March 15, 2011

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U.S. Department of Energy
SPRU Field Office SP23
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DOE CONTRACT NO. DE-AC05-06OR23100
SUBJECT: FINAL REPORT – INDEPENDENT VERIFICATION SURVEY ACTIVITIES AT THE SEPARATIONS PROCESS RESEARCH UNIT SITES, NISKAYUNA, NEW YORK – DCN 0496-SR-06-0

Dear Dr. Davis:

During the period of September 29, 2008 through August 25, 2010, the Oak Ridge Institute for Science and Education conducted verification survey activities of the Lower Level Land Area and North Field Project associated with the Separations Process Research Unit site. This final report consolidates the survey results from the previous five interim reports.

You may contact me via my information provided below or Phyllis Weaver at 865.576.5321 should you have any questions or require additional information.

Sincerely,

Evan M. Harpenau
Assistant Project Manager/Health Physicist
Independent Environmental Assessment and Verification

EMH:bf

Enclosure

c: File/0496

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FINAL REPORT – INDEPENDENT VERIFICATION SURVEY ACTIVITIES AT THE SEPARATIONS PROCESS RESEARCH UNIT SITES, NISKAYUNA, NEW YORK

E. M. Harpenau

Prepared for the
U.S. Department of Energy
The Oak Ridge Institute for Science and Education (ORISE) is a U.S. Department of Energy institute focusing on scientific initiatives to research health risks from occupational hazards, assess environmental cleanup, respond to radiation medical emergencies, support national security and emergency preparedness, and educate the next generation of scientists. ORISE is managed by Oak Ridge Associated Universities.

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INDEPENDENT VERIFICATION SURVEY ACTIVITIES
AT THE SEPARATIONS PROCESS RESEARCH UNIT SITES,
NISKAYUNA, NEW YORK

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Prepared for the
U.S. Department of Energy

FINAL REPORT

ORISE

MARCH 2011

This report is based on work performed by the Oak Ridge Institute for Science and Education under contract number DE-AC05-06OR23100 with the U.S. Department of Energy.
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AT THE SEPARATIONS PROCESS RESEARCH UNIT SITES,
NISKAYUNA, NEW YORK

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Date: 3/15/11

Date: 3/15/11
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ABBREVIATIONS AND ACRONYMS

BKG  background
cm   centimeter
COC  constituents of concern
cpm  counts per minute
CSAP/FSS Confirmation Sample Analysis Plan/Final Status Survey
DCGL derived concentration guideline level
DER  Division of Environmental Remediation
DOE  U.S. Department of Energy
FIDLER Field Instrument for the Detection of Low Energy Radiation
FSS  final status survey
GPS  Global Positioning System
in   inch
ISM  Integrated Safety Management
ISOCS In-Situ Object Counting System
ITP  Intercomparison Testing Program
IV   independent verification
JHA  Job Hazard Analyses
KAPL Knolls Atomic Power Laboratory
µg/kg microgram per kilogram
mg/kg milligram per kilogram
LLLA Lower Level Land Area
LLPL Lower Level Parking Lot
MAPEP Mixed Analyte Performance Evaluation Program
MARSSIM Multi-Agency Survey and Site Investigation Manual
MDC  minimum detectable concentration
NaI  sodium iodide
NFP  North Field Project
NIST National Institute for Standards and Technology
NRIP National Radiochemistry Intercomparison Program
NYSDEC New York State Department of Environmental Conservation
ORAU Oak Ridge Associated Universities
ORISE Oak Ridge Institute for Science and Education
pCi/g picocuries per gram
PSP  Project-Specific Plan
QAPP Quality Assurance Project Plan
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>RCRA</td>
<td>Resource Conservation and Recovery Act</td>
</tr>
<tr>
<td>ROCs</td>
<td>radionuclides of concern</td>
</tr>
<tr>
<td>RSS</td>
<td>ranked set sampling</td>
</tr>
<tr>
<td>SCOs</td>
<td>Soil Cleanup Objectives</td>
</tr>
<tr>
<td>SDSA</td>
<td>Slurry Drum Storage Area</td>
</tr>
<tr>
<td>SOF</td>
<td>sum of fractions</td>
</tr>
<tr>
<td>SPRU</td>
<td>Separations Process Research Unit</td>
</tr>
<tr>
<td>TAP</td>
<td>total absorption peak</td>
</tr>
<tr>
<td>VSP</td>
<td>Visual Sample Plan</td>
</tr>
</tbody>
</table>
1.0 INTRODUCTION AND SITE HISTORY

The Separations Process Research Unit (SPRU) complex located on the Knolls Atomic Power Laboratory (KAPL) site in Niskayuna, New York, was constructed in the late 1940s to research the chemical separation of plutonium and uranium (Figure A-1). SPRU operated as a laboratory scale research facility between February 1950 and October 1953. The research activities ceased following the successful development of the reduction oxidation and plutonium/uranium extraction processes. The oxidation and extraction processes were subsequently developed for large scale use by the Hanford and Savannah River sites (aRe 2008a). Decommissioning of the SPRU facilities began in October 1953 and continued through the 1990s.

The U.S. Department of Energy (DOE) used its authority under the Comprehensive Environmental Response, Compensation, and Liability Act to pursue cleanup of radioactivity and heavy metal chemicals of impacted soils in land areas utilized for SPRU operations. The objectives for this work included (1) restoring the Lower Level Land Area (LLLA) and North Field Project (NFP) area to a state suitable for reuse by KAPL in an area zoned for industrial and research use, (2) reducing surveillance and maintenance costs, and (3) reducing or eliminating the potential for future radiological and chemical releases from the SPRU land areas (aRe 2008b and 2009). After being awarded the contract, Accelerated Remediation Company began completing the last phases of decommissioning for the LLLA and NFP (Figure A-2). The contractor completed remediation of the former SPRU research operations in the LLLA and the NFP in late 2010.

The DOE Office of Environmental Management requested the services of Oak Ridge Institute for Science and Education (ORISE) to conduct independent verification (IV) of decontamination activities of specific land areas associated with the former SPRU operations. These verification activities included assessments of residual radiological contaminants in soil and in a few areas, provide verification that heavy metal chemical concentrations are within the established criteria. ORISE performed IV activities during the period of September 29, 2008 through August 25, 2010.
2.0 SITE DESCRIPTION

The SPRU complex, situated along the bank of the Mohawk River, occupied approximately 15 acres in the northwest corner of the 170 acre KAPL site. The LLLA at SPRU consisted of the Lower Level Parking Lot (LLPL), the Railroad Staging Area, the former K5 Retention Basin, and the former K6 and K7 waste storage pads (Re 2008a). These were subdivided into at least 40 final status survey (FSS) units designated as either Class 1, 2, or 3. During remediation and FSS activities, survey units were added and/or initial FSS unit boundaries were restructured (Figure A-3).

