u> 0.8	0.879	A
		Am-241		7.29 <u>+</u> 1.26	5.6 <u>+</u> 0.67	1.302	А
		Cm-244			3.6 <u>+</u> 0.27	1.419	W

TABLE 21 (Continued)

B&W Nuclear Environmental Services & DOE EML Cross Check Program Results for 2000

Sample ID No.	Media	Nuclide	Units	Reported Value	EML Value	Reported/EML Ratio	Performance Evaluation
		H-3		129.0 <u>+</u> 25.0	91.3 <u>+</u> 0.3	1.413	W
1		Co-60		75.1 <u>+</u> 3.4	73.7 <u>+</u> 2.9	1.019	А
		Sr-90		3.7 <u>+</u> 0.32	4.53 <u>+</u> 0.12	0.817	w
		Cs-137		66.6 <u>+</u> 2.2	67.0 <u>+</u> 3.5	0.994	А
		U-234		0.487 <u>+</u> 0.047	0.481 <u>+</u> 0.023	1.012	А
2009 WABX	Water Sample	Pu-238	Bq/Liter	0.725 <u>+</u> 0.056	0.786 <u>+</u> 0.011	0.922	А
		U-238		0.385 <u>+</u> 0.041	0.368 <u>+</u> 0.012	1.046	А
		Pu-239		0.58 <u>+</u> 0.048	0.591 <u>+</u> 0.021	0.981	А
		Am-241		1.23 <u>+</u> 0.14	1.192 <u>+</u> 0.045	1.032	А
		Gr-A	1	1190.0 <u>+</u> 50.0	1070.0 <u>+</u> 100.0	1.112	А
		Gr-B		899.0 <u>+</u> 36.0	950.0 <u>+</u> 90.0	0.946	A

Table 21 - Notes:

- (b) Presently under investigation.
- (c) A = Acceptable. Reported result falls within the limits.
 N = Not Acceptable. Reported result falls outside of the control limits.
 W = Warning Check for error.

⁽a) EML activity not available at this time.

APPENDIX G

DATA REPORTING AND ANALYSIS

Environmental samples frequently contain very little, if any, radioactivity. Even when very sensitive, state-of-the-art counting equipment is used, many of the sample count rates can not be differentiated from the background count rate or the count rate of the blank sample. When this occurs, the sample is said to have a radioactivity level or concentration at or below the sensitivity of the analysis method. In this case, the analysis result is reported as less than a numerical value, which corresponds to the sensitivity of the analysis method. Sensitivities are influenced by parameters such as sample volume, background or blank sample count rate and efficiency of the counting device.

The terms used to describe the sensitivity are the lower limit of detection (LLD) and minimum detectable concentration (MDC). For this report, these two terms are considered to be synonymous. They are defined as:

LLD (MDC) =
$$\frac{4.66 \text{ Sb}}{\text{E * V * 2.22 * Y * exp}^{-(\lambda \Delta t)}}$$

Where:

.		
Sb	=	the standard deviation of the background counting rate or the counting rate of a blank sample, as counts per minute,
Е	=	the counting efficiency of the equipment, as counts per disintegration,
V	=	the volume or mass of the sample, such as L, g or m ³ ,
2.22	=	the number of disintegrations per minute per picocurie,
Y	=	the chemical yield, if applicable,
λ	=	the radioactive decay constant for the particular radionuclide and
Δt	Ξ	the elapsed time between sample collection (or end of sample collection
		period) and counting.

The applicable LLD or MDC for each radionuclide and analysis is listed in Table 16. A large percentage of the 2000 sample results were reported as less than the LLD or MDC. Results which were reported as less than the LLD or MDC were not included in the calculations of averages, standard deviations and ranges (by station or group) in the text and tables of this report.

The data from samples that contained concentrations above the LLD or MDC were used in the calculations (averages, standard deviations and ranges) contained in this report. The individual results were generally reported to two significant figures. Each result also included a two-sigma counting uncertainty (95% confidence interval) to the same decimal place. At a minimum, a counting uncertainty equal to 10 percent of the measured concentration was reported. The counting uncertainties were not used in any statistical calculations in this report.

The data used in a few tables and all annual graphs were actual sample concentrations. The actual concentration is calculated by subtracting the background count rate or the count rate of a blank sample from the count rate of the sample. The net count rate is then converted to a net sample concentration which is either positive, negative or zero.

