

# CUY-90-14.90

PID 77332/85531

# **APPENDIX EC-11**

# **ESA Phase II Reports** (Contract Document)

State of Ohio
Department of Transportation
Jolene M. Molitoris, Director

Innerbelt Bridge
Construction Contract Group 1 (CCG1)

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This Appendix includes Phase II Environmental Site Assessment reports completed by the Department for this project. Please note that many of the reports come from the Cleveland Innerbelt Corridor report, which includes locations outside the footprint of Construction Contract Group 1. This Appendix only includes locations within the CCG1 project limits.

# PHASE II ENVIRONMENTAL SITE ASSESSMENT

# CUY-CLEVELAND INNERBELT COMMERCIAL ROAD ALIGNMENT PROJECT AREA (PID 77510) CLEVELAND, CUYAHOGA COUNTY, OHIO

### JANUARY 2010

### Prepared for:

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#### **EXECUTIVE SUMMARY**

HzW Environmental Consultants, LLC (HzW) was contracted by the Ohio Department of Transportation (ODOT) Office of Environmental Services (OES) to conduct a Phase II Environmental Site Assessment (ESA) of the CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area (PID 77510), Cleveland, Cuyahoga County, Ohio (herein referred to as the "Project Area"). This project was conducted in accordance with HzW's proposal dated November 3, 2009, which was authorized by ODOT OES on November 19, 2009. The purpose of the Phase II ESA investigation was to assess the unknown fill materials encountered during installation of geotechnical borings in July 2009 within the Project Area prior to property acquisition and/or the initiation of construction activities.

The CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area involves the realignment of Commercial Road between Canal Road and the proposed East Ninth Street Extension. In July 2009, geotechnical drilling crews installed a series of borings for the proposed realignment of Commercial Road. During installation of four (4) borings, drilling crews encountered unknown fill materials at depths of less than five (5) feet in three (3) borings and to a depth over 20 feet in one (1) boring. The unknown fill materials ranged in color from white to blue and odors were identified in several borings.

Phase II ESA activities were conducted in December 2009, and consisted of the installation of four (4) soil borings (designated HB-071, HB-072, HB-073 and HB-075) within the Project Area. The soil borings were installed adjacent to the geotechnical soil borings installed in July 2009. Subsurface materials encountered during boring installation consisted of non-native fill materials comprised of sandy clay, sand and sandy slag with varying amounts of gravel, slag, brick and wood fragments. Odors were identified at multiple depths in one (1) boring. No evidence of groundwater was encountered during boring installation.

Soil analytical results were compared against the Ohio Environmental Protection Agency's (EPA's) Voluntary Action Program (VAP) single-chemical Generic Direct Contact Soil Standards (GDCS) for commercial/industrial land use and construction/excavation activities and the State of Ohio's Bureau of Underground Storage Tank Regulations' (BUSTR) Petroleum-Contaminated Soil (PCS) Re-Use Levels. The regulatory standards referenced as part of this project are for comparative use only and may not be directly applicable to the Project Area.

Soil analytical results indicate that concentrations of several volatile organic compounds (VOCs), polynuclear aromatic hydrocarbon (PAH) compounds, one or more fractions of total petroleum hydrocarbons (TPH), total metals and/or total sulfide were detected in the soil samples collected within the Project Area. None of the concentrations of the constituents detected in soil samples exceeded the Ohio EPA's VAP GDCS for commercial/industrial land use or construction/excavation activities. However, concentrations of several constituents exceeded BUSTR's PCS Re-Use Levels, which consist of the following:

- ➤ Benzo(a)anthracene, benzo(a)pyrene, chrysene, indeno(1,2,3-cd)pyrene and naphthalene in HB-071 (8-10');
- ➤ Benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-071 (10-12');
- ➤ Indeno(1,2,3-cd)pyrene in HB-071 (16-18'), HB-072 (2-4') and HB-073 (0-2');
- ➤ Benzene, benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-072 (4-6'); and
- ➤ Benzo(a)anthracene, benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-075 (0-2') and HB-075 (2-4').

Therefore, based upon the above constituent concentrations exceeding BUSTR's PCS Re-Use Levels, a plan note should be included in construction documents for the proper handling, management and/or disposal of PCS in accordance with all applicable laws and regulations. The detected concentrations of total lead in HB-073 (0-2') and HB-075 (2-4') are 20 times the Toxicity Characteristic Leaching Procedure (TCLP) level for lead. In addition, although concentrations did not exceed VAP GDCS for commercial/industrial land use or construction/excavation activities, total cyanide was detected in six (6) soil samples and hexavalent chromium in one (1) soil sample. As a result, should construction plans/activities anticipate excavation and disposal of soils from these locations, properly characterize soils prior to disposal.

#### SIGNATURE PAGE

This Phase II Environmental Site Assessment report for the CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area (PID 77510) was prepared by HzW Environmental Consultants, LLC. Mr. Douglas M. Wetzel, Environmental Scientist, was the primary author of the report and Mr. John A. Zampino, Senior Geologist, reviewed the report. The signatures for Mr. Wetzel and Mr. Zampino are presented below.

Douglas M. Wetzel

**Environmental Scientist** 

#### PHASE II ENVIRONMENTAL SITE ASSESSMENT

CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area (PID 77510) Cleveland, Cuyahoga County, Ohio (HzW Project No. H09004-14)

#### 1.0 INTRODUCTION

The CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area (PID 77510) consists of realignment of Commercial Road between Canal Road and the proposed East Ninth Street Extension. A map showing the general location of the CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area is presented as **Figure 1**. The location of the proposed realignment of Commercial Road within the Project Area is presented in **Appendix A**.

In July 2009, geotechnical drilling crews installed a series of borings for the proposed realignment of Commercial Road. During installation of four (4) borings (designated B-071, B-072, B-073 and B-075, which are presented on the proposed realignment map of Commercial Road in Appendix A), drilling crews encountered materials described as:

- ➤ Having a "bad odor" or being an "unknown blue/white material" that "looked [corrosive]" to a depth of over 20 feet below ground surface in B-071;
- An "unknown blue material" at a depth of less than five (5) feet in B-072;
- An "unknown blue/green rock material" to a depth of less than five (5) feet in B-073; and
- A "nasty...blue/green material" from 1.5 to 4.5 feet in B-075, which "smells bad" and the drilling crew was "...not sure if [it was] safe to touch."

A review of historic Sanborn fire insurance maps provided to HzW by the Ohio Department of Transportation's (ODOT's) Office of Environmental Services (OES) indicated that the areas in question were historically occupied by railroad yards that were "full of tracks".

The purpose of the Phase II Environmental Site Assessment (ESA) investigation was to assess whether adverse subsurface impacts have occurred prior to property acquisition and/or the initiation of construction activities within the Project Area. Phase II ESA activities were conducted in accordance with the ODOT's April 2009 Environmental Site Assessment Guidelines for Phase II Environmental Site Assessments. Details regarding the sampling efforts and evaluation methods along with the corresponding findings of this study are presented in separate subsections of this report.

#### 2.0 BACKGROUND INFORMATION

The background geologic information pertaining to the Project Area is presented below.

#### 2.1 Physiography

According to the *Physiographic Regions of Ohio* map, published by the Ohio Department of Natural Resources (ODNR), the Project Area is located within the Erie Lake Plain of the Huron-Erie Lake Plains Section. The Erie Lake Plain is an Ice-Age lake basin separated from modern Lake Erie by shoreline cliffs and has major streams in deep gorges. Elevations of the Erie Lake Plain are generally of low relief between 570 and 800 feet. A copy of the Physiographic Map of Ohio is included as **Appendix B**.

#### 2.2 Topography

According to the 1994 Cleveland South, Ohio quadrangle 7.5-minute USGS topographic map, the topography of the Project Area varies from an elevation of approximately 630 feet above National Geodetic Vertical Datum (NGVD) in the southern portion to an elevation of 670 feet above NGVD in the northern portion. The locations in which soil borings were installed in the Project Area are nearly level. A copy of the 1994 Cleveland South, Ohio quadrangle USGS topographic map is included as **Appendix C**.

#### 2.3 Bedrock Geology

According to the *Geologic Map of Ohio*, published by the ODNR, bedrock beneath the Property consists of the Upper Devonian shales of the Cleveland, Chagrin and Huron groups. Bedrock is not exposed at the ground surface within the Project Area. A copy of the *Geologic Map of Ohio* is included as **Appendix D**.

#### 2.4 Bedrock Topography

According to the Revised 1996 Bedrock Topography Map of the USGS 7.5-minute Cleveland South, Ohio quadrangle published by the ODNR, bedrock beneath the Project Area is at an elevation of approximately 450 feet above NGVD. Bedrock within the vicinity of the Project Area slopes east/southeast. A copy of the Bedrock Topography Map is included as **Appendix E**.

#### 2.5 Glacial Geology

According to the *Glacial and Surficial Geology Map of Cuyahoga County*, published by the ODNR, the subsurface beneath the Project Area consists of Made Land. The areas of Made Land consist of reclaimed land, cut and fill, dumps, and continuous urban cover where 90 percent or more of the surface is covered with concrete, asphalt, building complexes, structures, or other manmade surfaces. A copy of the *Glacial and Surficial Geology Map of Cuyahoga County* is included as **Appendix F**.

#### 2.6 Hydrology

According to the *Principal Streams and Their Drainage Area Map*, published by the ODNR, the Project Area is located within the 809 square mile drainage basin of the Cuyahoga River. A copy of the *Principal Streams and Their Drainage Areas Map* is included as **Appendix G**.

#### 2.7 Hydrogeology

Based on local topographic conditions, local groundwater flow beneath the Project Area is anticipated to flow to the south/southwest towards the Cuyahoga River. However, actual groundwater flow direction is often influenced by factors such as underground structures, seasonal fluctuations, soil and bedrock geology, production wells and other factors beyond the scope of this study. According to the *Ground Water Resources Map of Cuyahoga County, Ohio*, published by the ODNR, two (2) hydrogeologic units underlie the Project Area. The majority of the Project Area is underlain by buried valleys that contain 200 to 300 feet of fine sand, silt and clay. Wells drilled within the buried valleys yield meager supplies (generally 3 to 10 gallons of groundwater per minute) unless thin, isolated sand and gravel lenses are encountered. The remaining portion of the Project Area is underlain by permeable sand and gravel deposits interbedded with silt and clay in a buried valley. Drilled wells within the permeable sand and gravel unit yield 100 to 300 gallons of groundwater per minute with yields as much as 250 gallons of groundwater per minute in locations where sufficient coarse material is present. A copy of the *Ground Water Resources Map of Cuyahoga County* is included in **Appendix H**.

#### 2.8 Soils

According to the *Soil Survey of Cuyahoga County, Ohio*, published by the United States Department of Agriculture, the Project Area is underlain by one (1) soil type, Urban land (Ub). The Urban land soil type consists of areas in where more than 80 percent of the surface is covered by asphalt, concrete, buildings or other manmade surfaces. A copy of the *Soil Survey* is included as **Appendix I**.

#### 2.9 Oil and Gas Wells

According to the Oil and Gas Well Map for the Cleveland South, Ohio 7.5-minute quadrangle published by the ODNR, no oil or gas wells are located within the Project Area. A copy of the Oil and Gas Well Map is included as **Appendix J**.

#### 3.0 FIELD ACTIVITIES AND SAMPLING PROCEDURES

#### 3.1 Sampling Methods

Phase II ESA activities were conducted on December 17, 2009, and consisted of the installation of four (4) soil borings (designated "HB-071," "HB-072," "HB-073" and "HB-075"). The soil borings were installed adjacent to the geotechnical soil borings installed in July 2009 and the designation of each soil boring was kept consistent with the geotechnical soil borings (e.g., HzW soil boring HB-071 was installed adjacent to geotechnical soil boring B-071). Visual evidence of two (2) geotechnical soil borings (HB-071 and HB-072) remained, which consisted of wooden stakes with the respective boring numbers and the auger hole. However, no visual evidence of the remaining two (2) geotechnical boring locations (HB-073 and HB-075) was identified. Therefore, in order to locate HB-073 and HB-075 and to verify the locations of HB-071 and HB-072, HzW utilized a Trimble® GeoXH<sup>TM</sup> Global Positioning System unit to locate the state plane coordinates for each boring, which were included on the geotechnical soil borings logs provided by ODOT OES. The locations of the soil borings on the Property are depicted on **Figure 2**.

All personnel on-site, or otherwise associated with the sample collection, were trained in accordance with Occupation Safety and Health Administration (OSHA) requirements, as stipulated under 29 CFR 1910.120. A copy of the site-specific Health and Safety Plan (HASP) prepared for use by all on-site personnel is included as **Appendix K**.

#### 3.2 Soil Boring Installation and Sample Collection

Soil samples were collected from ground surface to terminal depths. The soil borings were installed using hydraulic Geoprobe<sup>®</sup> direct push subsurface sampling techniques. Hydraulic Geoprobe<sup>®</sup> marco core direct push subsurface sampling techniques utilize a series of 5-foot long steel rods driven into the subsurface. Soil samples were collected using a five-foot long sample tube attached to five-foot long steel rods. The sample tube consists of a clean, disposable acetate (plastic) liner that is driven into the subsurface to obtain a core sample of the subsurface material.

Upon extraction from the soil, the plastic liner, with core intact, was removed from the sample tube. Each sample liner was initially split into two-foot intervals following sample collection and examined separately. Each two-foot soil sample was transferred to a clean, labeled sample container (provided by the laboratory) and placed in an ice cooler for preservation in the field. The sample intervals were characterized by a qualified environmental technician. Observations noted by the technician included the sample location/number, sample depth, sediment description, color, moisture content, odor, and presence or absence of contamination based on visual/olfactory observation. The observations were recorded on a boring log completed for each soil boring.

Following completion of soil sampling activities, all borings installed within the Project Area were filled with granular bentonite and hydrated. All equipment used during Phase II ESA sampling activities was decontaminated with a Liqui-Nox® and distilled water solution and triplerinsed with distilled water after each use to limit the potential for cross contamination.

#### 3.3 Sample Selection Methods

Soil samples were selected for laboratory analysis based upon visual and olfactory observations in the field and depths at which unknown materials were encountered by the geotechnical drilling crews as recorded in the geotechnical soil boring logs. The samples were shipped in a sample cooler, chilled to 4°C, under chain-of-custody documentation.

#### 3.4 Analytical Methods

The soil samples from each boring were submitted to TestAmerica Laboratories, Inc. (TestAmerica) of North Canton, Ohio, for analysis of volatile organic compounds (VOCs) by EPA Method 8260, polynuclear aromatic hydrocarbons (PAHs) by EPA Method 8270, total petroleum hydrocarbons (TPH) gasoline and diesel range organics by modified EPA Method 8015, total concentrations of arsenic, cadmium, chromium and lead by EPA Method 6010, total concentrations of cyanide by EPA Method 335.2, hexavalent chromium by EPA Method 7196, total sulfide by EPA Methods 9030/9034 and corrosivity (pH) by EPA Method 9045.

#### 3.5 Quality Assurance/Quality Control

HzW developed Quality Assurance/Quality Control (QA/QC) measures to ensure the Phase II ESA was conducted in accordance with consistent professional standards and specific ODOT requirements. The main elements of the QA/QC program include the following:

- Peer review of all project correspondence, notes, chain-of-titles, etc;
- ➤ Multi-layered report examination by the QA/QC team;
- > Daily involvement by HzW's project manager in all aspects of the project; and
- Regular discussion with technical personnel to review elements to be included in the reports; report format changes, and internal routing/review procedures.

Specific QA/QC procedures for this project included the following elements:

- 1. Review of the project Scope-of-Services between the HzW Project Manager and the field representatives;
- 2. Continual reference to the Phase II proposal by the field representatives during project implementation;
- 3. During field work, completion of thorough and accurate field notes (such as site drawings, information obtained from the property representative, and observations made during the site assessment);
- 4. During data evaluation, identifying appropriate and applicable reference standards with which to compare results, comparing results to field observations to identify any discrepancies; and
- 5. During report preparation:
  - completion of report in a format required by ODOT,
  - review of report by the author,
  - editing report, as needed,
  - review and editing of report by the Technical Editor, as needed,
  - review of report by the Project Manager, and
  - finalization of the report.

#### 4.0 PHASE II FINDINGS, DATA EVALUATION & REGULATORY INTERPRETATION

The findings of this Phase II ESA investigation are presented below. A discussion of the geology and hydrology of the Project Area based on soil boring logs, and the soil sample analytical results, are presented in separate subsections below.

#### 4.1 Boring Log Descriptions

Soil borings within the Project Area were installed to terminal depths of six (6) or 20 feet below ground surface (bgs). Subsurface materials encountered during boring installation consisted of non-native fill materials including brown, black and dark gray sandy clay, dark gray, black and brown sand, and dark brown, black and dark gray sandy slag with varying amounts of gravel, slag, brick and wood fragments. Slight to strong sulfur odors were identified in HB-071 (6-14' and 16-18'). No odors were identified in the remaining soil borings. Although damp soils were encountered in HB-073 and HB-075 (both of which terminated at six [6] feet bgs), no evidence of groundwater was encountered during boring installation. Copies of the boring logs for the borings installed within the Project Area are included as **Appendix L**.

#### 4.2 Soil Analytical Results

The analytical results of the soil samples collected from soil borings installed within the Project Area (HB-071, HB-072, HB-073 and HB-075) are presented in **Table 1**. The complete laboratory analytical report from TestAmerica, including the laboratory's internal QA/QC sample results, is included in **Appendix M**. The laboratory QA/QC reporting was determined to be acceptable with the exception of the following:

- ➤ The matrix spike/matrix spike duplicate (MS/MSD) for a batch of samples for VOC analysis had relative percent differences (RPDs) and recoveries outside acceptance limits. However, since the associated method blank and laboratory control sample was in control, no corrective action was necessary.
- ➤ The internal standard areas for VOC analysis were outside acceptance limits for samples HB-071 (10-12'), HB-071 (16-18') and HB-072 (2-4') due to matrix effects.
- ➤ Sample HB-071 (8-10') had elevated reporting limits due to tentatively identified compounds in the VOC analysis.
- ➤ Samples HB-071 (10-12'), HB-071 (16-18') and HB-072 (2-4') were reanalyzed at a dilution due to internal standard recoveries outside of acceptance limits, per Ohio VAP standards. Only compounds associated with internal standards that met criteria are reported from each analysis.
- ➤ Sample HB-075 (0-2') had elevated reporting limits in SVOC analysis due to matrix interferences.
- ➤ The batch QC for SVOC analysis batch 9353017 was unable to be reported due to repreparation of the parent sample and the original sample was not reported.
- ➤ The MS/MSD for a batch of samples for TPH analysis had recoveries outside acceptance limits. However, since the associated method blank and laboratory control sample was in control, no corrective action was necessary.
- ➤ The reported concentration of a TPH fraction in HB-075 (0-2') is flagged with a "J" as the results were between the method detection limit and the reporting limit. The possibility of false positive or mis-identification at these quantification levels exist. In analytical methods requiring confirmation of the analyte reported, confirmation was

- performed only down to the standard reporting limit. The acceptance criteria for QC samples may not be met at these quantification levels.
- ➤ The generic batch MS/MSDs for a TPH analysis batch was extracted and analyzed, but unable to be reported due to the laboratory system limitations.
- Sample HB-072 (4-6') for hexavalent chromium is suspect to have a reducing agent based on the results obtained from method of standard addition.
- ➤ The MS/MSD data for certain batches are not included in this report. The batch QC samples, which document the effect of a specific sample matrix on method performance, were not associated with a sample reported in this lot. The data, therefore, has no bearing on samples reported herein. In order to document compliance with the QC requirement for an MS/MSD per 20 environmental samples, a summary of sample/QC associations has been provided.

#### 4.3 Data Evaluation Criteria/Regulatory Discussion

Soil analytical results were compared against the Ohio Environmental Protection Agency's Voluntary Action Program (VAP) single-chemical Generic Direct Contact Soil Standards (GDCS) for commercial/industrial land use and construction/excavation activities and the State of Ohio's Bureau of Underground Storage Tank Regulations' (BUSTR) Petroleum-Contaminated Soil (PCS) Re-Use Levels. The GDCS values were developed to be protective of the environment and human health based on predictive models regarding potential exposures to adults from dermal contact with soil, inhalation of vapors and particles from soil, and ingestion of soil. The VAP and BUSTR comparative standards are included with the soil analytical results in **Table 1**.

The regulatory standards referenced as part of this project are for comparative use only and may not be directly applicable to the Project Area. The Ohio VAP and BUSTR standards referenced in this report apply only to sites that are participants in Ohio's Voluntary Action Program or sites regulated by BUSTR. However, because the Ohio Environmental Protection Agency and BUSTR recognize these standards as being protective of human health and the environment, they provide a useful tool for assessing environmental conditions within the Project Area.

According to soil analytical results, low concentrations of several VOCs consisting of acetone, benzene, carbon disulfide, ethylbenzene, methylene chloride, toluene and/or total xylenes were detected in soil samples HB-71 (10-12'), HB-71 (16-18'), HB-072 (2-4'), HB-072 (4-6'), HB-073 (0-2') and HB-075 (0-2'). None of the detected concentrations of VOCs exceeded VAP GDCS for commercial/industrial land use or construction/excavation activities. However, the detected concentration of benzene in HB-072 (4-6'), 0.027 milligrams per kilogram (mg/kg), exceeded the BUSTR PCS Re-Use Level of 0.015 mg/kg.

Concentrations of PAH compounds consisting of acenaphthene, acenaphthylene, anthracene, benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, benzo(g,h,i)perylene, benzo(k)fluoranthene, chrysene, dibenz(a,h)anthracene, fluoranthene, fluorene, indeno(1,2,3-cd)pyrene, 1-methylnaphthalene, 2-methylnaphthalene, naphthalene, phenanthrene and/or pyrene were detected in soil samples from all soil borings. Although none of the detected concentrations of PAH compounds exceeded VAP GDCS for commercial/industrial land use or construction/excavation activities, concentrations of several PAH compounds exceeded BUSTR PCS Re-Use Levels. The compounds exceeding Re-Use Levels consists of benzo(a)anthracene, benzo(a)pyrene, chrysene, indeno(1,2,3-cd)pryene and naphthalene in HB-071 (8-10'); benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-071 (10-12'); indeno(1,2,3-cd)pryene in HB-071 (16-18'), HB-072 (2-4') and HB-073 (0-2'); benzo(a)pyrene, chrysene and

indeno(1,2,3-cd)pyrene in HB-072 (4-6'); and benzo(a)anthracene, benzo(a)pryene, chrysene and indeno(1,2,3-cd)pyrene in HB-075 (0-2') and HB-075 (2-4').

Soil analytical results indicate that concentrations of one or more fractions of TPH were detected in the soil samples from all soil borings. None of the detected concentrations of TPH fractions exceeded VAP GDCS for commercial/industrial land use or construction/excavation activities or BUSTR PCS Re-Use Levels.

Concentrations of total arsenic, cadmium, chromium and/or lead were detected in soil samples from all soil borings. None of the detected concentrations of total metals exceeded VAP GDCS for commercial/industrial land use or construction/excavation activities. However, the detected concentrations of lead in HB-073 (0-2') and HB-075 (2-4'), 115 mg/kg and 107 mg/kg, respectively, are 20 times the Toxicity Characteristic Leaching Procedure (TCLP) level for lead (5.0 mg/kg).

According to soil analytical results, concentrations of total cyanide were detected in HB-071 (8-10'), HB-071 (10-12'), HB-071 (16-18'), HB-072 (2-4'), HB-072 (4-6') and HB-073 (0-2') and hexavalent chromium detected in HB-072 (4-6'). The detected concentrations of total cyanide or hexavalent chromium did not exceed VAP GDCS for commercial/industrial land use or construction/excavation activities.

No comparative standards currently exist for total sulfide or corrosivity (pH). Total sulfide concentrations in soil samples ranged from below detection limits in HB-075 (2-4') to 302 mg/kg in HB-073 (0-2'). Concentrations of pH ranged from 8.7 in HB-075 (2-4') to 11.5 in HB-075 (0-2').

#### 5.0 CONCLUSIONS AND RECOMMENDATIONS

The results of Phase II ESA activities conducted within the CUY-Cleveland Innerbelt, Commercial Road Alignment Project Area (PID 77510) indicate that concentrations of several VOCs, PAH compounds, one or more fractions of TPH, several total metals and total sulfide were detected in soil samples collected within the Project Area. None of the concentrations of the constituents detected in soil samples within the Project Area exceeded VAP GDCS for commercial/industrial land use or construction/excavation activities. However, concentrations of several constituents exceeded BUSTR's PCS Re-Use Levels, which consist of the following:

- ➤ Benzo(a)anthracene, benzo(a)pyrene, chrysene, indeno(1,2,3-cd)pyrene and naphthalene in HB-071 (8-10');
- ➤ Benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-071 (10-12');
- Indeno(1,2,3-cd)pyrene in HB-071 (16-18'), HB-072 (2-4') and HB-073 (0-2');
- ➤ Benzene, benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-072 (4-6'); and
- ➤ Benzo(a)anthracene, benzo(a)pyrene, chrysene and indeno(1,2,3-cd)pyrene in HB-075 (0-2') and HB-075 (2-4').

Based upon the above constituent concentrations exceeding BUSTR's PCS Re-Use Levels, a plan note should be included in construction documents for the proper handling, management and/or disposal of PCS in accordance with all applicable laws and regulations.

The detected concentrations of total lead in HB-073 (0-2') and HB-075 (2-4') are 20 times the TCLP level for lead. In addition, although concentrations did not exceed VAP GDCS for commercial/industrial land use or construction/excavation activities, total cyanide was detected in six (6) soil samples and hexavalent chromium in one (1) soil sample. As a result, should construction plans/activities anticipate excavation and disposal of soils from these locations, properly characterize soils prior to disposal.

TABLES

#### Table 1 **Summary of Soil Analytical Results** Commercial Road Alignment Project Area (PID 77510)

### Cleveland, Ohio

(All values presented in mg/kg, unless otherwise noted)

Sample ID		HB-071 (10-12')	HB-071 (16-18')	HB-072 (2-4')	HB-072 (4-6')	HB-073 (0-2')	HB-075 (0-2')	HB-075 (2-4')		1	
Sample Date	12/17/2009	12/17/2009	12/17/2009	12/17/2009	12/17/2009	12/17/2009	12/17/2009	12/17/2009	CIGDCS <sup>1</sup>	CEGDCS <sup>2</sup>	BUSTR Re-Use <sup>6</sup>
VOCs - EPA Method 8260		0.00									
Acetone	<23.0	0.036	< 0.023	< 0.023	< 0.023	<0.023	< 0.023	< 0.022	100,000	100,000	
Benzene	<5.7	< 0.006	< 0.006	< 0.006	0.027	<0.006	< 0.006	< 0.006	140	150	0.015
Carbon disulfide	<5.7	0.014	0.012	0.008	0.027	< 0.006	< 0.006	< 0.006	1,400	190	
Ethylbenzene	<5.7	< 0.006	< 0.028	< 0.028	< 0.006	0.011	< 0.006	< 0.006	230	230	4.55
Methylene chloride	<5.7	0.012	0.007	0.011	< 0.006	< 0.006	< 0.006	< 0.006	570	1,500	
Toluene	<5.7	0.007	< 0.028	< 0.028	0.044	0.006	< 0.006	< 0.006	520	520	4.91
Total Xylenes	<5.7	0.011	< 0.028	< 0.028	0.02	0.047	0.026	< 0.006	370	370	15.7
PAHs - EPA Method 8270											
Acenaphthene	7.0	1.3	0.9	0.11	0.14	0.082	0.19	0.57	56,000	440,000	
Acenaphthylene	< 0.38	0.27	0.12	0.14	0.34	0.15	0.1	0.072	170,000 <sup>3</sup>	51,000 4	
Anthracene	2.4	0.94	0.49	0.36	0.41	0.24	0.86	1.3	280,000	1,000,000	
Benzo(a)anthracene	2.2	1.5	0.81	0.84	1.3	1.1	2.9	2.9	76	680	2.2
Benzo(a)pyrene	2.2	1.3	0.66	0.63	1.2	1.0	2.1	2.5	7.7	69	1.1
Benzo(b)fluoranthene	3.3	2.2	1.0	1.1	2.1	1.5	2.5	3.2	77	690	5.53
Benzo(g,h,i)perylene	0.83	0.96	0.45	0.5	0.85	0.82	1.2	1.6	23,000 <sup>3</sup>	25,000 <sup>4</sup>	
Benzo(k)fluoranthene	0.57	0.83	0.45	0.39	0.75	0.82	0.94	1.0	770	6,900	1.97
Chrysene	2.2	1.5	0.42	0.84	1.5	0.42	2.6	2.6	7,600	69,000	1.27
Dibenz(a,h)anthracene	<0.38	0.38	0.18	0.15	0.26	0.29	0.46	0.5	7,000	69	0.94
Fluoranthene	8.8	4.0	2.2	1.5	2.1	1.6	5.1	5.5	37,000	290,000	0.94
	5.3	1.3	0.87	0.14	0.18	0.087	0.21	0.57		290,000	
Fluorene	1.4	0.94	0.44	0.46	0.76	0.6	1.1	1.5	37,000 77	690	0.15
Indeno(1,2,3-cd)pyrene 1-Methylnaphthalene	4.1	2.0	0.96	0.52	0.86	1.0	0.15	0.61	360	360	0.13
2-Methylnaphthalene	7.9	3.3	1.6	0.78	1.4	1.5	0.18	1.0	94,000 3	62,000 4	
Naphthalene	55.0	3.7	3.4	0.58	0.91	0.88	0.23	0.82	150	84	3.98
Phenanthrene	13.0	5.4	2.7	1.1	1.4	1.2	2.5	4.8	870,000 <sup>3</sup>	260,000 4	
Pyrene	5.6	3.1	1.8	1.4	2.1	1.5	6.1	5.4	28,000	220,000	
TPH - Modified EPA Method 8015											
C6-C12	2.7	0.17	0.15	< 0.11	0.7	0.21	0.78	< 0.11	1,000 5	1,000 5	1,000
C10-C20	260	170	72	44	82	120	100	48	2,000 5	2,000 5	2,000
C20-C34	250	220	150	140	220	130	610	110	5,000 5	5,000 5	5,000
Total Metals - EPA Method 6010											
Arsenic	10.0	11.3	9.7	8.6	9.3	9.3	4.7	10.4	82	420	
Cadmium	<0.23	0.31	<0.23	< 0.23	<0.23	0.33	0.23	<0.22	2,300	1,600	
	11.2	14.4	<0.23 <b>22.9</b>	<0.23 <b>10.6</b>	19.7	10.6	18.6	<b>8.9</b>	1,000,000		
Chromium Lead	38.8	65.9	55.7	27.0	38.4	115	21.3	107	1,800	1,000,000 750	
Leau	30.0	03.9	33.1	27.0	30.4	113	21.3	107	1,000	730	
Total Cyanide - EPA Method 335.2	0.8	2.1	0.73	2.9	2.1	4.6	< 0.57	< 0.55	59,000	39,000	
Hexavalent Chromium - EPA Method 7196	< 0.92	<0.92	<0.91	< 0.9	2.5	<0.9	<0.91	< 0.88	7,900	13,000	
Total Sulfide - EPA Methods 9030/9034	48.5	90.2	148	287	289	302	61.6	<33.1	NGS	NGS	
Corrosivity - EPA Method 9045 (unitless)	9.5	10.1	10.4	10.2	10.2	11.0	11.5	8.7	NGS	NGS	

**Bolded** values indicate laboratory detections

<sup>&</sup>lt;sup>1</sup>VAP Generic Direct Contact Soil Standards for Commercial/Industrial Land Use per OAC 3745-300-08(C)(3)(c), Table II

<sup>&</sup>lt;sup>2</sup>VAP Generic Direct Contact Soil Standards for Construction and Excavation Activities per OAC 3745-300-08(C)(3)(d), Table III

<sup>&</sup>lt;sup>3</sup>VAP Supplemental Direct Contact Soil Values for Commercial/Industrial Land Use, January 25, 2006

<sup>&</sup>lt;sup>4</sup>VAP Supplemental Direct Contact Soil Values for Construction and Excavation Activities, January 25, 2006

<sup>&</sup>lt;sup>5</sup>Petroleum standards are BUSTR standards per VAP Technical Guidance Compendium VA30008.09.001

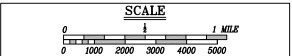
<sup>&</sup>lt;sup>6</sup>BUSTR's Petroleum Contaminated Soil Re-Use Levels per OAC 1301:7-9-16(D)(1), Table 1

<sup>=</sup> Value exceeds BUSTR's PCS Re-Use Levels

<sup>=</sup> J-qualified analytical results

		<b>FIGURES</b>

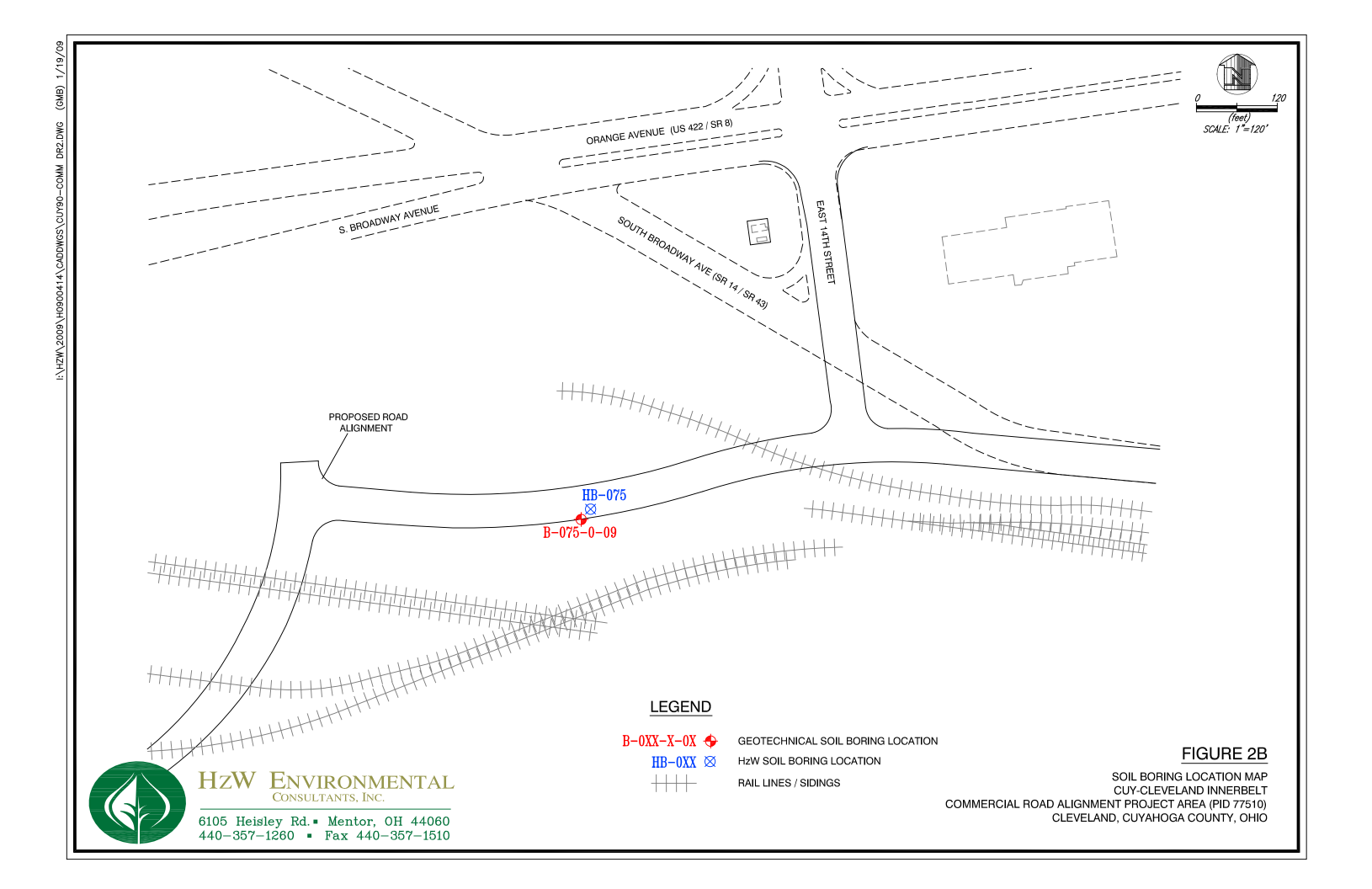




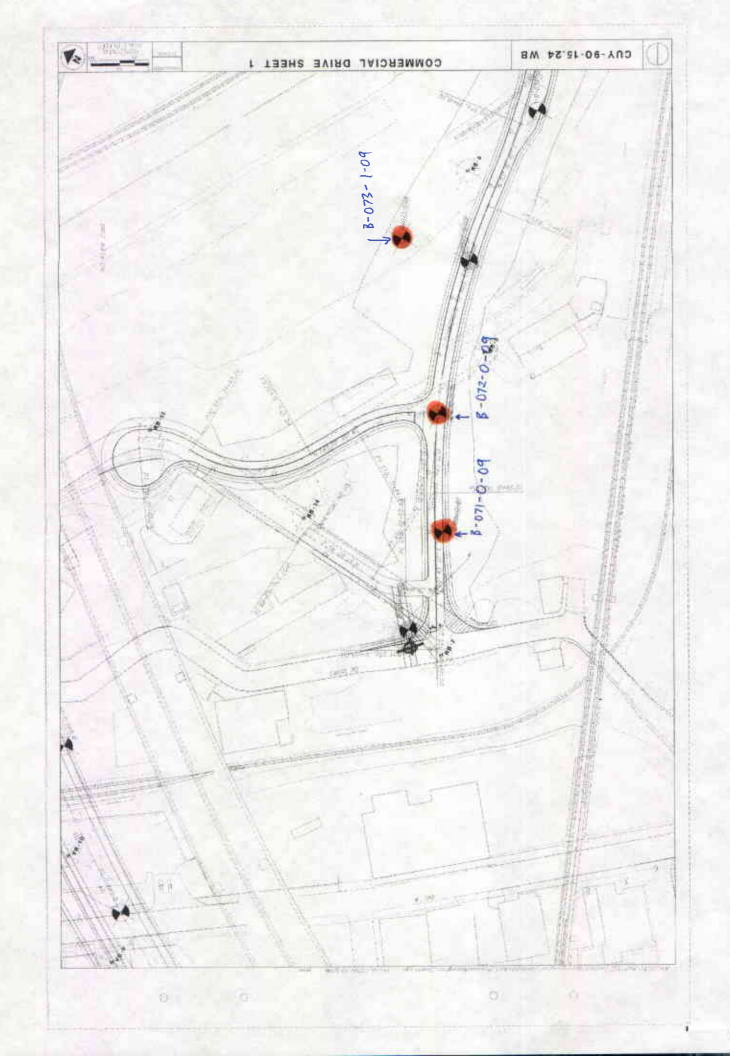


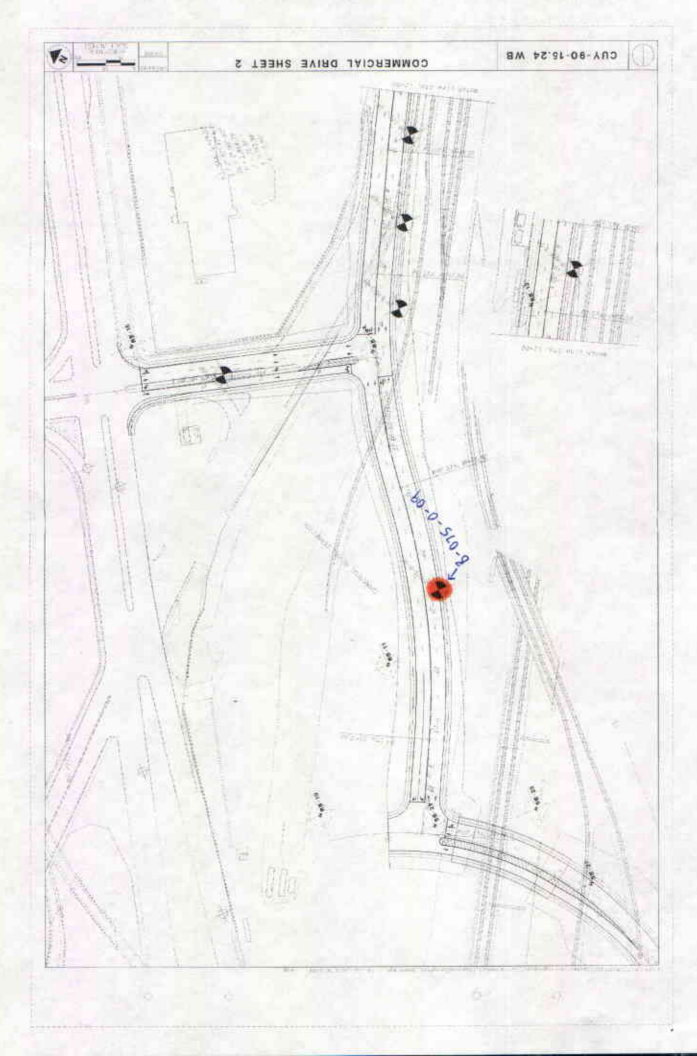
### FIGURE 1

SITE LOCATION MAP CUY-CLEVELAND INTERBELT COMMERCIAL ROAD ALIGNMENT PROJECT AREA (PID 77510) CLEVELAND, CUYAHOGA COUNTY, OHIO



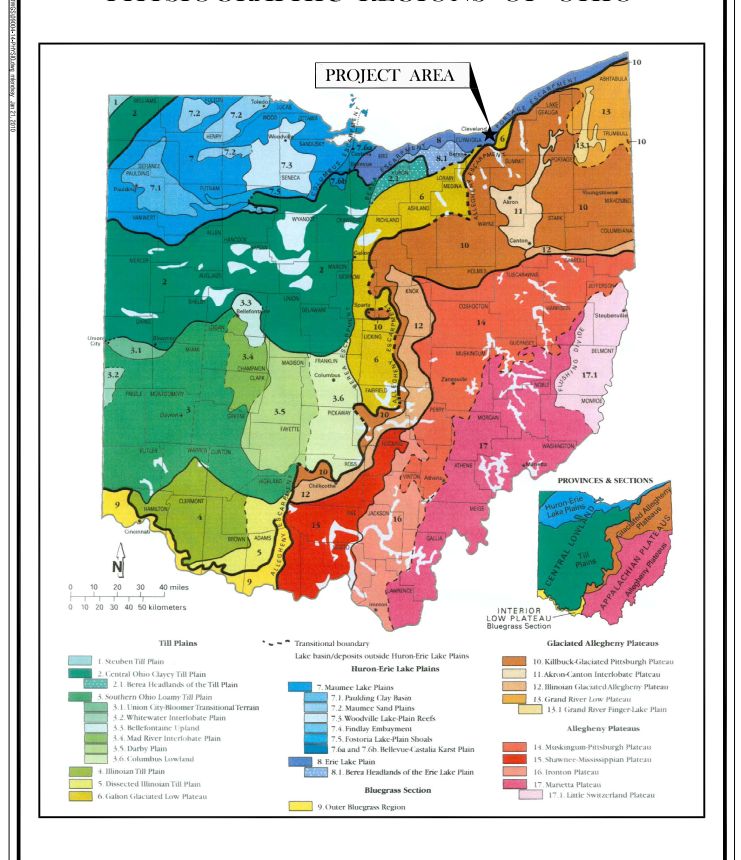
# APPENDIX A PROJECT PLANS





# APPENDIX B PHYSIOGRAPHIC MAP OF OHIO

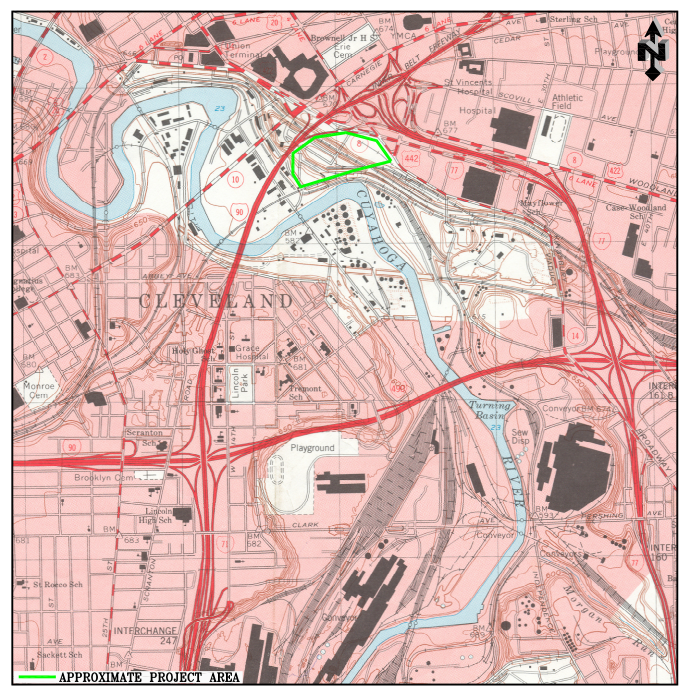
# PHYSIOGRAPHIC REGIONS OF OHIO

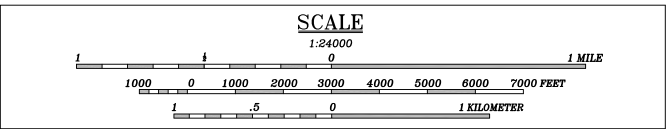




# APPENDIX C USGS TOPOGRAPHIC MAP

# USGS TOPOGRAPHIC MAP 1994 CLEVELAND SOUTH, OHIO QUADRANGLE

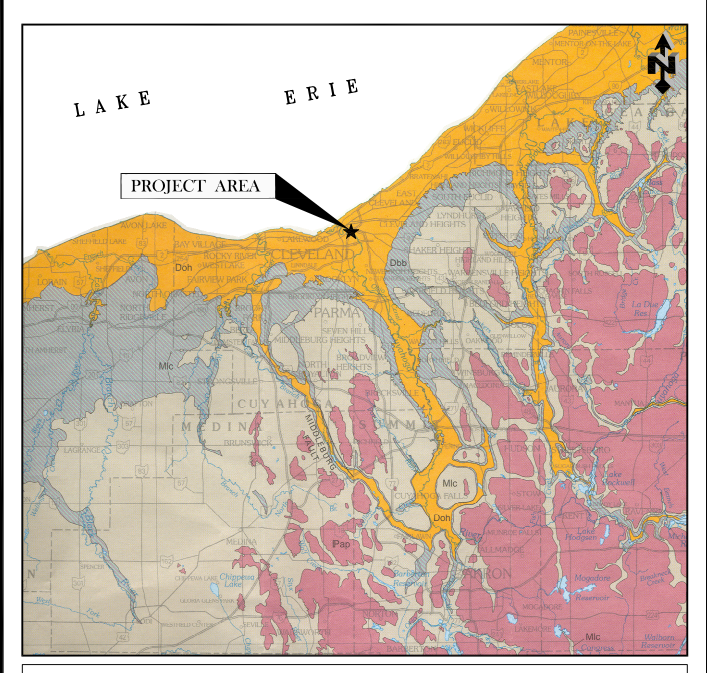






# APPENDIX D GEOLOGIC MAP OF OHIO

# 2006 BEDROCK GEOLOGIC MAP OF OHIO



## **LEGEND**

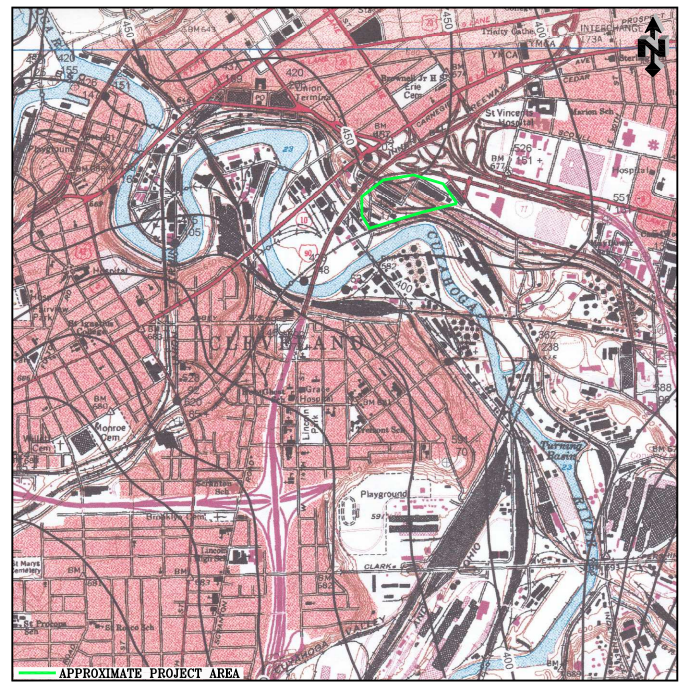
Doh

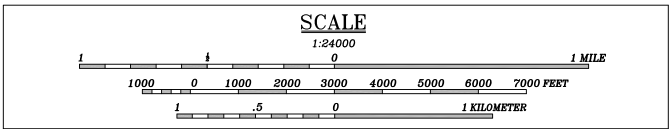
Ohio Shale (Upper Devonian) (mapped interval includes Olentangy Shale south of central Delaware County)—Unit consists generally of three members, in descending order: Cleveland, Chagrin, and Huron Members. Cleveland Member, shale; black; thickest in north-central portion of state; thins south and eastward; absent in northeastern portion of state. Chagrin Member, shale, siltstone, and very fine-grained sandstone; gray to greenish gray; thickest in northeastern portion of state; grades into underlying and overlying black shale members; thins southwestward, becomes Three Lick Bed in southern portion of state. Huron Member, shale; mostly black; carbonaceous; calcareous concretions common in lower portion. Clentangy Shale, mostly upper portion; thin; present but not mapped as separate unit south of central Delaware County; absent on Bellefontaine Outlier; see Olentangy Shale for description. Unit structurally deformed in Serpent Mound Impact Structure (see fig. 3).



# APPENDIX E BEDROCK TOPOGRAPHY MAP

## BEDROCK TOPOGRAPHY MAP 1996 CLEVELAND SOUTH, OHIO QUADRANGLE

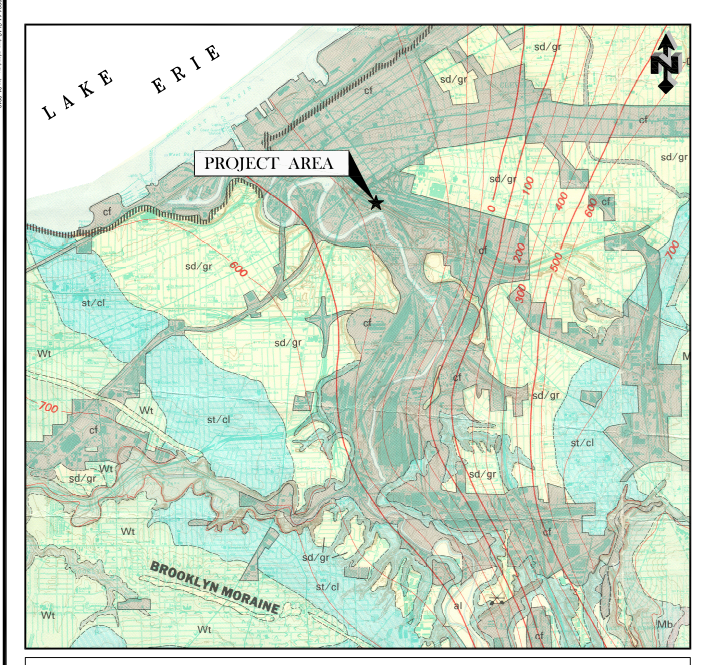






# APPENDIX F GLACIAL GEOLOGY MAP OF CUYAHOGA COUNTY

# GLACIAL & SURFICIAL GEOLOGY MAP OF OHIO



### **LEGEND**

cf

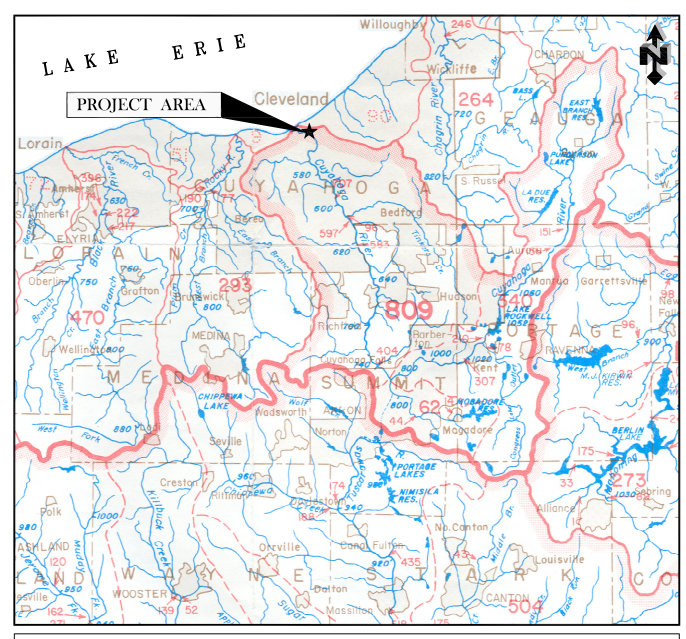
Made land. Areas of reclaimed land, cut and fill, dumps, and continuous urban cover where 90 percent or more of the surface is covered with concrete, asphalt, building complexes, structures, or other manmade surfaces; does not include urbanized areas where manmade surfaces are intricately associated with other types of cover



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# PRINCIPAL STREAMS & THEIR DRAINAGE AREAS MAP OF OHIO

# PRINCIPAL STREAMS AND DRAINAGE AREAS



# **LEGEND**

Areas of drainage basins, in square miles, are shown by red figures as follows:

Areas enclosed by shaded red lines 1757 Areas enclosed by unshaded red lines 313

Drainage areas above points indicated by arrows

Auxiliary land areas within the limits of the State.



Approximate low-water elevation in feet above sea level.



Reservoir

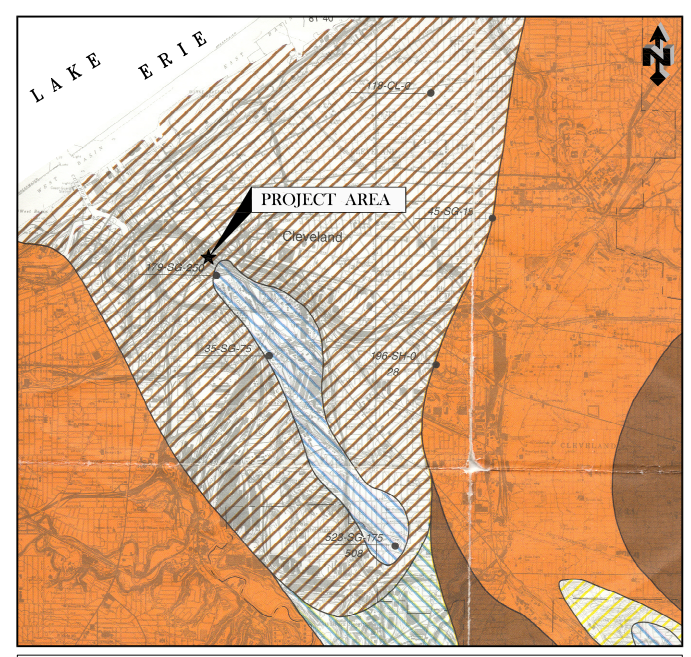


Selected urbanized areas, communities, and county seats chosen to aid in map orientation.



APPENDIX H
GROUND WATER RESOURCES MAP OF CUYAHOGA COUNTY

# GROUND WATER RESOURCES MAP 1992 CUYAHOGA COUNTY, OHIO



# **LEGEND**

#### AREAS IN WHICH 100 TO 300 GALLONS PER MINUTE MAY BE DEVELOPED



Good ground water areas. Permeable sand and gravel deposits interbedded with silt and clay lie in a buried valley. Yields of as much as 250 gallons per minute are available where sufficient coarse material is found. Exploratory drilling may be required to locate such deposits.

#### AREAS IN WHICH 3 TO 10 GALLONS PER MINUTE MAY BE DEVELOPED

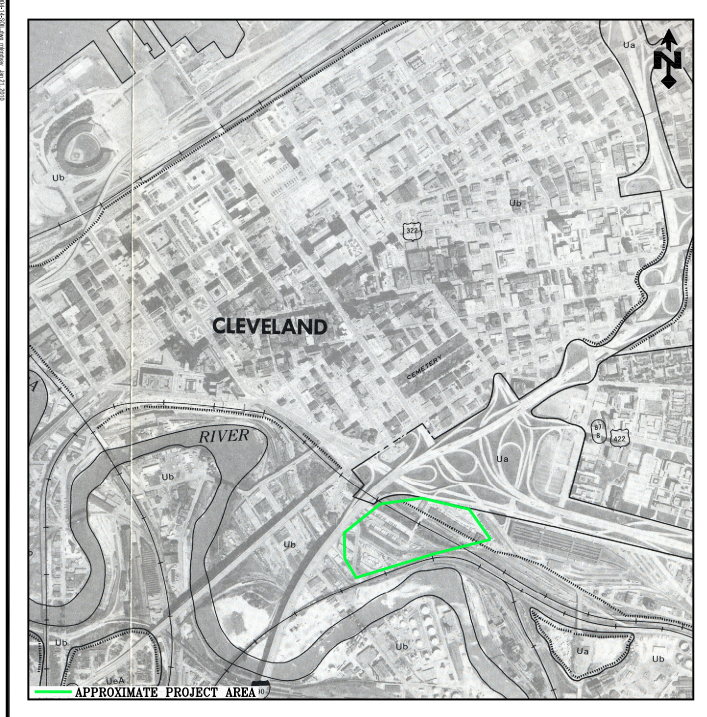


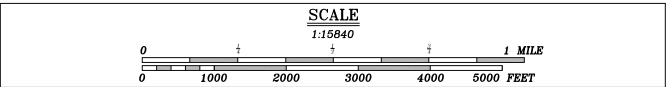
Buried valleys contain 200 to 300 feet of fine sand, silt, and clay. Drilled wells yield meager supplies unless encountering thin, isolated sand and gravel lenses.



# APPENDIX I SOIL SURVEY OF CUYAHOGA COUNTY

# SOIL SURVEY MAP 1980 CUYAHOGA COUNTY, OHIO

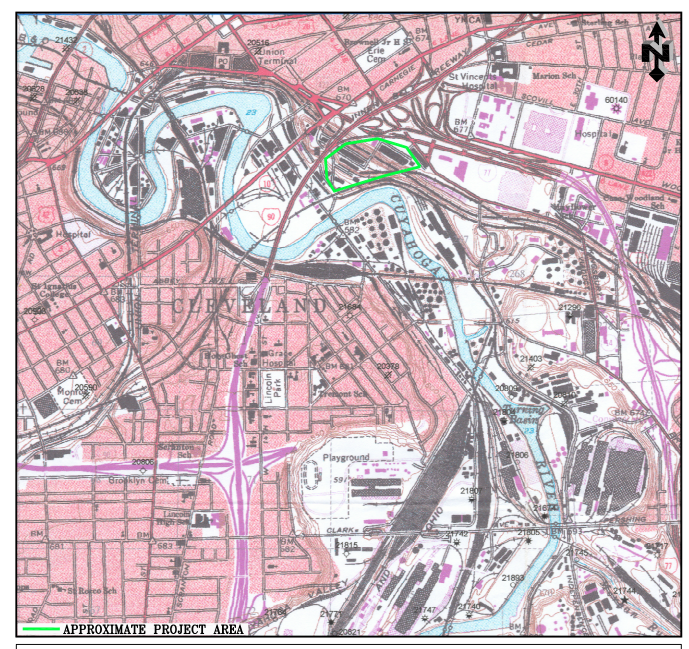






# APPENDIX J OIL AND GAS WELL MAP

# OIL AND GAS WELL MAP 2006 CLEVELAND SOUTH, OHIO QUADRANGLE



- **★** UNKNOWN STATUS
- □ BRINE FOR DUST CONTROL
- ☆ COALBED METHANE
- → DRY HOLE
- → DRY HOLE WITH GAS SHOW
- ightarrow DRY HOLE WITH OIL AND GAS SHOW
- ♦ DRY HOLE WITH OIL SHOW
- Ø EXPIRED PERMIT LOCATION
- ☆ GAS
- imes GAS WITH OIL SHOW
- → GAS SHOW

#### WELL SYMBOLS

- ₩ GAS AND OIL SHOW
- ♦ INJECTION
- → LOST HOLE
- ♥ OBSERVATION
- OIL AND GAS CONVERTED TO WATER
- OIL
- ☀ OIL AND GAS
- ₩ OIL WITH GAS SHOW
- OIL SHOW
- O PERMITTED LOCATION
- ⋈ PLUGGED BRINE FOR DUST CONTROL

- \* PLUGGED GAS WITH OIL SHOW

- \* PLUGGED OIL AND GAS
- → PLUGGED OIL WITH GAS SHOW
- RADIOACTIVE TOOL LOST IN HOLE
- \$ SOLUTION MINING
- GAS STORAGE
- X STRATIGRAPHY TEST
- WATER SUPPLY



# APPENDIX K SITE SPECIFIC HEALTH AND SAFETY PLAN

# SITE-SPECIFIC HEALTH AND SAFETY PLAN

# CUY-CLEVELAND INNERBELT COMMERCIAL ROAD ALIGNMENT (PID 77510) CLEVELAND, CUYAHOGA COUNTY, OHIO

December 2009

Prepared by
HzW Environmental Consultants, LLC
6105 Heisley Road
Mentor, Ohio 44060

# EMPLOYEE ACKNOWLEDGEMENT FORM

I have read this site-specific health and safety plan, understand the material presented, have been given an opportunity to ask questions and will abide by the provisions stated in this site-specific plan under which this project is to be implemented.

Site Manager:			
(Optional)	 		
HzW Project Manager:	 		
Contractor's Site Supervisor:(Optional)		· · · · · · · · · · · · · · · · · · ·	
Field Technician:	 		
Field Technician:			7.34
Field Technician:			
Field Technician:			

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# **HEALTH AND SAFETY PLAN**

#### 1.0 OBJECTIVE

The objective of this Health and Safety Plan (HASP) is to identify, evaluate, and control safety and health hazards, and provide for emergency response to hazardous materials which may be encountered in the course of these field activities. This HASP has been prepared as a proactive and precautionary measure to deal with unanticipated health and safety concerns encountered during field activities. This HASP is applicable to HzW personnel and will remain at the project site for the duration of the project.

# 2.0 RESPONSIBILITIES

All HzW personnel are responsible for continuous adherence to the health and safety procedures, covered in this HASP, during the performance of their work. No person may work in any manner that conflicts with the intent or the inherent safety and environmental precautions expressed in these procedures.

# 2.1 Project Manager

The Project Manager or his/her designee is ultimately responsible for ensuring that all project activities are completed in compliance with requirements set forth in this plan.

The Project Manager or his/her designee in conjunction with HzW's Health and Safety Department will be responsible for developing, modifying, amending, and/or deleting the site-specific provisions of this HASP. The HASP will be revised if warranted by changing site conditions.

The Project Manager or his/her designee and/or HzW's Health and Safety Department are the contacts for regulatory agencies on matters of health and safety.

The Project Manager or his/her designee responsibilities include:

- General health and safety program administration in the field;
- Updating equipment or procedures based on information obtained during field operations;
- · Establishing sampling and analysis parameters based on expected contaminants;
- · Stopping work as required to ensure the health and safety of personnel;
- Developing site-specific and project specific employee response plans as required, based on expected hazards; and
- Distributing copies of the HASP to the appropriate persons, and ensuring that all
  distributed copies are promptly updated to include modifications, amendments,
  and/or deletions.

# 2.2 HzW's Health and Safety Department

HzW's Health and Safety Department in conjunction with the Project Manager will assist in developing, modifying, amending, and/or deleting the site-specific provisions of this HASP. The HASP will be revised if warranted by changing site conditions.

HzW's Health and Safety Department and/or the Project Manager or his/her designee are the contacts for regulatory agencies on matters of health and safety.

HzW's Health and Safety Department responsibilities include:

- Determining initial and subsequent personnel protection requirements;
- Reviewing significant accidents and illnesses related to possible environmental causes, and implementing corrective actions; and
- Developing site-specific and project specific employee response plans as required based on expected hazards.

#### 2.3 Field Technicians

All field technicians working at the project site are required to comply with the provision of this HASP and all applicable federal, state, and local regulations. All field technicians will read and sign off on the Employee Acknowledgement Form included at the beginning of this HASP prior to arriving at the project site. A Field Technician's responsibilities include:

- Ensuring his/her own health and safety by completing tasks in a safe manner, and reporting any unsafe acts or conditions to their Project Manager or their designee;
- Monitoring themselves and their fellow employees for signs and symptoms of heat or cold stress and chemical exposure;
- · Maintaining the operation and calibration of monitoring equipment; and
- Assuring that adequate first aid is present on site.

Key HASP personnel for this project are listed in Appendix A.

## 3.0 SITE DESCRIPTION AND SCOPE OF WORK

In general, the site is located to the north and east of Commercial Drive and Canal Road in the city of Cleveland, Cuyahoga County, Ohio. The site is comprised of separate parcels/properties owned by Norfolk Southern Railway. Two (2) properties consist of vacant land and one (1) is currently developed as parking lot. A scope of services was developed based on the findings of geotechnical drilling conducted in July 2009. A copy of HzW's proposal is included as **Appendix B**.

# 4.0 SITE CHARACTERIZATION AND CHEMICAL HAZARD ASSESSMENT

During July 2009, geotechnical drilling crews installed a series of soil borings for the proposed realignment of Commercial Drive between Canal Road and the proposed East Ninth Street Extension within the CUY-Cleveland Innerbelt project area. The geotechnical drilling crews documented encountering materials with odors and varying colors indicative of being corrosive. Historic Sanborn fire insurance maps indicate that the location of the site was previously utilized as railroad yards. Therefore, there is a potential for field personnel to be exposed to hazardous substance and petroleum constituents during assessment activities. A list of worst-case scenario chemicals is presented in **Appendix C**.

#### 5.0 GENERAL WORK PRACTICES

- No food, beverages or tobacco products will be present, consumed, or used in areas where project activities are being conducted.
- Before eating, drinking or smoking, employees must wash their hands and remove any outer protective garments.
- First aid kits must be readily accessible at the project site.
- Fire extinguishers must be readily accessible at the project site. Where there is fire potential, fire extinguishers will be located in the adjacent area.
- Should there be any contaminated protective equipment, such as respirators, hoses, boots, etc., said equipment shall not be removed from the project site until such equipment has been cleaned, or properly packaged and labeled.
- Legible and understandable precautionary labels which comply with hazard communication requirements must be affixed prominently to containers of contaminated waste, debris and/or clothing.
- · Where work is being performed at or near roadways, orange vests must be worn.
- Removal of any contaminated soil from protective clothing or equipment by blowing, shaking, or any other means that disperse contaminants into the air is prohibited.
- All areas that have been determined as contaminated at the project site must be
  clearly marked as such. No personnel, equipment, etc., must be removed from the
  site until they have been properly decontaminated, according to applicable state and
  federal regulations.

# 6.0 PERSONAL PROTECTIVE EQUIPMENT

This section is provided as a proactive measure should unanticipated hazardous materials or wastes be encountered during the project. The items in this section are presented to give a broad spectrum of alternative personal protective equipment (PPE) options to address a worst case scenario. It is not likely that the more comprehensive of these options will need to be implemented.

This section outlines maintenance and storage of PPE, decontamination, donning and doffing procedures, inspection and monitoring effectiveness and limitation.

# 6.1 Respiratory Protection

- Only employees who have been trained to wear and maintain respirators properly shall be allowed to enter areas requiring respiratory protection.
- Only properly cleaned, maintained, National Institute of Occupational Safety and Health (NIOSH) approved respirators shall be used on site.
- Selection of respirators, as well as any decision regarding upgrading or downgrading of respiratory protection, will be made by HzW's Health and Safety Department.
- Used air-purifying cartridges shall be replaced at the end of each shift or when breakthrough is suspected.
- Positive and negative pressure tests shall be performed each time the respiratory is donned.
- Only employees who have been both fit tested and have had a medical surveillance examination within the last 12 months will be allowed to work in atmospheres where respirators are required. Contractors hired by HzW shall provide certificates of respirator fit testing completed within the last 12 months for each employee on site.
- Respirator users shall be instructed in the proper use and limitation of respirators.
- If an employee has difficulty in breathing during the fit test or during use, he/she shall be evaluated medically to determine if he/she can wear a respirator safely while performing assigned tasks.
- No employee shall be assigned to tasks requiring the use of respirators if, based upon the most recent medical surveillance examination, a physician determines that the health or safety of the employee will be impaired by respirator use.
- Contact lenses shall not be worn while using any type of respiratory protection.
- Respirators shall be cleaned and sanitized daily after use.
- Respirators shall be inspected during cleaning. Worn or deteriorated parts shall be replaced.
- Facial hair that might interfere with a good face-piece seal or proper operation of the respirator is prohibited.
- The Project Manager in conjunction with HzW's Health and Safety Department will review the respiratory protection program daily to ensure employees are properly wearing and maintaining their respirators and that the selected respiratory protection is adequately protecting the employees.

# 6.2 Levels of Protection

The following sections outline four (4) basic levels of PPE based on guidelines issued by NIOSH, the Occupational Safety and Health Administration (OSHA) and the United States Environmental Protection Agency (U.S. EPA). Selection of PPE is typically based upon the location and nature of the project, past use at the project site, the likelihood of encountering hazardous materials or waste, and any additional information gathered from previous sampling and analysis performed or sampling and

analysis performed as part of this HASP's Scope of Work. The PPE selected for this project site is outlined in **Appendix D**.

Specific levels of protection will be changed whenever additional information is obtained and/or site conditions and activities so dictate. Levels of protection can either be increased to the next higher level, or decreased to the next lower level. The decision to change levels of protection will be made by the Project Manager in conjunction with HzW's Health and Safety Department. If the need arises to protect health and safety, the Project Manager can upgrade protection levels without input from HzW's Health and Safety Department. However, the Project Manager must then discuss the decision with the Health and Safety Department as soon as feasible. Levels of protection will not be downgraded without prior approval from HzW's Health and Safety Department.

# 6.2.1 Level A Protection

Level A protection would be used when the greatest level of skin, respiratory and eye protection is required. This level of protection is typically required for fire fighting activities, oxygen deficient environments or immediately dangerous to life and health (IDLH) atmospheres. Level A Protection is not anticipated at this time for HzW personnel at the project site. The following equipment will be used for Level A protection:

- Full-face pressure demand SCBA (self-contained breathing apparatus) or a
  pressure demand (positive pressure) air line respirator with an escape
  bottle for IDLH atmospheres. (Assigned Protection Factor approximately
  10,000 and 2000, respectively)
- · Fully encapsulated chemical protective suit.
- · Gloves: inner and outer chemically resistant.
- Chemical resistant, safety-toe boots.
- Booties/disposable boot covers.
- Hard hat (under suit).

# 6.2.2 Level B Protection

Level B respiratory protection is the same as Level A respiratory protection. However, the protective clothing is different. Level B Protection is not anticipated at this time for HzW personnel at the project site. The following equipment will be used for Level B protection:

- Full-face, positive pressure SCBA or a pressure demand (positive pressure) airline respirator with an escape bottle for IDLH atmospheres.
- · Coated Tyvek (or other chemical resistant) coveralls with hood.
- Inner and outer chemical resistant gloves.
- Chemical resistant safety toe boots.
- Booties/disposable boot covers.

· Hard Hat.

# 6.2.3 Level C Protection

Level C Protection is mandatory for any personnel entering an area where the airborne contaminants exceed or may exceed OSHA PELs. <u>Level C</u>

<u>Protection is not anticipated at this time for HzW personnel at the project site.</u> A full-face air-purifying respirator may only be utilized if:

- . The chemical compounds have adequate warning properties;
- The personnel have passed qualitative fits tests for the particular mask as previous required in the HASP;
- The appropriate filter cartridges are used and their service limitation are not exceeded; and
- The project's operations will not encounter unknown compounds or excessive concentrations of known compounds.

Half face respirators will be used only when approved by the Project Manager in conjunction with HzW's Health and Safety Department.

The following equipment will be used for Level C protection:

- Full-faced, air purifying canister-equipped respirator;
- · Coated Tyvek® (or other chemical resistant) coverall with hood;
- · Chemical resistant, safety-toe boots;
- Booties/disposable boot covers;
- · Inner and outer chemical resistant gloves;
- Hard had (with goggles if half-face respirator is used);
- Respirator cartridges will be changed daily, and also upon the detection of any chemical odor by the worker.

#### 6.2.4 Level D Protection

Level D protection is the minimum level of protection required at the site.

Level D protection is anticipated at this time for HzW personnel the project site. The following equipment will be used for Level D protection:

- · Half-face air-purifying respirators, as necessary;
- Protective coveralls;
- · Safety glasses;
- · Safety-toe boots or shoes;
- · Gloves of an appropriate material;
- · Hard hat, as necessary; and
- Hearing protection, as necessary.

# 6.3 Using Personal Protective Equipment

All personnel at the project site will comply with the required PPE, according to established procedures in this HASP to minimize exposure potential. When leaving the project site, personal protective equipment will be removed according to these established procedures to minimize the potential for the spread of contamination.

# 6.3.1 Donning Procedure – Level D

- Put on protective coveralls;
- Put on boots:
- · Put on gloves;
- If hearing protection is required, put in earplugs;
- · Put on hardhat, if necessary; and
- Put on safety glasses or goggles.

# 6.3.2 Doffing Procedure – Level D

- Remove excess soil and/or other material from outer clothing and boots while at excavation area;
- Before leaving the project site, remove coveralls, boots, gloves, safety goggles and hard hat;
- If disposable coveralls are used, place them in the appropriate refuse container; and
- If coveralls require laundering, place in appropriate laundry receptacle.

#### 6.3.3 Donning Procedures – Levels C and B

- Remove street clothes and store in a clean location;
- Put on protective coveralls;
- Put on boots and boot covers and tape the coveralls;
- Don respirator and check for secure fit;
- Put on gloves;
- Tape the cuff of the gloves over the coveralls at the wrist;
- If hearing protection is required, put in earplugs;
- Put hood or head covering over the respirator:
- Put on hard hat, if necessary, over the hood and respirator (Tape to secure from falling off); and
- Put on any remaining protective equipment such as safety glasses or goggles.

# 6.3.4 Doffing Procedures – Levels C and B

At the discretion of HzW's Health and Safety Department and depending on the activities, one person may remain outside the project site to assist in decontamination of personnel leaving the site. Whenever a person requires decontamination, the following sequence of stations will be used:

Station 1. Exiting personnel will remove excess soil and/or other materials from their outer clothing and boots.

# Station 2. Required equipment at Station 2 shall include the following:

- Plastic lined receptacle
- Chair
- Clean, damp cloths
- Paper towels
- Plastic bags

At Station 2, personnel will wipe their respirators (if used), hard hats, and boots with clean, damp cloths and then remove those items. If the inner gloves are contaminated or appear to be dirty, they must be removed and replaced prior to wiping off equipment. All items removed are then hand-carried to the next station.

# Station 3. Required equipment at Station 3 shall include the following:

- Wash basin with soap and water
- Respirator sanitation station

At this Station, personnel will thoroughly wash their hands and face. Respirators will be sanitized and then placed in a clean, plastic ziplock bag. Lined waste receptacles containing disposable equipment, garments and PPE will be removed and disposed in accordance with RCRA regulations.

# 6.4 Personal Protective Equipment Selection

The level of PPE can be based on measurements of the work environment when such measurements can be made in real-time. When the assessment of the work environment depends on laboratory analysis of samples collected or past land use, then the selection of PPE will be made on the professional judgment of possible or expected exposures.

#### 7.0 SITE MONITORING

If determined necessary by field personnel, HzW's Project Manager or HzW's Health and Safety Department, monitoring of atmospheric and/or breathing zone atmospheric conditions will be performed during on-site activities using real-time instrumentation, a photoionization detector (PID) and a lower explosive limit (LEL) meter, to determine

total organic contaminant concentrations and/or the percentage of explosive gas vapors. Site monitoring of atmospheric conditions will not be conducted at the site during on-site activities. As necessary, monitoring of the breathing zone will be conducted prior to initiation of on-site activities and continue during activities. Site monitoring will be performed at the point of highest expected concentration with the sample media located at the ground surface and within the breathing zone at 4 to 6 feet above the ground surface.

The work area air monitoring program addressed in this section has been developed to aid in the selection of PPE and to document exposures to on-site personnel. Prior to commencement of project activities, PID scans will be utilized as a preliminary indication of site conditions. PID and LEL readings will also be taken in the breathing zone and over excavated soil as sampling activities proceed. PID and LEL reading times will be recorded on an air monitoring log. Monitoring may be decreased if the results prove to be negative or uniform below ½ of PEL for the chemicals of concern. Based on known site conditions, the monitoring program will take into account the following factors:

- Determining when peak concentrations may be encountered; and
- Determining when and where unusual contaminants may be present.

## 8.0 PERSONAL EXPOSURE AIR MONITORING

The personal exposure air monitoring program addressed in this section has been developed to aid in the selection of PPE and to document exposures to on-site personnel. Personal exposure air monitoring is not anticipated at this time. However, information pertaining to personal exposure air monitoring is presented below should personal exposure air monitoring be deemed necessary at the project site by the Project Manager and/or HzW's Health and Safety Department.

The frequency and duration of personnel monitoring will be at the discretion of HzW's Health and Safety Department. In situations where personnel exposure is being evaluated, the personnel sample will be collected in the breathing zone of the employee.

When applicable, site personnel will be notified of air sampling results as soon as they are available. Where samples require laboratory analysis, the results will be made known to site personnel the day following receipt of results.

The personal exposure air monitoring program may include real-time instrumentation (direct reading instruments) and/or integrated air sampling (personal sampling pump methods). Appropriate NIOSH or OSHA sampling and analytical procedures will be utilized for time-weighted average monitoring. All air monitoring equipment will be maintained and operated in accordance with manufacturers' recommendations. Real time instrument maintenance and calibration data will be recorded in the air-monitoring log. When applicable, calibration of sampling pumps will also be documented in the daily field notes.

# 8.1 Real-Time Sampling Methods

Real-time air monitoring will be conducted using the PID. Direct reading Draeger tubes for detection of specific contaminants may also be used periodically during the project.

## 8.2 Integrated Air Monitoring

Integrated air monitoring may be performed to determine;

- The presence of a specific contaminants or contaminants;
- Peak concentrations; and
- Time-weighted average concentrations.

Personal air monitoring will be conducted when high concentrations of volatile organics are anticipated. Air samples will be collected using Draeger tubes, diffusion type organic vapor monitor badges or 8-hour personal monitoring pumps. Air analysis will be done for targeted contaminants as determined by the Project Manager in conjunction with HzW's Health and Safety Department.

# 8.3 Noise Monitoring

Noise monitoring will not be conducted at the project site.

# 8.4 Equipment Tampering

On-site personnel must wear monitoring equipment and the required personal protective equipment. Refusal to wear appropriate equipment and/or intentional tampering with sampling apparatus will lead to disciplinary action and immediate dismissal from the project site.

# 8.5 Monitoring Record

The Project Manager will be responsible for establishing and maintaining records of all required monitoring as described below:

- Employee name and social security number;
- Date, time, pertinent task information and exposure information;
- · Type of PPE worn; and
- Engineering controls used to reduce exposure.

# 8.6 Notification

Employees will be notified of exposure in excess of the permissible exposure limit and will be provided with follow-up medical monitoring when required.

# 9.0 MEDICAL SURVEILLANCE EXAMINATION

All HzW field personnel will have successfully completed an initial and annual physical examination. The examination is designed to meet the requirements of 29 CFR 1920.120 for possible exposure to hazardous materials or waste. The medical surveillance examination consists of the following:

- Medical and occupational medical history and physical examination;
- · Visual test
- Urinalysis
- Audiogram
- Spirometry
- EKG (Age 40 and over)
- · dt Tetanus
- · Chest X-ray
- · CBC with differential
- 13 Blood Chemistry Tests
- Blood Lead and Zinc Protoporphrin
- Cholinesterase
- PCBs
- Mercury
- Cadmium

The following information is provided to the examining physician:

- Description of employee's duties;
- Anticipated chemical exposures and levels;
- · Description of PPE to be used; and
- Information from previous medical examinations.

Exit medical surveillance examinations are provided to HzW field personnel upon termination of employment.

A copy of the medical examination report is provided at the employee's request. The employee will be informed of any medical conditions that would result in work restriction.

#### 10.0 FIRST AID AND MEDICAL TREATMENT

All field personnel must report any near-miss incident, accident, injury, or illness to HzW's Health and Safety Department as well as their Project Manager. First aid will be rendered expeditiously by a person qualified to do so. The employee's Project Manager will complete an accident/injury report and conduct an investigation of the incident as soon as emergency conditions (if any) no longer exist and medical or first-aid treatment has been rendered. The investigation should follow completion of the accident/injury

report. HzW's Health and Safety Department shall promptly receive for review a copy of the accident/injury report, and the results of the incident investigation.

#### 11.0 MEDICAL RESTRICTION

When the examining physician determines a need to restrict a field employee from their job activities, that determination will be communicated to the employee's Project Manager as well as the Health and Safety Department. The Project Manager will ensure that the employee complies with the work restriction(s).

#### 12.0 MEDICAL RECORDS

Medical and exposure monitoring records will be maintained according to the requirements of 29 CFR 1910.120, and retained for a minimum of 30 years. Confidentiality of these records shall be maintained through retention.

#### 13.0 EMERGENCY PROCEDURES

Should the need for outside medical attention arise, St. Vincent Charity Hospital located in Cleveland, Ohio will be used. In addition, a map, as well as directions from the project site to the hospital, is also included as **Appendix E**. The City of Cleveland Police Department will be called to handle any security incidents at the project site.

The Project Manager will establish evacuation routes and assembly areas for the project site. All personnel entering the site will be informed of these routes and assembly areas.

Unusual events, activities, odors, and conditions will be reported to the Project Manager and subsequently HzW's Health and Safety Department. Emergency telephone numbers will be available in each motor vehicle at the project site, along with a map and directions to the nearest hospital. A list of emergency telephone numbers is included as **Appendix F**.

All incidents will be dealt with in a manner which minimizes health risks to project site workers, the environment and the local community. In the case of a medical emergency, paramedics will be summoned without delay.

#### 14.0 TRAINING

All field personnel shall have taken a 40-hour HAZWOPER training course with annual refreshers as well as first aid and CPR training. In addition, training shall cover Hazard Communication and Respiratory Protection.

All training shall be documented by a certificate signed by the instructor. A copy of each and every training certificate shall be maintained by HzW's Health and Safety Department. Subcontractors must provide to HzW's Health and Safety Department copies of certificates of training for all subcontractors personnel at the project site.

#### 15.0 SAFETY MEETINGS

The Project Manager shall conduct a safety meeting at the beginning of each shift, or whenever new employees arrive at the project site once the project commences. Topics to be discussed at these meetings include health and safety considerations for the day's activities, necessary protective equipment (as applicable), problems encountered, and new operations. Attendance records and meeting notes shall be maintained by the Project Manager.

# 16.0 MATERIAL SAFETY DATA SHEETS

This HASP includes Material Safety Data Sheets (MSDSs) for the chemicals listed in **Appendix C**. The MSDSs shall be maintained on site as part of this HASP, and shall be accessible to all employees. A copy of each chemical's MSDS is also included in **Appendix C**.

# APPENDIX A KEY HASP PERSONNEL

# The following is a list of potential personnel having responsibilities under this HASP:

Doug Wetzel Joan Sablar Steve Sablar Joshua Derico

Joshua Derico Joseph Harcher Project Manager

Health and Safety Representative

Senior Field Technician

Field Technician Field Technician

# APPENDIX B

HzW's PROPOSAL

#### ATTACHMENT 1

# SCOPE OF SERVICES for TARGETED PHASE II ENVIRONMENTAL SITE ASSESSMENT CUY-CLEVELAND INNERBELT, COMMERCIAL DRIVE ALIGNMENT PROJECT AREA

#### BACKGROUND

During July 2009, geotechnical drilling crews installed a series of borings for the proposed realignment of Commercial Drive between Canal Road and the proposed East Ninth Street Extension in the CUY-Cleveland Innerbelt project area. During installation of four borings (designated B-071, B-072, B-073 and B-075), the drilling crew encountered materials variously described as:

- ♦ Having a "bad odor" or being an "unknown blue/white material" that "looked [corrosive]" to a depth of over 20 feet below ground surface in B-071;
- ♦ An "unknown blue material" at a depth of less than 5 feet in B-072;
- ♦ An "unknown blue/green rock material" to a depth of less than 5 feet in B-073; and
- ♦ A "nasty...blue/green material" from 1.5 to 4.5 feet in B-075 which "smells bad" and the drilling crew was "...not sure if [it was] safe to touch."

A review of Sanborn fire insurance maps provided to HzW by ODOT's Office of Environmental Services (OES) indicated that the areas in question were historically occupied by railroad yards that were "full of tracks". Therefore, it is unclear what this discolored material could be.

In ODOT's Request for Task Order Proposal, it was requested that HzW install five soil borings to a depth of 20 feet. Based upon the above information, it would appear that the geotechnical drilling crews only encountered the material to a depth of 20 feet in one of the four borings. The material was encountered at a depth of less than 5 feet in three of the four geotechnical borings where this material was observed. Based upon a discussion with ODOT OES on November 3, 2009, it was agreed to reduce the number of borings from five (5) to four (4), and to reduce the depth at three of the four borings (those installed near locations B-072, B-073, and B-075) from 20 feet to 6 feet.

### SCOPE OF TARGETED PHASE II ENVIRONMENTAL SITE ASSESSMENT (ESA)

The proposed scope of investigation in the project area will consist of the following:

# Task A: Pre-Assessment Planning and Support

- Coordinating the location of proposed soil borings with the original geotechnical drilling crew, ODOT
  and affected property owners and/or tenants (as directed by ODOT), obtaining utility clearances, and
  otherwise obtaining clear access to soil boring locations.
- 2. Preparation of a site-specific health and safety plan (HASP) to be used by personnel responsible for implementing the targeted Phase II ESA. The HASP will be reviewed and signed by all personnel prior to initiation of field activities.

# Task B: Physical Site Assessment

- 1. Using Geoprobe® subsurface sampling techniques, install four (4) soil borings in the project area. One (in the general location of original geotechnical boring B-071) will penetrate to a terminal depth of twenty (20) feet below ground surface, while the other three borings (in the general locations of original geotechnical borings B-072, B-073, and B-075) will penetrate to a terminal depth of six (6) feet below ground surface. Soil samples will be collected continuously at 2-foot intervals in each boring from ground surface to terminal depth, with recovered materials logged in the field by a qualified field technician or field geologist.
- 2. In the event that any boring encounters the types of materials referenced by the geotechnical drilling crew, samples of the material will be collected and transferred to laboratory-supplied containers, and placed in an ice chest for preservation in the field. Based upon the geotechnical boring logs, HzW would anticipate encountering this material from between: depths of 0-14 feet at location B-071 (the strongest odor was noted at 8-10 feet) and perhaps near the terminal depth (20 feet); depths of 1-5 feet at location B-072; depths of 2-4 feet at location B-073; and depths of 1.5-4.5 feet at location B-075. Based upon this, it is anticipated that HzW would collect a maximum of: three (3) samples from the boring installed at location B-071; two (2) samples from the boring installed at location B-073; and two (2) samples from the boring installed at location B-075.
- 3. The eight (8) soil samples collected from these borings would be submitted to Test America Labs, Inc. of North Canton, Ohio, for analysis of: pH; hydrogen sulfide; arsenic, cadmium, chromium, hexavalent chromium, lead, and cyanide; volatile organic compounds (VOCs); semi-volatile organic compounds (SVOCs), and total petroleum hydrocarbons (TPH) by approved laboratory methods. Test America Labs, Inc. would be instructed in the proper preparation and handling of these samples, and to "hold" the samples for possible identification of tentatively identified compounds (TICs), which is included as a contingency analytical cost to this task order proposal.
- 4. Abandon all four (4) borings following installation using hydrated bentonite chips.

# Task C: Data Evaluation/Report Preparation/Coordination

This element will be conducted following completion of the physical site assessment portion of the Phase II ESA scope of work, and would consist of evaluating all analytical results upon receipt, and preparation of a report for submission to ODOT OES. This report will include a narrative of the circumstances surrounding the need and scope of Phase II ESA activities, site drawings showing the location of borings relative to on-site property features, and a discussion of the analytical results relative to comparable environmental standards in the State of Ohio. Three (3) copies of this report would be submitted to ODOT OES upon completion. In addition, this task includes time for coordination with ODOT District 12 and ODOT OES at ODOT District 12 Headquarters or via telephone to discuss the findings, the need for delineation of the materials encountered, and potential remedial options during anticipated construction activities in the CUY-Cleveland Innerbelt project area.

# APPENDIX C

# LIST OF POTENTIAL CHEMICAL CONTAMINANTS AT THE PROJECT SITE

# POTENTIAL CHEMICAL CONTAMINANTS AT THE PROJECT SITE

A listing of the chemicals that are likely to be encountered at the Property is presented below.

- ♦ Trichloroethene
- Benzene
- ♦ Benzo(a)pyrene
- ♦ Arsenic
- ♦ Cadmium
- Cyanide
- ♦ Lead

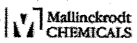
A copy of each of these chemicals Material Safety Data Sheet is included for review.

MSDS Number: T4940 \* \* \* \* \* Effective Date: 12/06/07 \* \* \* \* \* Supercedes: 08/01/05

MSDS

Material Safety Data Sheet

From: Mallinekrodt Baker, Inc. 222 Red School Lane Phillipsburg, NJ 08865





24 Hour Emergency Telephone: 908-459-2151 CHENTAEO: 1-880-424-8300

National Response in Canada CANUTEC: 619-406-4666

Outside U.S. and Canada Chemirec: 703-527-3887

NOTE: CHEMIFIEC, CANUTEC and National Response Center emergency numbers to be used only in the event of chemical emergencies, smothing a spill, leak, life, exposure or accident involving chemicals.

All non-emergency questions should be directed to Customer Service (1-800-582-2537) for assistance.

# TRICHLOROETHYLENE

# 1. Product Identification

Synonyms: Trichloroethene; TCE; acetylene trichloride; Ethinyl trichloride

CAS No.: 79-01-6

Molecular Weight: 131.39 Chemical Formula: C2HCl3

**Product Codes:** 

J.T. Baker: 5376, 9454, 9458, 9464, 9473

Mallinckrodt: 8600, 8633

# 2. Composition/Information on Ingredients

Ingredient	CAS No	Percent	Hazardous
Trichloroethylene	7901-6	100%	Yes

# 3. Hazards Identification

**Emergency Overview** 

WARNING! HARMFUL IF SWALLOWED OR INHALED. AFFECTS HEART, CENTRAL NERVOUS SYSTEM, LIVER AND KIDNEYS. CAUSES SEVERE SKIN IRRITATION. CAUSES IRRITATION TO EYES AND RESPIRATORY TRACT. SUSPECT CANCER HAZARD. MAY CAUSE CANCER. Risk of cancer depends on level and duration of exposure.

# SAF-T-DATA<sup>(tm)</sup> Ratings (Provided here for your convenience)

Health Rating: 2 - Moderate (Poison) Flammability Rating: 1 - Slight

Reactivity Rating: 1 - Slight Contact Rating: 3 - Severe

Lab Protective Equip: GOGGLES & SHIELD; LAB COAT & APRON; VENT HOOD; PROPER

**GLOVES** 

Storage Color Code: Blue (Health)

**Potential Health Effects** 

# Inhalation:

Vapors can irritate the respiratory tract. Causes depression of the central nervous system with symptoms of visual disturbances and mental confusion, incoordination, headache, nausea, euphoria, and dizziness. Inhalation of high concentrations could cause unconsciousness, heart effects, liver effects, kidney effects, and death.

# Ingestion:

Cases irritation to gastrointestinal tract. May also cause effects similar to inhalation. May cause coughing, abdominal pain, diarrhea, dizziness, pulmonary edema, unconsciousness. Kidney failure can result in severe cases. Estimated fatal dose is 3-5 ml/kg.

# **Skin Contact:**

Cause irritation, redness and pain. Can cause blistering. Continued skin contact has a defatting action and can produce rough, dry, red skin resulting in secondary infection.

#### Lye Contact:

Vapors may cause severe irritation with redness and pain. Splashes may cause eye damage.

# **Chronic Exposure:**

Chronic exposures may cause liver, kidney, central nervous system, and peripheral nervous system effects. Workers chronically exposed may exhibit central nervous system depression, intolerance to alcohol, and increased cardiac output. This material is linked to mutagenic effects in humans. This material is also a suspect carcinogen.

# **Aggravation of Pre-existing Conditions:**

Persons with pre-existing skin disorders, cardiovascular disorders, impaired liver or kidney or respiratory function, or central or peripheral nervous system disorders may be more susceptible to the effects of the substance.

# 4. First Aid Measures

#### Inhalation:

Remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. Call a physician.

# Ingestion:

Induce vomiting immediately as directed by medical personnel. Never give anything by mouth to an unconscious person. Call a physician.

## **Skin Contact:**

Immediately flush skin with plenty of soap and water for at least 15 minutes while removing contaminated clothing and shoes. Get medical attention. Wash clothing before reuse. Thoroughly clean shoes before reuse.

# **Eye Contact:**

Immediately flush eyes with plenty of water for at least 15 minutes, lifting lower and upper eyelids occasionally. Get medical attention immediately.

# Note to Physician:

Do not administer adrenaline or epinephrine to a victim of chlorinated solvent poisoning.

# 5. Fire Fighting Measures

#### Fire:

Autoignition temperature: 420C (788F) Flammable limits in air % by volume:

lel: 8; uel: 12.5 **Explosion:** 

A strong ignition source, e. g., a welding torch, can produce ignition. Sealed containers may rupture when heated.

# Fire Extinguishing Media:

Use water spray to keep fire exposed containers cool. If substance does ignite, use CO2, dry chemical or foam.

# **Special Information:**

In the event of a fire, wear full protective clothing and NIOSH-approved self-contained breathing apparatus with full facepiece operated in the pressure demand or other positive pressure mode. Combustion by-products include phosgene and hydrogen chloride gases. Structural firefighters' clothing provides only limited protection to the combustion products of this material.

# 6. Accidental Release Measures

Ventilate area of leak or spill. Remove all sources of ignition. Wear appropriate personal protective equipment as specified in Section 8. Isolate hazard area. Keep unnecessary and unprotected personnel from entering. Contain and recover liquid when possible. Use non-sparking tools and equipment. Collect liquid in an appropriate container or absorb with an inert material (e. g., vermiculite, dry sand, earth), and place in a chemical waste container. Do not use combustible materials, such as saw dust. Do not flush to sewer! US Regulations (CERCLA) require reporting spills and releases to soil, water and air in excess of reportable quantities. The toll free number for the US Coast Guard National Response Center is (800) 424-8802.

# 7. Handling and Storage

Keep in a tightly closed container, stored in a cool, dry, ventilated area. Protect against physical damage. Isolate from any source of heat or ignition. Isolate from incompatible substances. Containers of this material may be hazardous when empty since they retain product residues (vapors, liquid); observe all warnings and precautions listed for the product.

# 8. Exposure Controls/Personal Protection

# Airborne Exposure Limits:

Trichloroethylene:

-OSHA Permissible Exposure Limit (PEL):

100 ppm (TWA), 200 ppm (Ceiling),

300 ppm/5min/2hr (Max)

-ACGIH Threshold Limit Value (TLV):

10 ppm (TWA) 25 ppm (STEL); A2 Suspected Human Carcinogen.

# Ventilation System:

A system of local and/or general exhaust is recommended to keep employee exposures below the Airborne Exposure Limits. Local exhaust ventilation is generally preferred because it can control the emissions of the contaminant at its source, preventing dispersion of it into the general work area. Please refer to the ACGIH document, *Industrial Ventilation, A Manual of Recommended Practices*, most recent edition, for details.

# Personal Respirators (NIOSH Approved):

If the exposure limit is exceeded and engineering controls are not feasible, wear a supplied air, full-facepiece respirator, airlined hood, or full-facepiece self-contained breathing apparatus. Breathing air quality must meet the requirements of the OSHA respiratory protection standard (29CFR1910.134). This substance has poor warning properties. Where respirators are required, you must have a written program covering the basic requirements in the OSHA respirator standard. These include training, fit testing, medical approval, cleaning, maintenance, cartridge change schedules, etc. See 29CFR1910.134 for details.

### **Skin Protection:**

Wear impervious protective clothing, including boots, gloves, lab coat, apron or coveralls, as appropriate, to prevent skin contact. Neoprene is a recommended material for personal protective equipment.

# **Eye Protection:**

Use chemical safety goggles and/or a full face shield where splashing is possible. Maintain eye wash fountain and quick-drench facilities in work area.

# 9. Physical and Chemical Properties

### Appearance:

Clear, colorless liquid.

#### Odor:

Chloroform-like odor.

## Solubility:

Practically insoluble in water. Readily miscible in organic solvents.

# Specific Gravity:

1.47 @ 20C/4C

#### pH:

No information found.

% Volatiles by volume @ 21C (70F):

100

# **Boiling Point:**

87C (189F)

**Melting Point:** 

-73C (-99F)

Vapor Density (Air=1):

4.5

Vapor Pressure (mm Hg):

57.8 @ 20C (68F)

Evaporation Rate (BuAc=1):

No information found.

# 10. Stability and Reactivity

#### Stability:

Stable under ordinary conditions of use and storage. Will slowly decompose to hydrochloric acid when exposed to light and moisture.

#### **Hazardous Decomposition Products:**

May produce carbon monoxide, carbon dioxide, hydrogen chloride and phosgene when heated to decomposition.

#### **Hazardous Polymerization:**

Will not occur.

#### Incompatibilities:

Strong caustics and alkalis, strong oxidizers, chemically active metals, such as barium, lithium, sodium, magnesium, titanium and beryllium, liquid oxygen.

#### Conditions to Avoid:

Heat, flame, ignition sources, light, moisture, incompatibles

# 11. Toxicological Information

#### **Toxicological Data:**

Trichloroethylene: Oral rat LD50: 5650 mg/kg; investigated as a tumorigen, mutagen, reproductive effector.

### Reproductive Toxicity:

This material has been linked to mutagenic effects in humans.

\Cancer Lists\	NTD	Carcinogen	
Ingredient	Known	Anticipated	IARC Category
Trichloroethylene (79-01-6)	No	Yes	2A

# 12. Ecological Information

#### **Environmental Fate:**

When released into the soil, this material may leach into groundwater. When released into the soil, this material is expected to quickly evaporate. When released to water, this material is expected to quickly evaporate. This material has an experimentally-determined bioconcentration factor (BCF) of less than 100. This material is not expected to significantly bioaccumulate. When released into

the air, this material may be moderately degraded by reaction with photochemically produced hydroxyl radicals. When released into the air, this material is expected to have a half-life between 1 and 10 days.

#### **Environmental Toxicity:**

The LC50/96-hour values for fish are between 10 and 100 mg/l. This material is expected to be slightly toxic to aquatic life.

# 13. Disposal Considerations

Whatever cannot be saved for recovery or recycling should be handled as hazardous waste and sent to a RCRA approved incinerator or disposed in a RCRA approved waste facility. Processing, use or contamination of this product may change the waste management options. State and local disposal regulations may differ from federal disposal regulations. Dispose of container and unused contents in accordance with federal, state and local requirements.

# 14. Transport Information

Domestic (Land, D.O.T.)

Proper Shipping Name: TRICHLOROETHYLENE

Hazard Class: 6.1 UN/NA: UN1710 Packing Group: III

Information reported for product/size: 4L

International (Water, I.M.O.)

Proper Shipping Name: TRICHLOROETHYLENE

Hazard Class: 6.1 UN/NA: UN1710 Packing Group: III

Information reported for product/size: 4L

# 15. Regulatory Information

Chemical Inventory Status - Part 1\ Ingredient				 Australia
Trichloroethylene (79-01-6)	Yes	Yes	Yes	Yes
\Chemical Inventory Status - Part 2\				·
Ingredient	Korea		nada NDSL	Phil.
Trichloroethylene (79-01-6)	Yes	Yes	No	Yes
\Federal, State & International Regulati -SARA				313

Ingredient	RQ TP	Q List C	hemical Catg.
Trichloroethylene (79-01-6)	No No	Yes	No
\Federal, State & International Ingredient	Regulations CERCLA		SCA- 8 (d)
Trichloroethylene (79-01-6)	100	U228	No

Chemical Weapons Convention: No TSCA 12(b): No CDTA: No SARA 311/312: Acute: Yes Chronic: Yes Fire: No Pressure: No Reactivity: No (Pure / Liquid)

#### **WARNING:**

THIS PRODUCT CONTAINS A CHEMICAL(S) KNOWN TO THE STATE OF CALIFORNIA TO CAUSE CANCER.

Australian Hazchem Code: None allocated.

Poison Schedule: S6

WHMIS:

This MSDS has been prepared according to the hazard criteria of the Controlled Products Regulations (CPR) and the MSDS contains all of the information required by the CPR.

### 16. Other Information

NFPA Ratings: Health: 2 Flammability: 1 Reactivity: 0

Label Hazard Warning:

WARNING! HARMFUL IF SWALLOWED OR INHALED. AFFECTS HEART, CENTRAL NERVOUS SYSTEM, LIVER AND KIDNEYS. CAUSES SEVERE SKIN IRRITATION. CAUSES IRRITATION TO EYES AND RESPIRATORY TRACT. SUSPECT CANCER HAZARD. MAY CAUSE CANCER. Risk of cancer depends on level and duration of exposure.

#### **Label Precautions:**

Do not get in eyes, on skin, or on clothing.

Do not breathe vapor.

Keep container closed.

Use only with adequate ventilation.

Wash thoroughly after handling.

Keep away from heat and flame.

#### Label First Aid:

If swallowed, induce vomiting immediately as directed by medical personnel. Never give anything by mouth to an unconscious person. If inhaled, remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. In case of contact, immediately flush eyes or skin with plenty of water for at least 15 minutes. Remove contaminated clothing and shoes. Wash clothing before reuse. In all cases call a physician. Note to physician: Do not administer adrenaline or epinephrine to a victim of chlorinated solvent poisoning.

#### **Product Use:**

Laboratory Reagent.

#### **Revision Information:**

MSDS Section(s) changed since last revision of document include: 8.

#### Disclaimer:

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Prepared by: Environmental Health & Safety Phone Number: (314) 654-1600 (U.S.A.)

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# Material Safety Data Sheet Benzene

C# 02610

### Section 1 - Chemical Product and Company Identification

MSDS Name: Benzene

Catalog Numbers: AC167660000, AC167660010, AC167660025, AC167660250, AC167665000, AC168650250, AC295330000, AC295330010, AC295330025, AC295330250, AC296880000, AC296880010, AC296880025, AC296880250, AC610230010, AC610231000, AC611001000, B243-4, B243J, B245-4, B245-500, B411-1, B411-4, B412-1, S79920ACS

Synonyms: Benzol; Cyclohexatriene; Phenyl hydride.

**Company Identification:** 

Fisher Scientific 1 Reagent Lane Fair Lawn, NJ 07410

For information, call: 201-796-7100 Emergency Number: 201-796-7100

For CHEMTREC assistance, call: 800-424-9300

For International CHEMTREC assistance, call: 703-527-3887

### Section 2 - Composition, Information on Ingredients

CAS#	Chemical Name	Percent	EINECS/ELINCS
71-43-2	Benzene	> 99	200-753-7

### Section 3 - Hazards Identification

#### **EMERGENCY OVERVIEW**

Appearance: clear colorless liquid. Flash Point: -11 deg C.

**Danger!** Extremely flammable liquid and vapor. Vapor may cause flash fire. Harmful if swallowed, inhaled, or absorbed through the skin. Causes eye, skin, and respiratory tract irritation. Contains benzene. Benzene can cause cancer. Aspiration hazard if swallowed. Can enter lungs and cause damage. May cause blood abnormalities. May cause central nervous system effects.

**Target Organs:** Blood, central nervous system, respiratory system, eyes, bone marrow, immune system, skin.

#### **Potential Health Effects**

Eye: Causes eye irritation.

**Skin:** Causes skin irritation. Harmful if absorbed through the skin. Prolonged and/or repeated contact may cause defatting of the skin and dermatitis.

**Ingestion:** May cause central nervous system depression, characterized by excitement, followed by headache, dizziness, drowsiness, and nausea. Advanced stages may cause collapse, unconsciousness, coma and possible death due to respiratory failure. May cause effects similar to those for inhalation exposure. Aspiration of material into the lungs may cause chemical pneumonitis, which may be fatal.

halation: Causes respiratory tract irritation. May cause drowsiness, unconsciousness, and central ..ervous system depression. Exposure may lead to irreversible bone marrow injury. Exposure may lead to aplastic anemia. Potential symptoms of overexposure by inhalation are dizziness, headache, vomiting, visual disturbances, staggering gait, hilarity, fatigue, and other symptoms of CNS depression.

**Chronic:** May cause bone marrow abnormalities with damage to blood forming tissues. May cause anemia and other blood cell abnormalities. Chronic exposure to benzene has been associated with an increased incidence of leukemia and multiple myeloma (tumor composed of cells of the type normally found in the bone marrow). Immunodepressive effects have been reported. This substance has caused adverse roductive and fetal effects in laboratory animals.

### Section 4 - First Aid Measures

**Eyes:** In case of contact, immediately flush eyes with plenty of water for a t least 15 minutes. Get medical aid.

**Skin:** In case of contact, flush skin with plenty of water. Remove contaminated clothing and shoes. Get medical aid if irritation develops and persists. Wash clothing before reuse.

**Ingestion:** Potential for aspiration if swallowed. Get medical aid immediately. Do not induce vomiting unless directed to do so by medical personnel. Never give anything by mouth to an unconscious person. If vomiting occurs naturally, have victim lean forward.

**Inhalation:** If inhaled, remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. Get medical aid.

Notes to Physician: Treat symptomatically and supportively.

# Section 5 - Fire Fighting Measures

General Information: As in any fire, wear a self-contained breathing apparatus in pressure-demand, MSHA/NIOSH (approved or equivalent), and full protective gear. Use water spray to keep fire-exposed containers cool. Extremely flammable liquid and vapor. Vapor may cause flash fire. Approach fire from wind to avoid hazardous vapors and toxic decomposition products. Vapors are heavier than air and may avel to a source of ignition and flash back. Vapors can spread along the ground and collect in low or confined areas. This liquid floats on water and may travel to a source of ignition and spread fire. May accumulate static electricity.