The NFP included an area of 15 acres of relatively flat field and sparsely wooded area located within the southeastern boundary of the KAPL site (Figure A-4). The NFP site was designated as the Slurry Drum Storage Area (SDSA) for staging and temporary storage of the slurry generated in the extraction process (DOE 2006). In 1955, the bermed walls of the SDSA were bulldozed into the area as fill following the completion of SPRU operations. The contractor determined seventeen FSS units were necessary based on result of previous use and excavations.

3.0 OBJECTIVES

The objectives of IV were to confirm that remedial actions have been effective in meeting established release criteria and that documentation accurately and adequately describes the final radiological conditions and heavy metal chemical conditions (where required) of the LLLA and NFP.

4.0 PROCEDURES

ORISE site verification activities were performed during the period of September 29, 2008 through August 25, 2010 in accordance with Project-Specific Plans (PSPs) submitted to and approved by DOE, the ORISE Survey Procedures Manual, and the ORAU Quality Program Manual (ORISE 2008a and 2010a, 2008b, and ORAU 2009). ORISE activities included surface scans for both low and high energy gamma emitters, In-Situ Object Counting System (ISOCS) gamma measurements, observation of gamma walkover surveys, along with systematic and judgmental soil sample collection. Verification surveys were performed for FSS survey units 1 through 11, 13, and, 14 at the LLLA and units 101 through 110 and 112 through 116 at the NFP site. Survey unit information is summarized in Table 1 and in Figures A-3 and A-4.
### TABLE 1: SEPARATIONS PROCESS RESEARCH UNIT SURVEY UNIT SUMMARY

<table>
<thead>
<tr>
<th>Lower Level Land Area</th>
<th>North Field Project</th>
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<tbody>
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<td><strong>Survey Unit ID</strong></td>
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<tr>
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<tr>
<td>FSS 2</td>
<td>1</td>
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<td>FSS 13</td>
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</tr>
<tr>
<td>FSS 14</td>
<td>1</td>
</tr>
<tr>
<td>FSS L</td>
<td>1</td>
</tr>
<tr>
<td>FSS N</td>
<td>2</td>
</tr>
<tr>
<td>FSS N1</td>
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<tr>
<td>FSS P</td>
<td>2</td>
</tr>
<tr>
<td>FSS S</td>
<td>1</td>
</tr>
</tbody>
</table>

*Refer to Figures A-3 and A-4

### 4.1 DOCUMENT REVIEW

ORISE reviewed the Radiological Confirmation Sampling Analysis Plans/Final Status Surveys (CSAP/FSS) (aRe 2008b and 2009), for agreement with the Multi-Agency Radiation Survey and Site Investigation Manual (MARSSIM), Resource Conservation and Recovery Act (RCRA) work plan (aRe 2008a), and associated Quality Assurance Project Plan (QAPP) documents for technical content. ORISE also reviewed FSS data as it was provided by the contractor.

Overall, project documentation set forth the necessary requirements for the contractor to complete site remediation.

### 4.2 REFERENCE SYSTEM

ORISE used a global positioning system (GPS) for documenting survey area boundaries, tracking survey data, and identifying measurement and sampling locations. Typically, boundaries for the
specified survey areas were obtained from coordinates provided by the remediation contractor and uploaded to handheld GPS units. FSS unit boundaries would vary on occasion from original coordinates and adjustments would be made on-site. Once the survey area was bounded, GPS units were integrated with gamma radiation detectors coupled to ratemeter-scalers to collect position and gamma radiation count rate data during walkover scans. The specific geographic coordinate system required for documentation was the New York State Plane Coordinate System. GPS coordinates were typically accurate to within one meter of any given position data point.

4.3 SURFACE SCANS

High density scans for gamma radiation were performed in accessible areas at LLLA and the NFP sites. Scans were primarily performed using a 2-inch × 2-inch (2×2) sodium iodide (NaI) scintillation detector. Detectors were coupled to a ratemeter-scaler with audible output. Locations of suspect, elevated direct gamma levels were identified in real-time via the audio output. Additionally, the electronic data were plotted and reviewed to identify any anomalies. Scan results collected using 2×2 detectors for the LLLA and NFP are provided in Figures A-5 and A-6. Locations of suspect elevated direct gamma radiation were identified for further investigation, as necessary.

To gain a higher efficiency for detecting the low energy radionuclides of plutonium and americium, ORISE used the Field Instrument for the Detection of Low Energy Radiation (FIDLER) NaI scintillation detector (ORISE 2009a and b). FIDLER scans, in correlation with the surveys conducted with the 2x2 detector, were completed in several FSS units. The FIDLER scan data is illustrated in Figure A-7.

4.4 STATISTICAL SAMPLING DESIGN

Visual Sample Plan (VSP) software was used to generate random coordinates for gamma measurements and soil sampling. These measurement/sample points were down-loaded to the GPS and were based upon the reference grid system established by the site. These predetermined random field assessment and resultant soil sample locations were designed and generated based on the ranked set sampling (RSS) approach (EPA 2006).
The statistically-based RSS design was used to determine the number of random samples that would be required to estimate that the mean concentration of the analytes of concern do not exceed the soil cleanup objectives (SCOs) for a given population and compare to the site’s FSS results; however, it does not require the assumption of a normal distribution. The process combines random sampling with the use of professional judgment to select sampling locations. Professional judgment relies upon the ability to assess the relative magnitude of gamma radiation levels between randomly selected locations. In this case, the gamma count rate data collected at randomly selected locations provided the measurable field screening method that correlated with the relative concentrations of the gamma-emitting contaminants of concern. The count rate data obtained were then used to select a specific sampling location.

The RSS systematic planning process uses a replication method on a larger random population from which the locations for the resulting samples can be selected. Replication refers to the number of cycles ($r$) for performing a set size ($m$) of field measurement. The number of field assessment locations per cycle, is a function of the set size and is simply $m^2$. The number of field assessment locations will vary but is defined as $m^2 \times r$, or for example $3^2 \times 2 = 18$ for a particular survey unit. These measurements are grouped into cycle/sets and distributed in the survey area. The first set identification location is cycle 1 of set 1 at measurement location 1, designated as 1-1-1. Mapping the population of assessment locations uses color coding with a specific geometric shape to correspond to each RSS cycle.