There are several advantages of using actual sample concentrations. Biases in the data (averages, ranges, etc.), such as those caused by averaging only sample concentrations above the MDC, are eliminated. Missing data points on graphs also are eliminated. It should be noted that negative sample concentrations are important to the overall averages and trends in the data, but they have no physical significance. A negative sample concentration simply means that the background or blank sample count rate is greater than the sample.

Quality control results (interlaboratory and intralaboratory) were not statistically analyzed with other data. Including quality control data would introduce a bias at selected stations while providing little additional interpretive information.

APPENDIX H

REMP SAMPLE COLLECTION AND ANALYSIS METHODS

TABLE 22

SNEC Radiological Environmental Monitoring Program Summary of Sample Collection and Analysis Methods 2000

Analysis	Sample* Medium	Sampling Method	Approximate Sample Size Analyzed	Analysis Procedure Number	Procedure Abstract
Gr-Alpha	AP	Continuous weekly or biweekly air sampling through filter paper	1 filter (500-1000 Cubic Meters)	6510-IMP-4592.05 (AmerGen) BWXT-TP-316	Low background gas flow proportional counting
Gr-Beta	AP	Continuous weekly or biweekly air sampling through filter paper	1 filter (500-1000 Cubic Meters)	6510-IMP-4592.05 (AmerGen) BWXT-TP-316	Low background gas flow proportional counting
	GW	Quarterly grab sample	500 ml	6510-IMP-4592.01 (AmerGen) TBE-Westwood PRO-032-41	Sample evaporated on stainless steel planchet for low background gas flow proportional counting
Gamma Spectroscopy	AP	Quarterly composite of filter paper collected weekly or biweekly.	6 to 15 filters (6,900 - 9,300 Cubic Meters)	6510-IMP-4592.05 (AmerGen) 6510-OPS-4591.04 (AmerGen) BWXT-TP-398	Sample placed in counting container for gamma isotopic analysis
	BR	Annual grab sample	1 kg	6510-IMP-4592.03 (AmerGen) 6510-OPS-4591.04 (AmerGen) BWXT-TP-398	Edible portion placed in counting container for gamma isotopic analysis. Only root vegetables and fruits washed prior to analysis.
	GW	Quarterly grab sample which are collected with a gas displacement gromon system or from a faucet.	3.5 liters (if possible)	6510-IMP-4592.06 (AmerGen) 6510-OPS-4591.04 (AmerGen) BWXT-TP-398	Sample decanted and liquid portion placed in counting container for gamma isotopic analysis. Potable samples are mixed (not decanted) prior to analysis
			1 liter	TBE-Westwood Pro-042-5	Same as above

TABLE 22 (Continued)

SNEC Radiological Environmental Monitoring Program Summary of Sample Collection and Analysis Methods 2000

Analysis	Sample* Medium	Sampling Method	Approximate Sample Size Analyzed	Analysis Procedure Number	Procedure Abstract
Gamma Spectroscopy (Cont'd)	SD	Quarterly grab sample	1 kg (if possible)	6510-IMP-4592.04 (AmerGen) 6510-OPS-4591.04 (AmerGen) BWXT-TP-398	Dried and sieved sample placed in counting container for gamma isotopic analysis.
	SW	Quarterly grab samples	3.5 liters	6510-IMP-4592.06 (AmerGen) 6510-OPS-4591.04 (AmerGen) BWXT-TP-398	Sample placed in counting container for gamma isotopic analysis.
Tritium	GW	Quarterly grab sample	7-10 ml	6510-IMP-4592.02 (AmerGen) 6510-OPS-4591.05 (AmerGen) 6510-OPS-4591.08 (AmerGen) BWXT-TP-642	Sample is filtered, mixed with scintillation fluid for scintillation counting
	SW	Quarterly grab samples	2ml or 10 ml	TBE-Westwood PRO-052-2 PRO052-35	Same as above
			7-10 ml	6510-IMP-4592.02 (AmerGen) 6510-OPS-4591.05 (AmerGen) 6510-OPS-4591.08 (AmerGen) TBE-Westwood PRO-052-35	Sample filtered, mixed with scintillation fluid for scintillation counting. Distillation may be performed if impurities are found to be present.
Gamma (Direct Radiation)	ID	Dosimeters exchanged quarterly	2 TLDs/8 Elements	6610-OPS-4243.01 (AmerGen)	Thermoluminescent dosimetry using optical heating of crystals and PM tube for light measurement.
			1 TLD/4 Elements	TBE-Westwood PRO-342-17	Same as above
Sr-89, Sr-90	GW	Quarterly grab sample	1 liter	BWXT-TP-692 TBE-Westwood-PRO-032-128	