Extinguishing Media: Use water spray, dry chemical, carbon dioxide, or appropriate foam.

Flash Point: -11 deg C ( 12.20 deg F)

Autoignition Temperature: 498 deg C ( 928.40 deg F)

Explosion Limits, Lower: 1.3 vol %

**Upper:** 7.1 vol %

NFPA Rating: (estimated) Health: 2; Flammability: 3; Instability: 0

#### Section 6 - Accidental Release Measures

**General Information:** Use proper personal protective equipment as indicated in Section 8. **Spilis/Leaks:** Absorb spill with inert material (e.g. vermiculite, sand or earth), then place in suitable container. Avoid runoff into storm sewers and ditches which lead to waterways. Remove all sources of ignition. Provide ventilation. Approach spill from upwind. Use water spray to cool and disperse vapors, protect personnel, and dilute spills to form nonflammable mixtures.

### Section 7 - Handling and Storage

**Handling:** Wash thoroughly after handling. Remove contaminated clothing and wash before reuse. Ground and bond containers when transferring material. Avoid contact with eyes, skin, and clothing. Empty containers retain product residue, (liquid and/or vapor), and can be dangerous. Take precautionary

measures against static discharges. Keep container tightly closed. Do not pressurize, cut, weld, braze, solder, drill, grind, or expose empty containers to heat, sparks or open flames. Use only with adequate ventilation. Keep away from heat, sparks and flame. Avoid breathing vapor.

**Storage:** Keep away from sources of ignition. Store in a tightly closed container. Keep from contact with idizing materials. Store in a cool, dry, well-ventilated area away from incompatible substances.

### Section 8 - Exposure Controls, Personal Protection

**Engineering Controls:** Use process enclosure, local exhaust ventilation, or other engineering controls to control airborne levels below recommended exposure limits. Use explosion-proof ventilation equipment. Facilities storing or utilizing this material should be equipped with an eyewash facility and a safety shower. See 29CFR 1910.1028 for the regulatory requirements for the control of employee exposure to benzene.

**Exposure Limits** 

Chemical Name	ACGIH	NIOSH	OSHA - Final PELs
Benzene	0.5 ppm TWA; 2.5 ppm STEL; Skin - potential significant contribution to overall exposure by the cutaneous r oute	0.1 ppm TWA 500 ppm IDLH	1 ppm TWA; 10 ppm TWA

SHA Vacated PELs: Benzene: 10 ppm TWA (unless specified in 1910.1028)

Personal Protective Equipment
Eves: Wear chemical splash goggles.

Skin: Wear appropriate protective gloves to prevent skin exposure.

Clothing: Wear appropriate protective clothing to prevent skin exposure.

**Respirators:** A respiratory protection program that meets OSHA's 29 CFR 1910.134 and ANSI Z88.2 requirements or European Standard EN 149 must be followed whenever workplace conditions warrant respirator use.

# Section 9 - Physical and Chemical Properties

Physical State: Liquid
Appearance: clear colorless

Odor: sweetish odor - aromatic odor

pH: Not applicable.

Vapor Pressure: 75 mm Hg @ 20 deg C

Vapor Density: 2.8 (air=1)
Evaporation Rate:Not available.
Viscosity: 0.647mPa @ 20 deg C
Boiling Point: 80.1 deg C

Freezing/Melting Point: 5.5 deg C

**Pecomposition Temperature:** Not available.

Jubility: 0.180 g/100 ml @ 25°C

Specific Gravity/Density:0.8765 @ 20°C

Molecular Formula:C6H6 Molecular Weight:78.11

### Section 10 - Stability and Reactivity

emical Stability: Stable under normal temperatures and pressures. Conditions to Avoid: Ignition sources, excess heat, confined spaces. Incompatibilities with Other Materials: Strong oxidizing agents.

Hazardous Decomposition Products: Carbon monoxide, carbon dioxide.

Hazardous Polymerization: Has not been reported.

### Section 11 - Toxicological Information

RTECS#:

CAS# 71-43-2: CY1400000

LD50/LC50:

CAS# 71-43-2:

Dermal, guinea pig: LD50 = >9400 uL/kg; Draize test, rabbit, eye: 88 mg Moderate; Draize test, rabbit, eye: 2 mg/24H Severe; Draize test, rabbit, skin: 20 mg/24H Moderate; Inhalation, mouse: LC50 = 9980 ppm;

Inhalation, mouse: LC50 = 9980 ppm; Inhalation, mouse: LC50 = 24 mL/kg/2H; Inhalation, rat: LC50 = 10000 ppm/7H; Inhalation, rat: LC50 = 34 mL/kg/2H; Inhalation, rat: LC50 = 6.5 mL/kg/4H; Oral, mouse: LD50 = 4700 mg/kg; Oral, rat: LD50 = 930 mg/kg;

Oral, rat: LD50 = 950 fig/kg; Oral, rat: LD50 = 1 mL/kg;

Oral, rat: LD50 = 1800 Benzene is considered very toxic; probable human oral lethal dose would be 50-500 mg/kg. Human inhalation of approximately 20,000 ppm (2% in air) was fatal in 5-10 minutes. While percutaneous absorption of liquid benzene through intact human skin can be limited (e.g., 0.05% of the applied dose), the absorbed dose via direct dermal contact combined with that received from body surface exposure to benzene in workplace air is such that a substantial fraction (20-40%) of the total exposure is due to skin absorption.

#### Carcinogenicity:

CAS# 71-43-2:

ACGIH: A1 - Confirmed Human Carcinogen
 California: carcinogen, initial date 2/27/87

NTP: Known carcinogenIARC: Group 1 carcinogen

**Epidemiology:** IARC has concluded that epidemiological studies have established the relationship between benzene exposure and the development of acute myelogenous leukemia, and that there is sufficient evidence that benzene is carcinogenic to humans.

**Teratogenicity:** Inhalation, rat: TCLO = 50 ppm/24H (female 7-14 day(s) after conception) Effects on Embryo or Fetus - extra-embryonic structures (e.g., placenta, umbilical cord) and Effects on Embryo or Fetus - fetotoxicity (except death, e.g., stunted fetus).; Inhalation,mouse: TCLo = 5 ppm (female 6-15 day(s) after conception) Effects on Embryo or Fetus - cytological changes (including somatic cell genetic aterial) and Specific Developmental Abnormalities - blood and lymphatic systems (including spleen and arrow).

**Reproductive Effects:** Inhalation, rat: TCLO = 670 mg/m3/24H (female 15 day(s) pre-mating and female 1-22 day(s) after conception) female fertility index (e.g. # females pregnant per # sperm positive females; # females pregnant per # females mated).; Oral, mouse: TDLo = 12 gm/kg (female 6-15 day(s)

after conception) Fertility - post-implantation mortality (e.g. dead and/or resorbed implants per total number of implants).

Mutagenicity: DNA Inhibition: Human, Leukocyte = 2200 umol/L.; DNA Inhibition: Human, HeLa cell = 2200 umol/L.; Mutation Test Systems - not otherwise specified: Human, Lymphocyte = 5 umol/L.;

ogenetic Analysis: Inhalation, Human = 125 ppm/1Y.; Cytogenetic Analysis: Human, Leukocyte = 1

....nol/L/72H.; Cytogenetic Analysis: Human, Lymphocyte = 1 mg/L.

Neurotoxicity: See actual entry in RTECS for complete information.

**Other Studies:** 

# Section 12 - Ecological Information

**Ecotoxicity:** Fish: Mosquito Fish: TLm = 395 mg/L; 24 Hr; UnspecifiedFish: Goldfish: LC50 = 46 mg/L; 24 Hr; Modified ASTM D 1345Fish: Fathead Minnow: LC50 = 15.1 mg/L; 96 Hr; Flow-through at 25°C (pH 7.9-8.0)Fish: Rainbow trout: LC50 = 5.3 mg/L; 96 Hr; Flow-through at 25°C (pH 7.9-8.0)Fish: Bluegill/Sunfish: LD50 = 20 mg/L; 24-48 Hr; Unspecified If benzene is released to soil, it will be subject to rapid volatilization near the surface and that which does not evaporate will be highly to very highly mobile in the soil and may leach to groundwater. If benzene is released to water, it will be subject to rapid volatilization. It will not be expected to significantly adsorb to sediment, bioconcentrate in aquatic organisms or hydrolyze. It may be subject to biodegradation.

**Environmental:** If benzene is released to the atmosphere, it will exist predominantly in the vapor phase. Gas-phase benzene will not be subject to direct photolysis but it will react with photochemically produced hydroxyl radicals with a half-life of 13.4 days. The reaction time in polluted atmospheres which contain nitrogen oxides or sulfur dioxide is accelerated with the half-life being reported as 4-6 hours. Benzene is fairly soluble in water and is removed from the atmosphere in rain.

**Physical:** Products of photooxidation include phenol, nitrophenols, nitrobenzene, formic acid, and peroxyacetyl nitrate.

Other: No information available.

# Section 13 - Disposal Considerations

Chemical waste generators must determine whether a discarded chemical is classified as a hazardous waste. US EPA guidelines for the classification determination are listed in 40 CFR Parts 261.3. Additionally, waste generators must consult state and local hazardous waste regulations to ensure complete and accurate classification.

RCRA P-Series: None listed.

**RCRA U-Series:** 

CAS# 71-43-2: waste number U019 (Ignitable waste, Toxic waste).

### Section 14 - Transport Information

	US DOT	Canada TDG
Shipping Name:	BENZENE	BENZENE
Hazard Class:	3.	3
UN Number:	UN1114	UN1114
Packing Group:	II	II
Additional Info:		FLASHPOINT -11 C

# Section 15 - Regulatory Information

#### **US FEDERAL**

#### **TSCA**

CAS# 71-43-2 is listed on the TSCA inventory.

#### **Health & Safety Reporting List**

None of the chemicals are on the Health & Safety Reporting List.

#### uemical Test Rules

None of the chemicals in this product are under a Chemical Test Rule.

#### Section 12b

None of the chemicals are listed under TSCA Section 12b.

#### **TSCA Significant New Use Rule**

None of the chemicals in this material have a SNUR under TSCA.

#### **CERCLA Hazardous Substances and corresponding RQs**

CAS# 71-43-2: 10 lb final RQ (receives an adjustable RQ of 10 lbs based on potential carcinoge

#### SARA Section 302 Extremely Hazardous Substances

None of the chemicals in this product have a TPQ.

#### **SARA Codes**

CAS # 71-43-2: immediate, delayed, fire.

#### Section 313

This material contains Benzene (CAS# 71-43-2, > 99%), which is subject to the reporting requirements of Section 313 of SARA Title III and 40 CFR Part 373.

#### Clean Air Act:

CAS# 71-43-2 is listed as a hazardous air pollutant (HAP).

This material does not contain any Class 1 Ozone depletors.

This material does not contain any Class 2 Ozone depletors.

#### Clean Water Act:

CAS# 71-43-2 is listed as a Hazardous Substance under the CWA. CAS# 71-43-2 is listed as a Priority Pollutant under the Clean Water 
Act. CAS# 71-43-2 is listed as a Toxic Pollutant under the Clean Water 
Act.

#### OSHA:

None of the chemicals in this product are considered highly hazardous by OSHA.

#### \_ (ATE

CAS# 71-43-2 can be found on the following state right to know lists: California, New Jersey, Pennsylvania, Minnesota, Massachusetts.

#### California Prop 65

# The following statement(s) is(are) made in order to comply with the California Safe Drinking Water Act:

WARNING: This product contains Benzene, a chemical known to the state of California to cause cancer.

WARNING: This product contains Benzene, a chemical known to the state of California to cause male reproductive toxicity.

California No Significant Risk Level: CAS# 71-43-2: 6.4 æg/day NSRL (oral); 13 æg/day NSRL (inhalation)

### **European/International Regulations**

# European Labeling in Accordance with EC Directives

#### **Hazard Symbols:**

ΤF

#### **Risk Phrases:**

R 11 Highly flammable.

R 36/38 Irritating to eyes and skin.

R 45 May cause cancer.

R 46 May cause heritable genetic damage.

R 48/23/24/25 Toxic: danger of serious damage to health by prolonged exposure through inhalation, contact with skin and if swallowed.

R 65 Harmful: may cause lung damage if swallowed.

#### Safety Phrases:

S 45 In case of accident or if you feel unwell, seek medical advice immediately (show the label where possible). S 53 Avoid exposure - obtain special instructions before use.

#### **7K (Water Danger/Protection)**

CAS# 71-43-2: 3

#### Canada - DSL/NDSL

CAS# 71-43-2 is listed on Canada's DSL List.

#### Canada - WHMIS

This product has a WHMIS classification of B2, D2A, D2B.

This product has been classified in accordance with the hazard criteria of the Controlled Products Regulations and the MSDS contains all of the information required by those regulations.

#### **Canadian Ingredient Disclosure List**

CAS# 71-43-2 is listed on the Canadian Ingredient Disclosure List.

### Section 16 - Additional Information

MSDS Creation Date: 6/11/1999 Revision #8 Date: 3/15/2007

The information above is believed to be accurate and represents the best information currently available to us. However, we make no warranty of merchantability or any other warranty, express or implied, with respect to such information, and we assume no liability resulting from its use. Users should make their own investigations to determine the suitability of the information for their particular purposes. In no event shall Fisher be liable for any claims, losses, or damages of any third party or for lost profits or any special, indirect, incidental, consequential or exemplary damages, howsoever arising, even if Fisher has been advised of the possibility of such damages.

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# Material Safety Data Sheet

Benzo(a)pyrene, 98%

#### ACC# 37175

### Section 1 - Chemical Product and Company Identification

MSDS Name: Benzo(a)pyrene, 98%

Catalog Numbers: AC105600000, AC105600010, AC105601000

**Synonyms:** 3,4-Benzopyrene; 3,4-Benzpyrene.

Company Identification:
Acros Organics N.V.

One Reagent Lane Fair Lawn, NJ 07410

For information in North America, call: 800-ACROS-01 For emergencies in the US, call CHEMTREC: 800-424-9300

# Section 2 - Composition, Information on Ingredients

CAS#	Chemical Name	Percent	EINECS/ELINCS
50-32-8	Benzo[a]pyrene	98.0	200-028-5

Hazard Symbols: T Risk Phrases: 45 46

### Section 3 - Hazards Identification

#### **EMERGENCY OVERVIEW**

Appearance: slightly brown solid. Cancer suspect agent. Possible risk of harm to the unborn child.

**Caution!** The toxicological properties of this material have not been fully investigated. May cause eye and skin irritation. May cause respiratory and digestive tract irritation.

Target Organs: None.

#### **Potential Health Effects**

**Eye:** May cause eye irritation. **Skin:** May cause skin irritation.

**Ingestion:** May cause irritation of the digestive tract. The toxicological properties of this substance have

not been fully investigated.

Inhalation: May cause respiratory tract irritation. The toxicological properties of this substance have not

been fully investigated.

Chronic: May cause cancer in humans.

# Section 4 - First Aid Measures

**Eyes:** Flush eyes with plenty of water for at least 15 minutes, occasionally lifting the upper and lower eyelids. Immediately flush eyes with plenty of water for at least 15 minutes, occasionally lifting the upper and lower eyelids. Get medical aid.

**Skin:** Get medical aid. Flush skin with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Wash clothing before reuse.

Ingestion: Never give anything by mouth to an unconscious person. Get medical aid. Do NOT induce

vomiting. If conscious and alert, rinse mouth and drink 2-4 cupfuls of milk or water.

Inhalation: Remove from exposure and move to fresh air immediately. If not breathing, give artificial

respiration. If breathing is difficult, give oxygen. Get medical aid.

Notes to Physician: Treat symptomatically and supportively.

# Section 5 - Fire Fighting Measures

**General Information:** As in any fire, wear a self-contained breathing apparatus in pressure-demand, MSHA/NIOSH (approved or equivalent), and full protective gear. During a fire, irritating and highly toxic gases may be generated by thermal decomposition or combustion.

**Extinguishing Media:** In case of fire, use water, dry chemical, chemical foam, or alcohol-resistant foam.

Use agent most appropriate to extinguish fire. Use water spray, dry chemical, carbon dioxide, or

appropriate foam.

Flash Point: Not available.

**Autoignition Temperature:** Not available. **Explosion Limits, Lower:** Not available.

Upper: Not available.

NFPA Rating: (estimated) Health: ; Flammability: ; Instability:

#### Section 6 - Accidental Release Measures

**General Information:** Use proper personal protective equipment as indicated in Section 8. **Spills/Leaks:** Clean up spills immediately, observing precautions in the Protective Equipment section. Sweep up, then place into a suitable container for disposal. Avoid generating dusty conditions. Provide ventilation.

### Section 7 - Handling and Storage

**Handling:** Wash thoroughly after handling. Use with adequate ventilation. Minimize dust generation and accumulation. Avoid contact with eyes, skin, and clothing. Keep container tightly closed. Avoid ingestion and inhalation.

**Storage:** Store in a tightly closed container. Store in a cool, dry, well-ventilated area away from incompatible substances.

# Section 8 - Exposure Controls, Personal Protection

**Engineering Controls:** Facilities storing or utilizing this material should be equipped with an eyewash facility and a safety shower. Use adequate ventilation to keep airborne concentrations low. Use adequate general or local exhaust ventilation to keep airborne concentrations below the permissible exposure limits.

**Exposure Limits** 

Chemical Name ACGIH			OSHA - Final PELs	
Į	Benzo[a]pyrene	none listed	none listed	none listed

**OSHA Vacated PELs:** Benzo[a]pyrene: No OSHA Vacated PELs are listed for this chemical. **Personal Protective Equipment** 

Eyes: Wear appropriate protective eyeglasses or chemical safety goggles as described by OSHA's eye and

face protection regulations in 29 CFR 1910.133 or European Standard EN166.

Skin: Wear appropriate protective gloves to prevent skin exposure.

**Clothing:** Wear appropriate protective clothing to prevent skin exposure.

**Respirators:** A respiratory protection program that meets OSHA's 29 CFR 1910.134 and ANSI Z88.2 requirements or European Standard EN 149 must be followed whenever workplace conditions warrant a respirator's use. Wear a NIOSH/MSHA or European Standard EN 149 approved full-facepiece airline respirator in the positive pressure mode with emergency escape provisions.

### Section 9 - Physical and Chemical Properties

Physical State: Solid

**Appearance:** slightly brown **Odor:** faint aromatic odor

pH: Not available.

Vapor Pressure: Not available. Vapor Density: Not available. Evaporation Rate: Not available.

Viscosity: Not available.

**Boiling Point:** 495 deg C @ 760.00mm Hg **Freezing/Melting Point:**175 - 177 deg C **Decomposition Temperature:**Not available.

Solubility: 1.60x10-3 mg/l @25°C

Specific Gravity/Density:Not available.

Molecular Formula:C20H12 Molecular Weight:252.31

# Section 10 - Stability and Reactivity

Chemical Stability: Stable under normal temperatures and pressures.

Conditions to Avoid: Incompatible materials, dust generation.

Incompatibilities with Other Materials: Strong oxidizing agents.

Hazardous Decomposition Products: Carbon monoxide, irritating and toxic fumes and gases, carbon

dioxide, acrid smoke and fumes.

Hazardous Polymerization: Has not been reported.

### Section 11 - Toxicological Information

RTECS#:

CAS# 50-32-8: DJ3675000

LD50/LC50: Not available.

### Carcinogenicity:

CAS# 50-32-8:

**ACGIH:** A2 - Suspected Human Carcinogen **California:** carcinogen, initial date 7/1/87

NIOSH: potential occupational carcinogen (listed as Coal tar pitches)

NTP: Suspect carcinogen

**OSHA:** Possible Select carcinogen **IARC:** Group 2A carcinogen

**Epidemiology:** No information available. **Teratogenicity:** No information available.

Reproductive Effects: No information available.

**Neurotoxicity:** No information available. **Mutagenicity:** No information available.

Other Studies: No data available.

### Section 12 - Ecological Information

Ecotoxicity: No data available. No information available.

**Environmental:** No information found.

**Physical:** No information found. **Other:** No information available.

### Section 13 - Disposal Considerations

Chemical waste generators must determine whether a discarded chemical is classified as a hazardous waste. US EPA guidelines for the classification determination are listed in 40 CFR Parts 261.3. Additionally, waste generators must consult state and local hazardous waste regulations to ensure complete and accurate classification.

RCRA P-Series: None listed.

RCRA U-Series: CAS# 50-32-8: waste number U022.

### Section 14 - Transport Information

	US DOT	IATA	RID/ADR	IMO	Canada TDG
Shipping Name:	DOT regulated - small quantity provisions apply (see 49CFR173.4)				No information available.
Hazard Class:				÷ .	
UN Number:		•		·	
Packing Group:					

# Section 15 - Regulatory Information

#### **US FEDERAL**

#### TSCA

CAS# 50-32-8 is listed on the TSCA inventory.

#### **Health & Safety Reporting List**

None of the chemicals are on the Health & Safety Reporting List.

#### **Chemical Test Rules**

None of the chemicals in this product are under a Chemical Test Rule.

#### Section 12b

None of the chemicals are listed under TSCA Section 12b.

#### TSCA Significant New Use Rule

None of the chemicals in this material have a SNUR under TSCA.

#### SARA

#### **CERCLA Hazardous Substances and corresponding RQs**

CAS# 50-32-8: 1 lb final RQ; 0.454 kg final RQ

#### **SARA Section 302 Extremely Hazardous Substances**

None of the chemicals in this product have a TPQ.

#### **SARA Codes**

CAS # 50-32-8: acute, chronic.

#### Section 313

This material contains Benzo[a]pyrene (CAS# 50-32-8, 98 0%), which is subject to the reporting requirements of Section 313 of SARA Title III and 40 CFR Part 373.

#### Clean Air Act:

This material does not contain any hazardous air pollutants. This material does not contain any Class 1 Ozone depletors. This material does not contain any Class 2 Ozone depletors.

#### Clean Water Act:

None of the chemicals in this product are listed as Hazardous Substances under the CWA. CAS# 50-32-8 is listed as a Priority Pollutant under the Clean Water Act. None of the chemicals in this product are listed as Toxic Pollutants under the CWA.

#### OSHA:

None of the chemicals in this product are considered highly hazardous by OSHA.

#### STATE

CAS# 50-32-8 can be found on the following state right to know lists: California, New Jersey, Pennsylvania, Minnesota, Massachusetts.

The following statement(s) is(are) made in order to comply with the California Safe Drinking Water Act: WARNING: This product contains Benzo[a]pyrene, a chemical known to the state of California to cause cancer. California No Significant Risk Level: CAS# 50-32-8: 0.06 æg/day NSRL

# **European/International Regulations European Labeling in Accordance with EC Directives Hazard Symbols:**

Т

#### **Risk Phrases:**

R 45 May cause cancer.

R 46 May cause heritable genetic damage.

R 60 May impair fertility.

R 61 May cause harm to the unborn child.

#### **Safety Phrases:**

S 37 Wear suitable gloves.

S 45 In case of accident or if you feel unwell, seek medical advice immediately (show the label where possible).

S 53 Avoid exposure - obtain special instructions

S 28A After contact with skin, wash immediately with plenty of water.

#### WGK (Water Danger/Protection)

CAS# 50-32-8: No information available.

#### Canada - DSL/NDSL

CAS# 50-32-8 is listed on Canada's DSL List.

#### Canada - WHMIS

This product has a WHMIS classification of D2A.

#### **Canadian Ingredient Disclosure List**

CAS# 50-32-8 is listed on the Canadian Ingredient Disclosure List.

#### **Exposure Limits**

CAS# 50-32-8: OEL-AUSTRALIA; Carcinogen OEL-BELGIUM; Carcinogen OEL-FINLAND: TWA 0.01 mg/m3; Skin; Carcinogen OEL-FRANCE; Carcinogen OEL-GER

MANY; Carcinogen OEL-RUSSIA: STEL 0.00015 mg/m3; Carcinogen OEL-SWEDEN: TWA 0.005 mg/m3; STEL 0.03 mg/m3; Skin OEL IN BULGARIA, COLOMBIA, JORDA N, KOREA check ACGIH TLV OEL IN NEW ZEALAND, SINGAPORE, VIETNAM check ACGI TLV

### Section 16 - Additional Information

MSDS Creation Date: 9/02/1997 Revision #5 Date: 3/18/2003

The information above is believed to be accurate and represents the best information currently available to us. However, we make no warranty of merchantability or any other warranty, express or implied, with respect to such information, and we assume no liability resulting from its use. Users should make their own investigations to determine the suitability of the information for their particular purposes. In no event shall Fisher be liable for any claims, losses, or damages of any third party or for lost profits or any special, indirect, incidental, consequential or exemplary damages, howsoever arising, even if Fisher has been advised of the possibility of such damages.

• *	

#### MATERIAL SAFETY DATA SHEET

1CD GROUP METALS LLC **600 MADISON AVENUE NEW YORK, N.Y. 10022** 

TEL: 212-644-1500 FAX: 212-644-1480

FOR EMERGENCY CALL: CHEMTREC 1-800-424-9300

HMIS

HEALTH: 4	FLAMMABILITY: 0
REACTIVITY: 1	PROTECTION: H

#### SECTION A - PRODUCT INFORMATION

TRADE NAME: CAS NUMBER: ARSENIC METAL

**REVISION DATE: AUGUST 1, 2001** 

: 2MYMONY3 CHEMICAL FAMILY: 7440-38-2 ARSENIC MFTAL METALS GROUP 5a

FORMULA:

SECTION B - HAZARDOUS COMPONENTS

COMPONENT

CAS NO.

PELITLY

ARSENIC METAL

7440-38-2

100 %

0,01mg(As)m<sup>5</sup> OSHA TWA CFR 29

(1910.1018)

0.5mg/m3 OSHA TWA (AS INORGANIC COMPOUND

> 0.2mg/m3 ACGIH TWA 0.002mg/m³/15 min NIOSH 100mg/m3 IDLH-CARCINGGEN

#### SECTION C - PHYSICAL PROPERTIES

BOILING POINT (° C):

SUBLIMES @ 812

SPECIFIC GRAVITY: 5.727

FREEZING POINT (°): N/A

MELTING POINT (° C): VAPOR PRESSURE (mm Hgl:

814 @ 36 ATM

VAPOR DENSITY (AIR = 1):

1 mm @ 372 ° C

PERCENT VOLATILE (BY WT.):

N/A

N/A EVAPORATION RATE: N/A

SOLUBILITY IN WATER:

INSOLURI F

Ph ( % IN WATER): NONE

ODOR THRESHOLD:

APPEARANCE & ODOR:

N/A

SILVER GRAY CRYSTALS/NO ODOR AS METAL, AS COMPOUND ASH, HAS GARLIC

RODO

#### SECTION D - FIRE & EXPLOSION DATA

FLASH POINT (\* ):

FLAMMABLE LIMITS: **EXTENGUISHING MEDIA:**  LEL : (N/A) WATER: (X) UEL: (N/A) FOAM: (X)

CO<sub>z</sub>: (X)

AUTO IGNITION TEMP (° F): (N/A) DRY CHEMICAL: (XI

SPECIAL FIRE FIGHTING PROCEDURES:

RESTRICT PERSONS NOT WEARING PROTECTIVE EQUIPMENT FROM AREA. TRY TO SNUFF FIRE WITH SAND, DRY MEDIA, FOAM, OR CO2. IF NO OTHER OPTIONS AVAILABLE, USE WATER & ALWAYS WEAR SELF CONTAINED BREATHING APPARATUS OR MIOSH TOXIC

VAPOR RESPIRATOR.

UNUSUAL FIRE & EXPLOSION HAZARDS: HEATED ARSENIC IN CONTACT WITH ACID OR WATER VAPOR CAN PRODUCE HIGHLY TOXIC FUMES. ASENIC REACTS VIGOROUSLY WITH OXIDIZING MATERIALS.

JUCHT SPILL, CONTACT FXPI OSION HAZARD IN THE FORM OF DUST. IN THE EVENT OF A FIRE OR THE STATE DEPT. OF THE ENVIRONMENT & YOUR ENVIRONMENTAL PROTECTION AGENCY.

REGIONAL OFFICE OF THE FEDERAL

#### SECTION E - REACTIVITY DATA

STABILITY: STABLE

INCOMPATIBILITY: ACIDS, OXIDIZING AGENTS SUCH AS HALOGENS, PERCHLORATES, PEROXIDES,

PERMANGANATES, CHLORATES, NITRATES, HYDROCHLORIC ACID, SULFURIC ACID, NITRIC ACID, BROMINE AZIDE, DIRUBIDIUM ACETYLIDE, ZINC, NCI, NITRATES, Na.O.,

HEXAFLUORO ISOPROPYL IDEMEAMING LITHIUM

HAZARDOUS OF COMPOSITION PRODUCTS: ARSENIC FUMES, ARSINE, OTHER ARSENIC COMPOUNDS

HAZARDOUS POLYMERIZATION: WILL NOT OCCUR

CONDITIONS TO AVOID: AVOID OPEN CONTAINERS SITTING IN HUMID ATMOSPHERE

#### SECTION F - PERSONAL PROTECTIVE EQUIPMENT INFO

RESPIRATORY EQUIPMENT: NIOSH APPROVED TOXIC VAPOR RESPIRATOR TO PARTICULATE & FUME AIR LEVEL

PROTECTIVE GLOVES: NEOPRENE OR PLASTIC EYE PROTECTION: GOGGLES/GLASSES

VENTILATION: LOCAL EXHAUST/MECHANICAL (GENERAL) SCRUBBER OR TRAP IF POSSIBLE

OTHER PROTECTIVE EQUIPMENT: LAB COAT

#### SECTION G - HEALTH HAZARD DATA

THRESHOLD LIMIT VALUE: AIR 0.002mg/m3/15min, HUMAN TDLo 7857 mg/kg/55Y :SKN(:GIT ORAL)

PRIMARY ROUTES OF EXPOSURE: INHALATION/INGESTION/SKIN

ORAL LDso: NO, ORAL RAT TDLo 605ug/kg, ORAL-MAN TDLo 7857mg/kg/55y SKN

DERMAL IRRITATION-RABBIT: NO, SUBCUTANEOUS IMPLANT RABBIT TDLo 75mg/kg

EYE (RRITATION-RABBIT: NO

OSHA PEL: 0.01mg/m³

ACGIH TLV: 0,2mg/m3

EFFECTS OF OVEREXPOSURE: ORAL - NAUSEA, COLD SWEATS, VOMITING, DIARRHEA, BLOOD STOOLS, COLLAPSE.

SHOCK, LOSS OF APPETITE, CRAMPS, JAUNDICE, SKIN ABNORMALITIES

KNOWN EFFECT'S ON OTHER ILLNESSES: GASTROINTESTINAL, NERVOUS SYSTEM, LIVER & KIDNEY PROBLEME, AFTER EXPOSURE

HAVE URINE TEST.

LISTED CARCINOGEN: NONE (JOSHA (Y) NTP (Y) IARC (Y) OTHER (Y)

#### SECTION H - EMERGENCY & FIRST AID DATA

SKIN : FLUSH WITH SOAP WATER, AVOID RUBBING INTO THE SKIN. CONTACT PHYSICIAN IMMEDIATELY.

EYES: FLUSH WITH WATER FOR 15 MINUTES. CONTACT PHYSICIAN IMMEDIATELY.

INHALATION: REMOVE TO FRESH AIR. PROVIDE OXYGEN IF NECESSARY. CONTACT PHYSICIAN IMMEDIATELY.
INGESTION: TREATMENT WITH BASIDIMERCAPTOLI IS OF QUESTIONABLE EFFECTIVENESS IN TRIVALENT ARSENI

ESTION: TREATMENT WITH BASIDIMERCAPTOLI IS OF QUESTIONABLE EFFECTIVENESS IN TRIVALENT ARSENIC COMPOUNDS. INDUCE VOMITING & DO GASTRIC LAVAGE. GE! PERSONNEL TO HOSPITAL IMMEDIATELY. A PHYSICIAN CAN INITIATE AN EXCHANGE TRANSFUSION & DIALYSIS. ALSO

ADSORPTION & REMOVAL WITH ANIMAL BONE COAL OR Fe(OH),

#### SECTION I - SPILL & DISPOSAL INFORMATION

STEPS TO BE TAKEN IN CASE OF SPILL OR LEAK:

WEARING RESPIRATOR, GLOVES, GOGGLES, LAB COAT, GATHER UP CHUNKS, GRANULES, OR POWDER WITH VACUUM OR UTENSILS RESERVED FOR POISONOUS SOLIDS.

WASTE DISPOSAL INFORMATION:

SOLID WASTES SHOULD BE VITRIFIED, PLACED IN LABELLED CONTAINER & BURIED IN A EPA SUPERVISED FACILITY. ETCHING SOLUTIONS & CUTTING WASTES SHOULD BE PRECIPITATED. CEMENTED/VITRIFIED & BURIED IN METALIPLASTIC LABELLED CONTAINERS & BURIED IN EPA SUPERVISED FACILITY. PASS GAS THROUGH POTASSIUM PERMANGANATE, PRECIPITATE & TREAT AS ABOVE.

RCRA HAZARDOUS WASTE :NO () YES (X) RCRA #: () D004

CERCLA: NO () YES (X) RO (1 LB.)

FOLLOW ALL LOCAL, STATE AND FEDERAL INFORMATION AND REGULATIONS

#### SECTION J - OTHER REGULATORY INFORMATION

WE CERTIFY THAT ALL COMPONENTS OF THIS PRODUCT ARE REGISTERED UNDER THE REGULATIONS OF THE TOXIC SUBSTANCES CONTROL ACT.

SARA TITLE III, SECT. 313: LISTED (X) NOT LISTED ()

DOT REGULATED : YES: (X) NO: ()

8Q: (1 LB.) UN/NA NO.: (X) UN 1558

PROPER SHIPPING NAME: ARSENIC

**EMERGENCY RESPONSE GUIDE NO.: 53** 

HAZARD CLASSIFICATION: (X) POISON 6.1

LABEL: (X) POISON PG II

#### SECTION K - SPECIAL PRECAUTIONS

#### FOR INDUSTRIAL USE ONLY

HANDLING & STORAGE INFORMATION: PRIOR TO WORKING WITH ARSENIC, PERSONNEL SHOULD BE TRAINED IN PROPER HANDLING & STORAGE. STORE IN ORIGINAL PACKAGING IN COOL DRY AREA. WHEN HANDLING PLACE INTO INERT ATOMSPHERE IMMEDIATELY. WEAR RESPIRATORY PROTECTION, GLOVES & EYE PROTECTION. OTHER PRECAUTIONS : MINIMUM - HAVE QUARTERLY MEDICAL CHECKS INCLUDING URINE TESTS OF PERSONNEL WORKING WITH ARSENIC OR ARSENIC COMPOUNDS. DO NOT SMOKE OR EAT IN WORK AREA.

IN ACCORDANCE WITH GOOD PRACTICES OF PERSONAL HYGIENE, HANDLE WITH DUE CARE AND AVOID ANY UNNECESSARY CONTACT WITH THIS PRODUCT.

THIS INFORMATION IS BEING SUPPLIED TO YOU UNDER OSHA "RIGHT TO KNOW" REGULATION 29 CFR 1810.1260 AND IS OFFERED IN GOOD FAITH AS TYPICAL VALUES AND NOT AS PRODUCT SPECIFICATION. THE INFORMATION IS BELIEVED TO BE TRUE AND ACCURATE. NO WARRANTY, EXPRESSED OR INFIDED, REGARDING THE ACCURACY OF THIS DATA. THE HAZARD CONNECTED WITH USE OF THE MATERIAL. OR THE RESULTS TO BE OBTAINED FROM THE USE THEREOF, IS MADE. ICO GROUP METALS LIC AND ITS SUPPLIERS ASSUME NO RESPONSIBILITY FOR DAMAGE OR INJURY FROM THE USE OF THE PRODUCT DESCRIBED HEREIN.

ICD GROUP METALS LLC

MSDS Number: A7512 \* \* \* \* \* Effective Date: 02/16/06 \* \* \* \* \* Supercedes: 05/08/03



Material Safety Data Sheet

From: Mallinckrodt Baker, Inc. 222 Red School Lane Phillipsburg, NJ 08865



24 Hour Emergency Telephone: 908-859-2151

CHEMTREC: 1-800-424-9300

National Response in Canada CANUTEC: 613-996-6666

Outside U.S. And Canada Chemtrec: 703-527-3887

NOTE: CHEMTREC, CANUTEC and National Response Center emergency numbers to be used only in the event of chemical emergencies involving a spill, leak, fire, exposure or accident involving chemicals.

All non-emergency questions should be directed to Customer Service (1-800-582-2537) for assistance.

# ARSENIC TRIOXIDE

### 1. Product Identification

Synonyms: Arsenic (III) oxide; arsenic sesquioxide; arsenous trioxide, white arsenic

**CAS No.:** 1327-53-3

Molecular Weight: 197.84 Chemical Formula: As2O3

**Product Codes: 0061** 

# 2. Composition/Information on Ingredients

Ingredient	•	•	CAS No	Percent	Hazardous
		 			. , <del></del>
Arsenic Trioxide			132753-3	99 - 100%	Yes

### 3. Hazards Identification

**Emergency Overview** 

DANGER! MAY BE FATAL IF SWALLOWED OR INHALED. CANCER HAZARD. CONTAINS INORGANIC ARSENIC WHICH CAN CAUSE CANCER. Risk of cancer depends on duration and level of exposure. CAUSES IRRITATION TO SKIN, EYES AND RESPIRATORY TRACT. MAY CAUSE LIVER AND KIDNEY DAMAGE. USE ONLY WITH ADEQUATE VENTILATION AND RESPIRATORY EQUIPMENT.

# J.T. Baker SAF-T-DATA<sup>(tm)</sup> Ratings (Provided here for your convenience)

Health Rating: 4 - Extreme (Cancer Causing)

Flammability Rating: 0 - None Reactivity Rating: 1 - Slight Contact Rating: 1 - Slight

Lab Protective Equip: GOGGLES; LAB COAT; PROPER GLOVES

Storage Color Code: Blue (Health)

#### **Potential Health Effects**

#### Inhalation:

Arsenic may cause inflammation of the mucous membranes with cough and foamy sputum, restlessness, dyspnea, cyanosis, and rales. Symptoms like those from ingestion exposure may follow. May cause pulmonary edema.

### Ingestion:

Arsenic is highly toxic! May cause burning in esophagus, vomiting, and bloody diarrhea. Symptoms of cold and clammy skin, low blood pressure, weakness, headache, cramps, convulsions, and coma may follow. May cause damage to liver and kidneys. A suspected fetal toxin. Death may occur from circulatory failure. Estimated lethal dose 120 milligrams.

#### **Skin Contact:**

May cause irritation, symptoms including redness, itching, and pain.

#### **Eye Contact:**

May cause irritation with itching, burning, watering of eyes; may cause conjunctiva damage.

#### **Chronic Exposure:**

Arsenic on repeated or prolonged skin contact may cause bronzing of the skin, edema, dermatitis, and lesions. Repeated or prolonged inhalation of dust may cause damage to the nasal septum. Chronic exposure from inhalation or ingestion may cause hair and weight loss, a garlic odor to the breath and perspiration, excessive salivation and perspiration, central nervous system damage, hepatitis, gastrointestinal disturbances, cardiovascular damage, and kidney and liver damage. Arsenic compounds are known human carcinogens and may be teratogenic based on effects in laboratory animals.

#### Aggravation of Pre-existing Conditions:

No information found.

# 4. First Aid Measures

#### Inhalation:

Remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. Get medical attention.

#### Ingestion:

Induce vomiting immediately as directed by medical personnel. Never give anything by mouth to an unconscious person. Get medical attention immediately.

#### **Skin Contact:**

Immediately flush skin with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Get medical attention immediately. Wash clothing before reuse. Thoroughly clean shoes before reuse. Contaminated work clothes should be laundered by individuals who have been informed of the hazards of exposure to this substance.

**Eye Contact:** 

Immediately flush eyes with plenty of water for at least 15 minutes, lifting lower and upper eyelids occasionally. Get medical attention immediately.

Note to Physician:

If emesis if unsuccessful after two doses of Ipecac, consider gastric lavage. Monitor urine arsenic level. Alkalization of urine may help prevent disposition of red cell breakdown products in renal tubular cells. If acute exposure is significant, maintain high urine output and monitor volume status, preferably with central venous pressure line. Abdominal X-rays should be done routinely for all ingestions. Chelation therapy with BAL, followed by n-penicillamine is recommended, but specific dosing guidelines are not clearly established.

# 5. Fire Fighting Measures

#### Fire:

Not considered to be a fire hazard. Toxic fumes of arsenic trioxide and arsine may be formed in fire situations.

#### **Explosion:**

Not considered to be an explosion hazard.

### Fire Extinguishing Media:

Use any means suitable for extinguishing surrounding fire.

#### **Special Information:**

In the event of a fire, wear full protective clothing and NIOSH-approved self-contained breathing apparatus with full facepiece operated in the pressure demand or other positive pressure mode.

### 6. Accidental Release Measures

Ventilate area of leak or spill. Wear appropriate personal protective equipment as specified in Section 8. Spills: Sweep up and containerize for reclamation or disposal. Vacuuming or wet sweeping may be used to avoid dust dispersal. US Regulations (CERCLA) require reporting spills and releases to soil, water and air in excess of reportable quantities. The toll free number for the US Coast Guard National Response Center is (800) 424-8802.

# 7. Handling and Storage

Keep in a tightly closed container, stored in a cool, dry, ventilated area. Protect against physical damage. Isolate from incompatible substances. Wear special protective equipment (Sec. 8) for maintenance break-in or where exposures may exceed established exposure levels. Wash hands, face, forearms and neck when exiting restricted areas. Shower, dispose of outer clothing, change to clean garments at the end of the day. Avoid cross-contamination of street clothes. Wash hands before eating and do not eat, drink, or smoke in workplace. Containers of this material may be hazardous when empty since they retain product residues (dust, solids); observe all warnings and precautions listed for the product.

# 8. Exposure Controls/Personal Protection

**Airborne Exposure Limits:** 

-OSHA Permissible Exposure Limit (PEL):

10 ug(As)/m3 ppm (TWA)

-ACGIH Threshold Limit Value (TLV):

0.01 mg(As)/m3 (TWA),

listed as A1, confirmed human carcinogen.

Ventilation System:

A system of local and/or general exhaust is recommended to keep employee exposures below the Airborne Exposure Limits. Local exhaust ventilation is generally preferred because it can control the emissions of the contaminant at its source, preventing dispersion of it into the general work area. Please refer to the ACGIH document, *Industrial Ventilation*, *A Manual of Recommended Practices*, most recent edition, for details.

Personal Respirators (NIOSH Approved):

If the exposure limit is exceeded, a half-face high efficiency dust/mist respirator may be worn for up to ten times the exposure limit or the maximum use concentration specified by the appropriate regulatory agency or respirator supplier, whichever is lowest. A full-face piece high efficiency dust/mist respirator may be worn up to 50 times the exposure limit, or the maximum use concentration specified by the appropriate regulatory agency or respirator supplier, whichever is lowest. For emergencies or instances where the exposure levels are not known, use a full-facepiece positive-pressure, air-supplied respirator. WARNING: Air-purifying respirators do not protect workers in oxygen-deficient atmospheres.

#### Skin Protection:

Wear impervious protective clothing, including boots, gloves, lab coat, apron or coveralls, as appropriate, to prevent skin contact.

**Eye Protection:** 

Use chemical safety goggles and/or full face shield where dusting or splashing of solutions is possible. Maintain eye wash fountain and quick-drench facilities in work area.

#### **Other Control Measures:**

Any area where inorganic arsenic is stored, handled, used, etc., must be established as a 'Regulated Area' with controlled access, limited to authorized persons. Containers of inorganic arsenic and Regulated Areas must be labeled to show a CANCER SUSPECT AGENT is present. Eating, drinking, and smoking should not be permitted in areas where solids or liquids containing arsenic or lead compounds are handled, processed, or stored. See OSHA substance-specific standard for more information on personal protective equipment, engineering and work practice controls, medical surveillance, record keeping, and reporting requirements. (arsenic: 29 CFR 1910.1018; lead: 29 CFR 1910.1025).

# 9. Physical and Chemical Properties

Appearance:

Transparent crystals, or white powder.

Odor:

Odorless.

Solubility:

3.7 g/100 ml water @ 20C (68F)

**Specific Gravity:** 

3.74

pH:

No information found.

% Volatiles by volume @ 21C (70F):

0

**Boiling Point:** 

465C (869F)

**Melting Point:** 

315C (599F)

Vapor Density (Air=1):

No information found.

Vapor Pressure (mm Hg):

No information found.

Evaporation Rate (BuAc=1):

No information found.

# 10. Stability and Reactivity

Stability:

Stable under ordinary conditions of use and storage.

**Hazardous Decomposition Products:** 

Emits toxic fumes of arsenic when heated to decomposition.

Hazardous Polymerization:

Will not occur.

Incompatibilities:

Oxidizers, tannic acid, infusion cinchona and other vegetable astringent infusions and decoctions, iron solutions, rubidium carbide, chlorine trifluoride, fluorine, hydrogen fluoride, oxygen difluoride, acids, bases, sodium chlorate, zinc filings, other reactive metals and mercury. Corrosive to metals in the presence of moisture.

Conditions to Avoid:

Incompatibles.

# 11. Toxicological Information

Toxicological Data:

Oral rat LD50: 14.6 mg/kg; investigated as a mutagen, tumorigen, reproductive effector.

Reproductive Toxicity:

Has shown teratogenic effects in laboratory animals.

# 2. Ecological Information

**Environmental Fate:** 

When released into the soil, this material may biodegrade to a moderate extent. When released into water, this material may biodegrade to a moderate extent. This material is not expected to significantly bioaccumulate.

**Environmental Toxicity:** 

No information found.

# 13. Disposal Considerations

Whatever cannot be saved for recovery or recycling should be handled as hazardous waste and sent to a RCRA approved waste facility. Processing, use or contamination of this product may change the waste management options. State and local disposal regulations may differ from federal disposal regulations. Dispose of container and unused contents in accordance with federal, state and local requirements.

# 14. Transport Information

Domestic (Land, D.O.T.)

Proper Shipping Name: RQ, ARSENIC TRIOXIDE

Hazard Class: 6.1 UN/NA: UN1561 Packing Group: II

Information reported for product/size: 500G

International (Water, I.M.O.)

Proper Shipping Name: ARSENIC TRIOXIDE

Hazard Class: 6.1 UN/NA: UN1561 Packing Group: II

Information reported for product/size: 500G

International (Air, I.C.A.O.)

Proper Shipping Name: ARSENIC TRIOXIDE

Hazard Class: 6.1 UN/NA: UN1561 Packing Group: II

Information reported for product/size: 500G

# 15. Regulatory Information

\Chemical	Inventory	Status -	Part	1\				
Ingredient		•			TSCA	EC	Japan	Australia
Arsenic Trioxide	(1327-53-3)				Yes	Yes	Yes	Yes

Chemical Inventory Status - Part		Cana	da NDSL Phil.	•
Ingredient 				
Arsenic Trioxide (1327-53-3)			No Yes	
\Federal, State & International R	egulations-	Part 1\-		
			-SARA 313	
Ingredient	RQ TPQ	Llsį	Chemical C	alg
Arsenic Trioxide (1327-53-3)	1 100	No	Arsenic co	qm
\Federal, State & International R	tegulations-		-TSCA-	<del>-</del>
Ingredient	CERCLA			
Arsenic Trioxide (1327-53-3)	1.	P012	No	
	•			•

SARA 311/312: Acute: Yes Chronic: Yes

(Pure / Solid) Reactivity: No

#### **WARNING:**

THIS PRODUCT CONTAINS CHEMICALS KNOWN TO THE STATE OF CALIFORNIA TO CAUSE CANCER AND BIRTH DEFECTS OR OTHER REPRODUCTIVE HARM.

Australian Hazchem Code: 2Z

Poison Schedule: S6

WHMIS:

This MSDS has been prepared according to the hazard criteria of the Controlled Products Regulations (CPR) and the MSDS contains all of the information required by the CPR.

# 16. Other Information

NFPA Ratings: Health: 3 Flammability: 0 Reactivity: 0

Label Hazard Warning:

DANGER! MAY BE FATAL IF SWALLOWED OR INHALED. CANCER HAZARD. CONTAINS INORGANIC ARSENIC WHICH CAN CAUSE CANCER. Risk of cancer depends on duration and level of exposure. CAUSES IRRITATION TO SKIN, EYES AND RESPIRATORY TRACT. MAY CAUSE LIVER AND KIDNEY DAMAGE. USE ONLY WITH ADEOUATE VENTILATION AND RESPIRATORY EQUIPMENT.

#### Label Precautions:

Do not get in eyes, on skin, or on clothing.

Do not breathe dust.

Keep container closed.

Use only with adequate ventilation.

Wash thoroughly after handling.

#### Label First Aid:

If swallowed, induce vomiting immediately as directed by medical personnel. Never give anything by mouth to an unconscious person. Get medical attention immediately. If inhaled, remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. In case of contact, immediately flush eyes or skin with plenty of water for at least 15 minutes. Remove contaminated clothing and shoes. Wash clothing before reuse. In all cases, get medical attention.

**Product Use:** 

Laboratory Reagent.

**Revision Information:** 

No Changes.

Mallinckrodt Baker, Inc. provides the information contained herein in good faith but makes no representation as to its comprehensiveness or accuracy. This document is intended only as a guide to the appropriate precautionary handling of the material by a properly trained person using this product. Individuals receiving the information must exercise their independent judgment in determining its appropriateness for a particular purpose. MALLINCKRODT BAKER, INC. MAKES NO REPRESENTATIONS OR WARRANTIES, EITHER EXPRESS OR IMPLIED, INCLUDING WITHOUT LIMITATION ANY WARRANTIES OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE WITH RESPECT TO THE INFORMATION SET FORTH HEREIN OR THE PRODUCT TO WHICH THE INFORMATION REFERS. ACCORDINGLY, MALLINCKRODT BAKER, INC. WILL NOT BE RESPONSIBLE FOR DAMAGES RESULTING FROM USE OF OR RELIANCE UPON THIS INFORMATION.

Prepared by: Environmental Health & Safety Phone Number: (314) 654-1600 (U.S.A.)

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# Material Safety Data Sheet

Cadmium metal, granular

#### CC# 03720

### Section 1 - Chemical Product and Company Identification

MSDS Name: Cadmium metal, granular

Catalog Numbers: AC612135000, S79935, C3-500

Synonyms: None.

**Company Identification:** 

Fisher Scientific 1 Reagent Lane Fair Lawn, NJ 07410

For information, call: 201-796-7100 Emergency Number: 201-796-7100

For CHEMTREC assistance, call: 800-424-9300

For International CHEMTREC assistance, call: 703-527-3887

### Section 2 - Composition, Information on Ingredients

CAS#	Chemical Name	Percent	EINECS/ELINCS
7440-43-9	Cadmium	100	231-152-8

### Section 3 - Hazards Identification

#### **EMERGENCY OVERVIEW**

Appearance: silver white granules.

**Danger!** Flammable solid. May be fatal if inhaled. Harmful if swallowed. Causes eye, skin, and respiratory tract irritation. Contains cadmium. Cancer hazard. Avoid creating dust. Can cause lung and kidney disease. Inhalation of fumes may cause metal-fume fever. Air sensitive. May cause reproductive and fetal effects.

Target Organs: Blood, kidneys, liver, lungs, skeletal structures, prostate.

#### **Potential Health Effects**

Eye: Causes eye irritation. Skin: Causes skin irritation;

Ingestion: Harmful if swallowed. May cause gastrointestinal irritation with nausea, vomiting and diarrhea.

Ingestion may produce fluid loss, acute renal failure, and cardiopulmonary depression.

**Inhalation:** May be fatal if inhaled. Inhalation of fumes may cause metal fume fever, which is characterized by flu-like symptoms with metallic taste, fever, chills, cough, weakness, chest pain, muscle pain and increased white blood cell count. Damage may be delayed. May cause nausea, vomiting, abdominal pain, diarrhea, chest tightness, weakness, and delayed pulmonary edema. In humans inhalation causes proteinuria, an excess of protein in the urine.

**Chronic:** May cause respiratory tract cancer. Repeated inhalation may cause chronic bronchitis. Chronic inhalation may cause nasal septum ulceration and perforation. Cadmium and compounds may cause lung, wer and kidney damage and lung and prostate cancer in humans. May cause loss of smell, emphysema, nemia, bone demineralization, and lung fibrosis. The primary target organ for chronic cadmium disease is clearly the kidney.

#### Section 4 - First Aid Measures

res: Immediately flush eyes with plenty of water for at least 15 minutes, occasionally lifting the upper and lower eyelids. Get medical aid.

**Skin:** Get medical aid. Flush skin with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Wash clothing before reuse.

Ingestion: Do not induce vomiting. If victim is conscious and alert, give 2-4 cupfuls of milk or water.

Never give anything by mouth to an unconscious person. Get medical aid immediately.

**Inhalation:** POISON material. If inhaled, get medical aid immediately. Remove victim to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. Attempt rescue only after notifying at least one other person of the emergency and putting into effect established emergency procedures. Do not become a casualty yourself.

**Notes to Physician:** Administration of calcium disodium EDTA may be useful in acute poisoning with its use at the discretion of qualified medical personnel. Persons with kidney disease, chronic respiratory disease, liver disease, or skin disease may be at increased risk from exposure to this substance.

### Section 5 - Fire Fighting Measures

**General Information:** As in any fire, wear a self-contained breathing apparatus in pressure-demand, MSHA/NIOSH (approved or equivalent), and full protective gear. Material can spontaneously ignite (pyrophoric) when exposed to air at normal or slightly elevated temperatures. Dust can be an explosion hazard when exposed to heat or flame. Flammable solid. May burn rapidly with flare burning effect. May re-ignite after fire is extinguished. Dangerous fire hazard in the form of dust when exposed to heat or flame.

**Extinguishing Media:** Use dry sand, graphite powder, dry sodium chloride-based extinguishers.

ash Point: Not available.

Autoignition Temperature: Not available. Explosion Limits, Lower: Not available.

Upper: Not available.

NFPA Rating: (estimated) Health: 4; Flammability: 2; Instability: 1

### Section 6 - Accidental Release Measures

**General Information:** Use proper personal protective equipment as indicated in Section 8. **Spills/Leaks:** Vacuum or sweep up material and place into a suitable disposal container. Avoid generating dusty conditions. Remove all sources of ignition. Use a spark-proof tool. Provide ventilation. Place under an inert atmosphere.

# Section 7 - Handling and Storage

Handling: Wash thoroughly after handling. Wash thoroughly after handling. Remove contaminated clothing and wash before reuse. Minimize dust generation and accumulation. Use spark-proof tools and explosion proof equipment. Avoid contact with skin and eyes. Do not breathe dust, vapor, mist, or gas. Empty containers retain product residue, (liquid and/or vapor), and can be dangerous. Keep away from eat, sparks and flame. Do not ingest or inhale. Handle under an inert atmosphere. Store protected from \_ir. Use only in a chemical fume hood. Do not pressurize, cut, weld, braze, solder, drill, grind, or expose empty containers to heat, sparks or open flames.

Storage: Keep away from heat and flame. Keep away from sources of ignition. Store in a tightly closed

container. Store in a cool, dry, well-ventilated area away from incompatible substances. Do not expose to air. Store under an inert atmosphere.

# Section 8 - Exposure Controls, Personal Protection

**Engineering Controls:** Use explosion-proof ventilation equipment. Facilities storing or utilizing this material should be equipped with an eyewash facility and a safety shower. Use only under a chemical fume hood. See 29CFR 1910.1027 for regulations applying to all occupational exposures to cadmium and cadmium compounds, in all forms.

**Exposure Limits** 

Chemical Name	ACGIH	NIOSH	OSHA - Final PELs	
Cadmium	0.01 mg/m3 TWA; 0.002 mg/m3 TWA (respirable fraction)	9 mg/m3 IDLH (dust and fume)	0.2 mg/m3 TWA (dust); 0.1 mg/m3 TWA (fume); 0.6 mg/m3 Ceiling (dust); 0.3 mg/m3 Ceiling (fume); 2.5 æg/m3 Action Level; 5 æg/m3 TWA (Do not eat, drink or chew tobacco or gum or apply cosmetics in reg ulated areas. Carcinogen - dust can cause lung and kidney disease. See 29 CFR 1910.1027)	

OSHA Vacated PELs: Cadmium: No OSHA Vacated PELs are listed for this chemical.

**Personal Protective Equipment** 

Eyes: Wear appropriate protective eyeglasses or chemical safety goggles as described by OSHA's eye and acceprotection regulations in 29 CFR 1910.133 or European Standard EN166.

kin: Wear appropriate protective gloves to prevent skin exposure.

Clothing: Wear appropriate protective clothing to prevent skin exposure.

**Respirators:** A respiratory protection program that meets OSHA's 29 CFR 1910.134 and ANSI Z88.2 requirements or European Standard EN 149 must be followed whenever workplace conditions warrant respirator use.

### Section 9 - Physical and Chemical Properties

Physical State: Granules Appearance: silver white

Odor: odorless **pH:** Not available.

Vapor Pressure: Not applicable. Vapor Density: Not available. Evaporation Rate:Not applicable.

Viscosity: Not applicable.

**Boiling Point:** 765 deg C @ 760 mmHg **Freezing/Melting Point:** 321 deg C

Decomposition Temperature: Not available.

Solubility: Insoluble.

Specific Gravity/Density:8.64 @ 25°C

Molecular Formula:Cd olecular Weight:112.40

# Section 10 - Stability and Reactivity

**Chemical Stability:** Oxidizes when exposed to air. Easily tarnishes in moist air. Powder or liquid is rophoric. Contact with acid liberates gas.

Inditions to Avoid: Ignition sources, dust generation, excess heat, prolonged exposure to air.
Incompatibilities with Other Materials: Strong oxidizing agents, acids, sulfur, zinc, selenium, tellurium.

Hazardous Decomposition Products: Toxic cadmium oxide fumes.

Hazardous Polymerization: Has not been reported.

# Section 11 - Toxicological Information

RTECS#:

CAS# 7440-43-9: EU9800000

LD50/LC50: CAS# 7440-43-9:

Inhalation, rat: LC50 = 25 mg/m3/30M;

Oral, mouse: LD50 = 890 mg/kg; Oral, rat: LD50 = 2330 mg/kg;

### Carcinogenicity:

CAS# 7440-43-9:

ACGIH: A2 - Suspected Human Carcinogen
 California: carcinogen, initial date 10/1/87

NTP: Known carcinogenIARC: Group 1 carcinogen

**Epidemiology:** Occupational exposure to cadmium has been implicated in a significant increase in prostate and respiratory tract cancer. There is evidence of a significant excess of respiratory cancer deaths among a cohort of cadmium production workers, and concluded that cadmium and its compounds are potential carcinogens.

**Teratogenicity:** Oral, rat: TDLo = 155 mg/kg (male 13 week(s) pre-mating and female 13 week(s) pre-mating - 3 week(s) after conception) Effects on Newborn - growth statistics (e.g.%, reduced weight gain) and Effects on Newborn - behavioral.; Oral, rat: TDLo = 23 mg/kg (female 1-22 day(s) after conception) Specific Developmental Abnormalities - blood and lymphatic systems (including spleen and marrow).; Oral, mouse: TDLo = 1700 mg/kg (female 8-12 day(s) after conception) Effects on Newborn - viability index (e.g., # alive at day 4 per # born alive) and Effects on Newborn - growth statis

**Reproductive Effects:** Oral, rat: TDLo = 21500 ug/kg (multigenerations) Fertility - pre-implantation mortality (e.g. reduction in number of implants per female; total number of implants per corpora lutea).; Intraperitoneal, rat: TDLo = 1124 ug/kg (male 1 day(s) pre-mating) Paternal Effects - spermatogenesis (incl. genetic material, sperm morphology, motility, and count).

**Mutagenicity:** Micronucleus Test: Mouse, Embryo = 6 umol/L.; Cytogenetic Analysis: Hamster, Ovary = 1 umol/L.

Neurotoxicity: No information found

Other Studies:

# Section 12 - Ecological Information

**Ecotoxicity:** Fish: Rainbow trout: TLm = 30 ppm; 24 Hr; Hard waterFish: Striped bass: LC50 = 0.001

ppm; 24-48 Hr; Static bioassayFish: Fathead Minnow: TL50 = 7.2 ppm; 96 Hr; UnspecifiedFish:

Bluegill/Sunfish: LCO = 0.08 ppm; 96 Hr; Static bioassay (Hard water) No data available.

Environmental: Cadmium can enter the air from natural sources.

Physical: No information available. \* ther: No information available.

# Section 13 - Disposal Considerations

Chemical waste generators must determine whether a discarded chemical is classified as a hazardous waste. US EPA guidelines for the classification determination are listed in 40 CFR Parts 261.3. Additionally, waste generators must consult state and local hazardous waste regulations to ensure complete and accurate classification.

RCRA P-Series: None listed. RCRA U-Series: None listed.

# Section 14 - Transport Information

	US DOT	Canada TDG
Shipping Name:	TOXIC SOLIDS, FLAMMABLE, ORGANIC, N.O.S.	Toxic Solid, Flammable, Organic, N.O.S. (CADMIUM METAL)
Hazard Class:	6.1	6.1
UN Number:	UN2930	UN2930
Packing Group:	I	I

# Section 15 - Regulatory Information

### **US FEDERAL**

### **TSCA**

CAS# 7440-43-9 is listed on the TSCA inventory.

### **Health & Safety Reporting List**

None of the chemicals are on the Health & Safety Reporting List.

#### **Chemical Test Rules**

None of the chemicals in this product are under a Chemical Test Rule.

### Section 12b

None of the chemicals are listed under TSCA Section 12b.

### **TSCA Significant New Use Rule**

None of the chemicals in this material have a SNUR under TSCA.

### **CERCLA Hazardous Substances and corresponding RQs**

CAS# 7440-43-9: 10 lb final RQ (no reporting of releases of this hazardous substance is required

### **SARA Section 302 Extremely Hazardous Substances**

None of the chemicals in this product have a TPQ.

### **SARA Codes**

CAS # 7440-43-9: immediate, delayed, fire.

### Section 313

This material contains Cadmium (CAS# 7440-43-9, 100%), which is subject to the reporting requirements of Section 313 of SARA Title III and 40 CFR Part 373.

### Clean Air Act:

CAS# 7440-43-9 (listed as Chromium compounds) is listed as a hazardous air pollutant (HAP).

This material does not contain any Class 1 Ozone depletors.

This material does not contain any Class 2 Ozone depletors.

### Clean Water Act:

None of the chemicals in this product are listed as Hazardous Substances under the CWA. CAS#

7440-43-9 is listed as a Priority Pollutant under the Clean Toxic Pollutant under the Clean Water Act.

Water Act. CAS# 7440-43-9 is listed as a

### OSHA:

None of the chemicals in this product are considered highly hazardous by OSHA.

#### TATE

CAS# 7440-43-9 can be found on the following state right to know lists: California, New Jersey, Pennsylvania, Minnesota, Massachusetts.

### California Prop 65

The following statement(s) is(are) made in order to comply with the California Safe Drinking Water Act:

WARNING: This product contains Cadmium, a chemical known to the state of California to cause cancer. WARNING: This product contains Cadmium, a chemical known to the state of California to cause male reproductive toxicity.

California No Significant Risk Level: CAS# 7440-43-9: 0.05 æg/day NSRL (inhalation)

# European/International Regulations European Labeling in Accordance with EC Directives Hazard Symbols:

T+F

### **Risk Phrases:**

R 11 Highly flammable.

R 25 Toxic if swallowed.

R 26 Very toxic by inhalation.

R 45 May cause cancer.

### **Safety Phrases:**

S 36/37/39 Wear suitable protective clothing, gloves and eye/face protection.

S 45 In case of accident or if you feel unwell, seek medical advice immediately (show the label where possible).

S 53 Avoid exposure - obtain special instructions before use.

### WGK (Water Danger/Protection)

CAS# 7440-43-9: No information available.

### Canada - DSL/NDSL

CAS# 7440-43-9 is listed on Canada's DSL List.

### Canada - WHMIS

This product has a WHMIS classification of D1A, B4.

This product has been classified in accordance with the hazard criteria of the Controlled Products Regulations and the MSDS contains all of the information required by those regulations.

### **Canadian Ingredient Disclosure List**

CAS# 7440-43-9 is listed on the Canadian Ingredient Disclosure List.

# Section 16 - Additional Information

**MSDS Creation Date:** 6/28/1999 **Revision #6 Date:** 6/06/2006

The information above is believed to be accurate and represents the best information currently available to us. However, we make no warranty of merchantability or any other warranty, express or implied, with respect to such information, and we assume no liability resulting from its use. Users should make their own investigations to determine the suitability of the information for their particular purposes. In no event shall Fisher be liable for any claims, losses, or damages of any third party or for lost profits or any special, indirect, incidental, consequential or exemplary damages, howsoever arising, even if Fisher has been advised of the possibility of such damages.

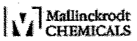
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MSDS Number: P5708 \* \* \* \* \* Effective Date: 02/21/07 \* \* \* \* \* Supercedes: 01/19/06

MSDS

### Material Safety Data Sheet

From: Mallinckrodt Baker, Inc. 222 Red School Lane Phillipsburg, NJ 08865





24 Hour Emergency Telephone: 906-459-2151 CHEMTREC: 1-890-424-9300

National Response in Canada CANUTEO: \$10.495-4666

Outside U.S. and Canada Chemirec: 703-527-3887

NOTE: CHEMIREO, CANUTEC and Matismal Response Camer emergency numbers to be used only in the event of chemical emergencies smooking a spill, leak, live, exposure or accident involving chemicals.

All non-emergency questions should be directed to Customer Service (1-500-552-2537) for assistance.

# POTASSIUM CYANIDE

# 1. Product Identification

Synonyms: Potassium cyanide, solid; hydrocyanic acid, potassium salt

CAS No.: 151-50-8

Molecular Weight: 65.12 Chemical Formula: KCN

Product Codes: J.T. Baker: 3080 Mallinckrodt: 6881

# 2. Composition/Information on Ingredients

Ingredient	CAS No	Percent	Hazardous
Potassium Cyanide	15150-8	96 - 100%	Yes

### 3. Hazards Identification

**Emergency Overview** 

POISON! DANGER! MAY BE FATAL IF SWALLOWED, INHALED OR ABSORBED THROUGH SKIN. CONTACT WITH ACIDS LIBERATES POISONOUS GAS. CAUSES

# BURNS TO SKIN, EYES, AND RESPIRATORY TRACT. AFFECTS BLOOD, CARDIOVASCULAR SYSTEM, CENTRAL NERVOUS SYSTEM AND THYROID.

SAF-T-DATA<sup>(tm)</sup> Ratings (Provided here for your convenience)

Health Rating: 4 - Extreme (Poison)

Flammability Rating: 0 - None Reactivity Rating: 2 - Moderate Contact Rating: 3 - Severe (Life)

Lab Protective Equip: GOGGLES & SHIELD; LAB COAT & APRON; VENT HOOD; PROPER

**GLOVES** 

Storage Color Code: Blue (Health)

### **Potential Health Effects**

In most cases, cyanide poisoning causes a deceptively healthy pink to red skin color. However, if a physical injury or lack of oxygen is involved, the skin color may be bluish. Reddening of the eyes and pupil dilation are symptoms of cyanide poisoning. Cyanosis (blue discoloration of the skin) tends to be associated with severe cyanide poisonings.

### Inhalation:

Corrosive to the respiratory tract. The substance inhibits cellular respiration and may cause blood, central nervous system, and thyroid changes. May cause headache, weakness, dizziness, labored breathing nausea and vomiting, which can be followed by weak and irregular heart beat, unconsciousness, convulsions, coma and death.

### Ingestion:

Highly Toxic! Corrosive to the gastro-intestinal tract with burning in the mouth and esophagus, and abdominal pain. Larger doses may produce sudden loss of consciousness and prompt death from respiratory arrest. Smaller but still lethal doses may prolong the illness for one or more hours. Bitter almonds odor may be noted on the breath or vomitus. Other symptoms may be similar to those noted for inhalation exposure.

### **Skin Contact:**

Corrosive. May cause severe pain and skin burns. Solutions are corrosive to the skin and eyes, and may cause deep ulcers which heal slowly. May be absorbed through the skin, with symptoms similar to those noted for inhalation.

### **Eye Contact:**

Corrosive. Symptoms may include redness, pain, blurred vision, and eye damage.

### **Chronic Exposure:**

Prolonged or repeated skin exposure may cause a "cyanide" rash and nasal sores.

### Aggravation of Pre-existing Conditions:

Workers using cyanides should have a preplacement and periodic medical exam. Those with history of central nervous system, thyroid, skin, heart or lung diseases may be more susceptible to the effects of this substance.

# 4. First Aid Measures

IN CASE OF CYANIDE POISONING, start first aid treatment immediately, then get medical attention. A cyanide antidote kit (amyl nitrite, sodium nitrite and sodium thiosulfate) should be available in any cyanide work area. Actions to be taken in case of cyanide poisoning should be

12/11/2007

planned and practiced before beginning work with cyanides. Oxygen and amyl nitrite can be given by a first responder before medical help arrives. Allow victim to inhale amyl nitrite for 15-30 seconds per minute until sodium nitrite and sodium thiosulfate can be administered intravenously (see Note to Physician). A new amyl nitrite ampule should be used every 3 minutes. If conscious but symptoms (nausea, difficult breathing, dizziness, etc.) are evident, give oxygen. If consciousness is impaired (non-responsiveness, slurred speech, confusion, drowsiness) or the patient is unconscious but breathing, give oxygen and amyl nitrite by means of a respirator. If not breathing, give oxygen and amyl nitrite immediately by means of a positive pressure respirator (artificial respiration).

### Inhalation:

If inhaled, remove to fresh air. Administer antidote kit and oxygen per pre-planned instructions if symptoms occur. Keep patient warm and at rest. Do not give mouth to mouth resuscitation.

### Ingestion:

If ingested, antidote kit and oxygen should be administered per above. If the patient is conscious, immediately give the patient activated charcoal slurry. Never give anything by mouth to an unconscious person. Do not induce vomiting as it could interfere with resuscitator use.

### **Skin Contact:**

Immediately flush skin with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Get medical attention immediately. Wash clothing before reuse. Thoroughly clean shoes before reuse. Administer antidote kit and oxygen per preplanned instructions if symptoms occur.

### **Eye Contact:**

Immediately flush eyes with plenty of water for at least 15 minutes, lifting lower and upper eyelids occasionally. Get medical attention immediately.

### Note to Physician:

If patient does not respond to amyl nitrite, inject intravenously with 10mL of a 3% solution of sodium nitrite at a rate of not more than 2.5 to 5 mL per minute. Once nitrite administration is complete, follow directly with 50 mL of a 25% solution of sodium thiosulfate at the same rate by the same route. Give victim oxygen and keep under observation. If exposure was severe, watch victim for 24-48 hours. If signs of cyanide poisoning persist or reappear, repeat nitrite and thiosulfate injections 1 hour later in 1/2 the original doses. Cyanocabalamin (B12), 1 mg intramuscularly, may speed recovery. Moderate cyanide exposures need be treated only by supportive measures such as bed rest and oxygen.

# 5. Fire Fighting Measures

#### Fire:

Not combustible, but upon decomposition or contact with acids, this material releases highly flammable and toxic hydrogen cyanide gas.

### **Explosion:**

Not considered an explosion hazard, but upon heating with chlorates or nitrites to 450C (842F) may cause an explosion. Violent explosion occurs if melted with nitrite salt. Sealed containers may rupture when heated.

### Fire Extinguishing Media:

Do Not use carbon dioxide. Carbon dioxide can react with this material in the presence of moisture to produce hydrogen cyanide. Water may be used on nearby fires not involving this material. Use alkali dry chemical. Water spray may be used to keep fire exposed containers cool. Reacts slowly with water to form hydrogen cyanide. Use any means suitable for extinguishing surrounding fire.

### **Special Information:**

In the event of a fire, wear full protective clothing and NIOSH-approved self-contained breathing

apparatus with full facepiece operated in the pressure demand or other positive pressure mode.

### ... Accidental Release Measures

Spills: Ventilate area of leak or spill. Allow only qualified personnel to handle spill. Clean-up personnel require protective clothing and respiratory protection from vapors. Collect material and place in a closed container for recovery or disposal. Do not flush to sewer! Decontaminate liquid or solid residues in spill area with sodium or calcium hypochlorite solution.

US Regulations (CERCLA) require reporting spills and releases to soil, water and air in excess of reportable quantities. The toll free number for the US Coast Guard National Response Center is (800) 424-8802.

# 7. Handling and Storage

Keep in a tightly closed container, stored in a cool, dry, ventilated area. Protect against physical damage. Separate from incompatibles. Workers must carefully follow good hygienic practices, including no eating, drinking, or smoking in workplace. Proper use and maintenance of protective equipment is essential. Workers using cyanide need preplacement and annual medical exams. Special training should be given to workers using cyanide. Containers of this material may be hazardous when empty since they retain product residues (dust, solids); observe all warnings and precautions listed for the product. Do not store near combustibles or flammables because subsequent fire fighting with water could lead to cyanide solution runoff. Do not store under sprinkler systems. All persons with the potential for cyanide poisoning should be trained to provide immediate First Aid using oxygen and amyl nitrite. A cyanide anitdote kit (amyl nitrite, sodium nitrite, and sodium thiosulfate) should be readily available in cyanide workplaces. The antidotes should be checked annually to ensure they are still within their shelf-lives. Identification of community hospital resources and emergency medical squads in order to equip and train them on handling cyanide emergencies is essential.

# 8. Exposure Controls/Personal Protection

### **Airborne Exposure Limits:**

-OSHA Permissible Exposure Limit (PEL):

5 mg/m3 skin (TWA) (as CN)

-ACGIH Threshold Limit Value (TLV):

5 mg/m3 (STEL) Ceiling, skin, as CN

### **Ventilation System:**

A system of local and/or general exhaust is recommended to keep employee exposures below the Airborne Exposure Limits. Local exhaust ventilation is generally preferred because it can control the emissions of the contaminant at its source, preventing dispersion of it into the general work area. Please refer to the ACGIH document, *Industrial Ventilation*, *A Manual of Recommended Practices*, most recent edition, for details.

Personal Respirators (NIOSH Approved):

If the exposure limit is exceeded and engineering controls are not feasible, wear a supplied air, full-

facepiece respirator, airlined hood, or full-facepiece self-contained breathing apparatus. Breathing air quality must meet the requirements of the OSHA respiratory protection standard (29CFR1910.134). This substance has poor warning properties.

### **Skin Protection:**

Wear impervious protective clothing, including boots, gloves, lab coat, apron or coveralls, as appropriate, to prevent skin contact.

### **Eye Protection:**

Use chemical safety goggles and/or full face shield where dusting or splashing of solutions is possible. Maintain eye wash fountain and quick-drench facilities in work area.

# 9. Physical and Chemical Properties

### Appearance:

White deliquescent granular solid.

Odor:

Bitter almonds.

Solubility:

Very soluble in water.

**Specific Gravity:** 

1.55 @ 20C/4C

pH:

11 (0.1 N aqueous solution)

% Volatiles by volume @ 21C (70F):

0

**Boiling Point:** 

1625C (2957F)

**Melting Point:** 

634C (1173F)

Vapor Density (Air=1):

No information found.

Vapor Pressure (mm Hg):

No information found.

Evaporation Rate (BuAc=1):

No information found.

# 10. Stability and Reactivity

### Stability:

Very stable when dry. Moisture will cause slow decomposition, releasing poisonous hydrogen cyanide gas.

### **Hazardous Decomposition Products:**

Emits toxic fumes of cyanide and oxides of nitrogen when heated to decomposition.

### Hazardous Polymerization:

Will not occur.

### Incompatibilities:

Strong acids and strong oxidizers. Reacts with acids to liberate toxic and flammable hydrogen cyanide gas. Water or weak alkaline solutions can produce dangerous amounts of hydrogen cyanide in confined areas. Can react with carbon dioxide in ordinary air to form hydrogen cyanide gas.

### Conditions to Avoid:

Heat, moisture, incompatibles.

# 11. Toxicological Information

Oral rat LD50: 6 mg/kg. Investigated as a mutagen, reproductive effector.

NTP	Carcinogen	
Known	Anticipated	IARC Category
No	No	None
	NTP Known	

# 12. Ecological Information

### **Environmental Fate:**

This material has an estimated bioconcentration factor (BCF) of less than 100. This material is not expected to significantly bioaccumulate.

### **Environmental Toxicity:**

This material is expected to be very toxic to aquatic life. This material is expected to be very toxic to terrestrial life.

# 13. Disposal Considerations

Cyanides must be oxidized to harmless waste before disposal. An alkaline solution (pH about 10) is treated with chlorine or commercial bleach in excess to decompose cyanide. When cyanide-free, it can be neutralized. Whatever cannot be saved for recovery or recycling should be handled as hazardous waste and sent to a RCRA approved waste facility. Processing, use or contamination of this product may change the waste management options. State and local disposal regulations may differ from federal disposal regulations. Dispose of container and unused contents in accordance with federal, state and local requirements.

# 14. Transport Information

Domestic (Land, D.O.T.)

Proper Shipping Name: RQ, POTASSIUM CYANIDE, SOLID

Hazard Class: 6.1 UN/NA: UN1680 Packing Group: I

Information reported for product/size: 12KG

International (Water, I.M.O.)

Proper Shipping Name: POTASSIUM CYANIDE, SOLID

Hazard Class: 6.1 UN/NA: UN1680 Packing Group: I

Information reported for product/size: 12KG

### International (Air, I.C.A.O.)

Proper Shipping Name: POTASSIUM CYANIDE, SOLID

Hazard Class: 6.1 UN/NA: UN1680 Packing Group: I

Information reported for product/size: 220LB

# 15. Regulatory Information

\Chemical Ingredient	·			TSCA	EC	Japan	Aust	ralia
Potassium Cyanide					Yes			
\Chemical	Inventory Stat	us - Part 2	2\			 nada		<del>-</del>
Ingredient		•			a DSL			
Potassium Cyanide				Yes	Yes	No	Ye	s
\Federal,	State & Intern	ational Re	gulati	ons-	Part 1	\		
	blace a moorn	dolonar no	-SARA	302-	 Li	SAR	A 313-	
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Ingredient	 (151-50-8)	ational Re	-SARA RQ  10	302- TPQ  100	Li  No Part 2 -RCRA-	SARU st Ch  Cya	A 313- emical anide SCA-	L Cat

Australian Hazchem Code: 4X

Poison Schedule: S7

WHMIS:

Reactivity: No

This MSDS has been prepared according to the hazard criteria of the Controlled Products Regulations (CPR) and the MSDS contains all of the information required by the CPR.

(Pure / Solid)

# 16. Other Information

NFPA Ratings: Health: 3 Flammability: 0 Reactivity: 0

Label Hazard Warning:

POISON! DANGER! MAY BE FATAL IF SWALLOWED, INHALED OR ABSORBED THROUGH SKIN. CONTACT WITH ACIDS LIBERATES POISONOUS GAS. CAUSES BURNS TO SKIN, EYES, AND RESPIRATORY TRACT. AFFECTS BLOOD, CARDIOVASCULAR SYSTEM, CENTRAL NERVOUS SYSTEM AND THYROID.

### **Label Precautions:**

Do not breathe dust.

Do not get in eyes, on skin, or on clothing.

Keep container closed.

Use only with adequate ventilation.

Wash thoroughly after handling.

### Label First Aid:

IN ALL CASES, GET MEDICAL ATTENTION IMMEDIATELY. KEEP A CYANIDE ANTIDOTE KIT (amyl nitrite, sodium nitrite and sodium thiosulfate) in area of product use or storage. First-aiders must take precautions to avoid contact with cyanide substance. If ingested, administer antidote kit and oxygen per pre-planned instructions. If the patient is conscious, immediately give the patient activated charcoal slurry. Never give anything by mouth to an unconscious person. Do not induce vomiting as it could interfere with resuscitator use. If inhaled, remove to fresh air. Administer antidote kit and oxygen per pre-planned instructions if symptoms occur. Keep patient warm and at rest. Do not give mouth to mouth resuscitation. In case of contact, immediately flush eyes or skin with plenty of water for at least 15 minutes while removing contaminated clothing and shoes. Wash clothing before reuse. Administer antidote kit and oxygen per preplanned instructions if symptoms occur.

### **Product Use:**

Laboratory Reagent.

**Revision Information:** 

MSDS Section(s) changed since last revision of document include: 14.

Prepared by: Environmental Health & Safety Phone Number: (314) 654-1600 (U.S.A.)

# **International Chemical Safety Cards**

# **CALCIUM CYANIDE**

**ICSC: 0407** 

CALCIUM CYANIDE Calcid

Calcyanide
Calcyan
C<sub>2</sub>CaN<sub>2</sub>/Ca(CN)<sub>2</sub>

Molecular mass: 92.1

CAS # 592-01-8 RTECS # EW0700000 ICSC # 0407 UN # 1575

EC # 020-002-00-5

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ SYMPTO		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Not combustible but for flammable gas on conwater or damp air.		NO open flames, NO sparks, a NO smoking. NO contact with water, carbon dioxide, acids. I contact with hot surfaces.	h	Powder, dry sand. NO hydrous agents. NO water. NO carbon dioxide.
EXPLOSION					
EXPOSURE	-	•	STRICT HYGIENE!		IN ALL CASES CONSULT A DOCTOR!
• INHALATION	Burning sensation. Co Dizziness. Headache. colouration of the skir breathing. Nausea. Sh breath. Unconsciousne Vomiting. Convulsion Death.	Red  1. Laboured  ortness of  ess.	Local exhaust or breathing protection.		Fresh air, rest. Half-upright position. Artificial respiration if indicated. Refer for medical attention. See Notes.
• SKIN	MAY BE ABSORBE burns. Pain. Itching. P (further see Inhalation	apules	Protective gloves. Protective clothing.		Remove contaminated clothes. Rinse and then wash skin with water and soap. Refer for medical attention. See Notes.
• EYES	Pain. Blurred vision. Possible permanent loss of vision. Severe deep burns.		Face shield or eye protection in combination with breathing protection if powder. Contact lenses should not be worn when working with this chemical.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
Confusion. Burning sense mouth. Numbness or tig throat. Salivation. Converted by paralysis (fill Inhalation).		ightness in vulsions	work.		Rinse mouth. Induce vomiting (ONLY IN CONSCIOUS PERSONS!). Refer for medical attention. See Notes.
SPILLAGE	E DISPOSAL		STORAGE	PAC	CKAGING & LABELLING
Evacuate danger are	a! Consult an expert!	Fireproof. Se	deparated from strong oxidants, Airtight. Do not transport with food a		

acids, food and feedstuffs. Cool. Dry.

Keep in a well-ventilated room.

feedstuffs.

T symbol R: 28-32

S: 7/8-23-36/37-45

UN Hazard Class: 6.1

To NOT wash away into sewer. Sweep

Carefully collect remainder, then remove

to safe place. Prevent contact with water

spilled substance into containers.

or moist substances (extra personal

protection: complete protective clothing including self-contained breathing apparatus).	UN Packing Group: I
S	EE IMPORTANT INFORMATION ON BACK
ICSC: 0407	repared in the context of cooperation between the International Programme on Chemical Safety & the Commission of e European Communities @ IPCS CEC 1993

# **International Chemical Safety Cards**

### CALCIUM CYANIDE

ICSC: 0407

CALCIUM	CYANIDE	
	PHYSICAL STATE; APPEARANCE: WHITE CRYSTALS OR POWDER. CRYSTALLINE POWDER, WITH CHARACTERISTIC ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation, through the skin or or by ingestion.
I M P	PHYSICAL DANGERS:	INHALATION RISK:  A harmful contamination of the air will be reached rather slowly on evaporation of this substance at 20° C when dispersed.
O R T A N T D A	CHEMICAL DANGERS: The substance decomposes on heating above 350°C producing toxic fumes (hydrogen cyanide, nitrous oxides). Reacts violently with water, moist air, carbon dioxide, acids, acidic salts producing highly toxic and flammable hydrogen cyanide. Reacts violently when heated with nitrites, nitrates, chlorates and perchlorates.	EFFECTS OF SHORT-TERM EXPOSURE: The substance is corrosive to the eyes, the skin and the respiratory tract. Inhalation of the substance may cause lung oedema (see Notes). The substance may cause effects on the nervous system, blood, heart and respiratory tract. Exposure at high level may result in death.
T A	OCCUPATIONAL EXPOSURE LIMITS (OELs): TLV (as CN): ppm; 5 mg/m <sup>3</sup> (skin) (ACGIH 1991-1992).	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE: Repeated or prolonged contact with skin may cause dermatitis. May cause reproductive toxicity in humans.
PHYSICAL PROPERTIES	Decomposes below melting point at >350°C°C Relative density (water = 1): 1.85	Solubility in water: reaction
ENVIRONMENTAL DATA	This substance may be hazardous to the environment organisms, soil.	; special attention should be given to water, aquatic
	NOTES	
become manifest until	odour warning when the exposure limit value is excee	hysical effort. Rest and medical observation are this substance; the appropriate means with instructions
	ADDITIONAL INFORMA	TION
ICSC: 0407	© IPCS, CEC, 1993	CALCIUM CYANIDE
1 Hu	leither the CEC or the IPCS nor any person acting on se which might be made of this information. This card eview Committee and may not reflect in all cases all	contains the collective views of the IPCS Peer

LEGAL NOTICE: legislation on the subject. The user should verify compliance of the cards with the relevant legislation in the country of use.

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MSDS Number: L2347 \* \* \* \* \* Effective Date: 07/05/07 \* \* \* \* \* Supercedes: 05/07/07



# MSDS Material Safety Data Sheet

From: Mallinckrodt Baker, Inc. 222 Red School Lane Phillipsburg, NJ 08865





24 Hour Emergency Telephone: 908 459-2161 CHEMITIEO: 1-600-424-6300

National Response in Canada CANUTEO: \$10-406-6666

Outside U.S. and Canada Chemirec: 703-527-3887

NOTE: CHEMIFIED, CANUTED and National Response Center emergency numbers to be used only in the event of chemical emergencies involving a spill, look, liee, exposure or accident involving obernicals.

All non-emergency questions should be directed to Customer Service (1-800-582-2537) for assistance.

# LEAD METAL

## 1. Product Identification

Synonyms: Granular lead, pigment metal; C.I. 77575

CAS No.: 7439-92-1

Molecular Weight: 207.19 Chemical Formula: Pb

**Product Codes:** 

J.T. Baker: 2256, 2266 Mallinckrodt: 5668

# 2. Composition/Information on Ingredients

Ingredient	CAS No	Percent	Hazardous
·			
Lead	743992-1	95 - 100%	Yes

# 3. Hazards Identification

**Emergency Overview** 

POISON! DANGER! MAY BE FATAL IF SWALLOWED OR INHALED. CAUSES IRRITATION TO SKIN, EYES AND RESPIRATORY TRACT. NEUROTOXIN. AFFECTS THE GUM TISSUE, CENTRAL NERVOUS SYSTEM, KIDNEYS, BLOOD AND REPRODUCTIVE SYSTEM. POSSIBLE CANCER HAZARD. MAY CAUSE CANCER BASED ON ANIMAL DATA. Risk of cancer depends on duration and level of exposure.

# SAF-T-DATA<sup>(tm)</sup> Ratings (Provided here for your convenience)

Health Rating: 3 - Severe (Cancer Causing) Flammability Rating: 3 - Severe (Flammable)

Reactivity Rating: 1 - Slight

Contact Rating: 2 - Moderate (Life)

Lab Protective Equip: GOGGLES & SHIELD; LAB COAT & APRON; VENT HOOD; PROPER

**GLOVES** 

Storage Color Code: Red (Flammable)

### **Potential Health Effects**

### Inhalation:

Lead can be absorbed through the respiratory system. Local irritation of bronchia and lungs can occur and, in cases of acute exposure, symptoms such as metallic taste, chest and abdominal pain, and increased lead blood levels may follow. See also Ingestion.

### Ingestion:

POISON! The symptoms of lead poisoning include abdominal pain and spasms, nausea, vomiting, headache. Acute poisoning can lead to muscle weakness, "lead line" on the gums, metallic taste, definite loss of appetite, insomnia, dizziness, high lead levels in blood and urine with shock, coma and death in extreme cases.

### Skin Contact:

Lead and lead compounds may be absorbed through the skin on prolonged exposure; the symptoms of lead poisoning described for ingestion exposure may occur. Contact over short periods may cause local irritation, redness and pain.

### **Eve Contact:**

Absorption can occur through eye tissues but the more common hazards are local irritation or abrasion.

### **Chronic Exposure:**

Lead is a cumulative poison and exposure even to small amounts can raise the body's content to toxic levels. The symptoms of chronic exposure are like those of ingestion poisoning; restlessness, irritability, visual disturbances, hypertension and gray facial color may also be noted.

### Aggravation of Pre-existing Conditions:

Persons with pre-existing kidney, nerve or circulatory disorders or with skin or eye problems may be more susceptible to the effects of this substance.

# 4. First Aid Measures

### Inhalation:

Remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. Get medical attention.

### Ingestion:

Induce vomiting immediately as directed by medical personnel. Never give anything by mouth to

LEAD METAL rage 2 01 9

an unconscious person. Get medical attention.

### Skin Contact:

Immediately flush skin with plenty of soap and water for at least 15 minutes. Remove contaminated clothing and shoes. Get medical attention. Wash clothing before reuse. Thoroughly clean shoes before reuse.

### **Eye Contact:**

Immediately flush eyes with plenty of water for at least 15 minutes, lifting lower and upper eyelids occasionally. Get medical attention immediately.

# 5. Fire Fighting Measures

### Fire:

Not considered to be a fire hazard. Powder/dust is flammable when heated or exposed to flame.

### **Explosion:**

Not considered to be an explosion hazard.

### Fire Extinguishing Media:

Use any means suitable for extinguishing surrounding fire. Do not allow water runoff to enter sewers or waterways.

### **Special Information:**

In the event of a fire, wear full protective clothing and NIOSH-approved self-contained breathing apparatus with full facepiece operated in the pressure demand or other positive pressure mode. Can produce toxic lead fumes at elevated temperatures and also react with oxidizing materials.

# **6.** Accidental Release Measures

Ventilate area of leak or spill. Wear appropriate personal protective equipment as specified in Section 8. Spills: Sweep up and containerize for reclamation or disposal. Vacuuming or wet sweeping may be used to avoid dust dispersal. US Regulations (CERCLA) require reporting spills and releases to soil, water and air in excess of reportable quantities. The toll free number for the US Coast Guard National Response Center is (800) 424-8802.

# 7. Handling and Storage

Keep in a tightly closed container, stored in a cool, dry, ventilated area. Protect against physical damage. Isolate from incompatible substances. Areas in which exposure to lead metal or lead compounds may occur should be identified by signs or appropriate means, and access to the area should be limited to authorized persons. Containers of this material may be hazardous when empty since they retain product residues (dust, solids); observe all warnings and precautions listed for the product.

# 8. Exposure Controls/Personal Protection

LEAD METAL Fage 4 01 8

### Airborne Exposure Limits:

For lead, metal and inorganic dusts and fumes, as Pb:

-OSHA Permissible Exposure Limit (PEL): 0.05 mg/m3 (TWA)

For lead, elemental and inorganic compounds, as Pb:

-ACGIH Threshold Limit Value (TLV): 0.05 mg/m3 (TWA), A3 animal carcinogen

ACGIH Biological Exposure Indices (BEI): 30 ug/100ml, notation B (see actual Indices for more information).

For lead, inorganic:

-NIOSH Recommended Exposure Limit (REL): 0.1 mg/m3 (TWA)

### **Ventilation System:**

A system of local and/or general exhaust is recommended to keep employee exposures below the Airborne Exposure Limits. Local exhaust ventilation is generally preferred because it can control the emissions of the contaminant at its source, preventing dispersion of it into the general work area. Please refer to the ACGIH document, *Industrial Ventilation*, *A Manual of Recommended Practices*, most recent edition, for details.

### Personal Respirators (NIOSH Approved):

If the exposure limit is exceeded and engineering controls are not feasible, a half-face high efficiency particulate respirator (NIOSH type N100 filter) may be worn for up to ten times the exposure limit or the maximum use concentration specified by the appropriate regulatory agency or respirator supplier, whichever is lowest. A full-face piece high efficiency particulate respirator (NIOSH type N100 filter) may be worn up to 50 times the exposure limit, or the maximum use concentration specified by the appropriate regulatory agency or respirator supplier, whichever is lowest. If oil particles (e.g. lubricants, cutting fluids, glycerine, etc.) are present, use a NIOSH type R or P filter. For emergencies or instances where the exposure levels are not known, use a full-facepiece positive-pressure, air-supplied respirator. WARNING: Air-purifying respirators do not protect workers in oxygen-deficient atmospheres.

### **Skin Protection:**

Wear impervious protective clothing, including boots, gloves, lab coat, apron or coveralls, as appropriate, to prevent skin contact.

### **Eye Protection:**

Use chemical safety goggles and/or full face shield where dusting or splashing of solutions is possible. Maintain eye wash fountain and quick-drench facilities in work area.

### **Other Control Measures:**

Eating, drinking, and smoking should not be permitted in areas where solids or liquids containing lead compounds are handled, processed, or stored. See OSHA substance-specific standard for more information on personal protective equipment, engineering and work practice controls, medical surveillance, record keeping, and reporting requirements. (29 CFR 1910.1025).

# 9. Physical and Chemical Properties

### Appearance:

Small, white to blue-gray metallic shot or granules.

Odor:

Odorless.

Solubility:

Insoluble in water.

Density:

11.34

pH:

No information found.

% Volatiles by volume @ 21C (70F):

0

**Boiling Point:** 

1740C (3164F)

Melting Point:

327.5C (622F)

Vapor Density (Air=1):

No information found.

Vapor Pressure (mm Hg):

1.77 @ 1000C (1832F)

Evaporation Rate (BuAc=1):

No information found.

# 10. Stability and Reactivity

### Stability:

Stable under ordinary conditions of use and storage.

### **Hazardous Decomposition Products:**

Does not decompose but toxic lead or lead oxide fumes may form at elevated temperatures.

### Hazardous Polymerization:

Will not occur.

### Incompatibilities:

Ammonium nitrate, chlorine trifluoride, hydrogen peroxide, sodium azide, zirconium, disodium acetylide, sodium acetylide and oxidants.

### Conditions to Avoid:

Heat, flames, ignition sources and incompatibles.

# 11. Toxicological Information

### **Toxicological Data:**

Investigated as a tumorigen, mutagen, reproductive effector.

### Reproductive Toxicity:

Lead and other smelter emissions are human reproductive hazards. (Chemical Council on Environmental Quality; Chemical Hazards to Human Reproduction, 1981).

### Carcinogenicity:

EPA / IRIS classification: Group B2 - Probable human carcinogen, sufficient animal evidence.

/Cancer Lists/		arcinogen	
Ingredient	Known	Anticipated	IARC Category
Lead (7439-92-1)	No	No	2B

# 12. Ecological Information

### **Environmental Fate:**

When released into the soil, this material is not expected to leach into groundwater. This material

may bioaccumulate to some extent.

Environmental Toxicity:

No information found.

# 13. Disposal Considerations

Whatever cannot be saved for recovery or recycling should be managed in an appropriate and approved waste facility. Although not a listed RCRA hazardous waste, this material may exhibit one or more characteristics of a hazardous waste and require appropriate analysis to determine specific disposal requirements. Processing, use or contamination of this product may change the waste management options. State and local disposal regulations may differ from federal disposal regulations. Dispose of container and unused contents in accordance with federal, state and local requirements.

# 14. Transport Information

Not regulated.

# 15. Regulatory Information

```
-----\Chemical Inventory Status - Part 1\-----
                                    TSCA EC Japan Australia
 Ingredient
 ________
                                     Yes Yes Yes
 Lead (7439-92-1)
 -----\Chemical Inventory Status - Part 2\-----
                                      --Canada--
                                    Korea DSL NDSL Phil.
 Ingredient
 Yes Yes No
 Lead (7439-92-1)
 -----\Federal, State & International Regulations - Part 1\------
                              -SARA 302- -----SARA 313-----
                                RQ TPQ List Chemical Catg.
 Ingredient
               _______
 Lead (7439-92-1)
 -----\Federal, State & International Regulations - Part 2\------
                                        -RCRA- -TSCA-
                                CERCLA
                                        261.33
                                              . 8 (d)
 Ingredient
 Lead (7439-92-1)
Chemical Weapons Convention: No TSCA 12(b): No CDTA: No
GARA 311/312: Acute: Yes Chronic: Yes Fire: No Pressure: No
 activity: No (Pure / Solid)
```

#### **WARNING:**

THIS PRODUCT CONTAINS CHEMICALS KNOWN TO THE STATE OF CALIFORNIA TO

LEAD METAL Page / of 8

### CAUSE CANCER AND BIRTH DEFECTS OR OTHER REPRODUCTIVE HARM.

Australian Hazchem Code: None allocated.

Poison Schedule: S6

WHMIS:

This MSDS has been prepared according to the hazard criteria of the Controlled Products Regulations (CPR) and the MSDS contains all of the information required by the CPR.

### 16. Other Information

NFPA Ratings: Health: 3 Flammability: 1 Reactivity: 0

Label Hazard Warning:

POISON! DANGER! MAY BE FATAL IF SWALLOWED OR INHALED. CAUSES IRRITATION TO SKIN, EYES AND RESPIRATORY TRACT. NEUROTOXIN. AFFECTS THE GUM TISSUE, CENTRAL NERVOUS SYSTEM, KIDNEYS, BLOOD AND REPRODUCTIVE SYSTEM. POSSIBLE CANCER HAZARD. MAY CAUSE CANCER BASED ON ANIMAL DATA. Risk of cancer depends on duration and level of exposure.

### **Label Precautions:**

Do not get in eyes, on skin, or on clothing.

Do not breathe dust.

Keep container closed.

Use only with adequate ventilation.

Wash thoroughly after handling.

### Label First Aid:

If swallowed, induce vomiting immediately as directed by medical personnel. Never give anything by mouth to an unconscious person. If inhaled, remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. In case of contact, immediately flush eyes or skin with plenty of water for at least 15 minutes. Remove contaminated clothing and shoes. Wash clothing before reuse. In all cases, get medical attention.

### **Product Use:**

Laboratory Reagent.

### **Revision Information:**

MSDS Section(s) changed since last revision of document include: 3.

Disclaimer:

\*

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Prepared by: Environmental Health & Safety

Phone Number: (314) 654-1600 (U.S.A.)

# APPENDIX D

# PERSONAL PROTECTIVE EQUIPMENT REQUIREMENTS

CUY-CLEVELAND INNERBELT COMMERCIAL ROAD ALIGNMENT (PID 77510) CLEVELAND, CUYAHOGA COUNTY, OHIO

### PERSONAL PROTECTION EQUIPMENT

The personal protective equipment requirements for HzW personnel for on-site activities were selected and based on previous land use and previous investigations. Therefore, unless more information is obtained regarding employee exposure at this specific site, such as personal or area air monitoring data or data from the PID indicates that a higher level of PPE is required, then the following PPE, at a minimum is required to be worn:

Respirator:

As necessary

Protective Clothing:

Protective Coveralls/As necessary

Gloves:

Nitrile

Boots:

Safety-toe boots or shoes

Hard Hat:

As necessary

Safety Glasses:

As necessary

Ear Protection:

Disposable, 30-32 dB noise reduction

\*\*ANY PPE REQUIREMENTS OF THE FACILITY SHALL SUPERSEDE THOSE PRESENTED IN THIS HASP\*\*.

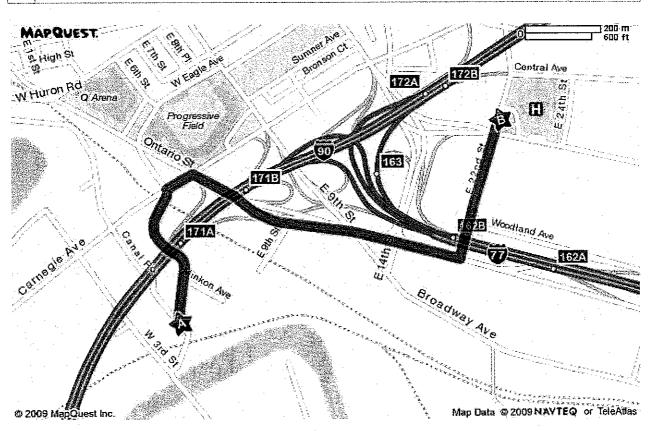
# APPENDIX E

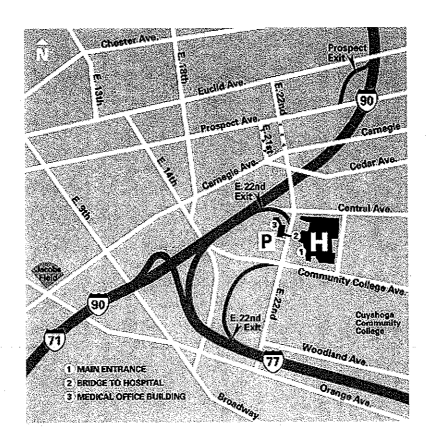
# LOCATION, MAP AND DIRECTIONS TO NEAREST HOSPITAL

# St. Vincent Charity Hospital 2351 East 22<sup>nd</sup> Street Cleveland, Ohio 44115 Phone: (216) 816-6200

### Directions:

1.	Go NORTH on COMMERCIAL ROAD towards CARNEGIE AVENUE	0.3 mile
2.	Turn RIGHT onto CARNEGIE AVENUE	
3.	Stay in <b>RIGHT LANE</b> , continue to intersection with <b>BROADWAY AVENUE</b> (US 422/SR 14/SR 43/SR 87)	0.1 mile
4.	Turn RIGHT onto BROADWAY AVENUE and stay in LEFT LANE	
5.	Continue on BROADWAY AVENUE to EAST 22ND STREET	0.5 mile
6.	Turn LEFT onto EAST 22 <sup>ND</sup> STREET	
7.	Continue to ST. VINCENT CHARITY HOSPITAL on RIGHT	0.3 mile





# APPENDIX F

# EMERGENCY INFORMATION AND PHONE NUMBERS

# **EMERGENCY INFORMATION AND PHONE NUMBERS**

OHIO EPA				
<ul> <li>NORTHEAST DISTRICT OFFICE (TWINSBURG)</li></ul>				
UNITED STATES EPA				
<ul> <li>CERCLA/RCRA HOTLINE 1-800-424-9346</li> <li>NATIONAL RESPONSE CENTER 1-800-424-8802</li> <li>EPCRA HOTLINE 1-800-535-0202</li> </ul>				
HAZMAT				
• EMERGENCY 911				
CITY OF CLEVELAND DEPT. OF PUBLIC HEALTH 1-216-664-2324				
<u>CLEVELAND FIRE DEPARTMENT</u>				
CLEVELAND POLICE DEPARTMENT 911 (non-emergency) 1-216-623-5000				
CITY OF CLEVELAND WATER DEPARTMENT 1-216-664-3060 (water main breaks/leaks)				
OHIO UTILITY PROTECTION SERVICE (OUPS)				

# APPENDIX L SOIL BORING LOGS

**Project No.:** H09004-14

Project: Phase II ESA

HzW Representative: JAD/JAH

Location: CUY-Innerbelt Commercial Road (PID 77510)



**Drill Date:** 12/17/2009

**Drilled By:** HzW Environmental

Drill Method: Hydrualic Direct Push

		Description		
Depth (feet)	Symbol		PID (ppm)	Remarks
0-		Ground Surface	4	
1-		Brown sandy CLAY w/ gravel, slag and brick fragments, dry, firm		
2-				
3-				
4-		Brown CLAY w/ sand, black slag and brick fragments, dry, firm	1	
5-				
6-	7.7.7.7.7	Black and dark brown sandy CLAY w/ slag and gravel, dry, firm, slight sulfur odor	1	
7-				
8-		Black and dark brown sandy SLAG w/ clay, gravel and wood fragments, strong sulfur odor		
10-				The 8-10 foot interval submitted for laboratory analysis
11-		Dark gray and brown sandy CLAY w/ gray slag, dry, firm, slight sulfur odor		The 10-12 foot interval submitted
12-				for laboratory analysis
13-				
14-	777	Dark gray and black SAND w/ slag, trace clay, dry, dense	-	
15-				
16-		Dark brown and gray sandy SLAG w/ trace clay, dry, dense, slight sulfur odor from 16-18'	-	
17-	• •			The 16-18 foot interval submitted for laboratory analysis
18-				
19-	• .			
20-		End of Bore		
21 –				

**Project No.:** H09004-14

Project: Phase II ESA

HzW Representative: JAD/JAH

Location: CUY-Innerbelt Commercial Road (PID 77510)



**Drill Date:** 12/17/2009

**Drilled By:** HzW Environmental

Drill Method: Hydrualic Direct Push

		Description		
Depth (feet)	Symbol		PID (ppm)	Remarks
0-		Ground Surface  Dak brown and black coarse sandy SLAG w/ trace clay, dry, loose		
1-		Dak brown and black coarse sailty SEAG w/ trace clay, try, roose		
	• • •			
2-		Dark brown and black sandy SLAG, dry, dense		
3-				The 2-4 foot interval submitted for laboratory analysis
4-		Black and dark gray sandy SLAG w/ additional large slag pieces, dry, dense		
5-				The 4-6 foot interval submitted for laboratory analysis
6-		End of Bore		
7-				

**Project No.:** H09004-14

Project: Phase II ESA

HzW Representative: JAD/JAH

Location: CUY-Innerbelt Commercial Road (PID 77510)



**Drill Date:** 12/17/2009

**Drilled By:** HzW Environmental

Drill Method: Hydrualic Direct Push

		Description		
Depth (feet)	Symbol		PID (ppm)	Remarks
0-	t mert	Ground Surface White and gray SLAG		
-		write and gray SLAG		
-		Brown and black sandy SLAG w/ brick fragments		Strong, unidentifable odor in 0-2' interval; the 0-2' interval submitted for laboratory analysis
2-		Brown and black coarse sandy SLAG w/ trace clay, brick and gravel, damp, dense		
3-				
4-				
5-		Brown medium SAND, damp, dense		
6-		End of Bore		
			1	

**Project No.:** H09004-14

Project: Phase II ESA

HzW Representative: JAD/JAH

Location: CUY-Innerbelt Commercial Road (PID 77510)



**Drill Date:** 12/17/2009

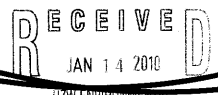
**Drilled By:** HzW Environmental

Drill Method: Hydrualic Direct Push

		Description		
Depth (feet)	Symbol		PID (ppm)	Remarks
0-		Ground Surface Brown coasre SAND w/ gray crushed slag, dry, loose		
_		, , , , , , , , , , , , , , , , , , ,		
1-				The 0-2 foot interval submitted for laboratory analysis
2-				
_				
3-				The 2-4 foot interval submitted for laboratory analysis
4-		Brown coarse SAND w/ gravel		
5-		Gray silty SAND, damp, soft		
6-	000 E 1110	End of Bore		
7-				

# APPNEDIX M LABORATORY ANALYTICAL REPORTS





HZW ENVIRUMENTAL CONSULTANTS (LC)

# Affidavit by Certified Lab Pursuant to OAC 3745-300-13(M)

TestAmerica Laboratories, Inc.