A one-minute static gamma count rate measurement was performed at each of the assessment locations per the survey unit requirement. The data points within a given cycle were then ranked as exhibiting either the lowest, medium, or highest gamma count in a set. The ranking resulted in the number of soil samples required for each FSS unit.

Twelve randomly generated locations were created in FSS 9 and FSS 10 using VSP software in lieu of the typical RSS method with a one minute static count collected at each location. This decision, approved by ORISE management and SPRU personnel, was prompted by a significant portion of FSS 9 not being accessible due to a large excavation filled with water and multiple smaller excavations scattered throughout the FSS unit. FSS 10 had multiple excavations as well as two large fractional distillation tanks contained within its boundary (ORISE 2010b).
4.5 **SOIL SAMPLING**

Surface soil samples (0 to 0.15 m) were collected at the determined RSS or random assessment locations. DOE also required chemical analysis of the LLLA samples be performed due to heavy metal chemical concentrations, primarily arsenic, observed in initial sampling efforts conducted by the decommissioning contractor. The chemical samples were obtained from undisturbed soils at each of the assessment locations. Thirty-six random radiological samples were collected from FSS units 2, 4, 9, 10, 13, and 14 in the LLLA and three additional judgmental samples based on scans (Figures A-8 and A-9). Judgmental samples were obtained from the locations that exhibited the maximum gamma activity levels observed during scans (ORISE 2009b and 2010b).

Chemical samples were collected in FSS units 1, 4, 9, and 10 using the RSS approach. Six judgmental samples from FSS 1 were collected at the direction of DOE to verify the contractor’s findings at specified locations. The remaining 18 samples were collected based either on process knowledge or the random/RSS sampling plan (Figure A-10). After the chemical verification samples had been collected, DOE determined that the random/RSS sampling approach for identifying heavy metal chemicals in FSS 13 and FSS 14 did not provide sufficient data to meet the criteria set in the RCRA Work Plan (aRc 2008a). Specific RCRA and the New York State Department of Environmental Conservation’s (NYSDEC) Division of Environmental Remediation (DER) DER-10 guidance (NYSDEC 2002), results from aRc data, and professional judgment were used to determine chemical sample locations in FSS units 13 and 14. It was decided that a sample be collected every 60 feet along the sidewalls of the excavation 15C resulting in 13 samples (ORISE 2010b). This resulted in the collection of 13 new samples from the two FSS units, four samples obtained from FSS 14 and nine from FSS 13 (Figure A-11). Figure A-12 depicts each sample type as it corresponds to a given location and the analytical results are provided in Table B-6.

In the NFP, six FSS units were sampled. A set of six RSS samples were collected from the FSS units 103, 109, 113, 114, 115, and 116. This resulted in 36 samples being collected for radiological analysis and no chemical sampling was requested (ORISE 2010c and 2010d). The location of samples within the NFP survey units are provided in Figure A-13. Several survey units were not selected for soil sampling because some areas had been backfilled prior to ORISE arriving on-site, process knowledge about radiological activities in specific areas was sufficient, or gamma walkover surveys did not indicate elevated radiation levels.
4.6 ISOCS MEASUREMENTS

The ISOCS portable gamma spectroscopy unit was used to identify and provide real time isotopic distribution at two locations identified via surface scans of high gamma activity in FSS units 103 and 115 (ORISE 2010c). The soil samples were counted for ten minutes and the spectra reviewed for the radionuclides of concern (ROCs) listed in Table 2. The locations of the ISOCS samples are shown in Figure A-14.

5.0 SOIL CLEANUP OBJECTIVES

The cleanup objectives for the ROCs at the LLLA and NFP are provided in Table 2 along with the respective derived concentration guideline levels (DCGLs) for each. The most prominent ROC identified has been cesium-137. The heavy metal chemical constituents of concern (COCs) and the respective SCOs are provided in Table 3 (aRe 2008a). The site goal was to return the soil to KAPL suitable for future use.
### TABLE 2:
**RADIONUCLIDES OF CONCERN DCGLS**  
**SEPARATIONS PROCESS RESEARCH UNIT**  
**NISKAYUNA, NEW YORK**

<table>
<thead>
<tr>
<th>ROC</th>
<th>DCGL (pCi/g)</th>
<th>ROC</th>
<th>DCGL (pCi/g)</th>
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<tbody>
<tr>
<td>Am-241</td>
<td>5.74E+02</td>
<td>Pu-241</td>
<td>1.91E+04</td>
</tr>
<tr>
<td>Cs-137</td>
<td>3.00E+01</td>
<td>Pm-147</td>
<td>1.59E+06</td>
</tr>
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<td>Co-60</td>
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<td>Sm-151</td>
<td>6.59E+06</td>
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<td>Pu-240</td>
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### TABLE 3:
**CHEMICAL CONSTITUENTS OF CONCERN**  
**SOIL CLEANUP OBJECTIVES**  
**SEPARATIONS PROCESS RESEARCH UNIT**  
**NISKAYUNA, NEW YORK**

<table>
<thead>
<tr>
<th>COC</th>
<th>SCO (mg/kg)a</th>
<th>COC</th>
<th>SCO (mg/kg)a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antimony</td>
<td>1b</td>
<td>Lead</td>
<td>400</td>
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<tr>
<td>Arsenic</td>
<td>16</td>
<td>Mercury</td>
<td>0.73</td>
</tr>
<tr>
<td>Cadmium</td>
<td>2.5</td>
<td>Silver</td>
<td>8.3</td>
</tr>
<tr>
<td>Trivalent Chromium</td>
<td>36</td>
<td>Thallium</td>
<td>2b</td>
</tr>
<tr>
<td>Cobalt</td>
<td>30c</td>
<td>Vanadium</td>
<td>150c</td>
</tr>
<tr>
<td>Copper</td>
<td>270</td>
<td>Zinc</td>
<td>2200</td>
</tr>
</tbody>
</table>

*a mg/kg = milligrams per kilogram.  
b Reporting Limit for ALS Laboratory Group.  
c TAGM 4046 recommended cleanup levels (aRe 2008a).
6.0 SAMPLE ANALYSIS AND DATA INTERPRETATION

Soil samples were returned to the ORISE laboratory in Oak Ridge, TN for radiological analysis and interpretation. Sample analyses were performed in accordance with the ORISE Laboratory Procedures Manual (ORISE 2010e). Soil samples were analyzed by gamma spectroscopy and alpha spectroscopy. The dominant ROC identified was cesium-137; however, gamma spectra were reviewed for any other gamma-emitting radionuclides. Alpha spectroscopy was also performed to provide supporting data to ensure the listed alpha emitting ROCs were below their respective DCGL. Spectroscopy results are reported in units of picocuries per gram (pCi/g). Analytical results were compared to the DCGLs provided in Table 2 (aRc 2008a).