TABLE 22 (Continued)

SNEC Radiological Environmental Monitoring Program Summary of Sample Collection and Analysis Methods 2000

NOTES

IDENTIFICATION KEY	APPROXIMATE SAMPLE SIZE COLLECTED PER STATION
AP = Air Particulate	1 Filter is approximately 500 cubic meters per week or biweekly
BR = Broad Leaf Vegetation	1 kg annually
GW = Ground Water	4 liters (if available) quarterly
ID = Immersion Dose (TLD)	2 TLDs / 8 elements quarterly
SD = Aquatic Sediment	1 kg semiannually
SW = Surface/Drinking Water	4 liters

APPENDIX I

TLD QUARTERLY DATA

TABLE 23

2000 TLD Quarterly Data mR/Std Month ± 2_{σ}

STATION	1 ST QUARTER	2 ND QUARTER	3 RD QUARTER	4 TH QUARTER
A1-5	5.6 ± 0.65	5.9 ± 0.3	5.9 ± 0.9	5.5 ± 0.5
B1-6	6.1 ± 0.57	6.0 ± 0.3	6.1 ± 0.9	5.7 ± 0.6
C1-9	5.6 ± 0.81	5.6 ± 0.3	5.5 ± 0.5	5.2 ± 0.5
C2-1	5.9 ± 0.32	6.2 ± 0.3	5.9 ± 0.8	6.1 ± 1.3
D1-4	6.2 ± 0.94	6.5 ± 0.2	6.3 ± 1.0	6.2 ± 0.6
D2-1	6.2 ± 0.63	6.2 ± 0.6	6.1 ± 0.5	5.9 ± 0.4
E1-7	5.3 ± 0.83	4.5 ± 0.2	5.5 ± 0.8	5.5 ± 0.6
E1-17	4.8 ± 0.64	5.7 ± 0.5	4.6 ± 0.4	4.4 ± 0.4
E2-1	8.0 ± 0.58	8.3 ± 0.8	8.6 ± 1.2	7.6 ± 0.4
E3-1	5.7 ± 0.46	6.0 ± 0.4	6.2 ± 0.9	6.1 ± 2.0
F1-2	6.6 ± 0.47	7.2 ± 0.4	7.0 ± 1.0	6.4 ± 0.3
G1-1	5.1 ± 0.39	5.2 ± 0.3	5.4 ± 0.7	5.1 ± 0.4
G1-2	5.9 ± 0.74	6.4 ± 0.2	6.4 ± 0.8	5.8 ± 0.4
G2-1	4.6 ± 0.41	5.0 ± 0.5	4.5 ± 0.4	4.8 ± 0.6
G10-2	5.9 ± 0.47	5.9 ± 0.4	5.9 ± 1.0	5.9 ± 0.4
H1-5	5.9 ± 0.35	6.1 ± 0.3	5.9 ± 0.5	5.6 ± 0.6
H2-1	5.9 ± 0.68	6.2 ± 0.4	6.2 ± 0.8	6.4 ± 1.7
H10-1	4.8 ± 0.39	5.1 ± 0.3	5.1 ± 0.5	5.2 ± 0.7
J1-1	5.1 ± 0.43	5.4 ± 0.3	5.5 ± 0.2	5.2 ± 0.3
K1-5	4.5 ± 0.49	4.5 ± 0.2	4.8 ± 0.6	4.9 ± 0.8
K1-8	5.5 ± 0.46	5.6 ± 0.4	5.9 ± 0.8	5.4 ± 0.4
L1-1	5.3 ± 0.85	5.7 ± 0.5	5.9 ± 0.5	5.9 ± 2.2
L2-1	5.3 ± 0.48	5.5 ± 0.2	5.7 ± 0.5	5.4 ± 0.6
M1-6	5.4 ± 0.64	5.7 ± 0.3	6.1 ± 1.1	5.2 ± 0.7
N1-4	5.4 ± 0.52	5.6 ± 0.3	5.9 ± 0.7	5.2 ± 0.5
P1-1	5.8 ± 0.53	5.9 ± 0.4	6.0 ± 0.4	5.6 ± 0.0
Q1-3	4.5 ± 0.54	4.7 ± 0.2	4.6 ± 0.6	4.6 ± 0.4
R1-1	5.4 ± 0.51	6.1 ± 0.6	5.8 ± 0.8	5.6 ± 0.5