State of Ohio

SS:

County of Stark

- I, Opal Davis-Johnson, being first duly sworn according to law, state that, to the best of my knowledge, information and belief:
- 1. I am an adult over the age of eighteen (18) years old and competent to testify herein.
- 2. I am employed by TestAmerica Laboratories, Inc. as Laboratory Director and authorized to submit this affidavit on behalf of TestAmerica North Canton.
- 3. The purpose of this submission is to support a request for a no further action letter or other aspects of a voluntary action, under Ohio's Voluntary Action Program (VAP) as set forth in Ohio Revised Code Chapter 3746 and Ohio Administrative Code (OAC) Chapter 3745-300.
- 4. TestAmerica North Canton performed analyses on behalf of **HZW Environmental Consultants** for a voluntary action at property known as **Commercial Road, located Northeast of Commercial & Canal Roads in Cleveland**
- 5. This affidavit applies to and is submitted with the following information, data, documents or reports for the property:

<u>Laboratory Report Number</u> **A9L180492** 

Report Date
January 11, 2010

- 6. TestAmerica North Canton was a VAP certified laboratory pursuant to OAC 3745-300-04 when it performed analyses referenced herein.
- 7. The analyses under this affidavit consist of certified data, as described by OAC 3745-300-04(B) with the exception of the analytes, parameter groups, or methods listed below:

Method:

Analyte/Compound:

9030B/9034

Sulfide

day of

9045

Corrosivity

- 8. TestAmerica North Canton performed the analyses within its current VAP certification. The laboratory was certified for each analyte, parameter group and method used at the time that it performed the analyses. The analyses were performed consistent with the laboratory's standard operating procedures and quality assurance program plan as approved under OAC 3745-300-04.
- 9. The information, data, documents, and reports identified under this affidavit are true, accurate, and complete.

Further affiant sayeth naught.

Signature of Affiant

Sworn to before me this 13

2010

JEFFREY C. SMITH

Notary Public, State of Ohio My Commission Expires Feb. 12, 2012 Jeffrey'Cl. Smith Notary Public

> Page 1 of 1 Revised 06/09



HZW ENVIRONMENTA CONSULTANTS, LLC

TestAmerica Laboratories, Inc.

# **ANALYTICAL REPORT**

PROJECT NO. H09004-14

COMMERCIAL RD.

Lot #: A9L180492

Doug Wetzel

HZW Environmental Consultants 6105 Heisley Rd Mentor, OH 44060

TESTAMERICA LABORATORIES, INC.

Nathan Pietras

Project Manager

nathan.pietras@testamericainc.com

Approved for release Nathan Pletras Project Manager 1/11/2010 3:20 PM

January 11, 2010



## CASE NARRATIVE

A9L180492

The following report contains the analytical results for eight solid samples submitted to TestAmerica North Canton by HZW Environmental Consultants from the Commercial Rd. Site, project number H09004-14. The samples were received December 18, 2009, according to documented sample acceptance procedures.

TestAmerica utilizes USEPA approved methods in all analytical work. The samples presented in this report were analyzed for the parameter(s) listed on the analytical methods summary page in accordance with the method(s) indicated. Preliminary results were provided to Doug Wetzel on January 04, 2010, and January 07, 2010. A summary of QC data for these analyses is included at the back of the report.

TestAmerica North Canton attests to the validity of the laboratory data generated by TestAmerica facilities reported herein. All analyses performed by TestAmerica facilities were done using established laboratory SOPs that incorporate QA/QC procedures described in the applicable methods. TestAmerica's operations groups have reviewed the data for compliance with the laboratory QA/QC plan, and data have been found to be compliant with laboratory protocols unless otherwise noted below.

All solid sample results are reported on an "as received" basis unless otherwise indicated by a dry weight adjustment footnote at the bottom of the analytical report page. The list of parameters which are never reported on a dry weight basis is included on the Sample Summary.

The test results in this report meet all NELAP requirements for parameters for which accreditation is required or available. Any exceptions to NELAP requirements are noted in this report. Pursuant to NELAP, this report may not be reproduced, except in full, without the written approval of the laboratory.

All parameters were evaluated to the reporting limit with the exception of Total Residue as Percent Solids, Extractable Petroleum Hydrocarbons and Semivolatile Organic Compounds by GC/MS which were evaluated to the method detection limit and include appropriate qualifiers.

Please refer to the Quality Control Elements Narrative following this case narrative for additional quality control information.

If you have any questions, please call the Project Manager, Nathan Pietras, at 330-497-9396.

This report is sequentially paginated. The final page of the report is labeled as "END OF REPORT."

# CASE NARRATIVE (continued)

# SUPPLEMENTAL QC INFORMATION

#### SAMPLE RECEIVING

The temperature of the cooler upon sample receipt was 0.0°C.

#### **GC/MS VOLATILES**

The matrix spike/matrix spike duplicate(s) for batch(es) 9356203 had RPD's and recoveries outside acceptance limits. However, since the associated method blank(s) and laboratory control sample(s) were in control, no corrective action was necessary.

The internal standard areas were outside acceptance limits for sample(s) HB-071(10-12'), HB-071(16-18') and HB-072(2-4') due to matrix effects. (Refer to IS report following this case narrative for additional detail.)

Sample(s) HB-071(8-10') had elevated reporting limits due to TICs.

Sample(s) HB-071(10-12'), HB-071(16-18') and HB-072(2-4') were reanalyzed at a dilution due to internal standard recoveries outside of acceptance limits, per Ohio VAP requirements. Only compounds associated with internal standards that met criteria are reported from each analysis.

There were no client requested Matrix Spike/Matrix Spike Duplicate (MS/MSD) samples in batch 9356563. Therefore, the laboratory has included a Laboratory Control Sample Duplicate (LCSD) in the QC batch. The LCSD recoveries, together with the LCS recoveries, are used to determine the reproducibility (precision) of the analytical system.

#### **GC/MS SEMIVOLATILES**

Sample HB-075(0-2') had elevated reporting limits due to matrix interferences.

The batch QC for batch 9353017 was unable to be reported due to re-preparation of the parent sample and the original sample was not reported.

# **CASE NARRATIVE (continued)**

#### PURGEABLE PETROLEUM HYDROCARBONS-8015

The matrix spike/matrix spike duplicate(s) for batch(es) 9363092 had recoveries outside acceptance limits. However, since the associated method blank(s) and laboratory control sample(s) were in control, no corrective action was necessary.

There were no client requested Matrix Spike/Matrix Spike Duplicate (MS/MSD) samples in batch 9354104. Therefore, the laboratory has included a Laboratory Control Sample Duplicate (LCSD) in the QC batch. The LCSD recoveries, together with the LCS recoveries, are used to determine the reproducibility (precision) of the analytical system.

#### EXTRACTABLE PETROLEUM HYDROCARBONS-8015

The sample(s) that contain results between the MDL and the RL were flagged with "J". There is a possibility of false positive or mis-identification at these quantitation levels. In analytical methods requiring confirmation of the analyte reported, confirmation was performed only down to the standard reporting limit (SRL). The acceptance criteria for OC samples may not be met at these quantitation levels.

The generic batch MS/MSDs for batch(es) 9353014 was extracted and analyzed. But unable to be reported due to LIMS limitations

#### **METALS**

The analytical results met the requirements of the laboratory's QA/QC program.

#### **GENERAL CHEMISTRY**

Sample HB-072(4-6') for Hexavalent Chromium is suspect to have a reducing agent based on the results obtained from method of standard addition.

The matrix spike/matrix spike duplicate data for batch(es) 9357388 and 9362384 are not included in this report. The batch QC samples, which document the effect of a specific sample matrix on method performance, were not associated with a sample reported in this lot. The data, therefore, has no bearing on the samples reported herein. In order to document compliance with the QC requirement for an MS/MSD per 20 environmental samples, a summary of sample/QC associations has been provided following this case narrative.

#### **OUALITY CONTROL ELEMENTS NARRATIVE**

TestAmerica conducts a quality assurance/quality control (QA/QC) program designed to provide scientifically valid and legally defensible data. Toward this end, several types of quality control indicators are incorporated into the QA/QC program, which is described in detail in QA Policy, QA-003. These indicators are introduced into the sample testing process to provide a mechanism for the assessment of the analytical data. Program or agency specific requirements take precedence over the requirements listed in this narrative.

#### **QC BATCH**

Environmental samples are taken through the testing process in groups called QUALITY CONTROL BATCHES (QC batches). A QC batch contains up to twenty environmental samples of a similar matrix (water, soil) that are processed using the same reagents and standards. TestAmerica North Canton requires that each environmental sample be associated with a QC batch.

Several quality control samples are included in each QC batch and are processed identically to the twenty environmental samples.

For SW846/RCRA methods, QC samples include a METHOD BLANK (MB), a LABORATORY CONTROL SAMPLE (LCS) and, where appropriate, a MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD) pair or a MATRIX SPIKE/SAMPLE DUPLICATE (MS/DU) pair. If there is insufficient sample to perform an MS/MSD or an MS/DU, then a LABORATORY CONTROL SAMPLE DUPLICATE (LCSD) is included in the QC batch.

For 600 series/CWA methods, QC samples include a METHOD BLANK (MB), a LABORATORY CONTROL SAMPLE (LCS) and, where appropriate, a MATRIX SPIKE (MS). An MS is prepared and analyzed at a 10% frequency for GC Methods and at a 5% frequency for GC/MS methods.

#### LABORATORY CONTROL SAMPLE

The Laboratory Control Sample is a QC sample that is created by adding known concentrations of a full or partial set of target analytes to a matrix similar to that of the environmental samples in the QC batch. Multi peak responders may not be included in the target spike list due to co-elution. The LCS analyte recovery results are used to monitor the analytical process and provide evidence that the laboratory is performing the method within acceptable guidelines. All control analytes indicated by a bold type in the LCS must meet acceptance criteria. Failure to meet the established recovery guidelines requires the repreparation and reanalysis of all samples in the QC batch. Comparison of only the failed parameters from the first batch are evaluated. The only exception to the rework requirement is that if the LCS recoveries are biased high and the associated sample is ND (non-detected) for the parameter(s) of interest, the batch is acceptable.

At times, a Laboratory Control Sample Duplicate (LCSD) is also included in the QC batch. An LCSD is a QC sample that is created and handled identically to the LCS. Analyte recovery data from the LCSD is assessed in the same way as that of the LCS. The LCSD recoveries, together with the LCS recoveries, are used to determine the reproducibility (precision) of the analytical system. Precision data are expressed as relative percent differences (RPDs). If the RPD fails for an LCS/LCSD and yet the recoveries are within acceptance criteria, the batch is still acceptable.

#### METHOD BLANK

The Method Blank is a QC sample consisting of all the reagents used in analyzing the environmental samples contained in the QC batch. Method Blank results are used to determine if interference or contamination in the analytical system could lead to the reporting of false positive data or elevated analyte concentrations. All target analytes must be below the reporting limits (RL) or the associated sample(s) must be ND except under the following circumstances:

• Common organic contaminants may be present at concentrations up to 5 times the reporting limits. Common metals contaminants may be present at concentrations up to 2 times the reporting limit, or the reported blank concentration must be twenty fold less than the concentration reported in the associated environmental samples. (See common laboratory contaminants listed in the table.)

Volatile (GC or GC/MS)	Semivolatile (GC/MS)	Metals ICP-MS	Metals ICP Trace
Methylene Chloride,	Phthalate Esters	Copper, Iron, Zinc,	Copper, Iron, Zinc, Lead
Acetone, 2-Butanone		Lead, Calcium,	
		Magnesium, Potassium,	
		Sodium, Barium,	
		Chromium, Manganese	

## QUALITY CONTROL ELEMENTS NARRATIVE (continued)

- Organic blanks will be accepted if compounds detected in the blank are present in the associated samples at levels 10 times the blank level. Inorganic blanks will be accepted if elements detected in the blank are present in the associated samples at 20 times the blank level.
- Blanks will be accepted if the compounds/elements detected are not present in any of the associated environmental samples.

Failure to meet these Method Blank criteria requires the repreparation and reanalysis of all samples in the QC batch.

#### MATRIX SPIKE/MATRIX SPIKE DUPLICATE

A Matrix Spike and a Matrix Spike Duplicate are a pair of environmental samples to which known concentrations of a full or partial set of target analytes are added. The MS/MSD results are determined in the same manner as the results of the environmental sample used to prepare the MS/MSD. The analyte recoveries and the relative percent differences (RPDs) of the recoveries are calculated and used to evaluate the effect of the sample matrix on the analytical results. Due to the potential variability of the matrix of each sample, the MS/MSD results may not have an immediate bearing on any samples except the one spiked; therefore, the associated batch MS/MSD may not reflect the same compounds as the samples contained in the analytical report. When these MS/MSD results fail to meet acceptance criteria, the data is evaluated. If the LCS is within acceptance criteria, the batch is considered acceptable.

For certain methods, a Matrix Spike/Sample Duplicate (MS/DU) may be included in the QC batch in place of the MS/MSD. For the parameters (i.e. pH, ignitability) where it is not possible to prepare a spiked sample, a Sample Duplicate may be included in the QC batch. However, a Sample Duplicate is less likely to provide usable precision statistics depending on the likelihood of finding concentrations below the standard reporting limit. When the Sample Duplicate result fails to meet acceptance criteria, the data is evaluated.

For certain methods (600 series methods/CWA), a Matrix Spike is required in place of a Matrix Spike/Matrix Spike Duplicate (MS/MSD) or Matrix Spike/Sample Duplicate (MS/DU).

The acceptance criteria do not apply to samples that are diluted.

#### SURROGATE COMPOUNDS

In addition to these batch-related QC indicators, each organic environmental and QC sample is spiked with surrogate compounds. Surrogates are organic chemicals that behave similarly to the analytes of interest and that are rarely present in the environment. Surrogate recoveries are used to monitor the individual performance of a sample in the analytical system.

If surrogate recoveries are biased high in the LCS, LCSD, or the Method Blank, and the associated sample(s) are ND, the batch is acceptable. Otherwise, if the LCS, LCSD, or Method Blank surrogate(s) fail to meet recovery criteria, the entire sample batch is reprepared and reanalyzed. If the surrogate recoveries are outside criteria for environmental samples, the samples will be reprepared and reanalyzed unless there is objective evidence of matrix interference or if the sample dilution is greater than the threshold outlined in the associated method SOP.

The acceptance criteria do not apply to samples that are diluted. All other surrogate recoveries will be reported.

For the GC/MS BNA methods, the surrogate criterion is that two of the three surrogates for each fraction must meet acceptance criteria. The third surrogate must have a recovery of ten percent or greater.

For the Pesticide and PCB methods, the surrogate criterion is that one of two surrogate compounds must meet acceptance criteria. The second surrogate must have a recovery of 10% or greater.



## TestAmerica Certifications and Approvals:

The laboratory is certified for the analytes listed on the documents below. These are available upon request. California (#01144CA), Connecticut (#PH-0590), Florida (#E87225),

Illinois (#200004), Kansas (#E10336), Minnesota (#39-999-348), New Jersey (#OH001), New York (#10975), Nevada (#OH-000482008A), OhioVAP (#CL0024), Pennsylvania (#008), West Virginia (#210), Wisconsin (#999518190), NAVY, ARMY, USDA Soil Permit

N:\QAQC\Customer Service\Narrative - Combined RCRA CWA 032609.doc

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Data File: \\cansvr11\dd\chem\MSV\a3ux8a.i\M91223A.b\UX81645.D

Report Date: 24-Dec-2009 14:30

#### TestAmerica North Canton

#### INTERNAL STANDARD COMPOUNDS AREA AND RT SUMMARY

Calibration Date: 23-DEC-2009

Client Smp ID: HB-071(10-12')

Calibration Time: 19:33

Level: LOW

Sample Type: SOIL

Instrument ID: a3ux8a.i

Lab File ID: UX81645.D

Lab Smp Id: LRCRC1AK

Analysis Type: VOA

Quant Type: ISTD

Operator: 402279

Method File: \\cansvr11\dd\chem\MSV\a3ux8a.i\M91223A.b\8260SUX8.m

Misc Info: M91223A,8260SUX8,1-8260.SUB,402279

		AREA I	IMIT		
COMPOUND	STANDARD	LOWER	UPPER	SAMPLE	%DIFF
=======================================	=======================================	<del></del>			======
1 Fluorobenzene	895502	447751	1791004	717055	-19.93
2 Chlorobenzene-d5	695105	347553	1390210	403978	-41.88
3 1,4-Dichlorobenze	375553	187777	751106	107352	-71.41
	1	1			

		RT I	LIMIT		
COMPOUND	STANDARD	LOWER	UPPER	SAMPLE	용DIFF
		=========	========		======
1 Fluorobenzene	5.31	4.81	5.81	5.30	-0.02
2 Chlorobenzene-d5	7.85	7.35	8.35	7.85	-0.01
3 1,4-Dichlorobenze	10.04	9.54	10.54	10.04	0.05
	l I		l		ll

AREA UPPER LIMIT = +100% of internal standard area.

AREA LOWER LIMIT = -50% of internal standard area.

RT UPPER LIMIT = + 0.50 minutes of internal standard RT.

RT LOWER LIMIT = - 0.50 minutes of internal standard RT.

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Data File: \\cansvr11\dd\chem\MSV\a3ux8a.i\M91223A.b\UX81630.D

Report Date: 24-Dec-2009 14:13

#### TestAmerica North Canton

#### INTERNAL STANDARD COMPOUNDS AREA AND RT SUMMARY

Instrument ID: a3ux8a.i Lab File ID: UX81630.D Lab Smp Id: LRCRE1AK

Analysis Type: VOA Quant Type: ISTD

Operator: 402279

Calibration Date: 23-DEC-2009 Calibration Time: 19:33 Client Smp ID: HB-071(10-12')

Level: LOW

Sample Type: SOIL

Method File: \\cansvrl1\dd\chem\MSV\a3ux8a.i\M91223A.b\8260SUX8.m

Misc Info: M91223A,8260SUX8,1-8260.SUB,402279

1		AREA I	TIMIT		
COMPOUND	STANDARD	LOWER	UPPER	SAMPLE	%DIFF
=====================================	=======================================	=======================================		=======	=====
1 Fluorobenzene	895502	447751	1791004	664514	-25.79
2 Chlorobenzene-d5	695105	347553	1390210	333803	-51.98 <
3 1,4-Dichlorobenze	375553	187777	751106	78423	-79.12 <
i i	ĺ	1			1

	]	RT L	IMIT		
COMPOUND	STANDARD	LOWER	UPPER	SAMPLE	%DIFF
====== <del>==============================</del>	========	========	========	=========	======
1 Fluorobenzene	5.31	4.81	5.81	5.31	0.05
2 Chlorobenzene-d5	7.85	7.35	8.35	7.85	-C.04
3 1,4-Dichlorobenze	10.04	9.54	10.54	10.04	0.03
1	1				1

AREA UPPER LIMIT = +100% of internal standard area.

AREA LOWER LIMIT = -50% of internal standard area.

RT UPPER LIMIT = + 0.50 minutes of internal standard RT.

RT LOWER LIMIT = - 0.50 minutes of internal standard RT.

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Data File: \\cansvr11\dd\chem\MSV\a3ux8a.i\M91223A.b\UX81631.D

Report Date: 24-Dec-2009 14:16

#### TestAmerica North Canton

#### INTERNAL STANDARD COMPOUNDS AREA AND RT SUMMARY

Instrument ID: a3ux8a.i Lab File ID: UX81631.D Lab Smp Id: LRCRH1AK Analysis Type: VOA

Analysis Type: VOA
Quant Type: ISTD
Operator: 402279

Method File: \\cansvr11\dd\chem\MSV\a3ux8a.1\ Misc Info: M91223A,8260SUX8,1-8260.SUB,402279

Calibration Date: 23-DEC-2009

Calibration Time: 19:33 Client Smp ID: HB-072(2-4')

Level: LOW Sample Type: SOIL

Method File: \\cansvr11\dd\chem\MSV\a3ux8a.i\M91223A.b\8260SUX8.m

	Ì	AREA I	TIMI	ĺ	
COMPOUND	STANDARD	LOWER	UPPER	SAMPLE	%DIFF
:	=======================================			=======================================	======
1 Fluorobenzene	895502	447751!	1791004	684851	-23.52
2 Chlorobenzene-d5	695105	347553	1390210	332213	-52.21 <
3 1,4-Dichlorobenze	375553	187777	751106	77254	-79.43 <
	I			1	1

		RT I	IMIT !		
COMPOUND	STANDARD	LOWER	UPPER	SAMPLE	%DIFF
======================================	=======================================	===== <b>====</b>			======
1 Fluorobenzene	5.31	4.81	5.81	5.31	0.05
2 Chlorobenzene-d5	7.85	7.35	8.35	7.85	0.04
3 1,4-Dichlorobenze	10.04	9.54	10.54	10.04	0.03
:	İ			j	I

AREA UPPER LIMIT = +100% of internal standard area. AREA LOWER LIMIT = -50% of internal standard area. RT UPPER LIMIT = +0.50 minutes of internal standard RT. RT LOWER LIMIT = -0.50 minutes of internal standard RT.

# TESTAMERICA LABORATORIES, INC.

# MS RUN NUMBER REVIEW

Lot ID	Smp#	Work Order	Batch	MS Run#	SDG	Prep Date	Method
A9L180470	001	LRA5L1AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	002	LRA5R1AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	003	LRA5T1AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	004	LRA5V1AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	005	LRA5W1AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	006	LRA5X1AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	007	LRA511AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	800	LRA531AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	009	LRA541AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	010	LRA551AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	011	LRA571AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180470	012	LRA591AE	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180492	001	LRCET1AL	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180492	002	LRCRC1AL	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L180492	003	LRCRE1AL	9357388	9357236		12/23/09	MCAWW 335.2 CLP-M
A9L220458	004	LRG481AH	9357388	9357236	9L03652	12/23/09	MCAWW 335.2 CLP-M
A9L220458	004	LRG481AL D	9357388	9357236	9L03652	12/23/09	MCAWW 335.2 CLP-M
A9L220458	004	LRG481AK S	9357388	9357236	9L03652	12/23/09	MCAWW 335.2 CLP-M
A9L170448	001	LQ8JA1AR	9357395	9357236		12/23/09	SW846 9012A
A9L150490	003	LQ4981AA	9362383	9362205		12/28/09	SW846 9012A
A9L150490	003	LQ4981AD D	9362383	9362205		12/28/09	SW846 9012A
A9L150490	003	LQ4981AC S	9362383	9362205		12/28/09	SW846 9012A
A9L180492	004	LRCRH1AL	9362384	9362205		12/28/09	MCAWW 335.2 CLP-M
A9L180492	005	LRCRK1AL	9362384	9362205		12/28/09	MCAWW 335.2 CLP-M
A9L180492	006	LRCRM1AL	9362384	9362205		12/28/09	MCAWW 335.2 CLP-M
A9L180492	007	LRCRP1AL	9362384	9362205		12/28/09	MCAWW 335.2 CLP-M
A9L180492	800	LRCRRIAL	9362384	9362205		12/28/09	MCAWW 335.2 CLP-M

## A9L180492

PARAMETER		RESULT	REPORTING LIMIT	UNITS	ANALYTICAL METHOD	
нв-071(8-10') 12/17	7/09 11:40 001					
C20-C34		250	31	mg/kg	SW846 8015B	
C10-C20		260	23	mg/kg	SW846 8015B	
Gasoline Rar (C6-C12)	nge Organics	2700	1200	ug/kg	SW846 8015A MOI	)
Arsenic		10.0	1.2	mg/kg	SW846 6010B	
Chromium		11.2	0.58	mg/kg	SW846 6010B	
Lead		38.8	0.35	mg/kg	SW846 6010B	
Benzo(a)anth	nracene	2200	380	ug/kg	SW846 8270C	
Benzo(b)fluo	oranthene	3300	380	ug/kg	SW846 8270C	
Benzo(k)fluo		570	380	ug/kg	SW846 8270C	
Benzo(a)pyre		2200	380	ug/kg	SW846 8270C	
Chrysene		2200	380	ug/kg	SW846 8270C	
Indeno(1,2,3	3-cd)pyrene	1400	380	ug/kg	SW846 8270C	
Naphthalene	, ou, p, 1 one	55000	380	ug/kg	SW846 8270C	
Acenaphthene	2	7000	380	ug/kg	SW846 8270C	
Anthracene		2400	380	ug/kg	SW846 8270C	
Benzo(ghi)pe	rvlene	830	380	ug/kg	SW846 8270C	
Fluoranthene		8800	380	ug/kg	SW846 8270C	
Fluorene		5300	380	ug/kg	SW846 8270C	
2-Methylnaph	nthalene	7900	380	ug/kg	SW846 8270C	
Phenanthrene		13000	380	ug/kg	SW846 8270C	
Pyrene	•	5600	380	ug/kg	SW846 8270C	
1-Methylnaph	thalene	4100	380	ug/kg	SW846 8270C	
Corrosivity	Itharene	9.5	300	No Units	SW846 9045A	
Percent Soli	de	86.8	10.0	%	MCAWW 160.3 MOI	`
Acid-soluble	:	48.5	34.6	mg/kg	SW846 9030B/903	
Total Cyanic		0.80	0.58	mg/kg	MCAWW 335.2 CLI	
HB-071(10-12') 12/1	17/09 11:40 002					
C20-C34		220	31	mg/kg	SW846 8015B	
C10-C20		170	23	mg/kg	SW846 8015B	
Gasoline Ran	ige Organics	170	120	ug/kg	SW846 8015A MOI	)
(C6-C12)	ige organico					
Arsenic		11.3	1.2	mg/kg	SW846 6010B	
Cadmium		0.31	0.23	mg/kg	SW846 6010B	
Chromium		14.4	0.58	mg/kg	SW846 6010B	
Lead		65.9	0.35	mg/kg	SW846 6010B	
Benzo(a)anth		1500	51	ug/kg	SW846 8270C	
Benzo(b)fluo		2200	51	ug/kg	SW846 8270C	
Benzo(k)fluc	ranthene	830	51	ug/kg	SW846 8270C	
Benzo(a)pyre	ene	1300	51	ug/kg	SW846 8270C	
Chrysene		1500	51	ug/kg	SW846 8270C	

# A9L180492

PARAMETER	RESULT	REPORTING LIMIT	UNITS	ANALYTICAL METHOD
HB-071(10-12') 12/17/09 11:40 002				
Dibenz(a,h)anthracene	380	51	ug/kg	SW846 8270C
Indeno(1,2,3-cd)pyrene	940	51	ug/kg	SW846 8270C
Naphthalene	3700	51	ug/kg	SW846 8270C
Acenaphthene	1300	51	ug/kg	SW846 8270C
Acenaphthylene	270	51	ug/kg	SW846 8270C
Anthracene	940	51	ug/kg	SW846 8270C
Benzo(ghi)perylene	960	51	ug/kg	SW846 8270C
Fluoranthene	4000	51	ug/kg	SW846 8270C
Fluorene	1300	51	ug/kg	SW846 8270C
2-Methylnaphthalene	3300	51	ug/kg	SW846 8270C
Phenanthrene	5400	51	ug/kg	SW846 8270C
Pyrene	3100	51	ug/kg	SW846 8270C
1-Methylnaphthalene	2000	51	ug/kg	SW846 8270C
Methylene chloride	12	5.8	ug/kg	SW846 8260A
Acetone	36	23	ug/kg	SW846 8260A
Carbon disulfide	14	5.8	ug/kg	SW846 8260A
Toluene	7.1	5.8	ug/kg	SW846 8260A
Xylenes (total)	11	5.8	ug/kg	SW846 8260A
Corrosivity	10.1		No Units	SW846 9045A
Percent Solids	86.8	10.0	8	MCAWW 160.3 MOD
Acid-soluble sulfide	90.2	34.6	mg/kg	SW846 9030B/9034
Total Cyanide	2.1	0.58	mg/kg	MCAWW 335.2 CLP-M
HB-071(16-18') 12/17/09 11:40 003				
C20-C34	150	31	mg/kg	SW846 8015B
C10-C20	72	23	mg/kg	SW846 8015B
Gasoline Range Organics (C6-C12)	150	110	ug/kg	SW846 8015A MOD
Arsenic	9.7	1.1	mg/kg	SW846 6010B
Chromium	22.9	0.57	mg/kg	SW846 6010B
Lead	55.7	0.34	mg/kg	SW846 6010B
Benzo(a)anthracene	810	30	ug/kg	SW846 8270C
Benzo(b)fluoranthene	1000	30	ug/kg	SW846 8270C
Benzo(k)fluoranthene	420	30	ug/kg	SW846 8270C
Benzo(a)pyrene	660	30	ug/k <b>g</b>	SW846 8270C
Chrysene	850	30	ug/kg	SW846 8270C
Dibenz(a,h)anthracene	180	30	ug/kg	SW846 8270C
Indeno(1,2,3-cd)pyrene	440	30	ug/kg	SW846 8270C
Naphthalene	3400	30	ug/kg	SW846 8270C
Acenaphthene	900	30	ug/kg	SW846 8270C
Acenaphthylene	120	30	ug/kg	SW846 8270C
Anthracene	490	30	ug/kg	SW846 8270C

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PARAMETER	RESULT	REPORTING LIMIT	UNITS	ANALYTICAL METHOD
HB-071(16-18') 12/17/09 11:40 003				
Benzo(ghi)perylene	450	30	ug/kg	SW846 8270C
Fluoranthene	2200	30	ug/kg	SW846 8270C
Fluorene	870	30	ug/kg	SW846 8270C
2-Methylnaphthalene	1600	30	ug/kg	SW846 8270C
Phenanthrene	2700	30	ug/kg	SW846 8270C
Pyrene	1800	30	ug/kg	SW846 8270C
1-Methylnaphthalene	960	30	ug/kg	SW846 8270C
Methylene chloride	7.3	5.7	ug/kg	SW846 8260A
Carbon disulfide	12	5.7	ug/kg	SW846 8260A
Corrosivity	10.4		No Units	SW846 9045A
Percent Solids	87.9	10.0	8	MCAWW 160.3 MOD
Acid-soluble sulfide	148	34.1	mg/kg	SW846 9030B/9034
Total Cyanide	0.73	0.57	mg/kg	MCAWW 335.2 CLP-M
HB-072(2-4') 12/17/09 12:20 004				
C20-C34	140	30	mg/kg	SW846 8015B
C10-C20	44	23	mg/kg	SW846 8015B
Arsenic	8.6	1.1	mg/kg	SW846 6010B
Chromium	10.6	0.56	mg/kg	SW846 6010B
Lead	27.0	0.34	mg/kg	SW846 6010B
Benzo(a)anthracene	840	7.5	ug/kg	SW846 8270C
Benzo(b) fluoranthene	1100	7.5	ug/kg	SW846 8270C
Benzo(k) fluoranthene	390	7.5	ug/kg	SW846 8270C
Benzo(a)pyrene	630	7.5	ug/kg	SW846 8270C
Chrysene	840	7.5	ug/kg	SW846 8270C
Dibenz(a,h)anthracene	150	7.5	ug/kg	SW846 8270C
Indeno(1,2,3-cd)pyrene	460	7.5	ug/kg	SW846 8270C
Naphthalene	580	7.5	ug/kg	SW846 8270C
Acenaphthene	110	7.5	ug/kg	SW846 8270C
Acenaphthylene	140	7.5	ug/kg	SW846 8270C
Anthracene	360	7.5	ug/kg	SW846 8270C
Benzo(ghi)perylene	500	7.5	ug/kg	SW846 8270C
Fluoranthene	1500	7.5	ug/kg	SW846 8270C
Fluorene	140	7.5	ug/kg	SW846 8270C
2-Methylnaphthalene	780	7.5	ug/kg	SW846 8270C
Phenanthrene	1100	7.5	ug/kg	SW846 8270C
Pyrene	1400	7.5	ug/kg	SW846 8270C
1-Methylnaphthalene	520	7.5	ug/kg	SW846 8270C
Methylene chloride	11	5.6	ug/kg	SW846 8260A
Carbon disulfide	8.0	5.6	ug/kg	SW846 8260A
Corrosivity	10.2		No Units	SW846 9045A
Percent Solids	88.7	10.0	8	MCAWW 160.3 MOD

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		REPORTIN	G	ANALYTICAL
PARAMETER	RESULT	LIMIT	<u>UNITS</u>	METHOD
HB-072(2-4') 12/17/09 12:20 004				
Acid-soluble sulfide	287	33.8	mg/kg	SW846 9030B/9034
Total Cyanide	2.9	0.56	mg/kg	MCAWW 335.2 CLP-M
HB-072(4-6') 12/17/09 12:20 005				
1115 072 (4 0 7 127 177 05 12120 005				
C20-C34	220	31	mg/kg	SW846 8015B
C10-C20	82	23	mg/kg	SW846 8015B
Gasoline Range Organics (C6-C12)	700	580	ug/kg	SW846 8015A MOD
Arsenic	9.3	1.2	mg/kg	SW846 6010B
Chromium	19.7	0.58	mg/kg	SW846 6010B
Lead	38.4	0.35	mg/kg	SW846 6010B
Benzo(a)anthracene	1300	15	ug/kg	SW846 8270C
Benzo(b)fluoranthene	2100	15	ug/kg	SW846 8270C
Benzo(k)fluoranthene	750	15	ug/kg	SW846 8270C
Benzo(a)pyrene	1200	15	ug/kg	SW846 8270C
Chrysene	1500	15	ug/kg	SW846 8270C
Dibenz(a,h)anthracene	260	15	ug/kg	SW846 8270C
Indeno(1,2,3-cd)pyrene	760	15	ug/kg	SW846 8270C
Naphthalene	910	15	ug/kg	SW846 8270C
Acenaphthene	140	15	ug/kg	SW846 8270C
Acenaphthylene	340	15	ug/kg	SW846 8270C
Anthracene	410	15	ug/kg	SW846 8270C
Benzo(ghi)perylene	850	<b>1</b> 5	ug/kg	SW846 8270C
Fluoranthene	2100	15	ug/kg	SW846 8270C
Fluorene	180	15	ug/kg	SW846 8270C
2-Methylnaphthalene	1400	15	ug/kg	SW846 8270C
Phenanthrene	1400	15	ug/kg	SW846 8270C
Pyrene	2100	15	ug/kg	SW846 8270C
1-Methylnaphthalene	860	15	ug/kg	SW846 8270C
Carbon disulfide	27	5.8	ug/kg	SW846 8260A
Benzene	27	5.8	ug/kg	SW846 8260A
Toluene	44	5.8	ug/kg	SW846 8260A
Xylenes (total)	20	5.8	ug/kg	SW846 8260A
Hexavalent	2.5	0.92	mg/kg	SW846 7196A
Chromium				
Corrosivity	10.2		No Units	SW846 9045A
Percent Solids	86.7	10.0	Se	MCAWW 160.3 MOD
Acid-soluble sulfide	289	34.6	mg/kg	SW846 9030B/9034
Total Cyanide	2.1	0.58	mg/kg	MCAWW 335.2 CLP-M

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PARAMETER	RESULT	REPORTING LIMIT	UNITS	ANALYTICAL METHOD
4,114,111,111,111				
HB-073(0-2') 12/17/09 12:50	006			
C20-C34	130	31	mg/kg	SW846 8015B
C10-C20	120	23	mg/kg	SW846 8015B
Gasoline Range Organi	cs 210	110	ug/kg	SW846 8015A MOD
(C6-C12)				
Arsenic	9.3	1.1	mg/kg	SW846 6010B
Cadmium	0.33	0.23	mg/kg	SW846 6010B
Chromium	10.6	0.57	mg/kg	SW846 6010B
Lead	115	0.34	mg/kg	SW846 6010B
Benzo(a)anthracene	1100	15	ug/kg	SW846 8270C
Benzo(b)fluoranthene	1500	15	ug/kg	SW846 8270C
Benzo(k)fluoranthene	420	15	ug/kg	SW846 8270C
Benzo(a)pyrene	1000	15	ug/kg	SW846 8270C
Chrysene	960	15	ug/kg	SW846 8270C
Dibenz(a,h)anthracene	290	15	ug/kg	SW846 8270C
Indeno(1,2,3-cd)pyren	e 600	15	ug/kg	SW846 8270C
Naphthalene	880	15	ug/kg	SW846 8270C
Acenaphthene	82	15	ug/kg	SW846 8270C
Acenaphthylene	150	15	ug/kg	SW846 8270C
Anthracene	240	15	ug/kg	SW846 8270C
Benzo(ghi)perylene	820	15	ug/kg	SW846 8270C
Fluoranthene	1600	15	ug/kg	SW846 8270C
Fluorene	87	15	ug/kg	SW846 8270C
2-Methylnaphthalene	1500	15	ug/kg	SW846 8270C
Phenanthrene	1200	15	ug/kg	SW846 8270C
Pyrene	1500	15	ug/kg	SW846 8270C
1-Methylnaphthalene	1000	15	ug/kg	SW846 8270C
Toluene	5.8	5.7	ug/kg	SW846 8260A
Ethylbenzene	11	5.7	ug/kg	SW846 8260A
Xylenes (total)	47	5.7	ug/kg	SW846 8260A
Corrosivity	11.0		No Units	SW846 9045A
Percent Solids	88.4	10.0	€	MCAWW 160.3 MOD
Acid-soluble sulfide	302	33.9	mg/kg	SW846 9030B/9034
Total Cyanide	4.6	0.57	mg/kg	MCAWW 335.2 CLP-M
HB-075(0-2') 12/17/09 13:20	007			
C20-C34	610	150	mg/kg	SW846 8015B
C10-C20	100 J	110	mg/kg	SW846 8015B
Gasoline Range Organi		570	ug/kg	SW846 8015A MOD
(C6-C12)				
Arsenic	4.7	1.1	mg/kg	SW846 6010B
Cadmium	0.23	0.23	mg/kg	SW846 6010B
Chromium	18.6	0.57	mg/kg	SW846 6010B

## A9L180492

			REPORTIN	īG	ANALYTICAL
	PARAMETER	RESULT	LIMIT	UNITS	METHOD
HB-07!	5(0-2') 12/17/09 13:20 007				
	Lead	21.3	0.34	mg/kg	SW846 6010B
	Benzo(a)anthracene	2900	76	ug/kg	SW846 8270C
	Benzo(b)fluoranthene	2500	76	ug/kg	SW846 8270C
	Benzo(k)fluoranthene	940	76	ug/kg	SW846 8270C
	Benzo(a)pyrene	2100	76	ug/kg	SW846 8270C
	Chrysene	2600	76	ug/kg	SW846 8270C
	Dibenz(a,h)anthracene	460	76	ug/kg	SW846 8270C
	Indeno(1,2,3-cd)pyrene	1100	76	ug/kg	SW846 8270C
	Naphthalene	230	76	ug/kg	SW846 8270C
	Acenaphthene	190	76	ug/kg	SW846 8270C
	Acenaphthylene	100	76	ug/kg	SW846 8270C
	Anthracene	860	76	ug/kg	SW846 8270C
	Benzo(ghi)perylene	1200	76	ug/kg	SW846 8270C
	Fluoranthene	5100	76	ug/kg	SW846 8270C
	Fluorene	210	76	ug/kg	SW846 8270C
	2-Methylnaphthalene	180	76	ug/kg	SW846 8270C
	Phenanthrene	2500	76	ug/kg	SW846 8270C
	Pyrene	6100	76	ug/kg	SW846 8270C
	1-Methylnaphthalene	150	76	ug/kg	SW846 8270C
	Xylenes (total)	26	5.7	ug/kg	SW846 8260A
	Corrosivity	11.5		No Units	SW846 9045A
	Percent Solids	87.9	10.0	%	MCAWW 160.3 MOD
	Acid-soluble sulfide	61.6	34.1	mg/kg	SW846 9030B/9034
нв-075	5(2-4') 12/17/09 13:20 008				
	C20-C34	110	15	mg/kg	SW846 8015B
	C10-C20	48	11	mg/kg	SW846 8015B
	Arsenic	10.4	1.1	mg/kg	SW846 6010B
	Chromium	8.9	0.55	mg/kg	SW846 6010B
	Lead	107	0.33	mg/kg	SW846 6010B
	Benzo(a)anthracene	2900	29	ug/kg	SW846 8270C
	Benzo(b)fluoranthene	3200	29	ug/kg	SW846 8270C
	Benzo(k)fluoranthene	1200	29	ug/kg	SW846 8270C
	Benzo(a)pyrene	2500	29	ug/kg	SW846 8270C
	Chrysene	2600	29	ug/kg	SW846 8270C
	Dibenz(a,h)anthracene	500	29	ug/kg	SW846 8270C
	Indeno(1,2,3-cd)pyrene	1500	29	ug/kg	SW846 8270C
	Naphthalene	820	29	ug/kg	SW846 8270C
	Acenaphthene	570	29	ug/kg	SW846 8270C
	Acenaphthylene	72	29	ug/kg	SW846 8270C
	Anthracene	1300	29	ug/kg	SW846 8270C
	Benzo(ghi)perylene	1600	29	ug/kg	SW846 8270C

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PARAMETER	RESULT	REPORTING LIMIT	UNITS	ANALYTICAL METHOD
HB-075(2-4') 12/17/09 13:20 008	·			
Fluoranthene	5500	29	ug/kg	SW846 8270C
Fluorene	570	29	ug/kg	SW846 8270C
2-Methylnaphthalene	1000	29	ug/kg	SW846 8270C
Phenanthrene	4800	29	ug/kg	SW846 8270C
Pyrene	5400	29	ug/kg	SW846 8270C
1-Methylnaphthalene	610	29	ug/kg	SW846 8270C
Corrosivity	8.7		No Units	SW846 9045A
Percent Solids	90.6	10.0	ક	MCAWW 160.3 MOD

# ANALYTICAL METHODS SUMMARY

## A9L180492

PARAMETER	ANALYTICAL METHOD
Corrosivity	SW846 9045A
Extractable Petroleum Hydrocarbons	SW846 8015B
Hexavalent Chromium	SW846 7196A
Semivolatile Organic Compounds by GC/MS	SW846 8270C
Sulfides, Total 9030B/9034	SW846 9030B/9034
Total Cyanide	MCAWW 335.2 CLP-M
Total Residue as Percent Solids	MCAWW 160.3 MOD
Trace Inductively Coupled Plasma (ICP) Metals	SW846 6010B
Volatile Organics	SW846 8015A MOD
Volatile Organics by GC/MS	SW846 8260A

## References:

MCAWW	"Methods for Chemical Analysis of Water and Wastes",
	EPA-600/4-79-020, March 1983 and subsequent revisions.
SW846	"Test Methods for Evaluating Solid Waste, Physical/Chemical
	Methods". Third Edition. November 1986 and its updates.

# SAMPLE SUMMARY

#### A9L180492

<u>WO #</u> .	SAMPLE#	CLIENT SAMPLE ID	SAMPLED DATE	SAMP TIME
LRCET	001	нв-071(8-10')	12/17/09	11:40
LRCRC	002	HB-071(10-12')	12/17/09	11:40
LRCRE	003	HB-071(16-18')	12/17/09	11:40
LRCRH	004	нв-072(2-4')	12/17/09	12:20
LRCRK	005	нв-072(4-6')	12/17/09	12:20
LRCRM	006	нв-073 (0-2')	12/17/09	12:50
LRCRP	007	HB-075(0-2')	12/17/09	13:20
LRCRR	008	HB-075(2-4')	12/17/09	13:20

## NOTE(S):

- The analytical results of the samples listed above are presented on the following pages.
- All calculations are performed before rounding to avoid round-off errors in calculated results.
- Results noted as "ND" were not detected at or above the stated limit.
- This report must not be reproduced, except in full, without the written approval of the laboratory.
- Results for the following parameters are never reported on a dry weight basis: color, corrosivity, density, flashpoint, ignitability, layers, odor, paint filter test, pH, porosity pressure, reactivity, redox potential, specific gravity, spot tests, solids, solubility, temperature, viscosity, and weight.

# Client Sample ID: HB-071(8-10')

## GC/MS Volatiles

Lot-Sample #: A9L180492-00 Date Sampled: 12/17/09 11: Prep Date: 12/20/09 Prep Batch #: 9356563		12/18/09	Matrix: SC
Dilution Factor: 19.92 % Moisture: 13	Method:	SW846 8260	A(
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Acetone	ND	23000	ug/kg
Benzene	ND	5700	ug/kg
Bromodichloromethane	ND	5700	ug/kg
Bromoform	ND	5700	ug/kg
Bromomethane	ND	11000	ug/kg
2-Butanone	ND	23000	ug/kg
Carbon disulfide	ND	5700	ug/kg
Carbon tetrachloride	ND	5700	ug/kg
Chlorobenzene	ND	5700	ug/kg
Dibromochloromethane	ND	5700	ug/kg
Chloroethane	ND	11000	ug/kg
Chloroform	ND	5700	ug/kg
Chloromethane	ND	11000	ug/kg
1,1-Dichloroethane	ND	5700	ug/kg
1,2-Dichloroethane	ND	5700	ug/kg
1,1-Dichloroethene	ND	5700	ug/kg
1,2-Dichloroethene (total)	ND	5700	ug/kg
1,2-Dichloropropane	ND	5700	ug/kg
cis-1,3-Dichloropropene	ND	5700	ug/kg
trans-1,3-Dichloropropene	ND	5700	ug/kg
Ethylbenzene	ND	5700	ug/kg
2-Hexanone	ND	23000	ug/kg
Methylene chloride	ND	5700	ug/kg
4-Methyl-2-pentanone	ND	23000	ug/kg
Styrene	ND	5700	ug/kg
1,1,2,2-Tetrachloroethane	ND	5700	ug/kg
Tetrachloroethene	ND	5700	ug/ <b>kg</b>
Toluene	ND	5700	ug/kg
1,1,1-Trichloroethane	ND	5700	ug/kg
1,1,2-Trichloroethane	ND	5700	ug/kg
Trichloroethene	ND	5700	ug/kg
Vinyl chloride	ND	2300	ug/kg
Xylenes (total)	ND	5700	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Dibromofluoromethane	74 DIL	(59 - 138)	
1,2-Dichloroethane-d4	0.0 DIL,*	(61 - 130)	
Toluene-d8	70 DIL	(60 - 143)	
4-Bromofluorobenzene	76 DIL	(47 - 158)	

(Continued on next page)

Client Sample ID: HB-071(8-10')

## GC/MS Volatiles

Lot-Sample #...: A9L180492-001 Work Order #...: LRCET1AK Matrix.....: SO

NOTE(S):

DLL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

<sup>\*</sup> Surrogate recovery is outside stated control limits.

Client Sample ID: HB-071(8-10')

#### GC/MS Semivolatiles

Lot-Sample #: A9L180492-001 Date Sampled: 12/17/09 11:40 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 50	Work Order #: Date Received: Analysis Date:	12/18/09	Matrix: SO
<b>% Moisture:</b> 13	Method:	SW846 8270	С
		REPORTING	
PARAMETER	RESULT	LIMIT	<u>UNITS</u>
Benzo(a)anthracene	2200	380	ug/kg
Benzo(b) fluoranthene	3300	380	ug/kg
Benzo(k) fluoranthene	570	380	ug/kg
Benzo(a)pyrene	2200	380	ug/kg
Chrysene	2200	380	ug/kg
Dibenz(a,h)anthracene	ND	380	ug/kg
Indeno(1,2,3-cd)pyrene	1400	380	ug/kg
Naphthalene	55000	380	ug/kg
Acenaphthene	7000	380	ug/kg
Acenaphthylene	ND	380	ug/kg
Anthracene	2400	380	ug/kg
Benzo(ghi)perylene	830	380	ug/kg
Fluoranthene	8800	380	ug/kg
Fluorene	5300	380	ug/kg
2-Methylnaphthalene	7900	380	ug/kg
Phenanthrene	13000	380	ug/kg
Pyrene	5600	380	ug/kg
1-Methylnaphthalene	4100	380	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	65 DIL	(24 - 112)	
2-Fluorobiphenyl	0.0 DIL,*	(34 - 110)	
Terphenyl-d14	87 DIL	(41 - 119)	
Phenol-d5	63 DIL	(28 - 110)	
2-Fluorophenol	61 DIL	(26 - 110)	
2,4,6-Tribromophenol	0.0 DIL,*	(10 - 118)	

 $<sup>\</sup>label{eq:def:DIL} \textbf{DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.}$ 

NOTE(S):

<sup>\*</sup> Surrogate recovery is outside stated control limits.

Client Sample ID: HB-071(8-10')

#### GC Volatiles

Lot-Sample #: A9L180492-00	Work Order #: LRCET1AJ	Matrix SO
----------------------------	------------------------	-----------

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09 Prep Date....: 12/19/09 Analysis Date..: 12/19/09

Prep Batch #...: 9354104

Dilution Factor: 10

**% Moisture....:** 13 **Method.....:** SW846 8015A MOD

REPORTING

PARAMETER RESULT LIMIT UNITS
Gasoline Range Organics 2700 1200 ug/kg

(C6-C12)

PERCENT RECOVERY
SURROGATE RECOVERY LIMITS
Trifluorotoluene 103 (10 - 150)

NOTE(S):

Client Sample ID: HB-071(8-10')

## GC Semivolatiles

Lot-Sample #: A9L180492-001			Matrix SO
Date Sampled: 12/17/09 11:40	Analysis Date:		
<b>-</b>	Analysis Date	12/23/09	
Prep Batch #: 9353014 Dilution Factor: 10			
% Moisture: 13	Method:	GM846 8015	D.
6 MOISCUIE: 13	rechod	20040 0013	
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
C20-C34	250	31	mg/kg
C10-C20	260	23	mg/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
C9 (nonane)	28 DIL	(10 - 110)	

NOTE (S):

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

# Client Sample ID: HB-071(8-10')

## TOTAL Metals

Lot-Sample #...: A9L180492-001 Matrix.....: S0

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

**% Moisture....:** 13

PARAMETER	RESULT	REPORTII LIMIT	NG <u>UNITS</u>	METHOD		ORK RDER #
Prep Batch #	: 9354052 10.0	1.2	mg/kg	SW846 6010B	12/21-12/23/09 L	RCET1AE
Arsenic	10.0	Dilution Fac			,	
Cadmium	ND	0.23 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 L	RCET1AF
Chromium	11.2	0.58 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 ь	RCET1AG
Lead	38.8	0.35 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 L	RCET1AH
NOTE(S):						

Client Sample ID: HB-071(8-10')

#### General Chemistry

Lot-Sample #...: A9L180492-001 Work Order #...: LRCET Matrix...... S0

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

% Moisture....: 13

PARAMETER	RESULT	RL	UNITS	METHOI	)	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide	48.5	34.6	mg/kg	SW846	9030B/9034	12/23/09	9357096
		Dilution Facto	or: 1				
Corrosivity	9.5		No Units	SW846	9045A	12/19/09	9353120
		Dilution Facto	or: 1				
Hexavalent Chromium	ND	0.92	mg/kg	SW846	7196A	12/23-12/24/09	9357236
		Dilution Facto	or: 1				
Percent Solids	86.8	10.0 Dilution Facto	<b>%</b> or: 1	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	0.80	0.58 Dilution Facto	mg/kg or: 1	MCAWW	335.2 CLP-	1 12/23/09	9357388

#### NOTE(S):

RL Reporting Limit

#### Client Sample ID: HB-071(10-12')

#### GC/MS Volatiles

Lot-Sample #...: A9L180492-002 Work Order #...: LRCRC1AK Matrix.....: S0

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09 Prep Date....: 12/24/09 Analysis Date..: 12/24/09

**Prep Batch #...:** 9362414

Dilution Factor: 1

**% Moisture....:** 13 **Method.....:** SW846 8260A

% Moisture: 13	Method:	SW846 8260	)A
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Chloromethane	ND	12	ug/kg
Bromomethane	ND	12	ug/kg
Vinyl chloride	ИD	2.3	ug/kg
Chloroethane	ND	12	ug/kg
Methylene chloride	12	5.8	ug/kg
Acetone	36	23	ug/kg
Carbon disulfide	14	5.8	ug/kg
1,1-Dichloroethene	ND	5.8	ug/kg
1,1-Dichloroethane	ND	5.8	ug/kg
1,2-Dichloroethene	ND	5.8	ug/kg
(total)			
Chloroform	ND	5.8	ug/kg
1,2-Dichloroethane	ND	5.8	ug/kg
2-Butanone	ND	23	ug/kg
1,1,1-Trichloroethane	ND	5.8	ug/kg
Carbon tetrachloride	ND	5.8	ug/kg
Bromodichloromethane	ND	5.8	ug/kg
1,2-Dichloropropane	ND	5.8	ug/kg
cis-1,3-Dichloropropene	ND	5.8	ug/kg
Trichloroethene	ND	5.8	ug/kg
Dibromochloromethane	ND	5.8	ug/kg
1,1,2-Trichloroethane	ND	5.8	ug/kg
Benzene	ND	5.8	ug/kg
trans-1,3-Dichloropropene	ND	5.8	ug/kg
Bromoform	ND	5.8	ug/kg
4-Methyl-2-pentanone	ND	23	ug/kg
2-Hexanone	ИD	23	ug/kg
Tetrachloroethene	ND	5.8	ug/kg
Toluene	7.1	5.8	ug/kg
Chlorobenzene	ND	5.8	ug/kg
Ethylbenzene	ND	5.8	ug/kg
Styrene	ND	5.8	ug/kg
Xylenes (total)	11	5.8	ug/kg
		D. D. C.	
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	_
Dibromofluoromethane	81	(59 - 138)	
1,2-Dichloroethane-d4	81	(61 - 130)	
Toluene-d8	83 63	(60 - 143)	
4-Bromofluorobenzene	0.3	(47 - 158)	

Client Sample ID: HB-071(10-12')

## GC/MS Volatiles

Lot-Sample #...: A9L180492-002 Work Order #...: LRCRC1AK Matrix.....: S0

NOTE(S):

# Client Sample ID: HB-071(10-12')

# GC/MS Volatiles

Lot-Sample #: A9L180492-002 Date Sampled: 12/17/09 11:40 Prep Date: 12/23/09 Prep Batch #: 9362414		12/18/09	<b>Matrix</b> : SO
Dilution Factor: 10 % Moisture: 13	Method:	CM846 8360	71
6 Moisture: 13	method:	5W040 020U	A
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
1,1,2,2-Tetrachloroethane	ND	58	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Dibromofluoromethane	94	(59 - 138)	
1,2-Dichloroethane-d4	88	(61 - 130)	
Toluene-d8	96	(60 - 143)	
4-Bromofluorobenzene	84	(47 - 158)	
NOTE(S):			

Client Sample ID: HB-071(10-12')

#### GC/MS Semivolatiles

Lot-Sample #: A9L180492-002 Date Sampled: 12/17/09 11:40 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 6.66		12/18/09	Matrix: SO
<b>% Moisture:</b> 13	Method:	SW846 8270	С
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Benzo(a)anthracene	1500	51	ug/kg
Benzo(b) fluoranthene	2200	51	ug/kg
Benzo(k)fluoranthene	830	51	ug/kg
Benzo(a)pyrene	1300	51	ug/kg
Chrysene	1500	51	ug/kg
Dibenz(a,h)anthracene	380	51	ug/kg
Indeno(1,2,3-cd)pyrene	940	51	ug/kg
Naphthalene	3700	51	ug/kg
Acenaphthene	1300	51	ug/kg
Acenaphthylene	270	51	ug/kg
Anthracene	940	51	ug/kg
Benzo(ghi)perylene	960	51	ug/kg
Fluoranthene	4000	51	ug/kg
Fluorene	1300	51	ug/kg
2-Methylnaphthalene	3300	51	ug/kg
Phenanthrene	5400	51	ug/kg
Pyrene	3100	51	ug/kg
1-Methylnaphthalene	2000	51	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	66 DIL	(24 - 112)	
2-Fluorobiphenyl	64 DIL	(34 - 110)	
Terphenyl-d14	91 DIL	(41 - 119)	
Phenol-d5	72 DIL	(28 - 110)	
2-Fluorophenol	79 DIL	(26 - 110)	
2,4,6-Tribromophenol	76 DIL	(10 - 118)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

# Client Sample ID: HB-071(10-12')

## GC Volatiles

Lot-Sample #: A9L180492-002 Date Sampled: 12/17/09 11:49 Prep Date: 12/23/09 Prep Batch #: 9358085 Dilution Factor: 1		12/18/09	Matrix: SO
% Moisture: 13	Method:	SW846 8015	A MOD
PARAMETER Gasoline Range Organics (C6-C12)	RESULT 170	REPORTING LIMIT 120	UNITS ug/kg
SURROGATE Trifluorotoluene	PERCENT RECOVERY 92	RECOVERY LIMITS (10 - 150)	

NOTE(S):

# Client Sample ID: HB-071(10-12')

## GC Semivolatiles

	12/17/09 11:40 12/19/09 9353014	Work Order #: Date Received: Analysis Date:	12/18/09	Matrix: SO
% Moisture:	13	Method:	SW846 8015	В
PARAMETER C20-C34 C10-C20		RESULT 220 170	REPORTING LIMIT 31 23	UNITS mg/kg mg/kg
SURROGATE C9 (nonane)		PERCENT RECOVERY 23 DIL	RECOVERY LIMITS (10 - 110)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

## Client Sample ID: HB-071(10-12')

#### TOTAL Metals

Lot-Sample #...: A9L180492-002 Matrix.....: S0

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

**% Moisture....:** 13

PARAMETER	RESULT	REPORTIN LIMIT	NG <u>U</u> NITS	METHOD	PREPARATION- WORK ANALYSIS DATE ORDER #
Prep Batch #.	9354052				
Arsenic	11.3	1.2	mg/kg	SW846 6010B	12/21-12/23/09 LRCRC1AE
		Dilution Fac	tor: 1		
Cadmium	0.31	0.23	mg/kg	SW846 6010B	12/21-12/23/09 LRCRC1AF
		Dilution Fac	tor: 1		
Chromium	14.4	0.58	mg/kg	SW846 6010B	12/21-12/23/09 LRCRC1AG
		Dilution Fac			
Lead	65.9	0.35	mg/kg	SW846 6010B	12/21-12/23/09 LRCRC1AH
		Dilution Fac	COL! I		
NOTE(S):					

Client Sample ID: HB-071(10-12')

# General Chemistry

Lot-Sample #...: A9L180492-002 Work Order #...: LRCRC Matrix.....: S0

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

**% Moisture....:** 13

PARAMETER	RESULT	RL	UNITS	METHOI	)	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide	90.2	34.6 Dilution Factor	mg/kg	SW846	9030B/903 <b>4</b>	12/23/09	9357096
Corrosivity	10.1	Dilution Facto	No Units	SW846	9045A	12/19/09	9353120
Hexavalent Chromium	ND	0.92	mg/kg	SW846	7196A	12/23-12/24/09	9357236
		Dilution Facto	or: 1				
Percent Solids	86.8	10.0 Dilution Facto	<b>%</b> or: 1	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	2.1	0.58 Dilution Facto	mg/kg or: 1	MCAWW	335.2 CLP-M	12/23/09	9357388

NOTE (S):

# Client Sample ID: HB-071(16-18')

## GC/MS Volatiles

Lot-Sample #:	A9L180492-003	Work Order #:	LRCRE1AK	Matrix S0
Date Sampled:	12/17/09 11:40	Date Received:	12/18/09	
Prep Date:	12/23/09	Analysis Date:	12/23/09	
Prep Batch #:	9362414			
Dilution Factor:	1			
% Moisture:	12	${\tt Method:}$	SW846 8260A	

PARAMETER         RESULT         LIMIT         UNITS           Chloromethane         ND         11         ug/kg           Bromomethane         ND         11         ug/kg           Vinyl chloride         ND         2.3         ug/kg           Chloroethane         ND         11         ug/kg           Methylene chloride         7.3         5.7         ug/kg           Acetone         ND         23         ug/kg           Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         Chloroform         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
Bromomethane         ND         11         ug/kg           Vinyl chloride         ND         2.3         ug/kg           Chloroethane         ND         11         ug/kg           Methylene chloride         7.3         5.7         ug/kg           Acetone         ND         23         ug/kg           Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,1-Dichloroethane         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         ND         5.7         ug/kg           1,2-Dichloroethane         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
Vinyl chloride         ND         2.3         ug/kg           Chloroethane         ND         11         ug/kg           Methylene chloride         7.3         5.7         ug/kg           Acetone         ND         23         ug/kg           Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,1-Dichloroethane         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         ND         5.7         ug/kg           1,2-Dichloroethane         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
Chloroethane         ND         11         ug/kg           Methylene chloride         7.3         5.7         ug/kg           Acetone         ND         23         ug/kg           Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,1-Dichloroethane         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         ND         5.7         ug/kg           1,2-Dichloroethane         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
Methylene chloride         7.3         5.7         ug/kg           Acetone         ND         23         ug/kg           Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,1-Dichloroethane         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         Chloroform         ND         5.7         ug/kg           1,2-Dichloroethane         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
Acetone         ND         23         ug/kg           Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,1-Dichloroethane         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         ND         5.7         ug/kg           1,2-Dichloroethane         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
Carbon disulfide         12         5.7         ug/kg           1,1-Dichloroethene         ND         5.7         ug/kg           1,1-Dichloroethane         ND         5.7         ug/kg           1,2-Dichloroethene         ND         5.7         ug/kg           (total)         V         5.7         ug/kg           Chloroform         ND         5.7         ug/kg           1,2-Dichloroethane         ND         5.7         ug/kg           2-Butanone         ND         23         ug/kg
1,1-Dichloroethene       ND       5.7       ug/kg         1,1-Dichloroethane       ND       5.7       ug/kg         1,2-Dichloroethene       ND       5.7       ug/kg         (total)       ND       5.7       ug/kg         Chloroform       ND       5.7       ug/kg         1,2-Dichloroethane       ND       5.7       ug/kg         2-Butanone       ND       23       ug/kg
1,1-Dichloroethane       ND       5.7       ug/kg         1,2-Dichloroethene       ND       5.7       ug/kg         (total)       ND       5.7       ug/kg         1,2-Dichloroethane       ND       5.7       ug/kg         2-Butanone       ND       23       ug/kg
1,2-Dichloroethene (total)       ND       5.7       ug/kg         Chloroform       ND       5.7       ug/kg         1,2-Dichloroethane       ND       5.7       ug/kg         2-Butanone       ND       23       ug/kg
(total)         Chloroform       ND       5.7       ug/kg         1,2-Dichloroethane       ND       5.7       ug/kg         2-Butanone       ND       23       ug/kg
ChloroformND5.7ug/kg1,2-DichloroethaneND5.7ug/kg2-ButanoneND23ug/kg
1,2-Dichloroethane ND 5.7 ug/kg 2-Butanone ND 23 ug/kg
2-Butanone ND 23 ug/kg
1,1,1-Trichloroethane ND 5.7 ug/kg
Carbon tetrachloride ND 5.7 ug/kg
Bromodichloromethane ND 5.7 ug/kg
1,2-Dichloropropane ND 5.7 ug/kg
cis-1,3-Dichloropropene ND 5.7 ug/kg
Trichloroethene ND 5.7 ug/kg
Benzene ND 5.7 ug/kg
4-Methyl-2-pentanone ND 23 ug/kg
PERCENT RECOVERY
SURROGATE RECOVERY LIMITS
Dibromofluoromethane 98 (59 - 138)
1,2-Dichloroethane-d4 88 (61 - 130)
Toluene-d8 78 (60 - 143)
4-Bromofluorobenzene 56 (47 - 158)

NOTE(S):

Client Sample ID: HB-071(16-18')

# GC/MS Volatiles

Prep Batch #: 9362414		12/18/09	Matrix: SO
Dilution Factor: 5 % Moisture: 12	Method:	SW846 8260	A
		REPORTING	
PARAMETER	RESULT	LIMIT	<u>UNITS</u>
Dibromochloromethane	ND	28	ug/kg
1,1,2-Trichloroethane	ND	28	ug/kg
trans-1,3-Dichloropropene	ND	28	ug/kg
Bromoform	ND	28	ug/kg
2-Hexanone	ND	110	ug/kg
Tetrachloroethene	ND	28	ug/kg
1,1,2,2-Tetrachloroethane	ND	28	ug/kg
Toluene	ND	28	ug/kg
Chlorobenzene	ND	28	ug/kg
Ethylbenzene	ND	28	ug/kg
Styrene	ND	28	ug/kg
Xylenes (total)	ND	28	ug/kg

	PERCENT	RECOVERY
SURROGATE	RECOVERY	LIMITS
Dibromofluoromethane	97	(59 - 138)
1,2-Dichloroethane-d4	86	(61 - 130)
Toluene-d8	92	(60 - 143)
4-Bromofluorobenzene	80	(47 - 158)

NOTE(S):

# Client Sample ID: HB-071(16-18')

#### GC/MS Semivolatiles

Lot-Sample #: A9L180492-003 Date Sampled: 12/17/09 11:40 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 4		12/18/09	Matrix: SO
% Moisture: 12	Method:	SW846 8270	С
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Benzo(a)anthracene	810	30	ug/kg
Benzo(b) fluoranthene	1000	30	ug/kg
Benzo(k) fluoranthene	420	30	ug/kg
Benzo(a)pyrene	660	30	ug/kg
Chrysene	850	30	ug/kg
Dibenz(a,h)anthracene	180	30	ug/kg
Indeno(1,2,3-cd)pyrene	440	30	ug/kg
Naphthalene	3400	30	ug/kg
Acenaphthene	900	30	ug/kg
Acenaphthylene	120	30	ug/kg
Anthracene	490	30	ug/kg
Benzo(ghi)perylene	450	30	ug/kg
Fluoranthene	2200	30	ug/kg
Fluorene	870	30	ug/kg
2-Methylnaphthalene	1600	30	ug/kg
Phenanthrene	2700	30	ug/kg
Pyrene	1800	30	ug/kg
1-Methylnaphthalene	960	30	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	63 DIL	(24 - 112)	
2-Fluorobiphenyl	61 DIL	(34 - 110)	
Terphenyl-d14	85 DIL	(41 - 119)	
Phenol-d5	73 DIL	(28 - 110)	
2-Fluorophenol	74 DIL	(26 - 110)	
2,4,6-Tribromophenol	58 DIL	(10 - 118)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

Client Sample ID: HB-071(16-18')

# GC Volatiles

Lot-Sample #: A9L180492-00 Date Sampled: 12/17/09 11: Prep Date: 12/23/09 Prep Batch #: 9358085 Dilution Factor: 1		12/18/09	Matrix: SO
% Moisture: 12	Method:	SW846 8015	SA MOD
PARAMETER Gasoline Range Organics (C6-C12)	RESULT 150	REPORTING LIMIT 110	UNITS ug/kg
SURROGATE Trifluorotoluene	PERCENT RECOVERY 90	RECOVERY LIMITS (10 - 150)	-

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

Client Sample ID: HB-071(16-18')

# GC Semivolatiles

	12/17/09 11:40 12/19/09 9353014	Work Order #: Date Received: Analysis Date:	12/18/09	<b>Matrix</b> : SO
% Moisture:	12	Method:	SW846 8015	В
PARAMETER C20-C34 C10-C20		RESULT 150 72	REPORTING LIMIT 31 23	UNITS mg/kg mg/kg
SURROGATE C9 (nonane)		PERCENT RECOVERY 21 DIL	RECOVERY LIMITS (10 - 110)	

NOTE(S):

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Client Sample ID: HB-071(16-18')

# TOTAL Metals

Lot-Sample #...: A9L180492-003 Matrix.....: S0

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

**% Moisture....:** 12

PARAMETER	RESULT	REPORTIN	NG UNITS	METHOD	PREPARATION- WORK ANALYSIS DATE ORDER #
Prep Batch # Arsenic	: 9354052 <b>9.7</b>	1.1 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRE1AE
Cadmium	ND	0.23 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRE1AF
Chromium	22.9	0.57 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRE1AG
Lead	55.7	0.34 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRE1AH
NOTE(S):					

Client Sample ID: HB-071(16-18')

# General Chemistry

Lot-Sample #...: A9L180492-003 Work Order #...: LRCRE Matrix...... SO

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

**% Moisture....:** 12

PARAMETER	RESULT	RL	UNITS	<u>METHOI</u>	D	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide		<b>34.1</b> Dilution Facto	mg/kg or: 1	SW846	9030B/9034	12/23/09	9357096
Corrosivity	10.4	Dilution Facto	No Units	SW846	9045A	12/19/09	9353120
Hexavalent Chromium	ND	0.91	mg/kg	SW846	7196A	12/23-12/24/09	9357236
Percent Solids	87.9	<b>10.0</b> Dilution Facto	<b>%</b> er: 1	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	0.73	0.57 Dilution Facto	<b>mg/kg</b> r: 1	MCAWW	335.2 CLP-M	12/23/09	9357388

# NOTE(S):

RL Reporting Limit

# Client Sample ID: HB-072(2-4')

# GC/MS Volatiles

Lot-Sample #: A9L180492-004  Date Sampled: 12/17/09 12:20  Prep Date: 12/23/09  Prep Batch #: 9362414		12/18/09	Matrix: SO
Dilution Factor: 1 % Moisture: 11	Method:	SW846 8260	A
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Chloromethane	ND	11	ug/kg
Bromomethane	ND	11	ug/kg
Vinyl chloride	ND	2.3	ug/kg
Chloroethane	ND	11	ug/kg
Methylene chloride	11	5.6	ug/kg
Acetone	ND	23	ug/kg
Carbon disulfide	8.0	5.6	ug/kg
1,1-Dichloroethene	ND	5.6	ug/kg
1,1-Dichloroethane	ND	5.6	ug/kg
1,2-Dichloroethene (total)	ND	5.6	ug/kg
Chloroform	ND	5.6	ug/kg
1,2-Dichloroethane	ND	5.6	ug/kg
2-Butanone	ND	23	ug/kg
1,1,1-Trichloroethane	ND	5.6	ug/kg
Carbon tetrachloride	ND	5.6	ug/kg
Bromodichloromethane	ND	5.6	ug/kg
1,2-Dichloropropane	ND	5.6	ug/kg
cis-1,3-Dichloropropene	ND	5.6	ug/kg
Trichloroethene	ND	5.6	ug/kg
Benzene	ND	5.6	ug/kg
4-Methyl-2-pentanone	ND	23	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Dibromofluoromethane	102	(59 - 138)	
1,2-Dichloroethane-d4	90	(61 - 130)	

81

57

NOTE(S):

Toluene-d8

4-Bromofluorobenzene

Results and reporting limits have been adjusted for dry weight.

(60 - 143)

(47 - 158)

Client Sample ID: HB-072(2-4')

#### GC/MS Volatiles

Lot-Sample #: A9L180492-004	Work Order #: LRCRH2AK	Matrix SO
-----------------------------	------------------------	-----------

Date Sampled...: 12/17/09 12:20 Date Received..: 12/18/09
Prep Date....: 12/24/09 Analysis Date..: 12/24/09

Prep Batch #...: 9362414

Dilution Factor: 5

**% Moisture....:** 11 **Method.....:** SW846 8260A

		REPORTIN	IG
PARAMETER	RESULT	LIMIT	UNITS
Dibromochloromethane	ND	28	ug/kg
1,1,2-Trichloroethane	ND	28	ug/kg
trans-1,3-Dichloropropene	ND	28	ug/kg
Bromoform	ND	28	ug/kg
2-Hexanone	ND	110	ug/kg
Tetrachloroethene	ND	28	ug/kg
1, 1, 2, 2-Tetrachloroethane	ND	28	ug/kg
Toluene	ND	28	ug/kg
Chlorobenzene	ND	28	ug/kg
Ethylbenzene	ND	28	ug/kg
Styrene	ND	28	ug/kg
Xylenes (total)	ND	28	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
m'1 63 .1	0.0		

	PERCENT	RECOVERY
SURROGATE	RECOVERY	LIMITS
Dibromofluoromethane	98	(59 - 138)
1,2-Dichloroethane-d4	90	(61 - 130)
Toluene-d8	94	(60 - 143)
4-Bromofluorobenzene	79	(47 - 158)

NOTE(S):

Client Sample ID: HB-072(2-4')

# GC/MS Semivolatiles

Lot-Sample #: A9L180492-004 Date Sampled: 12/17/09 12:20 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 1		12/18/09	Matrix SO
% Moisture: 11	Method:	SW846 8270	С
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Benzo(a) anthracene	840	7.5	ug/kg
Benzo(b) fluoranthene	1100	7.5	ug/kg
Benzo(k) fluoranthene	390	7.5	ug/kg
Benzo(a)pyrene	630	7.5	ug/kg
Chrysene	840	7.5	ug/kg
Dibenz(a,h)anthracene	150	7.5	ug/kg
Indeno(1,2,3-cd)pyrene	460	7.5	ug/kg
Naphthalene	580	7.5	ug/kg
Acenaphthene	110	7.5	ug/kg
Acenaphthylene	140	7.5	ug/kg
Anthracene	360	7.5	ug/kg
Benzo(ghi)perylene	500	7.5	ug/kg
Fluoranthene	1500	7.5	ug/kg
Fluorene	140	7.5	ug/kg
2-Methylnaphthalene	780	7.5	ug/kg
Phenanthrene	1100	7.5	ug/kg
Pyrene	1400	7.5	ug/kg
1-Methylnaphthalene	520	7.5	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	58	(24 - 112)	
2-Fluorobiphenyl	64	(34 - 110)	
Terphenyl-d14	86	(41 - 119)	
Phenol-d5	70	(28 - 110)	
2-Fluorophenol	72	(26 - 110)	
2,4,6-Tribromophenol	51	(10 - 118)	

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

Client Sample ID: HB-072(2-4')

# GC Volatiles

		Work Order #: Date Received:		Matrix S0
Prep Date: Prep Batch #:		Analysis Date:	12/23/09	
Dilution Factor:	<del>-</del>			
% Moisture:	11	Method:	SW846 8015	A MOD
			REPORTING	
PARAMETER		RESULT	LIMIT	UNITS
Gasoline Range On (C6-C12)	rganics	ND	110	ug/kg
		PERCENT	RECOVERY	

LIMITS

(10 - 150)

RECOVERY

NOTE(S):

SURROGATE

Trifluorotoluene

Client Sample ID: HB-072(2-4')

# GC Semivolatiles

Lot-Sample #: A9L180492-004 Date Sampled: 12/17/09 12:2 Prep Date: 12/19/09 Prep Batch #: 9353014 Dilution Factor: 10		12/18/09	Matrix: SO
% Moisture: 11	Method:	SW846 8015	В
PARAMETER C20-C34 C10-C20	RESULT 140 44	REPORTING LIMIT 30 23	UNITS mg/kg mg/kg
SURROGATE C9 (nonane)	PERCENT RECOVERY 24 DIL	RECOVERY LIMITS (10 - 110)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

# Client Sample ID: HB-072(2-4')

#### TOTAL Metals

Lot-Sample #...: A9L180492-004 Matrix.....: S0

Date Sampled...: 12/17/09 12:20 Date Received..: 12/18/09

**% Moisture....:** 11

PARAMETER	RESULT	REPORTIN LIMIT	G UNITS	METHOD	PREPARATION- ANALYSIS DATE	WORK ORDER #
Prep Batch #. Arsenic	.: 9354052 <b>8.6</b>	1.1 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09	LRCRH1AE
Cadmium	ND	0.23 Dilution Fact	mg/kg tor: 1	SW846 6010B	12/21-12/23/09	LRCRH1AF
Chromium	10.6	0.56 Dilution Fact	mg/kg	SW846 6010B	12/21-12/23/09	LRCRH1AG
Lead	27.0	0.34 Dilution Face	mg/kg tor: 1	SW846 6010B	12/21-12/23/09	LRCRH1AH
NOTE(S):						

Client Sample ID: HB-072(2-4')

# General Chemistry

Lot-Sample #...: A9L180492-004 Work Order #...: LRCRH Matrix.....: S0

Date Sampled...: 12/17/09 12:20 Date Received..: 12/18/09

**% Moisture....:** 11

PARAMETER	RESULT	RL	UNITS	<u>METHOI</u>	<u> </u>	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide	287	33.8 Dilution Factor	<b>mg/kg</b> or: 1	SW846	9030B/9034	12/23/09	9357096
Corrosivity	10.2	Dilution Facto	No Units	SW846	9045A	12/19/09	9353120
Hexavalent Chromium	ND	0.90	mg/kg	SW846	7196A	12/23-12/24/09	9357236
Percent Solids	88.7	10.0 Dilution Facto	ફ	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	2.9	0.56 Dilution Facto	mg/kg	MCAWW	335.2 CLP-M	12/28/09	9362384

# NOTE(S):

RL. Reporting Limit

# Client Sample ID: HB-072(4-6')

# GC/MS Volatiles

Lot-Sample #: A9L180492-005 Date Sampled: 12/17/09 12:20 Prep Date: 12/23/09 Prep Batch #: 9362414 Dilution Factor: 1		12/18/09	Matrix: SO
<b>% Moisture:</b> 13	Method:	SW846 8260	A
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Chloromethane	ND	12	ug/kg
Bromomethane	ND	12	ug/kg
Vinyl chloride	ND	2.3	ug/kg
Chloroethane	ND	12	ug/kg
Methylene chloride	ND	5.8	ug/kg
Acetone	ND	23	ug/kg
Carbon disulfide	27	5.8	ug/kg
1,1-Dichloroethene	ND	5.8	ug/kg
1,1-Dichloroethane	ND	5.8	ug/kg
1,2-Dichloroethene	ND	5.8	ug/kg
(total)			
Chloroform	ND	5.8	ug/kg
1,2-Dichloroethane	ND	5.8	ug/kg
2-Butanone	ND	23	ug/kg
1,1,1-Trichloroethane	ND	5.8	ug/kg
Carbon tetrachloride	ND	5.8	ug/kg
Bromodichloromethane	ND	5.8	ug/kg
1,2-Dichloropropane	ND	5.8	ug/kg
cis-1,3-Dichloropropene	ND	5.8	ug/kg
Trichloroethene	ND	5.8	ug/kg
Dibromochloromethane	ND	5.8	ug/kg
1,1,2-Trichloroethane	ND	5.8	ug/kg
Benzene	27	5.8	ug/kg
trans-1,3-Dichloropropene	ND	5.8	ug/kg
Bromoform	ND	5.8	ug/kg
4-Methyl-2-pentanone	ND	23	ug/kg
2-Hexanone	ND	23	ug/kg
Tetrachloroethene	ND	5.8	ug/kg
1,1,2,2-Tetrachloroethane	ND	5.8	ug/kg
Toluene	44	5.8	ug/kg
Chlorobenzene	ND	5.8	ug/kg
Ethylbenzene	ND	5.8	ug/kg
Styrene	ND	5.8	ug/kg
Xylenes (total)	20	5.8	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Dibromofluoromethane	91	(59 - 138)	
1,2-Dichloroethane-d4	81	(61 - 130)	
Toluene-d8	85	(60 - 143)	

(Continued on next page)

74

4-Bromofluorobenzene

(47 - 158)

Client Sample ID: HB-072(4-6')

GC/MS Volatiles

Lot-Sample #...: A9L180492-005 Work Order #...: LRCRK1AK Matrix...... S0

NOTE(S):

Client Sample ID: HB-072(4-6')

# GC/MS Semivolatiles

Lot-Sample #: A9L180492-005 Date Sampled: 12/17/09 12:20 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 2		12/18/09	Matrix S0
<b>% Moisture:</b> 13	Method:	SW846 8270	С
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Benzo(a) anthracene	1300	15	ug/kg
Benzo(b) fluoranthene	2100	15	ug/kg
Benzo(k) fluoranthene	750	15	ug/kg
Benzo(a)pyrene	1200	15	ug/kg
Chrysene	1500	15	ug/kg
Dibenz(a,h)anthracene	260	15	ug/kg
Indeno(1,2,3-cd)pyrene	760	15	ug/kg
Naphthalene	910	15	ug/kg
Acenaphthene	140	15	ug/kg
Acenaphthylene	340	15	ug/kg
Anthracene	410	15	ug/kg
Benzo(ghi)perylene	850	15	ug/kg
Fluoranthene	2100	15	ug/kg
Fluorene	180	15	ug/kg
2-Methylnaphthalene	1400	15	ug/kg
Phenanthrene	1400	15	ug/kg
Pyrene	2100	15	ug/kg
1-Methylnaphthalene	860	15	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	60 DIL	(24 - 112)	
2-Fluorobiphenyl	65 DIL	(34 - 110)	
Terphenyl-d14	82 DIL	(41 - 119)	
Phenol-d5	73 DIL	(28 - 110)	
2-Fluorophenol	70 DIL	(26 - 110)	
2,4,6-Tribromophenol	60 DIL	(10 - 118)	

 $\label{eq:def:DIL} \textbf{DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.}$ 

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

Client Sample ID: HB-072(4-6')

#### GC Volatiles

Lot-Sample #: A9L180492-005	Work Order #: LF	RCRK1AJ <b>Matrix</b> SO
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Date Sampled...: 12/17/09 12:20 Date Received..: 12/18/09 Analysis Date..: 12/23/09 Prep Date....: 12/23/09

Prep Batch #...: 9358085

Dilution Factor: 5

**% Moisture....:** 13 Method..... SW846 8015A MOD

REPORTING

RESULT PARAMETER LIMIT UNITS Gasoline Range Organics 700 580 ug/kg

(C6-C12)

PERCENT RECOVERY RECOVERY LIMITS

SURROGATE Trifluorotoluene 92 (10 - 150)

# NOTE(S):

Client Sample ID: HB-072(4-6')

# GC Semivolatiles

		Work Order #:		Matrix SO
Date Sampled:	12/17/09 12:20	Date Received:	12/18/09	
Prep Date:	12/19/09	Analysis Date:	12/23/09	
Prep Batch #:	9353014			
Dilution Factor:	10			
% Moisture:	13	Method:	SW846 8015	В
			REPORTING	
PARAMETER		RESULT	LIMIT	UNITS
C20-C34		220	31	mg/kg
C10-C20		82	23	mg/kg
		PERCENT	RECOVERY	
SURROGATE		RECOVERY	LIMITS	
C9 (nonane)		26 DIL	(10 - 110)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

# Client Sample ID: HB-072(4-6')

#### TOTAL Metals

Lot-Sample #...: A9L180492-005 Matrix.....: S0

Date Sampled...: 12/17/09 12:20 Date Received..: 12/18/09

**% Moisture....:** 13

PARAMETER	RESULT	REPORTIN LIMIT	IG <u>UNITS</u>	METHOD	PREPARATION- WORK ANALYSIS DATE ORDER #
Prep Batch #. Arsenic	: 9354052 <b>9.3</b>	1.2 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRK1AE
Cadmium	ND	0.23 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09 LRCRK1AF
Chromium	19.7	0.58 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRK1AG
Lead	38.4	0.35 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRK1AH
NOTE(S):	·				

Client Sample ID: HB-072(4-6')

# General Chemistry

Lot-Sample #...: A9L180492-005 Work Order #...: LRCRK Matrix.....: SO

Date Sampled...: 12/17/09 12:20 Date Received..: 12/18/09

**% Moisture....:** 13

PARAMETER	RESULT	RL	UNITS	METHO:	D	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide		34.6 lution Facto	mg/kg	SW846	9030B/9034	12/23/09	9357096
Corrosivity	10.2	lution Fact	No Units	SW846	9045A	12/19/09	9353120
Hexavalent Chromium	2.5	0.92	mg/kg	SW846	7196A	12/23-12/24/09	9357236
	D1.	lution Fact	or: 1				
Percent Solids	86.7	10.0 lution Facto	<b>%</b> or: 1	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	2.1	0.58 lution Facto	mg/kg	MCAWW	335.2 CLP-M	12/28/09	9362384

# NOTE(S):

RL Reporting Limit

# Client Sample ID: HB-073(0-2')

#### GC/MS Volatiles

Lot-Sample #: A9L180492-006	Work Order #: LRCRM1AK	Matrix SO
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Date Sampled...: 12/17/09 12:50 Date Received..: 12/18/09 Prep Date....: 12/23/09 Analysis Date..: 12/23/09

Prep Batch #...: 9362414

Dilution Factor: 1

**% Moisture....:** 12 **Method.....:** SW846 8260A

		DEDODETN	<b>.</b>
PARAMETER	RESULT	REPORTING LIMIT	UNITS
Chloromethane	ND	11	ug/kg
Bromomethane	ND	11	ug/kg
Vinyl chloride	ND	2.3	ug/kg
Chloroethane	ND	11	ug/kg
Methylene chloride	ND	5.7	ug/kg
Acetone	ND	23	ug/kg
Carbon disulfide	ND	5.7	ug/kg
1,1-Dichloroethene	ND	5.7	ug/kg
1,1-Dichloroethane	ND	5.7	ug/kg
1,2-Dichloroethene	ND	5.7	ug/kg
(total)			
Chloroform	ND	5.7	ug/kg
1,2-Dichloroethane	ND	5.7	ug/kg
2-Butanone	ND	23	ug/kg
1,1,1-Trichloroethane	ND	5.7	ug/kg
Carbon tetrachloride	ND	5.7	ug/kg
Bromodichloromethane	ND	5.7	ug/kg
1,2-Dichloropropane	ND	5.7	ug/kg
cis-1,3-Dichloropropene	ND	5.7	ug/kg
Trichloroethene	ND	5.7	ug/kg
Dibromochloromethane	ND	5.7	ug/kg
1,1,2-Trichloroethane	ND	5.7	ug/kg
Benzene	ND	5.7	ug/kg
trans-1,3-Dichloropropene	ND	5.7	ug/kg
Bromoform	ND	5.7	ug/kg
4-Methyl-2-pentanone	ND	23	ug/kg
2-Hexanone	ND	23	ug/kg
Tetrachloroethene	ND	5.7	ug/kg
1,1,2,2-Tetrachloroethane	ND	5.7	ug/kg
Toluene	5.8	5 <b>.7</b>	ug/kg
Chlorobenzene	ND	5.7	ug/kg
Ethylbenzene	11	5.7	ug/kg
Styrene	ND	5.7	ug/kg
Xylenes (total)	47	5.7	ug/kg
	PERCENT	RECOVERY	

	PERCENT	RECOVERY
SURROGATE	RECOVERY	LIMITS
Dibromofluoromethane	91	(59 - 138)
1,2-Dichloroethane-d4	85	(61 - 130)
Toluene-d8	90	(60 - 143)
4-Bromofluorobenzene	71	(47 - 158)

(Continued on next page)

Client Sample ID: HB-073(0-2')

# GC/MS Volatiles

Lot-Sample #: A9L180492-006	Work Order #: LRCRM1AK	Matrix: SO
NOTE(S):		

Client Sample ID: HB-073(0-2')

# GC/MS Semivolatiles

Lot-Sample #: A9L180492-006 Date Sampled: 12/17/09 12:50 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 2		12/18/09	Matrix S0
% Moisture: 12	Method:	SW846 8270	С
PARAMETER	RESULT	REPORTING LIMIT	UNITS
Benzo(a)anthracene	1100	15	ug/kg
Benzo(b) fluoranthene	1500	15	ug/kg
Benzo(k) fluoranthene	420	15	ug/kg
Benzo(a)pyrene	1000	15	ug/kg
Chrysene	960	15	ug/kg
Dibenz(a,h)anthracene	290	15	ug/kg
Indeno(1,2,3-cd)pyrene	600	15	ug/kg
Naphthalene	880	15	ug/kg
Acenaphthene	82	15	ug/kg
Acenaphthylene	150	15	ug/kg
Anthracene	240	15	ug/kg
Benzo(ghi)perylene	820	15	ug/kg
Fluoranthene	1600	15	ug/kg
Fluorene	87	15	ug/kg
2-Methylnaphthalene	1500	15	ug/kg
Phenanthrene	1200	15	ug/kg
Pyrene	1500	15	ug/kg
1-Methylnaphthalene	1000	15	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	68 DIL	(24 - 112)	
2-Fluorobiphenyl	66 DIL	(34 - 110)	
Terphenyl-d14	79 DIL	(41 - 119)	
Phenol-d5	67 DIL	(28 - 110)	
2-Fluorophenol	74 DIL	(26 - 110)	
2,4,6-Tribromophenol	48 DIL	(10 - 118)	

NOTE(S):

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Client Sample ID: HB-073(0-2')

# GC Volatiles

Lot-Sample #: A9L180492-006 Date Sampled: 12/17/09 12:50 Prep Date: 12/28/09 Prep Batch #: 9363092		12/18/09	<b>Matrix</b> : SO
Dilution Factor: 1 % Moisture: 12	Method:	SW846 8015	A MOD
PARAMETER	RESULT	REPORTING LIMIT	UNITS
Gasoline Range Organics (C6-C12)	210	110	ug/kg
SURROGATE	PERCENT RECOVERY	RECOVERY LIMITS	

(10 - 150)

99

NOTE(S):

Trifluorotoluene

Client Sample ID: HB-073(0-2')

# GC Semivolatiles

Lot-Sample #: Date Sampled: Prep Date: Prep Batch #: Dilution Factor:	12/17/09 12:50 12/19/09 9353014		12/18/09	Matrix: SO
% Moisture:	12	Method:	SW846 8015	В
PARAMETER C20-C34 C10-C20		RESULT 130 120	REPORTING LIMIT 31 23	UNITS mg/kg mg/kg
SURROGATE C9 (nonane)		PERCENT RECOVERY 34 DIL	RECOVERY LIMITS (10 - 110)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

# Client Sample ID: HB-073(0-2')

#### TOTAL Metals

Lot-Sample #...: A9L180492-006 Matrix.....: SO

Date Sampled...: 12/17/09 12:50 Date Received..: 12/18/09

**% Moisture....:** 12

PARAMETER	RESULT	REPORTIN	IG <u>UNITS</u>	METHOD	PREPARATION- WORK ANALYSIS DATE ORDER #
Prep Batch #	: 9354052				
Arsenic	9.3	1.1 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRM1AE
Cadmium	0.33	0.23 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRM1AF
Chromium	10.6	<b>0.57</b> Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRM1AG
Lead	115	0.34 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRM1AH
NOTE(S):					

Client Sample ID: HB-073(0-2')

# General Chemistry

Lot-Sample #...: A9L180492-006 Work Order #...: LRCRM Matrix.....: S0

Date Sampled...: 12/17/09 12:50 Date Received..: 12/18/09

**% Moisture....:** 12

PARAMETER	RESULT	RL	UNITS	METHOI	)	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide	302	33.9 Dilution Factor	mg/kg or: 1	SW846	9030B/903 <b>4</b>	12/23/09	9357096
Corrosivity	11.0	Dilution Facto	No Units	SW846	9045A	12/19/09	9353120
Hexavalent Chromium	ND	0.90	mg/kg	SW846	7196A	12/23-12/24/09	9357236
		Dilution Facto	or: 1				
Percent Solids	88.4	10.0 Dilution Facto	% or: 1	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	4.6	0.57 Dilution Facto	mg/kg er: 1	MCAWW	335.2 CLP-M	12/28/09	9362384
**************************************							

# NOTE(S):

RL Reporting Limit

Client Sample ID: HB-075(0-2')

#### GC/MS Volatiles

Lot-Sample #:	A9L180492-007	Work Order #: LRCRP1AK	<b>Matrix:</b> SO
Date Sampled :	12/17/09 13:20	Date Received . : 12/18/09	

Dilution Factor: 1

**% Moisture....:** 12 **Method.....:** SW846 8260A

REPORTING PARAMETER RESULT LIMIT UNITS	
Chloromethane ND 11 ug/kg	
Bromomethane ND 11 ug/kg	
Vinyl chloride ND 2.3 ug/kg	
Chloroethane ND 11 ug/kg	
Methylene chloride ND 5.7 ug/kg	
Acetone ND 23 ug/kg	
Carbon disulfide ND 5.7 ug/kg	
1,1-Dichloroethene ND 5.7 ug/kg	
1,1-Dichloroethane ND 5.7 ug/kg	
1,2-Dichloroethene ND 5.7 ug/kg	
(total)	
Chloroform ND 5.7 ug/kg	
1,2-Dichloroethane ND 5.7 ug/kg	
2-Butanone ND 23 ug/kg	
1,1,1-Trichloroethane ND 5.7 ug/kg	
Carbon tetrachloride ND 5.7 ug/kg	
Bromodichloromethane ND 5.7 ug/kg	
1,2-Dichloropropane ND 5.7 ug/kg	
cis-1,3-Dichloropropene ND 5.7 ug/kg	
Trichloroethene ND 5.7 ug/kg	
Dibromochloromethane ND 5.7 ug/kg	
1,1,2-Trichloroethane ND 5.7 ug/kg	
Benzene ND 5.7 ug/kg	
trans-1,3-Dichloropropene ND 5.7 ug/kg	
Bromoform ND 5.7 ug/kg	
4-Methyl-2-pentanone ND 23 ug/kg	
2-Hexanone ND 23 ug/kg	
Tetrachloroethene ND 5.7 ug/kg	
1,1,2,2-Tetrachloroethane ND 5.7 ug/kg	
Toluene ND 5.7 ug/kg	
Chlorobenzene ND 5.7 ug/kg	
Ethylbenzene ND 5.7 ug/kg	
Styrene ND 5.7 ug/kg	
Xylenes (total) 26 5.7 ug/kg	

	PERCENT	RECOVERY
SURROGATE	RECOVERY	LIMITS
Dibromofluoromethane	89	(59 - 138)
1,2-Dichloroethane-d4	83	(61 - 130)
Toluene-d8	102	(60 - 143)
4-Bromofluorobenzene	118	(47 - 158)

(Continued on next page)

Client Sample ID: HB-075(0-2')

GC/MS Volatiles

Lot-Sample #...: A9L180492-007 Work Order #...: LRCRP1AK Matrix...... S0

NOTE(S):

Client Sample ID: HB-075(0-2')

#### GC/MS Semivolatiles

Lot-Sample #: A9L180492-007 Date Sampled: 12/17/09 13:20 Prep Date: 12/19/09 Prep Batch #: 9353017 Dilution Factor: 10		12/18/09	Matrix: SO
% Moisture: 12	Method:	SW846 8270	С
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Benzo(a) anthracene	2900	76	ug/kg
Benzo(b) fluoranthene	2500	76	ug/kg
Benzo(k) fluoranthene	940	76	ug/kg
Benzo(a)pyrene	2100	76	ug/kg
Chrysene	2600	76	ug/kg
Dibenz(a,h)anthracene	460	76	ug/kg
Indeno(1,2,3-cd)pyrene	1100	76	ug/kg
Naphthalene	230	76	ug/kg
Acenaphthene	190	76	ug/kg
Acenaphthylene	100	76	ug/kg
Anthracene	860	76	ug/kg
Benzo(ghi)perylene	1200	76	ug/kg
Fluoranthene	5100	76	ug/kg
Fluorene	210	76	ug/kg
2-Methylnaphthalene	180	76	ug/kg
Phenanthrene	2500	76	ug/kg
Pyrene	6100	76	ug/kg
1-Methylnaphthalene	150	76	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Nitrobenzene-d5	99 DIL	(24 - 112)	
2-Fluorobiphenyl	70 DIL	(34 - 110)	
Terphenyl-d14	110 DIL	(41 - 119)	
Phenol-d5	73 DIL	(28 - 110)	
2-Fluorophenol	90 DIL	(26 - 110)	
2,4,6-Tribromophenol	64 DIL	(10 - 118)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

Client Sample ID: HB-075(0-2')

#### GC Volatiles

Lot-Sample #:	A9L180492-007	Work Order #:	LRCRP1AJ	Matrix SO
Date Sampled:	12/17/09 13:20	Date Received:	12/18/09	

Prep Date....: 12/23/09 Analysis Date..: 12/24/09

Prep Batch #...: 9358085

Dilution Factor: 5

**% Moisture....:** 12 **Method.....:** SW846 8015A MOD

REPORTING

PARAMETER RESULT LIMIT UNITS
Gasoline Range Organics 780 570 ug/kg

(C6-C12)

PERCENT RECOVERY

SURROGATE RECOVERY LIMITS

Trifluorotoluene 99 (10 - 150)

NOTE(S):

Client Sample ID: HB-075(0-2')

#### GC Semivolatiles

Lot-Sample #:	A9L180492-007	Work Order #:	LRCRP1AC	Matrix SO
Date Sampled:	12/17/09 13:20	Date Received:	12/18/09	
Prep Date:	12/19/09	Analysis Date:	12/23/09	
Prep Batch #:	9353014		•	•
Dilution Factor:	50			
% Moisture:	12	Method:	SW846 8015	В
			REPORTING	
PARAMETER		RESULT	LIMIT	UNITS
C20-C34		610	150	mg/kg
C10-C20		100 J	110	mg/kg
		PERCENT	RECOVERY	
SURROGATE		RECOVERY	LIMITS	
C9 (nonane)		35 DIL	(10 - 110)	

NOTE(S):

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

J Estimated result. Result is less than RL.