Samples collected to determine chemical (specifically the heavy metal: arsenic) concentrations were analyzed by ALS Laboratory Group in Fort Collins, Colorado, a DOE Oak Ridge Operations SMO approved facility. ALS performed analyses in accordance with EPA-approved Solid Waste-846 methods 6010 and 7471 (EPA 2007). The results were reported in units of micrograms per kilogram (μg/kg) and converted to units of milligrams per kilogram (mg/kg).

Radiological and chemical data were compared with either the approved site DCGLs or the SCOs, as appropriate, that are provided in Tables 2 and 3 (aRc 2008a).

7.0 FINDINGS AND RESULTS

The results of the verification surveys for FSS units 1 through 11, 13, and 14 at the SPRU LILLA project site, and units 101 through 110 and 112 through 116 at the NFP are provided below.

7.1 DOCUMENT REVIEWS

Several inconsistencies were noticed within co-related documents pertaining to clean up goals, contaminants of concern, sample confirmation, and implementation of the Data Quality Objective process (ORISE 2008c and d). These issues were appropriately addressed by the contractor through implementing revised plans and work processes.

7.2 SURFACE SCANS

ORISE performed high density scans of all accessible areas for the FSS units covered by the independent verification activities with the exception of FSS unit 7 and FSS unit 11. Initial
assessments of FSS unit 7 determined that the unit was not expected to contain residual contamination levels approaching the DCGLs, and therefore, received a low density scan. A significant portion of FSS 11 consisted of a steep hillside that was not easily accessible, and was also expected to have a low potential for contamination. As a result, a medium density scan of accessible areas was performed.

Additionally, ORISE was requested by the DOE to observe contractor personnel performing gamma walkover surveys. The overall walking speed of the surveyor was appropriate and the distance between the detector and ground was satisfactory. The observed surveyor kept the face of the detector pointed toward the ground during most of the swing and was not observed swinging the detector too high. However, it was noticed that the observed surveyor was not wearing headphones. Headphones should be worn while scanning in order to make it easier to detect a change in the audio output from the detector. Overall, the observed surveyor adequately performed gamma walkover surveys (ORISE 2009b).

The typical ambient background radiation levels observed with the NaI detectors ranged from 4,000 to 14,000 cpm. The a priori scan minimum detectable concentration (MDC) for a 2x2 detector is approximately 6.4 pCi/g for cesium-137 (estimated from Table 6.4 of NUREG-1507) which is less than the site specific DCGL of 30 pCi/g (NRC 1997). The areas identified as exhibiting elevated activity when compared to the rest of the FSS unit received additional investigative scans.

7.3 SOIL ANALYSIS

Gamma spectroscopy sample results for the independently verified FSS units are provided in Tables B-1 and B-2. Concentrations of the dominant ROC: cesium-137, ranged from -0.02 to 3.95 pCi/g. Alpha spectroscopy results for americium-241; plutonium-238, 239, and 240; uranium-234, 235, and 238 are reported in Tables B-3 and B-4. The two judgmental locations sampled and analyzed using the ISOCS detector, showed no ROCs above detectable levels. The radionuclide sum-of-fractions (SOF) for each sample location was evaluated and determined to be below the threshold of 1.0.

Analytical results for heavy metal chemicals in soil and the DOE requested RCRA samples are provided in Tables B-5 and B-6. The soil in six FSS units from the LLLA were sampled and analyzed for chemical content. The primary constituent of concern for chemicals was determined to
be the heavy metal: arsenic. Arsenic concentrations ranged from 4 to 170 mg/kg with six sample locations in excess of the SCO. Those locations were remediated after the IV results were sent to the SPRU site.

8.0 COMPARISION OF RESULTS WITH GUIDELINES

The final concentrations for the analytes of concern at SPRU’s LLLA and NFP meet the cleanup objectives as described in the revised CSAP. Radiological sample concentrations were less than 14% of the respective DCGL values for all radionuclides of concern.

Chemical concentration values in excess of the SCO for arsenic were identified in FSS units 1, 9, 10, and 14 during IV sampling activities (Tables B-5 and B-6). The sample locations that exceeded the SCOs were remediated by the site contractor and the post remediation sample results were submitted to the NYSDEC, Region 4. The NYSDEC issued approval statements after evaluating the sample results for FSS units 9 and 10 (NYSDEC 2009 and 2010).

9.0 CONCLUSION

During the period of September 29, 2008 through August 25, 2010, ORISE conducted verification survey activities including scans, sampling, and the observation of scan surveys of the Lower Level Parking Area and North Field Project at the SPRU site. Based on the scan and sample results of the ORISE verification survey activities and approval of additional remediation activities at the SPRU site, all independently verified FSS units meet the applicable derived concentration guideline level or site cleanup objectives for radiological and chemical contaminants.
10.0  REFERENCES


Oak Ridge Institute for Science and Education. Final Interim Report - Verification Survey Activities in Final Status Survey Units 7, 8, 9, 10, 11, 13, and 14 at the Separations Process Research Unit, Niskayuna, New York. Oak Ridge, Tennessee; May 26, 2010b.