# Client Sample ID: HB-075(0-2')

#### TOTAL Metals

Lot-Sample #...: A9L180492-007 Matrix.....: S0

Date Sampled...: 12/17/09 13:20 Date Received..: 12/18/09

**% Moisture....:** 12

PARAMETER	RESULT	REPORTIN LIMIT	G <u>UNITS</u>	METHOD	PREPARATION- WORK ANALYSIS DATE ORDER #
Prep Batch # Arsenic	: 9354052 <b>4.7</b>	1.1 Dilution Fac	<b>mg/kg</b> tor: 1	SW846 6010B	12/21-12/23/09 LRCRP1AE
Cadmium	0.23	0.23 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09 LRCRP1AF
Chromium	18.6	0.57 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09 LRCRP1AG
Lead	21.3	0.34 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRP1AH
NOTE(S):				· · · · · · · · · · · · · · · · · · ·	

Client Sample ID: HB-075(0-2')

# General Chemistry

Lot-Sample #...: A9L180492-007 Work Order #...: LRCRP Matrix.....: S0

Date Sampled...: 12/17/09 13:20 Date Received..: 12/18/09

**% Moisture....:** 12

PARAMETER	RESULT	RL	UNITS	METHO:	D	PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide		<b>34.1</b> Dilution Factor	mg/kg	SW846	9030B/9034	12/23/09	9357096
Corrosivity	<b>11.5</b>	Dilution Fact	No Units	SW846	9045A	12/19/09	9353120
Hexavalent Chromium	ND	0.91	mg/kg	SW846	7196A	12/23-12/24/09	9357236
	Ε	Dilution Fact	or: 1				
Percent Solids	87.9	10.0 Dilution Facto	<b>%</b> or: 1	MCAWW	160.3 MOD	12/22-12/23/09	9356363
Total Cyanide	ND E	0.57 Dilution Facto	mg/kg or: 1	MCAWW	335.2 CLP-M	12/28/09	9362384

NOTE (S):

RL Reporting Limit

# Client Sample ID: HB-075(2-4')

# GC/MS Volatiles

Date Sampled: 12/17/09 13:20 Prep Date: 12/21/09 Prep Batch #: 9356203	Work Order #: Date Received: Analysis Date:	12/18/09	Matrix S
Dilution Factor: 1 % Moisture: 9.4	Method:	SW846 8260	A
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
Chloromethane	ND	11	ug/kg
Bromomethane	ND	11	ug/kg
Vinyl chloride	ND	2.2	ug/kg
Chloroethane	ND	11	ug/kg
Methylene chloride	ND	5.5	ug/kg
Acetone	ND	22	ug/kg
Carbon disulfide	ND	5.5	ug/kg
1,1-Dichloroethene	ND	5.5	ug/kg
1,1-Dichloroethane	ND	5.5	ug/kg
1,2-Dichloroethene (total)	ND	5.5	ug/kg
Chloroform	ND	5.5	ug/kg
1,2-Dichloroethane	ND	5.5	ug/kg
2-Butanone	ND	22	ug/kg
1,1,1-Trichloroethane	ND	5.5	ug/kg
Carbon tetrachloride	ND	5.5	ug/kg
Bromodichloromethane	ND	5.5	ug/kg
1,2-Dichloropropane	ND	5.5	ug/kg
cis-1,3-Dichloropropene	ND	5.5	ug/kg
Trichloroethene	ND .	5.5	ug/kg
Dibromochloromethane	ND	5.5	ug/kg
1,1,2-Trichloroethane	ND	5.5	ug/kg
Benzene	ND	5.5	ug/kg
trans-1,3-Dichloropropene	ND	5.5	ug/kg
Bromoform	ND	5.5	ug/kg
4-Methyl-2-pentanone	ND	22	ug/kg
2-Hexanone	ND	22	ug/kg
Tetrachloroethene	ND	5.5	ug/kg
1,1,2,2-Tetrachloroethane	ND	5.5	ug/kg
Toluene	ND	5.5	ug/kg
Chlorobenzene	ND	5.5	ug/kg
Ethylbenzene	ND	5.5	ug/kg
Styrene	ND	5.5	ug/kg
Xylenes (total)	ND	5.5	ug/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	LIMITS	
Dibromofluoromethane	102	(59 - 138)	
1,2-Dichloroethane-d4	78	(61 - 130)	
Toluene-d8	103	(60 - 143)	

(Continued on next page)

Client Sample ID: HB-075(2-4')

#### GC/MS Volatiles

Lot-Sample #...: A9L180492-008 Work Order #...: LRCRR1AK Matrix.....: S0

NOTE(S):

Client Sample ID: HB-075(2-4')

# GC/MS Semivolatiles

Matrix..... S0

Lot-Sample #...: A9L180492-008 Work Order #...: LRCRR1AD

Prep Date:         12/19/09         Analysis Date         12/24/09           Prep Batch #:         9353017         Moisture         4           & Moisture:         9.4         Method         SW846 8270C           REPORTING           REPORTING           PARAMETER         RESULT         LIMIT         UNITS           Benzo(a) anthracene         2900         29         ug/kg           Benzo(b) fluoranthene         1200         29         ug/kg           Benzo(a) pyrene         2500         29         ug/kg           Benzo(a) pyrene         2600         29         ug/kg           Chrysene         2600         29         ug/kg           Dibenz (a, h) anthracene         500         29         ug/kg           Indeno (1,2,3-cd) pyrene         1500         29         ug/kg           Indeno (1,2,3-cd) pyrene         1500         29         ug/kg           Acenaphthene         570         29         ug/kg           Acenaphthylene         72         29         ug/kg           Anthracene         1300         29         ug/kg           Benzo (ghi)perylene         1600         29	Date Sampled: 12/17/09 13:20	Date Received:	12/18/09	
Prep Batch #: 9353017   Dilution Factor: 4   % Moisture: 9.4   Method	<del>-</del>			
Dilution Factor: 4	<del></del>	=		
PARAMETER				
DARAMETER   RESULT   LIMIT   UNITS	<b>% Moisture:</b> 9.4	Method:	SW846 8270	С
DARAMETER   RESULT   LIMIT   UNITS				
Benzo (a) anthracene         2900         29         ug/kg           Benzo (b) fluoranthene         3200         29         ug/kg           Benzo (k) fluoranthene         1200         29         ug/kg           Benzo (a) pyrene         2500         29         ug/kg           Chrysene         2600         29         ug/kg           Dibenz (a, h) anthracene         500         29         ug/kg           Dibenz (a, h) anthracene         500         29         ug/kg           Indeno (1, 2, 3-cd) pyrene         1500         29         ug/kg           Indeno (1, 2, 3-cd) pyrene         1500         29         ug/kg           Naphthalene         820         29         ug/kg           Acenaphthalene         570         29         ug/kg           Acenaphthylene         72         29         ug/kg           Acenaphthylene         72         29         ug/kg           Benzo (ghi) perylene         1600         29         ug/kg           Fluoranthene         570         29         ug/kg           Fluorene         570         29         ug/kg           Phenanthrene         4800         29         ug/kg           Percen			REPORTING	
Benzo (b) fluoranthene         3200         29         ug/kg           Benzo (k) fluoranthene         1200         29         ug/kg           Benzo (a) pyrene         2500         29         ug/kg           Chrysene         2600         29         ug/kg           Dibenz (a, h) anthracene         500         29         ug/kg           Indeno (1, 2, 3-cd) pyrene         1500         29         ug/kg           Indeno (1, 2, 3-cd) pyrene         1500         29         ug/kg           Naphthalene         820         29         ug/kg           Acenaphthylene         570         29         ug/kg           Acenaphthylene         72         29         ug/kg           Anthracene         1300         29         ug/kg           Benzo (ghi) perylene         1600         29         ug/kg           Fluoranthene         5500         29         ug/kg           Fluorene         570         29         ug/kg           Phenanthrene         4800         29         ug/kg           Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           SURROGATE <td< td=""><td>PARAMETER</td><td>RESULT</td><td><u>LIMIT</u></td><td><u>UNITS</u></td></td<>	PARAMETER	RESULT	<u>LIMIT</u>	<u>UNITS</u>
Benzo(k) fluoranthene       1200       29       ug/kg         Benzo(a) pyrene       2500       29       ug/kg         Chrysene       2600       29       ug/kg         Dibenz(a,h) anthracene       500       29       ug/kg         Indeno(1,2,3-cd) pyrene       1500       29       ug/kg         Naphthalene       820       29       ug/kg         Acenaphthylene       570       29       ug/kg         Acenaphthylene       72       29       ug/kg         Anthracene       1300       29       ug/kg         Benzo(ghi) perylene       1600       29       ug/kg         Fluoranthene       5500       29       ug/kg         Fluorene       570       29       ug/kg         2-Methylnaphthalene       1000       29       ug/kg         Phenanthrene       4800       29       ug/kg         Pyrene       5400       29       ug/kg         1-Methylnaphthalene       610       29       ug/kg         Nitrobenzene-d5       68       DIL       (24 - 112)         2-Fluorobiphenyl       56       DIL       (34 - 110)         Terphenyl-d14       87       DIL	Benzo(a)anthracene	2900	29	ug/kg
Benzo(a)pyrene       2500       29       ug/kg         Chrysene       2600       29       ug/kg         Dibenz(a,h)anthracene       500       29       ug/kg         Indeno(1,2,3-cd)pyrene       1500       29       ug/kg         Naphthalene       820       29       ug/kg         Acenaphthene       570       29       ug/kg         Acenaphthylene       72       29       ug/kg         Anthracene       1300       29       ug/kg         Benzo(ghi)perylene       1600       29       ug/kg         Fluoranthene       5500       29       ug/kg         Fluorene       570       29       ug/kg         2-Methylnaphthalene       1000       29       ug/kg         Pyrene       5400       29       ug/kg         1-Methylnaphthalene       610       29       ug/kg         Nitrobenzene-d5       68       DIL       (24 - 112)         2-Fluorobiphenyl       56       DIL       (34 - 110)         Terphenyl-d14       87       DIL       (41 - 119)         Phenol-d5       70       DIL       (28 - 110)         2-Fluorophenol       72       DIL	Benzo(b)fluoranthene	3200	29	ug/kg
Chrysene       2600       29       ug/kg         Dibenz(a,h)anthracene       500       29       ug/kg         Indeno(1,2,3-cd)pyrene       1500       29       ug/kg         Naphthalene       820       29       ug/kg         Acenaphthene       570       29       ug/kg         Acenaphthylene       72       29       ug/kg         Anthracene       1300       29       ug/kg         Benzo(ghi)perylene       1600       29       ug/kg         Fluoranthene       5500       29       ug/kg         Fluorene       570       29       ug/kg         2-Methylnaphthalene       1000       29       ug/kg         Phenanthrene       4800       29       ug/kg         Pyrene       5400       29       ug/kg         1-Methylnaphthalene       610       29       ug/kg         Nitrobenzene-d5       68       DIL       (24 - 112)         2-Fluorobiphenyl       56       DIL       (34 - 110)         Terphenyl-d14       87       DIL       (41 - 119)         Phenol-d5       70       DIL       (28 - 110)         2-Fluorophenol       72       DIL       (2	Benzo(k)fluoranthene	1200	29	ug/kg
Dibenz (a, h) anthracene         500         29         ug/kg           Indeno (1, 2, 3-cd) pyrene         1500         29         ug/kg           Naphthalene         820         29         ug/kg           Acenaphthene         570         29         ug/kg           Acenaphthylene         72         29         ug/kg           Anthracene         1300         29         ug/kg           Benzo (ghi) perylene         1600         29         ug/kg           Fluoranthene         5500         29         ug/kg           Fluorene         570         29         ug/kg           2-Methylnaphthalene         1000         29         ug/kg           Phenanthrene         4800         29         ug/kg           Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	Benzo(a)pyrene	2500	29	ug/kg
Indeno(1,2,3-cd)pyrene         1500         29         ug/kg           Naphthalene         820         29         ug/kg           Acenaphthene         570         29         ug/kg           Acenaphthylene         72         29         ug/kg           Anthracene         1300         29         ug/kg           Benzo(ghi)perylene         1600         29         ug/kg           Fluoranthene         5500         29         ug/kg           Fluorene         570         29         ug/kg           2-Methylnaphthalene         1000         29         ug/kg           Phenanthrene         4800         29         ug/kg           Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	Chrysene	2600	29	ug/kg
Naphthalene         820         29         ug/kg           Acenaphthene         570         29         ug/kg           Acenaphthylene         72         29         ug/kg           Anthracene         1300         29         ug/kg           Benzo (ghi) perylene         1600         29         ug/kg           Fluoranthene         5500         29         ug/kg           Fluorene         570         29         ug/kg           2-Methylnaphthalene         1000         29         ug/kg           Phenanthrene         4800         29         ug/kg           Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           PERCENT         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	Dibenz(a,h)anthracene	500	29	ug/kg
Acenaphthene 570 29 ug/kg Acenaphthylene 72 29 ug/kg Anthracene 1300 29 ug/kg Benzo(ghi)perylene 1600 29 ug/kg Fluoranthene 5500 29 ug/kg Fluorene 570 29 ug/kg 2-Methylnaphthalene 1000 29 ug/kg Phenanthrene 4800 29 ug/kg Pyrene 5400 29 ug/kg 1-Methylnaphthalene 610 29 ug/kg 1-Methylnaphthalene 610 29 ug/kg 1-Methylnaphthalene 610 (24 - 112) 2-Fluorobiphenyl 56 DIL (24 - 112) 2-Fluorobiphenyl 56 DIL (34 - 110) Terphenyl-d14 87 DIL (41 - 119) Phenol-d5 70 DIL (28 - 110) 2-Fluorophenol 72 DIL (26 - 110)	Indeno(1,2,3-cd)pyrene			ug/kg
Acenaphthylene 72 29 ug/kg Anthracene 1300 29 ug/kg Benzo(ghi)perylene 1600 29 ug/kg Fluoranthene 5500 29 ug/kg Fluorene 570 29 ug/kg 2-Methylnaphthalene 1000 29 ug/kg Phenanthrene 4800 29 ug/kg Pyrene 5400 29 ug/kg 1-Methylnaphthalene 610 29 ug/kg 1-Methylnaphthalene 610 29 ug/kg 1-Methylnaphthalene 610 (24 - 112) 2-Fluorobiphenyl 56 DIL (34 - 110) Terphenyl-d14 87 DIL (41 - 119) Phenol-d5 70 DIL (28 - 110) 2-Fluorophenol 72 DIL (26 - 110)	Naphthalene	820	29	ug/kg
Anthracene 1300 29 ug/kg Benzo(ghi)perylene 1600 29 ug/kg Fluoranthene 5500 29 ug/kg Fluorene 570 29 ug/kg 2-Methylnaphthalene 1000 29 ug/kg Phenanthrene 4800 29 ug/kg Pyrene 5400 29 ug/kg 1-Methylnaphthalene 610 29 ug/kg 1-Methylnaphthalene 610 29 ug/kg  PERCENT RECOVERY SURROGATE RECOVERY LIMITS Nitrobenzene-d5 68 DIL (24 - 112) 2-Fluorobiphenyl 56 DIL (34 - 110) Terphenyl-d14 87 DIL (41 - 119) Phenol-d5 70 DIL (28 - 110) 2-Fluorophenol 72 DIL (26 - 110)	Acenaphthene	570	29	ug/kg
Benzo(ghi)perylene         1600         29         ug/kg           Fluoranthene         5500         29         ug/kg           Fluorene         570         29         ug/kg           2-Methylnaphthalene         1000         29         ug/kg           Phenanthrene         4800         29         ug/kg           Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           PERCENT         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	Acenaphthylene	72	29	ug/kg
Fluoranthene       5500       29       ug/kg         Fluorene       570       29       ug/kg         2-Methylnaphthalene       1000       29       ug/kg         Phenanthrene       4800       29       ug/kg         Pyrene       5400       29       ug/kg         1-Methylnaphthalene       610       29       ug/kg         PERCENT       RECOVERY         SURROGATE       RECOVERY       LIMITS         Nitrobenzene-d5       68 DIL       (24 - 112)         2-Fluorobiphenyl       56 DIL       (34 - 110)         Terphenyl-d14       87 DIL       (41 - 119)         Phenol-d5       70 DIL       (28 - 110)         2-Fluorophenol       72 DIL       (26 - 110)	Anthracene	1300	29	ug/kg
Fluorene       570       29       ug/kg         2-Methylnaphthalene       1000       29       ug/kg         Phenanthrene       4800       29       ug/kg         Pyrene       5400       29       ug/kg         1-Methylnaphthalene       610       29       ug/kg         PERCENT       RECOVERY         SURROGATE       RECOVERY       LIMITS         Nitrobenzene-d5       68 DIL       (24 - 112)         2-Fluorobiphenyl       56 DIL       (34 - 110)         Terphenyl-d14       87 DIL       (41 - 119)         Phenol-d5       70 DIL       (28 - 110)         2-Fluorophenol       72 DIL       (26 - 110)	Benzo(ghi)perylene	1600	29	ug/kg
2-Methylnaphthalene       1000       29       ug/kg         Phenanthrene       4800       29       ug/kg         Pyrene       5400       29       ug/kg         1-Methylnaphthalene       610       29       ug/kg         PERCENT       RECOVERY         SURROGATE       RECOVERY       LIMITS         Nitrobenzene-d5       68 DIL       (24 - 112)         2-Fluorobiphenyl       56 DIL       (34 - 110)         Terphenyl-d14       87 DIL       (41 - 119)         Phenol-d5       70 DIL       (28 - 110)         2-Fluorophenol       72 DIL       (26 - 110)	Fluoranthene	5500	29	ug/kg
Phenanthrene         4800         29         ug/kg           Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           PERCENT         RECOVERY           SURROGATE         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	Fluorene	570	29	ug/kg
Pyrene         5400         29         ug/kg           1-Methylnaphthalene         610         29         ug/kg           PERCENT         RECOVERY           SURROGATE         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	2-Methylnaphthalene	1000	29	ug/kg
1-Methylnaphthalene         610         29         ug/kg           SURROGATE         PERCENT         RECOVERY           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	Phenanthrene	4800	29	ug/kg
PERCENT         RECOVERY           SURROGATE         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	<del>-</del>	5400	29	ug/kg
SURROGATE         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)	1-Methylnaphthalene	610	29	ug/kg
SURROGATE         RECOVERY         LIMITS           Nitrobenzene-d5         68 DIL         (24 - 112)           2-Fluorobiphenyl         56 DIL         (34 - 110)           Terphenyl-d14         87 DIL         (41 - 119)           Phenol-d5         70 DIL         (28 - 110)           2-Fluorophenol         72 DIL         (26 - 110)		PERCENT	RECOVERY	
Nitrobenzene-d5       68 DIL       (24 - 112)         2-Fluorobiphenyl       56 DIL       (34 - 110)         Terphenyl-d14       87 DIL       (41 - 119)         Phenol-d5       70 DIL       (28 - 110)         2-Fluorophenol       72 DIL       (26 - 110)	SURROGATE			
2-Fluorobiphenyl       56 DIL       (34 - 110)         Terphenyl-d14       87 DIL       (41 - 119)         Phenol-d5       70 DIL       (28 - 110)         2-Fluorophenol       72 DIL       (26 - 110)				
Terphenyl-d14 87 DIL (41 - 119) Phenol-d5 70 DIL (28 - 110) 2-Fluorophenol 72 DIL (26 - 110)				
Phenol-d5 70 DIL (28 - 110) 2-Fluorophenol 72 DIL (26 - 110)				
2-Fluorophenol 72 DIL (26 - 110)	<del>-</del>			
2,4,0 111510mophemor 44 515 (10 - 110)				
	2,4,0 IIIDIOMODIENOI	44 010	(10 110)	

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

Results and reporting limits have been adjusted for dry weight.

NOTE(S):

Client Sample ID: HB-075(2-4')

# GC Volatiles

Lot-Sample #: A9L180492-008 Date Sampled: 12/17/09 13:2 Prep Date: 12/23/09 Prep Batch #: 9358085 Dilution Factor: 1		12/18/09	Matrixso
<b>% Moisture:</b> 9.4	Method:	SW846 8015	A MOD
PARAMETER Gasoline Range Organics (C6-C12)	RESULT ND	REPORTING LIMIT 110	UNITS ug/kg
SURROGATE Trifluorotoluene	PERCENT RECOVERY 99	RECOVERY LIMITS (10 - 150)	

NOTE(S):

Client Sample ID: HB-075(2-4')

# GC Semivolatiles

Lot-Sample #: A9L180492-008	Work Order #:	LRCRR1AC	Matrix SO
Date Sampled: 12/17/09 13:20	Date Received:	12/18/09	
Prep Date: 12/19/09	Analysis Date:	12/23/09	
Prep Batch #: 9353014			
Dilution Factor: 5			
% Moisture: 9.4	Method:	SW846 8015	В
		REPORTING	
PARAMETER	RESULT	LIMIT	UNITS
C20-C34	110	15	mg/kg
C10-C20	48	11	mg/kg
	PERCENT	RECOVERY	
SURROGATE	RECOVERY	<u>LIMITS</u>	
C9 (nonane)	25 DIL	(10 - 110)	

NOTE(S):

DIL The concentration is estimated or not reported due to dilution or the presence of interfering analytes.

# Client Sample ID: HB-075(2-4')

# TOTAL Metals

Lot-Sample #...: A9L180492-008 Matrix.....: S0

Date Sampled...: 12/17/09 13:20 Date Received..: 12/18/09

**% Moisture....:** 9.4

PARAMETER	RESULT	REPORTIN LIMIT	G <u>UNITS</u>	METHOD	PREPARATION- WORK ANALYSIS DATE ORDER #
Prep Batch # Arsenic	: 9354052 1 <b>0.4</b>	1.1 Dilution Fac	mg/kg	SW846 6010B	12/21-12/23/09 LRCRR1AE
Cadmium	ND	0.22 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09 LRCRR1AF
Chromium	8.9	0.55 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09 LRCRR1AG
Lead	107	0.33 Dilution Fac	mg/kg tor: 1	SW846 6010B	12/21-12/23/09 LRCRR1AH
NOTE(S):					

Client Sample ID: HB-075(2-4')

# General Chemistry

Lot-Sample #...: A9L180492-008 Work Order #...: LRCRR Matrix.....: S0

Date Sampled...: 12/17/09 13:20 Date Received..: 12/18/09

**% Moisture....:** 9.4

PARAMETER	RESULT	RL	UNITS	METHO1	<b>D</b>		PREPARATION- ANALYSIS DATE	PREP BATCH #
Acid-soluble sulfide	ND	33.1 Dilution Fact	mg/kg or: 1	SW846	9030B/90	34	12/23/09	9357096
Corrosivity	8.7	Dilution Fact	No Units	SW846	9045A		12/19/09	9353120
Hexavalent Chromium	ND	0.88	mg/kg	SW846	7196A		12/23-12/24/09	9357236
		Dilution Fact	or: 1					
Percent Solids	90.6	10.0 Dilution Fact	<b>%</b> or: 1	MCAWW	160.3 MO	D	12/22-12/23/09	9356363
Total Cyanide	ND	0.55 Dilution Fact	mg/kg or: 1	MCAWW	335.2 CL	P-M	12/28/09	9362384

# NOTE(S):

RL Reporting Limit



# QUALITY CONTROL SECTION

# GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRGQR1AA Matrix.....: SOLID

MB Lot-Sample #: A9L220000-203

Prep Date....: 12/21/09
Prep Batch #...: 9356203

Analysis Date..: 12/21/09

Dilution Factor: 1

4-Bromofluorobenzene

### REPORTING

		KEFOKI.	140	
PARAMETER	RESULT	<u>LIMIT</u>	UNITS	METHOD
Chloromethane	ND	10	ug/kg	SW846 8260A
Bromomethane	ND	10	ug/kg	SW846 8260A
Vinyl chloride	ND	2.0	ug/kg	SW846 8260A
Chloroethane	ND	10	ug/kg	SW846 8260A
Methylene chloride	ND	5.0	ug/kg	SW846 8260A
Acetone	ND	20	ug/kg	SW846 8260A
Carbon disulfide	ND	5.0	ug/kg	SW846 8260A
1,1-Dichloroethene	ND	5.0	ug/kg	SW846 8260A
1,1-Dichloroethane	ND	5.0	ug/kg	SW846 8260A
1,2-Dichloroethene (total)	ND	5.0	ug/kg	SW846 8260A
Chloroform	ND	5.0	ug/kg	SW846 8260A
1,2-Dichloroethane	ND	5.0	ug/kg	SW846 8260A
2-Butanone	ND	20	ug/kg	SW846 8260A
1,1,1-Trichloroethane	ND	5.0	ug/kg	SW846 8260A
Carbon tetrachloride	ND	5.0	ug/kg	SW846 8260A
Bromodichloromethane	ND	5.0	ug/kg	SW846 8260A
1,2-Dichloropropane	ND	5.0	ug/kg	SW846 8260A
cis-1,3-Dichloropropene	ND	5.0	ug/kg	SW846 8260A
Trichloroethene	ND	5.0	ug/kg	SW846 8260A
Dibromochloromethane	ND	5.0	ug/kg	SW846 8260A
1,1,2-Trichloroethane	ND	5.0	ug/kg	SW846 8260A
Benzene	ND	5.0	ug/kg	SW846 8260A
trans-1,3-Dichloropropene	ND	5.0	ug/kg	SW846 8260A
Bromoform	ND	5.0	ug/kg	SW846 8260A
4-Methyl-2-pentanone	ND	20	ug/kg	SW846 8260A
2-Hexanone	ND	20	ug/kg	SW846 8260A
Tetrachloroethene	ND	5.0	ug/kg	SW846 8260A
1,1,2,2-Tetrachloroethane	ND	5.0	ug/kg	SW846 8260A
Toluene	ND	5.0	ug/kg	SW846 8260A
Chlorobenzene	ND	5.0	ug/kg	SW846 8260A
Ethylbenzene	ND	5.0	ug/kg	SW846 8260A
Styrene	ND	5.0	ug/kg	SW846 8260A
Xylenes (total)	ND	5.0	ug/kg	SW846 8260A
	PERCENT	RECOVER	Y	
SURROGATE	RECOVERY	LIMITS		
Dibromofluoromethane	99	(59 - 1		
1,2-Dichloroethane-d4	76	(61 - 13		
Toluene-d8	95	(60 - 14	-	
4 5	0.1	117 41		

(Continued on next page)

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(47 - 158)

# GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRGQR1AA Matrix.....: SOLID

NOTE(S):

### GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRH4C1AA Matrix.....: SOLID

MB Lot-Sample #: A9L220000-563

Prep Date....: 12/20/09

Dilution Factor: 1

Dibromofluoromethane

4-Bromofluorobenzene

Toluene-d8

1,2-Dichloroethane-d4

#### REPORTING PARAMETER RESULT LIMIT UNITS METHOD Acetone ND 1000 ug/kg SW846 8260A Benzene ND 250 ug/kg SW846 8260A Bromodichloromethane ND 250 ug/kg SW846 8260A Bromoform ND 250 ug/kg SW846 8260A Bromomethane ND 500 ug/kg SW846 8260A 2-Butanone ND 1000 SW846 8260A ug/kg Carbon disulfide ND 250 SW846 8260A ug/kg Carbon tetrachloride 250 ND ug/kg SW846 8260A Chlorobenzene ND 250 SW846 8260A ug/kg Dibromochloromethane ND 250 SW846 8260A ug/kg Chloroethane 500 ND ug/kg SW846 8260A Chloroform ND 250 ug/kg SW846 8260A Chloromethane ND 500 SW846 8260A ug/kg 1,1-Dichloroethane ND250 ug/kg SW846 8260A 1,2-Dichloroethane ND250 ug/kg SW846 8260A 1,1-Dichloroethene ND 250 SW846 8260A ug/kg 1,2-Dichloroethene ND250 ug/kg SW846 8260A (total) 1,2-Dichloropropane ND 250 SW846 8260A ug/kg cis-1,3-Dichloropropene ND 250 ug/kg SW846 8260A trans-1,3-Dichloropropene ND 250 ug/kg SW846 8260A Ethylbenzene ND 250 ug/kg SW846 8260A 2-Hexanone ND 1000 SW846 8260A ug/kg Methylene chloride ND 250 ug/kg SW846 8260A 4-Methyl-2-pentanone ND 1000 SW846 8260A ug/kg Styrene 250 ND SW846 8260A ug/kg 1,1,2,2-Tetrachloroethane ND 250 ua/ka SW846 8260A Tetrachloroethene ND 250 SW846 8260A ug/kg Toluene ND 250 ug/kg SW846 8260A 1,1,1-Trichloroethane ND 250 ug/kg SW846 8260A 1,1,2-Trichloroethane ND250 ug/kg SW846 8260A Trichloroethene ND 250 ug/kg SW846 8260A Vinyl chloride ND 100 ug/kg SW846 8260A Xylenes (total) ND250 ug/kg SW846 8260A PERCENT RECOVERY SURROGATE RECOVERY LIMITS

(Continued on next page)

(59 - 138)

(61 - 130)

(60 - 143)

(47 - 158)

100

105

97

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# GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRH4C1AA Matrix.....: SOLID

NOTE(S):

### GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRNH41AA Matrix.....: SOLID

**MB Lot-Sample #:** A9L280000-414

Prep Date....: 12/23/09

**Analysis Date..:** 12/23/09 **Prep Batch #...:** 9362414

Dilution Factor: 1

4-Bromofluorobenzene

		REPORTII		
PARAMETER	RESULT	LIMIT	UNITS	METHOD
Chloromethane	ND	10	ug/kg	SW846 8260A
Bromomethane	ND	10	ug/kg	SW846 8260A
Vinyl chloride	ND	2.0	ug/kg	SW846 8260A
Chloroethane	ND	10	ug/kg	SW846 8260A
Methylene chloride	ND	5.0	ug/kg	SW846 8260A
Acetone	ND	20	ug/kg	SW846 8260A
Carbon disulfide	ND	5.0	ug/kg	SW846 8260A
1,1-Dichloroethene	ND	5.0	ug/kg	SW846 8260A
1,1-Dichloroethane	ND	5.0	ug/kg	SW846 8260A
1,2-Dichloroethene	ND	5.0	ug/kg	SW846 8260A
(total)				
Chloroform	ND	5.0	ug/kg	SW846 8260A
1,2-Dichloroethane	ND	5.0	ug/kg	SW846 8260A
2-Butanone	ND	20	ug/kg	SW846 8260A
1,1,1-Trichloroethane	ND	5.0	ug/kg	SW846 8260A
Carbon tetrachloride	ND	5.0	ug/kg	SW846 8260A
Bromodichloromethane	ND	5.0	ug/kg	SW846 8260A
1,2-Dichloropropane	ND	5.0	ug/kg	SW846 8260A
cis-1,3-Dichloropropene	ND	5.0	ug/kg	SW846 8260A
Trichloroethene	ND	5.0	ug/kg	SW846 8260A
Dibromochloromethane	ND	5.0	ug/kg	SW846 8260A
1,1,2-Trichloroethane	ND	5.0	ug/kg	SW846 8260A
Benzene	ND	5.0	ug/kg	SW846 8260A
trans-1,3-Dichloropropene	ND	5.0	ug/kg	SW846 8260A
Bromoform	ND	5.0	ug/kg	SW846 8260A
4-Methyl-2-pentanone	ND	20	ug/kg	SW846 8260A
2-Hexanone	ND	20	ug/kg	SW846 8260A
Tetrachloroethene	ND	5.0	ug/kg	SW846 8260A
1,1,2,2-Tetrachloroethane	ND	5.0	ug/kg	SW846 8260A
Toluene	ND	5.0	ug/kg	SW846 8260A
Chlorobenzene	ND	5.0	ug/kg	SW846 8260A
Ethylbenzene	ND	5.0	ug/kg	SW846 8260A
Styrene	ND	5.0	ug/kg	SW846 8260A
Xylenes (total)	ND	5.0	ug/kg	SW846 8260A
	PERCENT	RECOVERY	,	
SURROGATE	RECOVERY	LIMITS		
Dibromofluoromethane	95	(59 - 13	81	
1,2-Dichloroethane-d4	89	(61 - 13		
Toluene-d8	96	(60 - 14		
	<i>J</i> 0	(00 - 14	: )	

(Continued on next page)

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# GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRNH41AA Matrix.....: SOLID

NOTE(S):

# GC/MS Semivolatiles

Client Lot #...: A9L180492 Work Order #...: LRDRM1AA Matrix.....: SOLID

MB Lot-Sample #: A9L190000-017

Prep Date....: 12/19/09
Prep Batch #...: 9353017

Analysis Date..: 12/24/09

Dilution Factor: 1

REP	ORT	ING
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PARAMETER	RESULT	LIMIT	UNITS	METHOD
Acenaphthene	ND	6.7	ug/kg	SW846 8270C
Acenaphthylene	ND	6.7	ug/kg	SW846 8270C
Anthracene	ND	6.7	ug/kg	SW846 8270C
Benzo(a)anthracene	ND	6.7	ug/kg	SW846 8270C
Benzo(b)fluoranthene	ND	6.7	ug/kg	SW846 8270C
Benzo(k)fluoranthene	ND	6.7	ug/kg	SW846 8270C
Benzo(ghi)perylene	ND	6.7	ug/kg	SW846 8270C
Benzo(a)pyrene	ND	6.7	ug/kg	SW846 8270C
Chrysene	ND	6.7	ug/kg	SW846 8270C
Dibenz(a,h)anthracene	ND	6.7	ug/kg	SW846 8270C
Fluoranthene	ND	6.7	ug/kg	SW846 8270C
Fluorene	ND	6.7	ug/kg	SW846 8270C
Indeno(1,2,3-cd)pyrene	ND	6.7	ug/kg	SW846 8270C
2-Methylnaphthalene	ND	6.7	ug/kg	SW846 8270C
Naphthalene	ND	6.7	ug/kg	SW846 8270C
Phenanthrene	ND	6.7	ug/kg	SW846 8270C
Pyrene	ND	6.7	ug/kg	SW846 8270C
	PERCENT	RECOVERS	Z	
SURROGATE	RECOVERY	LIMITS		
Nitrobenzene-d5	82	(24 - 11	12)	
2-Fluorobiphenyl	74	(34 - 11	LO)	
Terphenyl-d14	117	(41 - 11	L9)	
Phenol-d5	87	(28 - 11	LO)	
2-Fluorophenol	92	(26 - 11	LO)	
2,4,6-Tribromophenol	51	(10 - 11	L8)	

NOTE(S):

### GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LREOV1AA Matrix...... SOLID

MB Lot-Sample #: A9L200000-104

Prep Date....: 12/19/09
Analysis Date..: 12/19/09
Prep Batch #...: 9354104

Analysis Date..: 12/19/09 Prep Batch #...: 9354104 Dilution Factor: 1

,11401011 1400011 1

PARAMETER RESULT LIMIT UNITS METHOD

Gasoline Range Organics ND 100 ug/kg SW846 8015A MOD (C6-C12)

PERCENT RECOVERY

SURROGATERECOVERYLIMITSTrifluorotoluene99(10 - 150)

### GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LRMPJ1AA Matrix.....: SOLID

MB Lot-Sample #: A9L240000-085

Prep Date...: 12/23/09
Analysis Date.: 12/23/09
Prep Batch #..: 9358085

Dilution Factor: 1

REPORTING

PARAMETER RESULT LIMIT UNITS METHOD

Gasoline Range Organics ND 100 ug/kg SW846 8015A MOD (C6-C12)

PERCENT RECOVERY

SURROGATERECOVERYLIMITSTrifluorotoluene99(10 - 150)

# GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LRNPP1AA Matrix.....: SOLID

MB Lot-Sample #: A9L290000-092

Prep Date....: 12/28/09 Prep Batch #...: 9363092

Analysis Date..: 12/28/09

Dilution Factor: 1

REPORTING

PARAMETER RESULT LIMIT UNITS METHOD

Gasoline Range Organics

ND

100 ug/kg

SW846 8015A MOD

(C6-C12)

PERCENT RECOVERY

RECOVERY LIMITS

SURROGATE
Trifluorotoluene

101

(10 - 150)

### NOTE(S):

### GC Semivolatiles

Client Lot #...: A9L180492 Work Order #...: LRDRJ1AA Matrix.....: SOLID

MB Lot-Sample #: A9L190000-014

Prep Date....: 12/19/09
Prep Batch #...: 9353014

Analysis Date..: 12/23/09

Dilution Factor: 1

REPORTING

(10 - 110)

LIMIT RESULT PARAMETER UNITS METHOD C20-C34 2.7 mg/kg SW846 8015B  ${\rm ND}$ 2.0 SW846 8015B C10-C20 ND mg/kg PERCENT RECOVERY LIMITS SURROGATE RECOVERY

NOTE(S):

C9 (nonane)

Calculations are performed before rounding to avoid round-off errors in calculated results.

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### TOTAL Metals

PARAMETER	RESULT	REPORTING	G <u>UNITS</u>	METHOD	PREPARATION- WORK  ANALYSIS DATE ORDER #
MB Lot-Sample	#: A9L200000-	052 <b>Prep B</b>	atch #:	9354052	
Arsenic	ND	1.0	mg/kg	SW846 6010E	12/21-12/23/09 LREWX1AA
		Dilution Fact	or: 1		
Cadmium	ND	0.20	mg/kg	SW846 6010B	12/21-12/23/09 LREWX1AC
	1.2	Dilution Fact	2 2	2,,010	15, 21 10, 50, 65 2
Chromium	ND	0.50	mg/kg	SW846 6010E	12/21-12/23/09 LREWX1AD
		Dilution Fact	or: 1		
Lead	ND	0.30	mg/kg	SW846 6010E	12/21-12/23/09 LREWX1AE
		Dilution Fact	or: 1		
NOTE(S):					

# General Chemistry

Matrix..... SOLID

Client Lot #...: A9L180492

PARAMETER	RESULT	REPORTING	UNITS	METHOD	PREPARATION- ANALYSIS DATE	PREP <u>BATCH #</u>
Acid-soluble sulfic	de ND		mg/kg	MB Lot-Sample #: SW846 9030B/9034		9357096
Hexavalent Chromium		Work Order	#: LRJPH1AA	MB Lot-Sample #:	A9L230000-236	
CITE ONLE COM	ND	0.80 Dilution Fact		SW846 7196A	12/23-12/24/09	9357236
Percent Solids	ND		ક	MB Lot-Sample #: MCAWW 160.3 MOD	A9L220000-363 12/22-12/23/09	9356363
Total Cyanide	ND		mg/kg	MB Lot-Sample #: MCAWW 335.2 CLP-M		9357388
Total Cyanide	ND	Work Order 0.50 Dilution Fact	mg/kg	MB Lot-Sample #: MCAWW 335.2 CLP-M		9362384

Calculations are performed before rounding to avoid round-off errors in calculated results.

NOTE(S):

# GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRGQR1AC-LCS Matrix.....: SOLID

LCS Lot-Sample#: A9L220000-203 LRGQR1AD-LCSD

Prep Date....: 12/21/09 Analysis Date..: 12/21/09

Prep Batch #...: 9356203

Dilution Factor: 1

		DEGG!!ED!!	222	
	PERCENT	RECOVERY	RPD	MORITOD
PARAMETER	RECOVERY	LIMITS	RPD LIMITS	METHOD
1,1-Dichloroethene	107	(55 - 142)		SW846 8260A
	105	<b>(55 - 142)</b>	2.5 (0-27)	
Trichloroethene	104	(70 - 131)		SW846 8260A
	102	(70 - 131)	1.7 (0-23)	SW846 8260A
Benzene	99	<b>(75 - 129)</b>		SW846 8260A
	100	(75 - 129)	1.3 (0-20)	SW846 8260A
Toluene	99	(71 - 130)		SW846 8260A
1010000	101	(71 - 130)	2.5 (0-24)	SW846 8260A
Chlorobenzene	96	(75 - 127)		SW846 8260A
CHICLOSCALO	97	(75 - 127)	1.6 (0-22)	SW846 8260A
		•		
		PERCENT	RECOVERY	
SURROGATE		RECOVERY	LIMITS	
Dibromofluoromethane		100	(59 - 138)	
		104	(59 - 138)	
1,2-Dichloroethane-d4		78	(61 - 130)	
1,2 Dichiologonane da		78	(61 - 130)	
m-1 30		97	(60 - 143)	
Toluene-d8				
		101	(60 - 143)	
4-Bromofluorobenzene		96	(47 - 158)	
		95	(47 - 158)	

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

### GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRH4C1AC-LCS Matrix.....: SOLID

LCS Lot-Sample#: A9L220000-563 LRH4C1AD-LCSD

Prep Date....: 12/20/09 Analysis Date..: 12/22/09

Prep Batch #...: 9356563

Dilution Factor: 1

	PERCENT	RECOVERY	RPD	
PARAMETER	<u>RECOVERY</u>	<u>LIMITS</u>	RPD LIMITS	METHOD
Benzene	100	(75 - 129)		SW846 8260A
	92	<b>(75 - 129)</b>	7.9 (0-20)	SW846 8260A
Chlorobenzene	96	<b>(75 - 127)</b>		SW846 8260A
	91	<b>(75 - 127)</b>	5.4 (0-22)	SW846 8260A
1,1-Dichloroethene	100	(55 - <b>142</b> )		SW846 8260A
	91	(55 - 142)	9.4 (0-27)	SW846 8260A
Toluene	93	(71 - 130)		SW846 8260A
	88	(71 - 130)	5.2 (0-24)	SW846 8260A
Trichloroethene	96	(70 - 131)		SW846 8260A
	92	(70 - 131)	4.5 (0-23)	SW846 8260A
		PERCENT	RECOVERY	
SURROGATE		RECOVERY	LIMITS	
Dibromofluoromethane		111	(59 - 138)	
<u> </u>		104	(59 - 138)	
1,2-Dichloroethane-d4		111	(61 - 130)	
I, I DIGITO, O OCIONIO WI		106	(61 - 130)	
Toluene-d8		101	(60 - 143)	
Totache do		95	(60 - 143)	
4-Bromofluorobenzene		100	(47 - 158)	
4 DIOMOTIMOTODEHZEHE		95	(47 - 158)	
		95	(47 - 136)	

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

### GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRNH41AC-LCS Matrix...... SOLID

LCS Lot-Sample#: A9L280000-414 LRNH41AD-LCSD

Prep Date....: 12/23/09 Analysis Date..: 12/23/09

Prep Batch #...: 9362414

Dilution Factor: 1

	PERCENT	RECOVERY		RPD		
PARAMETER	RECOVERY	LIMITS	RPD	<u>LIMITS</u>	METHOD	
1,1-Dichloroethene	118	(55 - 142)			SW846	8260A
	124	<b>(55 - 142)</b>	4.9	(0-27)	SW846	8260A
Trichloroethene	107	(70 - 131)			SW846	8260A
	102	(70 - 131)	4.2	(0-23)	SW846	B260A
Benzene	103	(75 - 129)			SW846 8	3260A
	99	(75 - 129)	3.8	(0-20)	SW846	8260A
Toluene	100	(71 - 130)			SW846	3260A
	91	(71 - 130)	9.6	(0-24)	SW846 8	3260A
Chlorobenzene	99	(75 - 127)			SW846	3260A
	94	(75 - 127)	5.3	(0-22)	SW846 8	3260A
		PERCENT	RECOV	ERY		
SURROGATE		RECOVERY	LIMIT	S		
Dibromofluoromethane		96	(59 -	138)		
		98	(59 –	138)		
1,2-Dichloroethane-d4		87	(61 -	130)		
·		87	(61 -	·		
Toluene-d8		100	(60 -			
		96	(60 -			
4-Bromofluorobenzene		93	(47 -	·		
· ·		92	(47 -	•		
		-	,	,		

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

# GC/MS Semivolatiles

Client Lot #...: A9L180492 Work Order #...: LRDRM1AC Matrix.....: SOLID

LCS Lot-Sample#: A9L190000-017

Prep Date....: 12/19/09 Analysis Date..: 12/24/09

Prep Batch #...: 9353017

Dilution Factor: 1

	DEDGEME	DEGOTERN	
	PERCENT RECOVERY	RECOVERY LIMITS	METHOD
PARAMETER	63	(43 - 110)	SW846 8270C
1,2,4-Trichloro-	0.5	(45 - 110)	SW040 02/UC
benzene	0.2	(55 116)	OT 10 4 C 9 2 7 0 G
2,4-Dinitrotoluene	83	(55 - 116)	SW846 8270C
N-Nitrosodi-n-propyl- amine	75	(40 - 114)	SW846 8270C
1,4-Dichlorobenzene	64	(38 - 110)	SW846 8270C
Pentachlorophenol	5 <b>6</b>	(10 - 110)	SW846 8270C
Phenol	77	(39 - 110)	SW846 8270C
2-Chlorophenol	70	(39 - 110)	SW846 8270C
4-Chloro-3-methylphenol	84	(42 - 110)	SW846 8270C
4-Nitrophenol	85	(24 - 117)	SW846 8270C
Acenaphthene	71	(46 - 110)	SW846 8270C
Pyrene	89	(58 - 113)	SW846 8270C
		PERCENT	RECOVERY
SURROGATE	-	RECOVERY	<u>LIMITS</u>
Nitrobenzene-d5		72	(24 - 112)
2-Fluorobiphenyl		64	(34 - 110)
Terphenyl-d14		97	(41 - 119)
Phenol-d5		75	(28 - 110)
2-Fluorophenol		80	(26 - 110)
2,4,6-Tribromophenol		57	(10 - 118)

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

# GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LREOV1AC-LCS Matrix.....: SOLID

LCS Lot-Sample#: A9L200000-104 LRE0V1AD-LCSD

Prep Date....: 12/19/09 Analysis Date..: 12/19/09

Prep Batch #...: 9354104

Dilution Factor: 1

PARAMETER Gasoline Range Organics (C6-C12)	PERCENT <u>RECOVERY</u> 111	RECOVERY LIMITS (60 - 142)	RPD RPD LIMITS	METHOD SW846 8015A MOD
	109	(60 - 142)	1.8 (0-27)	SW846 8015A MOD
		PERCENT	RECOVERY	
SURROGATE		RECOVERY	LIMITS	

 SURROGATE
 RECOVERY
 LIMITS

 Trifluorotoluene
 104
 (10 - 150)

 101
 (10 - 150)

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results. Bold print denotes control parameters

# GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LRMPJ1AC-LCS Matrix.....: SOLID

LCS Lot-Sample#: A9L240000-085 LRMPJ1AD-LCSD

Prep Date....: 12/23/09 Analysis Date..: 12/23/09

**Prep Batch #...:** 9358085

Dilution Factor: 1

PARAMETER Gasoline Range Organics (C6-C12)	PERCENT RECOVERY 98 112	RECOVERY LIMITS (60 - 142) (60 - 142)	13	RPD LIMITS (0-27)	 8015A 8015A	
SURROGATE Trifluorotoluene		PERCENT RECOVERY 102	RECOV LIMIT (10 -	S		

104 (10 - 150)

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

# GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LRNPP1AC-LCS Matrix......: SOLID

LCS Lot-Sample#: A9L290000-092 LRNPP1AD-LCSD

Prep Date....: 12/28/09 Analysis Date..: 12/29/09

**Prep Batch #...:** 9363092

Dilution Factor: 1

PARAMETER Gasoline Range Organics (C6-C12)	PERCENT RECOVERY 106	RECOVERY LIMITS (60 - 142)	RPD LIMIT	SW846 8015A MOD
	108	(60 - 142)	2.3 (0-2)	) SW846 8015A MOD
		PERCENT	RECOVERY	
SURROGATE		RECOVERY	LIMITS	
Trifluorotoluene		102	(10 - 150)	
		101	(10 - 150)	

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

### GC Semivolatiles

Client Lot #...: A9L180492 Work Order #...: LRDRJ1AC Matrix.....: SOLID

LCS Lot-Sample#: A9L190000-014

Prep Date....: 12/19/09 Analysis Date..: 12/23/09

Prep Batch #...: 9353014

Dilution Factor: 1

PERCENT RECOVERY

PARAMETER RECOVERY LIMITS METHOD

TPH (as Diesel) 79 (47 - 138) SW846 8015B

SURROGATEPERCENTRECOVERYLIMITS

C9 (nonane) 34 (10 - 110)

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

# TOTAL Metals

Client Lot #:	A9L180492			Matrix	: SOLID
PARAMETER	PERCENT RECOVERY	RECOVERY LIMITS	METHOD	PREPARATION- ANALYSIS DATE	WORK ORDER #
LCS Lot-Sample#: Arsenic	A9L200000- 95	_	tch #: 9354052 SW846 6010B or: 1	12/21-12/23/09	LREWX1AF
Cadmium	97	(80 - 120) Dilution Facto	SW846 6010B or: 1	12/21-12/23/09	LREWX1AG
Chromium	95	(80 - 120) Dilution Facto	SW846 6010B or: 1	12/21-12/23/09	LREWX1AH
Lead	99	(80 - 120) Dilution Facto	SW846 6010B or: 1	12/21-12/23/09	LREWX1AJ
MOTE (C) ·					

# General Chemistry

PARAMETER Acid-soluble su	PERCENT RECOVERY 11fide 87	LIMITS METHOD ANALYSIS DATE Work Order #: LRKCN1AC LCS Lot-Sample#: A9L230000-	PREP BATCH # 096 9357096
Corrosivity	99	Work Order #: LREE21AA LCS Lot-Sample#: A9L190000- (97 - 103) SW846 9045A 12/19/09 Dilution Factor: 1	-120 9353120
Hexavalent Chromium		Work Order #: LRJPH1AC LCS Lot-Sample#: A9L230000-	-236
CIII OILI EIL	110	(80 - 120) SW846 7196A 12/23-12/24/09 Dilution Factor: 1	9357236
Total Cyanide	102	Work Order #: LRKK41AC LCS Lot-Sample#: A9L230000- (68 - 123) MCAWW 335.2 CLP-M 12/23/09 Dilution Factor: 1	-388 9357388
Total Cyanide	107	Work Order #: LRNER1AC LCS Lot-Sample#: A9L280000- (68 - 123) MCAWW 335.2 CLP-M 12/28/09 Dilution Factor: 1	-384 9362384

NOTE(S):

# GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRD521AF-MS Matrix.....: SOLID

MS Lot-Sample #: A9L190450-001 LRD521AG-MSD

Date Sampled...: 12/18/09 11:45 Date Received..: 12/19/09
Prep Date....: 12/21/09 Analysis Date..: 12/21/09

**Prep Batch #...:** 9356203

Dilution Factor: 0.92	% Moi:	sture: 19	)				
	PERCENT	RECOVERY		RPD			
PARAMETER	RECOVERY	LIMITS	RPD	LIMITS	METHOI	)	
1,1-Dichloroethene	97	(43 - 147)			SW846	8260A	
	88	(43 - 147)	16	(0-27)	SW846	8260A	
Trichloroethene	73	(46 - 143)			SW846	8260A	
	66	(46 - 143)	17	(0-23)	SW846	8260A	
Benzene	82	(55 - 138)			SW846	8260A	
	73	(55 - 138)	19	(0-20)	SW846	8260A	
Toluene	87	(46 - 147)			SW846	8260A	
	57 p	(46 - 147)	45	(0-24)	SW846	8260A	
Chlorobenzene	65	(49 - 139)			SW846	8260A	
	47 a,p	(49 – 139)	40	(0-22)	SW846	8260A	
		PERCENT		RECOVERY			
SURROGATE	_	RECOVERY		LIMITS	<del>.</del>		
Dibromofluoromethane		107		(59 - 138	•		
		104		(59 - 138	•		
1,2-Dichloroethane-d4		83		(61 - 130			
		82		(61 - 130	•		
Toluene-d8		113		(60 - 143	•		
		100		(60 - 143	•		
4-Bromofluorobenzene		174 *		(47 - 158	•		
		107		(47 - 158	)		

#### NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

- p Relative percent difference (RPD) is outside stated control limits.
- a Spiked analyte recovery is outside stated control limits.
- \* Surrogate recovery is outside stated control limits.

### GC/MS Volatiles

Client Lot #...: A9L180492 Work Order #...: LRKXH1AG-MS Matrix.....: SOLID

MS Lot-Sample #: A9L230573-001 LRKXH1AH-MSD

Date Sampled...: 12/23/09 10:30 Date Received..: 12/23/09 Prep Date....: 12/24/09 Analysis Date..: 12/24/09

Prep Batch #...: 9362414

Dilution Factor: 1 % Moisture....: 18

	PERCENT	RECOVERY		RPD		
PARAMETER	RECOVERY	LIMITS	RPD	LIMITS	METHO	)
1,1-Dichloroethene	108	(43 - 147)			SW846	8260A
•	108	(43 - 147)	0.06	(0-27)	SW846	8260A
Trichloroethene	93	(46 - 143)			SW846	8260A
	90	(46 - 143)	3.0	(0-23)	SW846	8260A
Benzene	93	(55 - 138)			SW846	8260A
	89	(55 - 138)	4.2	(0-20)	SW846	8260A
Toluene	88	(46 - 147)			SW846	8260A
	83	(46 - 147)	6.4	(0-24)	SW846	8260A
Chlorobenzene	83	<b>(49 - 139)</b>			SW846	8260A
	81	(49 - 139)	2.4	(0-22)	SW846	8260A
		PERCENT		RECOVERY		
SURROGATE	_	RECOVERY		LIMITS		
Dibromofluoromethane		96		(59 - 138	:)	
		98		(59 - 138	1)	
1,2-Dichloroethane-d4		84		(61 - 130	)	
		86		(61 - 130	)	
Toluene-d8		97		(60 - 143	)	
		100		(60 - 143	)	
4-Bromofluorobenzene		88		(47 - 158	)	
		93		(47 - 158	)	

#### NOTE (S)

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

### GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LRCRP1AQ-MS Matrix.....: S0

MS Lot-Sample #: A9L180492-007 LRCRP1AR-MSD

Date Sampled...: 12/17/09 13:20 Date Received..: 12/18/09 Prep Date....: 12/23/09 Analysis Date..: 12/24/09

**Prep Batch #...:** 9358085

Dilution Factor: 5

	PERCENT	RECOVERY	BDD	RPD LIMITS	METHOD
PARAMETER Gasoline Range Organics	RECOVERY 45	LIMITS (10 - 142)	RPD	LIMIT 15	SW846 8015A MOD
(C6-C12)	21	(10 - 142)	24	(0-94)	SW846 8015A MOD
		PERCENT		RECOVERY	
SURROGATE	_	RECOVERY		LIMITS	_
Trifluorotoluene		100		(10 - 150	)
		98		(10 - 150	1)

### NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

# GC Volatiles

Client Lot #...: A9L180492 Work Order #...: LQ8TD1AJ-MS Matrix.....: SOLID

**MS Lot-Sample #:** A9L170476-009 LQ8TD1AK-MSD

Date Sampled...: 12/16/09 15:40 Date Received..: 12/17/09 Prep Date....: 12/29/09 Analysis Date..: 12/29/09

Prep Batch #...: 9363092

	PERCENT	RECOVERY		RPD			
PARAMETER	RECOVERY	<u>LIMITS</u>	RPD_	LIMITS	METHOI	)	
Gasoline Range Organics (C6-C12)	112	(10 - 142)			SW846	8015A	MOD
	178 a	(10 - 142)	25	(0-94)	SW846	8015A	MOD
		PERCENT		RECOVERY			
SURROGATE	_	<u>RECOVERY</u>		LIMITS	_		
Trifluorotoluene		101		(10 - 150	)		
		101		(10 - 150	)		

### NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

a Spiked analyte recovery is outside stated control limits.

### TOTAL Metals

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

PARAMETER	PERCENT RECOVERY		RPD <u>LIMITS</u>	METHOD	PREPARATION- ANALYSIS DATE	WORK ORDER #				
MS Lot-Sample #: A9L180492-001 Prep Batch #: 9354052										
Arsenic	90	(75 - 125)		SW846 6010B	12/21-12/23/09	LRCET1AQ				
	89	(75 - 125) 0.73	(0-20)	SW846 6010B	12/21-12/23/09	LRCET1AR				
Dilution Factor: 1										
Cadmium		(75 - 125)		SW846 6010B	12/21-12/23/09	LRCET1AT				
	85	(75 - 125) 1.1	(0-20)	SW846 6010B	12/21-12/23/09	LRCET1AU				
Dilution Factor: 1										
Chromium	0.0	/7F 10F)		07:10.4.C. C.0.1.0.D.	10/01 10/03/00					
CHEOMIUM	89	(75 - 125)		SW846 6010B	12/21-12/23/09					
	93	(75 - 125) 2.8	/	SW846 6010B	12/21-12/23/09	LRCET1AW				
		Dilution Facto	or: 1							
Lead	87	/7E 10E)		ario 4.6 - 6.01.0 b	10/01/10/02/08	T D Opport 3 tr				
Lead	<del>-</del> ·	(75 - 125)		SW846 6010B	12/21-12/23/09					
	92	(75 - 125) 3.5		SW846 6010B	12/21-12/23/09	LRCET1A0				
		Dilution Facto	or: 1							

# NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

# General Chemistry

Client Lot #: A9L180492									
PARAMETER	PERCENT RECOVERY	RECOVERY LIMITS	RPD RPD LIMITS	METHOD	PREPARATION- PREP ANALYSIS DATE BATCH # 8 Moisture: 13				
Acid-soluble		(10 - 154) (10 - 154)		SW846 9030B/9034	ot-Sample #: A9L170448-001				
Cyanide, Tot	al 78 64	(50 - 134) (50 - 134)		Q4981AD-MSD MS L MCAWW 335.2 CLP-M MCAWW 335.2 CLP-M					
Total Cyanid	e 72 61	(50 - 134) (50 - 134)		RG481AL-MSD MS L MCAWW 335.2 CLP-M MCAWW 335.2 CLP-M					

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

### SAMPLE DUPLICATE EVALUATION REPORT

### General Chemistry

Client Lot #...: A9L180492 Work Order #...: LRCRR-SMP Matrix.....: SO

LRCRR-DUP

Date Sampled...: 12/17/09 13:20 Date Received..: 12/18/09

**% Moisture....:** 9.4

 PARAM RESULT
 RESULT
 UNITS
 RPD
 PREPARATION PREPA

Dilution Factor: 1

#### SAMPLE DUPLICATE EVALUATION REPORT

#### General Chemistry

Client Lot #...: A9L180492 Work Order #...: LQ1E4-SMP Matrix.....: SOLID

LQ1E4-DUP

Date Sampled...: 12/10/09 09:55 Date Received..: 12/11/09

**% Moisture....:** 19

 PARAM RESULT
 RESULT
 UNITS
 RPD
 METHOD
 ANALYSIS DATE
 BATCH #

 Percent Solids
 81.1
 79.6
 %
 1.8
 (0-20)
 MCAWW 160.3 MOD
 12/22-12/23/09
 9356363

Dilution Factor: 1

#### SAMPLE DUPLICATE EVALUATION REPORT

#### General Chemistry

Client Lot #...: A9L180492 Work Order #...: LRCRE-SMP Matrix.....: SO

LRCRE-DUP

Date Sampled...: 12/17/09 11:40 Date Received..: 12/18/09

**% Moisture....:** 12

 PARAM RESULT
 RESULT
 UNITS
 RPD
 METHOD
 ANALYSIS DATE
 BATCH #

 Percent Solids
 87.9
 88.7
 %
 0.94
 (0-20)
 MCAWW 160.3 MOD
 12/22-12/23/09
 9356363

Dilution Factor: 1

Chain of Custody Record
TestAmerica Laboratory location: North Cauton

12/18/69 1100	Company:	7	11:00 Received in Laboratory 6)	Date/Finne: -01/11:00	Company:	Religious A. Donier
Date/Fime:	Company:		Received by:	Date/Time:	Сотралу:	Relinquistred by:
Date/Time:	Company:		Received by:	Date/Time:	Company:	Relinquished by:
	or Months	Disposal By Lab Archive For	Sample Disposal ( A ree may be alreed [2]) C	Unknown	Skin Irritant Poissin B	Positile Lazard Identification    Non-Hazard   Flatimable
	メメメ	のイメメメ		*	V 15:20	HB-075 (2-41)
	メメメ	・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・	×		15:70	HB-075(0-21)
	メメメ	6 X X X X	<b>X</b>	<b>X</b>	12:50	HB-073 (0-2')
	××××	らメメメメ	×	<b>X</b>	12:30	146-072 (4-6')
OWLY	メメメ	Gメメメメ	×		12:20	HB-072 (2-4')
Cd Cx, Pb	メメメ	6 × × × ×	×	×	11:40	HB-071 (16-181)
include: As,	メメメメ	8 X X X X	×	~	1 11:40	HB-071 (10-121)
*Mutuals to	** **	6メメメ ス	×	×	12/17/09 11:46	HB-071 (8-10')
Sample Specific Notes / Special Instructions:	Tot To To M	Coy, You Syl TPt	H2SO4 HN03 HCI NaOH ZmAc/ NaOH Unpres Other:	Air Aqueous Sediment Solid Other:	Sample Date Sample Time	Sample Identification
# 1	ava al tal	(USIV CS CS (f - A	Martine and the state of the st			F0#
Harris Maria	lent Sul- Cyco (SX	why 826-82-	2 days		Shipping Tracking No:	Project Nambor:
	Cr fide wdi	(pH),0 70 fracs	2 weeks			COMMUNICITY Rd.
		)	TAT if different from below			(440) 357-1260
・	Analyses		· · · · · · · · · · · · · · · · · · ·	Wenv.com		Mentar ONID 44DEO
of 1 cocs	966-8296	(30) Abb-	Telephone:	00		6115 Heisley Road
COC No:	Nade Pietras	Nax F	Site Contact:		Down Wetzel	MA MAZY
TestAmerica Laboratories, Inc.	Te					Client Contact
STATE OF THE PERSON OF THE PER		Some Of VRP	NPDES RCRA	Wt₫	Regulatory program:	

TAL-0018 (1008)

gizzok, Toerémeksa Luborrelovine, Inc. All rights recerved. Toerémekoz & Deelge <sup>he</sup> ura trademarko of Toerémexico Luterrelovine. Inc.

TestAmerica Cooler	Receipt Form/Narrative L	ot Number: <u>A9L 8049</u> 2		
North Canton Facilit	********** * * * **********************			
Client H2W	Project	By: alana Margu		
	12/18/09 Opened on 12/18/07	(Signature)		
FedEx ☐ UPS ☐ DHL	☐ FAS ☐ Stetson ☐ Client Drop Off ☒ TestAn	nerica Courier Dother		
		Client Cooler 💢 Other		
1. Were custody seals o	n the outside of the cooler(s)? Yes 🔲 No 💢 🔝 Ir	ntact? Yes 🖺 No 🔲 NA 💢		
If YES, Quantity	Quantity Unsalvageable			
	n the outside of cooler(s) signed and dated?	Yes 🔲 No 🔲 NA 🔼		
Were custody seals o		Yes □ No 🏻		
if YES, are there any				
	attached to the cooler(s)?	Yes 🗌 No 🔯		
	company the sample(s)? Yes 🕅 No 🗌	Relinquished by client? Yes 📈 No 🗌		
4. Were the custody papers signed in the appropriate place?  Yes X No   Yes X No   Other				
5. Packing material used: Bubble Wrap  Foam  None  Other				
6. Cooler temperature upon receiptO °C See back of form for multiple coolers/temps				
METHOD: IR 🐧 Other 🗌				
	· • · · · · · · · · · · · · · · · · · ·	None 🔲		
7. Did all bottles arrive in good condition (Unbroken)?  8. Could all bottle labels be reconciled with the COC?  Yes ☒ No ☐  Yes ☒ No ☐				
		Yes 🛛 No 🗌		
	e correct pH upon receipt?	Yes ☐ No ☐ NA ☒ Yes ☒ No ☐		
10. were correct cone(s) 11. Were air bubbles >6 r	used for the test(s) indicated?			
	elved to perform indicated analyses?	Yes No No NA NA		
	ent in the cooler(s)? Yes 🔲 No 💢 Were VOA			
	Date by			
Concerning	Date by	via verbai [ ] voice iviaii [ ] Otriei [ ]		
14. CHAIN OF CUSTOD	·			
The following discrepance				
The relieving discrepance	oo oodan oo.			
<del> </del>				
15. SAMPLE CONDITION				
Sample(s)	were received after the i	recommended holding time had expired.		
Sample(s)		were received in a broken container.		
Sample(s)		bubble >6 mm in diameter. (Notify PM)		
16 SAMPLE PRESERV				
Sample(s)		ere further preserved in Sample		
	mended pH level(s). Nitric Acid Lot# 082509-HNO3, Sul DH; Hydrochloric Acid Lot# 092006-HCl; Sodium Hydroxia			
	on; nyarochione Ada Lot# 092006-HCI; Sodium nyaroxid at time was preservative added to sample(s)?	e dud Suc weatata Fot# 1001/0-		
Client ID	pH	Date Initials		
<u> </u>	P1	RWZ Unique		
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TestAmerica Cooler Rec	eipt Form/Narrative		
North Canton Facility Client ID	<u>pH</u>	Date	Initials
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			<del> </del>
			<del> </del>
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	•		
			-
Cooler #	Temp. °C	Method	Coolant
Codiei #			
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	mental hard and an analysis of the state of	PARTIE	
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	<del> </del>		<del> </del>



# END OF REPORT

## PHASE II ENVIRONMENTAL SITE ASSESSMENT

INNERBELT STUDY CLEVELAND, OHIO

CUY-CLEVELAND INNERBELT CORRIDOR PID NO. 77510



Prepared for

The Ohio Department of Transportation District 12 5500 Transportation Boulevard Garfield Heights, Ohio 44125

January 17, 2007



1375 Euclid Avenue Suite 600 Cleveland, Ohio 44115 216-622-2400

Project No. 15016633

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### **EXECUTIVE SUMMARY**

#### PROJECT DESCRIPTION

URS Corporation was retained by the Ohio Department of Transportation (ODOT) to conduct Phase II Environmental Site Assessments (ESAs) for select properties within the Cleveland Innerbelt Study. The Cleveland Innerbelt Study begins at the Interstate 71 (I-71) and Fulton Road interchange and extends north through the merge with Interstate 90 (I-90) across the Central Viaduct Bridge, which spans the Cuyahoga River, through the Central Interchange and terminates at the I-90 curve just east of the I-90 merge with State Route (SR) 2 (the Project Area). In addition, the Cleveland Innerbelt Study includes the portion of Interstate 77 (I-77) that begins at the Central Interchange and extends south to the I-77 and East 30<sup>th</sup> Street interchange. The Project Area, which is wholly located in the City of Cleveland, Cuyahoga County, spans approximately seven miles.

To determine the potential for encountering hazardous substances during the project, URS conducted an Environmental Site Assessment (ESA) Screening of the entire Project Area and Phase I ESAs of select properties within the Project Area. Based on the review of these documents, ODOT's Office of Environmental Services (OES) indicated further investigation via a Phase II ESA was warranted for twenty-four properties. A copy of the ODOT-OES' Inter-Office Communication is included in Appendix A. A Project Area and Phase II Site Location Map is included as Figure 1-1.

The Phase II ESA was conducted in accordance with ODOT Environmental Site Assessment Guidelines (ODOT, 1999). URS advanced up to eight soil borings/monitoring wells at each Site and submitted a soil sample from each boring for laboratory analysis. Soil data was compared to Ohio Environmental Protection Agency's (OEPA) Voluntary Action Program (VAP) generic standards for direct contact and the Bureau of Underground Storage Tanks Regulations (BUSTR) standards. Boring Location Maps are included as **Figures 6-1A through 6-24A**.

#### **CONCLUSIONS**

This Phase II ESA was conducted on 24 parcels of land located within the Cleveland Innerbelt Study Project Area. The Phase II ESA sites were selected because of known current or historic operations; known current or historic petroleum product and/or hazardous substance consumption, storage or use; or, the site is proposed for acquisition.

The objective of the Phase II ESA was to evaluate subsurface conditions relative to the current and former uses of the sites in the Project Area. The soil borings and monitoring wells were located in accessible areas of the site. The Project Area, which is wholly located in the City of Cleveland, Cuyahoga County, spans approximately seven miles.

Based on the analytical results, 14 sites listed on Table ES-1, Phase II Site Summary, have impacted soil and/or groundwater that exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels. Two sites did not have Phase II sampling conducted but are recommende to have Construction Plan Notes prepared as a result of past history. The remaining eight sites did not display contamination levels exceeding applicable standards.

	$\mathbf{Ph}$	Table ES-1 ase II Site Conclusion Sumn	nary
Site Number	Site	Address	Conclusion
2	Former Bauer Auto	3553 West 25 <sup>th</sup> Street	Plan Note Recommended
13	Former Glove Cleaning Service/Scranton Averell	2132-2150 West 15 <sup>th</sup> Street	No exceedances of applicable standards.
14	Bojacks Meats	2000 W. 14 <sup>th</sup> /1425 University	Impacted Groundwater
15	Leon Rudnick	1402-1408 Abbey Road	Plan Note Recommended
16	Wendell & Carroll Collins/1501 Companies	West 15 <sup>th</sup> Street	No exceedances of applicable standards.
17	Terminal Oil	308 Central Viaduct	Impacted Soil & Groundwater
18	Cleveland Fire Station	310 Carnegie	No exceedances of applicable standards.
19	Gillota Fuel Products	206-300 Central Viaduct	Impacted Soil & Groundwater
20	Earl Lee	2394 Canal Road	Impacted Soil & Groundwater
21	White Properties	1996 West 3 <sup>rd</sup> Street	Potential USTs on Site
22	Nova Properties	West 4 <sup>th</sup> Street	Impacted Soil & Groundwater
23	NS Railroad Building	840 Minkon Lane	Impacted Soil
24	James Vincent	2515 Canal Road	Impacted Soil
27	Meridian Properties/ Independent Towel	1802 Central Avenue	Potential USTs on Site and Impacted Groundwater
29	BP Gas Station	2701 Chester Avenue	Impacted Groundwater
33	State Industrial Products	3100 Hamilton Avenue	Impacted Groundwater
34	Former Teledyne Metal Finishing	1725 East 27th Street	Impacted Groundwater
42	CB Realty	2975 Superior Avenue	No exceedances of applicable standards.
43	KNC Building	2635 Payne Avenue	No exceedances of applicable standards.

## **EXECUTIVE SUMMARY**

	Ph	Table ES-1 ase II Site Conclusion Sumr	nary
Site Number	Site	Address	Conclusion
45	Harold Moss, Trustee	1748 East 27th Street	No exceedances of applicable standards.
51	Temp Craft Plastics	3960 South Marginal Road	Impacted Soil & Groundwater
53	Cleveland Fire Academy	3101 Lakeside Avenue	No exceedances of applicable standards.
57	Charles Martin	3501 Croton Avenue	No exceedances of applicable standards.
59	Parking Lot	Central Viaduct	Impacted Groundwater

#### 1.1 OBJECTIVE

The Cleveland Innerbelt Study has been undertaken by ODOT for the purpose of developing a strategy for the renewal of the transportation infrastructure in the I-71/I-90 corridor through downtown Cleveland. A Project Location Map is provided as **Figure 1-1**.

The pavements and structures comprising the existing Innerbelt through Cleveland are approaching the end of their service life. Further, the capacity of the existing Innerbelt is restricted by a series of interchanges and subsequent bottlenecks, which do not conform to current highway design standards, thereby contributing to traffic congestion, high accident rates, and traffic diversions through central city neighborhoods, which degrades the quality of life for Cleveland residents.

The objective of the Phase II ESA was to evaluate subsurface conditions relative to the current and former uses of the sites in the Project Area. The soil borings and monitoring wells were located in accessible areas of the site. The soil boring and monitoring well locations are depicted on **Figures 6-1A** through **6-24A**. The Project Area, which is wholly located in the City of Cleveland, Cuyahoga County, spans approximately seven miles.

#### 1.2 PROJECT DATES

The Phase I ESA was completed in December 2005. A Work Plan was developed and approved by ODOT in March 2006. The Phase II field activities began in July 2006. Analytical data review, the required Phase II determinations, and supporting documentation generated, were completed in December 2006. This report was finalized on December 31, 2006.

#### 1.3 REPORT ORGANIZATION

This report is organized into eight sections, including this Introduction. Sections 2.0 and 3.0 describe the Project Area background and physical setting. Section 4.0 summarizes the field investigation conducted in support of this Phase II. Sections 5.0 and 6.0 provide summary and interpretation of the collected data and are presented also on a site-by-site basis. The Phase II conclusions are included in Section 7.0. Section 8.0 provides a list of references used in the preparation of this report.

#### 2.1 SUMMARY OF PHASE I

The Environmental Site Assessment Screening and associated Phase I ESAs were conducted in accordance with the ODOT guidance. The Project Area encompasses commercial, industrial, institutional, municipal, and residential properties. The Screening process consisted of historical research, regulatory records review, and a Project Area reconnaissance. As a result of the Screening, the 33 properties listed below were recommended for Phase I ESAs.

	Pha	Table 2-1 se I ESA Summary	
Site Number	Site	Address	Rationale
2	Former Bauer Auto	3553 West 25th Street	LUST
	CENT	RAL VIADUCT - NORTH	
11	Stripmatic	1501 Abbey Road	LUST
13	Former Glove Cleaning Service/Scranton Averell	2132-2150 West 15th Street	Historic Operations
14	Bojacks Meats	1425 University/2000 West 14 <sup>th</sup> Street	Property Take
15	Leon Rudnick	1402-1408 Abbey Road	Property Take
16	Wendell & Carroll Collins/1501 Companies	West 15th Street	Property Take
17	Terminal Oil	308 Central Viaduct	UST
18	Cleveland Fire Station	310 Carnegie	Property Take
19	Gillota Fuel Products	206-300 Central Viaduct	Property Take
20	Earl Lee	2394 Canal Road	Property Take
21	White Properties	1996 West 3rd Street	Property Take
22	Nova Properties	West 4th Street	Property Take
23	NS Railroad Building	840 Minkon Lane	Property Take
24	James Vincent	2515 Canal Road	Property Take
	CEI	NTRAL INTERCHANGE	
27	Meridian Properties/Independent Towel	1802 Central Avenue	RCRA SQG, LUST
57	Charles Martin	3501 Croton	Property Take
58	JF Sanson	3561 Burwell	Property Take
<u></u>		TRENCH	
29	BP Gas Station	2701 Chester	UST

	Pha	Table 2-1 se I ESA Summary	
Site Number	Site	Address	Rationale.
33	State Industrial Products/ Information Systems Building	3100 Hamilton	RCRA SQG, LUST, OH Spills, PCBs
34	Former Teledyne Metal Finishing	1725 East 27th Street	RCRA TSD
41	1400 East 30th Partners	1400 East 30th Street	Property Take
42	CB Realty	2975 Superior	Property Take
43	KNC Building	2635 Payne	Property Take
44	EJ Investments	2630 Payne	Property Take
45	Harold Moss, Trustee	1748 East 27th Street	Property Take
46	Tri Building	2728 Euclid	Property Take
47	GKC Limited	2729 Prospect	Property Take
48	Parkwood Corporation	2829 Euclid	Property Take
49	Julius Sorma	3004 St. Clair	Property Take
	I	NNERBELT CURVE	
50	Cleveland Mounted Police	1150 East 38th Street	Property Take
51	Temp Craft Plastics	3960 South Marginal	Manufacturing
52	Architectural Real Estate	3000 Lakeside	Property Take
53	Cleveland Fire Academy	3101 Lakeside	Property Take

The Phase I ESA process conducted for the above-referenced sites consisted of additional historic research, regulatory agency file reviews, a reconnaissance of each site, and property owner interviews. As a result of the Phase I ESAs, 23 sites were recommended for Phase II ESAs. The findings of the Phase I ESA are presented the URS Environmental Site Assessment Screening and Phase I Environmental Site Assessment Report for the Innerbelt Study, dated December 2005.

#### 2.2 WORK PLAN

Per the ODOT Inter-Office Communication (IOC) dated December 13, 2005, 24 sites required further investigation via a Phase II ESA; therefore, a Work Plan was developed, which was approved by ODOT in March 2006.

In general, the Phase II ESA sites were selected because of known current or historic operations; known current or historic petroleum product and/or hazardous substance consumption, storage or use; or, the site is proposed for acquisition. To that end, the chemical parameters proposed for analysis were based on the notion that the sites fall into two categories; known petroleum product storage sites and other sites. In addition, based on conversations with ODOT Office of

Environmental Services personnel, monitoring wells were proposed for all potential acquisition sites. A master listing of Phase II sites is presented in Table 2-2 below.

Per the ODOT IOC dated February 23, 2006, the following sites were not investigated via a Phase II ESA; Site #2 - Former Bauer Auto and Site #15 - Leon Rudnick. Plan Notes will be developed for these sites. Site 59 was added per the IOC. The ODOT IOC is provided in **Appendix A**.

		Table 2-2 Phase II Master List	
Site Number	Site	Address.	Rationale
2	Former Bauer Auto	3553 West 25 <sup>th</sup> Street	LUST Site.
13	Former Glove Cleaning Service/Scranton Averell	2132-2150 West 15 <sup>th</sup> Street	LUST Site and Suspected Dumping.
14	Bojacks Meats	2000 W. 14 <sup>th</sup> /1425 University	UST Site and Chemical Use
15	Leon Rudnick	1402-1408 Abbey Road	Historic UST Site
16	Wendell & Carroll Collins/1501 Companies	West 15 <sup>th</sup> Street	Property Take
17	Terminal Oil	308 Central Viaduct	LUST Site, ASTs and Drum Storage.
18	Cleveland Fire Station	310 Carnegie	LUST Site.
19	Gillota Fuel Products	206-300 Central Viaduct	LUST and UST Site.
20	Earl Lee	2394 Canal Road	Historic UST Site and Chemical Use
21	White Properties	1996 West 3 <sup>rd</sup> Street	Historic UST Site.
22	Nova Properties	West 4 <sup>th</sup> Street	Historic UST and OHSPILLS Site, Asphalt Plant.
23	NS Railroad Building	840 Minkon Lane	Active UST Site and Waste Storage.
24	James Vincent	2515 Canal Road	Historic Operations
27	Meridian Properties/ Independent Towel	1802 Central Avenue	Historic UST and LUST Site.
29	BP Gas Station	2701 Chester Avenue	UST Site.
33	State Industrial Products	3100 Hamilton Avenue	OHSPILLS, LUST and UST Site.
34	Former Teledyne Metal Finishing	1725 East 27th Street	Historic UST and AST Site, Chemical Use.

		Table 2-2 Phase II Master List	
Site Number	Site	Address	Rationale
42	CB Realty	2975 Superior Avenue	Historic UST Site.
43	KNC Building	2635 Payne Avenue	Historic UST Site.
45	Harold Moss, Trustee	1748 East 27th Street	Historic UST and Dump Site.
51	Temp Craft Plastics	3960 South Marginal Road	Historic UST Site and Chemical Use.
53	Cleveland Fire Academy	3101 Lakeside Avenue	Historic UST and AST Site.
57	Charles Martin	3501 Croton Avenue	Property Take
59	Parking Lot	Central Viaduct	Historic UST Site (per ODOT)

Descriptions of the physical setting across the Project Area are presented in this section. Information is provided regarding topography, surface water drainage, regional geology and hydrogeology, and geologic and hydrogeologic conditions encountered during the Phase II ESA activities.

### 3.1 TOPOGRAPHY AND DRAINAGE

According to the 1994 Cleveland North and Cleveland South, Ohio, USGS 7.5 Minute Series Quadrangle Map, the Project Area elevations vary from approximately 680 feet above mean sea level (msl) for the majority of the Corridor to approximately 560 feet above msl in the Industrial Valley (USGS).

In heavily urbanized areas, the flow of surface water is predominantly controlled by the facility's and/or municipality's stormwater system. Since several catch basins were observed at the Project Area sites, surface water collected in sewer lines beneath the various sites likely flows into the public combined sanitary and storm water sewer system.

#### 3.2 GENERAL LAND USE

The Cleveland Innerbelt Study is located in a highly urbanized area. Land usage ranges from high density residential to predominantly industrial and commercial.

### 3.3 REGIONAL GEOLOGY

The Project Area is located in the Erie Lake Plain Physiographic Region of the Interior Low Plateau Physiographic Province in Ohio (Brockman). The Erie Lake Plain is characterized by ground surface that slopes gently toward Lake Erie and is dissected by re-entrant valleys, such as the Cuyahoga River Valley, draining northward to Lake Erie (Ford, 1987).

The Erie Lake Plain in this portion of Cleveland is underlain by subsurface soils consisting primarily of a considerable thickness of interbedded fine sand, silt, and clay deposits. The unconsolidated sediments are underlain by the Devonian-aged Ohio Shale Formation (ODNR, 1996a).

According to the Soil Survey of Cuyahoga County (USDA), the project area is classified as Urban Land. The Urban Land designation indicates that 80 percent of the surface is covered by manmade materials (e.g., buildings, concrete, and asphalt). The Erie Lake Plain in this portion of Cleveland is underlain by subsurface soils consisting primarily of Pleistocene to recent-aged sand and gravel. Surficial materials beneath the Property and vicinity likely consist of "made land" (e.g., fill material). The unconsolidated sediments are underlain by the Devonian-aged Ohio Shale Formation.

According to ODNR's Groundwater Pollution Potential Map of Cuyahoga County, the Project Area is located within an area of buried valley deposits. This area is characterized by thick deposits of sand and gravel that have been deposited in a former topographic low (a pre-glacial or

inter-glacial river valley) by glacial melt waters. Glacial till or recent alluvium often overlies that buried valley (ODNR, 1994).

#### 3.4 REGIONAL HYDROGEOLOGY

Potable water for the area surrounding the Project Area is obtained from Lake Erie and distributed by the City of Cleveland. Based on the availability of municipally supplied potable water, groundwater resources in the area are not likely to be developed for future potable use.

According to ODNR's Groundwater Resources Map of Cuyahoga County, the Project Area is located in an area characterized by buried valley which contains 200 to 300 feet of fine sand, silt, and clay. Drilled wells yield meager supplies unless encountering thin, isolated sand and gravel lenses (ODNR, 1979). Lake Erie is north of the Project Area. Based upon the local topography of the area, groundwater flow in the vicinity of the Project Area is assumed to be to the north, toward Lake Erie.

A groundwater well is located approximately 1½ mile southeast of the Project Area. This well was advanced approximately 179 feet into the sand and gravel aquifer. The production for this well is reported as approximately 250 gallon per minute.

The objective of the Phase II ESA was to evaluate subsurface conditions relative to the current and former uses of the sites in the Project Area.

#### 4.1 SITE RECONNAISSANCE

A site reconnaissance visit was conducted prior to initiating field activities. Site reconnaissance tasks included an assessment of proposed boring locations relative to potential obstructions and aboveground utilities, as well as an evaluation of these locations relative to site features (i.e. UST cavities, staining, etc.).

The location of each proposed boring or monitoring well was evaluated relative to subsurface and aboveground utilities. A minimum of two days before the start of boring activities, the Ohio Utilities Protection Service (OUPS) was contacted, by calling 800-362-2764 so that OUPS member utilities had the opportunity to mark their respective utility lines. Property owners were also interviewed concerning the locations of underground utilities, if possible.

The selected drilling subcontractor obtained and paid for all permits, applications, and other documents required by state and local authorities.

Following the completion of field activities, each site was restored as nearly as possible to its pre-construction condition. Unused or surplus materials, supplies, and waste material were removed from each sample location as the work is completed at that area. Stakes or flagging of monitoring wells, which were not located on concrete, remained near the monitoring wells for ease of locating during each sampling event.

A summary of proposed field activities is presented in Table 4-1, Proposed Phase II ESA Sampling.

### 4.2 SURFACE SOIL SAMPLING

Surface soil samples were collected at one site, as indicated in **Table 4-1**. The samples were collected from 0 to 2 feet below ground surface (bgs). The samples were collected via a sampling shovel, which was decontaminated between sample locations to minimize the potential for cross-contamination.

## 4.3 BOREHOLE AND MONITORING WELL INSTALLATION

Monitoring wells were installed in soil borings that were advanced using 4½-inch inner diameter (I.D.) hollow stem augers. During advancement of the soil boring, subsurface samples were continuously collected with a 2-foot long, 2-inch diameter split-spoon sampler in accordance with ASTM-D-1586 Standard Test Method for Penetration Test and Split-Barrel Sampling of Soils. Once sampling was initiated, the number of blows was recorded per six inches. The borings were advanced 5 to 10 feet into the first water-bearing zone encountered.

The split-spoon sampler was brought to the surface and opened. Soil samples were described in accordance with ASTM D-2488-90 Standard Practice for Description and Identification of Soils.

USCS group symbols were also used to describe soil samples. Special attention was given to zones where there was a high potential for contaminant transport (i.e., discoloration, silt zones). Information recorded included the borehole location, drilling information, sample description and sampling information such as sample intervals, recovery, and blow counts.

Once the soil boring was advanced to the terminal depth, monitoring well installation commenced. The monitoring well were installed in accordance with the Ohio Environmental Protection Agency's (OEPA) Technical Guidance Manual for Hydrogeologic Investigations and Groundwater Monitoring Programs and the Technical Guidelines for Well Construction and Ground Water Protection prepared by the State Coordinating Committee on Groundwater. The monitoring well was installed such that the screen intersected the top of the water-bearing unit so that seasonal water table fluctuations would not cause water levels to rise above or fall below the screened interval. The monitoring well was not be screened across more than one water-bearing unit.

The monitoring well casings consisted of 2-inch ID, Schedule 40, polyvinyl chloride (PVC) pipe with flush mount, threaded joints. The well screen consisted of a 10-foot section of 0.010-inch hermetically sealed, factory slotted screen, which would prevent 90 percent of the filter pack from entering the well. A flush threaded bottom plug was installed at the base of the screen as well. The annular space around the screened interval was filled with a 100 percent silica sand filter pack. The filter pack consisted of clean, sorted, well-rounded, acid-resistant, 100 percent silica sand that contains less than two percent flat particles. The filter pack was certified free of contaminants by the vendor. The filter pack consisted of Global # 7 sand, and extended at least two feet above the screened section within the borehole. The top of the sand pack was sounded to verify its depth during placement. Additional filter pack was placed as required to return the level of the pack to the top of the screen. The well was sounded until two feet of sand was measured above the top of the screen.

Once the sandpack was emplaced, a bentonite seal was installed to a minimum depth of two feet above the top of the sandpack and consisted of 100% sodium bentonite chips with a minimum dry bulk density of 70 pounds per cubic foot. The top of the seal was measured with a weighted tape to verify seal thickness. The remainder of the annulus was filled to the surface using a 100% sodium bentonite chips with a minimum dry bulk density of 70 pounds per cubic foot.

The monitoring well was completed with flush-mounted surface casings in paved areas and standpipes in unpaved areas. For the flush-mounts, a steel well vault cover was cemented in place over the well riser with a concrete pad at least two feet in diameter. The standpipes were also cemented in place with a concrete pad at least two feet in diameter. The monitoring well was secured as soon as possible after installation. A corrosion-resistant lock was placed on each monitoring well.

### 4.4 SOIL SAMPLING

Upon retrieval of the sampling barrel from the borehole, the lithology of the soils was recorded as described in the preceding sections. Then, a portion of the soil was placed into a resealable

bag and allowed to volatilize for approximately 15 minutes. The remaining portions of the soil were placed into laboratory-provided sampling jars.

After approximately 15 minutes, the tip of a Photo Ionization Detector (PID) pierced the bag and the headspace of the bag was taken. This headspace screening reading was recorded. This procedure was conducted on all the intervals of soils as the soil borings advance, with the exception of soils below the saturated zone.

One soil sample per soil boring and/or monitoring well was submitted for laboratory analyses. This sample was selected based on the headspace screening readings; the sample exhibiting the highest reading was submitted. If no readings are apparent, visual or olfactory indications of contamination were used as a secondary basis for sample selection. If these were absent, the sample was collected from the interval above the saturated zone. In the event groundwater was not encountered, the sample from the terminal depth of the boring was submitted.

#### 4.5 WELL DEVELOPMENT

Following completion of drilling, and no sooner than 24 hours after well installation, the monitoring well was developed by purging or bailing until the discharged water was relatively sediment free and the indicator parameters (pH, temperature, and specific conductance) reached stabilization criteria or until at least 6 well volumes were removed. Developing the well not only removes any sediment but also improves the hydraulic properties of the sand pack. Development procedures utilized are presented below:

- 1) The well was developed using decontaminated bailers or pumps. Care was taken not to introduce the equipment to contaminants during installation.
- Well development commences and continues until a minimum of three well volumes were removed and stabilization criteria were achieved. After each well volume was removed, stabilization criteria and turbidity were measured. Stabilization was achieved when variation in temperature was within ±1°C, pH was within ±0.1 units and electrical conductivity (EC) was within ±5 percent over at least three successive well volumes.
- 3) All measurements, the volume of water removed, and the discharge water color were recorded.

### 4.6 GROUNDWATER ELEVATION MEASUREMENT

Prior to sampling monitoring wells, the groundwater elevations were collected from all wells in the monitoring well network at each site within one calendar day. Measurements were taken after the well has been installed, developed, and the water level had recovered completely. After removal of the well cap, the well was allowed to equilibrate prior to groundwater level measurements.

The thickness of any phase-separated hydrocarbon in the monitoring well was measured with an electronic interface probe. Groundwater levels were measured from the notch located at the top of the well casing to the nearest 0.01-foot. The static water level was measured each time the

well was sampled, before any sampling equipment enters a well. The water levels were recorded for each well.

#### 4.7 SURVEY

URS utilized an area-wide survey of the innerbelt project that was conducted by the URS-Akron office. The Phase II sites were referenced into the survey and no site-specific survey data was collected for the Phase II inspections.

All surveying locations were reported as the distance in feet. The elevation of monitoring wells was surveyed comparative to ground surface elevation.

### 4.8 GROUNDWATER SAMPLING

Groundwater samples were collected using a submersible pump with a modified low flow (500 ml). When numerous monitoring wells were to be sampled in succession, those wells expected to have low levels of contamination or no contamination were sampled prior to those wells expected to have higher levels of contamination. This practice helps reduce the potential for cross contamination between wells. All sampling data was recorded.

The temperature, pH, EC, and turbidity of the purged water was measured and recorded. The sample was collected after the above parameters had stabilized. Stabilization was defined as three consecutive measurements of the parameters measured within these ranges: temperature (± 2°C), pH (± 0.1 SU), electrical conductivity (± 3 percent), and turbidity within 10 percent. Although turbidity readings should be at 10 NTUs or lower at the completion of purging, a maximum of six well volumes was purged prior to collecting the sample.

### 4.9 SAMPLING HANDLING

Samples were not be numbered contiguously for the Project; but contiguously for each Site. For example, a soil sample was identified by the Site Number, whether it came from a surface sample (SS), soil boring (SB), or monitoring well (MW), the number of that location, and the depth. Sample 57-MW2-0002 indicates the sample was collected at Site 57, Monitoring Well #2, from 0 to 2 feet bgs. A groundwater sample was identified by the Site Number, the monitoring well number and the date collected. Sample 57-MW2-021506 indicates the sample was collected at Site 57, Monitoring Well #2, on February 15, 2006.

Samples collected for quality assurance/quality control purposes were labeled as such. Field duplicates/replicates were labeled as described in the previous paragraph; however, a D was added to the end of the sample identification. Sample 57-MW2-0002D indicates the sample was collected at Site 57, Monitoring Well #2, from 0 to 2 feet bgs and was a duplicate sample.

Matrix spikes/matrix spike duplicates were labeled as described in the previous paragraph; however, a MS/MSD was added to the end of the sample identification. Sample 57-MW2-021506MS/MSD indicates the sample was collected at Site 57, Monitoring Well #2, on February 15, 2006 and was a matrix spike/matrix spike duplicate sample.

Trip blanks and equipment blanks were labeled as such and also referenced the date collected. Field Blank 021306 indicates a field blank was collected on February 13, 2006.

Immediately after the samples were labeled for off-site laboratory analysis, each sample bottle was sealed in a plastic bag and wrapped with shock-absorbent materials, such as bubble wrap, to prevent movement of sample containers during transport. Then, the samples were placed in a sturdy ice chest (cooler). The cooler was packed with resealable double-bagged ice packs and sealed with packaging tape.

The chain-of-custody sample log sheet(s) were filled out in indelible ink, placed in a resealable plastic bag, and taped to the inside lid of the cooler. Each sample contained in the cooler was specified on the chain-of-custody records by the sampling identification number. Sample containers were packaged to minimize potential breakage and to comply with Department of Transportation (DOT) requirements.

#### 4.10 SAMPLE ANALYSIS

The soils from sites with a history of petroleum product consumption, storage or use were analyzed for some and/or all of the following: volatile organic compounds (VOCs) via EPA Method SW-846 8260, semivolatile organic compounds (SVOCs) via EPA Method SW-846 8270, polyaromatic hydrocarbons (PAHs) via EPA Method SW-836 8310, total petroleum hydrocarbons (TPH) gasoline range organics (GRO) and diesel range organics (DRO) via EPA Method SW-846 8015, cyanide via EPA Method SW-846 9010/9012 and/or Resource Conservation and Recovery Act (RCRA) Metals via EPA Method SW-846 6010 and 7071.

The soils from sites without this history were analyzed for VOCs via EPA Method SW-846 8260, SVOCs via EPA Method SW-846 8270, PAHs via EPA Method SW-846 8310, cyanide via EPA Method SW-846 9010/9012 and/or RCRA Metals via EPA Methods SW-846 6010 and 7071.

The groundwater samples were analyzed for VOCs via EPA Method SW-846 8260, SVOCs via EPA Method SW-846 8270, PAHs via EPA Method SW-846 8310, and/or RCRA Metals via EPA Methods SW-846 6010 and 7071.

These analyses are further detailed in Table 4-1.

### 4.11 QUALITY ASSURANCE/QUALITY CONTROL

With the exception of the water level indicator and the interface probe, field instruments were calibrated at least once per day at the beginning of the day's activities.

Precision, accuracy, and potential contamination were measured in the field through the use of Equipment Blanks (drilling equipment only), Field Duplicates/Replicates, and Matrix Spike/Matrix Spike Duplicates. Equipment Blanks and Matrix Spike/Matrix Spike Duplicates were collected at the rate of one per 20 samples. Field Duplicates/Replicates were collected at the rate of one per 20 samples. Trip blanks were submitted daily; each cooler containing soil or groundwater samples for VOCs analysis had a trip blank.

#### 4.12 FIELD LOGS

During the advancement of the borings and monitoring wells, observations regarding lithology, potential contamination, headspace screening readings, and other pertinent information were recorded using electronic logging software. The units were downloaded at the end of each field day at the URS office in Cleveland, Ohio.

#### 4.13 DECONTAMINATION PROCEDURES

All equipment that directly or indirectly contacts samples was decontaminated before and after each use. Decontamination consisted of varying combinations of high-pressure hot water rinse, Liquinox® or Alconox® wash, and potable water rinse. Decontamination water was contained in 55-gallon drums and disposed properly, as discussed in **Section 4.14**.

Drilling, sampling, monitoring well installation and other equipment was decontaminated using the following procedures:

• Drill rig augers, drill rods, bits, etc. were steam cleaned before use and between borings.

• Soil and shallow groundwater sampling equipment was cleaned with an Alconox® or Liquinox® and potable water solution and rinsed with potable water.

• Electronic water level sounders and water quality probes were cleaned with an Alconox® or Liquinox® and potable water solution and rinsed with potable water.

The following procedures were used to decontaminate large pieces of equipment, such as casings, auger flights, pipe and rods. The external surfaces of equipment were washed with high-pressure hot water and Liquinox<sup>®</sup>, or equivalent laboratory-grade detergent, and if necessary, scrubbed until all visible dirt, grime, grease, oil, loose paint and/or rust flakes had been removed. The equipment was then be rinsed with potable water. The inside surfaces of casing, drill rod, and auger flights were also washed as described.

The following procedure was used to decontaminate sampling and drilling devices, such as split spoons, bailers, and augers, which could be hand-manipulated. For sampling and smaller drilling devices, the equipment was scrubbed with a solution of potable water and Liquinox<sup>®</sup>, or equivalent laboratory-grade detergent, and rinsed with potable water.

### 4.14 WASTE HANDLING AND DISPOSAL

Soil cuttings generated from boring installation and water generated from well development and purging were contained separately in 55-gallon drums. Decontamination water also was containerized in 55-gallon drums. The drums were labeled using a paint pen with date, project, boring or well designation, and type of waste (i.e., soil cuttings, purge water). In addition, non-hazardous drum labels were completed and affixed to each drum. All drums were staged in a selected location on each site. Disposal options were presented to ODOT following receipt of laboratory results.

Based on the analytical data received and the results of this assessment, soil and groundwater required special management. On November 13 through November 15, 2006, Enviroserve, of Cleveland, Ohio, removed 145 55-gallon drums containing the materials for off-site disposal. Copies of the disposal documentation are provided in **Appendix C**.

#### 4.15 GEOPHYSICAL SURVEY

In addition to subsurface investigations, a magnetometer survey was determined to be the most applicable geophysical survey technique for the Project. It is assumed that abandoned or orphaned USTs associated with these Sites likely were constructed of steel. The magnetometer survey relies on the ability of the equipment to detect a contrast in the magnetic properties of the target (in this case the steel USTs) and the surrounding soil. This contrast would be present if a UST constructed of ferrous (iron-containing) material were present. The magnetometer is insensitive to the fiberglass material used in present day petroleum USTs.

The survey was conducted using a Geometrics G-858 magnetometer operated with a two vertically oriented magnetometer sensors to collect gradient data. In this configuration, the magnetometer is capable of detecting small ferrous targets, such as a single drum to a depth of 18 feet or a UST at greater depths. Measurements were collected at 1.5-second intervals as the operator walks along parallel survey lines spaced 5-foot on center. Following completion of the survey, the data was transferred to a computer file for reduction.

Data reduction was conducted using software developed by Geometrics. The reduced data was plotted using color to denote areas of high and low magnetic gradients. Areas of high gradient would lie directly over ferrous objects. The color gradient plots then were compared to a site plan and field notes in an effort to correlate observed surface features to areas of high magnetic gradient.

The geophysical survey sites are identified on Table 4-1.

TABLES

Table-4-1 ODOT – Cleveland Innerbelt Study Proposed Phase II ESA Sampling

Site	Site	Address	Geophysical Survey	Surface Soils	Monitoring Wells	Samples/Analyses
2	Former Bauer Auto	3553 West 25 <sup>th</sup> Street	N/A	0	0	PLAN NOTE
13	Former Glove Cleaning Service/Scranton Averell	2132-2150 West 15 <sup>th</sup> Street	Yes	0	7	7 soil samples;1 per boring/VOCs, and SVOCs 7 groundwater samples/VOCs and SVOCs
14	Bojacks Meats	2000 W. 14 <sup>th</sup> /1425 University	Yes	0	9	6 soil samples; 1 per boring/VOCs, SVOCs, TPH, and Cyanide 6 groundwater samples/VOCs, SVOCs and Cyanide
15	Leon Rudnick	1402-1408 Abbey Road	Yes	0	0	PLAN NOTE
16	Wendell & Carroll Collins/1501 Companies	West 15 <sup>th</sup> Street	N/A	0	4	4 soil samples; 1 per boring/VOCs, SVOCs and RCRA Metals 4 groundwater samples/VOCs, SVOCs and RCRA Metals
17	Terminal Oil	308 Central Viaduct	Yes	0	4	4 soil samples; 1 per boring/VOCs, PAHs and TPH 4 groundwater samples/VOCs and PAHs
18	Cleveland Fire Station	310 Carnegie	Yes	0	3	3 soil samples;1 per boring/VOCs, PAHs and TPH 3 groundwater samples/VOCs and PAHs
19	Gillota Fuel Products	206-300 Central Viaduct	Yes	0	3	3 soil samples;1 per boring/VOCs, PAHs and TPH 3 groundwater samples/VOCs and PAHs
20	Earl Lee	2394 Canal Road	N/A	0	4	4 soil samples;1 per boring/VOCs, SVOCs, TPH, and RCRA Metals 4 groundwater samples/VOCs, SVOCs and RCRA Metals

Tabic=4-1 ODOT - Cleveland Innerbelt Study Proposed Phase II ESA Sampling

Site	Site	Address	Geophysical Survey	Surface Soils	Monitoring Wells	Samples/Analyses
21	White Properties	1996 West 3 <sup>rd</sup> Street	Yes	0	4	4 soil samples;1 per boring/VOCs, SVOCs, TPH, and RCRA Metals 4 groundwater samples/VOCs, SVOCs and RCRA Metals
22	Nova Properties	West 4 <sup>th</sup> Street	Yes	0	S	5 soil samples; 1 per boring/VOCs, SVOCs, TPH, and RCRA Metals 5 groundwater samples/VOCs, SVOCs and RCRA Metals
23	NS Railroad Building	840 Minkon Lane	Yes	9	4	6 surface soil samples/VOCs, SVOCs, TPH, and RCRA Metals 4 soil samples;1 per boring/VOCs, SVOCs, TPH, and RCRA Metals 4 groundwater samples/VOCs, SVOCs and RCRA Metals
24	James Vincent	2515 Canal Road	N/A	0	8	3 soil samples; 1 per boring/VOCs and SVOCs 3 groundwater samples/VOCs and SVOCs
27	Meridian Properties/ Independent Towel	1802 Central Avenue	Yes	0	4	4 soil samples;1 per boring/VOCs, SVOCs, TPH, and RCRA Metals 4 groundwater samples/VOCs, SVOCs and RCRA Metals
29	BP Gas Station	2701 Chester Avenue	N/A	0	4	4 soil samples;1 per boring/VOCs, PAHs and TPH 4 groundwater samples/VOCs and PAHs
33	State Industrial Products	3100 Hamilton Avenue	Yes	0	10	10 soil samples;1 per boring/VOCs, SVOCs, TPH, and RCRA Metals (TCL/TAL)  10 groundwater samples/VOCs, SVOCs and RCRA Metals (TCL/TAL)

Table 7-1 ODOT – Cleveland Innerbelt Study Proposed Phase II ESA Sampling

Site	Site	Address	Geophysical	Surface Soils	Monitoring Wells	Samples/Analyses
34	Former Teledyne Metal Finishing	1725 East 27th Street	No	0	3	3 soil samples; 1 per boring/VOCs, SVOCs, TPH, and RCRA Metals  3 groundwater samples/VOCs, SVOCs and RCRA Metals
42	CB Realty	2975 Superior Avenue	N/A	0	3	3 soil samples; 1 per boring/VOCs, SVOCs, TPH, and RCRA Metals  3 groundwater samples/VOCs, SVOCs and RCRA Metals
43	KNC Building	2635 Payne Avenue	Yes	0	3	3 soil samples;1 per boring/VOCs, PAHs and TPH 3 groundwater samples/VOCs and PAHs
45	Harold Moss, Trustee	1748 East 27th Street	Yes	0	3	3 soil samples; 1 per boring/VOCs, PAHs, TPH, and RCRA Metals 3 groundwater samples/VOCs, PAHs and RCRA Metals
51	Temp Craft Plastics	3960 South Marginal Road	N/A	0	4	4 soil samples; I per boring/VOCs and SVOCs 4 groundwater samples/VOCs and SVOCs
53	Cleveland Fire Academy	3101 Lakeside Avenue	N/A	0	4	4 soil samples;1 per boring/VOCs, SVOCs, TPH, and RCRA Metals 4 groundwater samples/VOCs, SVOCs and RCRA Metals
57	Charles Martin	3501 Croton Avenue	N/A	0	3	3 soil samples;1 per boring/VOCs and SVOCs 3 groundwater samples/VOCs and SVOCs
59	Parking Lot	Central Viaduct	Yes	0	E	3 soil samples;1 per boring/VOCs, SVOCs and TPH 3 groundwater samples/VOCs and SVOCs

# 5.1 DATA REVIEW AND LABORATORY CERTIFICATION DOCUMENTATION

All data generated by the analytical laboratory was reviewed by a URS chemist to evaluate and document data quality. A standard review includes assessment of supporting quality control (QC) parameters such as laboratory blank results, laboratory control sample recoveries, and other batch QC results, as well as detection limits, holding times, and information provided in the report narrative. A standard review does not include reconstruction of the analytical data.

The complete data assessment reports for each site, listing all data qualifications, are included in **Appendix B**. **Appendix B** also contains a compact disc (CD) of all the analytical data results, as reported by the laboratory and qualified by the data reviewer. All data was considered usable for decision making purposes.

**SECTIONSIX** 

**Phase II Findings** 

This section of the Phase II describes the analytical results of the sampling conducted during the previously-described field activities. These results are also interpreted and summarized to provide characterization of the Property for subsequent use in quantifying risk and, if needed, determining where remediation is required.

# 6.2 SITE 13 - FORMER GLOVE CLEANING SERVICE/SCANTON AVERELL

According to the Cleveland City Directories, the site was listed as Glove Cleaning Service Company in the 1954 through 1984 Directories. The site was identified as a LUST site in the EDR Report.

According to the Cleveland Fire Prevention Bureau files, the site was occupied by the Glove Cleaning Service Company (2130), Eyre Glove Cleaners (2140) and Indu-Sol Products Inc. (2150) in 1941. An entry in 1947 indicates a 500 to 700-gallon AST of solvent was to be installed at the 2130 West 15<sup>th</sup> Street address. A Fire Inspection Report, dated August 1952, indicates there were four USTs located at the 2130 site; however, the contents and quantities were not provided. A 3,000-gallon UST, which contained #4 fuel oil, was installed at the 2150 site in May 1951. A complaint was received at the Bureau on October 20, 1970; the Glove Cleaning Service Company reportedly was dumping Stoddard solvent in a vacant lot next to 2149 West 15<sup>th</sup> Street.

An application for UST removal was submitted on April 22, 1994. A 500-gallon UST, which contained dry cleaning fluid, was to be removed under the supervision of the Bureau. The UST reportedly was located north of the building. An NFA letter, dated January 30, 1996, was received for the incident. An application for UST removal was submitted on April 29, 1998. Two 4,000-gallon USTs, which contained heating oil, were to be removed under the supervision of the Bureau. The USTs reportedly were located to the west of the building. The USTs were removed by Cuyahoga Landmark. There was no further information regarding this closure in the file. An application for UST removal was submitted; however, the date was not provided. One 500-gallon kerosene and two 1,000-gallon cleaning fluid USTs were slated to be removed. The associated permit was dated February 10, 1999. All three USTs reportedly were located north of the building.

According to the BUSTR files, in August 1998, two 1,000-gallon cleaning fluid USTs and one 550-gallon kerosene UST were removed from the property. In the most recent correspondence, BUSTR sent a letter dated November 2001 requesting a Closure Assessment Report. Reportedly, no action has been taken by Scranton Averell, Inc. regarding this request.

According to the BUSTR files, a Closure Assessment for the removal of four 5,000-gallon USTs was submitted in October 2005. A Deficiency Letter was issued by BUSTR for the Closure on October 21, 2005. Scranton Averell indicated the USTs were not owned or operated by the company.

At the time of the URS Phase I reconnaissance, the site was occupied by TIG Products. A large concrete block building was observed in the central portion of the site. A gravel-covered area is located between this site and the adjacent Stripmatic site. Surface staining was observed on the site. No stressed vegetation and/or the storage or handling of hazardous materials was observed on the site.

### 6.2.1 Field Activities

A total of seven monitoring wells were proposed for Site 13. Groundwater was encountered in only one soil boring (MW02) during field activities; therefore, only one well was installed. A total of six soil borings (SB01, SB03 through SB07) were installed at Site 13. At the time of groundwater sampling, MW02 was dry and no groundwater sample could be obtained. One soil sample per boring was collected and analyzed for VOCs and SVOCs. A duplicate sample was collected from 13-SB01-1820. A Sample Location Map is included as **Figure 6-2A**.

# 6.2.2 Site-Specific Geology/Hydrogeology

The soils at Site 13 consisted primarily of sand with minor amounts of silty clay. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 13 on September 24, 2006. The monitoring well (MW02) was dry. Localized groundwater flow across Site 13 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 13 is likely to the south-southwest.

## 6.2.3 Geophysical Survey

URS conducted a geophysical survey at Site 13 in July 2006. The purpose of the survey was to locate, if present, any abandoned underground storage tanks (USTs).

The geophysical survey of Site 13 included the area west of the building to edge of a ravine, the partially paved area between the building and West 15<sup>th</sup> Street to the east, and portions of the neighboring property to the north. The presence of debris and equipment to the west of the building hindered establishment of a consistent survey grid. In addition, the presence of metallic sludge (reported by the current property owner) within the soil limited the sensitivity of the gradiometer. The survey transects were spaced five feet apart to provide adequate precession for the purpose of locating a small steel UST.

The survey identified three areas with anomalously high magnetic gradients. These areas are shown on **Figure 6-2B** as anomalies A, B and C as described below:

- A. A linear magnetic anomaly was observed parallel to the front of the building along West 15 th Street. This anomaly is likely the result of utilities and not related to a UST.
- B. Several broad areas of high magnetic gradients were detected to the rear of the property. Based on conversations with the property owner, these high gradients are likely related to buried metal and/or buried metallic sludge related to a former machine shop at this location. The presence of such materials would reduce the effectiveness of the geophysical survey.
- C. A single magnetic anomaly was observed north of the property that may represent a buried metallic object. This location was beyond the study area.

The results of this survey do not support the presence of a steel UST at Site 13. No further investigation is recommended to attempt to locate the UST.

### 6.2.4 Soil Analytical Results

No VOCs were detected in the soil samples submitted from Site 13.

A total of eleven SVOCs were detected in two (13-SB-1820D and 13-SB07-0608) of the eight soil samples submitted from Site 13. Concentration of anthracene (14 ug/kg), benzo(a)anthracene (15 ug/kg and 56 ug/kg), benzo(a)pyrene (16 ug/kg and 57 ug/kg), benzo(b)fluoranthene (21 ug/kg and 69 ug/kg), benzo(g,h,i)perylene (31 ug/kg), benzo(k)fluoranthene (29 ug/kg), chrysene (17 ug/kg and 64 ug/kg), fluoranthene (25 ug/kg and 120 ug/kg), indeno(1,2,3)pyrene (29 ug/kg), phenanthrene (67 ug/kg), and pyrene (22 ug/kg and 94 ug/kg) were detected in the soil samples submitted. All other SVOCs were below the detection limits.

The analytical results are presented in Table 6-2A.

# 6.2.5 Groundwater Analytical Results

No groundwater was encountered in MW02.

## 6.2.6 Comparison Standards

The analytical results were compared to the Ohio Environmental Protection Agency's (OEPAs) VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), and the Bureau of Underground Storage Tank Regulations (BUSTR) Closure Action Levels for Class 1 soils. The VAP and BUSTR standards are included on **Table 6-2A**.

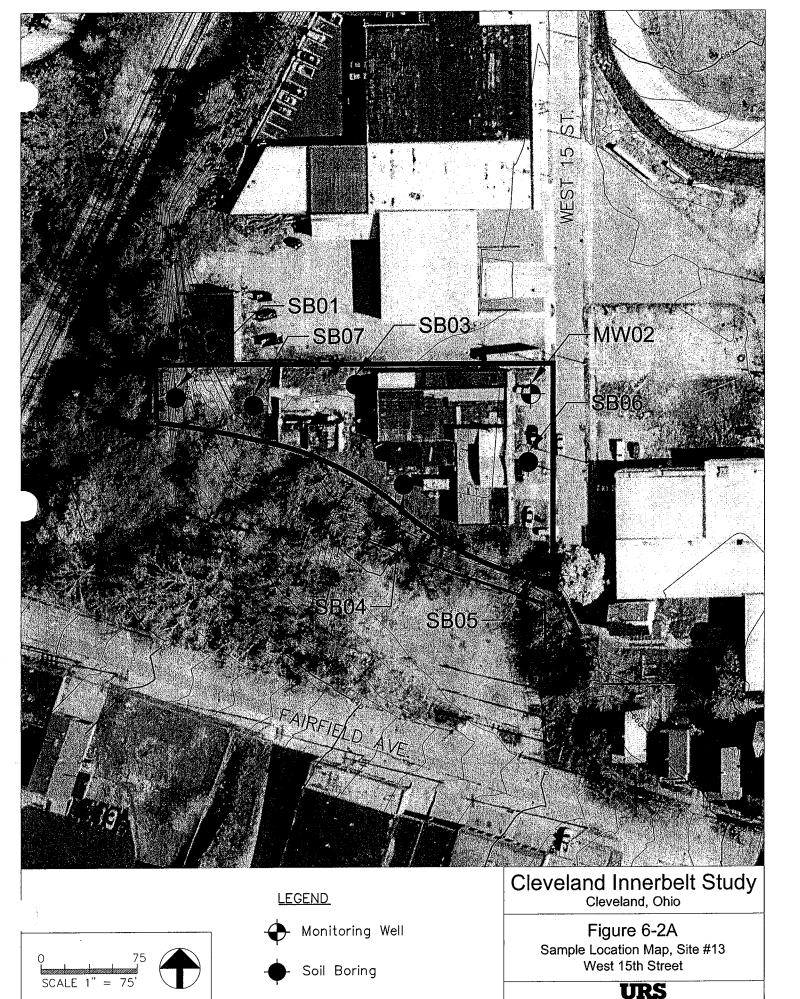
No VOCs were detected in the soil samples submitted from Site 13.

None of the eleven SVOCs detected in the soils at Site 13 exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, and/or BUSTR closure action levels.

### 6.2.7 Conclusions

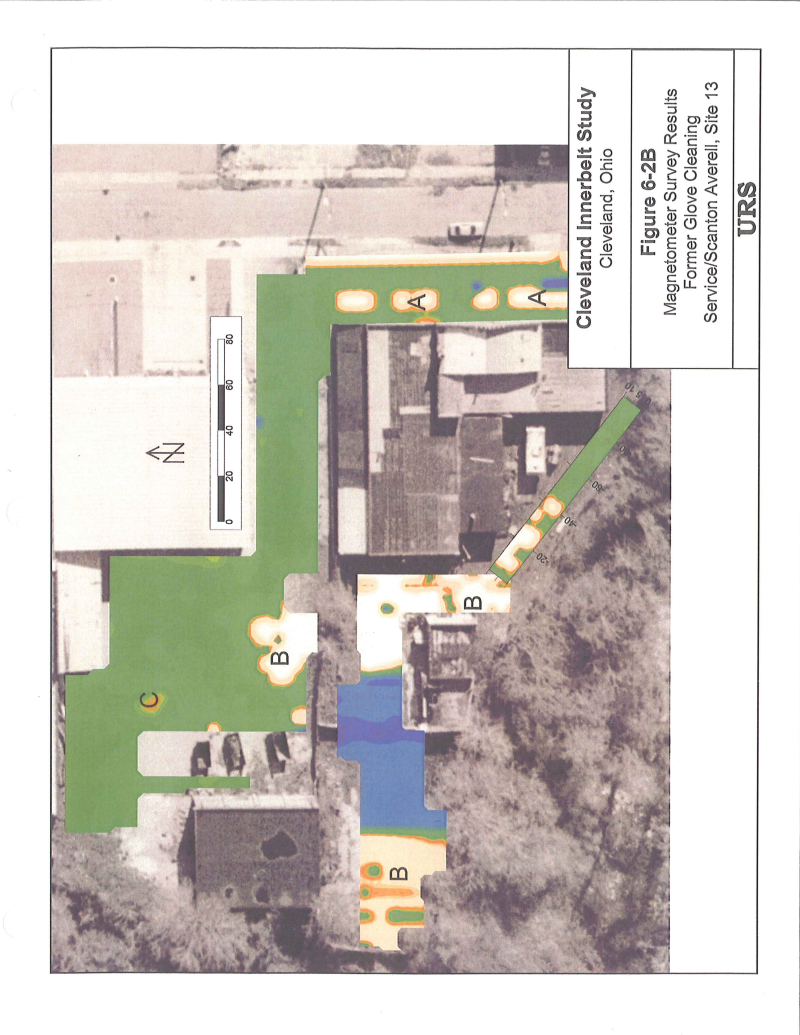
Based on the analytical results, it is unlikely that the soils at Site 13 would require special management during construction.

FIGURES



P:\0\0D0T\15016633\DWGs\Figures\Site 13\figure2-B.dwg User:Kontos

Dec 18, 2006 - 2:48pm



# TABLES

Site 13 - Former Glove Cleaning Service / Scranton Averell Summary of Detected Chemicals in Soil ODOT Innerbelt Study Cleveland, Ohio Table -- 2A

-- = Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were

(1) VAP Generic Direct Contact Soil Standard, Commercial/Industrial Land Use (2) VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

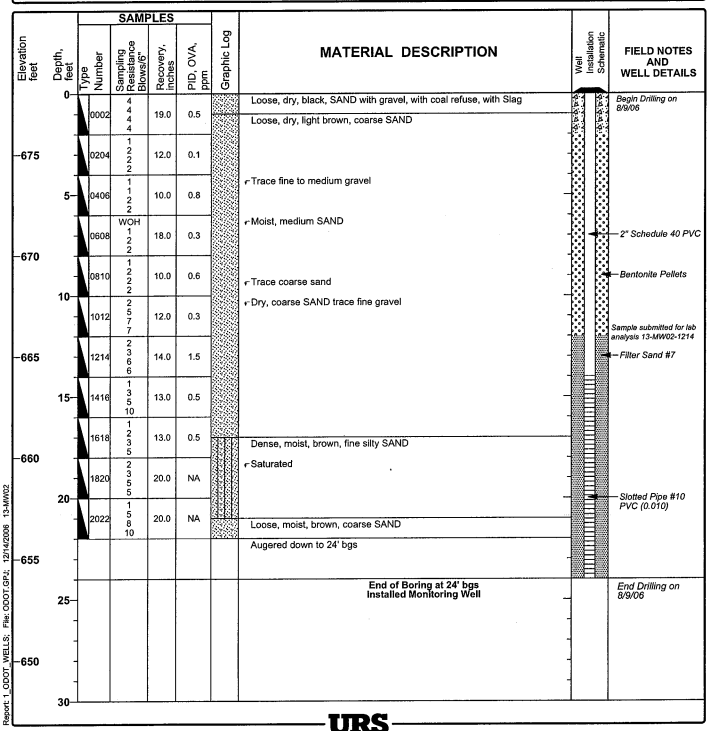
BORING LOGS

**Project: ODOT - Innerbelt Corridor** 

Project Location: Site 13
Project Number: 15016633

# **Log of Boring 13-MW02**

Date(s) Drilled 8/9/06	Logged By	J. Kaminski	Checked M. Wolff By
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 24.0′ bgs
Drill Rig Type LC-60	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 678'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Set monitoring well
Groundwater Level and Date Measured Well dry on 9/24/06			



Project: ODOT - Innerbelt Corridor
Project Location: Site 13
Project Number: 15016633

# Log of Boring 13-SB01

Date(s) Drilled and Installed 8/7/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth 30.0′ bgs
Sampling 2" Split Spoon	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 650'
Drill Rig Type: LC-60	Groundwater Not encountered Level(s)	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill bentonite	

Localic								
ſ			SAM	PLES		ļ		
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm		MATERIAL DESCRIPTION	FIELD NOTES
-650	0-		1				Dense, dry, black, fine and coarse SAND with gravel	Begin drilling on 8/7/06
		0002	3 3 1	12.0	34.4		Loose, dry, light brown, fine to medium SAND with, gravel	*****
	2-	0204	1 1 2 1	8.0	41.8			
-	4- - 6-	0406	2	5.0	47.1			
	8-	0608	3	16.0	47.3		Loose, moist, gray, coarse SAND with fine gravel	
-640	10 <del>-</del>	0810	5	24.0	50.4	177	Stiff, moist, brown, silty CLAY Loose, moist, gray, coarse SAND with fine gravel rDry	
	 12 <del>-</del> -	1012	3 7 8 8 4 7	16.0	50.9		rMedium SAND trace gravel	
	14	1214	9 8	20.0	45.2		r-No gravel	
-	- 16-	1416	2	22.0	275		Soft, wet, brown, silty CLAY	
	18	1618	3	22.0	>9999			Sample submitted for lab
-630	20-	1820	6	20.0	>9999		rCoarse SAND rFine to medium SAND	Sample submitted for lab analysis 13-SB01-1820
	22-	2022	6 3	22.0	29.3			
	24	2426	3	22.0	43.4		· •	*
	26	2628	4	20.0	41.1		- -	
	28- - -	2830	3 6 7	22.0	27		- -	
- -620	30- - 32-		6				End of Boring at 30' bgs	End drilling on 8/7/06
_	34							
· L							——— URS————	

Project: ODOT - Innerbelt Corridor
Project Location: Site 13
Project Number: 15016633

# Log of Boring 13-SB03

Date(s) Drilled 8/9/06 and Installed	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0' bgs
Sampling 2" Split Spoon	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 678'
Drill Rig Type: LC-60	Groundwater Not encountered Level(s)	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill bentonite	

Localic			CAMI	DI EC				
	-		SAM	LES				
Elevation feet	Depth, feet	l ype Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
	0-1	0002	4	27.0	0.0		Dense, dry, black, fine silty SAND with gravel  Medium dense, dry, brown, medium to coarse SAND trace fine gravel	Begin drilling on 8/9/06
_	2- - -	0204	1	12.0	1.0		_	
	4- - 6-	0406	3	13.0	1.2		FLoose -	
-670	8-	0608	3	20.0	0.3		Stiff, moist, brown, silty CLAY	
""	10-	0810	5	12.0	0.4		Loose, dry, brown, fine SAND Coarse SAND -	
	12-	1012	<b>4</b>	19.0	1.3		-rMoist	
-	14-	1214	13	19.0	1.3		rDry	
	16-	1416	1	22.0	2.3		- -	Sample submitted for laboratory analysis 13-SB03-1416
660	18-	1618	7	18.0	0.7		-	
	20-	1820	1 4 6 10	19.0	1.6			
	22-	2022	2 5 4 8	18.0	1.0		-	
_	- 24-	2224	4 6 7	20.0	1.0		rMoist, fine SAND	
202141721	26	2426	7	24.0	0.6		rDry, fine to medium SAND	
——————————————————————————————————————	28	2628	3 4 6 7 2 5	18.0	0.4			
	30	2830	5 6 5	20.0	0.0			
	32						End of Boring at 30' bgs	End drilling on 8/9/06
	34							
349							TIRS	

Project: ODOT - Innerbelt Corridor

Project Location: Site 13 Project Number: 15016633

# Log of Boring 13-SB04

Date(s) Drilled and Installed 8/7/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0′ bgs
Sampling 2" Split Spoon	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 679'
Drill Rig Type: LC-60	Groundwater Not encountered Level(s)	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill <b>bentonite</b>	

			SAME	PLES		<u> </u>		
Elevation feet	Depth, feet	l ype Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
	0-1	0002	2	17.0	12.4		Dense, dry, black, SAND with gravel, with silt	Begin drilling on 8/7/06
	2- - -	0204	1 1 1	7.0	13.4		Loose, dry, brown, fine to coarse SAND with gravel	
	4-	0406	1 2 2 2 2	12.0	13.0		rCoarse SAND	
	6	0608	2	12.0	15.3		rMoist, fine to medium SAND no gravel	
-670	8	0810	1 2 1 1	10.0	15			
	10-	1012	1 2 1	8.0	15.7		r∙Medium to coarse SAND	
	12-	1214	w w w 1	4.0	18.3			
	14	1416	1 1 1	8.0	19.3		Fine to medium SAND	
	=	1618	1 5 8 18	22.0	20.9	Ш	Very stiff, moist, brown, SILT	
-660	18-	1820	2 8 8 8	22.0	19.7		Medium dense, dry, brown, fine SAND	
	20	2022	2 8 9 12	17.0	2723		FBrown and gray, trace gravel	Sample submitted for lab analysis 13-SB04-2022
	22	2224	1 5 8 10	22.0	138		rNo gravel	
	24-	2426	8 8 8	20.0	762			
5	28-	2628	1 2 3 3	20.0	106			
<b>–650</b>	30	2830	2 8 10 12	26.0	70.6			
	32-						End of Boring at 30' bgs	End drilling on 8/7/06
	34-							
							URS	

Project: ODOT - Innerbelt Corridor
Project Location: Site 13
Project Number: 15016633

# Log of Boring 13-SB05

Date(s) Drilled 8/7/00 and Installed	Logge	d By	J. Kaminski	Reviewer	M. Wolff
Drilling Hollow S	tem Auger Drilling Contra		HAD, Inc.	Total Depth of Borehole	30.0′ bgs
Sampling 2" Split S	poon Drill Bi Size/T		4-1/4" ID HSA	Approximete Surface Elevation	676'
Drill Rig Type: LC-60	Ground Level(s	dwate	Not encountered	Hammer 140# Data	auto hammer
Boring Location: See Site	Map Boreho Backfil		bentonite		

			SAMI	PLES				
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
-		0002	2 5 3 4	18.0	11.1		Dense, dry, dark brown and black, fine SAND with gravel, with Slag	Begin drilling on 8/7/06
	2-	0204	1 2 3 3	10.0	22.9		Loose, dry, light brown, medium SAND with gravel	
-670	4- - 6-	0406	1 2 2 3	12.0	26		r-Medium to coarse SAND	
_670	8-	0608	1 4 2 2	12.0	28.3		rMoist	
	10-	0810	1 1 2 2	9.0	28.9		· -	
-	12-	1012	1 1 1	11.0	26.4		-	
	14	1214	1 2 3 3	14.0	25.6		- 	
-660	16	1416	6 4 3	12.0	30.9		FSoft, moist, brown, CLAY 2-inch seam  Medium to coarse SAND	Sample submitted for lab analysis 13-SB05-1416
	18	1618	3 10 10 8 2	14.0	28.7		rTrace gravel	
	20	1820	2 6 9 9	2	24.7		- -	-
-	22	2022	10 1	21.0	23.4		- Medium SAND	
	24-	2224	3 4 4 1 1	14.0	19.4		- -	
-650	26-	2426	4 7 1	17.0	14.4 		- -	
	28 <del>-</del> -	2830	4 8 9 2 9	14.0	7.1		- -	
<u> </u>	30-	N	11 13				End of Boring at 30' bgs	End drilling on 8/7/06
	32						<u>-</u>	8/7/06
	34						-	,
<u> </u>							URS	

Project: ODOT - Innerbelt Corridor
Project Location: Site 13

Project Number: 15016633

# Log of Boring 13-SB06

Date(s) Drilled 8/9/06 and Installed	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0′ bgs
Sampling 2" Split Spoon Method	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete 678'
Drill Rig Type: LC-60	Groundwater Not encountered Level(s)	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill <b>bentonite</b>	

				SAME	PLES				
	Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
		<b>1</b>	0002	1 3 5 12	12.0	1.2		Loose, dry, brown and black, SAND with gravel	Begin drilling on 8/9/06
	_	2- - -	0204	3 2 1 1	14.0	1.2		Medium dense, dry, light brown, coarse SAND with fine gravel	-
		4- - - 6-	0406	w w 1	5.0	1.2		rLoose	
	670	8	0608	1 1 2 2	14.0	1.3		≁Moist, with coarse gravel	
)	070	10-	0810	1 2 2 3	14.0	0.6			]
,		12-	1012	1 2 2 1	10.0	0.8		←Very moist, with fine gravel	
		14-	1214	1 2 2 3	20.0	0.5		Loose, dry to moist, brown, medium SAND	-
		16-	1416	1 3 5 6	18.0	0.8			1
	660	18-	1618	1 7 4 5	19.0	0.7			1
		 20	1820	3 5 7	15.0	1.1		←Medium to coarse SAND trace gravel	
		22	2022	3 6 10 12	22.0	0.6		r-Fine to medium SAND no gravel	-
3 13-SB06	-	24	2224	10 7 7	24.0	0.7		←With gravel	-
12/14/2006		26	2426	4 5 7	20.0	1.2			-
ODOT.GPJ;	-650	28	2628		24.0	5.6			Sample submitted for lab analysis 13-SB06-2628
ë		30	2830	5 9 7	19.0	4.4			-
30RINGS;		32						End of Boring at 30' bgs	- End drilling on 8/9/06
Report: 1_ODOT_BORINGS;	-	34							
Report: 1				l.	1			URS	

Project: ODOT - Innerbelt Corridor

Project Location: Site 13 Project Number: 15016633

# Log of Boring 13-SB07

Date(s) Dril and Installe		Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method	Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0' bgs
Sampling Method	2" Split Spoon	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 676'
Drill Rig Type:	LC-60	Groundwater Not encountered Level(s)	Hammer 140# auto hammer Data
Doring	See Site Map	Borehole Backfill <b>bentonite</b>	

			SAME	PLES				
Elevation feet	Depth, feet	ı ype Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
L	Ē	0002	3 6 5	12.0	10	607160716	ASPHALT  Dense, dry, black, gravelly SAND trace clay, trace Slag	Begin drilling on 8/7/06
	2-		1				Loose, dry, brown, coarse SAND with gravel	
	- 4-	0204	2 1 1	10.0	11.5			1
-670	- 6-	0406	1 2 3 2	12.0	12.5			
0,0	8-	0608	1 2 2 2	17.0	14.1			Sample submitted for lab analysis 13-SB07-0608
		0810	1 3 6 7	16.0	11.9		Stiff, moist, brown, SILT with fine to medium sand, with gravel	
	10 <del>-</del>  	1012	3 3 3 2	12.0	12.4		Loose, dry, light brown, fine to medium SAND with gravel	
	12-	1214	1 6 6	18.0	11		rNo gravel	
	14-	1416	1 6 8 6	20.0	11.8			
-660	16-	1618	2 6 11 6	19.0	11.3		r-Brown and gray	
	18— — 20—	1820	2 6 8 6	14.0	10.1			
_	22-	2022	2 4 6 6	20.0	7.8			-
	24-	2224	2 6 5 5	19.0	5.2		-rLight brown, medium SAND	
-650	26	2426	1 3 5 7	19.0	3.7			
	28-	2628	2 5 5	16.0	0.7			
	30	2830	1 6 8 6	16.0	0.6			-
_	32-						End of Boring at 30' bgs	End drilling on 8/8/06
l							URS	

DATA ASSESSMENT REPORT

# Data Assessment Report ODOT Innerbelt Study Site 13 - Former Glove Cleaning Service / Scranton Averell

Reviewer: P. Schuler Date: October 30, 2006

Eight soil samples were collected at the Former Glove Cleaning Service / Scranton Averell site in Cleveland, Ohio, on August 7 through 9, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Requested	Analyses <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC
A6H090255001	13-SB01-1820 <sup>(2)</sup>	08/07/2006	Soil	X	X
A6H090255002	13-SB05-1416	08/08/2006	Soil	X	X
A6H090255003	13-SB04-2022	08/08/2006	Soil	X	X
A6H090255004	13-SB01-1820D <sup>(2)</sup>	08/07/2006	Soil	X	X
A6H090255005	13-SB07-0608	08/08/2006	Soil	X	X
A6H090255006	13-SB06-2628	08/09/2006	Soil	X	X
A6H090255007	13-MW02-1214	08/09/2006	Soil	X	X
A6H120111001	13-SB03-1416	08/09/2006	Soil	X	X

<sup>(1)</sup> VOC = Volatile Organic Compounds [SW-846 Method 8260B] SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Positive detections for acetone in samples 13-SB06-2628 and 13-MW02-1214 and for bis(2-ethylhexyl)phthalate in samples 13-SB01-1820, 13-SB05-1416, 13-SB04-2022, 13-SB01-1820D, 13-SB07-0608, and 13-SB06-2628 were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks at similar concentrations.
- The semivolatile compound, bis(2-ethylhexyl)phthalate, was detected at concentrations ranging from 0.9 to 1.8 ug/L in all equipment blanks associated with project samples.

<sup>(2)</sup> Samples 13-SB01-1820 and 13-SB01-1820D are field duplicates.

Since this compound is considered a common laboratory contaminant, its presence in environmental samples at concentrations less than ten times the highest equipment blank concentration (converted to an equivalent soil concentration) is attributed to external contamination rather than actual site conditions. Therefore, the positive detection for bis(2-ethylhexyl)phthalate in sample 13-SB03-1416 was qualified as nondetect ("U") at the sample reporting limit, even though it was not detected in the associated method blank.

- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J"), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, the "J" qualifiers were retained with the numeric results.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 and 2-2.

Tabie 2-1
Analytical Data Summary
Site 13 Soil Volatiles
ODOT Innerbelt Study

		A6H090255001	A6H090255002	A6H090255003	A6H090255004	A6H090255005	A6H090255006	A6H090255007	A6H120111001
PARAMETER	UNITS	13-SB01-1820 08/07/2006	13-SB05-1416 08/08/2006	13-SB04-2022 08/08/2006	13-SB01-1820D 08/07/2006	13-SB07-0608 08/08/2006	13-SB06-2628 08/09/2006	13-MW02-1214 08/09/2006	13-SB03-1416 08/09/2006
Percent Solids	%	92.9	92.8	95.1	88	92.6	97	94.6	94.5
1,1,1-Trichloroethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
1,1,2,2-Tetrachloroethane	ng/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
1,1,2-Trichloroethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 U
1,1-Dichloroethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
1,1-Dichloroethene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
1,2,4-Trichlorobenzene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
1,2-Dibromo-3-chloropropane	ug/kg	J L	J L	J 1-	0 T	n T	10 U	11 C	11 0
1,2-Dibromoethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
1,2-Dichlorobenzene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
1,2-Dichloroethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
1,2-Dichloropropane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
1,3-Dichlorobenzene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
1,4-Dichlorobenzene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
2-Butanone	ug/kg	22 U	22 U	21 U	23 N	22 U	21 U	21 U	21 U
2-Hexanone	ug/kg	22 U	22 U	21 U	23 U	22 U	21 U	21 U	21 U
4-Methyl-2-pentanone	ug/kg	22 U	22 O	21 U	23 U	22 U	21 U	21 U	21 U
Acetone	ug/kg	22 U	22 U	21 U	23 U	22 U	21 U	21 U	21 U
Велхене	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Bromodichloromethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Bromoform	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5,4 U	5.2 U	5.3 ∪	5.3 U
Bromomethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 ∪	5.3 ∪	5.3 ∪
Carbon disulfide	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
Carbon tetrachloride	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
Chlorobenzene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
Chloroethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 U
Chloroform	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 ∪
Chloromethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 U
cis-1,2-Dichloroethene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
cis-1,3-Dichloropropene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Cyclohexane	ug/kg	11 U	11 U	U TI	D T	T ,	10 U	11 O	11 U
Dibromochloromethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 U
Dichlorodifluoromethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 U
Ethylbenzene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 ∪	5.3 U	5.3 U
Isopropylbenzene	ug/kg	5.4 U	5.4 U	5.3 ∪	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Methyl acetate	ug/kg	11 U	11 U	11 C	11 U	11 U	10 U	11 U	11 U

# Table-2.1 Analytical Data Summary Site 13 Soil Volatiles ODOT Innerbelt Study

PARAMETER	SLIND	A6H090255001 A6 13-SB01-1820 13 08/07/2006 0	A6H090255002 13-SB05-1416 08/08/2006	270000000000000000000000000000000000000	A6H090255004 13-SB01-1820D 08/07/2006	A6H090255003 A6H090255004 A6H090255005 13-SB04-2022 13-SB01-1820D 13-SB07-0608 08/08/2006 08/07/2006 08/08/2006	A6H090255006 13-SB06-2628 08/09/2006	A6H090255007 13-MW02-1214 08/09/2006	A6H120111001 13-SB03-1416 08/09/2006
Methyl tert-butyl ether	ug/kg	22 U	22 U	21 U	23 U	22 U	21 U	21 U	21 U
Methylcyclohexane	ug/kg	<b>T</b>	11 C	) -	n F	11 O	10 U	11 0	11 0
Methylene chloride	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Styrene	ug/kg	5.4 U	5.4 U	5.3 ∪	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
Tetrachloroethene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Toluene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
trans-1,2-Dichloroethene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 ∪
trans-1,3-Dichloropropene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Trichloroethene	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Trichlorofluoromethane	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 U	5.3 U
Vinyl chloride	ug/kg	5.4 U	5.4 U	5.3 U	5.7 U	5.4 U	5.2 U	5.3 ∪	5.3 U
Xylenes (total)	ug/kg	11 U	11 U	11 U	11 U	11 U	10 U	11 U	11 C

U=The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

Table-2-2 Analytical Data Summary Site 13 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	SINO	A6H090255001 13-SB01-1820 08/07/2006	A6H090255002 13-SB05-1416 08/08/2006	A6H090255003 13-SB04-2022 08/08/2006	A6H090255004 13-SB01-1820D 08/07/2006	A6H090255005 13-SB07-0608 08/08/2006	A6H090255006 13-SB06-2628 08/09/2006	A6H090255007 13-MW02-1214 08/09/2006	A6H120111001 13-SB03-1416 08/09/2006
1.1'-Biohenvi	ויט/גע	360 11	360 11	250 11	11 000	000	2000	0202/0000	000/2/2000
2 2'-oxyhis(1-Chloronronane)	5/ <sub>2</sub> / <sub>2</sub> / <sub>2</sub> / <sub>2</sub>		51 098					0.000	0.000
9.4 5. Trichlombonol	D 1	0 000	2000				0.00	0.000	0 000
2.4.5. Trichlorophonoi	5 1/25 5 1/25 5 1/25	0 000	0.000				340 0	320 0	320 0
2,4,0-11,011,011,01	מלועלו	0 000	O 005			_	340 0	320 N	320 N
2,4-Dichiorophenol	ug/kg	360 U	098 n	350 U	380 U	O 098	340 U	320 N	320 N
2,4-Dimethylphenol	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
2,4-Dinitrophenol	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
2,4-Dinitrotoluene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
2,6-Dinitrotoluene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
2-Chloronaphthalene	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
2-Chlorophenol	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
2-Methylnaphthalene	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
2-Methylphenol	ug/kg	09E	360 U	350 U	380 U	098 360	340 U	350 U	350 U
2-Nitroaniline	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
2-Nitrophenol	ug/kg	3e0 U	360 U	350 U	380 U	09E	340 U	350 U	350 U
3,3'-Dichlorobenzidine	ug/kg	360 U	O 098	350 U	380 U	O 098	340 U	350 U	350 U
3-Nitroaniline	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 ∪	350 U
4,6-Dinitro-2-methylphenol	ug/kg	360 U	O96	350 U	380 U	360 U	340 U	350 U	350 U
4-Bromophenyl phenyl ether	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
4-Chloro-3-methylphenol	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
4-Chloroaniline	ug/kg	360 U	O 098	350 U	380 U	360 U	340 U	350 U	350 U
4-Chlorophenyl phenyl ether	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
4-Methylphenol	ug/kg	360 U	O96	350 U	380 U	360 U	340 U	350 ∪	350 U
4-Nitroaniline	ug/kg	360 U	360 U	350 U	380 ∪	360 U	340 U	350 ∪	350 U
4-Nitrophenol	ug/kg	n 096	360 U	350 U	380 U	360 U	340 U	350 ∪	350 U
Acenaphthene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 ∪	350 U
Acenaphthylene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Acetophenone	ug/kg	72 U	72 U	70 U	76 U	72 U	N 69	O 07	70 U
Anthracene	ug/kg	360 U	360 U	350 U	380 U	14 J	340 U	350 U	350 U
Atrazine	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
Benzaldehyde	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Benzo(a)anthracene	ug/kg	360 U	360 U	350 U	15 J	56 J	340 U	350 U	350 U
Benzo(a)pyrene	ug/kg	360 U	360 U	350 U	16 J	57 J	340 U	350 U	350 U
Benzo(b)fluoranthene	ug/kg	360 U	360 U	350 U	21 J	r 69	340 U	350 U	350 U
Benzo(ghi)perylene	ug/kg	360 U	360 U	350 U	380 U	31 J	340 U	350 U	350 U
Benzo(k)fluoranthene	ug/kg	360 U	360 U	350 U	380 U	29 ט	340 U	350 U	350 U
bis(2-Chloroethoxy)methane	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U

Analytical Data Summary Site 13 Soil Semivolatiles ODOT Innerbelt Study

Tabre 2-2

		A6H090255001	A6H090255002	A6H090255003	A6H090255004	A6H090255005	A6H090255006	A6H090255007	A6H120111001
		13-SB01-1820	13-SB05-1416	13-SB04-2022	13-SB01-1820D	13-SB07-0608	13-SB06-2628	13-MW02-1214	13-SB03-1416
PAHAMETER	SLIND	08/02//2006	08/08/2006	08/08/2006	08/07/2006	08/08/2006	08/09/2006	08/09/2006	08/09/2006
bis(2-Chloroethyl) ether	ug/kg	O 098	360 U	350 U	380 U	360 U	340 U	350 U	350 U
bis(2-Ethylhexyl) phthalate	ug/kg	098	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Butyl benzyl phthalate	ug/kg	O 098	360 U	350 U	380 U	360 U	340 U	350 ∪	350 U
Caprolactam	ug/kg	O 098	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Carbazole	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
Chrysene	ug/kg	360 U	360 U	350 U	L 71	64 J	340 U	350 U	350 U
Dibenz(a,h)anthracene	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 ∪	350 U
Dibenzofuran	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 ∪	350 U
Diethyl phthalate	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
Dimethyl phthalate	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Di-n-butyl phthalate	ug/kg	O 098	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Di-n-octyl phthalate	ug/kg	O 098	360 U	350 U	380 U	O96	340 U	350 U	350 U
Fluoranthene	ug/kg	360 U	360 U	350 U	25 J	120 J	340 U	350 U	350 U
Fluorene	ug/kg	360 U	360 U	350 U	380 U	O98	340 U	350 U	350 U
Hexachlorobenzene	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
Hexachlorobutadiene	ug/kg	360 U	360 U	350 U	380 U	O96	340 U	350 U	350 U
Hexachlorocyclopentadiene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Hexachloroethane	ug/kg	360 U	360 U	350 U	380 U	O 098	340 U	350 U	350 U
Indeno(1,2,3-cd)pyrene	ug/kg	360 U	360 U	350 U	380 U	29 J	340 U	350 U	350 U
Isophorone	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Naphthalene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Nitrobenzene	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
N-Nitrosodi-n-propylamine	ug/kg	n 098	360 U	350 U	380 U	360 U	340 U	350 U	350 U
N-Nitrosodiphenylamine	ug/kg	O 098	∩ 096	350 U	380 U	360 U	340 U	350 U	350 U
Pentachlorophenol	ug/kg	O98	360 U	350 U	380 U	360 ∪	340 U	350 U	350 U
Phenanthrene	ug/kg	360 U	360 U	320 N	380 U	f 29	340 U	350 U	350 U
Phenol	ug/kg	360 U	360 U	350 U	380 U	360 U	340 U	350 U	350 U
Pyrene	ug/kg	360 U	360 U	350 U	22 J	94 J	340 U	. 350 U	350 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

### 6.3 SITE 14 – BOJACKS MEATS

According to the Cleveland City Directories, the site was listed as Paramount Fur Company from 1954 through 1974. Beginning in 1979 through 1999, the site was listed as Bojacks Meat and Poultry.

According to the Cleveland Fire Prevention Bureau files, the site was occupied by the J.R. Dyamond Terminal Warehouse in 1944. An entry in March 1952 indicates the site was occupied by the Paramount Fur Company, which utilized mixtures of carbon tetrachloride and apco thinner to clean furs. A Fire Inspection Report dated November 1967 indicates the Cuyahoga Chemical Company stored multiple drums of sodium cyanide, potassium cyanide, zinc cyanide, and copper cyanide in the warehouse.

A permit dated November 1951 indicates a gas station was located at 2000-2180 West 14<sup>th</sup> Street. There reportedly were three 3,000-gallon USTs associated with this operation.

At the time of the URS Phase I reconnaissance, this site was occupied by two large warehouse buildings. While surface staining was not observed, distressed vegetation was observed on the site.

### 6.3.1 Field Activities

A total of six monitoring wells were proposed for Site 14. During the Phase II site reconnaissance, it was determined that two of the proposed boring locations were inaccessible. One boring was proposed near the former loading docks, which were open beneath. The other proposed boring was located on a steep slope adjacent to a railroad siding.

Groundwater was encountered in three soil borings (MW01, MW02, and MW03) during field activities and monitoring wells were installed. One soil boring (SB04) was installed at Site 14. At the time of groundwater sampling, MW01 was dry and no groundwater sample could be obtained. Four soil and two groundwater samples were collected and analyzed for VOCs, SVOCs, TPH and/or Cyanide. A duplicate soil sample was collected from 14-SB04-2022 and a duplicate groundwater sample was collected from 14-MW02. A Sample Location Map is included as **Figure 6-3A**.

# 6.3.2 Site-Specific Geology/Hydrogeology

Soils at Site 14 consisted primarily of sand with minor amounts of silty clay and clay. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 14 on September 26, 2006. Monitoring well (MW01) was dry. Localized groundwater flow across Site 14 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 14 is to the north-northeast, towards the Cuyahoga River.

SECTIONS IX Phase II Findings

## 6.3.3 Geophysical Survey

URS conducted a geophysical survey at Site 14 in July 2006. Figure 6-3B indicates the results of the survey.

The geophysical survey of Site 14 included the paved surface of the property, northwest of the highway bridge. The survey transects were conducted in a north – south direction to approximately align transects along the earth's total magnetic field which provides the greatest sensitivity to buried ferrous objects such as steel underground storage tanks. Transects were spaced five feet apart and were walked in alternating directions. The presence of tall structures prevented the use of a global positioning system (GPS) for the establishment of coordinates.

No evidence of USTs were discovered as a result of the geophysical survey of Site 14.

# 6.3.4 Soil Analytical Results

No VOCs were detected in the soil samples submitted from Site 14.

A total of nineteen SVOCs were detected in four (14-MW012426, 14-MW02-0406, 14-MW03-1012, and 14-SB04-2022) of the five soil samples submitted from Site 14. Concentrations of 2-methylnaphthanene (21 ug/kg), acenaphthene (ranging from 17 ug/kg to 190 ug/kg), anthracene (ranging from 36 ug/kg to 280 ug/kg), benzo(a)anthracene (ranging from 11 ug/kg to 1,100 ug/kg), benzo(a)pyrene (ranging from 11 ug/kg to 1,000 ug/kg), benzo(b)fluoranthene (ranging from 16 ug/kg to 1,300 ug/kg), benzo(g,h,i)perylene (ranging from 8.9 ug/kg to 560 ug/kg), benzo(k)fluoranthene (ranging from 69 ug/kg to 560 ug/kg), caprolactam (48 ug/kg), chrysene (ranging from 22 ug/kg to 200 ug/kg), dibenz(a,h)anthracene (ranging from 24 ug/kg to 160 ug/kg), dibenzofuran (76 ug/kg), fluoranthene (ranging from 29 ug/kg to 2,300 ug/kg), fluorene (ranging from 13 ug/kg to 150 ug/kg), indeno(1,2,3)pyrene (ranging from 7.7 ug/kg to 520 ug/kg), naphthalene (ranging from 8.0 ug/kg to 15 ug/kg), phenanthrene (ranging from 21 ug/kg to 1,400 ug/kg), and pyrene (ranging from 21 ug/kg to 1,900 ug/kg) were detected in the soil samples submitted. All other SVOCs were below the detection limits.

Diesel range total petroleum hydrocarbons were detected in all of the soil samples. The highest concentrations were found in the heaviest range, which is consistent with the concentrations of SVOCs detected across the Property. Concentrations of the heavy petroleum fraction ranged from 8.5 to 40 mg/kg. Concentrations of the middle petroleum fraction ranged from 2.2 mg/kg to 6.7 mg/kg.

The analytical results are presented in Table 6-3A.

# 6.3.5 Groundwater Analytical Results

Three VOCs were detected in groundwater samples collected from Site 14. Concentrations of 2-butanone (ranging from 0.52 ug/L to 0.76 ug/L), bromodichloromethane (1.1 ug/L), and chloroform (ranging from 5.6 ug/L to 6.0 ug/L) were detected in the groundwater samples submitted. All other VOCs were below the detection limits.

Nine SVOCs were detected in the groundwater samples submitted from Site 14. Concentrations of benzo(a)anthracene (0.79 ug/L and 0.46 ug/L), benzo(a)pyrene (0.63 ug/L and 0.43 ug/L), benzo(b)fluoranthene (1.1 ug/L and 0.56 ug/L), benzo(g,h,i)perylene (0.52 ug/L), chrysene (0.75 ug/L and 0.51 ug/L), fluoranthene (2.0 ug/L and 1.2 ug/L), indeno(1,2,3)pyrene (0.95 ug/L), phenanthrene (1.1 ug/L and 0.59 ug/L), and pyrene (1.8 ug/L and 1.2 ug/L) were detected in the groundwater sample submitted from 14-MW02 and 14-MW02D, respectively. All other SVOCs were below the detection limits.

The analytical results are presented in Table 6-3B.

# 6.3.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on **Tables 6-3A** and **6-3B**.

There were no VOCs detected in any of the soil samples submitted from Site 14.

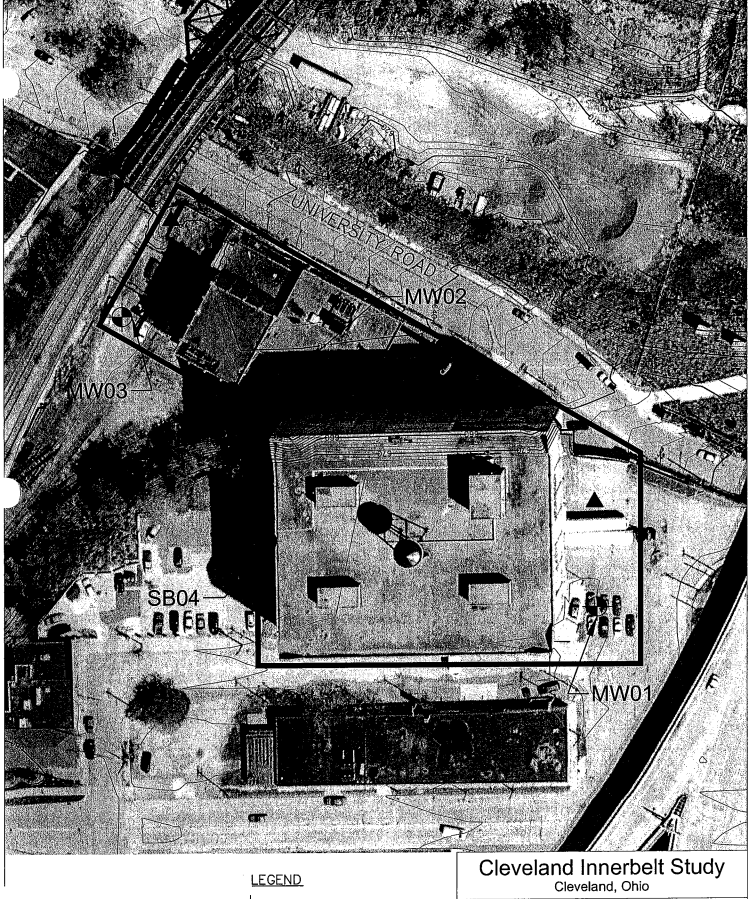
There were no SVOCs detected in any of the soil samples from Site 14, which exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, and/or the BUSTR closure action levels.

The concentration of benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, and indeno(1,2,3)pyrene detected in the groundwater samples from 14-MW02 and/or 14-MW02D exceeded the OEPA VAP standard for generic unrestricted potable use standards and/or BUSTR closure action levels.

### 6.3.7 Conclusions

Based on the analytical results, the soils at Site 14 are not expected to require special management during construction. However, the groundwater concentrations may require special disposal and/or worker protection protocols (plan note).

# FIGURES





Monitoring Well



Soil Boring

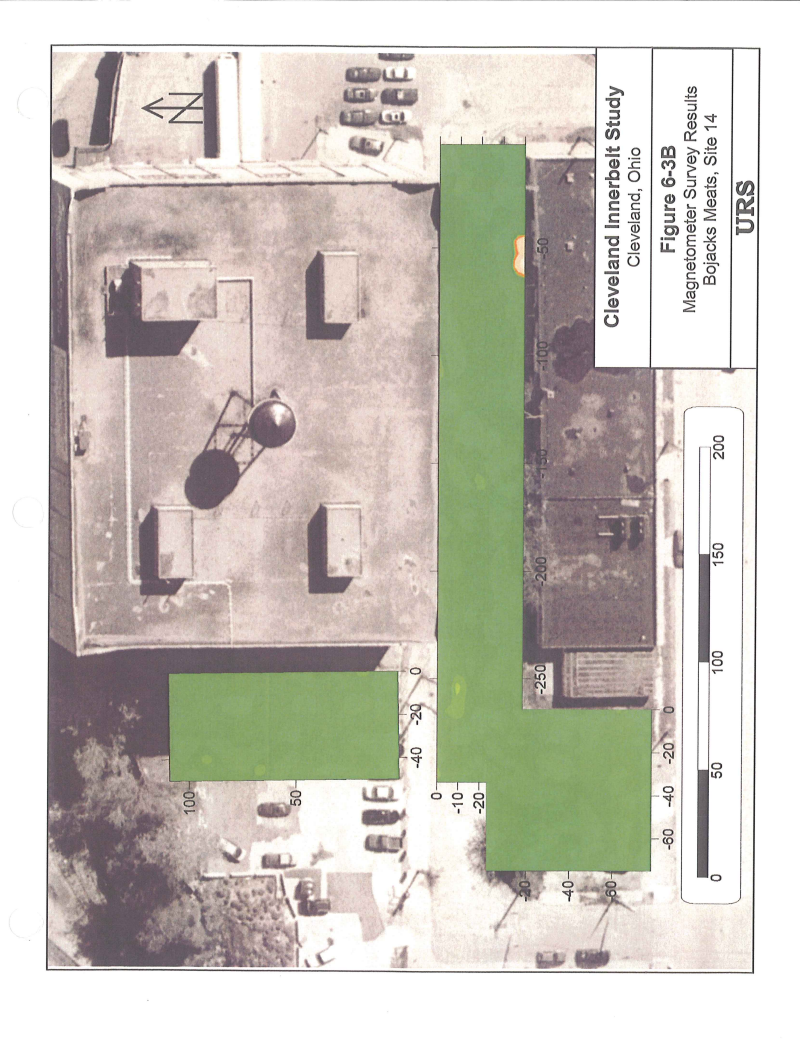


Abandoned Boring

# Figure 6-3A

Sample Location Map, Site #14 1425 University Road / 2000 W. 14TH Street

URS



**TABLES** 

Tab. A
Summary of Detected Chemicals in Soil
Site 14 - Bojacks Meats
ODOT Innerbelt Study
Cleveland, Ohio

W	NA	NA	W	MA	¥	WA	NA	NA	NA	Ą	¥.	NA	WA	NA	NA	NA	NA	NA	6.7	53
350 U	L 71	36 J	130 J	140 J	180 J	100 J	S8 J	350 U	22 J	160 J	24 J	350 U	290 J	13.5	85 J	ი 9.6	150 J	250 J	4.7	18
370 U	370 U	370 U	11.	10 J	16 J	8.9 კ	370 U	48 J	370 U	15 J	370 U	370 U	29 J	370 U	L 7.7	370 U	21.5	21.3	2.2	8.5
21.5	190 J	280 J	1100	1000	1300	560 J	560 J	710 U	200 J	1000	160 با	L 9Z	2300	150 J	520 J	15 J	1400	1900	22 U	4
O6E	36 J	91 J	190 J	150 J	200 J	P88	Ր 69	330 U			28 J	390 U	410	24 J	SO J	8.0 J	300 J	340 J	4.7	28
-	ı	1	11,000	1,100	11,000	1	110,000	1	•	1,100,000	1,100	1	•	ı	11,000	39,800	1	1	2,000	5,000
	530,000,000	1,000,000,000	810,000	81,000	810,000		8,100,000	ŀ	31,000,000	41,000,000	41,000	•	170,000,000	340,000,000	410,000	1,900,000	1	130,000,000		•
	180,000,000	880,000,000	63,000	6,300	63,000	1	630,000	1	10,000,000	6,700,000	6,700	1	33,000,000	120,000,000	67,000	530,000	1	25,000,000	•	
ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	mg/kg	mg/kg
2-Methylnaphthalene	Acenaphthene	Anthracene	Benzo(a)anthracene	Benzo(a)pyrene	Benzo(b)fluoranthene	Benzo(ghi)perylene	Benzo(k)fluoranthene	Caprolactam	Sarbazole	hrysene	ibenz(a,h)anthracene	ibenzofuran	luoranthene	luorene	ndeno(1,2,3-cd)pyrene	laphthalene	henanthrene	yrene	C10-C20	C20-C34
		ug/kg           390 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          36 J         190 J         370 U         17 J	ug/kg           390 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          36 J         190 J         370 U         17 J           ug/kg         880,000,000         1,000,000,000          81 J         280 J         370 U         36 J	ug/kg         180,000,000         530,000,000         -         81 J         36 J         190 J         370 U         17 J           ug/kg         880,000,000         1,000,000,000         -         81 J         280 J         370 U         17 J           ug/kg         63,000         810,000         11,000         11,000         110 J         110 J         110 J	ug/kg         180,000,000         530,000,000          81 J         21 J         370 U         350 U           ug/kg         180,000,000         1,000,000,000          81 J         280 J         370 U         17 J           ug/kg         63,000         810,000         11,000         11,000         1100         11 J         130 J           ug/kg         6,300         81,000         1,100         150 J         100         10 J         140 J	ug/kg         180,000,000         530,000,000          36 J         190 J         370 U         17 J           ug/kg         180,000,000         1,000,000,000          81 J         280 J         370 U         17 J           ug/kg         63,000         810,000         11,000         150 J         1100         11 J         130 J           ug/kg         6,300         810,000         11,000         200 J         1300         16 J         180 J	ug/kg         180,000,000         530,000,000          36 J         190 J         370 U         17 J           ug/kg         880,000,000         1,000,000,000          81 J         280 J         370 U         17 J           ug/kg         63,000         810,000         11,000         190 J         1100         11 J         130 J           ug/kg         6,300         81,000         1,100         150 J         160 J         140 J           ug/kg         6,300         810,000         11,000         200 J         16 J         180 J           ug/kg          88 J         560 J         8.9 J         100 J	ug/kg         180,000,000         530,000,000         -         -         36 J         190 J         370 U         17 J           ug/kg         180,000,000         1,000,000,000         -         81 J         280 J         370 U         17 J           ug/kg         63,000         810,000         11,000         190 J         11 J         130 J           ug/kg         6,300         81,000         11,000         200 J         1300         16 J         140 J           ug/kg         63,000         8,10,000         110,000         69 J         560 J         370 U         88 J	ug/kg            390 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          81 J         280 J         370 U         17 J           ug/kg         63,000         81,000         11,000         190 J         11 J         130 J           ug/kg         6,300         81,000         1,100         200 J         16 J         140 J           ug/kg         63,000         8,100,000         110,000         200 J         16 J         160 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         8,9 J         100 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         8,9 J         100 J           ug/kg           390 U         710 U         48 J         350 U	ug/kg         180,000,000         530,000,000         -         81 J         21 J         370 U         17 J           ug/kg         180,000,000         1,000,000,000         -         81 J         280 J         370 U         17 J           ug/kg         63,000         810,000         11,000         190 J         11 J         140 J         130 J           ug/kg         6,300         810,000         11,000         200 J         1000         16 J         180 J           ug/kg	ug/kg           390 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          81 J         280 J         370 U         17 J           ug/kg         880,000,000         1,000,000,000         11,000         190 J         11 J         130 J           ug/kg         6,300         810,000         11,000         200 J         100         16 J         180 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         89 J         100 J           ug/kg            390 U         710 U         48 J         350 U           ug/kg         10,000,000         31,000,000         1,100,000         180 J         160 J         160 J           ug/kg         6,700,000         41,000,000         1,100,000         180 J         160 J         150 J	ug/kg           390 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          81 J         190 J         370 U         17 J           ug/kg         63,000         810,000         11,000         190 J         1100         11 J         130 J           ug/kg         6,300         810,000         11,000         200 J         16 J         180 J         140 J           ug/kg         6,300         810,000         11,000         200 J         16 J         180 J         180 J           ug/kg         630,000         8.100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         6.700,000         31,000,000          -         88 J         350 U           ug/kg         6,700,000         31,000,000         1,100,000         180 J         370 U         48 J         350 U           ug/kg         6,700         41,000         1,100,000         180 J         160 J         370 U         22 J           e         ug/kg         6,700         41,000         1,100         28 J         160 J         370 U         24 J	ug/kg         —         —         —         350 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000         —         81 J         280 J         370 U         17 J           ug/kg         880,000,000         1,000,000,000         11,000         150 J         1100         11 J         130 J           ug/kg         6,300         810,000         1,100         200 J         1300         16 J         180 J           ug/kg         630,000         8100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         630,000         81,000,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         6,700,000         31,000,000         1,100,000         1,100,000         1,100,000         150 J         22 J           ug/kg         6,700,000         41,000,000         1,100,000         1,100,000         150 J         22 J           ug/kg         6,700         41,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000         1,100,000 <td< td=""><td>ug/kg        </td><td>ug/kg           350 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          81 J         190 J         370 U         17 J           ug/kg         63,000         1,000,000,000         11,000         190 J         1100         11 J         130 J           ug/kg         6,300         81,000         1,100         200 J         1300         16 J         180 J           ug/kg         63,000         810,000         11,000         200 J         100         10 J         140 J           ug/kg         63,000         8,100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         10,000,000         31,000,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         10,000,000         31,000,000         1100,000         350 U         22 J           ug/kg         6,700,000         41,000         1,100         28 J         160 J         370 U         24 J           ug/kg         6,700         41,000         1,100         2300         350 U         29 J         290 J           ug/kg</td><td>ug/kg             350 U         17 J           ug/kg         180,000,000         530,000,000          86 J         190 J         370 U         17 J           ug/kg         63,000         1,000,000,000         1,1000         150 J         1100         11 J         130 J           ug/kg         6,300         81,000         1,100         200 J         1300         16 J         140 J           ug/kg         6,300         81,000         1,100         200 J         1300         16 J         140 J           ug/kg         6,300         81,000         11,000         69 J         560 J         8.9 J         100 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         6,700,000         41,000,000         1,100,000         15 J         100 J         15 J         160 J           ug/kg         6,700         41,000,000         1,100         23 J         100 J         15 J         10 J</td><td>ug/kg        </td><td>ug/kg   </td><td>ug/kg         E30,000,000         E30 U         21 J         370 U         17 J           ug/kg         180,000,000         530,000,000         -         36 J         190 J         370 U         17 J           ug/kg         880,000,000         11,000         11,000         150 J         1100</td><td>ug/kg         630,000,000          980 U         21 J         370 U         360 U           ug/kg         680,000,000         1,000,000,000         11,000         181 J         280 J         370 U         17 J           ug/kg         63,000         810,000         1,100         190 J         1100         110 J         130 J           ug/kg         63,000         810,000         11,000         190 J         1100         1100 J         1100 J           ug/kg         63,000         810,000         11,000         69 J         560 J         370 U         48 J         100 J           ug/kg         6,700,000         11,000,0</td></td<>	ug/kg	ug/kg           350 U         21 J         370 U         350 U           ug/kg         180,000,000         530,000,000          81 J         190 J         370 U         17 J           ug/kg         63,000         1,000,000,000         11,000         190 J         1100         11 J         130 J           ug/kg         6,300         81,000         1,100         200 J         1300         16 J         180 J           ug/kg         63,000         810,000         11,000         200 J         100         10 J         140 J           ug/kg         63,000         8,100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         10,000,000         31,000,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         10,000,000         31,000,000         1100,000         350 U         22 J           ug/kg         6,700,000         41,000         1,100         28 J         160 J         370 U         24 J           ug/kg         6,700         41,000         1,100         2300         350 U         29 J         290 J           ug/kg	ug/kg             350 U         17 J           ug/kg         180,000,000         530,000,000          86 J         190 J         370 U         17 J           ug/kg         63,000         1,000,000,000         1,1000         150 J         1100         11 J         130 J           ug/kg         6,300         81,000         1,100         200 J         1300         16 J         140 J           ug/kg         6,300         81,000         1,100         200 J         1300         16 J         140 J           ug/kg         6,300         81,000         11,000         69 J         560 J         8.9 J         100 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         630,000         8,100,000         110,000         69 J         560 J         8.9 J         100 J           ug/kg         6,700,000         41,000,000         1,100,000         15 J         100 J         15 J         160 J           ug/kg         6,700         41,000,000         1,100         23 J         100 J         15 J         10 J	ug/kg	ug/kg	ug/kg         E30,000,000         E30 U         21 J         370 U         17 J           ug/kg         180,000,000         530,000,000         -         36 J         190 J         370 U         17 J           ug/kg         880,000,000         11,000         11,000         150 J         1100	ug/kg         630,000,000          980 U         21 J         370 U         360 U           ug/kg         680,000,000         1,000,000,000         11,000         181 J         280 J         370 U         17 J           ug/kg         63,000         810,000         1,100         190 J         1100         110 J         130 J           ug/kg         63,000         810,000         11,000         190 J         1100         1100 J         1100 J           ug/kg         63,000         810,000         11,000         69 J         560 J         370 U         48 J         100 J           ug/kg         6,700,000         11,000,0

-- Standard not available

 $\mathsf{U}$  = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

NA = Not analyzed

(1) VAP Generic Direct Contact Soil Standard, Commercial/Industrial Land Use

(2) VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Table 6-3B
Summary of Detected Chemicals in Groundwater
Site 14 - Bojacks Meats
ODOT Innerbelt Study
Cleveland, Ohio

۵	PARAMETER.	UNITS	VAP UPUS/ RDUPUS <sup>(1)</sup>	BUSTB Closure Action Level	14-MW-02 09/26/2006	14-MW-02D 09/26/2006	14-MW-03 09/26/2006
	2-Butanone	ng/L	9890	1	0.76 J	0.52 J	0.52 J
	Bromodichloromethane	ug/L	1	ı	1.0 U	1.0 U	7
	Chloroform	ug/L	50	**	5.6	5.9	6.0
	Benzo(a)anthracene	ng/L	ı	0.26	0.79	0.46	0.20
5	Benzo(a)pyrene	ug/L	0.2	0.2	0.63	0.43	0.20
SOC	Benzo(b)fluoranthene	ug/L		0.17		0.56	0.20 U
ΛS	Benzo(ghi)perylene	ug/L	1	•	0.52	0.20 U	0.20
	Chrysene	ng/L	47	47	0.75	0.51	0.20
	Fluoranthene	ug/L	370	1	2.0	1.2	0.20 ∪
	Indeno(1,2,3-cd)pyrene	ug/L		0.22	0.95	0.20 U	0.20 U
	Phenanthrene	ug/L		-	1.1	0.59	0.20
	Pyrene	ug/L	280	:	1.8	7.	0.20 U

-- = Standard not available

 $U \approx The$  analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

(1) VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

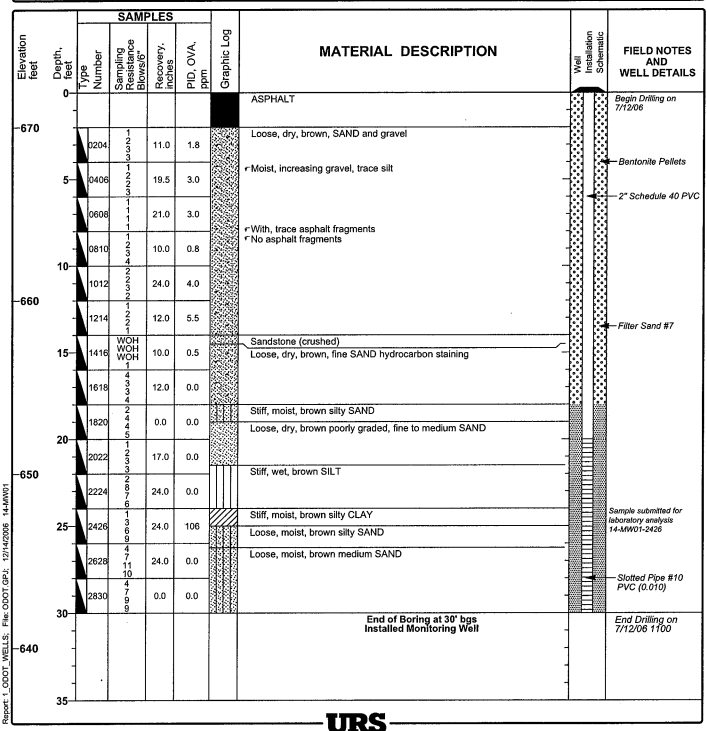
BORING LOGS

Project: ODOT - Innerbelt Corridor

Project Location: Site 14
Project Number: 15016633

# Log of Boring 14-MW01

Date(s) 7/12/06 Drilled 7/12/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# / 30" drop automatic	Total Depth of Borehole 30.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 672'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured Well dry on 9/26/06			

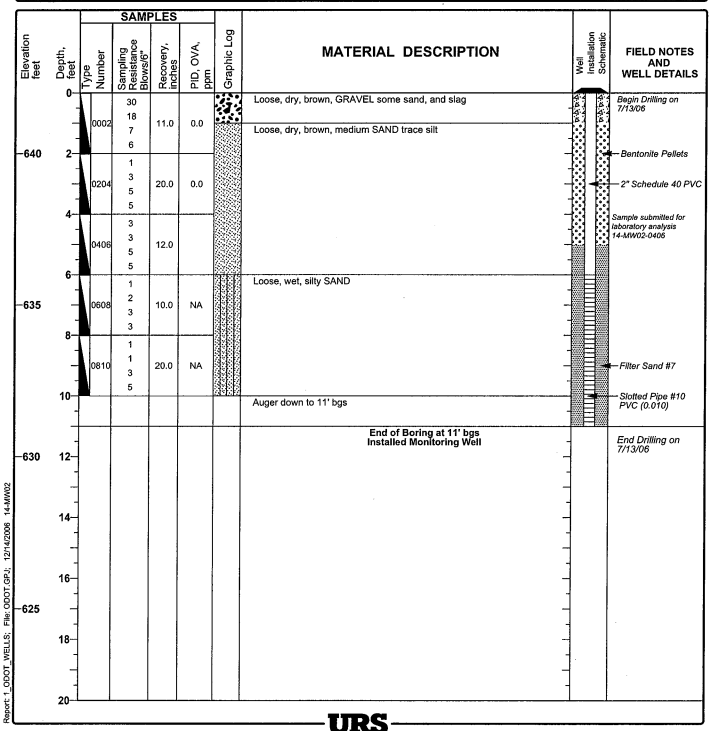


Project: ODOT - Innerbelt Corridor

Project Location: Site 14 Project Number: 15016633

## Log of Boring 14-MW02

Date(s) 7/13/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# / 30" drop automatic	Total Depth of Borehole 11.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 642'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 634.72 on 9/26/06			

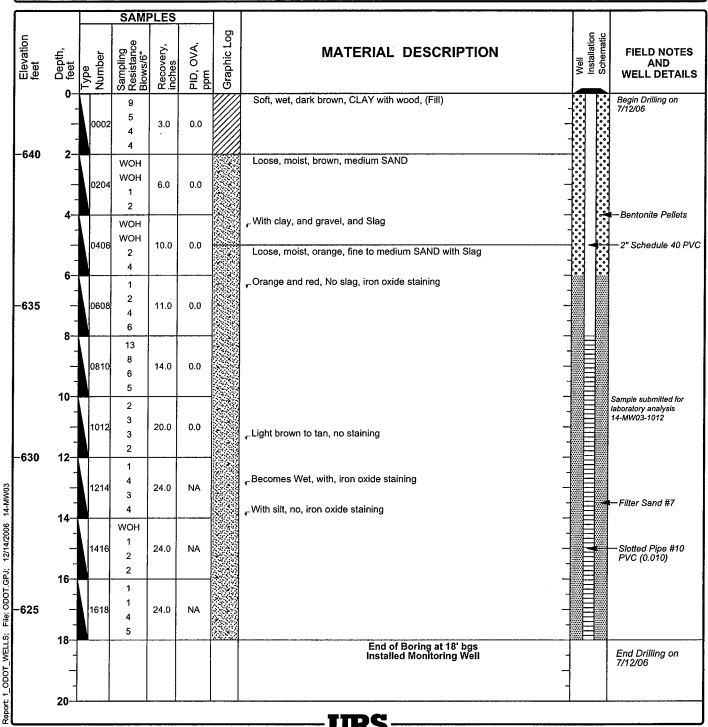


Project: ODOT - Innerbelt Corridor

Project Location: Site 14
Project Number: 15016633

## Log of Boring 14-MW03

Date(s) 7/12/06	Logged By J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer 140# auto hammer	Total Depth of Borehole 18.0′ bgs
Drill Rig CME-55 Type	Drilling Contractor HAD, Inc.	Approximate Ground Elevation 642'
Location See Site Map	Sampling 2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 626.86 on 9/26/06		



Project: ODOT - Innerbelt Corridor
Project Location: Site 14
Project Number: 15016633

## Log of Boring 14-SB04

Date(s) Drilled and Installed 8/10/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0' bgs
Sampling 2" Split Spoon Method	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 676'
Drill Rig Type: LC-60	Groundwater Not encountered Level(s)	Hammer 140# auto hammer
Boring See Site Map	Borehole Backfill <b>bentonite</b>	

1				SAM	LES				
	Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
	_	0-	0002	7	12.0	1.9		Loose, dry, black, gravelly SAND with Slag	Begin drilling on 8/10/06
		2-	0204	2222	16.0	0.4		Loose to medium dense, dry, brown, fine SAND	
	670	4-	0406	WOH 1 WOH 1	17.0	1.0		_	
	<b>−670</b>	6 <u>-</u> - 8-	0608	1 1 1	10.0	1.2		rTrace gravel	
		0- - 10-	0810	1 1 1	12.0	1.4			·
İ	-	10 -	1012	1 1 1	12.0	1.3		rNo gravel	
		14-	1214	1 3 6 8	12.0	1.0		rTrace gravel	
	-660	16-	1416	1 8 7 7	18.0	1.1		rCoarse SAND rNo gravel	
		18-	1618	1 5 6 7	20.0	2.1		: - -	
		20	1820	1 5 8 8	11.0	1.6		-	
	<del>-</del>	22-	2022	2 5 7 7	14.0	2.4		· -	Sample submitted for laboratory analysis 14-SB04-2022
3 14-SB0		24-	2224	1 2 3	22.0	0.7		Stiff, moist, brown, SILT  Dense, moist, brown, fine SAND trace, iron oxide staining	
12/14/2006 14-SB04	-650	26-	2426	1 3 3 4	15.0	0.9		בייייייייייייייייייייייייייייייייייייי	
ODOT.GPJ;		28-	2628	1 2 3 4	16.0	0.6		<u>-</u>	
File:		30	2830	2356	16.0	0.6			
30RINGS;	-	32-						End of Boring at 30' bgs	End drilling on 8/10/06
Report: 1_ODOT_BORINGS;		34-							
Report: 1		_1						URS	

DATA ASSESSMENT REPORT

#### Data Assessment Report ODOT Innerbelt Study Site 14 – Bojacks Meats

Reviewer: P. Schuler Date: November 12, 2006

Five soil samples, three groundwater samples, two equipment blanks, and two trip blanks were collected at the Bojacks Meats site in Cleveland, Ohio, on July 12 through September 26, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Re	Requested Analyses <sup>(1)</sup>		es <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC	TPH	CN
A6G140117001	14-MW01-2426	07/12/2006	Soil	X	X	X	X
A6G140117002	14-MW03-1012	07/12/2006	Soil	X	X	X	X
A6G140394001	14-MW02-0406	07/13/2006	Soil	X	X	X	X
A6H120114001	14-SB04-2022	08/10/2006	Soil	X	X	X	X
A6H120114002	14-SB04-2022D	08/10/2006	Soil			X	X
A6H120114003	14-SB04-EB	08/10/2006	Equip. Blank	X	X	X	X
A6H120114004	TB-081006	08/10/2006	Trip Blank	X			
A6I270118001	14-MW-02	09/26/2006	Groundwater	X	X		X
A6I270118002	TB-092606	09/26/2006	Trip Blank	X			
A6I270118003	14-MW-03	09/26/2006	Groundwater	X	X		X
A6I270118004	EB092506	09/25/2006	Equip. Blank	X	X		X
A6I270118005	14-MW-02D	09/26/2006	Groundwater	X	X		X

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B]

SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

CN = Total Cyanide [SW-846 Method 9012A]

(2) Samples 14-SB04-2022and 14-SB04-2022D are field duplicates (for selected analytes only).

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

Positive detections for methylene chloride in samples 14-MW01-2426, 14-MW03-1012,



and 14-MW02-0406, for acetone in sample 14-SB04-2022, for gasoline range organics in sample 14-MW02-0406, and for cyanide in samples 14-SB04-2022 and 14-SB04-2022D were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks at similar concentrations.

- Positive detections for methylene chloride in sample 14-SB04-2022 and for bis(2-ethylhexyl)phthalate in samples 14-MW-03 and 14-MW-02D were qualified as nondetect ("U") due to the presence of these analytes in the associated equipment blank or trip blank at a similar concentration.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J"), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, the "J" qualifiers were retained with the numeric results.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-5.

# Table 2-1 Analytical Data Summary Site 14 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140117001 14-MW01-2426 07/12/2006	A6G140117002 14-MW03-1012 07/12/2006	A6G140394001 14-MW02-0406 07/13/2006	A6H120114001 14-SB04-2022 08/10/2006	A6H120114003 14-SB04-EB 08/10/2006 ug/L	A6H120114004 TB-081006 08/10/2006 ug/L
1,1,1-Trichloroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 U
1,1,2-Trichloroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 U
1,1-Dichloroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,1-Dichloroethene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 ∪
1,2-Dibromo-3-chloropropane	ug/kg	12 U	11 U	11 U	10 U	2.0 ∪	2.0 U
1,2-Dibromoethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,2-Dichlorobenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,2-Dichloroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 U
1,2-Dichloropropane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
1,4-Dichlorobenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 U
2-Butanone	ug/kg	23 U	22 U	22 U	21 U	10 U	10 U
2-Hexanone	ug/kg	23 U	22 U	22 U	21 U	10 U	10 U
4-Methyl-2-pentanone	ug/kg	23 U	22 U	22 U	21 U	10 U	10 U
Acetone	ug/kg	23 U	22 U	22 U	21 U	10 U	10 U
Benzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Bromodichloromethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 U
Bromoform	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
romomethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
arbon disulfide	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Carbon tetrachloride	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Chlorobenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Chloroethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Chloroform	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Chloromethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
cis-1,2-Dichloroethene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 ∪	1.0 U
cis-1,3-Dichloropropene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Cyclohexane	ug/kg	12 U	11 U	11 U	10 U	1.0 U	1.0 U
Dibromochloromethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Dichlorodifluoromethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Ethylbenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Isopropylbenzene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Methyl acetate	ug/kg	12 U	11 U	11 U	10 U	10 U	10 U
Methyl tert-butyl ether	ug/kg	23 U	22 U	22 U	21 U	5.0 U	5.0 U
Methylcyclohexane	ug/kg	12 U	11 U	11 U	10 U	1.0 U	1.0 U
Methylene chloride	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	2.5	0.39 J
Styrene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Tetrachloroethene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Toluene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
trans-1,2-Dichloroethene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Trichloroethene	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Trichlorofluoromethane	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
Vinyl chloride	ug/kg	5.9 U	5.6 U	5.4 U	5.2 U	1.0 U	1.0 U
γlenes (total)	ug/kg	12 U	11 U	11 U	10 U	2.0 U	2.0 U

ne analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-2 Analytical Data Summary Site 14 Soil Semivolatiles ODOT Innerbelt Study

		A6G140117001 14-MW01-2426 07/12/2006	A6G140117002 14-MW03-1012 07/12/2006	A6G140394001 14-MW02-0406 07/13/2006	A6H120114001 14-SB04-2022 08/10/2006	A6H120114003 14-SB04-EB 08/10/2006 ug/L
PARAMETER	UNITS	weet in the file of the			!	
1,1'-Biphenyl	ug/kg	390 U	370 U	710 U	350 U	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/kg	390 U	370 U	710 U	350 U	1.0 U
2,4,5-Trichlorophenol	ug/kg	390 U	370 U	710 U	350 U	5.0 U
2,4,6-Trichlorophenol	ug/kg	390 U	370 U	710 U	350 U	5.0 U
2,4-Dichlorophenol	ug/kg	390 U	370 U	710 U	350 U	2.0 U
2,4-Dimethylphenol	ug/kg	390 U	370 U	710 U	350 U	2.0 U
2,4-Dinitrophenol	ug/kg	390 U	370 U	710 U	350 U	5.0 U
2,4-Dinitrotoluene	ug/kg	390 U	370 U	710 U	350 U	5.0 U
2,6-Dinitrotoluene	ug/kg	390 U	370 ∪	710 U	350 U	5.0 U
2-Chloronaphthalene	ug/kg	390 U	370 U	710 U	350 U	1.0 U
2-Chlorophenol	ug/kg	390 U	370 U	710 U	350 U	1.0 U
2-Methylnaphthalene	ug/kg	390 U	370 U	21 J	350 U	0.20 U
2-Methylphenol	ug/kg	390 U	370 U	710 U	350 U	1.0 U
2-Nitroaniline	ug/kg	390 U	370 U	710 U	350 U	2.0 U
2-Nitrophenol	ug/kg	390 U	370 U	710 U	350 U	2.0 U
3,3'-Dichlorobenzidine	ug/kg	390 U	370 U	710 U	350 U	5.0 U
3-Nitroaniline	ug/kg	390 U	370 U	710 U	350 U	2.0 U
4,6-Dinitro-2-methylphenol	ug/kg	390 U	370 U	710 U	350 U	5.0 U
4-Bromophenyl phenyl ether	ug/kg	390 U	370 U	710 U	350 U	2.0 U
4-Chloro-3-methylphenol	ug/kg	390 U	370 U	710 U	350 U	2.0 U
4-Chloroaniline	ug/kg	390 U	370 U	710 U	350 U	2.0 U
4-Chlorophenyl phenyl ether	ug/kg	390 U	370 U	710 U	350 U	2.0 U
4-Methylphenol	ug/kg	390 U	370 U	710 U	350 U	1.0 U
4-Nitroaniline	ug/kg	390 U	370 U	710 U	350 U	2.0 U
4-Nitrophenol	ug/kg	390 U	370 U	710 U	350 U	5.0 U
Acenaphthene	ug/kg	36 J	370 U	190 J	17 J	0.20 U
Acenaphthylene	ug/kg	390 U	370 U	710 U	350 U	0.20 U
Acetophenone	ug/kg	78 U	75 U	140 U	70 U	1.0 U
Anthracene	ug/kg	81 J	370 U	280 J	36 J	0.20 U
Atrazine	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Benzaldehyde	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Benzo(a)anthracene	ug/kg	190 J	11 J	1100	130 J	0.20 U
Benzo(a)pyrene	ug/kg	150 J	10 J	1000	140 J	0.20 U
Benzo(b)fluoranthene	ug/kg	200 J	16 J	1300	180 J	0.20 U
Benzo(ghi)perylene	ug/kg	88 J	8.9 J	560 J	100 J	0.20 U
Benzo(k)fluoranthene	ug/kg	69 J	370 U	560 J	88 J	0.20 U
bis(2-Chloroethoxy)methane	ug/kg	390 U	370 U	710 U	350 U	1.0 U
bis(2-Chloroethyl) ether	ug/kg	390 U	370 U	710 U	350 U	1.0 U
bis(2-Ethylhexyl) phthalate	ug/kg	390 U	370 U	710 U	350 U	1.4
Butyl benzyl phthalate	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Caprolactam	ug/kg	390 U	48 J	710 U	350 U	5.0 U
Carbazole	ug/kg	23 J	370 U	200 J	22 J	1.0 U
Chrysene	ug/kg	180 J	15 J	1000	160 J	0.36
Dibenz(a,h)anthracene	ug/kg	28 J	370 U	160 J	24 J	0.20 U
Dibenzofuran	ug/kg	390 U	370 U	76 J	350 U	1.0 U
Diethyl phthalate	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Dimethyl phthalate	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Di-n-butyl phthalate	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Di-n-octyl phthalate	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Fluoranthene	ug/kg ug/kg	410	29 J	2300	290 J	0.88
Fluorene	ug/kg	24 J	370 U	150 J	13 J	0.20 U

Table 2-2 Analytical Data Summary Site 14 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140117001 14-MW01-2426 07/12/2006	A6G140117002 14-MW03-1012 07/12/2006	A6G140394001 14-MW02-0406 07/13/2006	A6H120114001 14-SB04-2022 08/10/2006	A6H120114003 14-SB04-EB 08/10/2006 ug/L
Hexachlorobenzene	ug/kg	390 U	370 U	710 U	350 U	0.20 U
Hexachlorobutadiene	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Hexachlorocyclopentadiene	ug/kg	390 U	370 U	710 U	350 U	10 U
Hexachloroethane	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/kg	80 J	7.7 J	520 J	85 J	0.20 U
Isophorone	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Naphthalene	ug/kg	8.0 J	370 U	15 J	9.6 J	0.20 U
Nitrobenzene	ug/kg	390 U	370 U	710 U	350 U	1.0 U
N-Nitrosodi-n-propylamine	ug/kg	390 U	370 U	710 U	350 U	1.0 U
N-Nitrosodiphenylamine	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Pentachiorophenol	ug/kg	390 U	370 U	710 U	350 U	5.0 U
Phenanthrene	ug/kg	300 J	21 J	1400	150 J	0.71
Phenol	ug/kg	390 U	370 U	710 U	350 U	1.0 U
Pyrene	ug/kg	340 J	21 J	1900	250 J	0.57

# Table 2-3 Analytical Data Summary Site 14 Soil TPH and Cyanide ODOT Innerbelt Study

PARAMETER	UNITS	A6G140117001 14-MW01-2426 07/12/2006	A6G140117002 14-MW03-1012 07/12/2006	A6G140394001 14-MW02-0406 07/13/2006	A6H120114001 14-SB04-2022 08/10/2006	A6H120114002 14-SB04-2022D (field dup) 08/10/2006	A6H120114003 14-SB04-EB 08/10/2006 mg/L
Gasoline Range Organics (C6-C12)	ug/kg	120 U	110 U	110 U	100 U	110 U	33 J
C10-C20	mg/kg	4.7	2.2	22 U	4.7	6.7	50 U
C20-C34	mg/kg	28	8.5	40	18	29	80 U
Cyanide, Total	mg/kg	0.59 U	0.56 U	0.54 U	0.52 U	0.53 U	0.010 U
Percent Solids	%	85.3	89	92.4	95.3	94.3	

#### Table 2-4 Analytical Data Summary Site 14 Water VOCs & Cyanide ODOT Innerbelt Study

		A6I270118001 14-MW-02	A6I270118002 TB-092606	A6I270118003 14-MW-03	A6I270118004 EB092506	A6I270118005 14-MW-02D 09/26/2006
PARAMETER	UNITS	09/26/2006	09/26/2006	09/26/2006	09/25/2006	
1,1,1-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 Ų	1.0 U	1.0 U	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
2-Butanone	ug/L	0.76 J	10 U	0.52 J	10 U	0.52 J
2-Hexanone	ug/L	10 U	10 U	10 U	10 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U	10 U	10 U	10 U
Acetone	ug/L	10 U	10 U	10 U	10 U	10 U
Benzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U	1.1	1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chloroform	ug/L	5.6	1.0 U	6	1.0 U	5.9
Chloromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Dibromochloromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U	10 U	10 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Methylene chloride	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Styrene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
en en regiment de la companya de la companya de la companya de la companya de la companya de la companya de la	ug/L	1.0 U	0.73 J	1.0 U	1.0 U	1.0 U
Toluene trans-1,2-Dichloroethene	ug/L ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
and the contract of the contra	Water Or State of States	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	44.4		,	221 - 1 to 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	
Trichlorofluoromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
Cyanide, Total	mg/L	0.010 U	-	0.010 U	0.010 U	0.010 U

<sup>-- =</sup> Not analyzed.



U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-5 Analytical Data Summary Site 14 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I270118001 14-MW-02 09/26/2006	A6I270118003 14-MW-03 09/26/2006	A6I270118004 EB092506 09/25/2006	A6I270118005 14-MW-02D 09/26/2006
		· · · · · · · · · · · · · · · · · · ·	1 Jan 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		1.3 47 3
1,1'-Biphenyl	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2,4,5-Trichlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,4,6-Trichlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,4-Dichlorophenol	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
2,4-Dimethylphenol	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
2,4-Dinitrophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,4-Dinitrotoluene	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,6-Dinitrotoluene	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2-Chloronaphthalene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2-Chlorophenol	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2-Methylnaphthalene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
2-Methylphenol	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
2-Nitrophenol	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
3,3'-Dichlorobenzidine	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
3-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4,6-Dinitro-2-methylphenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
4-Bromophenyl phenyl ether	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Chloro-3-methylphenol	ug/L	2.0 U	2.0 ∪	2.0 U	2.0 U
4-Chloroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Chlorophenyl phenyl ether	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Methylphenol	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
4-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Nitrophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Acenaphthene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Acenaphthylene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Acetophenone	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Anthracene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Atrazine	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Benzaldehyde	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Benzo(a)anthracene	ug/L	0.79	0.20 U	0.20 U	0.46
Benzo(a)pyrene	ug/L	0.63	0.20 U	0.20 U	0.43
Benzo(b)fluoranthene	ug/L	1.1	0.20 U	0.20 U	0.56
Benzo(ghi)perylene	ug/L	0.52	0.20 U	0.20 U	0.20 U
Benzo(k)fluoranthene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
bis(2-Chloroethoxy)methane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
bis(2-Chloroethyl) ether	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
bis(2-Ethylhexyl) phthalate	ug/L	1.0 U	1.3 U	0.90 J	1.1 U
Butyl benzyl phthalate	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Caprolactam	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Carbazole	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Chrysene	ug/L	0.75	0.20 U	0.20 U	0.51
Dibenz(a,h)anthracene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Dibenzofuran	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Diethyl phthalate	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Dimethyl phthalate	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Di-n-butyl phthalate	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Di-n-octyl phthalate	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Fluoranthene	ug/L	2.0	0.20 U	0.20 U	1.2
Fluorene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Hexachlorobenzene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U

# Table 2-5 Analytical Data Summary Site 14 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I270118001 14-MW-02 09/26/2006	A6I270118003 14-MW-03 09/26/2006	A6I270118004 EB092506 09/25/2006	A6I270118005 14-MW-02D 09/26/2006
Hexachlorobutadiene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U	10 U	10 U	10 U
Hexachloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.95	0.20 U	0.20 U	0.20 U
Isophorone	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Naphthalene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Nitrobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Pentachlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Phenanthrene	ug/L	1.1	0.20 U	0.20 U	0.59
Phenol	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Pyrene	ug/L	1.8	0.20 U	0.20 U	1.2

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

#### 6.4 SITE 15 – LEON RUDNICK

According to the Cleveland City Directories, the site was listed as Baum Ice Cream from 1954 through 1979. According to the Cleveland Fire Prevention Bureau files, the site was occupied by Baum Ice Cream Company in 1944. An entry dated June 1950 indicates a fuel oil UST was to be installed in the tree lawn. A Fire Inspection Report dated December 1965 indicates a 1,000-gallon fuel oil UST was located on the site.

A Fire Inspection Report dated February 1987 indicates the 1,000-gallon UST was to be abandoned in place. On February 12, 1987, the UST reportedly was filled with a slurry mixture and abandoned in place, under the supervision of the Fire Department.

At the time of the URS Phase I reconnaissance, the site was occupied by one warehouse building, which was divided into three units. No surface staining, stressed vegetation and/or the storage and handling of hazardous substances were observed.

#### 6.4.1 Field Activities

No intrusive Phase II activities were requested to be conducted at this site. No information is available regarding site specific geology. A Site Location Map is included as **Figure 6-4A**.

#### 6.4.2 Geophysical Survey

URS conducted a geophysical survey at Site 15 in July 2006. The purpose of the survey was to locate, if present, abandoned underground storage tanks (USTs). Figure 6-4B indicates the results of the survey.

The geophysical survey at Site 15 included the paved and vegetated surfaces of the property and the pavement north of the property. The survey transects were conducted in a north – south direction, west of the building to align transects along the earth's total magnetic field which provides the greatest sensitivity to buried ferrous objects such as steel underground storage tanks. Due to the orientation of the roadway north of the property, transects in the roadway were conducted in an east-west direction. This orientation likely reduced the sensitivity of the survey. Transects were spaced five feet apart and were walked in alternating directions. The presence of tall structures prevented the use of a global positioning system (GPS) for the establishment of coordinates.

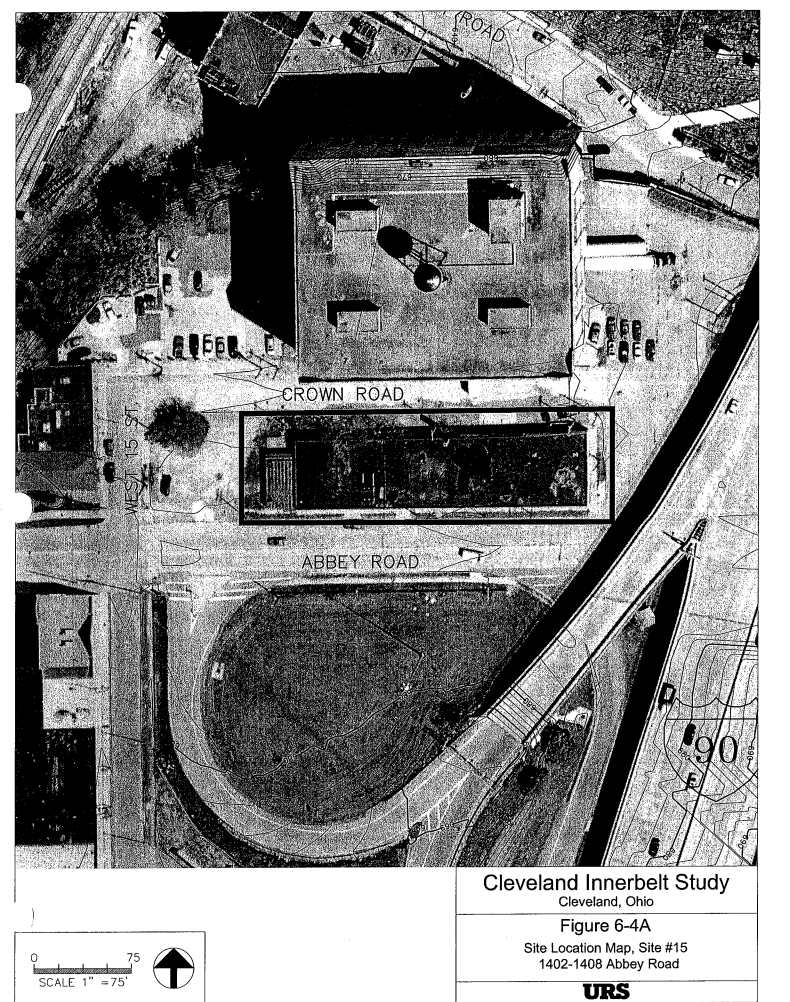
No evidence of a UST was discovered as a result of the geophysical survey at Site 15.

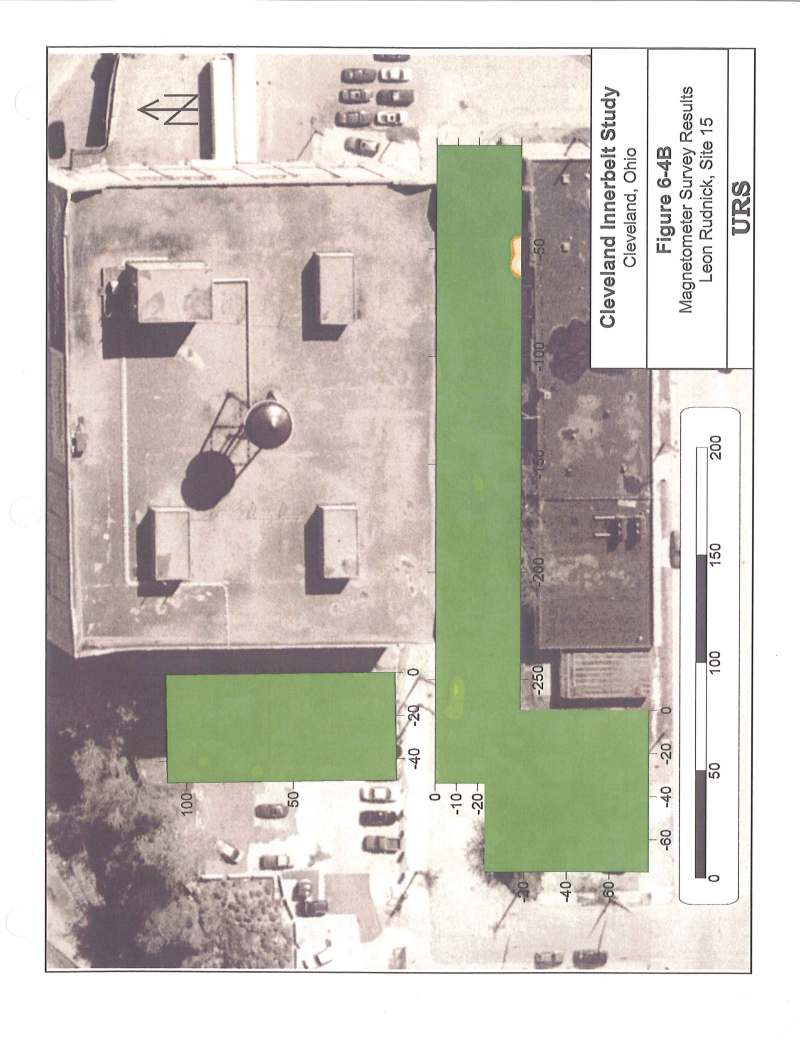
#### 6.4.3 Conclusions

It is unlikely the soil and groundwater at Site 15 would impede the planned construction; however, the project specifications could be revised to include a plan note indicating the potential for impacted soil and groundwater to be encountered during the course of excavation. The contractor conducting the proposed improvement project should be advised to plan

accordingly to protect the workers and to manage the exposure to soil and groundwater in the event such soil or groundwater is encountered.

**FIGURES** 





#### 6.5 SITE 16 - WENDELL & CARROLL COLLINS

According to the Cleveland Fire Prevention Bureau files, the site was occupied by Diamond Wheel Fabricating in 1999.

At the time of the URS Phase I reconnaissance, the site was occupied by a building and gravel-covered land. Surface staining was observed on the gravel-covered portion of the site.

#### 6.5.1 Field Activities

A total of four monitoring wells were scoped for Site 16. Groundwater was encountered in only one soil boring (MW01) during field activities. A total of three soil borings (SB02 through SB04) were installed at Site 16. At the time of groundwater sampling, MW01 was dry and no groundwater sample could be obtained. One soil sample per boring was collected and analyzed for VOCs, SVOCs, and RCRA Metals. A Sample Location Map is included as **Figure 6-5A**.

#### 6.5.2 Site-Specific Geology/Hydrogeology

Soils at Site 16 consisted primarily of sand with minor amounts of silty clay and clay interbedded. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 16 on September 26, 2006. Monitoring well (MW01) was dry. Localized groundwater flow across Site 16 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 16 is to the south-southwest.

#### 6.5.3 Soil Analytical Results

No VOCs were detected in the samples submitted from Site 16.

A total of eleven SVOCs were detected in two (16-SB02-1820 and 16-SB04-0810) of the four samples submitted from Site 16. Concentrations of benzo(a)anthracene (34 ug/kg), benzo(a)pyrene (37 ug/kg), benzo(b)fluoranthene (49 ug/kg), benzo(g,h,i)perylene (28 ug/kg), benzo(k)fluoranthene (22 ug/kg), bis(2-ethylhexyl)phthalate (480 ug/kg), chrysene (ranging from 8.7 ug/kg to 39 ug/kg), fluoranthene (ranging from 8.0 ug/kg to 68 ug/kg), indeno(1,2,3)pyrene (24 ug/kg), phenanthrene (29 ug/kg), and pyrene (52 ug/kg) were detected in the soil samples submitted. All other SVOCs were below the detection limits.

Six of the eight RCRA Metals were detected in the samples submitted from Site 16. Arsenic, ranging from 5.9 mg/kg to 8.5 mg/kg, was detected in all samples submitted. Barium, ranging from 12.0 mg/kg to 30.2 mg/kg, was detected in the all samples submitted. Cadmium, 0.15 mg/kg to 0.33 mg/kg, was detected in all samples submitted. Chromium, ranging from 4.9 mg/kg to 7.4 mg/kg, was detected in the samples submitted. Lead, ranging from 5.4 mg/kg to 33 mg/kg, was detected in the samples submitted. Mercury, 0.043 mg/kg, was detected in the sample submitted from 16-SB04-0810.

The analytical results are presented in Table 6-5A.

#### 6.5.4 Groundwater Analytical Results

No groundwater was encountered in MW02.

#### 6.5.5 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), and the BUSTR Closure Action Levels for Class 1 soils. The VAP and BUSTR standards are included on **Table 6-5A**.

No VOCs were detected in the samples submitted from Site 16.

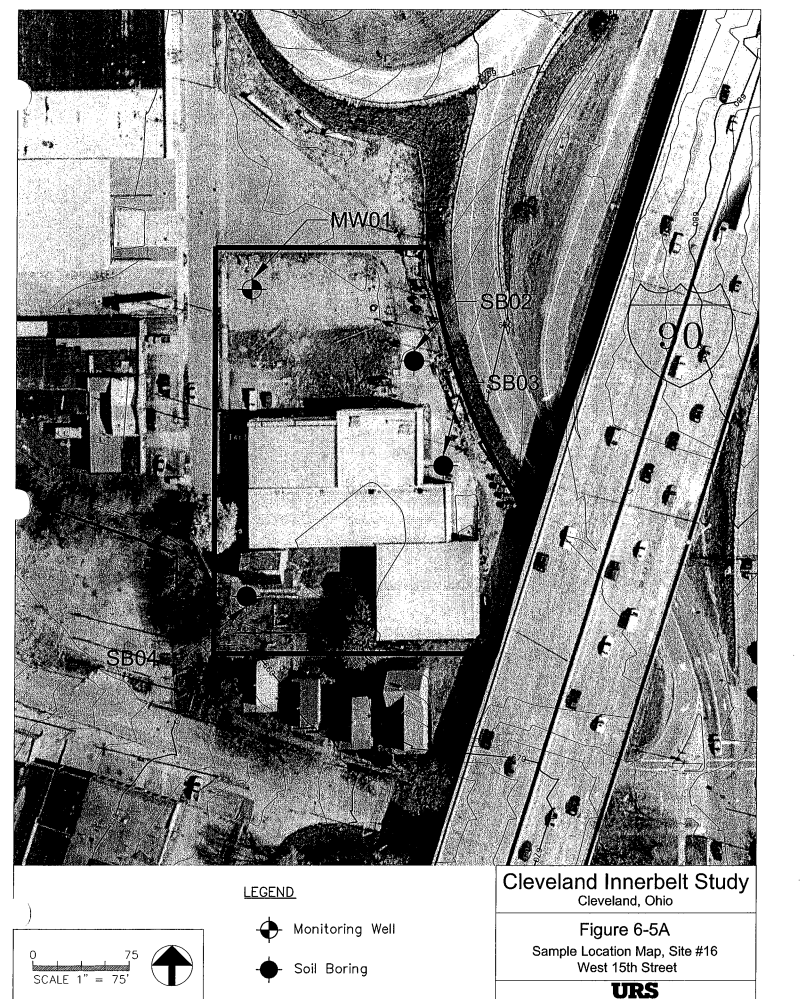
None of the eleven SVOCs detected at Site 16 exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or BUSTR closure action levels.

There were no RCRA Metals detected at Site 16 which exceeded the OEPA VAP standards for commercial and industrial land use or the construction and excavation worker activities category.

#### 6.5.6 Conclusions

Based on the analytical results, it is unlikely that the soils at Site 16 would require special management during construction.

FIGURES



## TABLES

# Table v-5A Summary of Detected Chemicals in Soil Site 16 - Wendell & Carroll Collins / 1501 Companies ODOT Innerbelt Study Cleveland, Ohio

74	PARAMETER	SLINO	VAP Commercial/ Industrial	VAP Construction Warken Standard	BUSTR Closure Action Level	16-WW01-1416 07/11/2006	16-SB02-1820 07/11/2006	16-SB03-2224 07/11/2006	16-SB04-0810 08/07/2006
	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	350 U	350 U	370 U	34 J
Ĭ.	Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	350 U	350 U	370 U	37 J
·	Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	350 U	350 U	370 U	49 J
ì	Benzo(ghi)perylene	ug/kg	CONTROL CONTRO	•	-	350 U	350 U	370 U	28 J
sO	Benzo(k)fluoranthene	ug/kg	630,000	8,100,000	110,000	350 U	350 U	370 U	22 J
ΟΛ	bis(2-Ethylhexyl) phthalate	ug/kg	230,000	230,000	The control of the co	350 U	350 U	370 U	480
S	Chrysene Chrysene Chrysene Chrysene Chrysene Chrysene	ug/kg	6,700,000	41,000,000	1,100,000	350 U	8.7 J	370 U	39 J
3	Fluoranthene	ug/kg	33,000,000	170,000,000	•	350 U	8.0 J	370 U	£ 89
	Indeno(1,2,3-cd)pyrene	ug/kg	67,000	410,000	11,000	350 U	350 U	370 U	24 J
	Phenanthrene	ug/kg	The second secon		1	350 U	320 ∩	370 U	29 J
	Pyrene	ug/kg	25,000,000	130,000,000	**	350 U	350 U	370 U	52 J
	Arsenic	mg/kg	80	210	•	5.9	7.1	7.5	8.5
	Barium	mg/kg	200,000	45,000	;	20.9 J	12.0 J	13.2 J	30.2
ials	Cadmium	mg/kg	23	420	TO SERVICE A CONTRACTOR OF THE SERVICE AND A	0.17	0.33	0.19 J	0.15 J
эM	Chromium	mg/kg	8,900	2,000	The standards and the standard	6.1	6.4	4.9	7.4
	Lead	mg/kg	The state of the s		THE LAND TO A PROPERTY AND STREET AND A STRE	5.4	6.7	6.3	33
	Mercury	mg/kg	300	84	No. 17, and 27 a	0.11 U	0.11 U	0.11 U	0.043 .1

-- = Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

(1) VAP Generic Direct Contact Soil Standard, Commercial/ Industrial Land Use

(2) VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

**BORING LOGS** 

Project: ODOT - Innerbelt Corridor

Project Location: Site 16
Project Number: 15016633

## Log of Boring 16-MW01

Date(s) 7/11/06 Drilled 7/11/06	Logged By	J. Kaminski	Checked M. Wolff By
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# / 30" drop automatic	Total Depth of Borehole 30.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 682'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured Well dry on 9/26/06			

ſ				SAME	PLES				Τ				
	Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION		Installation			
		-	0002	32 9 7 5	4.0	0.0	<b>***</b>	ASPHALT Brick, and Slag, (Fill)	A D D D	X 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Begin Drilling on 7/11/06		
Ī	-680	-	0204	4 2 2 2	5.0	1.0		Loose, moist, brown, well graded, SAND trace medium to coarse gravel					
		5-	0406	1 1 1 4	10.0	0.7		Loose, dry, brown, SAND and gravel, and Brick					
		-	0608	2 2 3 3	21.0	0.2		Loose, dry, brown, coarse SAND trace gravel, trace silt			Bentonite Pellets		
		-	0810	2 3 4 10	20.0	1.0		Soft, moist, brown, silty CLAY			2" Schedule 40 PVC		
		10-	1012	13 12 12 10	24.0	0.9	20872	Stiff, moist, brown, clayey SILT To silt  Very loose, dry, light brown, coarse SAND trace silt					
l	-670	1	1214	7 9 11 11	20.0	0.7		←With gravel					
		15-	1416	5 5 10 10	16.0	2.0		←No gravel, increasing silt			Sample submitted for laboratory analysis 16-MW01-1416		
		-			1618	1 5 5 8	8.0	0.4		←Trace shale fragments			
		-	1820	5 10 10 12	14.0	1.9		←Less silt, fine gravel	-				
		20— -	2022	1 2 2 3	24.0	1.1		Stiff, light brown and gray, fine SAND trace silt Soft, wet, dark brown, silty SAND					
16-MW01	-660	-	2224	2 3 5 7	10.0	1.4		Trace, iron oxide staining  Dense, moist, gray, silty SAND					
		25-	2426	3 5 8 9	14.0	1.0		←Becomes brownish gray  ←Trace, iron oxide staining			Filter Sand #7		
J; 12/14/2006		-	2628	3 7 7	24.0	1.2							
ODOT.GPJ;			2830	4 6 8 8	12.0	1.0					Slotted Pipe #10 PVC (0.010)		
LS; File: ODC		30					.J	End of Boring at 30' bgs Installed Monitoring Well		, 100	End Drilling on 7/11/06		
Report: 1_ODOT_WELLS;	-650	-							1		·		
port: 1_0[		35			<u>,</u>				1				
ď L								URS					

Project: ODOT - Innerbelt Corridor

Project Location: Site 16 Project Number: 15016633

## Log of Boring 16-SB02

Date(s) Drilled and Installed 7/11/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0′ bgs
Sampling 2" Split Spoon Method	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 678'
Drill Rig Type: CME-55	Groundwater Not Encountered Level(s)	Hammer Data 140# / 30" drop automatic
Boring See Site Map	Borehole Backfill bentonite	

			SAM	PLES						
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES		
	0-						CONCRETE and base	Begin drilling on 7/11/06		
_	2-	0204	8 7 5 3	24.0	0.1		Loose, dry, brown, silty SAND with gravel			
	4-	0406	1 3 5 5	24.0	1.6					
	6-	0608	4 4 4 3	24.0	2.9		Loose, dry, brown, coarse SAND with gravel, and silt			
-670	8-	0810	2 4 10 12	24.0	2.6		Medium stiff, dry, brown, silty CLAY  Medium stiff, moist, brown, silty CLAY			
	10-	1012	7 10 13 7	10.0	1.5		Trace sand Loose, dry, light brown, SAND and gravel			
-	12	1214	5 8 13 8	16.0	2.2		Loose, dry, gray, GRAVEL with sand			
	14	1416	4 10 8 6	24.0	3.5					
	16-	1618	1 3 3 3	17.0	1.8		Loose, dry, brown, medium SAND iron oxide staining			
-660	18	1820	2 2 3 3	12.0	4.3		,	Sample submitted for laboratory analysis 16-SB02-1820		
	20	2022	1 3 3 2	14.0	3.8		rNo staining			
-	22-	2224	3 3 3 3	14.0	1.5					
	24-	2426	3 2 3 4	12.0	2.5		-			
CEO.	26	2628	4 6 7 9	12.0	1.5					
-650	28- - 30-	2830	4 6 7 7	0.0	0.9		_			
	32-						End of Boring at 30' bgs	End drilling on 7/11/06 1500		
-	34						-			
	347						URS			

Project: ODOT - Innerbelt Corridor Project Location: Site 16

Project Number: 15016633

## Log of Boring 16-SB03

Date(s) Dr and Install	illed 7/11/06 ed	Logged By	J. Kaminski	Reviewer	M. Wolff
Drilling Method	Hollow Stem Auger	Drilling Contractor	HAD, Inc.	Total Depth of Borehole	30.0' bgs
Sampling Method	2" Split Spoon	Drill Bit Size/Type:	4-1/4" ID HSA	Approximete Surface Elevation	676'
Drill Rig Type:	CME-55	Groundwate Level(s)	er	Hammer 140#	/ 30" drop automatic
Boring Location:	See Site Map	Borehole Backfill	Set monitoring well		

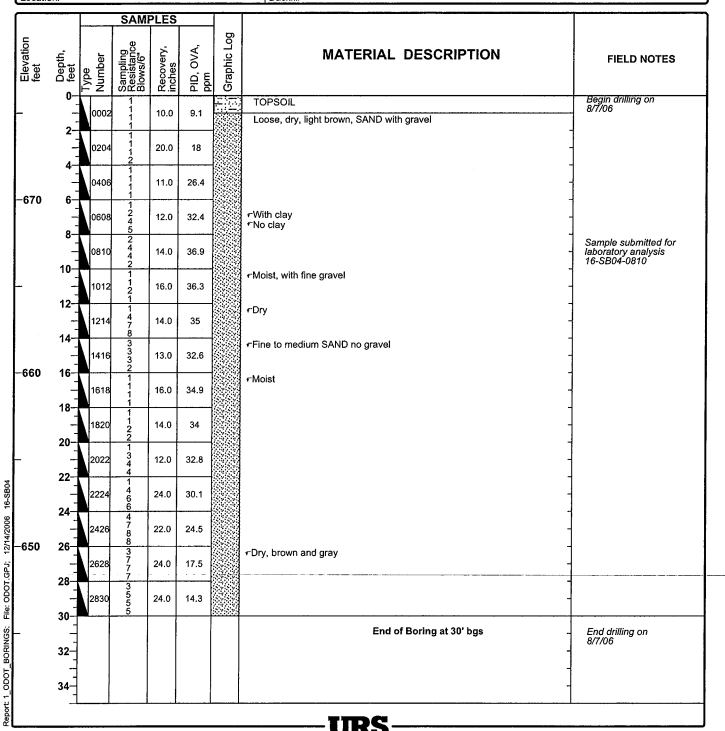
			SAM	PLES				
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
L	0-						CONCRETE	Begin drilling on 7/11/06
l	2		2				CRUSHED STONE	
	- - 4	0204	2 7 7 10	8.0	0.4		Hard, dry, brown and black, coarse clayey SAND and gravel, with Asphalt	-
-670	 6	0406	13	24.0	0.3		Loose, dry, brown, coarse SAND with gravel	 
	8-	0608	12	24.0	0.0		Hard, dry, brown, clayey SILT Soft, wet, Seam	-
	10-	0810	1 2 7 10	19.0	1.8		Hard, dry, brown, SILT trace fine sand	  -  -
-	12	1012	1 3 3 3	24.0	2.3		Loose, moist, gray coarse SAND	
	14	1214	4 4 3	17.0	2.1			
<b>–660</b>	16-	1416	3 4 5 5	20.0	1.8		Local day have fine to enading CAND trace of	
	18-	1618	3 2 3 3	13.0	2.9		Loose, dry, brown fine to medium SAND, trace silt	
	20	1820	3 3 2	19.0	1.5		rBecomes fine to medium SAND, trace silt	
_	22	2022	2 3 3 3	20.0	2.1			
-650	 24	2224	2 6 7 7	21.0	4.6			Sample submitted for laboratory analysis 16-SB03-2224
650	26	2426	2 7 7 8 7	21.0	2.0		rBecomes fine SAND	
	28-	2628	7 6 6	14.0	4.1		- ESSONIES INIC CANAD	
	30	2830	2 6 7 11	12.0	3.4		-	
-	32-						End of Boring at 30' bgs - - -	End drilling on 7/11/06 1230
	_1						—URS—	

**Project: ODOT - Innerbelt Corridor** 

Project Location: Site 16 Project Number: 15016633

## Log of Boring 16-SB04

Date(s) Drilled and Installed 8/7/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth 30.0′ bgs
Sampling 2" Split Spoon	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 676'
Drill Rig Type: LC-60	Groundwater Not Encountered Level(s)	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill <b>bentonite</b>	



DATA ASSESSMENT REPORT

# Data Assessment Report ODOT Innerbelt Study Site 16 – Wendell & Carroll Collins / 1501 Companies

Reviewer: P. Schuler Date: November 12, 2006

Four soil samples and one trip blank were collected at the Wendell & Carroll Collins / 1501 Companies site in Cleveland, Ohio, on July 11 and August 7, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Reque	ested Ana	lyses <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC	CN
A6G140116001	16-MW01-1416	07/11/2006	Soil	X	X	X
A6G140116002	16-SB02-1820	07/11/2006	Soil	X	X	X
A6G140116003	16-SB03-2224	07/11/2006	Soil	X	X	X
A6H090258001	16-SB04-0810	08/07/2006	Soil	X	X	X
A6H090258002	TB-080706	08/07/2006	Trip Blank	X		

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B] SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C] Met = RCRA Metals [SW-846 Methods 6010B/6020/7470A/7471A]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Positive detections for methylene chloride in samples 16-MW01-1416, 16-SB02-1820, and 16-SB03-2224 and for acetone in sample 16-SB04-0810 were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks and/or trip blanks at similar concentrations.
- The semivolatile compound, bis(2-ethylhexyl)phthalate, was detected at concentrations ranging from 0.9 to 1.8 ug/L in all equipment blanks associated with project samples. Since this compound is considered a common laboratory contaminant, its presence in environmental samples at concentrations less than ten times the highest equipment blank concentration (converted to an equivalent soil concentration) is attributed to external

contamination rather than actual site conditions. Therefore, the positive detections for bis(2-ethylhexyl)phthalate in samples 16-MW01-1416, 16-SB02-1820, and 16-SB03-2224 were qualified as nondetect ("U") at the sample reporting limit, even though it was not detected in the associated method blank.

- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J" for organics, and "B" for inorganics), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, organic "J" qualifiers were retained with the numeric results. The inorganic "B" qualifiers were changed to "J" qualifiers for the sake of consistency throughout the package.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-3.

Table 2-1
Analytical Data Summary
Site 16 Soil Volatiles
ODOT Innerbelt Study

PARAMETER	UNITS	A6G140116001 16-MW01-1416 07/11/2006	A6G140116002 16-SB02-1820 07/11/2006	A6G140116003 16-SB03-2224 07/11/2006	A6H090258001 16-SB04-0810 08/07/2006	UNITS	A6H090258002 TB-080706 08/07/2006
	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,1,1-Trichloroethane	ug/kg ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,1,2,2-Tetrachloroethane		5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,1,2-Trichloroethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,1-Dichloroethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,1-Dichloroethene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L ug/L	1.0 U
1,2,4-Trichlorobenzene	ug/kg		11 U	11 U	11 U	ug/L	2.0 U
1,2-Dibromo-3-chloropropane	ug/kg	11 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,2-Dibromoethane	ug/kg	5.3 U		!	5.4 U		1.0 U
1,2-Dichlorobenzene	ug/kg	5.3 U	5.4 U	5.6 U		ug/L	1
1,2-Dichloroethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,2-Dichloropropane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,3-Dichlorobenzene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
1,4-Dichlorobenzene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
2-Butanone	ug/kg	21 U	21 U	23 U	22 U	ug/L	10 U
2-Hexanone	ug/kg	21 U	21 U	23 U	22 U	ug/L	10 U
4-Methyl-2-pentanone	ug/kg	21 U	21 U	23 U	22 U	ug/L	10 U
Acetone	ug/kg	21 U	21 U	23 U	22 U	ug/L	10 U
Benzene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Bromodichloromethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Bromoform	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Bromomethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Carbon disulfide	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Carbon tetrachloride	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Chlorobenzene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Chloroethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Chloroform	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Chloromethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
cis-1,2-Dichloroethene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
cis-1,3-Dichloropropene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Cyclohexane	ug/kg	11 U	11 U	11 U	11 U	ug/L	1.0 U
Dibromochloromethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Dichlorodifluoromethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Ethylbenzene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Isopropylbenzene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Methyl acetate	ug/kg	11 U	11 U	11 U	11 U	ug/L	10 U
Methyl tert-butyl ether	ug/kg	21 U	21 U	23 U	22 U	ug/L	5.0 U
and a seed of the second of th	ug/kg	11 U	11 U	11 U	11 U	ug/L	1.0 U
Methylogophlorida	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Methylene chloride	ug/kg		5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Styrene	ug/kg	5.3 U 5.3 U	5.4 U	5.6 U	5.4 U	ug/L ug/L	1.0 U
Tetrachloroethene	ug/kg		5.4 U	5.6 U	5.4 U	ug/L ug/L	1.0 U
Toluene	ug/kg	5.3 U	1 g 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	The second of the first of the second of	5.4 U	*	1.0 U
trans-1,2-Dichloroethene	ug/kg	5.3-U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
trans-1,3-Dichloropropene	ug/kg	5.3 U	5.4 U	5.6 U	the second of the second of the second of	ug/L	
Trichloroethene	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Trichlorofluoromethane	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Vinyl chloride	ug/kg	5.3 U	5.4 U	5.6 U	5.4 U	ug/L	1.0 U
Xylenes (total)	ug/kg	11 U	11 U	11 U	11 U	ug/L	2.0 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

Table 2-2 Analytical Data Summary Site 16 Soil Semivolatiles ODOT Innerbelt Study

DADAMETED	UNITS	A6G140116001 16-MW01-1416 07/11/2006	A6G140116002 16-SB02-1820 07/11/2006	A6G140116003 16-SB03-2224 07/11/2006	A6H090258001 16-SB04-0810 08/07/2006
PARAMETER  1 11 Diphonul	ug/kg	350 U	350 U	370 U	360 U
1,1'-Biphenyl 2,2'-oxybis(1-Chloropropane)	ug/kg	350 U	350 U	370 U	360 U
<b>,</b> , , , , , , , , , , , , , , , , , ,	ug/kg ug/kg	350 U	350 U	370 U	360 U
2,4,5-Trichlorophenol	• •			370 U	360 U
2,4,6-Trichlorophenol	ug/kg	350 U	350 U	370 U	360 U
2,4-Dichlorophenol	ug/kg	350 U	350 U	1	l .
2,4-Dimethylphenol	ug/kg	350 U	350 U	370 U	360 U
2,4-Dinitrophenol	ug/kg	350 U	350 U	370 U	360 U
2,4-Dinitrotoluene	ug/kg	350 U	350 U	370 U	360 U
2,6-Dinitrotoluene	ug/kg	350 U	350 U	370 U	360 U
2-Chloronaphthalene	ug/kg	350 U	350 U	370 U	360 U
2-Chlorophenol	ug/kg	350 U	350 U	370 U	360 U
2-Methylnaphthalene	ug/kg	350 U	350 U	370 U	360 U
2-Methylphenol	ug/kg	350 U	350 U	370 U	360 U
2-Nitroaniline	ug/kg	350 U	350 U	370 U	360 U
2-Nitrophenol	ug/kg	350 U	350 U	370 U	360 U
3,3'-Dichlorobenzidine	ug/kg	350 U	350 U	370 U	360 U
3-Nitroaniline	ug/kg	350 U	350 U	370 U	360 U
4,6-Dinitro-2-methylphenol	ug/kg	350 U	350 U	370 U	360 U
4-Bromophenyl phenyl ether	ug/kg	350 U	350 U	370 U	360 U
4-Chloro-3-methylphenol	ug/kg	350 U	350 U	370 U	360 U
4-Chloroaniline	ug/kg	350 U	350 U	370 U	360 U
4-Chlorophenyl phenyl ether	ug/kg	350 U	350 U	370 U	360 U
4-Methylphenol	ug/kg	350 U	350 U	370 U	360 U
4-Nitroaniline	ug/kg	350 U	350 U	370 U	360 U
4-Nitrophenol	ug/kg	350 U	350 U	370 U	360 U
Acenaphthene	ug/kg	350 U	350 U	370 U	360 U
Acenaphthylene	ug/kg	350 U	350 U	370 U	360 U
Acetophenone	ug/kg	70 U	71 U	75 U	72 U
Anthracene	ug/kg	350 U	350 U	370 U	360 U
Atrazine	ug/kg	350 U	350 U	370 U	360 U
Benzaldehyde	ug/kg	350 U	350 U	370 U	360 U
Benzo(a)anthracene	ug/kg	350 U	350 U	370 U	34 J
Benzo(a)pyrene	ug/kg	350 U	350 U	370 U	37 J
Benzo(b)fluoranthene	ug/kg	350 U	350 U	370 U	49 J
Benzo(ghi)perylene	ug/kg	350 U	350 U	370 U	28 J
Benzo(k)fluoranthene	ug/kg	350 U	350 U	370 U	22 J
bis(2-Chloroethoxy)methane	ug/kg ug/kg	350 U	350 U	370 U	360 U
bis(2-Chloroethyl) ether	ug/kg ug/kg	350 U	350 U	370 U	360 U
and the control of th		350 U	350 U	370 U	480
bis(2-Ethylhexyl) phthalate	ug/kg	350 U	350 U	370 U	360 U
Butyl benzyl phthalate	ug/kg	350 U	350 U	370 U	360 U
Carbarala	ug/kg		350 U	370 U	360 U
Carbazole	ug/kg	350 U	A STATE OF THE STA	The state of the s	Service of the servic
Chrysene	ug/kg	350 U	8.7 J	370 U	39 J
Dibenz(a,h)anthracene	ug/kg	350 U	350 U	370 U	360 U
Dibenzofuran	ug/kg	350 U	350 U	370 U	360 U
Diethyl phthalate	ug/kg	350 U	350 U	370 U	360 U
Dimethyl phthalate	ug/kg	350 U	350 U	370 U	360 U
Di-n-butyl phthalate	ug/kg	350 U	350 U	370 U	360 U
Di-n-octyl phthalate	ug/kg	350 U	350 U	370 U	360 U
Fluoranthene	ug/kg	350 U	8.0 J	370 U	68 J
Fluorene	ug/kg	350 U	350 U	370 U	360 U

Table 2-2 Analytical Data Summary Site 16 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140116001 16-MW01-1416 07/11/2006	A6G140116002 16-SB02-1820 07/11/2006	A6G140116003 16-SB03-2224 07/11/2006	A6H090258001 16-SB04-0810 08/07/2006
Hexachlorobenzene	ug/kg	350 U	350 U	370 U	360 U
Hexachlorobutadiene	ug/kg	350 U	350 U	370 U	360 U
Hexachlorocyclopentadiene	ug/kg	350 U	350 U	370 U	360 U
Hexachloroethane	ug/kg	350 U	350 U	370 U	360 U
Indeno(1,2,3-cd)pyrene	ug/kg	350 U	350 U	370 U	24 J
Isophorone	ug/kg	350 U	350 U	370 U	360 U
Naphthalene	ug/kg	350 U	350 U	370 U	360 U
Nitrobenzene	ug/kg	350 U	350 U	370 U	360 U
N-Nitrosodi-n-propylamine	ug/kg	350 U	350 U	370 U	360 U
N-Nitrosodiphenylamine	ug/kg	350 U	350 U	370 U	360 U
Pentachlorophenol	ug/kg	350 U	350 U	370 U	360 U
Phenanthrene	ug/kg	350 U	350 U	370 U	29 J
Phenol	ug/kg	350 U	350 U	370 U	360 U
Pyrene	ug/kg	350 U	350 U	370 U	52 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-3 Analytical Data Summary Site 16 Soil Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6G140116001 16-MW01-1416 07/11/2006	A6G140116002 16-SB02-1820 07/11/2006	A6G140116003 16-SB03-2224 07/11/2006	A6H090258001 16-SB04-0810 08/07/2006
Percent Solids	%	94.7	93.2	88.5	92.7
Arsenic	mg/kg	5.9	7.1	7.5	8.5
Barium	mg/kg	20.9 J	12.0 J	13.2 J	30.2
Cadmium	mg/kg	0.17 J	0.33	0.19 J	0.15 J
Chromium	mg/kg	6.1	6.4	4.9	7.4
Lead	mg/kg	5.4	6.7	6.3	33
Selenium	mg/kg	0.53 U	0.54 U	0.56 U	0.54 U
Silver	mg/kg	0.53 U	0.54 U	0.56 U	0.54 U
Mercury	mg/kg	0.11 U	0.11 U	0.11 U	0.043 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

### 6.6 SITE 17 - TERMINAL OIL

According to the Cleveland City Directories, the site was listed as O'Brien Cartage from 1954 through 1989. The site was listed as Gillota Fuel Products in the 1994 and 1999 City Directories. The site was identified as a LUST site in the EDR Report.