APPENDIX A
FIGURES
Figure A-1: Separations Process Research Unit (SPRU) at the Knolls Atomic Power Laboratory
Figure A-2: SPRU Plot Plans for the Lower Level Land Area and North Field Project
Figure A-3: SPRU Lower Level Land Area Final Status Survey Units
Figure A-4: North Field Project Final Status Survey Units
Figure A-5: Lower Level Land Area Surface Scans with 2x2 Detector
Figure A-6: North Field Project Surface Scans with 2x2 Detector
Figure A-7: Lower Level Land Area Surface Scans with FIDLER Detector
Figure A-8: Radiological RSS Sample Locations for the Lower Level Land Area
Figure A-9: Radiological Judgmental Sample Locations for the Lower Level Land Area
Figure A-10: Chemical Sample Locations for the Lower Level Land Area
Figure A-11: RCRA Sample Locations for the Lower Level Land Area
Figure A-12: RCRA Sample Location Type for the Lower Level Land Area
Figure A-13: Radiological RSS Sample Locations for the North Field Project
Figure A-14: Judgmental Radiological ISOCS Sample Locations for the North Field Project
### TABLE B-1:
SUMMARIZED RADIONUCLIDE CONCENTRATION RANGES IN SOIL BY GAMMA SPECTROSCOPY
LOWER LEVEL LAND AREA
SEPARATIONS PROCESS RESEARCH UNIT
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample Area</th>
<th>Radionuclide Concentration in Soil Samples (pCi/g)</th>
<th>Co-60</th>
<th>Cs-137</th>
<th>Eu-152</th>
<th>Eu-154</th>
<th>Eu-155</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lower Level Parking Lot</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FSS 2</td>
<td></td>
<td>-0.02 to 0.01</td>
<td>0.06 to 0.96</td>
<td>-0.90 to 0.06</td>
<td>-0.41 to 0.30</td>
<td>-0.03 to 0.10</td>
</tr>
<tr>
<td>FSS 4</td>
<td></td>
<td>-0.04 to 0.04</td>
<td>0.25 to 3.83</td>
<td>0.01 to 0.10</td>
<td>-0.03 to 0.03</td>
<td>-0.01 to 0.14</td>
</tr>
<tr>
<td>FSS 9</td>
<td></td>
<td>-0.03 to 0.00</td>
<td>0.08 to 3.95</td>
<td>-0.07 to 0.05</td>
<td>-0.03 to 0.03</td>
<td>-0.06 to 0.23</td>
</tr>
<tr>
<td>FSS 10</td>
<td></td>
<td>-0.04 to 0.05</td>
<td>0.20 to 0.38</td>
<td>-0.06 to 0.05</td>
<td>-0.02 to 0.01</td>
<td>0.02 to 0.08</td>
</tr>
<tr>
<td>FSS 13</td>
<td></td>
<td>-0.03 to 0.05</td>
<td>0.04 to 0.97</td>
<td>-0.05 to 0.00</td>
<td>-0.06 to 0.03</td>
<td>0.02 to 0.12</td>
</tr>
<tr>
<td>FSS 14</td>
<td></td>
<td>-0.03 to 0.03</td>
<td>0.03 to 0.35</td>
<td>-0.02 to 0.01</td>
<td>-0.02 to 0.02</td>
<td>-0.04 to 0.06</td>
</tr>
</tbody>
</table>

*Refer to Figures A-8 and A-9

### TABLE B-2:
SUMMARIZED RADIONUCLIDE CONCENTRATION RANGES IN SOIL BY GAMMA SPECTROSCOPY
NORTH FIELD PROJECT
SEPARATIONS PROCESS RESEARCH UNIT
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample Area</th>
<th>Radionuclide Concentration in Soil Samples (pCi/g)</th>
<th>Co-60</th>
<th>Cs-137</th>
<th>Eu-152</th>
<th>Eu-154</th>
<th>Eu-155</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>North Field Project</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FSS 103</td>
<td></td>
<td>-0.06 to 0.04</td>
<td>-0.02 to 0.36</td>
<td>-0.06 to 0.04</td>
<td>-0.02 to 0.01</td>
<td>-0.01 to 0.15</td>
</tr>
<tr>
<td>FSS 109</td>
<td></td>
<td>-0.05 to 0.03</td>
<td>0.04 to 2.07</td>
<td>-0.05 to 0.00</td>
<td>-0.03 to 0.01</td>
<td>0.00 to 0.05</td>
</tr>
<tr>
<td>FSS 113</td>
<td></td>
<td>-0.06 to 0.02</td>
<td>0.18 to 1.36</td>
<td>-0.04 to 0.04</td>
<td>-0.01 to 0.02</td>
<td>0.01 to 0.10</td>
</tr>
<tr>
<td>FSS 114</td>
<td></td>
<td>-0.05 to 0.07</td>
<td>0.21 to 0.42</td>
<td>-0.05 to 0.02</td>
<td>-0.02 to 0.01</td>
<td>-0.05 to 0.12</td>
</tr>
<tr>
<td>FSS 115</td>
<td></td>
<td>-0.04 to 0.03</td>
<td>0.14 to 0.48</td>
<td>-0.04 to 0.01</td>
<td>-0.02 to 0.02</td>
<td>-0.04 to 0.08</td>
</tr>
<tr>
<td>FSS 116</td>
<td></td>
<td>-0.02 to 0.02</td>
<td>0.20 to 0.30</td>
<td>-0.05 to 0.07</td>
<td>-0.02 to 0.02</td>
<td>-0.01 to 0.12</td>
</tr>
</tbody>
</table>

*Refer to Figures A-13.
TABLE B-3:
SUMMARIZED RADIONUCLIDE CONCENTRATION RANGES IN SOIL BY ALPHA SPECTROSCOPY
SEPARATIONS PROCESS RESEARCH UNIT
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample Area</th>
<th>Am-241</th>
<th>Pu-238</th>
<th>Pu-239/240</th>
<th>U-234</th>
<th>U-235</th>
<th>U-238</th>
<th>Total U^b</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lower Level Land Area</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FSS 2</td>
<td>0.00 to 0.04</td>
<td>-0.03 to 0.01</td>
<td>0.00 to 0.15</td>
<td>0.81 to 0.99</td>
<td>0.04 to 0.06</td>
<td>0.82 to 1.03</td>
<td>1.67 to 2.06</td>
</tr>
<tr>
<td>FSS 4</td>
<td>-0.02 to 0.04</td>
<td>-0.01 to 0.00</td>
<td>0.01 to 0.16</td>
<td>0.57 to 2.35</td>
<td>0.03 to 0.11</td>
<td>0.55 to 2.31</td>
<td>1.16 to 4.77</td>
</tr>
<tr>
<td>FSS 9</td>
<td>0.01 to 0.03</td>
<td>0.00 to 0.05</td>
<td>0.01 to 0.02</td>
<td>0.33 to 3.97</td>
<td>0.01 to 0.19</td>
<td>0.37 to 3.85</td>
<td>0.71 to 8.01</td>
</tr>
<tr>
<td>FSS 10</td>
<td>0.00 to 0.03</td>
<td>0.01 to 0.05</td>
<td>0.01 to 0.02</td>
<td>0.55 to 1.09</td>
<td>0.02 to 0.08</td>
<td>0.55 to 1.12</td>
<td>1.12 to 2.24</td>
</tr>
<tr>
<td>FSS 13</td>
<td>-0.02 to 0.02</td>
<td>0.00 to 0.03</td>
<td>0.01 to 0.03</td>
<td>0.52 to 1.15</td>
<td>0.03 to 0.08</td>
<td>0.48 to 1.09</td>
<td>1.03 to 2.32</td>
</tr>
<tr>
<td>FSS 14</td>
<td>0.00 to 0.02</td>
<td>0.01 to 0.04</td>
<td>0.01 to 0.03</td>
<td>0.65 to 1.45</td>
<td>0.03 to 0.06</td>
<td>0.70 to 1.38</td>
<td>1.37 to 2.88</td>
</tr>
</tbody>
</table>