According to the Cleveland Fire Prevention Bureau files, the site was occupied by O'Brien Cartage in 1940. According to a permit dated October 18, 1955, a 2,000-gallon gasoline UST was installed at the site by the Oil Equipment Service Company. The 2,000-gallon UST reportedly was installed to replace a 1,000-gallon UST, which had leaked. A permit dated March 7, 1957 indicates approximately 6,000-gallons of gasoline were stored at the site in USTs. In 1966, the site was occupied by O'Brien and Nye Cartage Company. A permit dated September 27, 1974 indicates a 6,000-gallon fuel oil UST was to be installed at the site.

A letter dated September 28, 1993 from Tank Tech Environmental to the Fire Prevention Bureau indicates BUSTR recognized Norfolk and Western Railway as the UST owner. The BUSTR registration form indicates there are two 6,000-gallon diesel USTs at the site. The operator of the steel USTs was listed as Gillota Fuel Products; however, the owner was listed Norfolk and Western Railway. A letter dated March 24, 1995 from BUSTR to Norfolk Southern indicates a UST Closure Report had been received and the extent of the release had been defined. The letter stipulates Norfolk Southern was required to submit a Remedial Action Plan to describe how the soil and/or groundwater at the site would be cleaned up.

At the time of the URS Phase I reconnaissance, the site was occupied by Terminal Oil. The site appeared to be utilized for the storage of tanker trucks. Several ASTs and multiple 55-gallon storage drums were observed on the site. Surface staining was observed.

### 6.6.1 Field Activities

A total of four monitoring wells were proposed for Site 17. Groundwater was encountered in three of the four soil borings during field activities, monitoring wells were installed and one soil boring (SB01) was advanced at Site 17. One soil sample per boring was collected and analyzed for VOCs, PAHs, and TPH. A duplicate soil sample was collected from 17-MW04-0204. A Sample Location Map is included as **Figure 6-6A**.

Groundwater samples were collected and analyzed for VOCs and PAHs, from MW02, MW03, and MW04.

### 6.6.2 Site-Specific Geology/Hydrogeology

Soils at Site 17 consisted primarily of sand with minor amounts of silty sand and clayey sand. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 17 on August 21, 2006. Localized groundwater flow across Site 17 is likely influenced by both natural features and urban development,

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including paved surfaces, buildings, and underground utilities. The general flow direction at Site 17 is to the west-southwest, towards the Cuyahoga River.

### 6.6.3 Geophysical Survey

The geophysical survey at Site 17 included the asphalt-paved parking and drive surfaces in three general areas surrounding the main building.

- The northern area included the parking spaces directly north of the main building.
- A parking lot east of the building, measuring approximately 50 by 60 feet.
- The parking area and drive south and southwest of the main building

The survey transects were conducted in a north – south direction to approximately align transects along the earth's total magnetic field, which provides the greatest sensitivity to buried ferrous objects such as steel underground storage tanks. Transects were spaced five feet apart. The presence of buildings adjacent to surveyed areas prevented the use of a global positioning system (GPS) to obtain corresponding latitude and longitude datum necessitating the need to paint survey points on the pavement.

The survey north of the main building identified four areas of anomalously high magnetic gradients. These areas are shown on Figure 6-6B as anomalies A and B as described below:

- A. A gradient anomaly was observed along the drive, west of the pump island. This anomaly is likely the result of metallic objects on the surface at this location and not related to a UST.
- B. An anomaly was detected immediately west of the gas station building that was attributed to a parked vehicle that was present at the time of the survey.

The survey east of the building indicated the presence of linear anomalies that appear to correspond to specific transects, possibly due to instrument error.

South of the building, a series of high gradient anomalies were detected along a paved surface that appeared to be designed for heavy traffic. The anomalies observed along the paved surface were assumed to be related to reinforcing steel.

The results of this survey do not support the presence of a steel UST at Site 17. No further investigation is recommended to attempt to locate the UST.

### 6.6.4 Soil Analytical Results

A total of thirteen VOCs were detected in the five soil samples submitted from Site 17. The chemicals detected in the samples submitted were 1,2-dichlorobenzene (0.60 ug/kg), 1,3-dichlorobenzene (0.50 ug/kg), 1,4-dichlorobenzene (0.66 ug/kg), 2-butanone (ranging from 5.0 ug/kg to 50 ug/kg), benzene (ranging from 1.2 ug/kg to 290 ug/kg), carbon disulfide (ranging

**Phase II Findings** 

from 1.3 ug/kg to 3.2 ug/kg), cyclohexane (ranging from 0.73 ug/kg to 180 ug/kg), ethylbenzene (ranging from 1.2 ug/kg and 2,300 ug/kg), isopropylbenzene (ranging from 0.26 ug/kg to 380 ug/kg), methylcyclohexane (ranging from 0.88 ug/kg to 350 ug/kg), methylene chloride (7.1 ug/kg), toluene (ranging from 0.74 ug/kg to 3,000 ug/kg), and total xylenes (ranging from 1.5 ug/kg to 16,000 ug/kg). All other VOCs were below the detection limits.

A total of seventeen PAHs were detected in the five samples submitted from Site 17. Concentrations of 2-methylnaphthanene (ranging from 120 ug/kg to 1,700 ug/kg), acenaphthene (ranging from 100 ug/kg to 550 ug/kg), acenaphthylene (ranging from 59 ug/kg to 160 ug/kg), anthracene (ranging from 240 ug/kg to 2,400 ug/kg), benzo(a)anthracene (ranging from 41 ug/kg to 4,100 ug/kg), benzo(a)pyrene (ranging from 37 ug/kg to 3,500 ug/kg), benzo(b)fluoranthene (ranging from 52 ug/kg to 4,500 ug/kg), benzo(g,h,i)perylene (ranging from 25 ug/kg to 2,000 ug/kg), benzo(k)fluoranthene (ranging from 250 ug/kg to 1,500 ug/kg), chrysene (ranging from 41 ug/kg to 3,900 ug/kg), dibenz(a,h)anthracene (ranging from 66 ug/kg to 540 ug/kg), fluoranthene (ranging from 85 ug/kg to 11,000 ug/kg), fluorene (ranging from 140 ug/kg to 820 ug/kg), indeno(1,2,3)pyrene (ranging from 220 ug/kg to 1,900 ug/kg), naphthalene (ranging from 84 ug/kg to 2,600 ug/kg), phenanthrene (ranging from 42 ug/kg to 8,100 ug/kg), and pyrene (ranging from 68 ug/kg to 8,100 ug/kg) were detected in the soil samples submitted.

Gasoline and diesel range TPH were detected in the soil samples from Site 17. The highest concentrations were found in the gasoline range, which is consistent with the concentrations of VOCs detected across Site 17. Concentrations of the light petroleum fraction ranged from 330 ug/kg to 250,000 ug/kg. Concentrations of the middle petroleum fraction ranged from 17 mg/kg to 41 mg/kg. Concentrations of the heavy petroleum fraction ranged from 15 mg/kg to 260 mg/kg.

The analytical results are presented in Table 6-6A.

## 6.6.5 Groundwater Analytical Results

Nine VOCs were detected in the three groundwater samples collected from Site 17. Concentrations of 2-butanone (0.59 ug/L), acetone (1.0 ug/L), benzene (ranging from 14 ug/L to 3,700 ug/L), cyclohexane (72 ug/L), ethylbenzene (1,000 ug/L), isopropylbenzene (45 ug/L), methyl tert-butyl ether (ranging from 3.7 ug/L to 220 ug/L), toluene (ranging from 1.1 ug/L to 2,800 ug/L), and total xylenes (ranging from 0.95 ug/L to 6,500 ug/L) were detected in the groundwater samples submitted. All other VOCs were below the detection limits.

Thirteen PAHs were detected in the groundwater samples submitted from Site 17. Concentrations of 2-methylnaphthanene (0.22 ug/L and 25 ug/L), anthracene (0.35 ug/L), benzo(a)anthracene (0.42 ug/L and 0.77 ug/L), benzo(a)pyrene (0.58 ug/L), benzo(b)fluoranthene (0.70 ug/L), benzo(g,h,i)perylene (0.35 ug/L), benzo(k)fluoranthene (0.31 ug/L), chrysene (0.60 ug/L), fluoranthene (0.62 ug/L and 1.6 ug/L), indeno(1,2,3)pyrene (0.3 ug/L), naphthalene (0.22 ug/L and 43 ug/L), phenanthrene (0.72 ug/L and 1.2 ug/L), and pyrene (0.49 ug/L and 1.4 ug/L) were detected in the groundwater samples submitted from 17-MW02 and 17-MW03. All other PAHs were below the detection limits.

The analytical results are presented in Table 6-6B.

### 6.6.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on **Tables 6-6A** and **6-6B**.

The concentration of benzene (290 ug/kg) and total xylenes (16,000 ug/kg) detected in sample 17-SB01-2628 exceeded the BUSTR closure action levels. No other VOCs detected exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

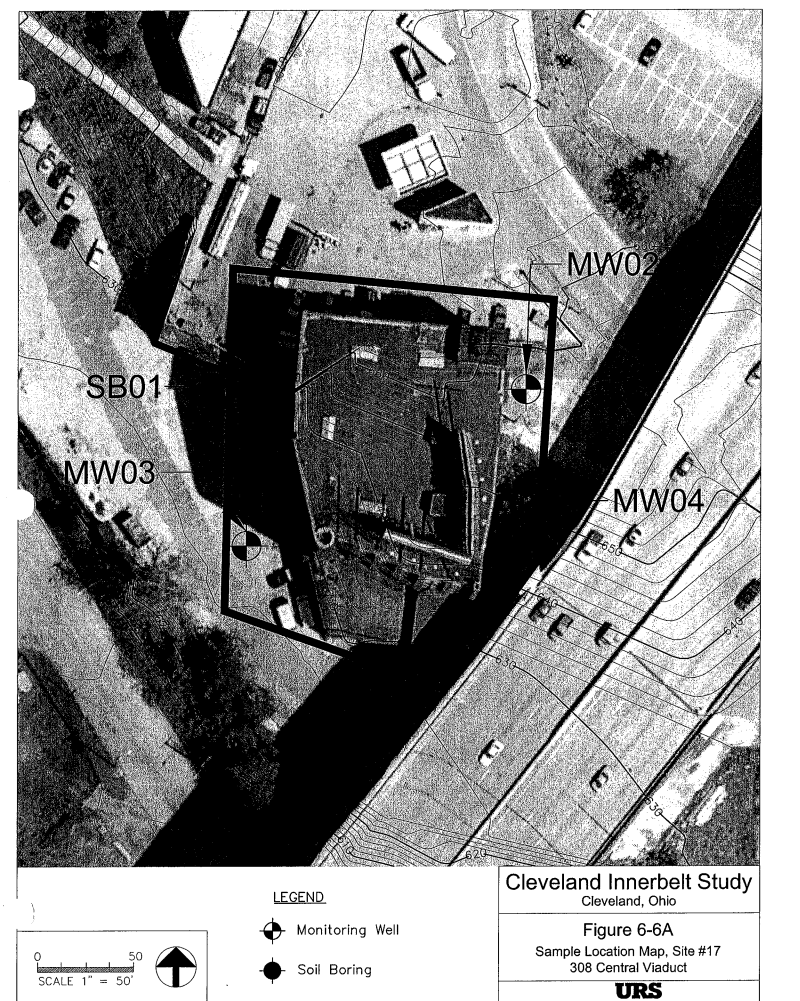
The concentrations of benzo(a)pyrene detected in samples 17-MW03-0406, 17-MW04-0204, and 17-MW04-0204D exceeded the BUSTR closure action levels. No other PAHs detected exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

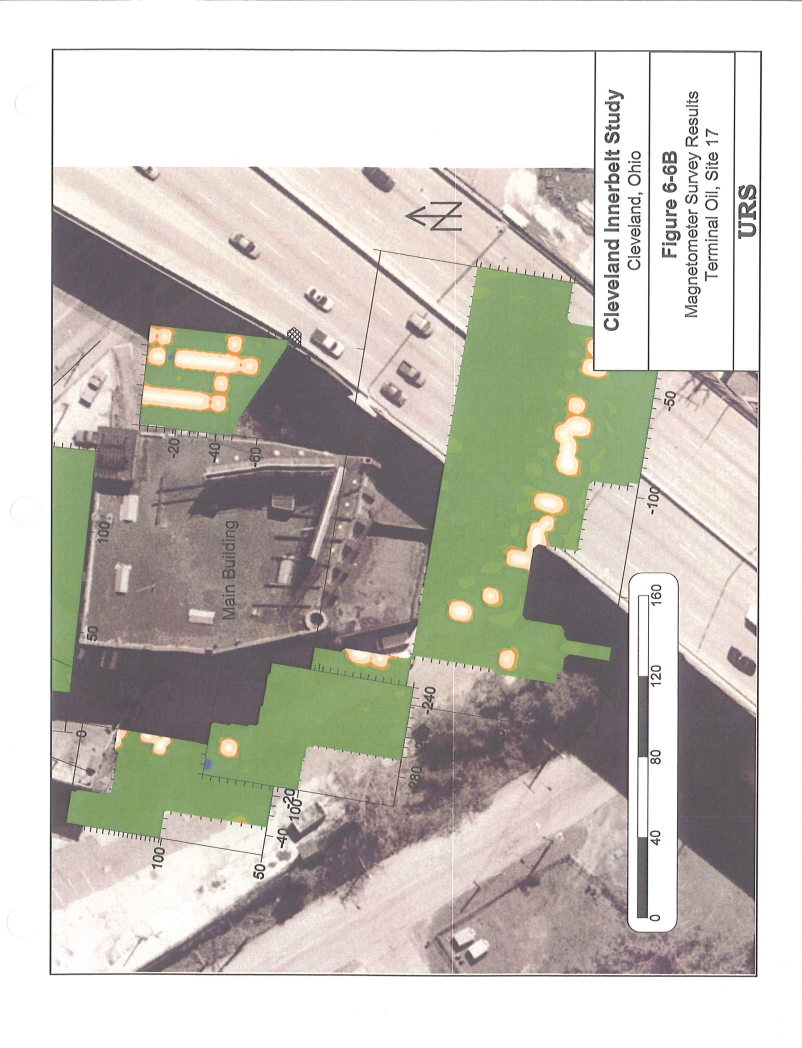
The concentrations of benzene, ethylbenzene, methyl tert-butyl ether, toluene, benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, and indeno(1,2,3-cd)pyrene detected in the groundwater samples 17-MW02, 17-MW03, and 17-MW04 exceeded the OEPA VAP standard for generic unrestricted potable use standards and BUSTR closure action levels.

#### 6.6.7 Conclusions

Based on the analytical results, the soil and groundwater at Site 17 may require special disposal and/or worker protection protocols (plan note) during construction activities.

**FIGURES** 





### TABLES

Table—JA
Summary of Detected Chemicals in Soil
Site 17 - Terminal Oil
ODOT Innerbelt Study
Cleveland, Ohio

roberszene ugkg 370,000 370,000 6.3 UJ 5.6 U 0.68 J 0.00 orderszene ugkg 370,000 5.20,000 6.3 UJ 5.6 U 0.55 U 0.68 J 0.00 orderszene ugkg 77,160,000 5.00 0.00 1.49 1.13 J	Ž	PARAMETER	UNITS	VAP Commercial/ #Industrial === Standard <sup>(1)</sup>	VAP Construction  E.Worker  F. Standard <sup>2</sup>	BUSTIF Closure Action Level:	.17-MW02-2022 08/15/2006	17-MW03-0406 08/15/2006	17-MW04-0204 08/15/2006	17-MW04-0204-D 08/15/2006	17-SB01-2628 08/14/2006
1,2-Chelotocheragene   ug/kg   77,0000   240,000     65 UU   56 U   0.65 J   0.65 J     1,4-Chelotocheragene   ug/kg   77,0000   51,00000     65 U   13 J   13 J     1,1-Chelotocheragene   ug/kg   77,0000   720,000     140   173 J   13 J     1,1-Chelotocheragene   ug/kg   77,0000   720,000     140   173 J   13 J     1,1-Chelotocheragene   ug/kg   77,0000   720,000     140   0.73 J   13 J     1,1-Chelotocheragene   ug/kg   77,0000   720,000     140   0.73 J   13 J     1,1-Chelotocheragene   ug/kg   77,0000   22,0000   45,500   65 U   65 U     1,1-Chelotocheragene   ug/kg   73,0000   22,0000   45,500   63 U   65 U     1,1-Chelotocheragene   ug/kg   73,0000   22,0000   15,700   15,700   15,700     1,1-Chelotocheragene   ug/kg   75,0000   22,0000   15,700   15,700   15,700   15,700     1,1-Chelotocheragene   ug/kg   75,0000   10,000,000   15,7		1,2-Dichlorobenzene	ug/kg	370,000	370,000	1	6.3 W	5.6 U	0.60 J	6.3 W	0.049
1.   1.   1.   1.   1.   1.   1.   1.		1,3-Dichlorobenzene	ug/kg	240,000	240,000		6.3 UJ	5.6 U	0.50 J	6.3 UJ	670 U
2-Butanone         ug/kg         77,600,000         80,000,000         —         50         13 J         50 J           Denration         ug/kg         77,600,000         730,000         —         23 J         1.5 J         1.3 J           Contron disultide         ug/kg         720,000         720,000         —         23 J         1.3 J         1.3 J           Cyclobezana         ug/kg         220,000         220,000         45,500         6.5 J         6.5 U         6.5 U           Entyleanzane         ug/kg         220,000         220,000         45,500         6.5 U         6.5 U         6.5 U           Methylordolexane         ug/kg         520,000         2.300,000         45,100         6.5 U         6.5 U         6.5 U           Methyloredolexane         ug/kg         520,000         2.300,000         1.500,000         2.3 J         2.5 J         2.5 J           Methyloredolexane         ug/kg         520,000         1.500,000         1.500,000         1.500         3.2 J         2.5 J         3.5 J           Acchieve (dull)         100         1.500         1.500         1.500         1.500         1.500         1.500         1.500         1.500         1.500         1.500 </td <th></th> <td>1,4-Dichlorobenzene</td> <td>ug/kg</td> <td>470,000</td> <td>5,300,000</td> <td></td> <td>6.3 W</td> <td>5.6 U</td> <td>0.66 J</td> <td>6.3 UJ</td> <td>0.029</td>		1,4-Dichlorobenzene	ug/kg	470,000	5,300,000		6.3 W	5.6 U	0.66 J	6.3 UJ	0.029
Bencree         Ug/kg         100,000         310,000         149         12 J         16 J         13 J         13 J           Carbon idestifice         Ug/kg         720,000         750,000         -         -         13 J         13 J         13 J           Carbon idestifice         Ug/kg         230,000         820,000         6.3 U         6.5 U <th></th> <td>2-Butanone</td> <td>ug/kg</td> <td>71,600,000</td> <td>80,000,000</td> <td>1</td> <td>20</td> <td>13 J</td> <td>5.0 J</td> <td>6.1 J</td> <td>2700 U</td>		2-Butanone	ug/kg	71,600,000	80,000,000	1	20	13 J	5.0 J	6.1 J	2700 U
Carbon disultide         ug/kg         720,000         720,000         - 140         52.3         13.2         113.3           Entylebrezine         ug/kg         200,000         200,000         45,500         6.5 U         6.5 U         6.5 U           Entylebrezine         ug/kg         200,000         200,000         45,500         6.5 U         6.5 U         6.5 U           Rehtlyden carbonide         ug/kg         1,300,000         2,200,000         - 500,000         6.5 U         1.4 U         0.88 J           Adentyles choices         ug/kg         1,200,000         2,200,000         - 500,000         6.5 U         6.5 U         6.5 U           Adentyles choices         ug/kg         1,200,000         1,500,000         1,500         1.4 J         0.88 J           Adentyles choices         1,500		Benzene	ug/kg	100,000	310,000	149	1.2 J	1.6 J	1.3 J	4.1 J	290 J
Cyclobrexine         Ug/kg         2-0.00         4-0.00         6.3 U         6.5 U         6.5 U           Eth/berzene         Ug/kg         850,000         850,000         45,500         6.3 U         6.5 U         6.5 U           Repropylebrazine         Ug/kg         1300,000         2,230,000          5.9 U         6.5 U         6.5 U           Methyleracine         Ug/kg         1,300,000         2,230,000          5.9 U         6.5 U         6.5 U           Acenaciphthere         Ug/kg         1,300,000         2,230,000         1.5 U         2.9 J         2.5 J         2.5 J           Acenaciphthere         Ug/kg         180,000         1,000          1.00 J         4.0 J         1.00 J         2.5 J	S	Carbon disulfide	ug/kg	720,000	720,000	1	2.3 J	3.2 J	1.3 J	1.8 J	0.079
Ethylheerizene         Ug/kg         220,000         45,500         6.3 U         5.6 U         6.5 U           Reptroplearizene         Ug/kg         1,300,000         2,300,000         -         5.8 U         6.5 U         6.5 U           Methylpicalchezane         Ug/kg         1,300,000         2,300,000         -         -         5.8 U         6.5 U         6.5 U           Methylpica chloride         Ug/kg         1,300,000         2,300,000         -         6.3 U         2.5 J         2.5 J           Acenaphthaliere chloride         Ug/kg         150,000         1,000         1,100         1,100 J         2.9 J         2.5 J         2.5 J           Acenaphthaliere chloride         Ug/kg         150,000         1,000         -         1,00 J         2.9 J         2.5 J         2.5 J           Acenaphthaliere         Ug/kg         150,000         1,100         -         1,100 J         450 J         350 J         1,100 J           Acenaphthylere         Ug/kg         1,200,000         1,100         -         1,100 J         450 J         1,100 J           Acenaphthylere         1,000         1,100         1,100 J         1,100 J         1,100 J         1,100 J         1,100 J         1,1	00	Cyclohexane	ug/kg	-	-	1	140	0.73 ປ	13 U	0.93 კ	180 J
Septicipy/lentratione         Ug/kg         860,000         860,000         —         6.5 U         6.5 U         6.5 U           Mehrly/cochokazine         Ug/kg         1.300,000         2.300,000         2.300,000         —         6.3 U         6.5 U         6.5 U           Toluene         Ug/kg         1.300,000         5.20,000         15,700         1.5 J         2.5 J         2.5 J           Xylenes (total)         Ug/kg         160,000         15,700         1.5 J         2.5 J         2.5 J           Xylenes (total)         Ug/kg         180,000,000         550,000         —         1.00 J         450 J         2.5 J           Acenaphthylene         Ug/kg         180,000,000         1,000,000         —         240 J         150 J         2.5 J           Acenaphthylene         Ug/kg         180,000,000         1,100         410 J         150 J         150 J         150 J           Acenaphthylene         Ug/kg         180,000,000         1,100         1,100         410 J         150 J <td< td=""><th>٨</th><td>Ethylbenzene</td><td>ug/kg</td><td>230,000</td><td>230,000</td><td>45,500</td><td>6.3 U</td><td>5.6 U</td><td>6.5 U</td><td>1.2 J</td><td>2300</td></td<>	٨	Ethylbenzene	ug/kg	230,000	230,000	45,500	6.3 U	5.6 U	6.5 U	1.2 J	2300
Methylicyclohexane         ug/kg            59         14 J         0.88 J           Methylicyclohexane         ug/kg         1,300,000         2,300,000         2,300,000         2,300,000         2,300,000         2,300,000         2,300,000         2,300,000         2,300,000         2,26 J         2.6 J         2.6 J         2.6 J         2.5 J         2.6 J <t< td=""><th></th><td>Isopropylbenzene</td><td>ug/kg</td><td>860,000</td><td>860,000</td><td></td><td>0.26 J</td><td>0.51 J</td><td>6.5 U</td><td>0.37 J</td><td>380 J</td></t<>		Isopropylbenzene	ug/kg	860,000	860,000		0.26 J	0.51 J	6.5 U	0.37 J	380 J
Methylene artioride         ug/kg         1,390,000         2,300,000         49,100         0,74 J         2.9 J         2.5 J         2.5 J           Toluene         ug/kg         550,000         520,000         1,5,70         1,5 J         2.9 J         2.5 J         2.5 J           Zykanes (data)         ug/kg         160,000         160,000         160,000          120 J         2.9 J         2.5 J           Acenaphthylene         ug/kg         180,000,000         1,000,000          190 J         450 J         350 J           Acenaphthylene         ug/kg         880,000,000         1,000,000         1,000 J         240 J         1900 J         160 J           Acenaphthylene         ug/kg         83,000         1,100 J         490 J         1900 J         1100 J           Berczo(a)lancanthene         ug/kg         6,300         1,100 J         490 J         1900 J         4500 J           Berczo(phlucanthene         ug/kg         6,300         41,000 O         1,100 O         490 J         1900 J         240 J         1500 J           Berczo(phlucanthene         ug/kg         6,300         41,000 O         1,100 OO         220 J         1900 J         240 J         1500 J		Methylcyclohexane	ug/kg	The state of the s	1	-	59	1.4 J	0.88 J	1.6 J	350 J
Vylennes (total)         ug/kg         520,000         520,000         15 J         2.5 J         2.5 J           Xylennes (total)         ug/kg         160,000         15,700         1.5 J         2.9 J         2.5 J           Acentaphtrajente         ug/kg         160,000         550,000          120 J         270 J         380 J           Acentaphtryjene         ug/kg         180,000,000         1,000,000,000          240 J         460 J         160 J           Anthracene         ug/kg         63,000         11,000         490 J         1900 J         1100 J           Benzzólphyrene         ug/kg         63,000         11,000         490 J         1900 J         4500 J           Benzzólphiloranthene         ug/kg         63,000         11,000         490 J         1900 J         4500 J           Benzzólphiloranthene         ug/kg         63,000         41,000,000         11,000         250 J         80 J         5000 J           Benzzólphiloranthene         ug/kg         67,000         41,000,000         1,100,000         250 J         80 J         500 J           Benzzólphiloranthene         ug/kg         67,000         41,000,000         1,100,000         250 J <t< td=""><th></th><td>Methylene chloride</td><td>ug/kg</td><td>1,300,000</td><td>2,300,000</td><td></td><td>6.3 U</td><td>5.6 U</td><td>6.5 U</td><td>7.7</td><td>O 029</td></t<>		Methylene chloride	ug/kg	1,300,000	2,300,000		6.3 U	5.6 U	6.5 U	7.7	O 029
Xylenes (total)         ug/kg         160,000         15,700         15 J         32 J         2.6 J           Z-Methylmaphthalene         ug/kg         160,000         530,000,000		Toluene	ug/kg	520,000	520,000	49,100	0.74 ט	2.9 J	2.5 J	6.6	3000
2-Methylinaphthalene         ug/kg	_	Xylenes (total)	ug/kg	160,000	160,000	15,700	1.5 J	3.2 J	2.6 J	9.6 J	16000
Acenaphthene         ug/kg         180,000,000         530,000,000         -         100 J         450 J         550 J           Acenaphthene         ug/kg         880,000,000         1,000,000,000         -         240 J         160 J         160 J           Anthracene         ug/kg         63,000         11,000         490 J         1900         4500 J           Benzo(a)pyrene         ug/kg         63,000         810,000         11,000         480 J         1900         4500 J           Benzo(a)pyrene         ug/kg         63,000         810,000         11,000         480 J         1900         4500 J           Benzo(b)filocranthene         ug/kg         630,000         11,000         250 J         840 J         1500 J           Chrysene         ug/kg         6,700,000         41,00,000         1,100         66 J         280 J         1500 J           Chrysene         ug/kg         6,700         41,000,000         1,100         66 J         280 J         1900 J           Fluorene         ug/kg         6,700         41,000,000         1,100         66 J         280 J         1900 J           Fluorene         ug/kg         67,000         1,000,000         1,100         66 J <th></th> <td>2-Methylnaphthalene</td> <td>ug/kg</td> <td>1</td> <td></td> <td></td> <td>120 J</td> <td>270 J</td> <td>360 J</td> <td>320 J</td> <td>1700</td>		2-Methylnaphthalene	ug/kg	1			120 J	270 J	360 J	320 J	1700
Acenaphthylene         ug/kg         —         —         —         970 J         160 J           Anthracene         ug/kg         63,000         1,000,000,000         —         240 J         970 J         2400 J           Benzo(a)pytrane         ug/kg         63,000         81,000         1,100         490 J         1900         4100 J           Benzo(a)pytrane         ug/kg         63,000         810,000         11,000         490 J         1900         4500 J           Benzo(b)filoranthene         ug/kg         63,000         810,000         110,000         250 J         840 J         1500 J           Chrysene         ug/kg         6,700,000         41,000,000         1,100,000         460 J         1700 J         3900 J           Pluoranthene         ug/kg         6,700,000         170,000,000         —         220 J         840 J         540 J           Pluoranthene         ug/kg         6,700,000         170,000,000         —         140 J         480 J         540 J           Pluoranthene         ug/kg         6,700,000         170,000,000         —         140 J         480 J         540 J           Phenanthrene         ug/kg         530,000         1,900,000         —<		Acenaphthene	ug/kg	180,000,000	530,000,000		100 ე	450 ש	550 J	260 J	1100 U
Anthracene         ug/kg         880,000,000         1,000,000,000          240 J         970 J         2400 J           Benzo(a)anthracene         ug/kg         63,000         81,000         1,100         490 J         1900         4100 J           Benzo(a)pyrene         ug/kg         63,000         81,000         1,100         480 J         1900         4500           Benzo(phlloranthene         ug/kg         63,000         81,00,000         11,000         250 J         840 J         2000 J           Chrysene         ug/kg         6,700,000         41,000,000         1,100,000         250 J         840 J         540 J           Pluoranthene         ug/kg         6,700         41,000,000          1200         1           Pluoranthene         ug/kg         6,700         41,000,000          140 J         850 J         540 J           Fluoranthene         ug/kg         120,000,000         11,000         220 J         840 J         1900 J           Phenanthrene         ug/kg         55,000,000         19,000,000         220 J         840 J         1900 J           Pyrane         ug/kg         55,000,000         19,000,000         19,000,000         19,000,000		Acenaphthylene	ug/kg	-		1	830 U	1900 U	160 J	59 J	1100 U
Benzo(a)anthracene         ug/kg         63,000         810,000         11,000         490 J         1900         4100 J         5300 J         5300 J         5300 J         410 J         1600 J         3500 J         5300 J         5300 J         5300 J         5300 J         5300 J         5300 J         5400 J         5500 J <th< td=""><th></th><td>Anthracene</td><td>ug/kg</td><td>880,000,000</td><td>1,000,000,000</td><td>**************************************</td><td>240 J</td><td>6 OZ6</td><td>2400 J</td><td>f 069</td><td>1100 U</td></th<>		Anthracene	ug/kg	880,000,000	1,000,000,000	**************************************	240 J	6 OZ6	2400 J	f 069	1100 U
Benzo(a)pyrene         ug/kg         6,300         81,000         1,100         480 J         1600 J         3500 J           Benzo(h)fluoranthene         ug/kg         63,000         810,000         11,000         480 J         1900         4500           Benzo(h)fluoranthene         ug/kg		Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	490 J	1900	4100 J	2300	41 J
Benzo(b)filtoranthene         ug/kg         63,000         810,000         11,000         480 J         1900         4500           Benzo(ghi)perylene         ug/kg		Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	410 J	1600 J	3500 J	2000 J	37 J
Benzo(ghi)perylene         ug/kg         -         -         -         240 J         910 J         2000 J           Benzo(k)fluoranthene         ug/kg         630,000         4,100,000         1,100,000         250 J         840 J         1500 J           Chrysene         ug/kg         6,700,000         41,000,000         -         100         540 J         3900 J           Dibenz(a, h)anthracene         ug/kg         6,700         41,000         -         1200         4500         11000           Fluoranthene         ug/kg         120,000,000         340,000,000         -         140 J         480 J         820 J           Indeno(1,2,3-cd)pyrene         ug/kg         67,000         410,000         -         140 J         480 J         820 J           Naphthralene         ug/kg         530,000         1,900,000         -         -         770 J         250 J           Phenanthrene         ug/kg         25,000,000         130,000,000         -         -         770 J         3500           Gasoline Range Organics (Ge-C12)         ug/kg         -         -         -         770 J         36 J         38 J           C20-C23         mg/kg         -         -         - <th></th> <td>Benzo(b)fluoranthene</td> <td>ug/kg</td> <td>63,000</td> <td>810,000</td> <td>11,000</td> <td>480 J</td> <td>1900</td> <td>4500</td> <td>2500</td> <td>52 J</td>		Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	480 J	1900	4500	2500	52 J
Benzo(k)fluoranthene         ug/kg         630,000         8,100,000         110,000         250 J         840 J         1500 J           Chrysene         ug/kg         6,700,000         41,000,000         1,100,000         66 J         280 J         540 J           Dibenz(a, h)authracene         ug/kg         6,700         170,000,000         -         1200         4500         11000           Fluoranthene         ug/kg         120,000,000         340,000,000         -         140 J         480 J         820 J           Indenot(1,2,3-cd)pyrene         ug/kg         67,000         11,000,000         39,800         84 J         270 J         260 J           Naphthalene         ug/kg         530,000         1,900,000         -         710 J         3500         8100           Phenanthrene         ug/kg         25,000,000         130,000,000         -         -         710 J         3500         8100           Pyrene         ug/kg         25,000,000         130,000,000         -         -         1,000,000         1600         770         130 U           Gasoline Range Organics (C6-C12)         ug/kg         -         -         -         -         170 J         36 J         36 J	s	Benzo(ghi)perylene	ug/kg	-	All the state of t	The state of the s	240 J	910 J	2000 J	1300 J	25 J
Chrysene         ug/kg         6,700,000         41,000,000         1,100,000         460 J         17700 J         3900 J           Dibenz(a,h)authracene         ug/kg         6,700         41,000         1,100         66 J         280 J         540 J           Fluorenthene         ug/kg         120,000,000         170,000,000         -         1200         4500         11000           Pluorene         ug/kg         120,000,000         340,000,000         -         140 J         850 J         1900 J           Indeno(1,2,3-cd)pyrene         ug/kg         530,000         1,900,000         39,800         84 J         270 J         260 J           Phenanthrene         ug/kg          -         710 J         3500         8100           Pyrene         ug/kg         25,000,000         130,000,000         -         870         3400         8100           Gasoline Range Organics (C6-C12)         ug/kg         -         -         -         1,000,000         -         -         1,000,000         3400         8100         -           C20-C24         mg/kg         -         -         -         -         -         -         -         -         -         - <t< td=""><th>H∀</th><td>Benzo(k)fluoranthene</td><td>ug/kg</td><td>630,000</td><td>8,100,000</td><td>110,000</td><td>250 J</td><td>840 J</td><td>1500 J</td><td>1000 J</td><td>1100 U</td></t<>	H∀	Benzo(k)fluoranthene	ug/kg	630,000	8,100,000	110,000	250 J	840 J	1500 J	1000 J	1100 U
Dibenz(a,h)anthracene         ug/kg         6,700         41,000         1,100         66 J         280 J         540 J           Fluorenthene         ug/kg         33,000,000         170,000,000         -         1200         4500         11000           Fluorene         ug/kg         120,000,000         340,000,000         -         140 J         480 J         820 J           Indeno(1,2,3-cd)pyrene         ug/kg         67,000         410,000         11,000         220 J         850 J         1900 J           Naphthalene         ug/kg          -         710 J         3500         8100           Pyrene         ug/kg          -         870         3400         8100           Gasoline Range Organics (C6-C12)         ug/kg          -         1,000,000         1600         770         130 U           C10-C20         mg/kg          -         2,000         17 J         36 J         38 J           C20-C34         mg/kg          -         -         2,000         17 J         120 J         260 J		Chysene	ug/kg	6,700,000	41,000,000	1,100,000	460 J	1700 J	3900 J	2100	. <del>1</del>
Fluoranthene         ug/kg         33,000,000         170,000,000         -         1200         4500         11000           Fluorene         ug/kg         120,000,000         340,000,000         -         140 J         480 J         820 J           Indeno(1,2,3-cd)pyrene         ug/kg         67,000         410,000         11,000         220 J         850 J         1900 J           Naphthalene         ug/kg         530,000         1,900,000         39,800         84 J         270 J         260 J           Pyrene         ug/kg          710 J         3500         8100           Gasoline Range Organics (C6-C12)         ug/kg          1,000,000         1600         770         130 U           C10-C20         mg/kg          2,000         17 J         36 J         38 J           C20-C34         mg/kg          5,000         10 J         120 J         260 J		Dibenz(a,h)anthracene	ug/kg	6,700	41,000	1,100	P 99	280 J	540 J	360 J	1100 U
Fluorene         ug/kg         120,000,000         340,000,000         —         140 J         480 J         820 J           Indenof1,2,3-cd)pyrene         ug/kg         67,000         410,000         11,000         220 J         850 J         1900 J           Naphthalene         ug/kg         530,000         1,900,000         39,800         84 J         270 J         260 J           Pyrene         ug/kg         25,000,000         130,000,000         -         870         3400         8100           Gasoline Range Organics (C6-C12)         ug/kg         -         1,000,000         1600         770         130 U           C10-C20         mg/kg         -         2,000         17 J         36 J         38 J           C20-C34         mg/kg         -         -         5,000         80 J         120 J         260 J		Fluoranthene	ug/kg	33,000,000	170,000,000	and the second s	1200	4500	11000	4800	85 J
Indenot (1,2,3-cd)pyrene         ug/kg         67,000         410,000         11,000         220 J         850 J         1900 J           Naphthalene         ug/kg         530,000         1,900,000         39,800         84 J         270 J         260 J           Phenanthrene         ug/kg         25,000,000         130,000,000         -         870         3400         8100           Gasoline Range Organics (C6-C12)         ug/kg         -         1,000,000         1600         770         130 U           C10-C20         mg/kg         -         2,000         17 J         36 J         38 J           C20-C34         mg/kg         -         -         5,000         80 J         120 J         260 J		Fluorene	ug/kg	120,000,000	340,000,000	accuses our establishment in traductive description and other	ل 140	480 J	820 J	270 J	1100 U
Naphthalene         ug/kg         530,000         1,900,000         39,800         84 J         270 J         260 J           Phenanthrene         ug/kg           870         8100         8100           Pyrene          870         3400         8100         8100           Gasoline Range Organics (C6-C12)         ug/kg          1,000,000         1600         770         130 U           C10-C20         mg/kg          -         2,000         17 J         36 J         38 J           C20-C34         mg/kg          -         5,000         80 J         120 J         260 J		Indeno(1,2,3-cd)pyrene	ug/kg	67,000	410,000	11,000	220 J	850 J	1900 J	1200 J	1100 U
Phenanthrene         Ug/kg         25,000,000         130,000,000         -         870         3400         8100           Pyrene         Gasoline Range Organics (Ce-C12)         ug/kg         -         1,000,000         1600         770         130 U           C10-C20         mg/kg         -         2,000         17 J         36 J         38 J           C20-C34         mg/kg         -         5,000         80 J         120 J         260 J		Naphthalene	ug/kg	530,000	1,900,000	39,800	84 J	270 J	260 J	280 J	2600
Pyrene         ug/kg         25,000,000         130,000,000         -         870         3400         8100           Gasoline Range Organics (C6-C12)         ug/kg         -         1,000,000         1600         770         130 U           C10-C20         mg/kg         -         -         2,000         17 J         36 J         38 J           C20-C34         mg/kg         -         5,000         80 J         120 J         260 J		Phenanthrene	ug/kg	The state of the s		1	710 J	3500	8100	2600	45 J
Gasoline Range Organics (C6-C12)         ug/kg          1,000,000         1600         770           C10-C20         mg/kg          2,000         17 J         36 J           C20-C34         mg/kg          5,000         80 J         120 J		Pyrene	ug/kg	25,000,000	130,000,000	1	870	3400	8100	3800	F 89
C10-C20 17 J 36 J 36 J C20-C34 80 J 120 J	ŀ	Gasoline Range Organics (C6-C12)	ug/kg	1		1,000,000	1600	077	130 U	330	250000
C20-C34 80 J 120 J	ЧďĮ	C10-C20	mg/kg	1	1	2,000	ر 17	36 J	38 T	36 J	41 J
		C20-C34	mg/kg	1	1	5,000	80 J	120 א	260 J	15 J	130 J

<sup>-- =</sup> Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial/ Industrial Land Use

<sup>(2)</sup> VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Tabbew6B
Summary of Detected Chemicals in Water
Site 17 - Terminal Oil
ODOT Innerbelt Study
Cleveland, Ohio

۵	PARAMETER	UNITS	VAPIUPUS/	BUSTRICIOSURA Action Level	17-WW-02 08/22/2006	17-MW-03 08/21/2006	17-MW-04 08/21/2006
	2-Butanone	J/6n	0089	-	1200 U	100 U	0.59 J
	Acetone	ug/L	1600	•	1200 U	100 U	1.0 J
	Benzene	ug/L	ß	5	3700	10 C	14
S(	Cyclohexane	ug/L		•	72 J	: 0 2	1.0 U
00/	Ethylbenzene	ug/L	700	700	1100	10 U	) 0 (1
١	Isopropylbenzene	ug/L	1300	ACTIVITY OF THE POST OF THE PO	45 J	10 U	1.0 U
	Methyl tert-butyl ether	ug/L	9	40	620 U	220	3.7 J
	Toluene	ug/L	1000	1000	2800	10 U	; <b>;</b>
	Xylenes (total)	ug/L	10000	10000	6500	20 U	0.95 J
	2-Methylnaphthalene	ng/L	:	1	25	0.22	0.20 U
	Anthracene	ng/L	2600	American de la companya de la compan	0.40 U	0.35	0.20 U
	Benzo(a)anthracene	ng/L	1	0.264	0.42	0.77	0.20
	Benzo(a)pyrene	ng/L	0.2	0.2	0.40 U	0.58	0.20 U
	Benzo(b)fluoranthene	ng/L		0.179	0.40 U	0.7	0.20 U
S	Benzo(ghi)perylene	ng/L	•		0.40 U	0.35	0.20 U
IАЧ	Benzo(k)fluoranthene	ng/L	***	1.79	0.40 U	0.31	0.20
	Chrysene	ng/L	47	47	0.40 U	9.0	0.20 U
	Fluoranthene	ng/L	370	•	0.62	1.6	0.20 U
	Indeno(1,2,3-cd)pyrene	ng/L	The state of the s	0.23	0.40 U	0.3	0.20 U
	Naphthalene	ug/L	140	140	43	0.22	0.20 U
	Phenanthrene	ug/L	-	•	0.72	1.2	0.20 U
	Pyrene	ng/L	280	-	0.49	1.4	11 08 0

<sup>-- =</sup> Standard not available

U=The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

<sup>(1)</sup> VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

## BORING LOGS

Project Location: Site 17
Project Number: 15016633

## Log of Boring 17-MW02

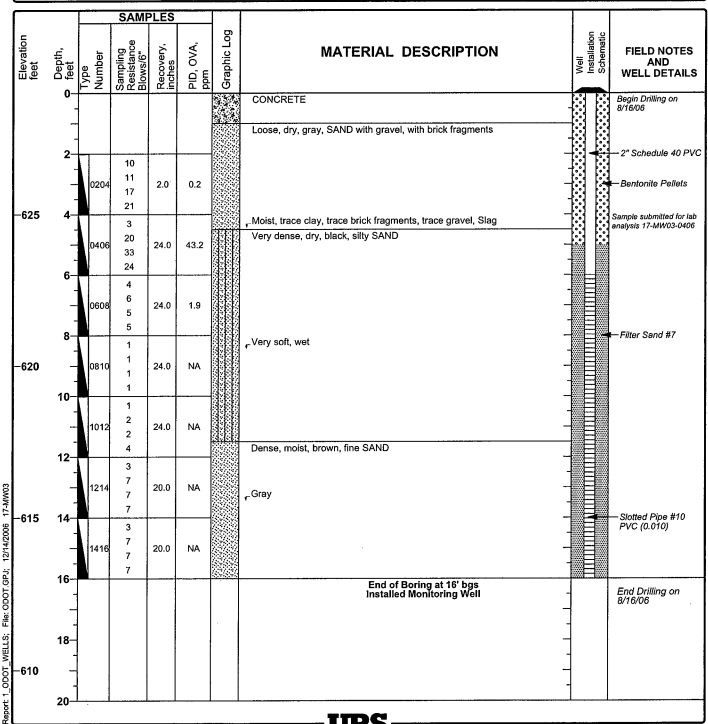
Date(s) 8/15/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 30.0′ bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 658'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 630.15 on 8/21/06			

Cana D	ate Mea	ourou						=		
			SAME	LES						
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm		MATERIAL DESCRIPTION	Ilew 1	Installation Schematic	FIELD NOTES AND WELL DETAILS
	_						CONCRETE	:		Begin Drilling on 8/15/06
	_						Loose, dry, black and brown, SAND with gravel, with Slag	-		
	-	0204	. 8	10.0	1.2		Will be delegated			-2" Schedule 40 PVC
	5	0406	<u>5</u>	12.0	0.9		r-With brick fragments			
-650	-	0608	3	14.0	0.6		M. U. and the state beauty and track a course CAND trace fine			
	- 10–	0810	3	20.0	0.6		Medium dense, dry, dark brown, medium to coarse SAND trace fine gravel			
	-	1012	<u>6</u>	11.0	0.3		rTrace gravel  rMoist to wet, with rock fragments, with brick fragments, with gravel			
	-	1214	4	18.0	0.4		- FMoist			
	15-	1416	5	7.0	0.4	77777	Medium dense, moist, black, fine clayey SAND			t—Bentonite Pellets
-640	-	1618	1 2 2 2 WOH	22.0	2.9			- No.		
	- 20	1820	1 1 1	24.0	14.8		rLoose, moist to wet, with gravel			Sample submitted for lab analysis 17-MW02-2022
	-	2022	3	24.0	15.1		r Saturated			analysis 17-WVV02-2022
17-MW02	-	2224	1 2 2	24.0	15.1		ரDense, moist	-		
14/2006	<b>25</b>	2426	2	24.0	NA ———			-		Filter Sand #7
	-	2628	6	18.0	NA		√Wood Loose, saturated, brown, medium SAND	Γ.		— Slotted Pipe #10 PVC (0.010)
le: ODOI	30	2830	2 3 2	11.0	NA		End of Boring at 30' bgs			End Drilling on
Report 1_ODOT_WELLS; File: ODOT.GFJ; 12/14/2006 17-MW02 9 9 9 9 00	- -						Installed Monitoring Well	1		8/15/06
ort: 1_ODO	35–									
Rep							URS		_	

Project Location: Site 17
Project Number: 15016633

## Log of Boring 17-MW03

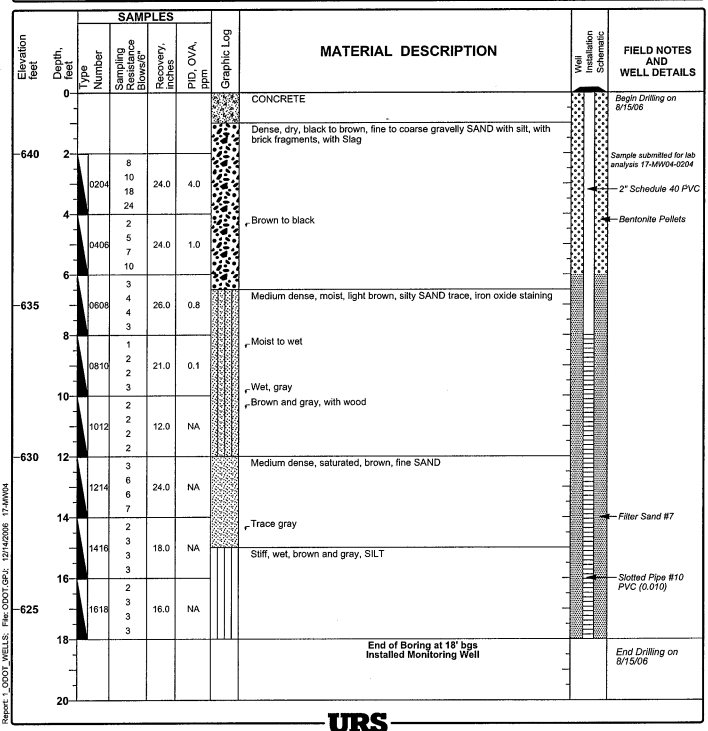
Total Depth of Borehole 16.0' bgs  Approximate cool
Approximate
Ground Elevation 629'
Borehole Set monitoring well



Project Location: Site 17
Project Number: 15016633

### Log of Boring 17-MW04

Date(s) 8/15/06	Logged By J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer 140# auto hammer	Total Depth of Borehole 18.0' bgs
Drill Rig Type CME-55	Drilling Contractor HAD, Inc.	Approximate Ground Elevation 642'
Location See Site Map	Sampling 2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 631.46 on 8/21/06		



Project Location: Site 17
Project Number: 15016633

## Log of Boring 17-SB01

Date(s) Drilled and Installed 8/14/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 30.0' bgs
Sampling 2" Split Spoon Method	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 668'
Drill Rig Type: CME-55	Groundwater NA .	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill <b>bentonite</b>	

	Locatio								
				SAME	PLES				
	Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
		0-						ASPHALT	Begin drilling on 8/14/06
		F						Loose, dry, brown and black, fine to coarse SAND with gravel, trace brick fragments, trace Slag	
	<del>-</del>	2-	0204	WOH 2 1 1	0.0	NA		brick fragments, trace Stag	
		4- - 6-	0406	WOH 1 WOH 1	3.0	1.0		rNo brick fragments, no Slag -	
	-660	8-	0608	1 2 2 5	20.0	0.0		Stiff, moist, brown, sandy CLAY With sand seams -	
	000	10-	0810	3 6 7 7	24.0	0.0		Loose to medium dense, dry, brown, fine to coarse SAND -	
		12-	1012	3 6 7 7	24.0	0.8		rLoose -	
	_	14-	1214	3 5 6	24.0	1.4		- 	
		16-	1416	2 4 8 8	20.0	2.0			
	_650	18-	1618	3 8 6 8	24.0	188		\times Dense, wet, light brown, SILT trace fine sand Loose, dry, light brown, fine to medium SAND	
	-030	20-	1820	2 5 5 8	16.0	186		rMoist, gray, trace, iron oxide staining	
		20	2022	2 5 6 9	24.0	602		r-Medium to coarse SAND	
17-SB01	_	24	2224	5 8 8	19.0	648		rDense, gray	
12/14/2006		26-	2426	3 6 5 7	24.0	1542		rDense, moist, fine SAND	
DOT.GPJ; 12	640	28	2628	4 7 9 10	17.0	3073		r-Dry	Sample submitted for lab analysis 17-SB01-2628
File: ODOT	040	30	2820	WOH 1 8 8	10.0	1061			
Report: 1_ODOT_BORINGS;		32						End of Boring at 30' bgs -	End drilling on 8/14/06
t: 1_ODOT	_	34							
Report								URS	

DATA ASSESSMENT REPORT

### Data Assessment Report ODOT Innerbelt Study Site 17 – Terminal Oil

Reviewer: P. Schuler Date: November 16, 2006

Five soil samples, three groundwater samples, and two trip blanks were collected at the Terminal Oil site in Cleveland, Ohio, from July 14 through August 21, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Reques	sted Ana	lyses <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	PAH	TPH
A6H160369001	17-SB01-2628	08/14/2006	Soil	X	X	X
A6H160369002	17-MW02-2022	08/15/2006	Soil	X	X	X
A6H160369003	17-MW04-0204	08/15/2006	Soil	X	X	X
A6H160369004	17-MW04-0204-D	08/15/2006	Soil	X	X	X
A6H160369005	17-MW03-0406	08/15/2006	Soil	X	X	X
A6H160369006	TB-081406	08/15/2006	Trip Blank	X		
A6H230305001	17-MW-04	08/21/2006	Groundwater	X	X	
A6H230305002	17-MW-03	08/21/2006	Groundwater	X	X	
A6H230305003	17-MW-02	08/22/2006	Groundwater	X	X	
A6H230305004	TRIP BLANK	08/21/2006	Trip Blank	X		

<sup>(1)</sup> VOC = Volatile Organic Compounds [SW-846 Method 8260B]

PAH = Polynuclear Aromatic Hydrocarbons [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

(2) Samples 17-MW04-0204 and 17-MW04-0204-D are field duplicates.

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate (MS/MSD) recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

Positive detections for acetone in samples 17-MW04-0204, 17-MW04-0204-D, 17-MW03-0406, and 17-MW-03, for methylene chloride in samples 17-SB01-2628, 17-MW02-2022, 17-MW04-0204-D, and 17-MW-03, for 1,2,4-trichlorobenzene in sample 17-MW04-0204, and for gasoline range organics in sample 17-MW04-0204 were

- qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks and/or trip blanksat similar concentrations.
- The laboratory "B" flags on the C<sub>20</sub>-C<sub>34</sub> diesel range organics results for several samples, indicating that the analyte was detected in the method blank, were removed in the final data set because the sample results were greater than five times the blank concentration. The sample results are considered representative of site conditions and any contribution due to external contamination is negligible.
- One or more volatile internal standard responses were outside of the acceptance range in samples 17-MW02-2022, 17-MW04-0204, and 17-MW04-0204D. The results for all volatile analytes quantified from each of the noncompliant internal standards were qualified as estimated ("J" or "UJ").
- The C<sub>10</sub>-C<sub>20</sub> and C<sub>20</sub>-C<sub>34</sub> diesel range organics results for all soil samples were qualified as estimated ("J") due to a continuing calibration standard response above the upper QC limit. Results may be biased high.
- The C<sub>10</sub>-C<sub>20</sub> and C<sub>20</sub>-C<sub>34</sub> diesel range organics results for sample 17-MW02-2022 were qualified as estimated ("J") due to MS/MSD recoveries outside of the QC acceptance limits.
- The positive results for C<sub>20</sub>-C<sub>34</sub> diesel range organics in field duplicate samples 17-MW04-0204 and 17-MW04-0204-D were qualified as estimated ("J") due to poor precision between the results.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J"), in accordance with USEPA Contract Laboratory Program (CLP) conventions. The "J" qualifiers were retained with the numeric results in the final data set.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-4.

# Table 2-1 Analytical Data Summary Site 17 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6H160369001 17-SB01-2628 08/14/2006	A6H160369002 17-MW02-2022 08/15/2006	A6H160369003 17-MW04-0204 08/15/2006	A6H160369004 17-MW04-0204-D 08/15/2006	A6H160369005 17-MW03-0406 08/15/2006	A6H160369006 TB-081406 08/15/2007 ug/L
1,1,1-Trichloroethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/kg	670 U	6.3 UJ	6.5 UJ	6.3 UJ	5.6 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,1,2-Trichloroethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,1-Dichloroethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
e for a comment of	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,1-Dichloroethene 1,2,4-Trichlorobenzene	ug/kg	670 U	6.3 UJ	6.5 UJ	6.3 UJ	5.6 U	1.0 U
and the second second	ug/kg	1300 U	13 UJ	13 UJ	13 UJ	11 U	2.0 U
1,2-Dibromo-3-chloropropane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,2-Dibromoethane		670 U	6.3 UJ	0.60 J	6.3, UJ	5.6 U	1.0 U
1,2-Dichlorobenzene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,2-Dichloroethane	ug/kg	and the second of the second o	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
1,2-Dichloropropane	ug/kg	670 U	6.3 UJ	0.50 J	6.3 UJ	5.6 U	1.0 U
1,3-Dichlorobenzene	ug/kg	670 U	6.3 UJ	0.50 J	6.3 UJ	5.6 U	1.0 U
1,4-Dichlorobenzene	ug/kg	670 U		5.0 J	6.3 UJ	3.6 U	1.0 U
2-Butanone	ug/kg	2700 U	50	and the second of the second of the second	the second of the second of the	The state of the s	
2-Hexanone	ug/kg	2700 U	25 U	26 U	25 U	23 U	10 U
4-Methyl-2-pentanone	ug/kg	2700 U	25 U	26 U	25 U	23 U	10 U
Acetone	ug/kg	2700 U	25 U	27 U	34 U	53 U	10 U
Benzene	ug/kg	290 J	1.2 J	1.3 J	4.1 J	1.6 J	1.0 U
Bromodichloromethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Bromoform	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
omomethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
rbon disulfide	ug/kg	670 U	2.3 J	1.3 J	1.8 J	3.2 J	1.0 U
Carbon tetrachloride	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Chlorobenzene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 ∪
Chloroethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Chloroform	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Chloromethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
cis-1,2-Dichloroethene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
cis-1,3-Dichloropropene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Cyclohexane	ug/kg	180 J	140	13 U	0.93 J	0.73 J	1.0 U
Dibromochloromethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Dichlorodifluoromethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Ethylbenzene	ug/kg	2300	6.3 U	6.5 U	1.2 J	5.6 U	1.0 U
Isopropylbenzene	ug/kg	380 J	0.26 J	6.5 U	0.37 J	0.51 J	1.0 U
Methyl acetate	ug/kg	1300 U	13 U	13 U	13 U	11 U	10 U
Methyl tert-butyl ether	ug/kg	2700 U	25 U	26 U	25 U	23 U	5.0 U
Methylcyclohexane	ug/kg	350 J	59	0.88 J	1.6 J	1.4 J	1.0 U
Methylene chloride	ug/kg	670 U	6.3 U	6.5 U	7.1 U	5.6 U	0.68 J
Styrene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Tetrachloroethene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Toluene	ug/kg	3000	0.74 J	2.5 J	9.9	2.9 J	1.0 U
trans-1,2-Dichloroethene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
trans-1,3-Dichloropropene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Trichloroethene	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Trichlorofluoromethane	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
Vinyl chloride	ug/kg	670 U	6.3 U	6.5 U	6.3 U	5.6 U	1.0 U
\enes (total)	ug/kg	16000	1.5 J	2.6 J	9.6 J	3.2 J	2.0 U

lenes (total) ug/kg 16000 1.5 he analyte was analyzed for, but was not detected. Value shown is the reporting limit.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

### Table 2-2 Analytical Data Summary Site 17 Soil PAHs and TPH ODOT Innerbelt Study

PARAMETER	UNITS	A6H160369001 17-SB01-2628 08/14/2006	A6H160369002 17-MW02-2022 08/15/2006	A6H160369003 17-MW04-0204 08/15/2006	A6H160369004 17-MW04-0204-D 08/15/2006	A6H160369005 17-MW03-0406 08/15/2006
2-Methylnaphthalene	ug/kg	1700	120 J	360 J	320 J	270 J
Acenaphthene	ug/kg	1100 U	100 J	550 J	260 J	450 J
Acenaphthylene	ug/kg	1100 U	830 U	160 J	59 J	1900 U
Anthracene	ug/kg	1100 U	240 J	2400 J	690 J	970 J
Benzo(a)anthracene	ug/kg	41 J	490 J	4100 J	2300	1900
Benzo(a)pyrene	ug/kg	37 J	410 J	3500 J	2000 J	1600 J
Benzo(b)fluoranthene	ug/kg	52 J	480 J	4500	2500	1900
Benzo(ghi)perylene	ug/kg	25 ქ	240 J	2000 J	1300 J	910 J
Benzo(k)fluoranthene	ug/kg	1100 U	250 J	1500 J	1000 J	840 J
Chrysene	ug/kg	41 J	460 J	3900 J	2100	1700 J
Dibenz(a,h)anthracene	ug/kg	1100 U	66 J	540 J	360 J	280 J
Fluoranthene	ug/kg	85 J	1200	11000	4800	4500
Fluorene	ug/kg	1100 U	140 J	820 J	270 J	480 J
Indeno(1,2,3-cd)pyrene	ug/kg	1100 U	220 J	1900 J	1200 J	850 J
Naphthalene	ug/kg	2600	84 J	260 J	280 J	270 J
Phenanthrene	ug/kg	42 J	710 J	8100	2600	3500
Pyrene	ug/kg	68 J	870	8100	3800	3400
Gasoline Range Organics (C6-C	12) ug/kg	250000	1600	130 U	330	770
C10-C20	mg/kg	41 J	17 J	38 J	36 J	36 J
C20-C34	mg/kg	130 J	80 J	260 J	15 J	120 J
Percent Solids	%	74.2	79.7	76.7	78.8	88.8

<sup>)</sup> U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

 $<sup>\</sup>frac{1}{2}$  J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-3 Analytical Data Summary Site 17 Water Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6H230305001 17-MW-04 08/21/2006	A6H230305002 17-MW-03 08/21/2006	A6H230305003 17-MW-02 08/22/2006	A6H230305004 TRIP BLANK 08/21/2006
1,1,1-Trichloroethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	10 U	120 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	10 U	120 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	20 U	250 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	10 U	120 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	10 U	120 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	10 U	120 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	10 U	120 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	10 U	120 U	1.0 U
2-Butanone	ug/L	0.59 J	100 U	1200 U	10 U
2-Hexanone	ug/L	10 U	100 U	1200 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	100 U	1200 U	10 U
Acetone	ug/L	1.0 J	100 U	1200 U	10 U
Benzene	ug/L	14	10 U	3700	1.0 U
Bromodichloromethane	ug/L	1.0 U	10 U	120 U	1.0 U
Bromoform	ug/L	1.0 U	10 U	120 U	1.0 U
Bromomethane	ug/L	1.0 U	10 U	120 U	1.0 U
Carbon disulfide	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	10 U	120 U	1.0 U
Chlorobenzene	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
Chloroethane	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
Chloroform	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
Chloromethane	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	10 U	120 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	10 U	120 U	1.0 U
Cyclohexane	ug/L	1.0 U	10 U	72 J	1.0 U
Dibromochloromethane	ug/L	1.0 U	10 U	120 U	1.0 U
Dichlorodifluoromethane	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
Ethylbenzene	ug/L ug/L	1.0 U	10 U	1100	1.0 U
and the second of the second o	ug/L ug/L	1.0 U	10 U	45 J	1.0 U
Isopropylbenzene	ug/L ug/L	10 U	100 U	1200 U	10 U
Methyl acetate	ug/L ug/L	3.7 J	220	620 U	5.0 U
Methylorelehovens	ug/L ug/L	1.0 U	10 U	120 U	1.0 U
Methylcyclohexane		1.0 U	10 U	120 U	1.0 U
Methylene chloride	ug/L	1.0 U	10 U	120 U	1.0 U
Styrene	ug/L		10 U	120 U	1.0 U
Tetrachloroethene	ug/L	1.0 U		2800	1.0 U
Toluene	ug/L	1.1	10 U		a product of the first open and the
trans-1,2-Dichloroethene	ug/L	1.0 U	10 U	120 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	10 U	120 U	1.0 U
Trichloroethene	ug/L	1.0 U	10 U	120 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	10 U	120 U	1.0 U
Vinyl chloride	ug/L	1.0 U	10 U	120 U	1.0 U
Xylenes (total)	ug/L	0.95 J	20 U	6500	2.0 U

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-4 Analytical Data Summary Site 17 Water PAHs ODOT Innerbelt Study

PARAMETER	UNITS	A6H230305001 17-MW-04 08/21/2006	A6H230305002 17-MW-03 08/21/2006	A6H230305003 17-MW-02 08/22/2006
2-Methylnaphthalene	ug/L	0.20 U	0.22	25
Acenaphthene	ug/L	0.20 U	0.20 U	0.40 U
Acenaphthylene	ug/L	0.20 U	0.20 U	0.40 U
Anthracene	ug/L	0.20 U	0.35	0.40 U
Benzo(a)anthracene	ug/L	0.20 U	0.77	0.42
Benzo(a)pyrene	ug/L	0.20 U	0.58	0.40 U
Benzo(b)fluoranthene	ug/L	0.20 U	0.7	0.40 U
Benzo(ghi)perylene	ug/L	0.20 U	0.35	0.40 U
Benzo(k)fluoranthene	ug/L	0.20 U	0.31	0.40 U
Chrysene	ug/L	0.20 U	0.64	0.40 U
Dibenz(a,h)anthracene	ug/L	0.20 U	0.20 U	0.40 U
Fluoranthene	ug/L	0.20 U	1.6	0.62
Fluorene	ug/L	0.20 U	0.20 U	0.40 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U	0.3	0.40 U
Naphthalene	ug/L	0.20 U	0.22	43
Phenanthrene	ug/L	0.20 U	1.2	0.72
Pyrene	ug/L	0.20 U	1.4	0.49

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

### 6.7 SITE 18 - CLEVELAND FIRE STATION

According to the Cleveland City Directories, the site was listed as City of Cleveland from 1964 through 1999. The site was identified as a LUST site in the EDR Report.

According to the Cleveland Fire Prevention Bureau files, the site was occupied by Fire Station #28 in 1963. According to a Fire Inspection Report dated June 13, 1963, a 1,500-gallon gasoline UST was located at the site. A Report dated March 28, 1984 indicates the Fire Department Hazardous Materials Unit responded to a call regarding gasoline odors in the basement of the building. Approximately 25 gallons of gasoline were noted in the UST. A Report dated July 12, 1985 indicates gasoline odors were noted in the basement again. At this time, it was determined the 1,500-gallon UST should be removed from the ground, as well as an abandoned 2,000-gallon UST.

On September 16, 1965, a 550-gallon gasoline UST was installed. This UST reportedly was to be utilized for the emergency generator. A Report dated December 8, 1998 indicates gasoline odors were reported in the basement of the building. The odors were traced to a public sewer catch basin west of the building. Approximately 14 gallons of product/water were removed and disposed of. A permit dated May 23, 2003 indicates a 250-gallon diesel UST was removed from the site. The UST reportedly was utilized in association with the emergency generator.

According to the BUSTR files, a diesel UST was removed from the site in May 2003. The UST Closure Report was received in July 2003. A NFA Letter was issued for the closure on July 28, 2003.

At the time of the URS Phase I reconnaissance, the site was observed to be a City of Cleveland Fire Department Fire Station. No surface staining, stressed vegetation and/or the storage and handling of hazardous substances were observed.

#### 6.7.1 Field Activities

A total of three monitoring wells were proposed for Site 18. Groundwater was encountered in two soil borings (MW01 and MW02) during field activities and monitoring wells were installed. One soil borings (SB03) was installed at Site 18. Three soil and two groundwater samples were collected and analyzed for VOCs, PAHs, and/or TPH. A Sample Location Map is included as **Figure 6-3A**.

### 6.7.2 Site-Specific Geology/Hydrogeology

Soils at Site 18 consisted primarily of sand and silty sand with minor amounts of silt and silty clay. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 18 on August 22, 2006. Localized groundwater flow across Site 18 is likely influenced by both natural features and urban development,

**Phase II Findings** 

including paved surfaces, buildings, and underground utilities. The general flow direction at Site 18 is to the west, towards the Cuyahoga River.

### 6.7.3 Geophysical Survey

The geophysical survey at Site 18 included the asphalt and concrete-paved parking and drive surfaces southeast of the building.

• The southeastern area included the paved area approximately 200 feet east towards the bend in the Central Viaduct.

The survey transects were conducted in a north – south direction to approximately align transects along the earth's total magnetic field, which provides the greatest sensitivity to buried ferrous objects such as steel underground storage tanks. Transects were spaced five feet apart. The presence of buildings adjacent to surveyed areas prevented the use of a global positioning system (GPS) to obtain corresponding latitude and longitude datum necessitating the need to paint survey points on the pavement.

The survey southeast of the main building identified one area of anomalously high magnetic gradients. This area is shown on Figure 6-7B as anomaly B as described below:

B. An anomaly was detected immediately west of the gas station building that was attributed to a parked vehicle that was present at the time of the survey.

The results of this survey do not support the presence of a steel UST at Site 18. No further investigation is recommended to attempt to locate any USTs.

### 6.7.4 Soil Analytical Results

A total of six VOCs were detected in a soil sample (18-MW01-0406) submitted from Site 18. The chemicals detected in the sample submitted were 2-butanone (2.4 ug/kg), carbon disulfide (0.62 ug/kg), ethylbenzene (0.62 ug/kg), methyl tert-butyl ether (0.34 ug/kg), toluene (0.35 ug/kg), and total xylenes (4.0 ug/kg). All other VOCs were below the detection limits.

A total of seventeen PAHs were detected in a sample (18-MW01-0406) submitted from Site 18. Concentration of 2-methylnaphthanene (19 ug/kg), acenaphthene (17 ug/kg), acenaphthylene (12 ug/kg), anthracene (57 ug/kg), benzo(a)anthracene (200 ug/kg), benzo(a)pyrene (200 ug/kg), benzo(b)fluoranthene (270 ug/kg), benzo(g,h,i)perylene (140 ug/kg), benzo(k)fluoranthene (91 ug/kg), chrysene (210 ug/kg), dibenz(a,h)anthracene (36 ug/kg), fluoranthene (430 ug/kg), fluorene (17 ug/kg), indeno(1,2,3)pyrene (120 ug/kg), naphthalene (13 ug/kg), phenanthrene (200 ug/kg), and pyrene (350 ug/kg) were detected in the soil sample submitted.

Diesel range total petroleum hydrocarbons were detected in the soil samples from Site 18, which is consistent with the concentrations of PAHs detected across Site 18. Concentrations of the middle petroleum fraction ranged from 1.7 mg/kg to 15 mg/kg. Concentrations of the heavy petroleum fraction ranged from 6.0 mg/kg to 55 mg/kg.

The analytical results are presented in Table 6-7A.

### 6.7.5 Groundwater Analytical Results

Eight VOCs were detected in groundwater samples collected from Site 18. Concentrations of 2-butanone (0.60 ug/L), acetone (2.7 ug/L), benzene (2.7 ug/L), cyclohexane (0.15 ug/L), ethylbenzene (0.43 ug/L), methyl tert-butyl ether (5 ug/L), toluene (0.51 ug/L and 0.58 ug/L), and total xylenes (1.4 ug/L) were detected in the groundwater samples submitted. All other VOCs were below the detection limits.

No PAHs were detected in the groundwater samples submitted from Site 18.

The analytical results are presented in Table 6-7B.

### 6.7.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on **Tables 6-7A** and **6-7B**.

None of the VOCs, PAHs or TPH concentrations detected in the soils at Site 18 exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or BUSTR closure action levels.

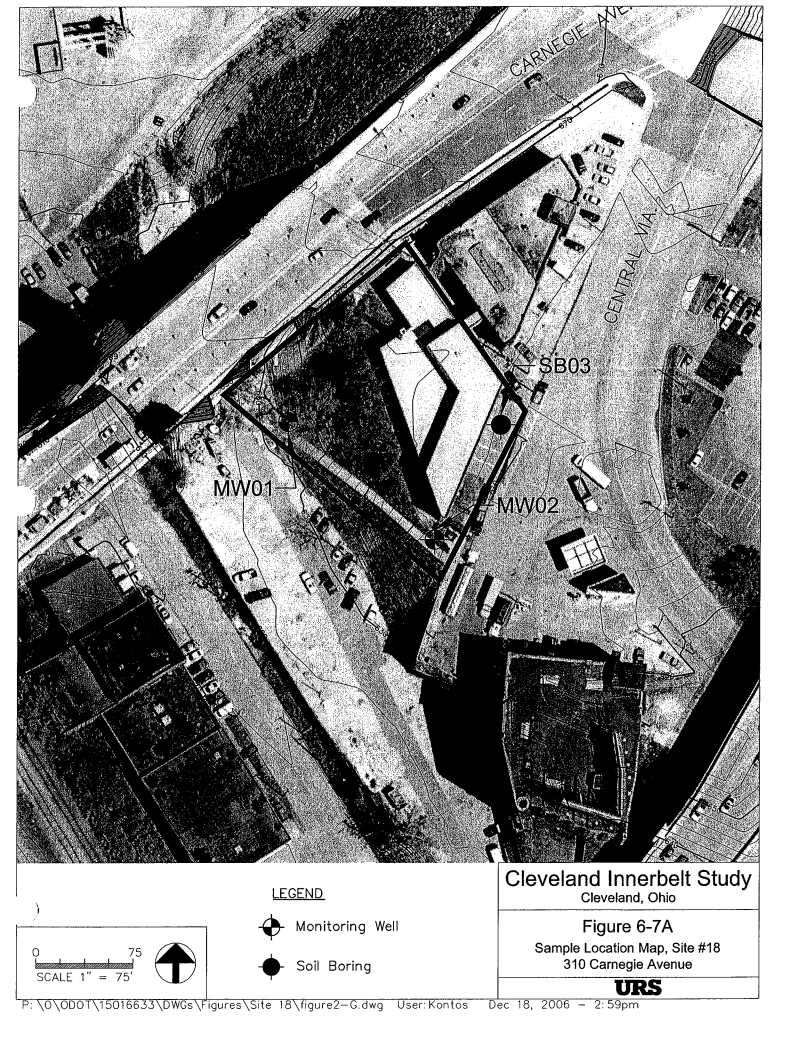
There were no PAHs detected in any of the groundwater samples submitted from Site 18.

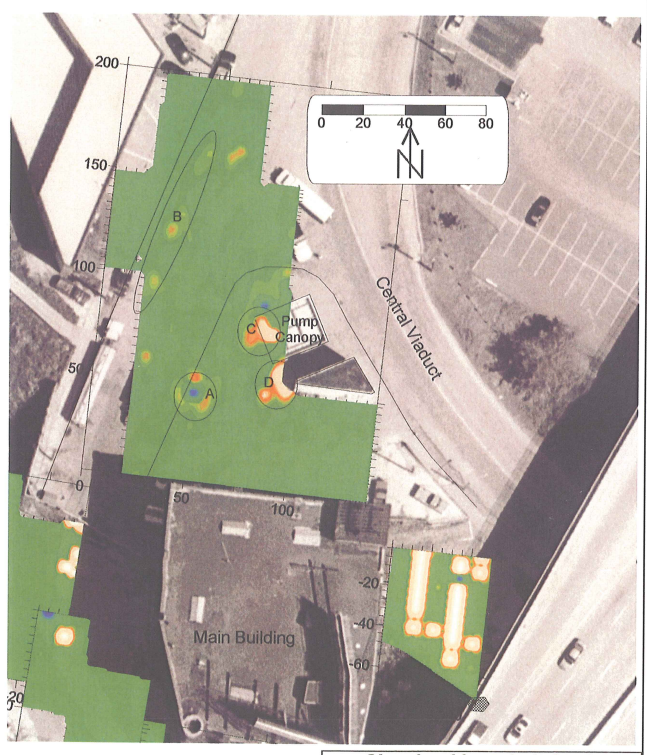
None of the VOCs concentrations detected at Site 18 exceeded the OEPA VAP standard for generic unrestricted potable use standards and/or BUSTR closure action levels.

#### 6.7.7 Conclusions

Based on the analytical results, it is unlikely that the soils and groundwater at Site 18 would require special management during construction.

**FIGURES** 





## Cleveland Innerbelt Study Cleveland, Ohio

### Figure 6-7B

Magnetometer Survey Results Cleveland Fire Station, Site 18

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**TABLES** 

Table V-7A
Summary of Detected Chemicals in Soil
Site 18 - Cleveland Fire Station
ODOT Innerbelt Study
Cleveland, Ohio

	20 Bit 12		VAP Commercial	VAE construction		18-MW01-0406 08/16/2006	18-MW02-1416 08/17/2006	18-SB03-1618 08/17/2006
PA	PARAMETER	UNITS	Standard!!?	Standard <sup>(2)</sup>	Action Level			) 
	2-Butanone	ug/kg	71,600,000	80,000,000	The state of the s	2.4 J	22 U	21 U
;	Carbon disulfide	ug/kg	720,000	720,000	•	0.62 J	5.5 U	5.3 U
sე(	Ethylbenzene	ug/kg	230,000	230,000	45,500	0.62 კ	5.5 U	5.3 U
ΟΛ	Methyl tert-butyl ether	ug/kg	7,200,000	7,200,000	470	0.34 J	22 U	21 U
	Toluene	ug/kg	520,000	520,000	49,100	0.35 J	5.5 U	5.3 U
	Xylenes (total)	ug/kg	160,000	160,000	15,700	4.0 J	11 U	11 0
	2-Methylnaphthalene	ug/kg		•		19 J	360 U	350 U
	Acenaphthene	ug/kg	180,000,000	530,000,000	1	17 J	360 U	350 U
	Acenaphthylene	ug/kg	•	•	•	12 J	360 U	350 U
	Anthracene	ug/kg	880,000,000	1,000,000,000		57 J	360 ∪	350 U
	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	200 J	360 U	320 U
•	Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	200 J	360 U	350 U
1	Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	270 J	360 U	350 U
S	Benzo(ghi)perylene	ug/kg		•	- The state of the	140 J	360 U	350 U
H∀c	Benzo(k)fluoranthene	ug/kg	630,000	8,100,000	110,000	91	360 U	350 U
<u>, , , , , , , , , , , , , , , , , , , </u>	Chrysene	ug/kg	6,700,000	41,000,000	1,100,000	210 J	360 U	350 U
	Dibenz(a,h)anthracene	ug/kg	6,700	41,000	1,100	36 J	360 U	350 U
	Fluoranthene	ug/kg	33,000,000	170,000,000	•	430	360 U	350 U
	Fluorene	ug/kg	120,000,000	340,000,000	100000000000000000000000000000000000000	17 J	360 U	350 U
	Indeno(1,2,3-cd)pyrene	ug/kg	67,000	410,000	11,000	120 J	360 U	350 U
	Naphthalene	ug/kg	530,000	1,900,000	39,800	13 J	360 U	350 U
	Phenanthrene Present September 2015 Present Pr	ug/kg				200 J	360 U	350 U
	Pyrene	ug/kg	25,000,000	130,000,000	-	350 J	360 U	350 U
Hd	C10-C20	mg/kg			2,000	15	1.7 ل	2.3 J
T	C20-C34	mg/kg	1	1	5,000	55	6.0	10 C

<sup>-- =</sup> Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial/ Industrial Land Use

<sup>(2)</sup> VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Table 6-7B
Summary of Detected Chemicals in Water
Site 18 - Cleveland Fire Station
ODOT Innerbelt Study
Cleveland, Ohio

	PARAMETER	SLIND	VAP upus/ BDUPUS <sup>(1)</sup>	BUSTR Closure Action Level	18-MW-01 08/22/2006	18-MW-02 08/22/2006
1L	2-Butanone	ug/L	6800	1	0.60 ل	10 U
	Acetone	ng/L	1600		2.7 J	10 U
	Benzene	ng/L	ري د	2	1.0 U	2.7
	Cyclohexane	ng/L	ì	1	1.0 U	0.15 J
, ;	Ethylbenzene	ug/L	200	200	1.0 U	0.43 J
	yl ether	ng/L		40	9	2.0 ∪
		ug/L	1000	1000	0.51 J	0.58 J
		ug/L	-	10000	2.0 U	1.4 J

-- = Standard not available

 $\mathsf{U}=\mathsf{The}$  analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

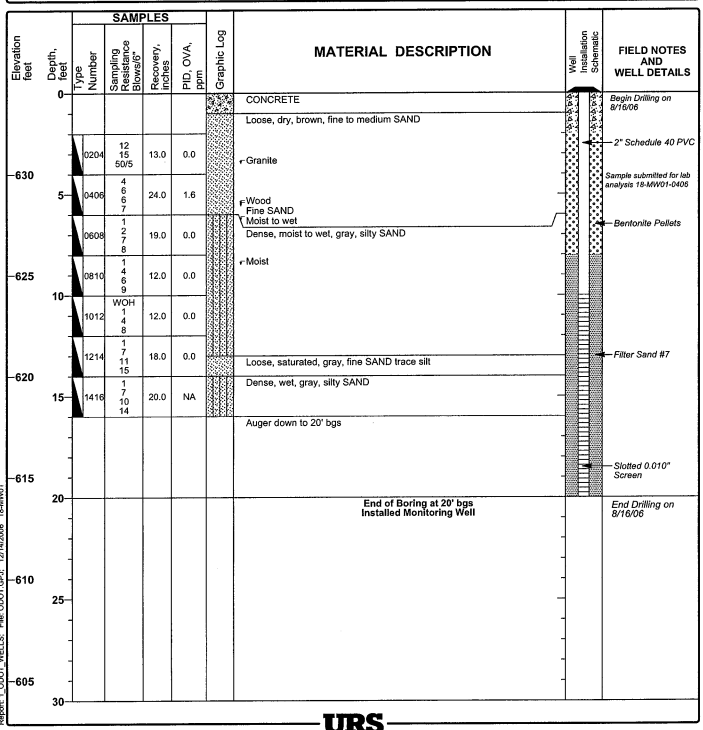
(1) VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

**BORING LOGS** 

Project Location: Site 18
Project Number: 15016633

### Log of Boring 18-MW01

Date(s) B/16/06 /	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 20.0′ bgs
Drill Rig Type LC-60	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 634'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 623.10 on 8/22/06			



Project Location: Site 18
Project Number: 15016633

## Log of Boring 18-MW02

Date(s) 8/17/06	Logged J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data 140# auto hammer	Total Depth of Borehole 28.0′ bgs
Drill Rig Type LC-60	Drilling Contractor HAD, Inc.	Approximate Ground Elevation 666'
Location See Site Map	Sampling Method(s) 2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 642.37 on 8/22/06		

	-		SAME	PLES						
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	Well		FIELD NOTES AND WELL DETAILS
-665	ا						CONCRETE	4 6 6 6 6	4	Begin Drilling on 8/17/06
_003							Loose, dry, brown, fine SAND	4	4	
	-	0204	1 2 4 4	12.0	0.5				4.4.4.4.	
-660	5-	0406	1 2 3 3	6.0	313		r Medium to coarse SAND trace fine gravel			
	-	0608	1 2 5 8	16.0	224					2" Schedule 40 PVC
	10-	0810	WOH 5 11 3	7.0	48.9		r-With coarse gravel -			
-655		1012	WOH 2 2	17.0	58.6					Bentonite Pellets
***			3				Stiff, moist, brown, SILT Stiff, moist, brown, silty CLAY			
		1214	1 2 2 4	24.0	155		Soft, wet  Loose, dry to moist, brown, fine SAND			
	15-	1416	WOH 1 2 2	12.0	571		ா1-Inch layer of moist to wet, with silt 			Sample submitted for lab analysis 18-MW02-1416
<b>⊢650</b>	1	1618	WOH 4 7 7	10.0	152		Loose, dry, brown, fine SAND			
	20-	1820	WOH 3 5 7	14.0	227		Fine to medium SAND			
-645	20	2022	WOH 2 5 4	8.0	108					
		2224	WOH 2 3 4	24.0	NA		- Saturated			
-645 -640	25-	2426	WOH 2 2 3	24.0	NA		→ Stiff, moist, brown, SILT			
040	-						Stiff, wet, brown, silty CLAY			— Slotted 0.010" Screen
	†					.1.1/2	End of Boring at 28' bgs Installed Monitoring Well			End Drilling on 8/17/06
	30						URS		1	

Project: ODOT - Innerbelt Corridor
Project Location: Site 18
Project Number: 15016633

## Log of Boring 18-SB03

Date(s) Drilled and Installed 8/17/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth 30.0' bgs
Sampling 2" Split Spoon Method	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 666'
Drill Rig Type: LC-60	Groundwater NA Level(s)	Hammer 140# auto hammer Data
Boring See Site Map	Borehole Backfill <b>bentonite</b>	

			SAMI	PLES				
Elevation feet	Depth, P feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm		MATERIAL DESCRIPTION	FIELD NOTES
-	-						CONCRETE  Loose, dry, brown, medium to coarse SAND with medium to coarse	Begin drilling on 8/17/06
	<b>2</b> —	0204	8 2 3	8.0	0.0		gravel	-
	4-	0406	4 6 3 6 11	12.0	0.4			-
-660	6	0608	11	13.0	0.2		rFine to medium SAND with fine gravel	-
	8-	0810	2 6 7	20.0	3.1		rMedium to coarse SAND trace gravel	
L	10-	1012	- 6 2 5 7	18.0	2.3	Ш	Dense, dry, brown, silty SAND	
	12-	1012	7	10.0	2.5		Dense, dry, brown, silty SAND	
	14-	1214	2 6 7 13 WOH	20.0	1.0		Loose, dry, brown and gray, fine to medium SAND Medium dense, brown, fine SAND	-
	-	1416	1 4	10.0	1.3		rMedium dense, brown, fine SAND	-
<b>−650</b>	16-	1618	MÖH	22.0	6.3			Sample submitted for lab analysis 18-SB03-1618
	18	1820	WOH 5 9 11	18.0	0.2			
-	20-	2022	WOH 3 7 7	12.0	2.7			
	22- - 24-	2224	WOH 3 7 7	12.0	2.7			
-640	24-	2426	WOH 4 3 7	20.0	4.6			
-040	20- - 28-	2628	8	12.0	2.5			
	30-	2830	WOH 3 4 7	10.0	1.8			
	32-						End of Boring at 30' bgs	End drilling on 8/17/06
1	34-							
	J			L				<u> </u>
							——URS———	

DATA ASSESSMENT REPORT

#### **Data Assessment Report ODOT Innerbelt Study** Site 18 - Cleveland Fire Station

Reviewer: P. Schuler **Date: November 16, 2006** 

Five soil samples, three groundwater samples, and two trip blanks were collected at the Cleveland Fire Station site at 310 Carnegie Avenue in Cleveland, Ohio, from July 16 through 22, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1 Sample and Analysis Summary

		Sample		Reques	sted Ana	lyses <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	PAH	TPH
A6H160361001	18-MW01-0406	08/16/2006	Soil	X	X	X
A6H170327001	18-SB03-1618	08/17/2006	Soil	X	X	X
A6H170327002	18-MW02-1416	08/17/2006	Soil	X	X	X
A6H230309001	18-MW-01	08/22/2006	Groundwater	X	X	
A6H230309002	18-MW-02	08/22/2006	Groundwater	X	X	

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B]

= Polynuclear Aromatic Hydrocarbons [SW-846 Method 8270C] PAH

= Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (OC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate (MS/MSD) recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Positive detections for acetone in samples 18-MW01-0406 and for methylene chloride in all soil samples were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks and/or trip blanks at similar concentrations (the trip blank associated with the soil samples was logged in with samples from another site).
- The laboratory "B" flag on the C<sub>20</sub>-C<sub>34</sub> diesel range organics result for sample 18-MW01-0406, indicating that the analyte was detected in the method blank, was removed in the final data set because the sample result was greater than five times the blank concentration. The sample result is considered representative of site conditions and any contribution due to external contamination is negligible.

- One internal standard response was outside of the acceptance range in sample 18-MW01-0406. The results for all volatile analytes quantified from the noncompliant internal standard were qualified as estimated ("J" or "UJ").
- The C<sub>10</sub>-C<sub>20</sub> and C<sub>20</sub>-C<sub>34</sub> diesel range organics results for sample 18-MW01-0406 were qualified as estimated ("J") due to MS/MSD recoveries outside of the QC acceptance limits.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J"), in accordance with USEPA Contract Laboratory Program (CLP) conventions. The "J" qualifiers were retained with the numeric results in the final data set.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-4.

# Table 2-1 Analytical Data Summary Site 18 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6H160361001 18-MW01-0406 08/16/2006	A6H170327001 18-SB03-1618 08/17/2006	A6H170327002 18-MW02-1416 08/17/2006
1,1,1-Trichloroethane	ug/kg	5.8 U	5.3 U	5.5 U
1,1,2,2-Tetrachloroethane	ug/kg	5.8 UJ	5.3 U	5.5 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.8 U	5.3 U	5.5 U
1,1,2-Trichloroethane	ug/kg	5.8 U	5.3 U	5.5 U
1,1-Dichloroethane	ug/kg	5.8 U	5.3 U	5.5 U
1.1-Dichloroethene	ug/kg	5.8 U	5.3 U	5.5 U
1,2,4-Trichlorobenzene	ug/kg ug/kg	5.8 UJ	5.3 U	5.5 U
1,2-Dibromo-3-chloropropane	ug/kg	12 UJ	11 U	11 U
in the commence were set the free contract of the	1	5.8 U	5.3 U	5.5 U
1,2-Dibromoethane	ug/kg			5.5 U
1,2-Dichlorobenzene	ug/kg	5.8 UJ	5.3 U 5.3 U	5.5 U
1,2-Dichloroethane	ug/kg	5.8 U		
1,2-Dichloropropane	ug/kg	5.8 U	5.3 U	5.5 U
1,3-Dichlorobenzene	ug/kg	5.8 UJ	5.3 U	5.5 U
1,4-Dichlorobenzene	ug/kg	5.8 UJ	5.3 U	5.5 U
2-Butanone	ug/kg	2.4 J	21 U	22 U
2-Hexanone	ug/kg	23 U	21 U	22 U
4-Methyl-2-pentanone	ug/kg	23 U	21 U	22 U
Acetone	ug/kg	30 U	21 U	22 U
Benzene	ug/kg	5.8 U	5.3 U	5.5 U
Bromodichloromethane	ug/kg	5.8 U	5.3 U	5.5 U
Bromoform	ug/kg	5.8 U	5.3 U	5.5 U
Bromomethane	ug/kg	5.8 U	5.3 U	5.5 U
Carbon disulfide	ug/kg	0.62 J	5.3 U	5.5 U
Carbon tetrachloride	ug/kg	5.8 U	5.3 U	5.5 U
Chlorobenzene	ug/kg	5.8 U	5.3 U	5.5 U
Chloroethane	ug/kg	5.8 U	5.3 U	5.5 U
Chloroform	ug/kg	5.8 U	5.3 U	5.5 U
Chloromethane	ug/kg	5.8 U	5.3 U	5.5 U
cis-1,2-Dichloroethene	ug/kg	5.8 U	5.3 U	5.5 U
cis-1,3-Dichloropropene	ug/kg	5.8 U	5.3 U	5.5 U
Cyclohexane	ug/kg	12 U	11 U	11 U
Dibromochloromethane	ug/kg	5.8 U	5.3 U	5.5 U
Dichlorodifluoromethane	ug/kg	5.8 U	5.3 U	5.5 U
Ethylbenzene	ug/kg	0.62 J	5.3 U	5.5 U
Isopropylbenzene	ug/kg	5.8 U	5.3 U	5.5 U
The second of th	ug/kg ug/kg	12 U	11 U	11 U
Methyl acetate Methyl tert-butyl ether	ug/kg ug/kg	0.34 J	21 U	22 U
the set as the second of the s		12 U	11 U	11 U
Methylcyclohexane	ug/kg	and the second second second second second	and the second second	
Methylene chloride	ug/kg	5.8 U	7.6 U	8.9 U
Styrene	ug/kg	5.8 U	5.3 U	5.5 U
Tetrachloroethene	ug/kg	5.8 U	5.3 U	5.5 U
Toluene	ug/kg	0.35 J	5.3 U	5.5 U
trans-1,2-Dichloroethene	ug/kg	5.8 U	5.3 U	5.5 U
trans-1,3-Dichloropropene	ug/kg	5.8 U	5.3 U	5.5 U
Trichloroethene	ug/kg	5.8 U	5.3 U	5.5 U
Trichlorofluoromethane	ug/kg	5.8 U	5.3 U	5.5 U
Vinyl chloride	ug/kg	5.8 U	5.3 U	5.5 U
Xylenes (total)	ug/kg	4.0 J	11 U	11 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

#### Table 2-2 Analytical Data Summary Site 18 Soil PAHs and TPH ODOT Innerbelt Study

PARAMETER	UNITS	A6H160361001 18-MW01-0406 08/16/2006	A6H170327001 18-SB03-1618 08/17/2006	A6H170327002 18-MW02-1416 08/17/2006
2-Methylnaphthalene	ug/kg	19 J	350 U	360 U
Acenaphthene	ug/kg	17 J	350 U	360 U
Acenaphthylene	ug/kg	12 J	350 U	360 U
Anthracene	ug/kg	57 J	350 U	360 U
Benzo(a)anthracene	ug/kg	200 J	350 U	360 U
Benzo(a)pyrene	ug/kg	200 J	350 U	360 U
Benzo(b)fluoranthene	ug/kg	270 J	350 U	360 U
Benzo(ghi)perylene	ug/kg	140 J	350 U	360 U
Benzo(k)fluoranthene	ug/kg	91 J	350 U	360 U
Chrysene	ug/kg	210 J	350 U	360 U
Dibenz(a,h)anthracene	ug/kg	36 J	350 U	360 U
Fluoranthene	ug/kg	430	350 U	360 U
Fluorene	ug/kg	17 J	350 U	360 U
Indeno(1,2,3-cd)pyrene	ug/kg	120 J	350 U	360 U
Naphthalene	ug/kg	13 J	350 U	360 U
Phenanthrene	ug/kg	200 J	350 U	360 U
Pyrene	ug/kg	350 J	350 U	360 U
Gasoline Range Organics (C6-C12)	ug/kg	120 U	110 U	110 U
C10-C20	mg/kg	15	2.3 J	1.7 J
C20-C34	mg/kg	55	10 J	6.0
Percent Solids	%	86.4	94.8	91

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

# Table 2-3 Analytical Data Summary Site 18 Water Volatiles ODOT Innerbelt Study

		A6H230309001 18-MW-01	A6H230309002 18-MW-02
PARAMETER	UNITS	08/22/2006	08/22/2006
1,1,1-Trichloroethane	ug/L	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U
2-Butanone	ug/L	0.60 J	10 U
2-Hexanone	ug/L	10 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U
Acetone	ug/L	2.7 J	10 U
Benzene	ug/L	1.0 U	2.7
Bromodichloromethane	ug/L ug/L	1.0 U	1,0 U
The Control of the Co		1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U
Carbon tetrachloride	ug/L		1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U
Chlorostane	ug/L	1.0 U	1.0 U
Chloroform	ug/L	1.0 U	
Chloromethane	ug/L	1.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	0.15 J
Dibromochloromethane	ug/L	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	0.43 J
Isopropylbenzene	ug/L	1.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U
Methyl tert-butyl ether	ug/L	5.0	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U
Methylene chloride	ug/L	1.0 U	1.0 U
Styrene	ug/L	1.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U
Toluene	ug/L	0.51 J	0.58 J
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	1.4 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

 $J = \mbox{Estimated}$  concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-4 Analytical Data Summary Site 18 Water PAHs ODOT Innerbelt Study

PARAMETER	UNITS	A6H230309001 18-MW-01 08/22/2006	A6H230309002 18-MW-02 08/22/2006
2-Methylnaphthalene	ug/L	0.20 U	0.20 U
Acenaphthene	ug/L	0.20 U	0.20 U
Acenaphthylene	ug/L	0.20 U	0.20 U
Anthracene	ug/L	0.20 U	0.20 U
Benzo(a)anthracene	ug/L	0.20 U	0.20 U
Benzo(a)pyrene	ug/L	0.20 U	0.20 U
Benzo(b)fluoranthene	ug/L	0.20 U	0.20 U
Benzo(ghi)perylene	ug/L	0.20 U	0.20 U
Benzo(k)fluoranthene	ug/L	0.20 U	0.20 U
Chrysene	ug/L	0.20 U	0.20 U
Dibenz(a,h)anthracene	ug/L	0.20 U	0.20 U
Fluoranthene	ug/L	0.20 U	0.20 U
Fluorene	ug/L	0.20 U	0.20 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U	0.20 U
Naphthalene	ug/L	0.20 U	0.20 U
Phenanthrene	ug/L	0.20 U	0.20 U
Pyrene	ug/L	0.20 U	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

 $J = \mbox{Estimated}$  concentration because the result was below the sample reporting limit or quality control criteria were not met.

#### 6.8 SITE 19 - GILLOTA FUEL PRODUCTS

According to the Cleveland City Directories, the site was listed as various warehouses and a gas station from 1954 through 1999. The site was identified as LUST, RCRIS-SQG and UST site in the EDR Report.

According to the Cleveland Fire Prevention Bureau files, the site was occupied by a furniture company (206) and a gas station (300) in 1950. A Permit dated February 1952 indicates a 1,000-gallon gasoline UST was located on the 206 property. A Permit dated August 1952 indicates there were approximately 13,600 gallons of gasoline stored on the 300 site in two USTs. A report dated October 2, 1964 indicates part of the site was a vacant service station. There reportedly were at least three USTs associated with the site, which had been vacant for approximately 1.5 years.

A Permit dated November 1984 indicates two 10,000-gallon USTs of #2 fuel oil and three 12,000-gallon USTs of #1 fuel oil were to be installed on the site. According to the Permit, the USTs are located east of the existing building. A Tank Inventory for the site, dated January 1, 1992, indicates there were 18 USTs containing various amounts of petroleum products.

A letter dated July 10, 1992 from Centerior Energy to Gillota Fuel indicates the adjacent CEI property had been impacted by petroleum products emanating from the site. The letter concludes that they expected Gillota to manage the regulatory implications. In September 1992, four USTs; two 1,500-gallon, one 12,000-gallon and one 20,000-gallon, were removed from the site. At this time four USTs were installed to replace the removed USTs. A Permit, dated September 16, 1992, was filed to remove six USTs from the adjacent CEI property. The USTs were four 6,000-gallon used oil and two 8,000-gallon used oil. The Bureau reports associated with this removal indicated petroleum product was observed throughout the excavations. According to the BUSTR files, over-excavation of the soils associated with the USTs was conducted; approximately 5,666 tons of soil were removed. This incident attained NFA status on October 25, 1993; however, it was noted in the file the site had the potential to be re-contaminated by seepage water. The seepage water reportedly was emanating from a bridge abutment; the contaminants were thought to be associated with USTs operated on the adjacent property.

A Permit dated October 5, 1993 was issued for the removal of two 6,000-gallon USTs from the site. A Permit dated May 11, 1995 was issued for the removal of three 12,000-gallon fuel oil USTs, two 10,000-gallon gasoline USTs, and one 1,000-gallon fuel oil UST from the 300 site. A Closure Report was written for the removal of the six USTs and submitted to BUSTR requesting No Further Action. NFA status was granted for the removal of the six USTs on October 3, 1995.

A Permit dated July 14, 2000 was issued for the installation of UST piping around two 8,000-gallon gasoline/diesel USTs, one 6,000-gallon gasoline UST and one 4,000-gallon kerosene UST. A Permit dated August 9, 2002 indicates there are two USTs located at the 300 site; one 8,000-gallon and one 6,000-gallon, which both contain gasoline. A Permit dated August 9, 2002 indicates there are four USTs; one 8,000-gallon and two 6,000-gallon, which contain diesel and one 4,000-gallon kerosene.

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At the time of the URS Phase I reconnaissance, the site was a Marathon Gas Station and an empty warehouse. An underground storage tank cavity was observed on the gas station property. Surface staining and a petroleum odor were discerned on the gas station property, as well.

#### 6.8.1 Field Activities

A total of three monitoring wells were proposed for Site 19. During the Phase II field activities, obstructions were encountered at approximately four feet bgs. Attempts were made to relocate the borings, but due to underground utilities and obstructions, these borings were abandoned. One of the abandoned borings was located southwest of the retail building. The other abandoned boring was located north of the pump island along Central Viaduct. One boring was successfully advanced.

Groundwater was encountered in one soil boring (MW01) during field activities and monitoring wells were installed. One soil and one groundwater sample were collected and analyzed for VOCs, PAHs, and/or TPH. A Sample Location Map is included as **Figure 6-8A**.

#### 6.8.2 Site-Specific Geology/Hydrogeology

Soils at Site 19 consisted primarily of sand with interbedded silty sand and silty clay. Bedrock was not encountered in the soil borings, which were advanced to 35 feet bgs.

A groundwater elevation was measured at Site 19 on August 22, 2006. Localized groundwater flow across Site 19 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 19 is to the south-southeast, towards the Cuyahoga River.

### 6.8.3 Geophysical Survey

The geophysical survey at Site 19 included the asphalt-paved surfaces around the pump island and building.

• The northern area included the parking spaces directly north of the main building, the dispenser island and extended 200 feet north into the intersection with Central Viaduct.

The survey transects were conducted in a north – south direction to approximately align transects along the earth's total magnetic field, which provides the greatest sensitivity to buried ferrous objects such as steel underground storage tanks. Transects were spaced five feet apart. The presence of buildings adjacent to surveyed areas prevented the use of a global positioning system (GPS) to obtain corresponding latitude and longitude datum necessitating the need to paint survey points on the pavement.

The survey north of the main building identified four areas of anomalously high magnetic gradients. These areas are shown on Figure 6-8B as anomalies C and D as described below:

C. An anomaly was detected immediately west of the pump island that was attributed to the pump island.

D. An elongated anomaly was detected along the street. It is assumed that this anomaly is related to a buried utility line.

The results of this survey do not support the presence of a steel UST at Site 19. No further investigation is recommended to attempt to locate any USTs.

#### 6.8.4 Soil Analytical Results

A total of seven VOCs were detected in the soil sample submitted from Site 19. The chemicals detected in the samples submitted were benzene (410 ug/kg), cyclohexane (1,100 ug/kg), ethylbenzene (26,000 ug/kg), isopropylbenzene (4,200 ug/kg), methylcyclohexane (2,600 ug/kg), toluene (16,000 ug/kg), and total xylenes (130,000 ug/kg). All other VOCs were below the detection limits.

A total of six PAHs were detected in the soil sample submitted from Site 19. Concentrations of 2-methylnaphthanene (9,500 ug/kg), benzo(a)anthracene (95 ug/kg), fluoranthene (160 ug/kg), naphthalene (7,000 ug/kg), phenanthrene (120 ug/kg), and pyrene (140 ug/kg) were detected in the soil sample submitted.

Gasoline and diesel range total petroleum hydrocarbons were detected in the soil samples from Site 19. The highest concentrations were found in the gasoline range, which is consistent with the concentrations of VOCs detected across Site 19. The concentration of the light petroleum fraction was 77,000 ug/kg. The concentration of the middle petroleum fraction was 200 mg/kg and the concentration of the heavy petroleum fraction was 33 mg/kg.

The analytical results are presented in **Table 6-8A**.