^aRefer to Figures A-8 and A-9.
^bTotal U is calculated by U-234 + U-235 + U-238.
### TABLE B-4:
SUMMARIZED RADIONUCLIDE CONCENTRATION
RANGES IN SOIL BY ALPHA SPECTROSCOPY
SEPARATIONS PROCESS RESEARCH UNIT
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample Area¹</th>
<th>Am-241</th>
<th>Pu-238</th>
<th>Pu-239/240</th>
<th>U-234</th>
<th>U-235</th>
<th>U-238</th>
<th>Total U⁵</th>
</tr>
</thead>
<tbody>
<tr>
<td>North Field Project</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FSS 103</td>
<td>0.00 to 0.01</td>
<td>0.00 to 0.02</td>
<td>0.00 to 0.02</td>
<td>0.74 to 0.93</td>
<td>0.03 to 0.05</td>
<td>0.77 to 0.95</td>
<td>1.56 to 1.91</td>
</tr>
<tr>
<td>FSS 109</td>
<td>-0.05 to 0.03</td>
<td>0.01 to 0.03</td>
<td>0.00 to 0.03</td>
<td>0.66 to 1.23</td>
<td>-0.08 to 0.22</td>
<td>0.66 to 1.23</td>
<td>1.24 to 2.56</td>
</tr>
<tr>
<td>FSS 113</td>
<td>-0.06 to 0.04</td>
<td>0.00 to 0.01</td>
<td>0.01 to 0.03</td>
<td>0.69 to 1.03</td>
<td>-0.06 to 0.19</td>
<td>0.69 to 1.03</td>
<td>1.32 to 2.21</td>
</tr>
<tr>
<td>FSS 114</td>
<td>-0.09 to 0.05</td>
<td>0.01 to 0.02</td>
<td>0.00 to 0.02</td>
<td>0.69 to 1.38</td>
<td>-0.20 to 0.19</td>
<td>0.69 to 1.38</td>
<td>1.40 to 2.95</td>
</tr>
<tr>
<td>FSS 115</td>
<td>-0.01 to 0.01</td>
<td>-0.01 to 0.01</td>
<td>0.00 to 0.04</td>
<td>0.76 to 1.11</td>
<td>0.03 to 0.06</td>
<td>0.81 to 0.97</td>
<td>1.61 to 2.11</td>
</tr>
<tr>
<td>FSS 116</td>
<td>-0.09 to 0.04</td>
<td>0.00 to 0.02</td>
<td>0.01 to 0.02</td>
<td>0.79 to 1.49</td>
<td>-0.05 to 0.14</td>
<td>0.79 to 1.49</td>
<td>1.61 to 3.10</td>
</tr>
</tbody>
</table>

---

¹Refer to Figures A-13.

⁵Total U is calculated by U-234 + U-235 + U-238
### TABLE B-5:  
CONCENTRATIONS OF CHEMICALS IN SOIL SAMPLES  
SEPARATIONS PROCESS RESEARCH UNIT  
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample ID&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Contaminant of Concern (mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sb</td>
</tr>
<tr>
<td><strong>FSS 1</strong></td>
<td></td>
</tr>
<tr>
<td>S007</td>
<td>1.4 U&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>S008</td>
<td>0.5 J</td>
</tr>
<tr>
<td>S009</td>
<td>1.5 UJ</td>
</tr>
<tr>
<td>S010</td>
<td>1.1 UJ</td>
</tr>
<tr>
<td>S011</td>
<td>1.3 UJ</td>
</tr>
<tr>
<td>S012</td>
<td>1.2 UJ</td>
</tr>
<tr>
<td><strong>FSS 4</strong></td>
<td></td>
</tr>
<tr>
<td>S019</td>
<td>1.1 BJ</td>
</tr>
<tr>
<td>S020</td>
<td>1.3 BJ</td>
</tr>
<tr>
<td>S021</td>
<td>0.49 BJ</td>
</tr>
<tr>
<td>S022</td>
<td>0.29 BJ</td>
</tr>
<tr>
<td>S023</td>
<td>1.2 BJ</td>
</tr>
<tr>
<td>S024</td>
<td>1.4 BJ</td>
</tr>
<tr>
<td><strong>FSS 10</strong></td>
<td></td>
</tr>
<tr>
<td>S039</td>
<td>0.99 BJ</td>
</tr>
<tr>
<td>S040</td>
<td>1.2 J</td>
</tr>
<tr>
<td>S041</td>
<td>1.3 J</td>
</tr>
<tr>
<td>S042</td>
<td>1.2 J</td>
</tr>
<tr>
<td>S043</td>
<td>0.83 BJ</td>
</tr>
<tr>
<td>S044</td>
<td>0.87 BJ</td>
</tr>
</tbody>
</table>
TABLE B-5:
CONCENTRATIONS OF CHEMICALS IN SOIL SAMPLES
SEPARATIONS PROCESS RESEARCH UNIT
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample ID&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Contaminant of Concern (mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sb</td>
</tr>
<tr>
<td>FSS 9</td>
<td></td>
</tr>
<tr>
<td>S045</td>
<td>1.7 N&lt;sup&gt;j&lt;/sup&gt;</td>
</tr>
<tr>
<td>S046</td>
<td>1.4 J</td>
</tr>
<tr>
<td>S047</td>
<td>1.1 BJ</td>
</tr>
<tr>
<td>S048</td>
<td>1.2 BJ</td>
</tr>
<tr>
<td>S049</td>
<td>0.3 UJ</td>
</tr>
<tr>
<td>S050</td>
<td>1 BJ</td>
</tr>
</tbody>
</table>

<sup>a</sup>Refer to Figure A-10.

<sup>b</sup>U = Compound was analyzed for but not detected. The associated numerical value is the estimated sample quantitation limit which is included and corrected for dilution and percent moisture.

<sup>j</sup>J = Indicates an estimated value. This flag is used when estimating a concentration for tentatively identified compounds where a 1:1 response is assumed or when the mass spectral data indicate the presence of a compound that meets the identification criteria but the result is less than the specified detection limit but greater than zero.

<sup>k</sup>B = Value was less than the CRDL (Contract Required Detection Limit) or RRL (Required Reporting Limit) specified, but greater than or equal to the IDL (Instrument Detection Limit)/MDL (Method Detection Limit).

<sup>n</sup>N = Spiked sample recovery not within control limits. A post spike is analyzed for all ICP analyses with the matrix spike or spike duplicate fail and the native sample concentration is less than four times the spike added concentration.
TABLE B-6:
RCRA SAMPLES FOR CHEMICALS IN SOIL SAMPLES
SEPARATIONS PROCESS RESEARCH UNIT
NISKAYUNA, NEW YORK

<table>
<thead>
<tr>
<th>Sample ID&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Contaminant of Concern (mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sb</td>
</tr>
<tr>
<td>S076 1.1 B&lt;sup&gt;b&lt;/sup&gt;J&lt;sup&gt;c&lt;/sup&gt;</td>
<td>24</td>
</tr>
<tr>
<td>S077 1.3 J</td>
<td>5.7</td>
</tr>
<tr>
<td>S078 1.1 J</td>
<td>6.7</td>
</tr>
<tr>
<td>S079 0.59 BJ</td>
<td>5</td>
</tr>
<tr>
<td>S080 1.0 BJ</td>
<td>5.9</td>
</tr>
<tr>
<td>S081 0.79 BJ</td>
<td>6.8</td>
</tr>
<tr>
<td>S082 0.92 BJ</td>
<td>4.5</td>
</tr>
<tr>
<td>S083 0.74 BJ</td>
<td>4.3</td>
</tr>
<tr>
<td>S084 1.5 BJ</td>
<td>7.3</td>
</tr>
<tr>
<td>S085 0.75 BJ</td>
<td>4.6</td>
</tr>
<tr>
<td>S086 1.5 J</td>
<td>4.7</td>
</tr>
<tr>
<td>S087 0.67 BJ</td>
<td>5.3</td>
</tr>
<tr>
<td>S088 1.0 BJ</td>
<td>7.1</td>
</tr>
</tbody>
</table>

<sup>a</sup>Refer to Figure A-11.

<sup>b</sup>B = Value was less than the CRDL (Contract Required Detection Limit) or RRL (Required Reporting Limit) specified, but greater than or equal to the IDL. (Instrument Detection Limit)/MDL. (Method Detection Limit)

<sup>c</sup>J = Indicates an estimated value. This flag is used when estimating a concentration for tentatively identified compounds where a 1:1 response is assumed or when the mass spectral data indicate the presence of a compound that meets the identification criteria but the result is less than the specified detection limit but greater than zero.

<sup>d</sup>U = Compound was analyzed for but not detected. The associated numerical value is the estimated sample quantitation limit which is included and corrected for dilution and percent moisture.
APPENDIX C
MAJOR INSTRUMENTATION
The display of a specific product is not to be construed as an endorsement of the product or its manufacturer by the author or his employer.

C.1 Scanning and Measurement Instrument/Detector Combinations

Ludlum Scintillation Detector Model SPA-3, Crystal: 2-in. x 2-in. (Ludlum Measurements, Inc., Sweetwater, TX)
Coupled to:
Ludlum Ratemeter-Scaler Model 2221 (Ludlum Measurements, Inc., Sweetwater, TX)
Coupled to:
Trimble GeoXH Receiver and Data Logger (Trimble Navigation Limited, Sunnyvale, CA)

Bicron Field Instrument Detector for Low Energy Radiation (FIDLER) Model G5 (Bicron Corporation, Newburg, OH)
Coupled to:
Ludlum Ratemeter-Scaler Model 2221 (Ludlum Measurements, Inc., Sweetwater, TX)
Coupled to:
Trimble GeoXH Receiver and Data Logger (Trimble Navigation Limited, Sunnyvale, CA)

High Purity Germanium Detector Model GR3819 (Canberra, Meriden, CT)
Used in conjunction with:
In-Situ Object Counting System (ISOCS) Calibration Software (Canberra, Meriden, CT) and Inspector -2000 Multichannel Analyzer with Dell Workstation

C.2 Laboratory Analytical Instrumentation

High-Purity Extended Range Intrinsic Detector CANBERRA/Tennelec Model No: ERVDS30-25195 (Canberra, Meriden, CT)
Used in conjunction with:
Lead Shield Model G-11 (Nuclear Lead, Oak Ridge, TN), Apex Gamma Software (Canberra, Meriden, CT) and Multichannel Analyzer with Dell Workstation

High-Purity Extended Range Intrinsic Detector Model No. GMX-45200-5 (AMETEK/ORTEC, Oak Ridge, TN)
Used in conjunction with:
Lead Shield Model SPG-16-K8 (Nuclear Data), Apex Gamma Software (Canberra, Meriden, CT) and Multichannel Analyzer with Dell Workstation

High-Purity Germanium Detector Model GMX-30-P4, 30% Eff. (AMETEK/ORTEC, Oak Ridge, TN)
Used in conjunction with:
Lead Shield Model G-16 (Gamma Products, Palos Hills, IL), Apex Gamma Software (Canberra, Meriden, CT) and Multichannel Analyzer with Dell Workstation
Alpha Spectroscopy System with 7401VR Spectrometers (Canberra, Meriden, CT) and TC256 Spectrometers (Tennelec/Canberra, Meriden, CT), Apex Alpha Software (Canberra, Meriden, CT) and Multichannel Analyzer with Dell Workstation.
APPENDIX D
SURVEY AND ANALYTICAL PROCEDURES
D.1 PROJECT HEALTH AND SAFETY

The survey and sampling procedures were evaluated to ensure that any hazards inherent to the procedures themselves were addressed in current job hazard analyses (JHAs). All survey and laboratory activities were conducted in accordance with ORISE health and safety and radiation protection procedures.