### 6.8.5 Groundwater Analytical Results

Four VOCs were detected in the groundwater sample collected from Site 19. Concentrations of benzene (21,000 ug/L), methyl tert-butyl ether (1,100 ug/L), toluene (1,000 ug/L), and total xylenes (450 ug/L) were detected in the groundwater sample submitted. All other VOCs were below the detection limits.

Four PAHs were detected in the groundwater sample submitted from Site 19. Concentrations of 2-methylnaphthanene (2.6 ug/L), acenaphthene (0.28 ug/L), naphthalene (4.2 ug/L), and phenanthrene (0.48 ug/L) were detected in the groundwater sample submitted from Site 19. All other PAHs were below the detection limits.

The analytical results are presented in Table 6-8B.

### 6.8.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic

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Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on **Tables 6-8A** and **6-8B**.

The concentration of benzene (410 ug/kg) and total xylenes (130,000 ug/kg) detected in soil sample 19-MW01-2628 exceeded the BUSTR closure action levels. No other VOCs detected exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

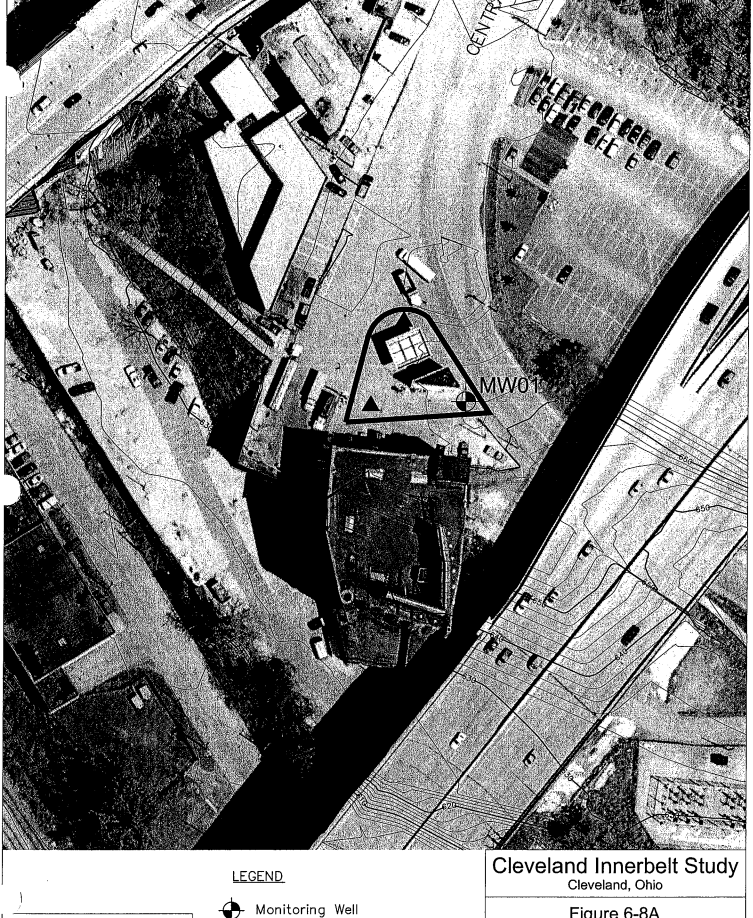
No PAHs detected exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

The concentrations of benzene, methyl tert-butyl ether, and toluene detected in groundwater sample 19-MW01 exceeded the OEPA VAP standard for generic unrestricted potable use standards and BUSTR closure action levels.

#### 6.8.7 Conclusions

Based on the analytical results, the soil and groundwater at Site 19 may require special disposal and/or worker protection protocols (plan note) during construction activities.

**FIGURES** 

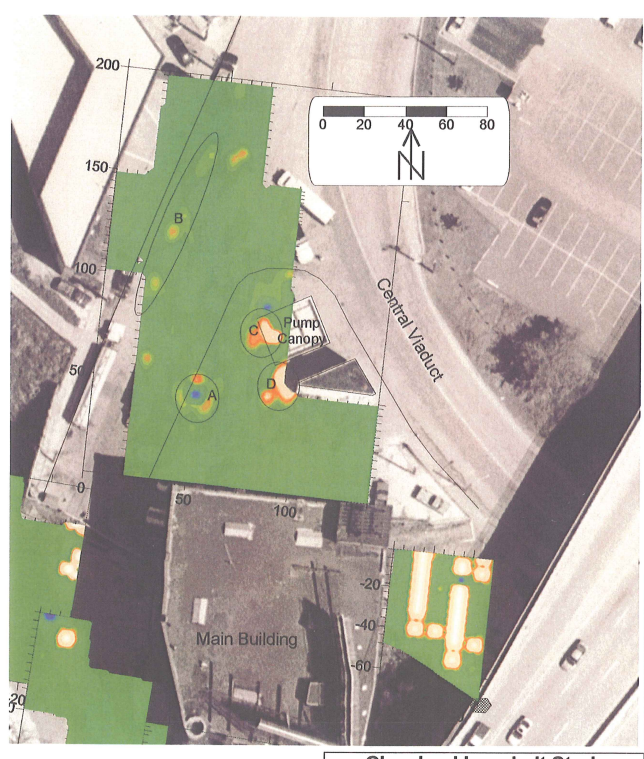


Abandoned Boring

### Figure 6-8A

Sample Location Map, Site #19 206-300 Central Viaduct

Dec 18, 2006 - 3:03pm



Cleveland Innerbelt Study
Cleveland, Ohio

# Figure 6-8B

Magnetometer Survey Results Gillota Fuel Products, Site 19

**URS** 

**TABLES** 

Summary of Detected Chemicals in Soil Site 19 - Gillota Fuel Products **ODOT Innerbelt Study** Cleveland, Ohio Table v-8A

			VAP Commercial/ Commercial/	VAP Construction	BUSTRE Closure Action	19-MW01-2628 08/16/2006
9	PARAMETER	UNITS	Standard 🖰 🐇	Standard <sup>2</sup>	Level	
	Вептення в верой в вер	ug/kg	100,000	310,000	149	410 J
<del>`</del>	Cyclohexane	ug/kg	•	•	1	1100 J
S(	Ethylbenzene	ug/kg	230,000	230,000	45,500	26000
00/	Isopropylbenzene	ug/kg	860,000	860,000	1	4200
\	Methylcyclohexane	ug/kg		-	-	2600 J
	Toluene	ug/kg	520,000	520,000	49,100	16000
	Xylenes (total)	ug/kg	160,000	160,000	15,700	130000
	2-Methylnaphthalene	ug/kg	1	1	1	9500
	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	ր 56
sH/	Fluoranthene	ug/kg	33,000,000	170,000,000	ı	160 J
/d	Naphthalene	ug/kg	530,000	1,900,000	39,800	2000
	Phenanthrene	ug/kg		1	1	120 J
	Pyrene	ug/kg	25,000,000	130,000,000	-	140 J
ŀ	Gasoline Range Organics (C6-C12)	ug/kg	and any state of the control of the		1,000,000	77000
ldΤ	C10-C20	mg/kg	•	1	2,000	200
	C20-C34	mg/kg	:	1	5,000	33

-- = Standard not available

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

(1) VAP Generic Direct Contact Soil Standard, Commercial/Industrial Land Use (2) VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Summary of Detected Chemicals in Water Site 19 - Gillota Fuel Products ODOT Innerbelt Study Cleveland, Ohio Tabl

19-MW-01 Action 08/22/2006	21000	S. M. San C. Charles		10000 450 J	2.6	0.28	4.2	0.48
BUSTR VAPUPUS Closura Act RDUPUS <sup>(1)</sup> Level	S Control of the Cont	40 40			:	089	140	•
UNITS	ng/L	ug/L	ug/L	ug/L	ug/t	ng/L	ng/L	ug/L
PARAMETER	Benzene	Methyl tert-butyl ether	tours continued to the continued of the	Xylenes (total)	2-Methylnaphthalene	Acenaphthene	Naphthalene	Phenanthrene
7	Г	sO	OΛ		Γ	s	lΑq	

-- = Standard not available

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

(1) VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

**BORING LOGS** 

**Project: ODOT - Innerbelt Corridor** 

Project Location: Site 19
Project Number: 15016633

# Log of Boring 19-MW01

Sheet 1 of 2

Date(s) 8/17/06 Drilled 8/17/06	Logged By	J. Kaminski	Checked By M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 35.0′ bgs
Drill Rig LC-60	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation <b>664</b>
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 632.20 on 8/22/06			

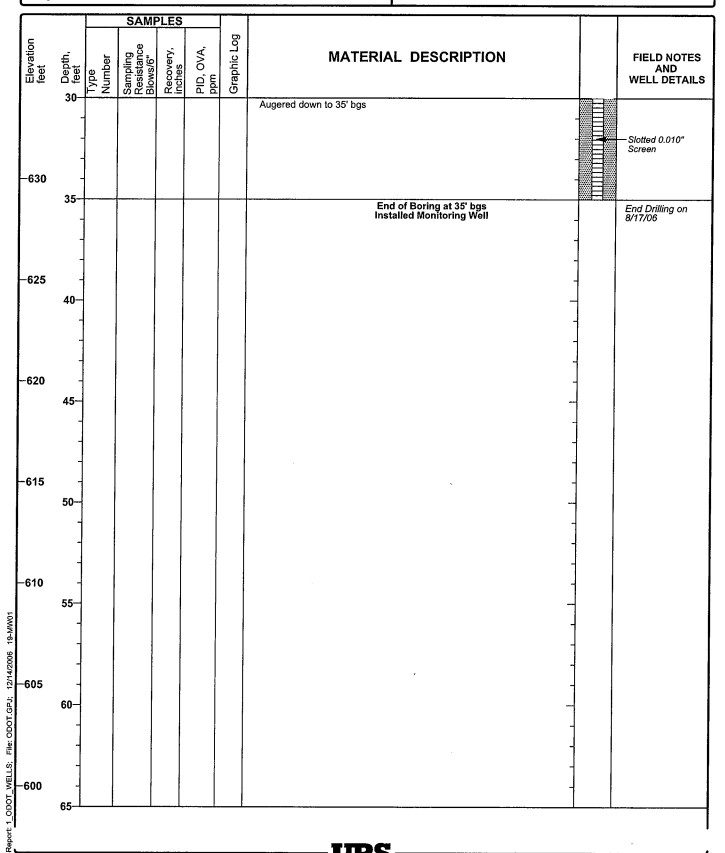
1			Γ	SAMI	PLES				Т		···
	Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"		PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION		Installation Schematic	FIELD NOTES AND WELL DETAILS
		- -		NA NA NA NA	NA	NA		CONCRETE  Loose, dry, brown, SAND with brick fragments, with rock fragments, with gravel	4444	4444	Begin Drilling on 8/17/06
	660	_	0204	27 4 5 3	14.0	1.6		Loose, dry, brown, medium SAND			
	660	5-	0406	2 5 5 2	8.0	1.4		ℯWith wood, no rock fragments –			2" Cabadula 40 DVC
		-	0608	WOH 1 2 2	11.0	0.0		Loose to medium dense, dry, brown, fine SAND			— 2" Schedule 40 PVC l ⊢Bentonite Pellets
	-655	10-	0810	WOH 1 2 3	6.0	0.4		rFine to medium SAND			
		10	1012	2	20.0	1.1		-Loose, medium SAND			
	650	- 1	1214	WOH 2 4 4	10.0	14.7		Loose, dry, dark gray, medium to coarse SAND trace, hydrocarbon odor			
	000	15-	1416	WOH 3 6 6	12.0	188		≁Medium SAND –			
		-	1618	1 3 5 7	8.0	211		Dense, moist to wet, brown, silty SAND -			
19-MW01	-645	20-	1820	WOH 6 6 9	24.0	412		Loose, dry, brown and gray, fine SAND			
12/14/2006 19-N		-	2022	WOH 7 8 10	11.0	322		- We diese des des des des des des des des des			
GPJ;	-640	-	2224	1 6 7 9	14.0	673		Medium dense, dry, brown, silty SAND with, hydrocarbon odor			
File: ODOT.		25-	2426	7 10 11	14.0	1688					Sample submitted for lab
_wells;			2628	8 14 16	16.0	>10000		<del>-</del>			analysis 19-MW01-2628
Report: 1_ODOT_WELLS;	-635	30	2830	4 10 12	10.0	>10000		r-Saturated			Filter Sand #7
8								URS			

**Project: ODOT - Innerbelt Corridor** 

Project Location: Site 19
Project Number: 15016633

# Log of Boring 19-MW01

Sheet 2 of 2



DATA ASSESSMENT REPORT

#### Data Assessment Report ODOT Innerbelt Study Site 19 – Gillota Fuel Products

Reviewer: P. Schuler Date: November 16, 2006

One soil sample and one groundwater sample were collected at the Gillota Fuel Products site in Cleveland, Ohio, on July 16 and 22, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Requested Analyses <sup>(1)</sup>				
Laboratory ID	Sample ID	Date	Matrix	VOC	PAH	TPH		
A6H160402001	19-MW01-2628	08/16/2006	Soil	X	X	X		
A6H230315001	19-MW-01	08/22/2006	Groundwater	X	X			

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B]

PAH = Polynuclear Aromatic Hydrocarbons [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate (MS/MSD) recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- The positive detection for methylene chloride in sample 19-MW01-2628 was qualified as nondetect ("U") due to the presence of methylene chloride in the associated method blank.
- The laboratory "B" flag on the C<sub>20</sub>-C<sub>34</sub> diesel range organics result for sample 19-MW01-2628, indicating that the analyte was detected in the method blank, was removed in the final data set because the sample result was greater than five times the blank concentration. The sample result is considered representative of site conditions and any contribution due to external contamination is negligible.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J"), in accordance with USEPA Contract Laboratory Program (CLP) conventions. The "J" qualifiers were retained with the numeric results in the final data set.

• All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-4.

# Table 2-1 Analytical Data Summary Site 19 Soil Volatiles ODOT Innerbelt Study

		A6H160402001 19-MW01-2628
PARAMETER	UNITS	08/16/2006
1,1,1-Trichloroethane	ug/kg	3000 U
1,1,2,2-Tetrachloroethane	ug/kg	3000 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	3000 U
1,1,2-Trichloroethane	ug/kg	3000 U
1,1-Dichloroethane	ug/kg	3000 U
1,1-Dichloroethene	ug/kg	3000 U
1,2,4-Trichlorobenzene	ug/kg	3000 U
1,2-Dibromo-3-chloropropane	ug/kg	5900 U
1,2-Dibromoethane	ug/kg	3000 U
1,2-Dichlorobenzene	ug/kg	3000 U
1,2-Dichloroethane	ug/kg	3000 U
1,2-Dichloropropane	ug/kg	3000 U
1,3-Dichlorobenzene	ug/kg	3000 U
1,4-Dichlorobenzene	ug/kg	3000 U
2-Butanone	ug/kg ug/kg	12000 U
2-Hexanone	ug/kg	12000 U
4-Methyl-2-pentanone	ug/kg	12000 U
Acetone	ug/kg	12000 U
Benzene	ug/kg	410 J
Bromodichloromethane	ug/kg	3000 U
Bromoform	ug/kg	3000 U
Bromomethane	ug/kg ug/kg	3000 U
Carbon disulfide	ug/kg	3000 U
Carbon tetrachloride		3000 U
Chlorobenzene	ug/kg	3000 U
Chloroethane	ug/kg	3000 U
Chloroform	ug/kg	3000 U
Chloromethane	ug/kg	3000 U
cis-1,2-Dichloroethene	ug/kg	3000 U
cis-1,3-Dichloropropene	ug/kg	3000 U
Cyclohexane	ug/kg	1100 J
Dibromochloromethane	ug/kg	3000 U
Dichlorodifluoromethane	ug/kg	3000 U
Consider the Consideration of	ug/kg	and the contract of the contra
Ethylbenzene Isopropylbenzene	ug/kg	26000
ALMER A MILLION TO CONTROL OF THE CO	ug/kg	4200
Methyl acetate	ug/kg	5900 U
Methyl tert-butyl ether	ug/kg	12000 U <b>2600 J</b>
Methylcyclohexane	ug/kg	
Methylene chloride	ug/kg	3000 U
Styrene	ug/kg	3000 U
Telvene	ug/kg	3000 U
Toluene	ug/kg	16000
trans-1,2-Dichloroethene	ug/kg	3000 U
trans-1,3-Dichloropropene	ug/kg	3000 U
Trichloroethene	ug/kg	3000 U
Trichlorofluoromethane	ug/kg	3000 U
Vinyl chloride	ug/kg	3000 U
Xylenes (total)	ug/kg	130000

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-2 Analytical Data Summary Site 19 Soil PAHs and TPH ODOT Innerbelt Study

PARAMETER	UNITS	A6H160402001 19-MW01-2628 08/16/2006
2-Methylnaphthalene	ug/kg	9500
Acenaphthene	ug/kg	4000 U
Acenaphthylene	ug/kg	4000 U
Anthracene	ug/kg	4000 U
Benzo(a)anthracene	ug/kg	95 J
Benzo(a)pyrene	ug/kg	4000 U
Benzo(b)fluoranthene	ug/kg	4000 U
Benzo(ghi)perylene	ug/kg	4000 U
Benzo(k)fluoranthene	ug/kg	4000 U
Chrysene	ug/kg	4000 U
Dibenz(a,h)anthracene	ug/kg	4000 U
Fluoranthene	ug/kg	160 J
Fluorene	ug/kg	4000 U
Indeno(1,2,3-cd)pyrene	ug/kg	4000 U
Naphthalene	ug/kg	7000
Phenanthrene	ug/kg	120 J
Pyrene	ug/kg	140 J
Gasoline Range Organics (C6-C12)	ug/kg	77000
C10-C20	mg/kg	200
C20-C34	mg/kg	33
Percent Solids	%	83.3

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

#### Table 2-3 Analytical Data Summary Site 19 Water Volatiles ODOT Innerbelt Study

		A6H230315001 19-MW-01
PARAMETER	UNITS	08/22/2006
1,1,1-Trichloroethane	ug/L	710 U
1,1,2,2-Tetrachloroethane	ug/L	710 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	710 U
1,1,2-Trichloroethane	ug/L	710 U
1,1-Dichloroethane	ug/L	710 U
1,1-Dichloroethene	ug/L	710 U
1,2,4-Trichlorobenzene	ug/L	710 U
1,2-Dibromo-3-chloropropane	ug/L	1400 U
1,2-Dibromoethane	ug/L	710 U
1,2-Dichlorobenzene	ug/L	710 U
1,2-Dichloroethane	ug/L	710 U
1,2-Dichloropropane	ug/L	710 U
1,3-Dichlorobenzene	ug/L	710 U
1,4-Dichlorobenzene	ug/L	710 U
2-Butanone	ug/L	7100 U
2-Hexanone	ug/L	7100 U
4-Methyl-2-pentanone	ug/L	7100 U
Acetone	ug/L	7100 U
Benzene	ug/L	21000
Bromodichloromethane	ug/L	710 U
Bromoform		710 U
Bromomethane	ug/L	710 U
The property of the second of	ug/L	and the state of the second
Carbon disulfide	ug/L	710 U
Carbon tetrachloride	ug/L	710 U
Chlorothers	ug/L	710 U
Chloroethane	ug/L	710 U
Chloroform	ug/L	710 U
Chloromethane	ug/L	710 U
cis-1,2-Dichloroethene	ug/L	710 U
cis-1,3-Dichloropropene	ug/L	710 U
Cyclohexane	ug/L	710 U
Dibromochloromethane	ug/L	710 U
Dichlorodifluoromethane	ug/L	710 U
Ethylbenzene	ug/L	710 U
Isopropylbenzene	ug/L	710 U
Methyl acetate	ug/L	7100 U
Methyl tert-butyl ether	ug/L	1100 J
Methylcyclohexane	ug/L	710 U
Methylene chloride	ug/L	710 U
Styrene	ug/L	710 U
Tetrachloroethene	ug/L	710 U
Toluene	ug/L	1000
trans-1,2-Dichloroethene	ug/L	710 U
trans-1,3-Dichloropropene	ug/L	710 U
Trichloroethene	ug/L	710 U
Trichlorofluoromethane	ug/L	710 U
Vinyl chloride	ug/L	710 U
Xylenes (total)	ug/L	450 J

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-4 Analytical Data Summary Site 19 Water PAHs ODOT Innerbelt Study

PARAMETER	UNITS	A6H230315001 19-MW-01 08/22/2006
2-Methylnaphthalene	ug/L	2.6
Acenaphthene	ug/L	0.28
Acenaphthylene	ug/L	0.20 U
Anthracene	ug/L	0.20 U
Benzo(a)anthracene	ug/L	0.20 U
Benzo(a)pyrene	ug/L	0.20 U
Benzo(b)fluoranthene	ug/L	0.20 U
Benzo(ghi)perylene	ug/L	0.20 U
Benzo(k)fluoranthene	ug/L	0.20 U
Chrysene	ug/L	0.20 U
Dibenz(a,h)anthracene	ug/L	0.20 U
Fluoranthene	ug/L	0.20 U
Fluorene	ug/L	0.20 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U
Naphthalene	ug/L	4.2
Phenanthrene	ug/L	0.48
Pyrene	ug/L	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

#### 6.9 SITE 20 - EARL LEE

The Cleveland City Directories indicate the site was listed as Allis Chemical Company from 1954 through 1979. The site was listed as W & W Meats in the 1984 through 1999 Directories.

According to the Cleveland Fire Prevention Bureau files, a Permit dated October 10, 1951 indicates the site was occupied by Allis Chemical Inc., which stored flammable materials in an outside shed. In addition, there was a 1,000-gallon kerosene AST. A Permit dated July 17, 1957 indicates a 4,000-gallon gasoline UST was installed at the site, which was occupied by the Elnor Provision Company.

A BUSTR permit dated January 1989 indicates the 4,000-gallon UST was present. The site was occupied by W & W Meats Incorporated. A Permit dated March 16, 1990 was issued for the removal of the UST. According to the BUSTR files, a NFA Letter was issued for the closure on March 23, 1992.

At the time of the URS Phase I reconnaissance, the site was vacant. No surface staining, stressed vegetation and/or the handling and storage of hazardous materials were observed on the site.

#### 6.9.1 Field Activities

A total of four monitoring wells were proposed for Site 20. Groundwater was encountered in all four soil borings (MW01, MW02, MW03, and MW04) during field activities and monitoring wells were installed. Four soil and four groundwater samples were collected and analyzed for VOCs, SVOCs, TPH and RCRA Metals. A Sample Location Map is included as Figure 6-9A.

### 6.9.2 Site-Specific Geology/Hydrogeology

Soils at Site 20 consisted primarily of sand and sandy clay with minor amounts of silty clay and clay. Brick fragments and hydrocarbon odors were observed in several of the borings. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 20 on September 25, 2006. Localized groundwater flow across Site 20 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 20 is to the west-southwest, towards the Cuyahoga River.

### 6.9.3 Soil Analytical Results

A total of ten VOCs were detected in the four soil samples submitted from Site 20. The chemicals detected in the samples submitted were 2-butanone (ranging from 3.3 ug/kg to 50 ug/kg), acetone (81 ug/kg), benzene (0.59 ug/kg and 2.9 ug/kg), carbon disulfide (ranging from 0.33 ug/kg to 12 ug/kg), cyclohexane (200 ug/kg), ethylbenzene (0.68 ug/kg), isopropylbenzene (ranging from 0.33 ug/kg to 1,800 ug/kg), methylcyclohexane (ranging from 2.4 ug/kg to 890 ug/kg), toluene (0.63 ug/kg and 1.3 ug/kg), and total xylenes (ranging from 2.3 ug/kg to 43 ug/kg). All other VOCs were below the detection limits.

**Phase II Findings** 

A total of nineteen SVOCs were detected in the five samples submitted from Site 20. Concentrations of 2-methylnaphthanene (ranging from 29 ug/kg to 4,300 ug/kg), acenaphthene (ranging from 88 ug/kg to 7,100 ug/kg), acenaphthylene (190 ug/kg), anthracene (ranging from 17 ug/kg to 12,000 ug/kg), benzo(a)anthracene (ranging from 48 ug/kg to 16,000 ug/kg), benzo(a)pyrene (ranging from 51 ug/kg to 13,000 ug/kg), benzo(b)fluoranthene (ranging from 68 ug/kg to 17,000 ug/kg), benzo(g,h,i)perylene (ranging from 36 ug/kg to 7,900 ug/kg), benzo(k)fluoranthene (ranging from 28 ug/kg to 5,500 ug/kg), carbazole (ranging from 95 ug/kg to 4,200 ug/kg), chrysene (ranging from 52 ug/kg to 14,000 ug/kg), dibenz(a,h)anthracene (ranging from 8.8 ug/kg to 2,000 ug/kg), dibenzofuran (510 ug/kg and 5,300 ug/kg), fluoranthene (ranging from 85 ug/kg to 40,000 ug/kg), fluorene (ranging from 11 ug/kg to 6,400 ug/kg), indeno(1,2,3)pyrene (ranging from 34 ug/kg to 6,600 ug/kg), naphthalene (ranging from 23 ug/kg to 7,200 ug/kg), phenanthrene (ranging from 48 ug/kg to 42,000 ug/kg), and pyrene (ranging from 82 ug/kg to 32,000 ug/kg) were detected in the soil samples submitted.

Gasoline and diesel range total petroleum hydrocarbons were detected in the soil samples from Site 20. The highest concentrations were found in the gasoline range, which is consistent with the concentrations of VOCs detected across Site 20. Concentrations of the light petroleum fraction ranged from 240 ug/kg to 280,000 ug/kg. Concentrations of the middle petroleum fraction ranged from 28 mg/kg to 800 mg/kg. Concentrations of the heavy petroleum fraction ranged from 8.4 mg/kg to 1,800 mg/kg.

Seven of the eight RCRA Metals were detected in the soil samples submitted from Site 20. Arsenic, ranging from 5.9 mg/kg to 12.8 mg/kg, was detected in all samples submitted. Barium, ranging from 24.3 mg/kg to 97.9 mg/kg, was detected in all samples submitted. Cadmium, 0.12 mg/kg to 0.59 mg/kg, was detected in all samples submitted. Chromium, ranging from 6.7 mg/kg to 12.2 mg/kg, was detected in the samples submitted. Lead, ranging from 22.9 mg/kg to 446 mg/kg, was detected in the samples submitted. Selenium, 0.72 mg/kg and 0.8 mg/kg, was detected in samples 20-MW02-1012 and 20-MW04-1416. Mercury, ranging from 0.17 mg/kg to 0.54 mg/kg, was detected in all samples submitted.

The analytical results are presented in Table 6-9A.

## 6.9.4 Groundwater Analytical Results

Ten VOCs were detected in groundwater samples collected from Site 20. Concentrations of 1,1-dichloroethene (0.27 ug/L), 2-butanone (ranging from 1.4 ug/L to 5.5 ug/L), acetone (3.2 ug/L), bromodichloromethane (3.1 ug/L), chloroform (15 ug/L), cyclohexane (7.7 ug/L and 150 ug/L), dibromochloromethane (0.42 ug/L), ethylbenzene (1.2 ug/L), isopropylbenzene (0.34 ug/L and 190 ug/L), and methylcyclohexane (19 ug/L and 36 ug/L) were detected in the groundwater samples submitted. All other VOCs were below the detection limits.

Five SVOCs were detected in the groundwater samples submitted from Site 20. Concentrations of acenaphthene (0.38 ug/L), anthracene (0.51 ug/L), benzo(a)anthracene (0.63 ug/L), benzo(b)fluoranthene (0.51 ug/L), and chrysene (0.55 ug/L) were detected in the groundwater samples submitted from 17-MW02 and 17-MW03. All other SVOCs were below the detection limits.

Seven of the eight RCRA Metals were detected in the samples submitted from Site 20. Arsenic, ranging from 23.1 ug/L to 94.8 ug/L, was detected in samples 20-MW03 and 20-MW04. Barium, ranging from 32.8 ug/L to 658 ug/L, was detected in all samples submitted. Cadmium, 0.85 ug/L, was detected in sample 20-MW03. Chromium, ranging from 3.1 ug/L to 32 ug/L, was detected in all the samples submitted. Lead, 4.4 ug/L and 522 ug/L, was detected in 20-MW01 and 20-MW03. Selenium, 2.6 ug/L, was detected in sample 20-MW01. Mercury, 0.54 ug/L, was detected in 20-MW03.

The analytical results are presented in Table 6-9B.

#### 6.9.5 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on Tables 6-9A and 6-9B.

None of the VOCs or TPH detected in the soils at Site 20 exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or BUSTR closure action levels.

The concentrations of benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, and dibenz(a,h)anthracene detected in soil sample 20-MW03-1012 exceeded the OEPA VAP standards for commercial and industrial land use and/or BUSTR closure action levels.

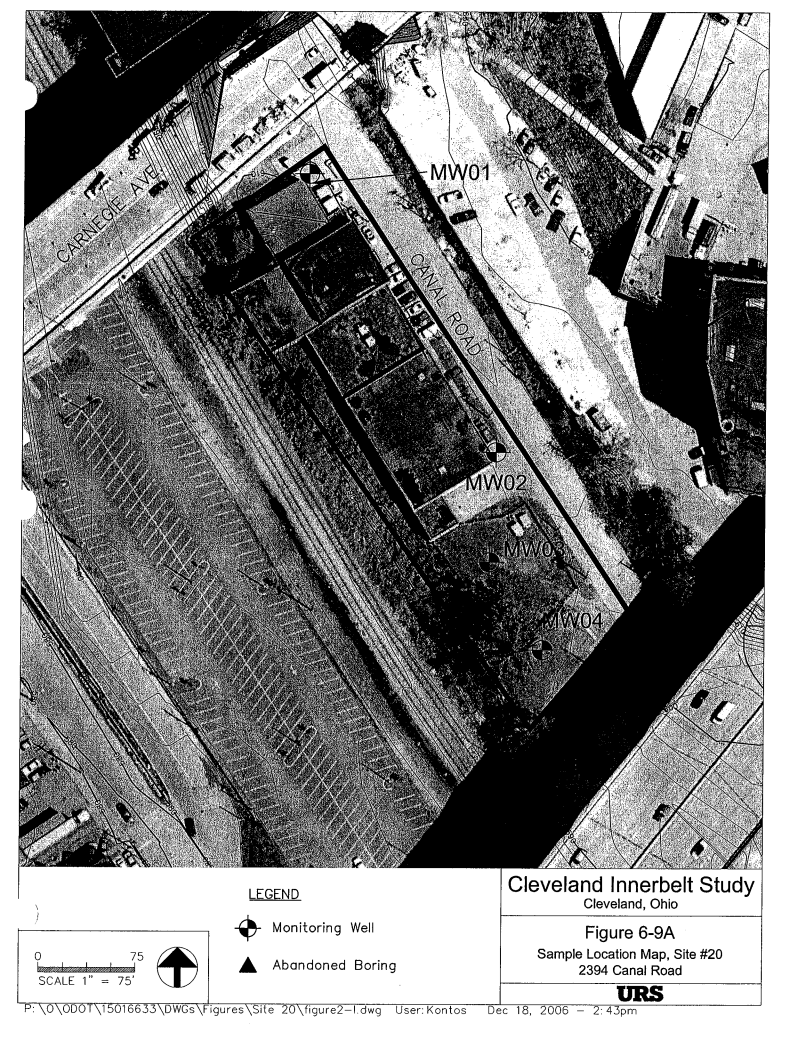
The concentration of lead detected in soil sample 20-MW03-1012 exceeded the OEPA VAP standards for commercial and industrial land use and the construction and excavation worker activities category action levels.

The concentrations of benzo(a)anthracene, benzo(b)fluoranthene, and lead detected in groundwater sample 20-MW03 exceeded the OEPA VAP standard for generic unrestricted potable use standards and/or BUSTR closure action levels.

#### 6.9.6 Conclusions

Based on the analytical results, the soil and groundwater at Site 20 may require special disposal and/or worker protection protocols (plan note) during construction activities.

**FIGURES** 



**TABLES** 

# Summary of Detected Chemicals in Soil Site 20 - Earl Lee Table

**ODOT Innerbelt Study** Cleveland, Ohio

2-Buttenere         up/96         71,000,000         0.00,000         - 60         4 24 1         4 24 1         4 4 1         4 23 1         4 64 1           Percheric districte         up/96         10,000         210,000         149 0         17,000	Δ.	PARAMETER	UNITS	VAP Commercial/ Findustrial Standard <sup>1)</sup>	VAP Construction Worker Standard	Elosure Action Crosure Action Crovel	20-MW01-1618 07/18/2006	20-MW02-1012 07/18/2006	20-MW03-1012 07/18/2006	20-MW04-1416 08/11/2006
Active the particular properties of the particular		2-Butanone	ug/kg	71,600,000	80,000,000		20	47 J	3.3 J	46 J
Percentage   1979   100,000   110,000   1149   618 J   61 U   61 U		Acetone	ug/kg	100,000,000	100,000,000	National and the control of the cont	27 U	240 U	24 U	. E
Carbon distillation         uging         720,000         720,000         —         51,1         61,1         0.33 J           Entrylentrane         uging         220,000         220,000         45,500         68,1         60 J         12 U           Entrylentrane         uging         220,000         220,000         45,500         68,1         0.0         12 U         2.4 J           Entrylentrane         uging         520,000         520,000         4,10         13 J         60 U         0.63 J           Animation         uging         520,000         520,000         4,10         13 J         60 U         0.63 J           Animation         uging         520,000         520,000         4,10         13 J         60 U         0.63 J           Animation         uging         520,000         50,000,000         15,700         2.4 J         170 U         2.3 J         400 U         0.63 J           Emraciol prime         uging         56,000         11,000         130 J         170 J         120 U		Benzene	ug/kg	100,000	310,000	149	0.59 J	n 09	6.1 U	2.9 J
Chylchesane         Ug/kg         2.0.0         2.0.0         4.5.0         14 U         12 U         12 U           Chylchesane         Ug/kg         2.00,000         2.0.0         4.5.0         6.5.1         6.0 U         0.68 J           Beopropytherane         Ug/kg         2.00,000         2.0.0         4.5.00         4.5.0         6.0 U         0.0.8         1           Reportorytherane         Ug/kg         2.0.0         2.0.0         2.0.0         1.3 J         6.0 U         0.0.8         1           Administration         Ug/kg         1.0.0         1.0.0         1.1.0         2.4 J         1.2 U         2.3 J           Administration         Ug/kg         1.0.0         1.0.0         1.1.0         2.0.0         2.0.0           Administration         Ug/kg         1.0.0         1.1.0         2.0.0         2.0.0         2.0.0           Administration         Ug/kg         6.3.00         810,000         1.1.00         2.0.0         2.0.0         1.2.0         2.0.0           Bearzo(plinoranthene         Ug/kg         6.3.00         810,000         1.1.00         2.0.0         2.0.0         2.0.0         2.0.0         2.0.0         2.0.0         2.0.0         2.0.0		Carbon disulfide	ug/kg	720,000	720,000		5.1 J	6.1	0.33 J	12 J
Exception   September   Sept	s)(	Cyclohexane	ug/kg	-	The state of the s		14 U	120 U	12 U	200
Newtykerscripters   Ug/kg   S80,000   S60,000   S60,000   S70,000   S70   S70   S74   S70   S7	٥Λ	Ethylbenzene	ug/kg	230,000	230,000	45,500	0.8.0	∩ 09	0.68 J	n 09
Math/Accideration   Ug/kg   S20,000   14,1   13,1   100   0.63 J     Administration   Ug/kg   S20,000   150,000   14,1   150   10   0.63 J     Administration   Ug/kg   S20,000,000   150,000   1,10		Isopropylbenzene	ug/kg	860,000	860,000	er teller i de en de teller de en de en de en de en de en de en de en de en de en de en de en de en de en de e	0.53 J	8.8	0.33 J	1800
Tolleane         Ug/kg         \$20,000         \$20,000         \$20,000         13 J         60 U         0.68 J           Yelenes (total)         ug/kg         160,000         15,700         15,700         24 J         70 U         22 J           Zehen/indathriane         ug/kg         160,000         550,000,000         550,000,000         100         29 J         400 U         2000         1           Acentaphthylene         ug/kg         860,000,000         11,000         2000         1         100         2000         1           Anthracene         ug/kg         860,000,000         11,000         2000         400 U         20000 U         1           Bennzo(a)priene         ug/kg         630,000         810,000         11,000         230         44 J         1000 U           Bennzo(a)priene         ug/kg         630,000         810,000         11,000         280 J         48 J         1000 J           Bennzo(a)priene         ug/kg         630,000         810,000         11,000         280 J         400 J         2000 J           Bennzo(b)fulcoranthene         ug/kg         630,000         810,000         11,000         220 J         400 J         1000 J           Christani		Methylcyclohexane	ug/kg		. No. 1950 Commented and the comment of the comment	to and the control of	14 U	210	2.4 J	068
Xidense (total)         up/kg         160,000         157,00         2.4 J         120 U         2.3 J           Ackenspfritaelne         ug/kg         180,000,000         530,000         —         860 J         2.9 J         4500 J           Ackenspfritaelne         ug/kg         880,000         1,000,000,000         —         190 J         400 U         20000 U           Ackenspfritaelne         ug/kg         880,000         1,000,000,000         1,100         48 J         15000 J           Benzo(sjamtracene         ug/kg         880,000         810,000         1,100         1800         48 J         15000 J           Benzo(sjamtracene         ug/kg         630,00         810,000         1,100         1800         51 J         1000 J           Benzo(sjamtracene         ug/kg         630,00         810,000         1,100         180         55 J         17000 J           Benzo(sjamtracene         ug/kg         630,00         8100,000         1,100         220         22 J         17000 J           Benzo(sjamtracene         ug/kg         670,000         1,100         220         22 J         1400 J           Grafazzole         ug/kg         6,700,000         1,1000         220         2		Toluene	ug/kg	520,000	520,000	49,100	1.3 J	n 09	0.63 J	n 09
2-Mathylapithalene         Ug/kg         16.000,000         —         —         866 J         29 J         4400 J         7700 J           Acentaphthalene         Ug/kg         16.000,000         1.000,000         —         460 J         7700 J         7700 J           Achthalene         Ug/kg         653,000         110,000         2200 J         460 J         2700 J         4700 J         47		Xylenes (total)	ug/kg	160,000	160,000	15,700	2.4 J	120 U	2.3 J	43 J
Acentaphthene         ug/kg         190,000,000         550,000,000		2-Methylnaphthalene	ug/kg	:	-	1	F 098	29 J	4300 J	87 J
Accomply this land         Accomply by graph of the control of		Acenaphthene	ug/kg	180,000,000	530,000,000	VIII COMPANIAN AND AND AND AND AND AND AND AND AND A	410 J	400 U	7100 J	S 8 9
Anthracene         Anthracene         Jug/49         880,000,000         1,000         2300         17,1         12000 J           Benzo(a)pyrane         ug/49         65,000         81,000         11,000         2300         81,0         13000 J           Benzo(a)pyrane         ug/49         65,000         81,000         11,000         22100         86 J         77000 J           Benzo(a)pyrane         ug/49         65,000         81,000,000         11,000         38 J         7300 J           Benzo(a)pyrane         ug/49         65,000         81,000,000         11,000         38 J         7300 J           Chysene         ug/40         6,700,000         81,000,000         1,100         38 J         7300 J           Chysene         ug/40         6,700,000         41,000         1,100         38 J         7300 J           Dibenzoluran         ug/40         6,700,000         1,100,000         2200         38 J         7300 J           Dibenzoluran         ug/40         6,700,000         1,100,000         1,100         38 J         7300 J           Fluoranthene         ug/40         6,700,000         1,100,000         1,100,000         34 J         400 J         1,100 J		Acenaphthylene	ug/kg		Administration of the steel steel and the steel	ee Corporited And Training and Control (1990)	190 1	400 U	20000 U	1600 U
Benzo(a)authracene         ug/kg         65.000         810,000         11,000         2300         46         1 15000         1           Benzo(a)authracene         ug/kg         65.300         810,000         11,000         2300         66 J         17000 J           Benzo(din)perviene         ug/kg         65.300         810,000         11,000         86 J         17000 J           Benzo(fill)perviene         ug/kg         650,000         8,100,000         11,000         2200         82 J         7900 J           Chysene         ug/kg         6,700,000         41,000,000         1,100,000         2200         220         220 J         420 J         42000 J           Chysene         ug/kg         6,700,000         41,000         1,100         220         220         220         220         220         2200 J         4200 J         42000 J         14000 J         42000 J         14000 J         1100         1100         1100         1100         1100 J         1100         1100         1100         1100         1100         1100         1100         1100         1100         1100         1100         1100         1110         1110         1111         1110         1110         1111 <td< td=""><td></td><td>Anthracene</td><td>ug/kg</td><td>880,000,000</td><td>1,000,000,000</td><td></td><td>840 J</td><td>17 J</td><td>12000 ל</td><td>f 02</td></td<>		Anthracene	ug/kg	880,000,000	1,000,000,000		840 J	17 J	12000 ל	f 02
Benzo(a)pyrene         ug/kg         6.300         81,000         1,100         1800         51 J         13000 J           Benzo(a)pyrene         ug/kg         63,000         81,000         11,000         890 J         36 J         77000 J           Benzo(k)fluoranthene         ug/kg         630,000         81,00,000         11,000         380 J         38 J         7800 J           Carbazole         ug/kg         6,700,000         41,000,000         1,100,000         2200         220         420 J         4200 J           Chysene         ug/kg         6,700,000         41,000,000         1,100         220         220         1,400 J         4200 J           Dibenzofuran         ug/kg         6,700         41,000,000          510 J         400 U         5300 J           Dibenzofuran         ug/kg         6,700         41,000         1,100         230 J         400 U         5300 J           Dibenzofuran         ug/kg         33,000,000         170,000,000          510 J         400 U         5300 J           Fluoranthene         ug/kg         55,000         11,000,000          510 U         5300 J           Fluoranthene         ug/kg <t< td=""><td></td><td>Benzo(a)anthracene</td><td>ug/kg</td><td>63,000</td><td>810,000</td><td>11,000</td><td>2300</td><td>48 J</td><td>16000 J</td><td>130 J</td></t<>		Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	2300	48 J	16000 J	130 J
Benzo(b)fluoranthene         Ug/kg         65,000         810,000         11,000         2100         66 J         17000 J           Benzo(b)fluoranthene         Ug/kg         650,000         810,000         110,000         36 J         7800 J         7800 J           Benzo(k)fluoranthene         Ug/kg         650,000         81,000,000         11,00,000         2200         52 J         4200 J         4200 J           Chysene         Ug/kg         6,700         41,000,000         1,100,000         2200         52 J         4200 J         4200 J           Dibenzor(al.n)anthracene         Ug/kg         6,700         41,000,000         1,100,000         220 J         88 J         2000 J           Dibenzor(al.n)anthracene         Ug/kg         6,700         41,000,000         1,100,000         220 J         88 J         2000 J           Dibenzor(al.n)anthracene         Ug/kg         120,000,000         1,100,000         1,11         4000 U         5500 J           Fluoranthracene         Ug/kg         120,000         1,100,000         1,11         4200 U         1,200 U           Fluoranthracene         Ug/kg         120,000         1,100,000         1,11         4200 U         1,200 U           Pharanthracene		Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	1800	51 J	13000 J	110 J
Benzo(ghil)penylene         ug/kg           890 J         36 J         7900 J           Carbaze(khiloranthene         ug/kg         630,000         8,100,000         110,000         2200         52 J         4200 J           Chysene         ug/kg         (7,700,000)         41,000,000         1,100,000         2200         52 J         14000 J           Dibenz(a,h)anthracene         ug/kg         (7,700,000)         177,000,000         1,100,000         2200         52 J         14000 J           Dibenz(a,h)anthracene         ug/kg         (7,700,000)         177,000,000         1,100,000         220 J         8.8 J         2000 J           Dibenz(a,b)anthracene         ug/kg         (7,700)         41,000         1,100,000         220 J         11 J         6600 J           Fucrene         ug/kg         120,000,000         1,500,000         230 J         11 J         6600 J         1,200 J           Naphthalene         ug/kg         550,000         1,500,000         230 J         12 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J         1,20 J		Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	2100	F 89	17000 J	130 J
Benzol(yflucranthene         ug/kg         630,000         8,100,000         110,000         990 J         28 J         5500 J           Cardazole         ug/kg         1,000,000         31,000,000         1,100         2200         52 J         14000 J         14000 J           Chrysene         ug/kg         6,700         41,000,000         1,100         2200 J         5200 J         14000 J         5300 J           Dibenzoluran         ug/kg         33,000,000         1,70,000,000         -         -         5100 J         400 U         5300 J           Fluorente         ug/kg         33,000,000         340,000,000         -         -         5100 J         4000 U         5300 J           Fluorente         ug/kg         122,000,000         340,000,000         -         -         4000 U         48 J         4000 U           Fluorente         ug/kg         125,000,000         190,000         11,000         220 J         11 J         620 J         11 J         6400 J           Phenauthrene         ug/kg         550,000         130,000,000         -         -         11 J         4200 J         120 J		Benzo(ghi)perylene	ug/kg		The state of the s	THE THE PROPERTY OF THE PROPER	F 068	36 J	ر 2900	76 J
Carbazole         ug/kg         (1,000,000         31,000,000         1,100,000         2200         52 J         400 U         4200 J           Chrysene         ug/kg         6,700         41,000,000         1,100,000         2200         52 J         14000 J           Diberacta,hanthracene         ug/kg	sე	Benzo(k)fluoranthene	ug/kg	630,000	8,100,000	110,000	P 066	28 J	5500 J	95 J
Chrysene         Ug/kg         6,700,000         41,000,000         1,100         220         52 J         14000 J           Dibenz(a,h)anthracene         Ug/kg         6,700         41,000         1,100         230 J         8.8 J         2000 J           Fluctenzofuran         Ug/kg         6,700         170,000,000         -         620 J         111         6400 J           Fluctene         Ug/kg         120,000,000         340,000         -         620 J         111         6400 J           Fluctene         Ug/kg         67,000         1,900,000         -         620 J         11         6400 J           Phenanthrene         Ug/kg         530,000         1,900,000         38,000         790 J         23 J         7200 J           Phenanthrene         Ug/kg         55,000         19,000,000         10,000         33,000         33,000         1200 J           Phenanthrene         Ug/kg         55,000,000         130,000,000         -         20,00         48 J         7200 J           Phenanthrene         Ug/kg         25,000,000         130,000,000         -         1000         82 J         33,000           Gasoline Range Organics (CG-C12)         Ug/kg         25,000,000	٥٨	Carbazole	ug/kg	10,000,000	31,000,000	●●	430 J	400 U	4200 J	. 65
Dibenz(a,h)anthracene         ug/kg         6,700         41,000         1,100         230 J         8.8 J         2000 J           Plucenz(uan persoduran problemsoduran problemsoduran problemsodurant probl	s	Chrysene	ug/kg	6,700,000	41,000,000	1,100,000	2200	52 J	14000 J	130 J
Dibenzofuran         ug/kg         3.3000,000         170,000,000         -         510 J         400 U         5300 J           Fluoranthane         ug/kg         120,000,000         340,000,000         -         5100         85 J         40000           Fluoranthane         ug/kg         120,000,000         1,000,000         39,800         33 J         6400 J           Indenof (12,3-cd)pyrane         ug/kg         67,000         1,900,000         39,800         48 J         7200 J           Phenanthrane         ug/kg         25,000,000         130,000,000         -         4000         82 J         3200           Phenanthrane         ug/kg         25,000,000         130,000,000         -         4000         82 J         3200           Phenanthrane         ug/kg         25,000,000         130,000,000         -         4000         82 J         3200           Gasoline Range Organics (Ce-C12)         ug/kg         -         -         1,000,000         240         180 J         28           C10-C20         mg/kg         80         -         -         5,000         120 U         12.8           Barium         mg/kg         8,900         2,000         -         -		Dibenz(a,h)anthracene	ug/kg	6,700	41,000	1,100	230 J	8.8	2000 J	1600 U
Flucranthene         ug/kg         33,000,000         170,000,000         5100         85 J         40000           Flucrene         ug/kg         120,000,000         340,000,000         11,000         850 J         34 J         6400 J           Indenof (12.3-cd)pyrene         ug/kg         67,000         410,000         11,000         820 J         34 J         6600 J           Naphthalene         ug/kg         530,000         1,900,000         39,800         780 J         48 J         4200 J           Phenanthrene         ug/kg         25,000,000         130,000,000         -         4000         82 J         3200 J           Gascline Range Organics (C6-C12)         ug/kg         -         -         4000         82 J         3200 J           Gascline Range Organics (C6-C12)         ug/kg         -         -         1,000,000         240         180 J         84         12.8         100           C20-C24         mg/kg         200,000         45,000         -         5,000         111         59         12.8         12.8           Barlum         mg/kg         770         420         -         -         6.7         9.45         6.7         9.8           Chomium		Dibenzofuran	ug/kg	1		***	510 J	400 U	5300 J	1600 U
Fluorene         ug/kg         120,000,000         340,000,000         11,000         820 J         11 J         6400 J           Indenof12.3-cd)pyrene         ug/kg         57,000         410,000         39,800         790 J         23 J         7200 J           Phenanthrene         ug/kg         530,000         1,900,000         39,800         48 J         4200           Phenanthrene         ug/kg         25,000,000         130,000,000         -         4000         82 J         7200 J           Gasolline Range Organics (C6-C12)         ug/kg         25,000,000         130,000,000         240         180         28         100           C20-C20         ng/kg         -         -         2,000         300         240         12.8         100           C20-C30         ng/kg         200,000         45,000         -         5,000         11.0         5.9         12.8         100           Arsenic         mg/kg         200,000         45,000         -         94.5         24.3         9.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8         12.8		Fluoranthene	ug/kg	33,000,000	170,000,000		5100	85 J	40000	330 J
Indenof 12.3-cd)pyrene         ug/kg         67,000         410,000         11,000         820 J         34 J         6600 J           Naphthalene         Ug/kg         530,000         1,900,000         39,800         730 J         23 J         7200 J           Phenanthrane         Ug/kg         25,000,000         130,000,000         -         4200         48 J         42000           Pyrene         Ug/kg         25,000,000         130,000,000         -         4000         82 J         32000           Gasoline Range Organics (C6-C12)         Ug/kg         25,000,000         1,000,000         240         160000         120 U           C10-C20         C10-C20         300         240         160000         120 U         120 U           C10-C20         Msykg         80         210         -         5,000         11.1         5.9         12.8           Barium         mg/kg         200,000         45,000         -         11.1         5.9         446           Cadmium         mg/kg         8.900         2,000         -         12.2         6.7         9.8           Lead         mg/kg         15,000         -         -         -         9.67         9.7 <td></td> <td>Fluorene</td> <td>ug/kg</td> <td>120,000,000</td> <td>340,000,000</td> <td>earth, occupation on the department of the control</td> <td>620 J</td> <td>7 =</td> <td>6400 J</td> <td>. f 26</td>		Fluorene	ug/kg	120,000,000	340,000,000	earth, occupation on the department of the control	620 J	7 =	6400 J	. f 26
Naphthalene         Ug/kg         530,000         1,900,000         39,800         790 J         23 J         7200 J           Phenanthrene         Ug/kg           4200         48 J         42000           Pyrene         Ug/kg         25,000,000         130,000,000          4000         82 J         32000           Gasoline Range Organics (C6-C12)         Ug/kg           1,000,000         240         160000         120 U           C10-C20         mg/kg           5,000         11,1         5.9         12.8           Arsenic         mg/kg         200,000         45,000          94.5         24.3         97.9           Cadmium         mg/kg         200,000         45,000          94.5         24.3         97.9           Chromium         mg/kg         8,900         2,000           94.5         6.7         9.8           Lead         mg/kg         15,000          0.51         0.12 J         0.68 U         0.72 J         9.8           Lead         mg/kg         15,000           0.68 U         0.77 D		Indeno(1,2,3-cd)pyrene	ug/kg	67,000	410,000	11,000	820 J	34 J	r 0099	7 99
Phenanthrene         Ug/kg         25,000,000         130,000,000         -         4200         48 J         42000           Pyrene         Ug/kg         25,000,000         130,000,000         -         4000         82 J         32000           Gasoline Range Organics (C6-C12)         Ug/kg         -         -         -         2,000         300         28         100           C10-C20         C20-C34         mg/kg         -         -         -         2,000         8.4         580         100           Arsenic         mg/kg         200,000         45,000         -         94.5         24.3         97.9           Cadmium         mg/kg         770         420         -         6.7         9.8           Lead         mg/kg         8,900         2,000         -         94.5         6.7         9.8           Lead         mg/kg         15,000         4,300         -         -         9.5         6.7         9.8           Lead         mg/kg         15,000         4,300         -         -         9.5         9.8         9.6           Mercury         mg/kg         15,000         -         -         9.5         9.5		Naphthalene	ug/kg	530,000	1,900,000	39,800	ر 290 J	23 J	7200 J	1600 U
Pyrene         Ug/kg         25,000,000         130,000,000         —         4000         82 J         32000           Gasoline Range Organics (C6-C12)         Ug/kg         —         —         —         2,000         240         160000         120 U         1           C10-C20         C20-C34         mg/kg         —         —         —         2,000         1800         84         580         100           Arsenic         mg/kg         200,000         45,000         —         94.5         24.3         97.9         97.9           Cadmium         mg/kg         770         420         —         0.51         0.12 J         0.59         77.9         9.8           Lead         mg/kg         15,000         4,300         —         —         94.5         24.3         97.9         97.9           Chromium         mg/kg         8,900         2,000         —         —         122 G.7         6.7         9.8           Lead         mg/kg         15,000         4,300         —         —         9.5         9.6         9.8           Mercury         mg/kg         15,000         —         —         9.5         6.7         9.8		Phenanthrene	ug/kg	er den en elle det i Report de de la compressión		•	4200	48 J	42000	300 1
Gasoline Range Organics (C6-C12)         ug/kg         -         -         1,000,000         240         160000         120 U           C10-C20         C10-C20         300         28         100           C20-C34         mg/kg         -         -         -         5,000         180         8.4         580           Arsenic         mg/kg         80         210         -         94.5         5.9         12.8           Barium         mg/kg         200,000         45,000         -         94.5         24.3         97.9           Cadmium         mg/kg         770         420         -         0.51         0.12 J         0.59           Chromium         mg/kg         8,900         2,000         -         94.5         6.7         9.8           Lead         mg/kg         15,000         4,300         -         -         9.67         9.8           Selenium         mg/kg         15,000         4,300         -         0.68 U         0.77         0.61 U           Mercury         mg/kg         300         84         -         0.54         0.076 J         0.17		Pyrene	ug/kg	25,000,000	130,000,000	1	4000	82 J	32000	250 J
Cito-C20         mg/kg          2,000         300         28         100           C20-C34         mg/kg          5,000         1800         8.4         580           Arsenic         mg/kg         200,000         45,000          94.5         24.3         97.9           Barium         mg/kg         770         420          94.5         24.3         97.9           Chromium         mg/kg         8,900         2,000          12.2         6.7         9.8           Lead         mg/kg         15,000         4,300          176         22.9         446           Selenium         mg/kg         15,000         4,300          0,68 U         0,772         0,61 U           Mercury         mg/kg         300         84          0,54         0,076 J         0,177	Н	Gasoline Range Organics (C6-C12)	ug/kg	Produktioner of a substitute of statety stockage	Charles all a Activities and the Landsconnection of the Control of	1,000,000	240	160000	120 U	280000
C20-C34         mg/kg           5,000         1800         8.4         580           Arsenic         mg/kg         80         210          94.5         24.3         97.9           Barium         mg/kg         770         45,000          94.5         24.3         97.9           Cadmium         mg/kg         770         420          94.5         0.59         6.7         9.8           Lead         mg/kg         8.900         2,000          12.2         6.7         9.8           Selenium         mg/kg         15,000         4,300          0.68 U         0.72         0.61 U           Mercury         mg/kg         300         84          0.54         0.076 J         0.17	ďΤ	C10-C20	mg/kg	***		2,000	300	28	\$	008
Arsenic         mg/kg         80         210         -         11.1         5.9         12.8         12.8           Barlum         mg/kg         200,000         45,000         -         94.5         24.3         97.9         97.9           Cadmium         mg/kg         770         420         -         0.51         0.12 J         0.59         0.59           Chromium         mg/kg         8,900         2,000         -         12.2         6.7         9.8           Lead         mg/kg         15,000         4,300         -         0.68 U         0.72         0.61 U           Mercury         mg/kg         300         84         -         0.54         0.076 J         0.17		C20-C34	mg/kg	1		5,000	1800	8.4	580	140
Barlum         mg/kg         200,000         45,000         94.5         24.3         97.9           Cadmium         mg/kg         770         420         -         0.51         0.12 J         0.59           Chromium         mg/kg         8,900         2,000         -         12.2         6.7         9.8           Lead         mg/kg         15,000         4,300         -         0.68 U         0.72         0.61 U           Mercury         mg/kg         300         84         -         0.54         0.076 J         0.17		Arsenic	mg/kg	80	210	-	11.1	5.9	12.8	11.7
Cadmium         mg/kg         770         420          0.51         0.12 J         0.59           Chromium         mg/kg         8,900         2,000          12.2         6.7         9.8           Lead         mg/kg         15,000         4,300          0.68 U         0.72         0.61 U           Mercury         mg/kg         300         84         0.076 J         0.076 J         0.17	- 9	Barlum	mg/kg	200,000	45,000		94.5	24.3	97.9	43
Chromium         mg/kg         8,900         2,000         -         12.2         6.7         9.8           Lead         mg/kg         -         176         22.9         446           Selenium         mg/kg         15,000         4,300         -         0.68 U         0.72         0.61 U           Mercury         mg/kg         300         84         0.076 J         0.076 J         0.17	ΙΓS	Cadmium	mg/kg	440	420	•	0.51	0.12 J	0.59	0.29
Lead         mg/kg          176         22.9         446           Selenium         mg/kg         15,000         4,300          0.68 U         0.72         0.61 U           Mercury         mg/kg         300         84          0.54         0.076 J         0.17	/TE	Chromium	mg/kg	8,900	2,000	1	12.2	6.7	8.6	8.6
n mg/kg 300 84 - 0.54 0.076 J 0.17	IM	Lead	mg/kg	•		Windows on Section 1111 Sections	176	22.9	446	43.7
mg/kg 300 84 0.54 0.076 J 0.17		Selenium	mg/kg	15,000	4,300		0.68 U	0.72	0.61 U	0.8
		Mercury	mg/kg	300	84	· contract of the season of th	0.54	0.076 J	0.17	0.2

<sup>-- =</sup> Standard not available
U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.
J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial/ Industrial Land Use

<sup>(2)</sup> VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Table-6-9B
Summary of Detected Chemicals in Water
Site 20 - Earl Lee
ODOT Innerbelt Study
Cleveland, Ohio

o.	Parameter	UNITS	VAPJUPUS/ RECPUS <sup>(1)</sup>	Closura Action	-20-MW-01 09/22/2006	20-MW-02 09/25/2006	20-MW-03 09/25/2006	, 20-MW-04 09/25/2006
	1,1-Dichloroethene	T/6n	7	-	1.0 U	1.0 U	0.27 J	5.0 U
	2-Butanone	ug/L	6800	1	10 U	1.4 J	5.5 J	4.9 J
	Acetone	ug/L	1600	The section of the se	000 University of the Company of the	10 U	3.2 J	50 0
		ug/L	1	A real Video del logicolo in accordina del propositiona d	3.1	1.0 U	1.0 U	5.0 11
sე(		ng/L	50	And the second s	13	1.0 U	1.0 U	5 0 1
	Cyclohexane	ug/L	The state of the s	And the first of the control of the	1.0 U	1.0 U	7.7	150
	Dibromochloromethane	ug/L	19		0.42 ט	1.0 U	1.0 U	5.0 11
	Ethylbenzene	ug/L	700	200	1.0 U	1.0 U	) O.T	1.2.1
	Isopropylbenzene	ng/L	1300	Special Company of the Company of th	1.0 U	1.0 U	0.34 J	190
	Methylcyclohexane	ng/L	ì	•-	1.0 U	1.0 U	19	36
	Acenaphthene	ug/L	680	1	0.20 U	0.20 U	0.38	0.80
so	- 1	ug/L	2600		0.20 U	0.20 U	0.51	0.80
OΛ		ug/L		0.264	0.20 U	0.20 U	0.63	0.80
S	ŝ	ug/L	The state of the s	0.179	0.20 U	0.20 U	0.51	0.80 U
	Chrysene	ug/L	47	47	0.20 U	0.20 U	0.55	0.80
	Arsenic	ug/L	20	1	10.0 U	10.0 U	23.1	94.8
	Barium	ug/L	2000	•	32.8 J	622	287	929
STY	Cadmium	ug/L	S	COR Patrick City 140 districts for the control of t	2.0 U	2.0 U	0.85	2.0 U
/T3	Chromium	ng/L	100		11.8	3.1 J	32	4.0 ک
M	Lead	ug/L	15	The second secon	4.4	3.0 U	522	3.0 U
	Selenium	ng/L	50	-	2.6 J	5.0 U	5.0 U	5.0 U
	Mercury	ng/L	2		0.20 U	0.20 U	0.54	0.20

<sup>-- =</sup> Standard not available

 $<sup>\</sup>mathsf{U} = \mathsf{The}$  analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

<sup>(1)</sup> VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

BORING LOGS

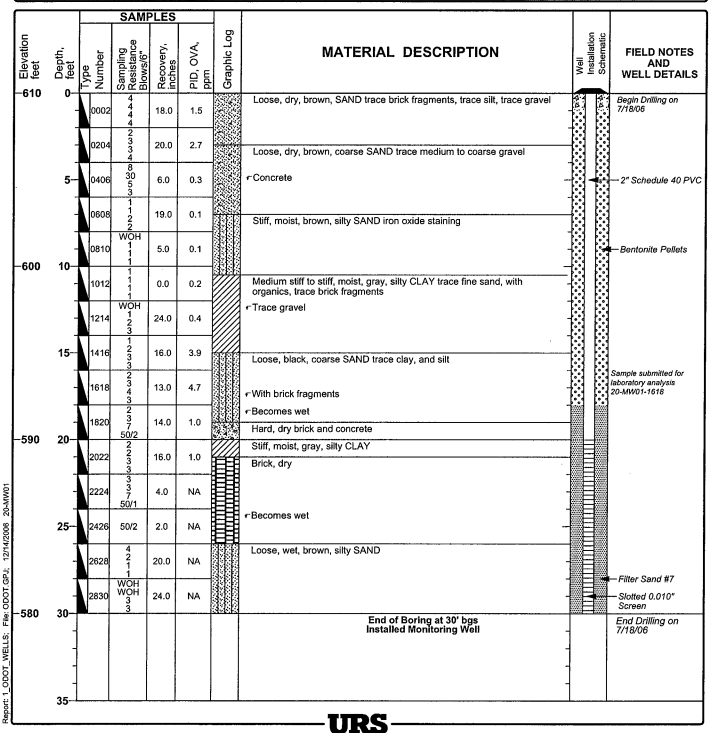
**Project: ODOT - Innerbelt Corridor** 

Project Location: Site 20
Project Number: 15016633

### Log of Boring 20-MW01

Sheet 1 of 1

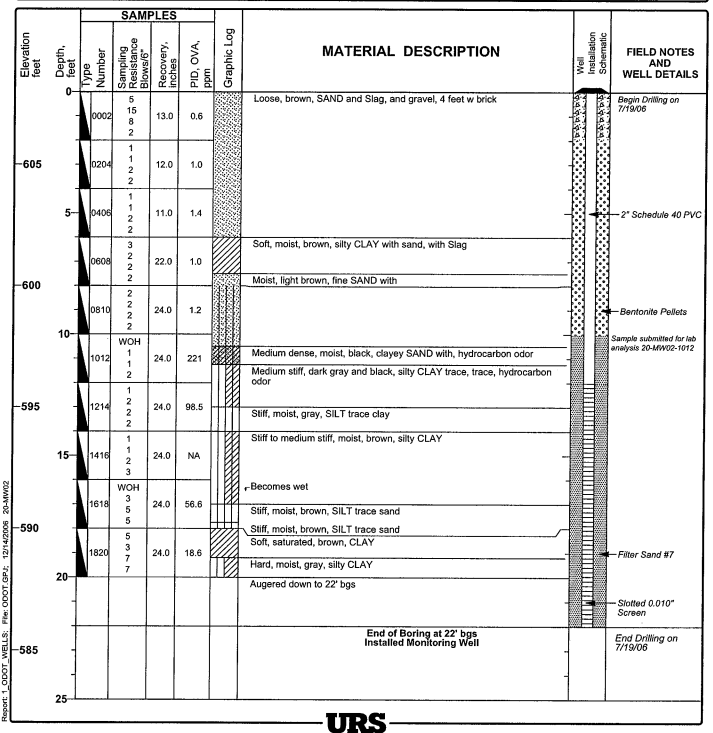
Date(s) 7/18/06 Drilled 7/18/06	Logged By	J. Kaminski	Checked M. Wolff By
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# / 30" drop automatic	Total Depth of Borehole 30.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 610'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 586.72 on 9/25/06			



Project Location: Site 20 Project Number: 15016633

## Log of Boring 20-MW02

Date(s) 7/19/06 Drilled 7/19/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140 lb Hammer 30 inches	Total Depth of Borehole 22.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 608'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Set monitoring well
Groundwater Level and Date Measured 596.08 on 9/25/06			

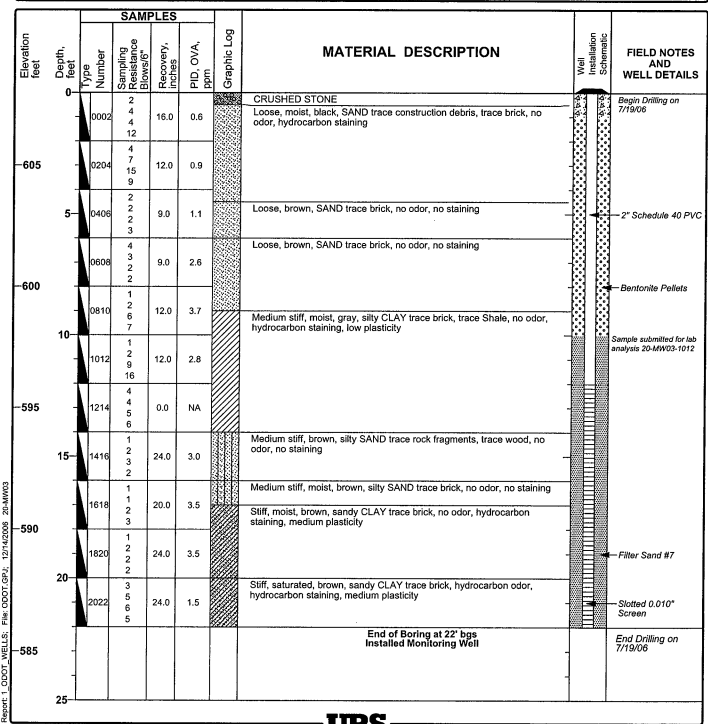


**Project Location: Site 20 Project Number:** 15016633

ODOT

### Log of Boring 20-MW03

Date(s) 7/19/06 Drilled 7/19/06	Logged By	X. Sotelo	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140 lb Hammer 30 inches	Total Depth of Borehole 22.0′ bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 608'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Set monitoring well
Groundwater Level and Date Measured 589.02 on 9/25/06			



Project Location: Site 20 Project Number: 15016633

## Log of Boring 20-MW04

Date(s) 8/11/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 24.0′ bgs
Drill Rig Type <b>LC-60</b>	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 604'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 588.57 on 9/25/06			

1				SAM	PLES		ĺ		1		
	Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"			Graphic Log	MATERIAL DESCRIPTION	Well	Schematic	FIELD NOTES AND WELL DETAILS
		-	0002	4	12.0	0.7		Loose, dry, dark brown, SAND with gravel, with brick fragments, and Slag	4.	4.4.4.4.	Begin Drilling on 8/11/06
	600		0204	5 20 20 20	19.0	1.5		Hard, dry, brown, silty CLAY with sand, with brick fragments, and gravel			
	000	5-	0406	50/5	14.0	2.1	* 4	CONCRETE Loose, dry, brown, coarse SAND with gravel, with Slag			2" Schedule 40 PVC
		-	0608	6	22.0	2.0		Fine to medium SAND			
,	595	10-	0810	5	20.0	1.6		Loose, dry, black and dark brown, SAND with gravel, with Slag			Bentonite Pellets
		-	1012	3	4.0	1.2					
	-590	-	1214	7 5 7 10	13.0	192		Dense, dry, brown, fine SAND  -Moist to wet, black, hydrocarbon odor			Sample submitted for lab
		15-	1416	7	20.0	>10000	TVZ	Dense, dry, brown, fine SAND  -Moist, gray, silty CLAY 3-Inch layer  -Trace clay, with gravel			analysis 20-MW04-1416
		-	1618	WOH 1 2 3 WOH	12.0	NA		Stiff, wet, gray and brown, silty CLAY trace sand, trace, iron oxide staining			
20-MW04	-585	- 20-	1820	2 4 4 WOH	12.0	NA		r-Hard, dry, no sand			−Filter Sand #7
12/14/2006 20-		-	2022	2 2 6 WOH	10.0	NA		Hard, dry, brown, clayey SILT			Slotted 0.010" Screen
GPJ;	-580	-	2224	2 4 7	11.0	NA					End Dvilling on
LS; File: ODOT		25-		·		į		End of Boring at 24' bgs Installed Monitoring Well			End Drilling on 8/11/06
Report: 1_ODOT_WELLS;	-575	-				į		- -			
Report: 1		30						URS		_	

DATA ASSESSMENT REPORT

#### Data Assessment Report ODOT Innerbelt Study Site 20 – Earl Lee

Reviewer: P. Schuler Date: November 12, 2006

Four soil samples and four groundwater samples were collected at the Earl Lee site at 2394 Canal Road in Cleveland, Ohio, from July 18 through September 25, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Re	Requested Analyses <sup>(1)</sup>			
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC	TPH	Met	
A6G190357001	20-MW01-1618	07/18/2006	Soil	X	X	X	X	
A6G190357002	20-MW02-1012	07/18/2006	Soil	X	X	X	X	
A6G190357003	20-MW03-1012	07/18/2006	Soil	X	X	X	X	
A6H120109001	20-MW04-1416	08/11/2006	Soil	X	X	X	X	
A6I220386001	20-MW-01	09/22/2006	Groundwater	X	X		X	
A6I270124001	20-MW-02	09/25/2006	Groundwater	X	X		X	
A6I270124002	20-MW-04	09/25/2006	Groundwater	X	X		X	
A6I270124003	20-MW-03	09/25/2006	Groundwater	X	X	!	X	

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B] SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

Met = RCRA Metals [SW-846 Methods 6010B/6020/7470A/7471A]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Positive detections for methylene chloride in sample 20-MW01-1618, for acetone in samples 20-MW02-1012 and 20-MW03-1012, and for bis(2-ethylhexyl)phthalate in samples 20-MW02-1012 and 20-MW-01 were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks at similar concentrations.
- Trip blanks submitted with the 08/11/06 and 09/25/06 samples were logged in and

reported with samples from other sites. Trace amounts of methylene chloride and toluene were detected in one trip blank each. Therefore, the positive methylene chloride result for sample 20-MW04-1416 and the positive toluene results in samples 20-MW-04 and 20-MW-03 were qualified as nondetect ("U"), since the sample concentrations were not significantly higher than those in the blanks.

- The semivolatile compound, bis(2-ethylhexyl)phthalate, was detected at concentrations ranging from 0.9 to 1.8 ug/L in all equipment blanks associated with project samples. Since this compound is considered a common laboratory contaminant, its presence in environmental samples at concentrations less than ten times the highest equipment blank concentration is attributed to external contamination rather than actual site conditions. Therefore, the positive detection for bis(2-ethylhexyl)phthalate in sample 20-MW-02 was qualified as nondetect ("U") at the value reported, even though it was not detected in the associated method blank.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J" for organics, and "B" for inorganics), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, organic "J" qualifiers were retained with the numeric results. The inorganic "B" qualifiers were changed to "J" qualifiers for the sake of consistency throughout the package.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-7.

# Table 2-1 Analytical Data Summary Site 20 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G190357001 20-MW01-1618 07/18/2006	A6G190357002 20-MW02-1012 07/18/2006	A6G190357003 20-MW03-1012 07/18/2006	A6H120109001 20-MW04-1416 08/11/2006
1.1.1-Trichloroethane	ug/kg	6.8 U	60 U	6.1 U	60 U
1,1,2,2-Tetrachloroethane	ug/kg	6.8 U	60 U	6.1 U	60 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U
1,1,2-Trichloroethane	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U
1,1-Dichloroethane	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U
1,1-Dichloroethene	ug/kg ug/kg	6.8 U	60 U	6.1 U	1
1,2,4-Trichlorobenzene	ug/kg	6.8 U	60 U	6.1 U	60 U
1,2-Dibromo-3-chloropropane	ug/kg	14 U	120 U	12 U	120 U
1.2-Dibromoethane	ug/kg	6.8 U	60 U	6.1 U	60 U
1,2-Dichlorobenzene	ug/kg ug/kg	6.8 U	60 U	6.1 U	
1,2-Dichloroethane	or 🛊 is now Program in the	6.8 U	60 U	6.1 U	60 U
The form the same of the same	ug/kg	1	1 7 7 7	•	60 U
1,2-Dichloropropane	ug/kg	6.8 U 6.8 U	60 U	6.1 U	60 U
1,3-Dichlorobenzene 1,4-Dichlorobenzene	ug/kg	6.8 U	60 U	6.1 U 6.1 U	60 U
1,4-Dicnioropenzene 2-Butanone	ug/kg	5.8 U	47 J	3.3 J	60 U
2-Butanone 2-Hexanone	ug/kg	27 U	240 U	3.3 J 24 U	46 J
The state of the s	ug/kg	27 U	240 U	24 U	240 U
4-Methyl-2-pentanone Acetone	ug/kg	27 U	240 U	24 U	240 U
Benzene	ug/kg	0.59 J	60 U	**	81 J
Bromodichloromethane	ug/kg ug/kg	6.8 U	60 U	6.1 U 6.1 U	2.9 J
AND THE RESERVE OF THE PROPERTY OF THE PROPERT		6.8 U		6.1 U	60 U
Bromoform	ug/kg	6.8 U	60 U	6.1 U	60 U
Bromomethane	ug/kg	5.1 J	6.1 J	and the second second second	60 U
Carbon disulfide Carbon tetrachloride	ug/kg	6.8 U	60 U	<b>0.33 J</b> 6.1 U	12 J
Chlorobenzene	ug/kg	6.8 U	60 U	6.1 U	60 U
Chloroethane	ug/kg	6.8 U	60 U	6.1 U	60 U
Chloroform	ug/kg	6.8 U	60 U	6.1 U	60 U
Chloromethane	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U 60 U
cis-1,2-Dichloroethene	ug/kg	6.8 U	60 U	6.1 U	1
cis-1,3-Dichloropropene	ug/kg	6.8 U	60 U	6.1 U	60 U 60 U
Cyclohexane	ug/kg	14 U	120 U	12 U	The state of the s
Dibromochloromethane	ug/kg	6.8 U	60 U	6.1 U	<b>200</b> 60 U
Dichlorodifluoromethane	ug/kg	6.8 U	60 U	6.1 U	60 U
Ethylbenzene	ug/kg	6.8 U	60 U	0.68 J	60 U
Isopropylbenzene	ug/kg	0.53 J	8.8 J	0.33 J	1800
Methyl acetate	ug/kg	14 U	120 U	12 U	120 U
Methyl tert-butyl ether	ug/kg	27 U	240 U	12 U	240 U
Methylcyclohexane	ug/kg	14 U	210	2.4 J	890
Methylene chloride	ug/kg	9.3 U	60 U	6.1 U	60 U
Styrene	ug/kg	6.8 U	60 U	6.1 U	60 U
Tetrachloroethene	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U
Toluene	ug/kg	1.3 J	60 U	0.63 J	60 U
trans-1,2-Dichloroethene	ug/kg	6.8 U	60 U	6.1 U	60 U
trans-1,3-Dichloropropene	ug/kg	6.8 U	60 U	6.1 U	60 U
Trichloroethene	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U
Trichlorofluoromethane	ug/kg ug/kg	6.8 U	60 U	6.1 U	60 U
Vinyl chloride	ug/kg ug/kg	6.8 U	60 U	the state of the s	the transfer of the state of th
	1			6.1 U	60 U
Xylenes (total)	ug/kg	2.4 J	120 U	2.3 J	43 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-2 Analytical Data Summary Site 20 Soil Semivolatiles ODOT Innerbelt Study

		A6G190357001 20-MW01-1618	A6G190357002 20-MW02-1012	A6G190357003 20-MW03-1012	A6H120109001 20-MW04-1416
PARAMETER	UNITS	07/18/2006	07/18/2006	07/18/2006	08/11/2006
1,1'-Biphenyl	ug/kg	1800 U	400 U	20000 U	1600 U
2,2'-oxybis(1-Chloropropane)	ug/kg	1800 U	400 U	20000 U	1600 U
2,4,5-Trichlorophenol	ug/kg	1800 U	400 U	20000 U	1600 U
2,4,6-Trichlorophenol	ug/kg	1800 U	400 U	20000 U	1600 U
2,4-Dichlorophenol	ug/kg	1800 U	400 U	20000 U	1600 U
2,4-Dimethylphenol	ug/kg	1800 U	400 U	20000 U	1600 U
2,4-Dinitrophenol	ug/kg	1800 U	400 U	20000 U	1600 U
2,4-Dinitrotoluene	ug/kg	1800 U	400 U	20000 U	1600 U
2,6-Dinitrotoluene	ug/kg	1800 U	400 U	20000 U	1600 U
2-Chloronaphthalene	ug/kg	1800 U	400 U	20000 U	1600 U
2-Chlorophenol	ug/kg	1800 U	400 U	20000 U	1600 U
2-Methylnaphthalene	ug/kg	860 J	29 J	4300 J	87 J
2-Methylphenol	ug/kg	1800 U	400 U	20000 U	1600 U
2-Nitroaniline	ug/kg	1800 U	400 U	20000 U	1600 U
2-Nitrophenol	ug/kg	1800 U	400 U	20000 U	1600 U
3,3'-Dichlorobenzidine	ug/kg	1800 U	400 U	20000 U	1600 U
3-Nitroaniline	ug/kg	1800 U	400 U	20000 U	1600 U
4,6-Dinitro-2-methylphenol	ug/kg	1800 U	400 U	20000 U	1600 U
4-Bromophenyl phenyl ether	ug/kg	1800 U	400 U	20000 U	1600 U
4-Chloro-3-methylphenol	ug/kg	1800 U	400 U	20000 U	1600 U
4-Chloroaniline	ug/kg	1800 U	400 U	20000 U	1600 U
4-Chlorophenyl phenyl ether	ug/kg	1800 U	400 U	20000 U	1600 U
4-Methylphenol	ug/kg	1800 U	400 U	20000 U	1600 U
4-Nitroaniline	ug/kg	1800 U	400 U	20000 U	1600 U
4-Nitrophenol	ug/kg	1800 U	400 U	20000 U	1600 U
Acenaphthene	ug/kg	410 J	400 U	7100 J	88 J
Acenaphthylene	ug/kg	190 J	400 U	20000 U	1600 U
Acetophenone	ug/kg	360 U	80 U	4000 U	320 U
Anthracene	ug/kg	840 J	17 J	12000 J	70 J
Atrazine	ug/kg	1800 U	400 U	20000 U	1600 U
Benzaldehyde	ug/kg	1800 U	400 U	20000 U	1600 U
Benzo(a)anthracene	ug/kg	2300	48 J	16000 J	130 J
Benzo(a)pyrene	ug/kg	1800	51 J	13000 J	110 J
Benzo(b)fluoranthene	ug/kg	2100	68 J	17000 J	130 J
Benzo(ghi)perylene	ug/kg	890 J	36 J	7900 J	76 J
Benzo(k)fluoranthene	ug/kg	990 J	28 J	5500 J	92 J
bis(2-Chloroethoxy)methane	ug/kg	1800 U	400 U	20000 U	1600 U
bis(2-Chloroethyl) ether	ug/kg	1800 U	400 U	20000 U	1600 U
bis(2-Ethylhexyl) phthalate	ug/kg	1800 U	400 U	20000 U	1600 U
Butyl benzyl phthalate	ug/kg	1800 U	400 U	20000 U	1600 U
Caprolactam	ug/kg	1800 U	400 U	20000 U	1600 U
Carbazole	ug/kg	430 J	400 U	4200 J	95 J
Chrysene	ug/kg	2200	52 J	14000 J	130 J
Dibenz(a,h)anthracene	ug/kg	230 J	8.8 J	2000 J	1600 U
Dibenzofuran	ug/kg	510 J	400 U	5300 J	1600 U
Diethyl phthalate	ug/kg	1800 U	400 U	20000 U	1600 U
Dimethyl phthalate	ug/kg	1800 U	400 U	20000 U	1600 U
Di-n-butyl phthalate	ug/kg	1800 U	400 U	20000 U	1600 U
Di-n-octyl phthalate	ug/kg	1800 U	400 U	20000 U	1600 U
Fluoranthene	ug/kg	5100	85 J	40000	330 J
Fluorene	ug/kg	620 J	11 J	6400 J	97 J

Table 2-2 Analytical Data Summary Site 20 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G190357001 20-MW01-1618 07/18/2006	A6G190357002 20-MW02-1012 07/18/2006	A6G190357003 20-MW03-1012 07/18/2006	A6H120109001 20-MW04-1416 08/11/2006
Hexachlorobenzene	ug/kg	1800 U	400 U	20000 U	1600 U
Hexachlorobutadiene	ug/kg	1800 U	400 U	20000 U	1600 U
Hexachlorocyclopentadiene	ug/kg	1800 U	400 U	20000 U	1600 U
Hexachloroethane	ug/kg	1800 U	400 U	20000 U	1600 U
Indeno(1,2,3-cd)pyrene	ug/kg	820 J	34 J	6600 J	66 J
Isophorone	ug/kg	1800 U	400 U	20000 U	1600 U
Naphthalene	ug/kg	790 J	23 J	7200 J	1600 U
Nitrobenzene	ug/kg	1800 U	400 U	20000 U	1600 U
N-Nitrosodi-n-propylamine	ug/kg	1800 U	400 U	20000 U	1600 U
N-Nitrosodiphenylamine	ug/kg	1800 U	400 U	20000 U	1600 U
Pentachlorophenol	ug/kg	1800 U	400 U	20000 U	1600 U
Phenanthrene	ug/kg	4200	48 J	42000	300 J
Phenol	ug/kg	1800 U	400 U	20000 U	1600 U
Pyrene	ug/kg	4000	82 J	32000	250 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-3 Analytical Data Summary Site 20 Soil TPH ODOT Innerbelt Study

PARAMETER	UNITS	A6G190357001 20-MW01-1618 07/18/2006	A6G190357002 20-MW02-1012 07/18/2006	A6G190357003 20-MW03-1012 07/18/2006	A6H120109001 20-MW04-1416 08/11/2006
Gasoline Range Organics (C6-C12)	ug/kg	240	160000	120 U	280000
C10-C20	mg/kg	300	28	100	800
C20-C34	mg/kg	1800	8.4	580	140

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

# Table 2-4 Analytical Data Summary Site 20 Soil Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6G190357001 20-MW01-1618 07/18/2006	A6G190357002 20-MW02-1012 07/18/2006	A6G190357003 20-MW03-1012 07/18/2006	A6H120109001 20-MW04-1416 08/11/2006
Percent Solids	%	73.3	83.4	82.5	83.8
Arsenic	mg/kg	11.1	5.9	12.8	11.7
Barium	mg/kg	94.5	24.3	97.9	43
Cadmium	mg/kg	0.51	0.12 J	0.59	0.29
Chromium	mg/kg	12.2	6.7	9.8	8.6
Lead	mg/kg	176	22.9	446	43.7
Selenium	mg/kg	0.68 U	0.72	0.61 U	0.8
Silver	mg/kg	0.68 U	0.60 U	0.61 U	0.60 U
Mercury	mg/kg	0.54	0.076 J	0.17	0.2

 $<sup>\</sup>ensuremath{\mathsf{U}}$  =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-5
Analytical Data Summary
Site 20 Water Volatiles
ODOT Innerbelt Study

1,1,1-Trichloroethane 1,1,2,2-Tetrachloroethane 1,1,2-Trichloro-1,2,2-trifluoroethane 1,1,2-Trichloroethane 1,1-Dichloroethane 1,1-Dichloroethene 1,2,4-Trichlorobenzene 1,2-Dibromo-3-chloropropane 1,2-Dibromoethane 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloroethane 1,2-Dichloropropane 1,3-Dichloropropane 1,3-Dichlorobenzene	ug/L ug/L ug/L ug/L ug/L ug/L ug/L ug/L	1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 2.0 U 1.0 U	1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 2.0 U	5.0 U 5.0 U 5.0 U 5.0 U 5.0 U 5.0 U 5.0 U	1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 0.27 J 1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane 1,1,2-Trichloroethane 1,1-Dichloroethane 1,1-Dichloroethene 1,2,4-Trichlorobenzene 1,2-Dibromo-3-chloropropane 1,2-Dibromoethane 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L ug/L ug/L ug/L ug/L ug/L	1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 2.0 U 1.0 U	1.0 U 1.0 U 1.0 U 1.0 U 1.0 U 2.0 U	5.0 U 5.0 U 5.0 U 5.0 U 5.0 U	1.0 U 1.0 U 1.0 U <b>0.27 J</b>
1,1,2-Trichloroethane 1,1-Dichloroethane 1,1-Dichloroethene 1,2,4-Trichlorobenzene 1,2-Dibromo-3-chloropropane 1,2-Dichlorobenzene 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L ug/L ug/L ug/L ug/L	1.0 U 1.0 U 1.0 U 1.0 U 2.0 U 1.0 U 1.0 U	1.0 U 1.0 U 1.0 U 1.0 U 2.0 U	5.0 U 5.0 U 5.0 U 5.0 U	1.0 U 1.0 U <b>0.27 J</b>
1,1-Dichloroethane 1,1-Dichloroethene 1,2,4-Trichlorobenzene 1,2-Dibromo-3-chloropropane 1,2-Dichlorobenzene 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L ug/L ug/L ug/L	1.0 U 1.0 U 1.0 U 2.0 U 1.0 U 1.0 U	1.0 U 1.0 U 1.0 U 2.0 U	5.0 U 5.0 U 5.0 U	1.0 U <b>0.27 J</b>
1,1-Dichloroethene 1,2,4-Trichlorobenzene 1,2-Dibromo-3-chloropropane 1,2-Dibromoethane 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L ug/L ug/L ug/L	1.0 U 1.0 U 2.0 U 1.0 U 1.0 U	1.0 U 1.0 U 2.0 U	5.0 U 5.0 U	0.27 J
1,2,4-Trichlorobenzene 1,2-Dibromo-3-chloropropane 1,2-Dibromoethane 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L ug/L ug/L	1.0 U 2.0 U 1.0 U 1.0 U	1.0 U 2.0 U	5.0 U	
1,2-Dibromo-3-chloropropane 1,2-Dibromoethane 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L ug/L	2.0 U 1.0 U 1.0 U	2.0 U		1.0 U
1,2-Dibromoethane 1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L ug/L	1.0 U 1.0 U		10 U	
1,2-Dichlorobenzene 1,2-Dichloroethane 1,2-Dichloropropane	ug/L ug/L	1.0 U	40.1		2.0 U
1,2-Dichloroethane 1,2-Dichloropropane	ug/L		1.0 U	5.0 U	1.0 U
1,2-Dichloropropane		40.11	1.0 U	5.0 U	1.0 U
ngi ang kanada ni nikata kanada kanada ang kanada ka		1.0 U	1.0 U	5.0 U	1.0 U
1,3-Dichlorobenzene		1.0 U	1.0 U	5.0 U	1.0 U
	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
2-Butanone	ug/L	10 U	1.4 J	4.9 J	5.5 J
2-Hexanone	ug/L	10 U	10 U	50 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U	50 U	10 U
Acetone	ug/L	10 U	10 U	50 U	3.2 J
Benzene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Bromodichloromethane	ug/L	3.1	1.0 U	5.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Chloroethane	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Chloroform	ug/L	15	1.0 U	5.0 U	1.0 U
Chloromethane	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	1.0 U	150	7.7
Dibromochloromethane	ug/L	0.42 J	1.0 U	5.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U	1.2 J	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U	190	0.34 J
Methyl acetate	ug/L	10 U	10 U	50 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U	25 U	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U	36	19
Methylene chloride	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Styrene	ug/L ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Toluene	ug/L ug/L	1.0 U	1.0 U	5.0 U	1.0 U
The first of the state of the s		1.0 U	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		The second of the second of the second of
trans-1,2-Dichloroethene trans-1,3-Dichloropropene	ug/L		1.0 U	5.0 U	1.0 U
the state of the second section and the second section is the second section of the second section in the second section is the second section in the second section in the second section is section in the second section in the second section is section in the second section in the second section is section in the section in the second section in the second section is section in the second section in the second section is section in the second section in the second section is section in the second section in the second section is section in the section in the section in the section is section in the section in the section is section in the section in the section is section in the section in the section is section in the section in the section is section in the section in the section in the section is section in the secti	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U	5.0 U	1.0 U
Vinyl chloride  Xylenes (total)	ug/L ug/L	1.0 U 2.0 U	1.0 U 2.0 U	5.0 U 10 U	1.0 U 2.0 U

 $<sup>\</sup>mbox{\bf U}$  =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

#### Table 2-6 Analytical Data Summary Site 20 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A61220386001 20-MW-01 09/22/2006	A6I270124001 20-MW-02 09/25/2006	A6I270124002 20-MW-04 09/25/2006	A6I270124003 20-MW-03 09/25/2006
1,1'-Biphenyl	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
2,4,5-Trichlorophenol	ug/L	5.0 U	5.0 U	20 U	5.0 U
2,4,6-Trichlorophenol	ug/L	5.0 U	5.0 U	20 U	5.0 U
2,4-Dichlorophenol	ug/L	2.0 U	2.0 U	8.0 U	2.0 U
2,4-Dimethylphenol	ug/L	2.0 U	2.0 U	8.0 U	2.0 ∪
2,4-Dinitrophenol	ug/L	5.0 U	5.0 U	20 U	5.0 U
2,4-Dinitrotoluene	ug/L	5.0 U	5.0 U	20 U	5.0 U
2,6-Dinitrotoluene	ug/L	5.0 U	5.0 U	20 U	5.0 U
2-Chloronaphthalene	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
2-Chlorophenol	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
2-Methylnaphthalene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
2-Methylphenol	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
2-Nitroaniline	ug/L	2.0 U	2.0 U	8.0 U	I'
2-Nitrophenol	ug/L ug/L	2.0 U	2.0 U	8.0 U	2.0 U 2.0 U
3,3'-Dichlorobenzidine	ug/L ug/L	5.0 U	2.0 U	20 U	5.0 U
3-Nitroaniline	ug/L	2.0 U	2.0 U	8.0 U	2.0 U
4,6-Dinitro-2-methylphenol	ug/L	5.0 U	5.0 U	20 U	5.0 U
4-Bromophenyl phenyl ether	ug/L	2.0 U	2.0 U	8.0 U	the state of the s
4-Chloro-3-methylphenol	ug/L	2.0 U	2.0 U	8.0 U	2.0 U
4-Chloroaniline	ug/L	2.0 U	2.0 U	the state of the s	2.0 U
4-Chlorophenyl phenyl ether		2.0 U		8.0 U	2.0 U
The state of the s	ug/L	And the second second second second second second	2.0 U	8.0 U	2.0 U
4-Methylphenol 4-Nitroaniline	ug/L	1.0 U 2.0 U	1.0 U	4.0 U	1.0 U
the state of the s	ug/L	and the second s	2.0 U	8.0 U	2.0 U
4-Nitrophenol	ug/L	5.0 U	5.0 U	20 U	5.0 U
Acenaphthulana	ug/L	0.20 U	0.20 U	0.80 U	0.38
Acetaphthylene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
Acetophenone	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Anthracene	ug/L	0.20 U	0.20 U	0.80 U	0.51
Atrazine	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Benzaldehyde	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Benzo(a)anthracene	ug/L	0.20 U	0.20 U	0.80 U	0.63
Benzo(a)pyrene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
Benzo(b)fluoranthene	ug/L	0.20 U	0.20 U	0.80 U	0.51
Benzo(ghi)perylene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
Benzo(k)fluoranthene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
bis(2-Chloroethoxy)methane	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
bis(2-Chloroethyl) ether	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
bis(2-Ethylhexyl) phthalate	ug/L	1.8 U	1.3 U	4.0 U	1.0 U
Butyl benzyl phthalate	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Caprolactam	ug/L	5.0 U	5.0 U	20 U	5.0 U
Carbazole	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Chrysene	ug/L	0.20 U	0.20 U	0.80 U	0.55
Dibenz(a,h)anthracene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
Dibenzofuran	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Diethyl phthalate	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Dimethyl phthalate	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Di-n-butyl phthalate	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Di-n-octyl phthalate	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Fluoranthene	ug/L	0.20 U	0.20 U	0.80 U	2.0
Fluorene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
Hexachlorobenzene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U

Table 2-6
Analytical Data Summary
Site 20 Water Semivolatiles
ODOT Innerbelt Study

PARAMETER	UNITS	A6I220386001 20-MW-01 09/22/2006	A6I270124001 20-MW-02 09/25/2006	A6I270124002 20-MW-04 09/25/2006	A6I270124003 20-MW-03 09/25/2006
Hexachlorobutadiene	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U	10 U	40 U	10 U
Hexachloroethane	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U	0.20 U	0.80 U	0.20 U
Isophorone	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Naphthalene	ug/L	0.20 U	0.20 U	0.80 U	0.39
Nitrobenzene	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Pentachlorophenol	ug/L	5.0 U	5.0 U	20 U	5.0 U
Phenanthrene	ug/L	0.20 ∪	0.20 U	0.80 U	1.7
Phenol	ug/L	1.0 U	1.0 U	4.0 U	1.0 U
Pyrene	ug/L	0.20 U	0.20 U	0.80 U	1.8

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

# Table 2-7 Analytical Data Summary Site 20 Water Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6I220386001 20-MW-01 09/22/2006	A61270124001 20-MW-02 09/25/2006	A6I270124002 20-MW-04 09/25/2006	A6I270124003 20-MW-03 09/25/2006
Arsenic	ug/L	10.0 U	10.0 U	94.8	23.1
Barium	ug/L	32.8 J	622	658	287
Cadmium	ug/L	2.0 U	2.0 U	2.0 U	0.85 J
Chromium	ug/L	11.8	3.1 J	4.0 J	32
Lead	ug/L	4.4	3.0 U	3.0 U	522
Selenium	ug/L	2.6 J	5.0 U	5.0 U	5.0 U
Silver	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Mercury	ug/L	0.20 U	0.20 U	0.20 U	0.54

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

**Phase II Findings** 

### 6.10 SITE 21 - WHITE PROPERTIES

According to the Cleveland City Directories, the site was listed as Whitman Jackson from 1954 through 1984. There were no further listings for the site.

According to the Cleveland Fire Prevention Bureau files, a Permit dated November 27, 1953 indicates a 500-gallon UST with unknown contents was installed at the site. In addition, there were dip tanks at the site, associated with the wood treatment operations. A Permit dated November 28, 1955, indicates a 1,000-gallon UST of gasoline was installed at the site. A Report dated August 10, 1965 indicates there is a 1,000-gallon gasoline UST, 900 1-gallon turpentine containers, 2,000 1-gallon liquid floor wax containers, and 250 gallons of spray paint in 8-ounce cans stored at the site. A Report dated March 25, 1983 indicates a 500-gallon steel UST was removed from the site. The UST was located near the northwest corner of the building. A letter was sent to the Mr. Andy Lakowitz of Global Reserve from the Bureau, indicating the USTs on the site had not been properly abandoned.

At the time of the Phase I reconnaissance, the site was a commercial warehouse. No surface staining, stressed vegetation and/or the handling and storage of hazardous materials were observed on the site. Multiple 55-gallon drums were observed on an adjacent property.

#### 6.10.1 Field Activities

A total of four monitoring wells were proposed for Site 21. During the Phase II site reconnaissance, it was determined that one of the proposed boring locations was inaccessible; therefore, only three monitoring wells were installed. The boring was to be located adjacent to the northeast corner of the building. The overhead and underground utilities located in the area of the site made this proposed well inaccessible.

Groundwater was encountered in three soil borings (MW01, MW02, and MW03) during field activities and monitoring wells were installed. At the time of groundwater sampling, MW01 was dry and no groundwater sample could be obtained. Three soil and two groundwater samples were collected and analyzed for VOCs, SVOCs, TPH and RCRA Metals. A Sample Location Map is included as **Figure 6-10A**.

# 6.10.2 Site-Specific Geology/Hydrogeology

Soils at Site 21 consisted primarily of sand underlain by silty clay and clay with minor amounts of organics. Bedrock was not encountered in any of the soil borings, which were advanced to 16 feet bgs.

Groundwater elevations were measured at Site 21 on September 21, 2006. Monitoring well (MW01) was dry. Localized groundwater flow across Site 21 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 21 is to the north-northwest, towards the Cuyahoga River.

**Phase II Findings** 

#### 6.10.3 Geophysical Survey

The geophysical survey at Site 21 included the asphalt-paved parking and drive surfaces northeast, northwest, and west of the building. The survey transects were completed in four subsurveys that were aligned parallel or perpendicular to the building. The presence of buildings adjacent to surveyed areas prevented the use of a global positioning system (GPS) to obtain corresponding latitude and longitude datum necessitating the need to paint survey points on the pavement.

The survey identified five areas of anomalously high magnetic gradients. These areas are shown on **Figure 6-10B** as anomalies A through E as described below:

- A. A gradient anomaly was observed in the loading dock area. This anomaly is likely due to the presence of a steel dumpster being used to dispose of roofing material at the time of the survey.
- B. An anomaly was detected near the center of the survey conducted northeast of the building. Based on shape of the anomaly and the presence of pipes protruding from the asphalt, this anomaly likely represents a UST.
- C. Several localized magnetic anomalies were detected that were assumed to be related to surface objects and are unlikely related to a UST.
- D. Areas of high magnetic gradients were observed in the northwest corner of the Site that may relate to one or more USTs. Their detection was partially obscured by a high background gradient that was assumed to be related to recent paving.
- E. An elongated magnetic anomaly was observed in the loading doc area. This anomaly may be related to buried pipes.

Based on the results of the geophysical survey, several probable USTs were detected.

### 6.10.4 Soil Analytical Results

A total of six VOCs were detected in the three soil samples submitted from Site 21. The chemicals detected in the samples submitted were 2-butanone (7.4 ug/kg), acetone (40 mg/kg), benzene (0.53 ug/kg), carbon disulfide (ranging from 0.38 ug/kg to 1.3 ug/kg), toluene (0.38 ug/kg), and trichlorofluoromethane (4.8 ug/kg). All other VOCs were below the detection limits.

A total of seventeen SVOCs were detected in the three samples submitted from Site 21. Concentrations of 2-methylnaphthanene (60 ug/kg and 61 ug/kg), acenaphthene (17 ug/kg and 98 ug/kg), acenaphthylene (ranging from 9.5 ug/kg to 36 ug/kg), anthracene (ranging from 15 ug/kg to 190 ug/kg), benzaldehyde (140 ug/kg), benzo(a)anthracene (ranging from 23 ug/kg to 600 ug/kg), benzo(a)pyrene (ranging from 17 ug/kg to 510 ug/kg), benzo(b)fluoranthene (ranging from 20 ug/kg to 690 ug/kg), benzo(g,h,i)perylene (ranging from 53 ug/kg to 330 ug/kg), benzo(k)fluoranthene (ranging from 48 ug/kg to 260 ug/kg), bis(2-ethylhexyl)phthalate (21 ug/kg and 51 ug/kg), butyl benzyl phthalate (24 ug/kg),carbazole (53 ug/kg), chrysene (ranging from 27

ug/kg to 570 ug/kg), dibenz(a,h)anthracene (17 ug/kg and 86 ug/kg), dibenzofuran (23 ug/kg and 44 ug/kg), and di-n-butyl phthalate (68 ug/kg) were detected in the soil samples submitted.

Diesel range total petroleum hydrocarbons were detected in the soil samples from Site 21. The highest concentrations were found in the diesel range, which is consistent with the concentrations of SVOCs detected across Site 21. Concentrations of the middle petroleum fraction ranged from 3.7 mg/kg to 14 mg/kg. Concentrations of the heavy petroleum fraction ranged from 16 mg/kg to 53 mg/kg.

Seven of the eight RCRA Metals were detected in the samples submitted from Site 21. Arsenic, ranging from 6.2 mg/kg to 9.5 mg/kg, was detected in all samples submitted. Barium, ranging from 25.2 mg/kg to 78.8 mg/kg, was detected in the all samples submitted. Cadmium, 0.095 mg/kg to 0.5 mg/kg, was detected in all samples submitted. Chromium, ranging from 5 mg/kg to 17.3 mg/kg, was detected in the samples submitted. Lead, ranging from 28.1 mg/kg to 156 mg/kg, was detected in the samples submitted. Selenium, 0.52 mg/kg and 2 mg/kg, was detected in samples submitted. Mercury, ranging from 0.091 mg/kg to 0.62 mg/kg, was detected in the samples submitted.

The analytical results are presented in Table 6-10A.

### 6.10.5 Groundwater Analytical Results

Four VOCs were detected in groundwater samples collected from Site 21. Concentrations of 1,1-dichloroethene (0.0.67 ug/L), acetone (1.1 ug/L), chloromethane (0.15 ug/L), and methyl tertbutyl ether (0.27 ug//L and 0.84 ug/L) were detected in the groundwater samples submitted. All other VOCs were below the detection limits.

Four SVOCs were detected in the groundwater samples submitted from Site 21. Concentrations of caprolactam (0.70 ug/L), diethyl phthalate (0.89 ug/L), dimethyl phthalate (0.95 ug/L), and di-n-octyl phthalate (1.3 ug/L) were detected in the groundwater sample submitted from 21-MW03. All other SVOCs were below the detection limits.

Three of the eight RCRA metals were detected in the samples submitted from Site 21. Barium, ranging from 119 ug/L to 135 ug/L, was detected in all samples submitted. Cadmium, 10.8 ug/L, was detected in sample 21-MW03. Selenium, 3.3 ug/L, was detected in sample 21-MW03.

The analytical results are presented in Table 6-10B.

### 6.10.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on Table 6-10A and 6-10B.

**SECTIONSIX** 

**Phase II Findings** 

There were no VOCs, SVOCs, TPH, and RCRA metals detected in any of the soil samples from Site 21, which exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

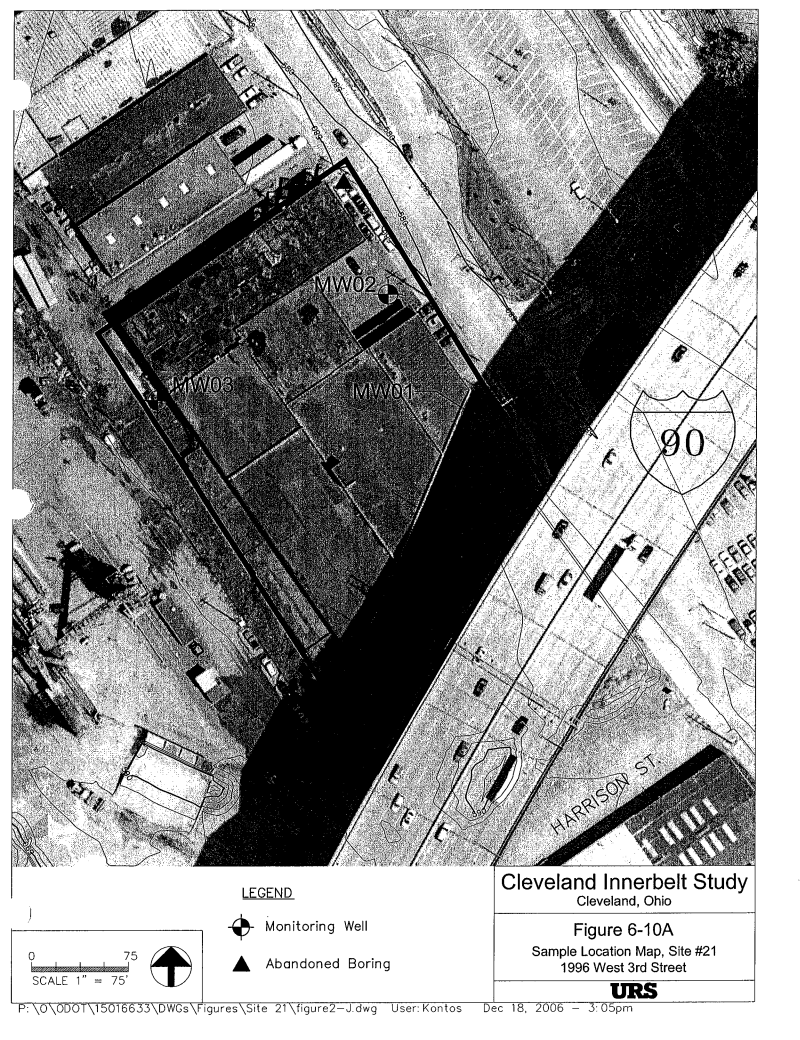
There were no VOCs, SVOCs, or RCRA metals detected in any of the groundwater samples from Site 21, which exceeded the OEPA VAP standard for generic unrestricted potable use standards or BUSTR closure action levels.

#### 6.10.7 Conclusions

Based on the results of the geophysical survey, several probable USTs were detected.

Based on the analytical results, it is unlikely that the soils and groundwater at Site 21 would require special management during construction.