Pre-survey activities included an overview of potential health and safety issues. SPRU representatives provided site-specific safety awareness training for each individual ORISE survey effort. In-process and verification surveys were performed according to the ORISE generic health and safety plan, site-specific Integrated Safety Management (ISM) pre-job hazard checklist, and safety procedures discussed during the on-site training.

D.2 QUALITY ASSURANCE

Analytical and field survey activities were conducted in accordance with procedures from the following ORAU and ORISE documents:

- Survey Procedures Manual (ORISE 2008b)
- Laboratory Procedures Manual (ORISE 2010e)
- Quality Program Manual (ORAU 2009)

The procedures contained in these manuals were developed to meet the requirements of 10 CFR 830 Subpart A, Quality Assurance Requirements, Department of Energy Order 414.1C, Quality Assurance, and the U.S. Nuclear Regulatory Commission, Quality Assurance Manual for the Office of Nuclear Material Safety and Safeguards, and contain measures to assess processes during their performance.

Quality control procedures include:

- Daily instrument background and check-source measurements to confirm that equipment operation is within acceptable statistical fluctuations.

- Participation in Mixed Analyte Performance Evaluation Program (MAPEP), National Institute for Standards and Technology (NIST) Radiochemistry Intercomparison Program (NRIP), and Intercomparison Testing Program (ITP) Laboratory Quality
Assurance Programs.

- Training and certification of all individuals performing procedures.
- Periodic internal and external audits.

D.3 Calibration

Calibration of all field and laboratory instrumentation was based on standards/sources, traceable to NIST, when such standards/sources were available. In cases where they were not available, standards of an industry-recognized organization were used.

D.4 Survey Procedures

D.4.1 Surface Scans

Scans for elevated gamma radiation were performed by passing the detector slowly over the surface. The distance between the detector and surface was maintained at a nominal of about 1 to 5 centimeters (cm). NaI scintillation detectors were coupled to GPS units that enabled real-time recording of position in one-second intervals. Identification of elevated radiation levels was based on increases in the audible signal from the instrument. Positioning data files were downloaded from field data loggers for plotting using commercially available software (http://trl.trimble.com/docushare/dsweb/Get/Document-261826/GeoExpl2005_100A_GSG_ENG.pdf).

The scan minimum detectable concentrations (MDCs) for the NaI scintillation detector for the contaminants of concern in surface soil were obtained directly from NUREG-1507 when available or estimated using the calculation approach described in NUREG-1507. A typical NaI 2-inch by 2-inch detector MDC is 6.4 pCi/g for cesium-137. Audible increases in the activity rate are investigated by ORISE. It is standard procedure for ORISE staff to pause and investigate any locations where gamma radiation is distinguishable from background levels.

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D.4.2 Soil Sampling

Approximately 0.5 to 1 kilogram of soil was collected at each sample location. Collected samples were placed in plastic bags, sealed, and labeled in accordance with ORISE survey procedures.

D.5 Radiological Analysis

D.5.1 Detection Limits

Detection limits, referred to as MDC, were based on $3 + 4.65 \times \text{standard deviation of the background count}$, because of variations in background levels, measurement efficiencies, and contributions from other radionuclides in samples, the detection limits differ from sample to sample and instrument to instrument.

D.5.2 Alpha Spectroscopy

Soil samples were homogenized then dissolved by a combination of potassium hydroxide fluoride and pyrosulfate fusions. The fusion cakes are dissolved and all alpha emitters are co-precipitated on barium sulfate. The barium sulfate is re-dissolved and the COC is separated from the other actinides by extraction chromatography using Eichrom resins, co-precipitated with a cerium fluoride carrier, and analyzed using ion implanted detectors, alpha spectrometers, and multichannel analyzer.

An alpha spectroscopy detector system calculates an MDC for each individual isotope per sample based on the detector background, counting efficiency, yield, and quantity. An MDC is printed out with each sample results. The typical MDC for a 1,000-minute count time was 0.02 pCi/g. Total absorption peaks (TAPs) used for determining the activities of radionuclides of concern and the typical associated MDCs were:

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>TAP (MeV)</th>
<th>MDC (pCi/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Am-241</td>
<td>5.486</td>
<td>0.05</td>
</tr>
<tr>
<td>Pu-238</td>
<td>5.499</td>
<td>0.04</td>
</tr>
<tr>
<td>Pu-239/240</td>
<td>5.155</td>
<td>0.02</td>
</tr>
<tr>
<td>U-234</td>
<td>4.775</td>
<td>0.02</td>
</tr>
<tr>
<td>U-235</td>
<td>4.396</td>
<td>0.02</td>
</tr>
<tr>
<td>U-238</td>
<td>4.190</td>
<td>0.10</td>
</tr>
</tbody>
</table>

*Spectra were also reviewed for other identifiable TAPs.*
D.5.3 Gamma Spectroscopy

Samples of soil were dried, mixed, crushed, and/or homogenized as necessary, and a portion sealed in a 0.5-liter Marinelli beaker or other appropriate container. The quantity placed in the beaker was chosen to reproduce the calibrated counting geometry. Net material weights were determined and the samples counted using intrinsic germanium detectors coupled to a pulse height analyzer system. Background and Compton stripping, peak search, peak identification, and concentration calculations were performed using the computer capabilities inherent in the analyzer system. All TAPs associated with the radionuclides of concern were reviewed for consistency of activity. Total absorption peaks used for determining the activities of radionuclides of concern and the typical associated MDCs for a one-hour count time were:

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>TAP (MeV)</th>
<th>MDC (pCi/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cs-137</td>
<td>0.662</td>
<td>0.05</td>
</tr>
<tr>
<td>Co-60</td>
<td>1.170</td>
<td>0.06</td>
</tr>
<tr>
<td>Eu-152</td>
<td>0.344</td>
<td>0.30</td>
</tr>
<tr>
<td>Eu-154</td>
<td>0.723</td>
<td>0.30</td>
</tr>
<tr>
<td>Eu-155</td>
<td>0.105</td>
<td>0.14</td>
</tr>
</tbody>
</table>

*Spectra were also reviewed for other identifiable TAPs.

D.5.4 Uncertainties

The uncertainties associated with the analytical data presented in the tables of this report represent the total propagated uncertainties for those data. These uncertainties were calculated based on both the gross sample count levels and the associated background count level.