**FIGURES** 





**TABLES** 

Summary of Detected Chemicals in Soil Site 21 - White Properties ODOT Innerbelt Study Cleveland, Ohio Table v=10A

ARAMETER	UNITS	VAP commercial Industrial Standard	VAP Construction Worker Standard <sup>©</sup>	BUSTR Closure Action Level	21-MW01-0406 08/10/2006	21-MW02-0406 07/19/2006	21-MW03-0204 08/10/2006
2-Butanone	ug/kg	71,600,000	80,000,000	•	26 U	24 U	7.4 J
Acetone Annexacionamente proposition and annotation	ug/kg	100,000,000	100,000,000	•	26 U	24 U	40
Benzene State production of the state of the	ug/kg	100,000	310,000	149	6.4 U	5.9 ∪	0.53 J
Carbon disulfide	ug/kg	720,000	720,000	•••	0.38 J	0.95 J	1.3 J
Toluene	ug/kg	520,000	520,000	49,100	6.4 U	5.9 U	0.38 J
Trichlorofluoromethane	ug/kg	2,000,000	2,000,000		4.8 J	5.9 U	5.7 U
2-Methylnaphthalene	ng/kg	***	•	1	Ր 09	390 U	61 J
Acenaphthene	ug/kg	180,000,000	530,000,000		7 86	17.0	380 0
Acenaphthylene	ug/kg		4	••	36 J	068 n	ر 1 3.6
Anthracene	ug/kg	880,000,000	1,000,000,000	The state of the s	190 J	15.J	29 J
Benzaldehyde	ug/kg	-	-	-	430 U	140 J	380 U
Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	009	23 J	91.7
Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	510	L 71	72 J
Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	069	20 J	100
Benzo(ghi)perylene	ug/kg	•	-		330 J	390 U	<b>₽</b>
Benzo(k)fluoranthene	ug/kg	630,000	8,100,000	110,000	260 J	390 U	48 J
bis(2-Ethylhexyl) phthalate	ug/kg	230,000	230,000		51 J	21.5	380 U
Butyl benzyl phthalate	ug/kg	220,000	220,000	ı	24 J	390 U	380 U
Carbazole	ug/kg	10,000,000	31,000,000		53 J	OSE	380 U
Chrysene	ug/kg	6,700,000	41,000,000	1,100,000	570	27 J	92 J
Dibenz(a,h)anthracene	ug/kg	6,700	41,000	1,100	P 98	390 U	17.0
Dibenzofuran	ug/kg	The state of the s		1	44 J	OSE	23 J
Di-n-butyi phthalate	ug/kg	100,000	100,000	-	68 J	390 U	380 ∪
C10-C20	mg/kg	1		2,000	26 U	3.7	14 J
C20-C34	mg/kg	1	1	5,000	53	16	38
Arsenic	mg/kg	80	210	\$ 100 mm of 100 mm	9.5	6.3	6.2
Barium	mg/kg	200,000	45,000	-	78.8	25.2	46.7
Cadmium	mg/kg	022	420	The state of the s	0.5	0.095 J	0.31
Chromium	mg/kg	8,900	2,000	The the contract of the contra	17.3	ъ	7.4
Lead	mg/kg	And the property of the Constitution and phonone is a constitute of	See Section 1. Million of the Conference of the	The state of the s	156	28.1	87.1
Selenium	mg/kg	15,000	4,300		7	0.59 U	0.52 J
Silver	mg/kg	15,000	4,300	The state of the s	0.64 U	0.59 U	0.57 U
Mercury	mg/kg	300	84	-	0.62	ບ.091	0.44

<sup>-- =</sup> Standard not available

U= The analyte was analyzed for, but was not detected. Value shown is the reporting limit. J= Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial/ Industrial Land Use (2) VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Table v=10B
Summary of Detected Chemicals in Water
Site 21 - White Properties
ODOT Innerbelt Study
Cleveland, Ohio

	PARAMETER	UNITS	TVAP UPUS/T RDUPUS <sup>(1)</sup>	BUSTR Closure Action Level	21-MW-02 09/21/2006	21-MW-03 09/21/2006
	1,1-Dichloroethane	ng/L	1400	1	L 79.0	1.0 U
\$၂(	Acetone	ug/Ľ	1600	;	1.1 J	10 U
ΛC	Chloromethane		•	The self-time information for the self-time of the self-t	0.15 ປ	1.0 U
	Methyl tert-butyl ether	ug/L	40	40	0.27 J	0.84 J
\$	Caprolactam	ng/L	*		5.0 U	ر 07.0
OC	Diethyl phthalate		13000		1.0 U	D 68.0
ΛS	Dimethyl phthalate	ug/L	•		1.0 U	0.95 J
	Di-n-octyl phthalate	ug/L		The state of the s	1.0 U	1.3
ירצ	Barium	ng/L	2000	1	119	135
AT:	Lead	ng/L	15	*** *** *** *** *** *** *** *** *** **	3.0 U	10.8
IW	Selenium	ug/L	20	The state of the s	5.0 U	3.3 J. 8.5

-- = Standard not available

 $\mathsf{U} = \mathsf{The}$  analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

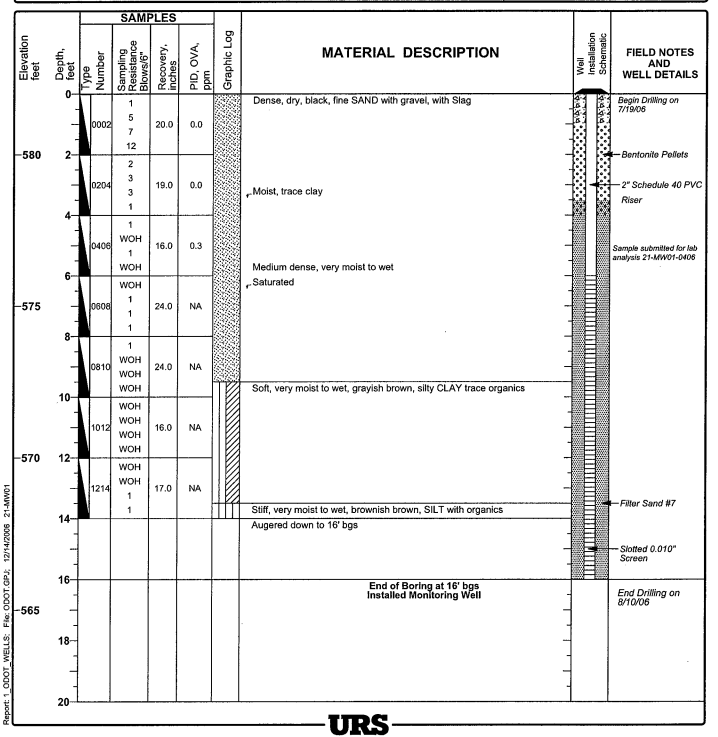
(1) VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

**BORING LOGS** 

Project Location: Site 21
Project Number: 15016633

## Log of Boring 21-MW01

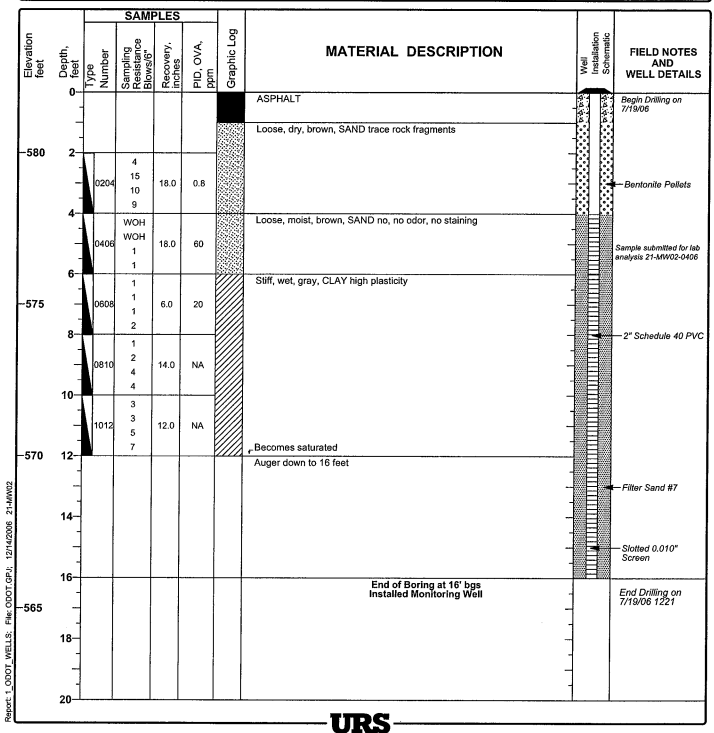
Date(s) 7/19/06 Drilled 7/19/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 16.0' bgs
Drill Rig Type LC-60	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 582
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured Well dry on 9/21/06			



Project Location: Site 21
Project Number: 15016633

## Log of Boring 21-MW02

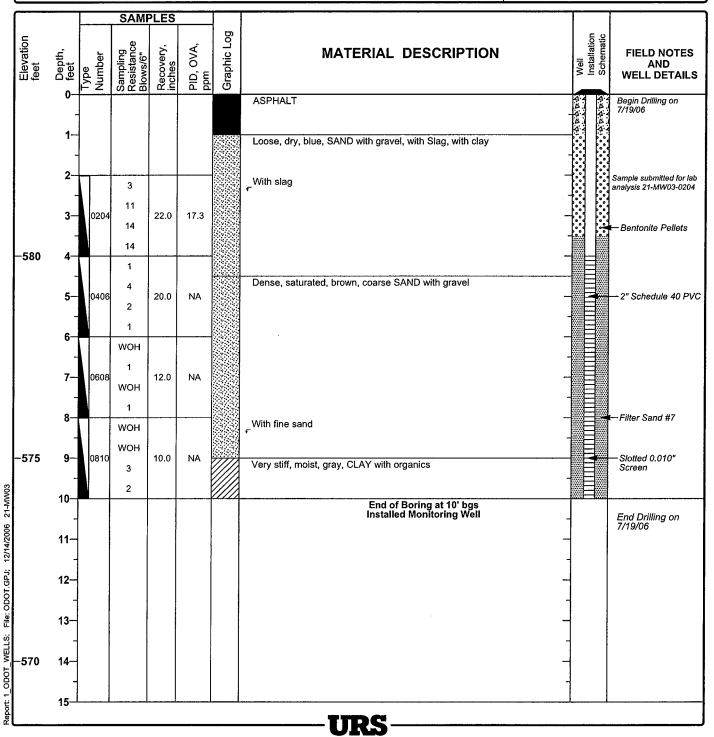
Date(s) 7/19/06 Drilled 7/19/06	Logged By X. Sotelo	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer 140# auto hammer	Total Depth of Borehole 16.0' bgs
Drill Rig Type CME-55	Drilling Contractor HAD, Inc.	Approximate Ground Elevation 582'
Location See Site Map	Sampling Method(s) 2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 578.96 on 9/21/06		



Project Location: Site 21
Project Number: 15016633

## Log of Boring 21-MW03

Date(s) Drilled 7/19/06	Logged J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data 140# auto hammer	Total Depth of Borehole 10.0' bgs
Drill Rig Type LC-60	Drilling Contractor HAD, Inc.	Approximate Ground Elevation 584'
Location See Site Map	Sampling Method(s) 2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 580.21 on 9/21/06		



DATA ASSESSMENT REPORT

#### Data Assessment Report ODOT Innerbelt Study Site 21 – White Properties

Reviewer: P. Schuler Date: November 13, 2006

Three soil samples and two groundwater samples were collected at the White Properties site at 1996 W. 3<sup>rd</sup> Street in Cleveland, Ohio, from July 19 through September 21, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample			Requested Analyses <sup>(1)</sup>				
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC	TPH	Met		
A6G210344001	21-MW02-0406	07/19/2006	Soil	X	X	X	X		
A6H120110001	21-MW01-0406	08/10/2006	Soil	X	X	X	X		
A6H120110002	21-MW03-0204	08/10/2006	Soil	X	X	X	X		
A6I220383001	21-MW-03	09/21/2006	Groundwater	X	X		X		
A6I220383002	21-MW-02	09/21/2006	Groundwater	X	X		X		

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B] SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

Met = RCRA Metals [SW-846 Methods 6010B/6020/7470A/7471A]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Positive detections for 2-butanone in sample 21-MW02-0406 and for gasoline range organics in sample 21-MW03-0204 were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks at similar concentrations.
- Trip blanks submitted with the Site 21 samples were logged in and reported with samples from other sites. Trace amounts of methylene chloride and acetone were detected in one or more trip blank each. Therefore, the positive methylene chloride results for samples 21-MW03-0204 and 21-MW-03 and the positive acetone result in sample 21-MW-03

- were qualified as nondetect ("U"), since the sample concentrations were not significantly higher than those in the associated blanks.
- The semivolatile compound, bis(2-ethylhexyl)phthalate, was detected at concentrations ranging from 0.9 to 1.8 ug/L in all equipment blanks associated with project samples. Since this compound is considered a common laboratory contaminant, its presence in environmental samples at concentrations less than ten times the highest equipment blank concentration is attributed to external contamination rather than actual site conditions. Therefore, the positive detections for bis(2-ethylhexyl)phthalate in samples 21-MW02-0406 and 21-MW01-0406 were qualified as nondetect ("U") at the value reported, even though it was not detected in the associated method blanks.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J" for organics, and "B" for inorganics), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, organic "J" qualifiers were retained with the numeric results. The inorganic "B" qualifiers were changed to "J" qualifiers for the sake of consistency throughout the package.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-7.

#### Table 2-1 Analytical Data Summary Site 21 Soil Volatiles ODOT Innerbelt Study

		A6G210344001 21-MW02-0406	A6H120110001 21-MW01-0406	A6H120110002 21-MW03-0204
PARAMETER	UNITS	07/19/2006	08/10/2006	08/10/2006
1,1,1-Trichloroethane	ug/kg	5.9 U	6.4 U	5.7 U
1,1,2,2-Tetrachloroethane	ug/kg	5.9 U	6.4 U	5.7 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.9 U	6.4 U	5.7 U
1,1,2-Trichloroethane	ug/kg	5.9 U	6.4 U	5.7 U
1,1-Dichloroethane	ug/kg	5.9 U	6.4 U	5.7 U
1,1-Dichloroethene	ug/kg	5.9 U	6.4 U	5.7 U
1,2,4-Trichlorobenzene	ug/kg	5.9 U	6.4 U	5.7 U
1,2-Dibromo-3-chloropropane	ug/kg	12 U	13 U	11 U
1,2-Dibromoethane	ug/kg	5.9 U	6.4 U	5.7 U
1,2-Dichlorobenzene	ug/kg	5.9 ∪	6.4 U	5.7 U
1,2-Dichloroethane	ug/kg	5.9 U	6.4 U	5.7 U
1,2-Dichloropropane	ug/kg	5.9 U	6.4 U	5.7 U
1,3-Dichlorobenzene	ug/kg	5.9 U	6.4 U	5.7 U
1,4-Dichlorobenzene	ug/kg	5.9 U	6.4 U	5.7 U
2-Butanone	ug/kg	24 U	26 U	7.4 J
2-Hexanone	ug/kg	24 U	26 U	23 U
4-Methyl-2-pentanone	ug/kg	24 U	26 U	23 U
Acetone	ug/kg	24 U	26 U	40
Benzene	ug/kg	5.9 U	6.4 U	0.53 J
Bromodichloromethane	ug/kg	5.9 U	6.4 U	5.7 U
Bromoform	ug/kg	5.9 U	6.4 U	5.7 U
Bromomethane	ug/kg	5.9 U	6.4 U	5.7 U
Carbon disulfide	ug/kg	0.95 J	0.38 J	1.3 J
Carbon tetrachloride	ug/kg	5.9 U	6.4 U	5.7 U
Chlorobenzene	ug/kg	5.9 U	6.4 U	5.7 U
Chloroethane	ug/kg	5.9 U	6.4 U	5.7 U
Chloroform	ug/kg	5.9 U	6.4 U	5.7 U
Chloromethane	ug/kg	5.9 U	6.4 U	5.7 U
cis-1,2-Dichloroethene	ug/kg	5.9 U	6.4 U	5.7 U
cis-1,3-Dichloropropene	ug/kg	5.9 U	6.4 U	5.7 U
Cyclohexane	ug/kg	12 U	13 U	11 U
Dibromochloromethane	ug/kg	5.9 U	6.4 U	5.7 U
Dichlorodifluoromethane	ug/kg	5.9 U	6.4 U	5.7 U
Ethylbenzene	ug/kg	5.9 U	6.4 U	5.7 U
Isopropylbenzene	ug/kg	5.9 U	6.4 U	5.7 U
Methyl acetate	ug/kg	12 U	13 U	11 U
Methyl tert-butyl ether	ug/kg ug/kg	24 U	26 U	23 U
Methylcyclohexane	ug/kg ug/kg	12 U	13 U	11 U
Methylene chloride	ug/kg ug/kg	5.9 U	6.4 U	5.7 U
Styrene	ug/kg ug/kg	5.9 U	6.4 U	5.7 U
Tetrachloroethene	ug/kg ug/kg	5.9 U	6.4 U	5.7 U
Toluene	ug/kg ug/kg	5.9 U	6.4 U	0.38 J
trans-1,2-Dichloroethene		processors and the second second second second	6.4 U	5.7 U
commence and the commence of t	ug/kg	5.9 U		
trans-1,3-Dichloropropene	ug/kg	5.9 U	6.4 U	5.7 U
Trichloroftharanthana	ug/kg	5.9 U	6.4 U	5.7 U
Trichlorofluoromethane	ug/kg	5.9 U	4.8 J	5.7 U
Vinyl chloride	ug/kg	5.9 U	6.4 U	5.7 U
Xylenes (total)	ug/kg	12 U	13 U	11 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-2 Analytical Data Summary Site 21 Soil Semivolatiles ODOT Innerbelt Study

		A6G210344001 21-MW02-0406	A6H120110001 21-MW01-0406	A6H120110002 21-MW03-0204
PARAMETER	UNITS	07/19/2006	08/10/2006	08/10/2006
1,1'-Biphenyl	ug/kg	390 U	430 U	380 U
2,2'-oxybis(1-Chloropropane)	ug/kg	390 U	430 U	380 U
2,4,5-Trichlorophenol	ug/kg	390 U	430 U	380 U
2,4,6-Trichlorophenol	ug/kg	390 U	430 U	380 U
2,4-Dichlorophenol	ug/kg	390 U	430 U	380 U
2,4-Dimethylphenol	ug/kg	390 U	430 U	380 U
2,4-Dinitrophenol	ug/kg	390 U	430 U	380 U
2,4-Dinitrotoluene	ug/kg	390 U	430 U	380 U
2,6-Dinitrotoluene	ug/kg	390 U	430 U	380 U
2-Chloronaphthalene	ug/kg	390 U	430 U	380 U
2-Chlorophenol	ug/kg	390 U	430 U	380 U
2-Methylnaphthalene	ug/kg	390 U	60 J	61 J
2-Methylphenol	ug/kg	390 U	430 U	380 U
2-Nitroaniline	ug/kg	390 U	430 U	380 U
2-Nitrophenol	ug/kg	390 U	430 U	380 U
3,3'-Dichlorobenzidine	ug/kg	390 U	430 U	380 U
3-Nitroaniline	ug/kg	390 U	430 U	380 U
4,6-Dinitro-2-methylphenol	ug/kg	390 U	430 U	380 U
4-Bromophenyl phenyl ether	ug/kg	390 U	430 U	380 U
4-Chloro-3-methylphenol	ug/kg	390 U	430 U	380 U
4-Chloroaniline	ug/kg	390 U	430 U	380 U
4-Chlorophenyl phenyl ether	ug/kg	390 U	430 U	380 U
4-Methylphenol	ug/kg	390 U	430 U	the same to the sa
4-Nitroaniline		390 U	1	380 U
***************************************	ug/kg	390 U	430 U	380 U
4-Nitrophenol	ug/kg	l de la distribución de	430 U	380 U
Acenaphthele	ug/kg	17 J	98 J	380 U
Acenaphthylene	ug/kg	390 U	36 J	9.5 J
Acetophenone	ug/kg	79 U	86 U	76 U
Anthracene	ug/kg	15 J	190 J	29 J
Atrazine	ug/kg	390 U	430 U	380 U
Benzaldehyde	ug/kg	140 J	430 U	380 U
Benzo(a)anthracene	ug/kg	23 J	600	91 J
Benzo(a)pyrene	ug/kg	17 J	510	72 J
Benzo(b)fluoranthene	ug/kg	20 J	690	100 J
Benzo(ghi)perylene	ug/kg	390 U	330 J	53 J
Benzo(k)fluoranthene	ug/kg	390 U	260 J	48 J
bis(2-Chloroethoxy)methane	ug/kg	390 U	430 U	380 U
bis(2-Chloroethyl) ether	ug/kg	390 U	430 U	380 U
bis(2-Ethylhexyl) phthalate	ug/kg	21 J	51 J	380 U
Butyl benzyl phthalate	ug/kg	390 U	24 J	380 U
Caprolactam	ug/kg	390 U	430 U	380 U
Carbazole	ug/kg	390 U	53 J	380 U
Chrysene	ug/kg	27 J	570	92 J
Dibenz(a,h)anthracene	ug/kg	390 U	86 J	17 J
Dibenzofuran	ug/kg	390 U	44 J	23 J
Diethyl phthalate	ug/kg	390 U	430 U	380 U
Dimethyl phthalate	ug/kg	390 U	430 U	380 U
Di-n-butyl phthalate	ug/kg	390 U	68 J	380 U
Di-n-octyl phthalate	ug/kg	390 U	430 U	380 U
Fluoranthene	ug/kg	61 J	1100	150 J
Fluorene	ug/kg	15 J	66 J	12 J

Table 2-2
Analytical Data Summary
Site 21 Soil Semivolatiles
ODOT Innerbelt Study

PARAMETER	UNITS	A6G210344001 21-MW02-0406 07/19/2006	A6H120110001 21-MW01-0406 08/10/2006	A6H120110002 21-MW03-0204 08/10/2006
Hexachlorobenzene	ug/kg	390 U	430 U	380 U
Hexachlorobutadiene	ug/kg	390 U	430 U	380 U
Hexachlorocyclopentadiene	ug/kg	390 U	430 U	380 ∪
Hexachloroethane	ug/kg	390 U	430 U	380 U
Indeno(1,2,3-cd)pyrene	ug/kg	390 U	290 J	44 J
Isophorone	ug/kg	390 U	430 U	380 U
Naphthalene	ug/kg	390 U	60 J	28 J
Nitrobenzene	ug/kg	390 U	430 U	380 U
N-Nitrosodi-n-propylamine	ug/kg	390 U	430 U	380 U
N-Nitrosodiphenylamine	ug/kg	390 U	430 U	380 U
Pentachlorophenol	ug/kg	390 U	430 U	380 U
Phenanthrene	ug/kg	99 J	570	120 J
Phenol	ug/kg	390 U	430 U	380 U
Pyrene	ug/kg	58 J	950	130 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-3 Analytical Data Summary Site 21 Soil TPH ODOT Innerbelt Study

PARAMETER	UNITS	A6G210344001 21-MW02-0406 07/19/2006	A6H120110001 21-MW01-0406 08/10/2006	A6H120110002 21-MW03-0204 08/10/2006
Gasoline Range Organics (C6-C12)	ug/kg	120 U	130 U	110 U
C10-C20	mg/kg	3.7	26 U	14 J
C20-C34	mg/kg	16	53	38

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

## Table 2-4 Analytical Data Summary Site 21 Soil Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6G210344001 21-MW02-0406 07/19/2006	A6H120110001 21-MW01-0406 08/10/2006	A6H120110002 21-MW03-0204 08/10/2006
Percent Solids	%	84.4	77.6	87.4
Arsenic	mg/kg	6.3	9.5	6.2
Barium	mg/kg	25.2	78.8	46.7
Cadmium	mg/kg	0.095 J	0.5	0.31
Chromium	mg/kg	5.0	17.3	7.4
Lead	mg/kg	28.1	156	87.1
Selenium	mg/kg	0.59 U	2.0	0.52 J
Silver	mg/kg	0.59 U	0.64 U	0.57 U
Mercury	mg/kg	0.091 J	0.62	0.44

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met

## Table 2-5 Analytical Data Summary Site 21 Water Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I220383001 21-MW-03 09/21/2006	A6I220383002 21-MW-02 09/21/2006
1.1.1-Trichloroethane	ug/L	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	0.67 J
1,1-Dichloroethene	ug/L	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U
2-Butanone	ug/L	10 U	10 U
2-Hexanone	ug/L	10 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U
Acetone	ug/L	10 U	1.1 J
Benzene	ug/L	1.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U
Carbon disulfide	ug/L ug/L	1.0 U	1.0 U
Carbon tetrachloride		1.0 U	1.0 U
Carbon tetrachionde Chlorobenzene	ug/L ug/L	1.0 U	1.0 U
Chloroethane	ug/L ug/L	1.0 U	1.0 U
Chloroform	ug/L	1.0 U	1.0 U
Chloromethane	ug/L	1.0 U	0.15 J
cis-1,2-Dichloroethene	ug/L ug/L	1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Cyclohexane		1.0 U	1.0 U
Dibromochloromethane	ug/L ug/L	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L ug/L	1.0 U	1.0 U
Ethylbenzene	ug/L ug/L	1.0 U	1.0 U
Isopropylbenzene	ug/L ug/L	1.0 U	1.0 U
The state of the s	ug/L ug/L	1.0 U	10 U
Methyl text build other		0.84 J	0.27 J
Methyl tert-butyl ether	ug/L	1.0 U	1.0 U
Methylcyclohexane	ug/L	The state of the s	1.0 U
Methylene chloride	ug/L	1.0 U	exercise and a second second
Styrene	ug/L	1.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U
Toluene	ug/L	1.0 U	1.0 U
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	2.0 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.



 $J = \mbox{Estimated concentration}$  because the result was below the sample reporting limit or quality control criteria were not met.

## Table 2-6 Analytical Data Summary Site 21 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I220383001 21-MW-03 09/21/2006	A6I220383002 21-MW-02 09/21/2006
PARAMETER  1,1'-Biphenyl	ug/L	1.0 U	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/L ug/L	1.0 U	1.0 U
2,4,5-Trichlorophenol	ug/L	5.0 U	5.0 U
	ug/L ug/L	5.0 U	5.0 U
2,4,6-Trichlorophenol	-	2.0 U	2.0 U
2,4-Dichlorophenol	ug/L	2.0 U	2.0 U
2,4-Dimethylphenol	ug/L	2.0 U	5.0 U
2,4-Dinitrophenol	ug/L	5.0 U	5.0 U
2,4-Dinitrotoluene	ug/L	5.0 U	5.0 U
2,6-Dinitrotoluene	ug/L	1.0 U	1.0 U
2-Chloronaphthalene	ug/L	the state of the state of	1.0 U
2-Chlorophenol	ug/L	1.0 U	0.20 U
2-Methylnaphthalene	ug/L	0.20 U 1.0 U	1.0 U
2-Methylphenol	ug/L		1.0 U 2.0 U
2-Nitroaniline	ug/L	2.0 U	
2-Nitrophenol	ug/L	2.0 U	2.0 U
3,3'-Dichlorobenzidine	ug/L	5.0 U	5.0 U
3-Nitroaniline	ug/L	2.0 U	2.0 U
4,6-Dinitro-2-methylphenol	ug/L	5.0 U	5.0 U
4-Bromophenyl phenyl ether	ug/L	2.0 U	2.0 U
4-Chloro-3-methylphenol	ug/L	2.0 U	2.0 U
4-Chloroaniline	ug/L	2.0 U	2.0 U
4-Chlorophenyl phenyl ether	ug/L	2.0 U	2.0 U
4-Methylphenol	ug/L	1.0 U	1.0 U
4-Nitroaniline	ug/L	2.0 U	2.0 U
4-Nitrophenol	ug/L	5.0 U	5.0 U
Acenaphthene	ug/L	0.20 U	0.20 U
Acenaphthylene	ug/L	0.20 U	0.20 U
Acetophenone	ug/L	1.0 U	1.0 U
Anthracene	ug/L	0.20 U	0.20 U
Atrazine	ug/L	1.0 U	1.0 U
Benzaldehyde	ug/L	1.0 U	1.0 U
Benzo(a)anthracene	ug/L	0.20 U	0.20 U
Benzo(a)pyrene	ug/L	0.20 U	0.20 U
Benzo(b)fluoranthene	ug/L	0.20 U	0.20 U
Benzo(ghi)perylene	ug/L	0.20 U	0.20 U
Benzo(k)fluoranthene	ug/L	0.20 U	0.20 U
bis(2-Chloroethoxy)methane	ug/L	1.0 U	1.0 U
bis(2-Chloroethyl) ether	ug/L	1.0 U	1.0 U
bis(2-Ethylhexyl) phthalate	ug/L	1.7 U	2.6 U
Butyl benzyl phthalate	ug/L	1.0 U	1.0 U
Caprolactam	ug/L	0.70 J	5.0 U
Carbazole	ug/L	1.0 U	1.0 U
Chrysene	ug/L	0.20 U	0.20 U
Dibenz(a,h)anthracene	ug/L	0.20 U	0.20 U
Dibenzofuran	ug/L	1.0 U	1.0 U
Diethyl phthalate	ug/L	0.89 J	1.0 U
Dimethyl phthalate	ug/L	0.95 J	1.0 U
Di-n-butyl phthalate	ug/L	1.0 U	1.0 U
Di-n-octyl phthalate	ug/L	1.3	1.0 U
Fluoranthene	ug/L	0.20 U	0.20 U
Fluorene	ug/L	0.20 U	0.20 U



### Table 2-6 Analytical Data Summary Site 21 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I220383001 21-MW-03 09/21/2006	A6I220383002 21-MW-02 09/21/2006
Hexachlorobenzene	ug/L	0.20 U	0.20 U
Hexachlorobutadiene	ug/L	1.0 U	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U	10 U
Hexachloroethane	ug/L	1.0 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U	0.20 U
Isophorone	ug/L	1.0 U	1.0 U
Naphthalene	ug/L	0.20 U	0.20 U
Nitrobenzene	ug/L	1.0 U	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U	1.0 U
Pentachlorophenol	ug/L	5.0 U	5.0 U
Phenanthrene	ug/L	0.20 U	0.20 U
Phenol	ug/L	1.0 U	1.0 U
Pyrene	ug/L	0.20 U	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

 $J = \mbox{Estimated}$  concentration because the result was below the sample reporting limit or quality control criteria were not met.

## Table 2-7 Analytical Data Summary Site 21 Water Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6I220383001 21-MW-03 09/21/2006	A61220383002 21-MW-02 09/21/2006
Arsenic	ug/L	10.0 U	10.0 U
Barium	ug/L	135	119
Cadmium	ug/L	2.0 U	2.0 U
Chromium	ug/L	5.0 U	5.0 U
Lead	ug/L	10.8	3.0 U
Selenium	ug/L	3.3 J	5.0 U
Silver	ug/L	5.0 U	5.0 U
Mercury	ug/L	0.20 U	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

 $<sup>{\</sup>sf J}$  = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

#### 6.11 SITE 22 - NOVA PROPERTIES

According to the Cleveland City Directories, the site was listed as the National Engineering Company from 1954 through 1964. There were no further listings for the site. This site was identified as an OHSPILLS site in the EDR Report.

According to the Cleveland Fire Prevention Bureau files, in October 1953, two 1,000-gallon USTs; gasoline and diesel, were installed at the site. In 1965, the site was owned by the New York Central Rail Road and the ANCO Corporation. The last entry in the file indicates the on-site building was razed in 1974; however, the USTs did not appear to have been properly abandoned.

At the time of the URS Phase I reconnaissance, the site was occupied by an asphalt plant, which is operated by Kenmore Construction. Large chunks of asphalt were observed in piles on the site. Surface staining was observed in multiple locations on the site.

#### 6.11.1 Field Activities

A total of five monitoring wells were scoped for Site 22. Groundwater was encountered in all soil borings; however, one property owner (Kenmore Construction Company, Inc.) would not allow the installation of one monitoring well. Therefore, four monitoring wells (MW02 through MW05) and one soil boring (SB01) were installed. A groundwater sample was collected from SB01 with a bailer inside the augers. Five soil and groundwater samples were collected and analyzed for VOCs, SVOCs, TPH and RCRA Metals. A duplicate groundwater sample was collected from 14-MW05. A Sample Location Map is included as **Figure 6-11A**.

#### 6.11.2 Site-Specific Geology/Hydrogeology

Soils at Site 22 consisted primarily of sand and silty sand with minor amounts of clay, slag, and gravel. Bedrock was not encountered in any of the soil borings, which were advanced to 12 feet bgs.

Groundwater elevations were measured at Site 22 on September 20, 2006. Localized groundwater flow across Site 22 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 22 is to the west-southwest, towards the Cuyahoga River.

#### 6.11.3 Geophysical Survey

During the Phase II reconnaissance, the northeastern portion of the site was occupied by an asphalt plant, which was operated by Kenmore Construction. The eastern portion of the site was occupied by Lafarge North America aggregate operations. Large chunks of asphalt and aggregate were observed in piles across the site. Based on the variability in facility operations and the nature of the material and equipment utilized, a geophysical survey of Site 22 was not possible.

**Phase II Findings** 

#### 6.11.4 Soil Analytical Results

A total of seven VOCs were detected in the soil samples submitted from Site 22. The chemicals detected in the samples submitted were 1,3-dichlorobenzene (1.0 ug/kg), 2-butanone (ranging from 15 ug/kg to 35 ug/kg), acetone (ranging from 28 ug/kg to 190 ug/kg), carbon disulfide (ranging from 0.70 ug/kg to 8.2 ug/kg), methylene chloride (2.3 ug/kg), 2,4-dinitrophenol (130 ug/kg), and 2,6-dinitrotoluene (140 ug/kg). All other VOCs were below the detection limits.

A total of 20 SVOCs were detected in the samples submitted from Site 22. Concentrations of 2-methylnaphthalene (ranging from 400 ug/kg to 760 ug/kg), acenaphthylene (210 ug/kg), anthracene (42 ug/kg and 550 ug/kg), benzaldahyde (27 ug/kg), benzo(a)anthracene (150 ug/kg and 1,600 ug/kg), benzo(a)pyrene (ranging from 19 ug/kg to 1,600 ug/kg), benzo(b)fluoranthene (ranging from 230 ug/kg to 2,200 ug/kg), benzo(g,h,i)perylene (190 ug/kg and 830 ug/kg), benzo(k)fluoranthene (ranging from 18 ug/kg to 790 ug/kg), caprolactam (51 ug/kg), carbazole (220 ug/kg), chrysene (ranging from 180 ug/kg to 1,800 ug/kg), dibenzofuran (85 ug/kg), fluoranthene (ranging from 240 ug/kg to 3,900 ug/kg), fluorene (260 ug/kg), indeno(1,2,3)pyrene (110 ug/kg and 850 ug/kg), isophorone (200 ug/kg), naphthalene (260 ug/kg and 460 ug/kg), phenanthrene (ranging from 240 ug/kg to 1,800 ug/kg), and pyrene (ranging from 240 ug/kg to 3,100 ug/kg) were detected in the soil samples submitted.

Gasoline and diesel range total petroleum hydrocarbons were detected in the soil samples from Site 22. The highest concentrations were found in the diesel range, which is consistent with the concentrations of SVOCs detected across Site 22. A concentration of the light petroleum fraction from 22-MW02-0204 was 12,000 ug/kg. Concentrations of the middle petroleum fraction ranged from 2.3 mg/kg to 1,800 mg/kg. Concentrations of the heavy petroleum fraction ranged from 10 mg/kg to 3,000 mg/kg.

Seven of the eight RCRA Metals were detected in the samples submitted from Site 22. Arsenic, ranging from 2.3 mg/kg to 8.5 mg/kg, was detected in all samples submitted. Barium, ranging from 8.9 mg/kg to 96.4 mg/kg, was detected in the all samples submitted. Cadmium, 0.042 mg/kg to 0.18 mg/kg, was detected in all samples submitted. Chromium, ranging from 3.5 mg/kg to 10.5 mg/kg, was detected in the samples submitted. Lead, ranging from 7.1 mg/kg to 48 mg/kg, was detected in the samples submitted. Selenium, 0.40 mg/kg, was detected in sample 22-SB01-0204. Mercury, ranging from 0.017 mg/kg to 0.32 mg/kg, was detected in the samples submitted.

The analytical results are presented in Table 6-11A.

#### 6.11.5 Groundwater Analytical Results

A total of three VOCs were detected in four of the six groundwater samples submitted from Site 22. The chemicals detected in the samples submitted were 1,1-dichloroethane (0.42 ug/L), 2-butanone (ranging from 0.64 ug/L to 0.97 ug/L), and acetone (ranging from 1.2 ug/L to 3.1 ug/L). All other VOCs were below the detection limits.

A total of eighteen SVOCs were detected in the six samples submitted from Site 22. Concentration of 4-methylphenol (1.0 ug/L), acenaphthene (ranging from 0.23 ug/L to 0.85 ug/L), anthracene (0.23 ug/L and 0.28 ug/L), atrazine (0.71 ug/L), benzo(a)anthracene (0.24 ug/L and 0.63 ug/L), benzo(a)pyrene (ranging from 0.65 ug/L to 1.0 ug/L), benzo(b)fluoranthene (ranging from 0.84 ug/L to 1.0 ug/L), benzo(g,h,i)perylene (0.49 ug/L), benzo(k)fluoranthene (ranging from 0.37 ug/L to 0.94 ug/L), caprolactam (ranging from 0.65 ug/L to 0.77 ug/L), chrysene (0.23 ug/L and 0.74 ug/L), diethyl phthalate (0.86 ug/L), di-n-butyl phthalate (0.79 ug/L), fluoranthene (ranging from 0.25 ug/L to 1.6 ug/L), indeno(1,2,3)pyrene (0.43 ug/L), naphthalene (0.31 ug/L), phenanthrene (0.26 ug/L and 0.68 ug/L), and pyrene (ranging from 0.17 ug/L to 1.1 ug/L) were detected in the groundwater samples submitted.

All eight RCRA Metals were detected in the samples submitted from Site 22. Arsenic, ranging from 4.5 ug/L to 193 ug/L, was detected in all samples submitted. Barium, ranging from 57.1 ug/L to 1,410 ug/L, was detected in all samples submitted. Cadmium, ranging from 3.1 ug/L to 28.2 ug/L, was detected in all samples submitted. Chromium, ranging from 5.3 ug/L to 117 ug/L, was detected in all samples submitted. Lead, ranging from 1.7 ug/L to 6,970 ug/L, was detected in the samples submitted. Selenium, 2.8 ug/L and 9.4 ug/L, was detected in sample 22-MW03 and 22-MW04. Silver was detected in 22-SB01 at 2.2 ug/L. Mercury, 0.11 ug/L and 14.7 ug/L, was detected in 22-SB01 and 22-MW02.

The analytical results are presented in Table 6-11B.

#### 6.11.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on **Table 6-11A** and **6-11B**.

There were no VOCs, TPH, or RCRA metals detected in any of the soil samples from Site 22, which exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

The concentrations of benzo(a)pyrene (1,600 mg/kg) detected in sample 22-SB01-0204 exceeded the BUSTR closure action levels. No other SVOCs detected exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

There were no VOCs detected in any of the groundwater samples from Site 22, which exceeded the OEPA VAP standard for generic unrestricted potable use standards or BUSTR closure action levels.

**Phase II Findings** 

#### **SECTIONSIX**

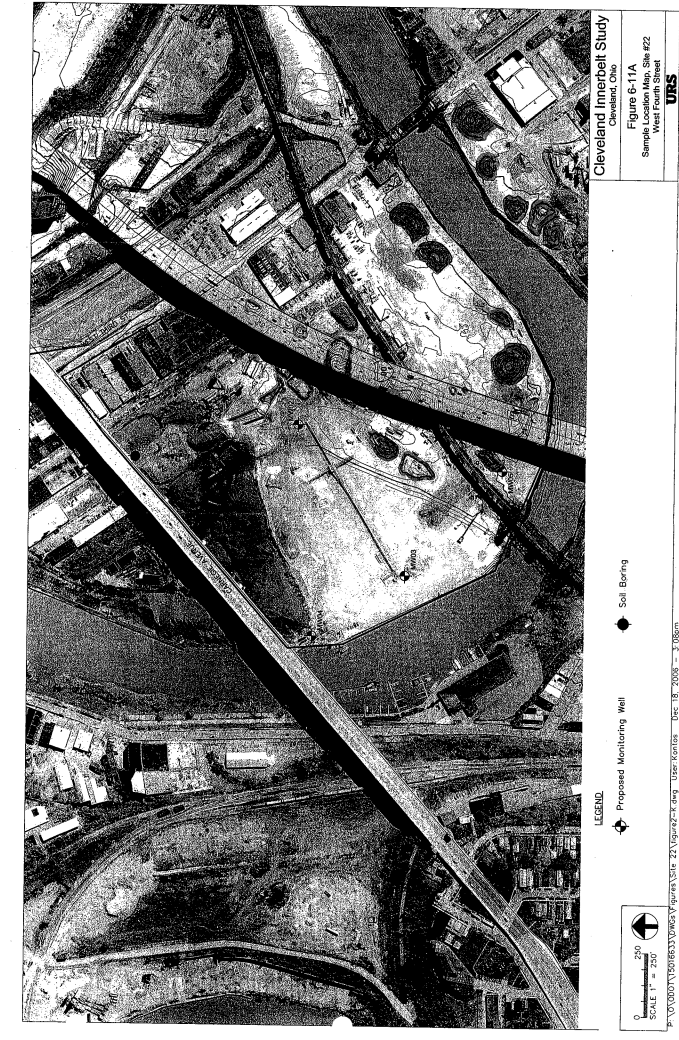
The concentrations of benzo(a)pyrene, benzo(b)fluoranthene, and indeno(1,2,3-cd)pyrene detected in samples 22-MW02, 22-MW04, and 22-SB01 exceeded the OEPA VAP standard for generic unrestricted potable use standards and BUSTR closure action levels.

Eight metals were detected in the groundwater samples (Table 6-11B). Lead was the only metal detected above UPUS. Lead was detected at 6,970 ug/L in a sample collected from 22-SB01 (UPUS is 15 ug/L). The groundwater sample from 22-SB01 was collected from an open boring.

#### 6.11.7 Conclusions

Based on the analytical results, the soil and groundwater at Site 22 may require special disposal and/or worker protection protocols (plan note) during construction activities.





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**TABLES** 

### Summary of Detected Chemicals in Soil Site 22 - Nova Properties ODOT Innerbelt Study Cleveland, Ohio Table w-11A

	Part of the second seco		VAP Commercial/ Industrial (**	Construction	BUSTR 6	22-MW02-0204	22-MW03-0406	22-MW04-0608	22-MW05-0204	22-SB01-0204
	PARAMETER	UNITS	Standard	Standard	Section 1	08/31/2006	08/31/2006	09/01/2006	09/01/2006	08/31/2006
	1,3-Dichlorobenzene	ug/kg	240,000	240,000	1	1.0 J	5.8 UJ	6.1 UU	5.7 U	5.7 W
	2-Butanone	ug/kg	71,600,000	80,000,000	CENTRALINGUAL THAT NAMED IN THE PROPERTY OF TH	77 U	15.0	25 J	35	23 U
s(	Acetone	ug/kg	100,000,000	100,000,000	The second control of the second control of	29 J	80	190 J	190	78
00/	1	ug/kg	720,000	720,000	-	2.0 J	1.2 J	8.2 J	0.71 J	0.70 J
١	- 1	ug/kg	1,300,000	2,300,000	-	19 U	5.8 U	2.3 J	5.7 U	5.7 U
	2,4-Dinitrophenol	ug/kg		-	The state of the s	7600 W	U 0//	1600 U	130 J	3800 U
	2,6-Dinitrotoluene	ug/kg	2,900,000	8,800,000	**************************************	7600 W	U 077	1600 U	140	3800 11
	2-Methylnaphthalene	ug/kg	1	ì	1	7600 UJ	400 J	1600 U	370 U	J. 09Z
	Acenaphthylene	ug/kg	1	1		7600 UJ	J 0/4	1600 U	370 U	2.072
	Anthracene	ug/kg	880,000,000	1,000,000,000	1	7600 UJ	42 J	1600 U	370 U	550
	Benzaldehyde	ug/kg		-	manufic days of all one of companying the discussion of the companying the discussion of the companying the com	7600 UJ	U 077	1600 U	27 ي	3800 0
	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	7600 U	150 J	1600 U	370 U	1600
	Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	360 J	170 5	1600 U	19	1600 J
	Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	430	230 J	1600 U	370 U	2200 J
	Benzo(ghi)perylene	ug/kg	1	-	The state of the s	7600 U	190 ປ	1600 U	370 U	830 J
s	ŝ	ug/kg	000'089	8,100,000	110,000	7600 U	C 29	1600 U	18 J	F 062
00	Caprolactam	ug/kg	•	-	-	7600 UJ	770 U	1600 U	51.0	3800 U
۸S	ş	ug/kg	10,000,000	31,000,000	A proprieto produce per compression per construction per	7600 UJ	U 077	1600 U	370 U	220 J
	Chrysene and Society of an another processing	ug/kg	6,700,000	41,000,000	1,100,000	1300 J	180 J	1600 U	370 U	1800 J
	Dibenzofuran	ug/kg	1	1	And the section of th	7600 UJ	. SS J	1600 U	370 U	3800 U
	Fluoranthene	ug/kg	33,000,000	170,000,000	E	640 J	240 J	1600 U	370 U	3900
	Fluorene	ug/kg	120,000,000	340,000,000	1	7600 UJ	O 077	1600 U	370 U	260 J
	Indeno(1,2,3-cd)pyrene	ug/kg	62,000	410,000	11,000	7600 U	110 J	1600 U	370 U	850 J
	Isophorone	ug/kg	4,600,000	4,600,000	1	7600 UJ	U 077	1600 U	200 J	3800 U
	Naphthalene	ug/kg	530,000	1,900,000	39,800	7600 UJ	260 J	1600 U	370 U	460 J
	Phenanthrene	ug/kg	1	-	The state of the s	350 J	240 J	1600 U	370 U	1800 J
	Pyrene	ug/kg	25,000,000	130,000,000	-	830 J	240 J	1600 U	370 U	3100 J
Н	Gasoline Range Organics (C6-C	ug/kg	- Commence of the Commence of		1,000,000	12000	120 U	120 U	110 U	110 U
qT	C10-C20	mg/kg			2,000	1800	9.7 J	2.3 J	2.4	110 U
I	C20-C34	mg/kg	1	-	5,000	3000	98	20	2	710
	Arsenic	mg/kg	08	210	•	8.4	5.7	8.5	4.9	2.3
•	Barium	mg/kg	200,000	45,000		96.4	40	28.9	8.9 J	49.1
37V	Cadmium	mg/kg	220	420	1	0.17 ئ	0.18 J	0.074 J	0.18 J	0.042 J
/T3	Chromium	mg/kg	8,900	2,000		10.5	5.7	7.2	4.3	3.5
W	Lead	mg/kg		1	1	48	17.5	12	9.2	6
	Selenium	mg/kg	15,000	4,300	1:	0.58 U	0.58 U	0.61 U	U 250	0.40 J
	Mercury	mg/kg	300	84	1	0.32	0.073 J	0.018 J	0.030 J	0.017 J
	Standard not excellable									

-- = Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.
J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and

precisely measure the analyte in the sample.

(1) VAP Generic Direct Contact Soil Standard, Corrunercial/Industrial Land Use (2) VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Summary of Detected Chemicals in Water Site 22 - Nova Properties **ODOT Innerbelt Study** Cleveland, Ohio Table v-11B

<u> </u>	s	00/		SAOCS									METALS																
PARAMETER	1,1-Dichloroethane	2-Butanone	Acetone	4-Methylphenol	Acenaphthene	Anthracene	Atrazine	Benzo(a)anthracene	Benzo(a)pyrene	Benzo(b)fluoranthene	Benzo(ghi)perylene	Benzo(k)fluoranthene	Caprolactam	Chrysene	Diethyl phthalate	Di-n-butyl phthalate	Fluoranthene	Indeno(1,2,3-cd)pyrene	Naphthalene	Phenanthrene	Pyrene	Arsenic	Barium	Cadmium	Chromium	Lead	Selenium	Silver	Mercury
UNITS	ug/L	ug/L	ng/L	ug/L	ug/L	ug/L	ug/L	ug/L	ng/L	ng/L	ng/L	ng/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	ng/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	ng/L	ng/L	ng/L
VAP UPUS/	1400	0890	1600	78	680	2600	က	A CARLONIA MANAGEMENT OF THE ACTION	0.2	The state of the s	The state of the s	and the second s	-	47	13000	1400	370	The state of the s	140		280	20	2000	ß	100	15	20	78	8
BUSTR Closura Action	993666 per pili har i schild open prej hestyren derbit in gest (perseger				•	3	1	0.264	0.2	0.179	-	1.79	1	47	•	- Constitution of Constitution	•	0.23	140	1	•	the state of the s		The state of the s	- Control of the Cont	And the second s		•	:
22-MW-02 09/20/2006	1.0 U	10 U	10 U	1.0 U	0.85	0.20 U	1.0 U	0.24	1.0	1.0	0.20 U	0.94	5.0 U	0.23	1.0 U	1.0 U	0.4	0.20 U	0.20 U	0.26	0.46	7.5 J	214	2.0 U	5.5	20.4	5.0 U	5.0 U	0.11 ט
22-MW-03 09/20/2006	1.0 U	10 U	10 U	1.0 U	0.20 U	0.20 U	1.0 U	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U	0.67 J	0.20 U	1.0 U	1.0 U	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U	4.5 J	57.1 J	2.0 U	4.9 J	3.0 U	9.4	5.0 U	0.20 U
22-MW-04 09/20/2006	1.0 U	0.73 J	1.2 J	1.0	0.84	0.23	1.0 U	0.20 U	0.97	0.95	0.20 U	0.88	5.0 U	0.20 U	1.0 U	ს 67.0	0.25	0.20 U	0.20 U	0.20 U	0.17 J	6.8 J	337	2.0 U	5.3	1.7 J	2.8 J	5.0 U	0.20 ∪
22-MW-05 09/20/2006	1.0 U	0.97 J	2.3 J	1.0 U	0.20 U	0.20 U	1.0 U	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U	0.77 J	0.20 U	D.86 J	1.0 U	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U	50.0 U	855	28.2 J	25.0 U	53.1 J	25.0 U	25.0 U	0.20 U
22-MW-05D 09/20/2006	1.0 U	0.64 J	3.1 J	1.0 U	0.20 U	0.20 U	0.71 <sub>.</sub> J	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U	0.65 J	0.20 U	0.86 J	1.0 U	0.20 U	0.20 U	0.20	0.20 U	0.20 U	20.0 U	688	6.7 J	10.0 U	18.8 J	10.0 U	10.0 U	0.20
22-SB01-083106 08/31/2006	0.42 J	0 1	10 U	1.0 U	0.23	0.28	1.0 U	0.63	0.65	0.84	0.49	0.37	0.65 J	0.74	1.0 U	1.0 U	- 1.6	0.43	0.31	0.68	1:1	193	1410	3.1	117	0269	5.0 U	2.2 J	14.7

<sup>-- =</sup> Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit. J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

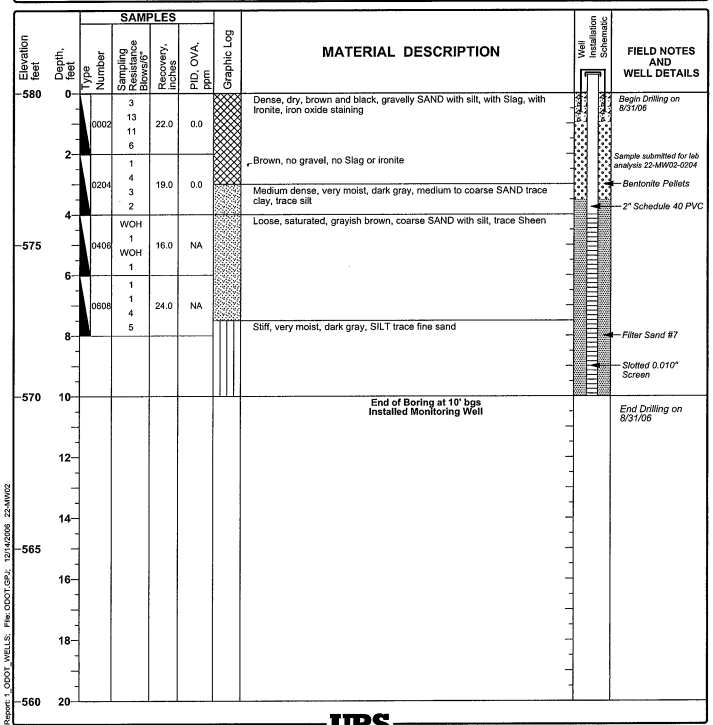
<sup>(1)</sup> VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

**BORING LOGS** 

Project Location: Site 22
Project Number: 15016633

#### Log of Boring 22-MW02

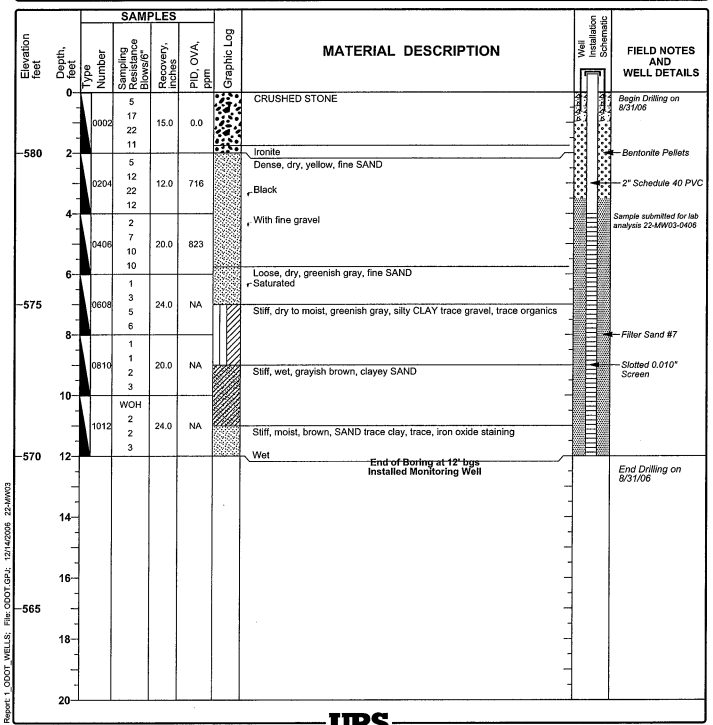
Date(s) 8/31/06	Logged J. Kaminski By	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer 140# auto hammer	Total Depth of Borehole 10.0' bgs
Drill Rig LC-60 Type	Drilling Contractor HAD, Inc.	Approximate Ground Elevation 580'
Location See Site Map	Sampling Method(s) 2" Split Spoon	Borehole Completion Set monitoring well



Project Location: Site 22 Project Number: 15016633

#### Log of Boring 22-MW03

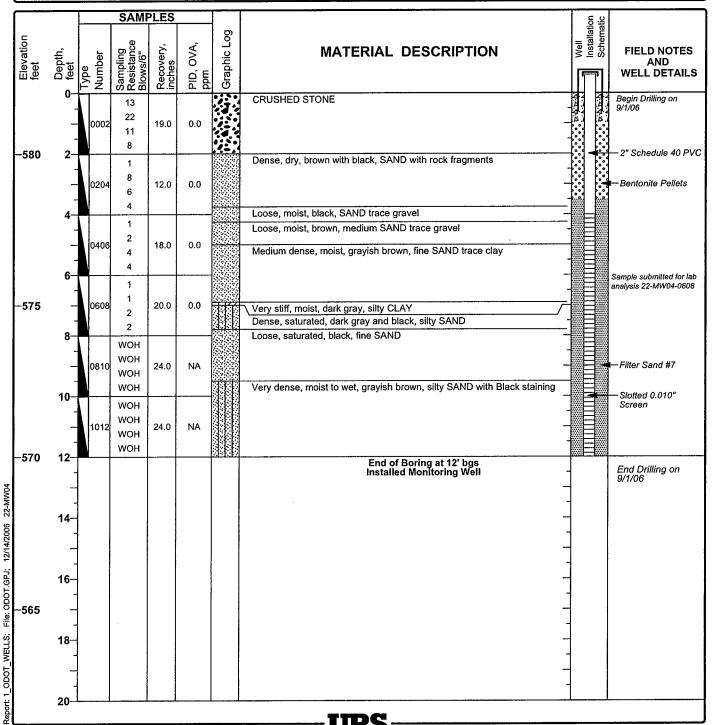
Date(s) 8/31/06 Drilled 8/31/06	Logged By	J. Kaminski	Checked By M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 12.0′ bgs
Drill Rig LC-60 Type LC-60	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 582'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 595.80 on 9/20/06			



Project Location: Site 22 Project Number: 15016633

#### Log of Boring 22-MW04

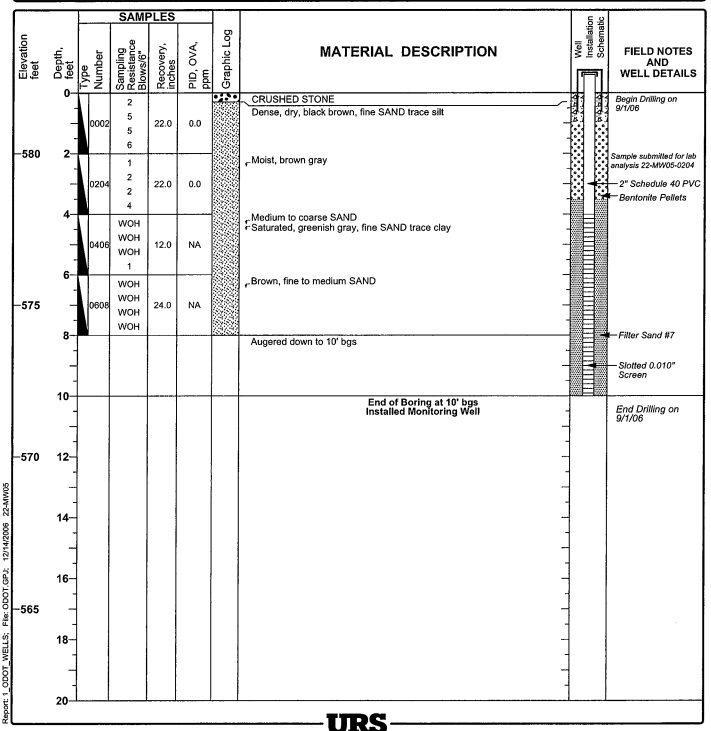
Date(s) 9/1/06 Drilled 9/1/06	Logged J.	Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer 14 Data	40# auto hammer	Total Depth of Borehole 12.0′ bgs
Drill Rig LC-60 Type	Drilling Contractor H	AD, Inc.	Approximate Ground Elevation 582
Location See Site Map	Sampling 2" Method(s)	' Split Spoon	Borehole Set monitoring well



Project Location: Site 22 Project Number: 15016633

#### Log of Boring 22-MW05

Date(s) 9/1/06 Drilled 9/1/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 10.0' bgs
Drill Rig Type LC-60	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 582
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 599.55 on 9/20/06			



**Project Location: Site 22** Project Number: 15016633

### Log of Boring 22-SB01

Date(s) Drilled and Installed 8/31/06	Logged By J. Kaminski	Reviewer M. Wolff
Drilling Method Hollow Stem Auger	Drilling Contractor HAD, Inc.	Total Depth of Borehole 10.0' bgs
Sampling 2" Split Spoon Method	Drill Bit Size/Type: 4-1/4" ID HSA	Approximete Surface Elevation 582'
Drill Rig Type: LC-60	Groundwater NA Level(s)	Hammer 140# auto hammer
Boring Location: See Site Map	Borehole Backfill <b>bentonite</b>	

			CARA	n re		1		1
			SAMI	LES		_		
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	FIELD NOTES
		0002	6	22.0	0.0		Loose to medium dense, dry, black, fine to coarse silty SAND with Crushed asphalt	Begin drilling on 8/31/06
-580	2-	0204	3 5 7 5	20.0	7.5		Dense, dry to moist, brown, fine SAND trace gravel	Sample submitted for lab analysis 22-SB01-0204
	4	0406	1 1 1 3	12.0	NA		r-Wet  Loose, saturated, brown, coarse SAND trace fine gravel	
-575	6	0608	1 2 1	6.0	NA		-With silt, layers -Black stained	
	8 - - 10	0810	1 2 1 1	1.0	NA		- - -	
E70	-						End of Boring at 10' bgs	End drilling on 8/31/06
-570	12-						-	
GPJ; 12/14/200	14-						- - -	
-565 -565	16— - - -						- - -	
Report 1_0D01_BORINGS; File: 0D01.GFJ; 12/14/2006 22:SB01  G	18 - -					:	- - -	
Кероп.	20-1						URS	

DATA ASSESSMENT REPORT

#### Data Assessment Report ODOT Innerbelt Study Site 22 – Nova Properties

Reviewer: P. Schuler Date: November 13, 2006

Five soil samples, six groundwater samples, and one trip blank were collected at the Nova Properties site on W. 4<sup>th</sup> Street in Cleveland, Ohio, from August 31 through September 20, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Re	quested .	Analyse	es <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC	TPH	Met
A6I010325001	22-SB01-0204	08/31/2006	Soil	X	X	X	X
A6I010325002	22-MW02-0204	08/31/2006	Soil	X	X	X	X
A6I010325003	22-MW03-0406	08/31/2006	Soil	X	X	X	X
A6I010325004	22-MW04-0608	09/01/2006	Soil	X	X	X	X
A6I010325005	22-MW05-0204	09/01/2006	Soil	X	X	X	X
A6I010325006	22-SB01-083106	08/31/2006	Groundwater	X	X		X
A6I220382001	22-MW-02	09/20/2006	Groundwater	X	X		X
A6I220382002	22-MW-05	09/20/2006	Groundwater	X	X		X
A6I220382003	22-MW-05D	09/20/2006	Groundwater	X	X		X
A6I220382004	22-MW-03	09/20/2006	Groundwater	X	X		X
A6I220382005	22-MW-04	09/20/2006	Groundwater	X	X		X
A6I220382006	TB-092006	09/20/2006	Trip Blank	X			

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B]

SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

Met = RCRA Metals [SW-846 Methods 6010B/6020/7470A/7471A]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

Positive detections for bis(2-ethylhexyl)phthalate in soil sample 22-SB01-0204 and in all six groundwater samples, as well as gasoline range organics in samples 22-SB01-0204, 22-MW03-0406, 22-MW04-0608, and 22-MW05-0204, were qualified as nondetect ("U") due to the presence of the analytes in the method blanks at similar concentrations.

- Positive detections for toluene in water samples 22-MW-02 and 22-MW-04 were qualified as nondetect ("U") due to the presence of toluene in the associated method trip blank at a similar concentration.
- The trip blank submitted with the 08/31/06 and 09/01/06 samples were logged in and reported with samples from other sites. A trace amount of toluene was detected in the trip blank. Therefore, the positive toluene result for sample 22-MW04-0608 was qualified as nondetect ("U"), since the sample concentrations were not significantly higher than those in the associated blanks.
- The laboratory "J" flags on selected chromium and barium soil sample results, indicating that the analytes were detected in the method blank, were removed in the final data set, since the sample results were greater than five times the blank concentrations. The sample results are considered representative of site conditions and any contribution due to external contamination is negligible.
- One or more volatile internal standard responses were outside of the acceptance range in samples 22-SB01-0204, 22-MW03-0406, and 22-MW04-0608. The results for all volatile analytes quantified from each of the noncompliant internal standards were qualified as estimated ("J" or "UJ").
- The responses for four semivolatile internal standards were outside of the acceptance range in sample 22-MW02-0204. The results for all semivolatile analytes quantified from each of the noncompliant internal standards were qualified as estimated ("J" or "UJ").
- The results for cadmium and lead in field duplicate samples 22-MW-05 and 22-MW-05D were qualified as estimated ("J") due to poor precision between the results.
- The lab reported some metals results with a "G" qualifier, indicating that a dilution was required due to matix interferences. The "G" qualifiers were removed in the final data set, but the sample reporting limits are elevated accordingly.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J" for organics, and "B" for inorganics), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, organic "J" qualifiers were retained with the numeric results. The inorganic "B" qualifiers were changed to "J" qualifiers for the sake of consistency throughout the package.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-7.

### Table 2-1 Analytical Data Summary Site 22 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I010325001 22-SB01-0204 08/31/2006	A6I010325002 22-MW02-0204 08/31/2006	A6I010325003 22-MW03-0406 08/31/2006	A6I010325004 22-MW04-0608 09/01/2006	A6I010325005 22-MW05-0204 09/01/2006
1,1,1-Trichloroethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1.1.2.2-Tetrachloroethane	ug/kg	5.7 UJ	19 U	5.8 UJ	6.1 UJ	5.7 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,1,2-Trichloroethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,1-Dichloroethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,1-Dichloroethene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,2,4-Trichlorobenzene	ug/kg	5.7 UJ	19 U	5.8 UJ	6.1 UJ	5.7 U
1,2-Dibromo-3-chloropropane	ug/kg	11 UJ	38 U	12 UJ	12 UJ	11 U
1,2-Dibromoethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,2-Dichlorobenzene	ug/kg	5.7 UJ	19 U	5.8 UJ	6.1 UJ	5.7 U
1,2-Dichloroethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,2-Dichloropropane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
1,3-Dichlorobenzene	ug/kg	5.7 UJ	1.0 J	5.8 UJ	6.1 UJ	5.7 U
1,4-Dichlorobenzene	ug/kg	5.7 UJ	19 U	5.8 UJ	6.1 UJ	5.7 U
2-Butanone	ug/kg	23 U	77 U	15 J	25 J	35
2-Hexanone	ug/kg	23 U	77 U	23 U	25 UJ	23 U
4-Methyl-2-pentanone	ug/kg	23 UJ	77 U	23 UJ	25 UJ	23 U
Acetone	ug/kg	28	29 J	80	190 J	190
Benzene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Bromodichloromethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Bromoform	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Bromomethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Carbon disulfide	ug/kg	0.70 J	2.0 J	1.2 J	8.2 J	0.71 J
Carbon tetrachloride	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Chlorobenzene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Chloroethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Chloroform	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Chloromethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
cis-1,2-Dichloroethene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
cis-1,3-Dichloropropene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Cyclohexane	ug/kg	11 U	38 U	12 U	12 UJ	11 U
Dibromochloromethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Dichlorodifluoromethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Ethylbenzene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Isopropylbenzene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Methyl acetate	ug/kg	11 U	38 U	12 U	12 UJ	11 U
Methyl tert-butyl ether	ug/kg	23 U	77 U	23 U	25 UJ	23 U
Methylcyclohexane	ug/kg	11 U	38 U	12 U	12 UJ	11 U
Methylene chloride	ug/kg	5.7 U	19 U	5.8 U	2.3 J	5.7 U
Styrene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Tetrachloroethene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Toluene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
trans-1,2-Dichloroethene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
trans-1,3-Dichloropropene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Trichloroethene	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Trichlorofluoromethane	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Vinyl chloride	ug/kg	5.7 U	19 U	5.8 U	6.1 UJ	5.7 U
Xylenes (total)	ug/kg	11 U	38 U	12 U	12 UJ	11 U

<sup>m /U</sup> =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-2 Analytical Data Summary Site 22 Soil Semivolatiles ODOT Innerbelt Study

		A6I010325001 22-SB01-0204	A6I010325002 22-MW02-0204	A6I010325003 22-MW03-0406	A6I010325004 22-MW04-0608	A6I010325005 22-MW05-0204
PARAMETER	UNITS	08/31/2006	08/31/2006	08/31/2006	09/01/2006	09/01/2006
1,1'-Biphenyl	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,2'-oxybis(1-Chloropropane)	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,4,5-Trichlorophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,4,6-Trichlorophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,4-Dichlorophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,4-Dimethylphenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,4-Dinitrophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	130 J
2,4-Dinitrotoluene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2,6-Dinitrotoluene	ug/kg	3800 U	7600 UJ	770 U	1600 U	140 J
2-Chloronaphthalene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2-Chlorophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2-Methylnaphthalene	ug/kg	760 J	7600 UJ	400 J	1600 U	370 U
2-Methylphenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2-Nitroaniline	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
2-Nitrophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
3,3'-Dichlorobenzidine	ug/kg	3800 U	7600 U	770 U	1600 U	370 U
3-Nitroaniline	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4,6-Dinitro-2-methylphenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Bromophenyl phenyl ether	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Chloro-3-methylphenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Chloroaniline	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Chlorophenyl phenyl ether	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Methylphenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Nitroaniline	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
4-Nitrophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Acenaphthene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 ∪
Acenaphthylene	ug/kg	210 J	7600 UJ	770 U	1600 U	370 U
Acetophenone	ug/kg	760 U	1500 UJ	150 U	330 U	76 U
Anthracene	ug/kg	550 J	7600 UJ	42 J	1600 U	370 U
Atrazine	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Benzaldehyde	ug/kg	3800 U	7600 UJ	770 U	1600 U	27 J
Benzo(a)anthracene	ug/kg	1600 J	7600 U	150 J	1600 U	370 U
Benzo(a)pyrene	ug/kg	1600 J	360 J	170 J	1600 U	19 J
Benzo(b)fluoranthene	ug/kg	2200 J	430 J	230 J	1600 U	370 U
Benzo(ghi)perylene	ug/kg	830 J	7600 U	190 J	1600 U	370 U
Benzo(k)fluoranthene	ug/kg	790 J	7600 U	67 J	1600 U	18 J
bis(2-Chloroethoxy)methane	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
bis(2-Chloroethyl) ether	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
bis(2-Ethylhexyl) phthalate	ug/kg	3800 U	7600 U	770 U	1600 U	370 U
Butyl benzyl phthalate	ug/kg	3800 U	7600 U	770 U	1600 U	370 U
Caprolactam	ug/kg	3800 U	7600 UJ	770 U	1600 U	51 J
Carbazole	ug/kg	220 J	7600 UJ	770 U	1600 U	370 U
Chrysene	ug/kg	1800 J	1300 J	180 J	1600 U	370 U
Dibenz(a,h)anthracene	ug/kg	3800 U	7600 U	770 U	1600 U	370 U
Dibenzofuran	ug/kg	3800 U	7600 UJ	85 J	1600 U	370 U
Diethyl phthalate	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Dimethyl phthalate	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Di-n-butyl phthalate	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Di-n-octyl phthalate	ug/kg	3800 U	7600 U	770 U	1600 U	370 U
Fluoranthene	ug/kg	3900	640 J	240 J	1600 U	370 U
Fluorene	ug/kg	260 J	7600 UJ	770 U	1600 U	370 ∪
Hexachlorobenzene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U

#### Table 2-2 Analytical Data Summary Site 22 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I010325001 22-SB01-0204 08/31/2006	A6I010325002 22-MW02-0204 08/31/2006	A6I010325003 22-MW03-0406 08/31/2006	A6I010325004 22-MW04-0608 09/01/2006	A6I010325005 22-MW05-0204 09/01/2006
Hexachlorobutadiene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Hexachlorocyclopentadiene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Hexachloroethane	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Indeno(1,2,3-cd)pyrene	ug/kg	850 J	7600 U	110 J	1600 U	370 U
Isophorone	ug/kg	3800 U	7600 UJ	770 U	1600 U	200 J
Naphthalene	ug/kg	460 J	7600 UJ	260 J	1600 U	370 U
Nitrobenzene	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
N-Nitrosodi-n-propylamine	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
N-Nitrosodiphenylamine	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Pentachlorophenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Phenanthrene	ug/kg	1800 J	350 J	240 J	1600 U	370 U
Phenol	ug/kg	3800 U	7600 UJ	770 U	1600 U	370 U
Pyrene	ug/kg	3100 J	830 J	240 J	1600 U	370 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

### Table 2-3 Analytical Data Summary Site 22 Soil TPH ODOT Innerbelt Study

#### A6I010325001 A6I010325002 A6I010325003 A6I010325004 A6I010325005 22-SB01-0204 22-MW02-0204 22-MW03-0406 22-MW04-0608 22-MW05-0204 09/01/2006 08/31/2006 08/31/2006 08/31/2006 09/01/2006 PARAMETER UNITS Gasoline Range Organics (C6-C12) ug/kg 110 U 12000 120 U 120 U 110 U C10-C20 mg/kg 110 U 1800 9.7 J 2.3 J 2.4 C20-C34 mg/kg 710 3000 86 20 10

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

## Table 2-4 Analytical Data Summary Site 22 Soil Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6I010325001 22-SB01-0204 08/31/2006	A6I010325002 22-MW02-0204 08/31/2006	A6I010325003 22-MW03-0406 08/31/2006	A6I010325004 22-MW04-0608 09/01/2006	A6I010325005 22-MW05-0204 09/01/2006
Percent Solids	%	88	86.8	86.2	81.5	88.2
Arsenic	mg/kg	2.3	8.4	5.7	8.5	4.9
Barium	mg/kg	49.1	96.4	40	28.9	8.9 J
Cadmium	mg/kg	0.042 J	0.17 J	0.18 J	0.074 J	0.18 J
Chromium	mg/kg	3.5	10.5	5.7	7.2	4.3
Lead	mg/kg	9.0	48	17.5	7.1	9.2
Selenium	mg/kg	0.40 J	0.58 U	0.58 U	0.61 U	0.57 U
Silver	mg/kg	0.57 U	0.58 U	0.58 U	0.61 U	0.57 U
Mercury	mg/kg	0.017 J	0.32	0.073 J	0.018 J	0.030 J

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

# Table 2-5 Analytical Data Summary Site 22 Water Volatiles ODOT Innerbelt Study

		A6I010325006 22-SB01-083106	A6I220382001 22-MW-02	A61220382002 22-MW-05	A61220382003 22-MW-05D	A6I220382004 22-MW-03
PARAMETER	UNITS	08/31/2006	09/20/2006	09/20/2006	09/20/2006	09/20/2006
1,1,1-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,1-Dichloroethane	ug/L	0.42 J	1.0 U	1.0 U	1.0 U	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 Ų	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
2-Butanone	ug/L	10 U	10 U	0.97 J	0.64 J	10 U
2-Hexanone	ug/L	10 U	10 U	10 U	10 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U	10 U	10 U	10 U
Acetone	ug/L	10 U	10 U	2.3 J	3.1 J	10 U
Benzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chloroethane	ug/L ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chloroform	5 S S S S S S S S S S S S S S S S S S S	1.0 U	1.0 U	1.0 U	1.0 U	the contract of the second section is
Chloromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
A COLOR DE CONTRACTO DE CASA DE CONTRACTO DE	ug/L	1.0 U	1.0 U	1.0 U		1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U			1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	A 200 DESCRIPTION OF THE PARTY	1.0 U	1.0 U	1.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Dibromochloromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U	10 U	10 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	, 1.0 U
Methylene chloride	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Styrene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Toluene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	1,0 U	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

## Table 2-5 Analytical Data Summary Site 22 Water Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I220382005 22-MW-04 09/20/2006	A6I220382006 TB-092006 09/20/2006
1,1,1-Trichloroethane	ug/L	1.0 U	1.0 U
• •	ug/L ug/L	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane		1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L		- "
1,1-Dichloroethane	ug/L	1.0 U	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U
2-Butanone	ug/L	0.73 J	10 U
2-Hexanone	ug/L	10 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U
Acetone	ug/L	1.2 J	10 U
Benzene	ug/L	1.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U
Carbon disulfide	ug/L_	1.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U
Chloroethane	ug/L	1.0 U	1.0 U
Chloroform	ug/L	1.0 U	1.0 U
Chloromethane	ug/L	1.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	1.0 U
Dibromochloromethane	ug/L	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U
Methylcyclohexane	ug/L ug/L	1.0 U	1.0 U
Methylene chloride	ug/L	1.0 U	1.0 U
and the contract of the contra	ug/L ug/L	1.0 U	1.0 U
Styrene Tetrachloroethene	ug/L ug/L	1.0 U	1.0 U
A CONTRACTOR OF THE CONTRACTOR		1.0 U	1.0
Toluene	ug/L	and the second of the second of the second of	en a company of the second
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	0.57 J

U =The analyte was analyzed for, but was not detected. Valu



 $<sup>{\</sup>bf J}={\bf E}{\bf s}$ timated concentration because the result was below th

Table 2-6 Analytical Data Summary Site 22 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I010325006 22-SB01-083106 08/31/2006	A6I220382001 22-MW-02 09/20/2006	A61220382002 22-MW-05 09/20/2006	A61220382003 22-MW-05D 09/20/2006	A61220382004 22-MW-03 09/20/2006
1,1'-Biphenyl	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
2,4,5-Trichlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
2,4,6-Trichlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
The first of the second of the	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
2,4-Dichlorophenol	ug/L ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
2,4-Dimethylphenol	ug/L ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
2,4-Dinitrophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
2,4-Dinitrotoluene	ug/L ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
2,6-Dinitrotoluene		1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
2-Chloronaphthalene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
2-Chlorophenol	ug/L		0.20 U	0.20 U	0.20 U	0.20 U
2-Methylnaphthalene	ug/L	0.20 U		1.0 U	1.0 U	1.0 U
2-Methylphenol	ug/L	1.0 U	1.0 U		2.0 U	2.0 U
2-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
2-Nitrophenol	ug/L	2.0 U	2.0 U	2.0 U		4.
3,3'-Dichlorobenzidine	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
3-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
4,6-Dinitro-2-methylphenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
4-Bromophenyl phenyl ether	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
4-Chloro-3-methylphenol	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
4-Chloroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
4-Chlorophenyl phenyl ether	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
4-Methylphenol	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
4-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U	2.0 U	2.0 U
4-Nitrophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
Acenaphthene	ug/L	0.23	0.85	0.20 U	0.20 U	0.20 U
Acenaphthylene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U
Acetophenone	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Anthracene	ug/L	0.28	0.20 U	0.20 U	0.20 U	0.20 U
Atrazine	ug/L	1.0 U	1.0 U	1.0 U	0.71 J	1.0 U
Benzaldehyde	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Benzo(a)anthracene	ug/L	0.63	0.24	0.20 U	0.20 U	0.20 U
Benzo(a)pyrene	ug/L	0.65	1	0.20 U	0.20 U	0.20 U
Benzo(b)fluoranthene	ug/L	0.84	1	0.20 U	0.20 U	0.20 U
Benzo(ghi)perylene	ug/L	0.49	0.20 U	0.20 U	0.20 U	0.20 U
Benzo(k)fluoranthene	ug/L	0.37	0.94	0.20 U	0.20 U	0.20 U
bis(2-Chloroethoxy)methane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
bis(2-Chloroethyl) ether	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
bis(2-Ethylhexyl) phthalate	ug/L	1.8 U	3.9 U	3.1 ∪	1.6 U	1.2 U
Butyl benzyl phthalate	ug/L	1.0 U	1.0 U	1.0 ∪	1.0 U	1.0 U
Caprolactam	ug/L	0.65 J	5.0 U	0.77 J	0.65 J	0.67 J
Carbazole	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Chrysene	ug/L	0.74	0.23	0.20 U	0.20 U	0.20 U
and the second of the second o	ug/L	0.20 U	0.20 U	0.20 U	0.20 U	0.20 U
Dibenz(a,h)anthracene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Dibenzofuran Diethyl obtholote	ug/L ug/L	1.0 U	1.0 U	0.86 J	0.86 J	1.0 U
Diethyl phthalate	1	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Dimethyl phthalate	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Di-n-butyl phthalate	ug/L	10.00	1.0 U	1.0 U	1.0 U	1.0 U
Di-n-octyl phthalate	ug/L	1.0 U		the state of the s	0.20 U	0.20 U
Fluoranthene	ug/L	1.6	0.4	0.20 U		
Fluorene	ug/L	0.20 U	0.20 U 0.20 U	0.20 U 0.20 U	0.20 U 0.20 U	0.20 U 0.20 U

Table 2-6 Analytical Data Summary Site 22 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6I010325006 22-SB01-083106 08/31/2006	A6I220382001 22-MW-02 09/20/2006	A6I220382002 22-MW-05 09/20/2006	A6I220382003 22-MW-05D 09/20/2006	A6I220382004 22-MW-03 09/20/2006
Hexachlorobutadiene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U	10 U	10 U	10 U	10 U
Hexachloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.43	0.20 U	0.20 U	0.20 U	0,20 U
Isophorone	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Naphthalene	ug/L	0.31	0.20 U	0.20 U	0.20 U	0.20 U
Nitrobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U	1.0 U	1.0 ∪	1.0 U	1.0 U
Pentachlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U	5.0 U
Phenanthrene	ug/L	0.68	0.26	0.20 U	0.20 U	0.20 U
Phenol	ug/L	1.0 U	1.0 U	1.0 U	1.0 U	1.0 U
Pyrene	ug/L	1.1	0.46	0.20 U	0.20 U	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-6 Analytical Data Summary Site 22 Water Semivolatiles ODOT Innerbelt Study

		A61220382005 22-MW-04
PARAMETER	UNITS	09/20/2006
1,1'-Biphenyl	ug/L	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/L	1.0 U
2,4,5-Trichlorophenol	ug/L	5.0 U
2,4,6-Trichlorophenol	ug/L	5.0 U
2,4-Dichlorophenol	ug/L	2.0 U
2,4-Dimethylphenol	ug/L	2.0 U
2,4-Dinitrophenol	ug/L	5.0 U
2,4-Dinitrotoluene	1	5.0 U
2,4-Dinitrotoluene	ug/L	5.0 U
	ug/L	l .
2-Chloronaphthalene	ug/L	1.0 U
2-Chlorophenol	ug/L	1.0 U
2-Methylnaphthalene	ug/L	0.20 U
2-Methylphenol	ug/L	1.0 U
2-Nitroaniline	ug/L	2.0 U
2-Nitrophenol	ug/L	2.0 U
3,3'-Dichlorobenzidine	ug/L	5.0 U
3-Nitroaniline	ug/L	2.0 U
4,6-Dinitro-2-methylphenol	ug/L	5.0 U
4-Bromophenyl phenyl ether	ug/L	2.0 U
4-Chloro-3-methylphenol	ug/L	2.0 U
4-Chloroaniline	ug/L	2.0 U
4-Chlorophenyl phenyl ether	ug/L	2.0 U
4-Methylphenol	ug/L	1
4-Nitroaniline	ug/L	2.0 U
4-Nitrophenol	ug/L	5.0 U
Acenaphthene	ug/L	0.84
Acenaphthylene	ug/L	0.20 U
Acetophenone	ug/L	1.0 U
Anthracene	ug/L	0.23
Atrazine	ug/L	1.0 U
Benzaldehyde	ug/L	1.0 U
and the matter of the control of the		0.20 U
Benzo(a)anthracene	ug/L	0.20 0
Benzo(a)pyrene	ug/L	and the second
Benzo(b)fluoranthene	ug/L	0.95
Benzo(ghi)perylene	ug/L	0.20 U
Benzo(k)fluoranthene	ug/L	0.88
bis(2-Chloroethoxy)methane	ug/L	1.0 U
bis(2-Chloroethyl) ether	ug/L	1.0 U
bis(2-Ethylhexyl) phthalate	ug/L	2.4 U
Butyl benzyl phthalate	ug/L	1.0 U
Caprolactam	ug/L	5.0 U
Carbazole	ug/L	1.0 U
Chrysene	ug/L	0.20 U
Dibenz(a,h)anthracene	ug/L	0.20 U
Dibenzofuran	ug/L	1.0 U
Diethyl phthalate	ug/L	1.0 U
Dimethyl phthalate	ug/L	1.0 U
Di-n-butyl phthalate	ug/L	0.79 J
Di-n-octyl phthalate	ug/L	1.0 U
Fluoranthene	ug/L	0.25
Fluorene	ug/L	0.20 U
Hexachlorobenzene	ug/L	0.20 U

Table 2-6 Analytical Data Summary Site 22 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A61220382005 22-MW-04 09/20/2006
Hexachlorobutadiene	ug/L	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U
Hexachloroethane	ug/L	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U
Isophorone	ug/L	1.0 U
Naphthalene	ug/L	0.20 U
Nitrobenzene	ug/L	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U
Pentachlorophenol	ug/L	5.0 U
Phenanthrene	ug/L	0.20 U
Phenol	ug/L	1.0 U
Pyrene	ug/L	0.17 J

U =The analyte was analyzed for, but was not detected

J = Estimated concentration because the result was be

### Table 2-7 Analytical Data Summary Site 22 Water Metals ODOT Innerbelt Study

PARAMETER	UNITS	A6I010325006 22-SB01-083106 08/31/2006	A6I220382001 22-MW-02 09/20/2006	A61220382002 22-MW-05 09/20/2006	A6I220382003 22-MW-05D 09/20/2006	A61220382004 22-MW-03 09/20/2006	A6I220382005 22-MW-04 09/20/2006
Arsenic	ug/L	193	7.5 J	50.0 U	20.0 U	4.5 J	6.8 J
Barium	ug/L	1410	214	855	688	57.1 J	337
Cadmium	ug/L	3.1	2.0 U	28.2 J	6.7 J	2.0 U	2.0 U
Chromium	ug/L	117	5.5	25.0 G U	10.0 G U	4.9 J	5.3
Lead	ug/L	6970	20.4	53.1 J	18.8 J	3.0 U	1.7 J
Selenium	ug/L	5.0 U	5.0 U	25.0 U	10.0 U	9.4	2.8 J
Silver	ug/L	2.2 J	5.0 U	25.0 U	10.0 U	5.0 U	5.0 U
Mercury	ug/L	14.7	0.11 J	0.20 U	0.20 ∪	0.20 U	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

#### 6.12 SITE 23 - NS RAILROAD BUILDING

The Cleveland City Directories for 1954 through 1999, in five-year intervals, were reviewed for information regarding the site. There were no listings for the site.

According to the Cleveland Fire Prevention Bureau files, the site is currently occupied by a storage building for the Rail Road. There reportedly is a 500-gallon gasoline UST on the site.

At the time of the URS Phase I reconnaissance, the site was occupied by a railroad storage building. Multiple piles of tires, roofing materials, lumber, and construction debris were observed on the site. Surface staining and oily sheens were observed on the ground. Drums, which were labeled to contain soil cuttings, were located on the site.

#### 6.12.1 Field Activities

A total of four monitoring wells and six surface samples were proposed for Site 23. Groundwater was encountered in all soil borings during field activities and monitoring wells were installed. A total of eleven soil samples (surface and subsurface) and four groundwater samples were collected and analyzed for VOCs, SVOCs, TPH and RCRA Metals. A duplicate soil sample was collected from 23-MW01-2224. A Sample Location Map is included as **Figure 6-12A**.

#### 6.12.2 Site-Specific Geology/Hydrogeology

Soils at Site 23 consisted primarily of sand, silty sand, and silty clay with minor amounts of slag and gravel. Bedrock was not encountered in any of the soil borings, which were advanced to 30 feet bgs.

Groundwater elevations were measured at Site 23 on August 10, 2006. Localized groundwater flow across Site 23 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 23 is to the south-southwest, towards the Cuyahoga River.

#### 6.12.3 Geophysical Survey

The geophysical survey at Site 23 included the asphalt-paved parking and drive surfaces northeast and northwest of the building. The remaining property was fenced and inaccessible. The investigation was completed in a single set of transects aligned parallel or perpendicular to the building in a northeast-southwest orientation. Paint marks were plotted on the pavement to align the transects.

The survey identified four areas of anomalously high magnetic gradients. These areas are shown on **Figure 6-12B** as anomalies A through D as described below:

A. The anomaly labeled "A" is related to a manhole.

**SECTIONSIX** 

- B. Several anomalies were detected along the northwest edge of the building and the fence to the rear of the building. These anomalies are assumed to be related to utility connections at the building and the steel, chain-linked fence.
- C. An anomaly was noted at the footer for of a former AST and is likely related to the presence of reinforcing steel.
- D. An anomaly was observed at the southeastern portion of the Site that is assumed to be related to a water line and water connections.

No evidence of a UST was discovered as a result of the geophysical survey at Site 23.

### 6.12.4 Soil Analytical Results

A total of eighteen VOCs were detected in five soil samples submitted from Site 23. The chemicals detected in the samples submitted were 1,1,1-trichlorothane (25 ug/kg), 1,1-dichloroethene (1.8 ug/kg), 1,2,4-trichlorobenzene (6.0 ug/kg), 1,2-dichlorobenzene (1.6 ug/kg), 1,3-dichlorobenzene (1.6 ug/kg), 1,4-dichlorobenzene (1.9 ug/kg), 2-butanone (ranging from 1.5 ug/kg to 8.8 ug/kg), acetone (ranging from 4.5 ug/kg to 45 ug/kg), benzene (ranging from 0.37 ug/kg to 1.5 ug/kg), carbon disulfide (ranging from 0.57 ug/kg to 11 ug/kg), cyclohexane (0.86 ug/kg and 1.5 ug/kg), ethylbenzene (ranging from 1.6 ug/kg and 11 ug/kg), isopropylbenzene (ranging from 0.80 ug/kg to 15 ug/kg), methylcyclohexane (2.7 ug/kg and 4.5 ug/kg), methylene chloride (5.5 ug/kg and 7.9 ug/kg), styrene (1.6 ug/kg), toluene (ranging from 0.45 ug/kg to 6.5 ug/kg), and total xylenes (ranging from 4.5 ug/kg to 25 ug/kg). All other VOCs were below the detection limits.

A total of 27 SVOCs were detected in the samples submitted from Site 23. Concentrations of 1,1'-biphenyl (ranging from 40 ug/kg to 2,800 ug/kg), 2-methylnaphthalene (ranging from 41 ug/kg to 11,000 ug/kg), 4-methylphenol / 3-methylphenol (72 ug/kg), acenaphthene (ranging from 12 ug/kg to 21,000 ug/kg), acenaphthylene (ranging from 24 ug/kg to 140 ug/kg), acetophenone (380 ug/kg), anthracene (ranging from 18 ug/kg to 16,000 ug/kg), benzaldahyde (2,900 ug/kg), benzo(a)anthracene (ranging from 59 ug/kg to 11,000 ug/kg), benzo(a)pyrene (ranging from 57 ug/kg to 5,300 ug/kg), benzo(b)fluoranthene (ranging from 84 ug/kg to 8,000 ug/kg), benzo(g,h,i)perylene (ranging from 49 ug/kg to 1,900 ug/kg), benzo(k)fluoranthene (ranging from 30 ug/kg to 3,200 ug/kg), bis(2-ethylhexyl)phthalate (ranging from 31 ug/kg to 5,600 ug/kg), caprolactam (45 ug/kg), carbazole (ranging from 33 ug/kg to 10,000 ug/kg), chrysene (ranging from 69 ug/kg to 9,500 ug/kg), dibenz(a,h)anthracene (ranging from 13 ug/kg to 320 ug/kg), dibenzofuran (ranging from 130 ug/kg to 13,000 ug/kg), di-n-butyl phthalate (120 ug/kg), fluoranthene (ranging from 110 ug/kg to 46,000 ug/kg), fluorene (ranging from 28 ug/kg to 22,000 ug/kg), indeno(1,2,3)pyrene (ranging from 38 ug/kg to 1,800 ug/kg), naphthalene (ranging from 16 ug/kg to 31,000 ug/kg), phenanthrene (ranging from 76 ug/kg to 74,000 ug/kg), phenol (220 ug/kg), and pyrene (ranging from 100 ug/kg to 34,000 ug/kg) were detected in the soil samples submitted.

Gasoline and diesel range total petroleum hydrocarbons were detected in the soil samples from Site 23. The highest concentrations were found in the diesel range, which is consistent with the

**Phase II Findings** 

concentrations of SVOCs detected across Site 23. A concentration of the light petroleum fraction from 23-MW01-2224 was 88 ug/kg and from 23-MW02-1416 was 5,600 ug/kg. Concentrations of the middle petroleum fraction ranged from 7.2 mg/kg to 100 mg/kg. Concentrations of the heavy petroleum fraction ranged from 16 mg/kg to 2,000 mg/kg.

Seven of the eight RCRA Metals were detected in the samples submitted from Site 23. Arsenic, ranging from 2.6 mg/kg to 18.6 mg/kg, was detected in all the samples submitted. Barium, ranging from 47 mg/kg to 311 mg/kg, was detected in all the samples submitted. Cadmium, 0.20 mg/kg to 2.5 mg/kg, was detected in all the samples submitted. Chromium, ranging from 5.5 mg/kg to 364 mg/kg, was detected in all the samples submitted. Lead, ranging from 6.6 mg/kg to 761 mg/kg, was detected in all the samples submitted. Silver, 0.50 mg/kg, was detected in sample 23-SS01. Mercury, ranging from 0.020 mg/kg to 0.16 mg/kg, was detected in all the samples submitted.

The analytical results are presented in Table 6-12A.

## 6.12.5 Groundwater Analytical Results

A total of four VOCs were detected in three of the four groundwater samples submitted from Site 23. The chemicals detected in the samples submitted were 1,1,1-trichloroethane (62 ug/L), 1,1-dichloroethane (1.1 ug/L and 4.9 ug/L), 1,1-dichloroethene (0.47 ug/L), and toluene (0.25 ug/L). All other VOCs were below the detection limits.

A total of two SVOCs were detected in 23-MW02 from Site 23. Concentrations of 2-methylnaphthalene (0.35 ug/L) and acenaphthene (0.62 ug/L) were detected in the groundwater sample.

Five of the eight RCRA metals were detected in the samples submitted from Site 23. Arsenic, ranging from 5.7 ug/L to 6.1 ug/L, was detected in the samples submitted. Barium, ranging from 47.4 ug/L to 59.5 ug/L, was detected in the samples submitted. Chromium, ranging from 3.5 ug/L to 4.0 ug/L, was detected in the samples submitted. Lead, ranging from 2.0 ug/L to 3 ug/L, was detected in the samples submitted. Selenium, 5.3 ug/L, was detected in sample 23-MW04.

The analytical results are presented in Table 6-12B.

## 6.12.6 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on Table 6-12A and 6-12B.

There were no VOCs, TPH, or RCRA metals detected in any of the soil samples from Site 23, which exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

**Phase II Findings** 

The concentrations of benzo(a)anthracene, benzo(a)pyrene, and dibenz(a,h)anthracene deteced from samples 23-MW02-1416, 23-SS01, and 23-SS06 exceeded the BUSTR closure action levels. No other SVOCs detected exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

There were no VOCs, SVOCs, or RCRA metals detected in any of the groundwater samples from Site 23, which exceeded the OEPA VAP standard for generic unrestricted potable use standards or BUSTR closure action levels.

#### 6.12.7 Conclusions

Based on the analytical results, the soil at Site 23 may require special disposal and/or worker protection protocols (plan note) during construction activities.







TABLES

Table v=12A
Summary of Detected Chemicals in Soil
Site 23 - NS Railroad Building
ODOT Innerbelt Study
Cleveland, Ohio

			VAP	VAP		POGO EUMAN GO	d roco roman co	25.17 COMMO 200	CFOF-COMMITTEE	22-MAY04-1012
PA	PARAMETER	UNITS	Commercial Industrial S	Constitueitons  Verker Standard	Glosure Action	23-WW01-2224 07/13/2006	23-MW0I-2224-D 07/13/2006	07/13/2006	25-WW05-1012 07/14/2006	07/14/2006
	1,1,1-Trichloroethane	ug/kg	1,400,000	1,400,000	-	6.1 U	6.3 U	28 U	5.9 U	25
	1,1-Dichloroethane	ug/kg	2,300,000	2,300,000	•	6.1 ∪	6.3 ∪	28 U	5.9 U	1.8 J
ř	1,2,4-Trichlorobenzene	ug/kg	The state of the s	The state of the s	-	6.1 U	6.3 UJ	6.0 J	5.9 UJ	5.8 ∪
·	1,2-Dichlorobenzene	ug/kg	370,000	370,000	•	6.1 U	6.3 W	1.6 J	5.9 UJ	5.8 U
	1,3-Dichlorobenzene	ug/kg	240,000	240,000	e e	6.1 U	6.3 UJ	1.6 J	5.9 UJ	5.8 ∪
<i>.</i>	1,4-Dichlorobenzene	ug/kg	470,000	5,300,000		6.1 U	6.3 UJ	1.9 J	5.9 UJ	5.8 U
	2-Butanone	ug/kg	71,600,000	80,000,000	-	1.9 J	25 U	7.1 J	8.8 J	1.5 J
	Acetone	ug/kg	100,000,000	100,000,000	•	25 U	12 J	110 U	45	4.5 J
so	Benzene	ug/kg	100,000	310,000	149	6.1 ∪	1.5 J	1.4 J	0.86 J	5.8 U
ΟΛ	Carbon disulfide	ug/kg	720,000	720,000		6.1 U	1.7 J	11 J	2.4 J	5.8 ∪
	Cyclohexane	ug/kg			•	D 21	1.5 J	55 U	0.86 კ	12 U
	Ethylbenzene	ug/kg	230,000	230,000	45,500	6.1 U	2.5 J	11 (	1.6 J	5.8 U
	Isopropylbenzene	ug/kg	860,000	860,000	ı	6.1 U	0.80 J	ו5 ל	7.5 J	5.8 U
	Methylcyclohexane	ug/kg	1			12 U	4.5 J	SS ∪	2.7 J	12 U
	Methylene chloride	ug/kg	1,300,000	2,300,000	1	6.1 U	6.3 U	28 U	7.9	5.5 J
	Styrene	ug/kg	1,700,000	1,700,000		6.1 U	6.3 UJ	1.6 J	5.9 UJ	5.8 U
	Toluene	ug/kg	520,000	520,000	49,100	6.1 U	13 J	6.5 J	3.6 J	0.45 J
	Xylenes (total)	ug/kg	160,000	160,000	15,700	12 U	7.0 J	25 J	4.5 J	12 U
	1,1'-Biphenyl	ug/kg		•		810 U	40 J	2800 J	C 06Z	380 U
	2-Methylnaphthalene	ug/kg	1		•	820	260	11000 J	5200	41 J
	4-Methylphenol / 3-Methylphenol	ug/kg		•	1	810 U	72 J	18000 U	1500 U	380 U
	Acenaphthene	ug/kg	180,000,000	530,000,000	1	810 U	43 J	21000	1500 U	12 J
	Acenaphthylene	ug/kg			**************************************	810 U	35 J	18000 U	1500 U	380 U
•	Acetophenone	ug/kg	290,000,000	870,000,000		160 U	84 U	3700 U	310 U	U 77
	Anthracene	ug/kg	880,000,000	1,000,000,000		260 J	120 J	ا 16000	ל 190	18 J
sc	Benzaldehyde	ug/kg		-		810 U	410 U	18000 U	1500 U	380 U
00/	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	890	460	11000 J	560 J	59 J
S	Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	810	420	5300 J	480 J	27 א
	Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	1000	089	୮ 0008	P 089	84 J
	Benzo(ghi)perylene	ug/kg		1	an ea an Tomark John La Couran de La Andréa de Andréa de La Courant de L	460 J	ב 280	ا 1900	310 J	49 J
,	Benzo(k)fluoranthene	ug/kg	630,000	8,100,000	110,000	440 J	210 J	3200 J	230 J	30 P
	bis(2-Ethylhexyl) phthalate	ug/kg	230,000	230,000	1	810 U	7 88 38	18000 U	1500 U	38 1
	Caprolactam	ug/kg	1	1	1	810 U	410 U	18000 U	1500 U	380 U
	Carbazole	ug/kg	10,000,000	31,000,000		120 J	et J	10000	150 J	380 U
	Chrysene	ug/kg	6,700,000	41,000,000	1,100,000	006	460	9500 J	640 J	r 69

Table v=12A
Summary of Detected Chemicals in Soil
Site 23 - NS Railroad Building
ODOT Innerbelt Study
Cleveland, Ohio

			VAP Commercial/	VAP Construction	BUSTR	23-MW01-2224	23-MW01-2224-D	23-MW02-1416	23-MW03-1012	23-MW04-1012
<u> </u>	PARAMETER	UNITS	Industrial Standard <sup>()</sup>	- Worker - Standard <sup>(2)</sup>	Closure Action	07/13/2006	07/13/2006	07/13/2006	07/14/2006	07/14/2006
	Dibenz(a,h)anthracene	ug/kg	002'9	41,000	1,100	150 J	76 J	18000 U	1500 U	13 J
	Dibenzofuran	ug/kg	et peur mark et a volume et plate, de menerale de		-	180 J	130 J	13000 J	950 J	380 U
	Di-n-butyi phthalate	ug/kg	100,000	100,000	-	810 U	410 U	18000 U	1500 U	380 U
s	3	ug/kg	33,000,000	170,000,000	-	1800	880	46000	840 J	110 J
<b>0</b> 0.	ź	ug/kg	120,000,000	340,000,000		P 66	52 J	22000	120 J	0 08E
۸s		ug/kg	67,000	410,000	11,000	440 J	260 J	1800 J	260 J	38 J
	Naphthalene	ug/kg	530,000	1,900,000	39,800	550 J	400 J	31000	3000	16 J
	Phenanthrene	ug/kg	1	1	1	1100	290	74000	1500	ر 9 <i>2</i>
	Phenol	ug/kg	1,000,000,000	510,000,000	1	810 U	410 U	18000 U	1500 U	380 U
	Pyrene	ug/kg	25,000,000	130,000,000	1	1500	750	34000	870 J	100 7
Н	Gasoline Range Organics (C6-C12)	ug/kg	1 00 000	1	1,000,000	120 U	7 88 8	5600	140 U	120 U
lЧT	C10-C20	mg/kg		-	2,000	25	12 J	88	49	7.2
$\mathbb{I}$	C20-C34	mg/kg	1	1	5,000	130	52	130	310	16
	Arsenic	mg/kg	80	210	•	18.6 J	15.6 J	5.5 J	14.4	4.6
	Barium	mg/kg	200,000	45,000	1	160	133	45.7	311	13.8 J
STY	Second and the second	mg/kg	770	420	•	1.7 J	1.2 J	0.27 J	1.6	0.20
/T3	nium	mg/kg	8,900	2,000	1	20.8	17.3	5.5	45.7	6.7
M	Lead	mg/kg	-	•	1	221	212	16.2	128	9.9
	Silver	mg/kg	15,000	4,300	1	0.61 U	0.63 U	0.55 U	0.59 U	0.58 U
	Mercury	mg/kg	300	84		0.16	0.073 J	0.031 J	0.089 J	0.12 U

<sup>-- =</sup> Standard not available

 $U \approx The$  analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not

met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial Industrial Land Use

<sup>(2)</sup> VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Tabl A
Summary of Detecter-Chemicals in Soil
Site 23 - NS Railroad Building
ODOT Innerbelt Study
Cleveland, Ohio

			VAP	VAP							
			Commercial/	Construction: Worker	BUSTIR Flosting Action	23-SS01 07/14/2006	23-SS02 07/14/2006	23-5503	23-SS04 07/14/2006	23-SS05 07/14/2006	23-SS06 07/14/2006
	PARAMETER	UNITS	*Standard <sup>(1)</sup>	Standard <sup>(2)</sup>	Section 1						
	1,1,1-Trichloroethane	ug/kg	1,400,000	1,400,000	•	7.6 U	5.8 U	5.3 U	5.3 U	0.9	6.4 U
	1,1-Dichloroethane	ug/kg	2,300,000	2,300,000	The state of the s	7.6 U	5.8 U	5.3 U	5.3 U	0.9 0.9	6.4 U
	1,2,4-Trichlorobenzene	ug/kg	And the section with the section of the section with the section of the section o	Property Section of control Section Assessment Sect	- Andrews Andr	7.6 UJ	5.8 W	5.3 U	5.3 U	0.0 O	6.4 UJ
	1,2-Dichlorobenzene	ug/kg	370,000	370,000	-	7.6 UJ	5.8 UJ	5.3 U	5.3 U	0.0 U	6.4 UJ
	1,3-Dichlorobenzene	ug/kg	240,000	240,000		7.6 UJ	5.8 UJ	5.3 U	5.3 U	0.0 U	6.4 UJ
	1,4-Dichlorobenzene	ug/kg	470,000	5,300,000	en en en en en en en en en en en en en e	7.6 UJ	5.8 UJ	5.3 U	5.3 ∪	6.0 U	6.4 UJ
	2-Butanone	ug/kg	71,600,000	80,000,000		31 U	23 ∪	21 0	21 U	24 U	1.7 J
	Acetone	ug/kg	100,000,000	100,000,000	1	31 C	23 ∪	2	21 U	24 ∪	26 U
SO	Benzene	ug/kg	100,000	310,000	149	7.6 U	0.37 J	5.3 U	5.3 U	0.0 U	6.4 U
OΛ	Carbon disulfide	ug/kg	720,000	720,000		7.6 U	5.8 ∪	5.3 U	5.3 U	6.0 U	0.57 J
	Cyclohexane	ug/kg	•		••	15 U	12 U	1 U	11 O	12 U	13 U
	Ethylbenzene	ug/kg	230,000	230,000	45,500	7.6 U	5.8 W	5.3 U	5.3 U	6.0 U	6.4 U
	Isopropylbenzene	ug/kg	860,000	860,000	•	7.6 U	5.8 UJ	5.3 ∪	5.3 ∪	6.0 U	1.0 J
	Methylcyclohexane	ug/kg	-	-	••	15 U	12 U	11 U	- 1 - 1	12 U	13 U
	Methylene chloride	ug/kg	1,300,000	2,300,000	B B	7.6 U	6.7	5.3 U	5.3 U	6.0 U	6.4 U
	Styrene	ug/kg	1,700,000	1,700,000		7.6 U	5.8 UJ	5.3 ∪	5.3 U	6.0 U	6.4 U
	Toluene	ug/kg	520,000	520,000	49,100	7.6 U	1.3 J	5.3 ∪	5.3 U	6.0 U	6.4 U
	Xylenes (total)	ug/kg	160,000	160,000	15,700	15 U	± €	11 O	11 U	12 U	13 U
	1,1'-Biphenyl	ug/kg	:	1	;	2000 U	19000 U	350 UJ	1400 U	400 U	1100 U
	2-Methylnaphthalene	ug/kg	. 1	,		480 J	€800	350 UJ	68 J	67 J	100 J
	4-Methylphenol / 3-Methylphenol	ug/kg	1	1		2000 U	19000 UJ	350 UJ	1400 U	400 U	1100 U
	Acenaphthene	ug/kg	180,000,000	530,000,000	1	120 J	19000 U	27 J	28 J	14 J	57 J
	Acenaphthylene	ug/kg	The state of the s	-	•	140 J	19000 U	24 J	37 J	400 U	54 J
	Acetophenone	ug/kg	290,000,000	870,000,000		380 J	3900 UJ	20 €	280 U	80 U	210 U
	Anthracene	ug/kg	880,000,000	1,000,000,000	1	350 J	19000 UJ	Ր 96	130 J	51 ט	260 J
sç	Benzaldehyde	ug/kg		-		2900	19000 UJ	350 UJ	1400 U	400 U	1100 U
OΛ	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	1600 J	600 ع	370 J	560 J	330 1	1300
S	Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	1500 J	520 J	380 J	710 J	540	1400
	Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	2800	1100 J	620 J	Ր 086	089	2100
	Benzo(ghi)perylene	ug/kg		1		1200 J	830 J	300 J	570 J	450	1000
	Benzo(k)fluoranthene	ug/kg	000'089	8,100,000	110,000	L 07e	19000 U	230 J	340 J	330 J	760 J
	bis(2-Ethylhexyl) phthalate	ug/kg	230,000	230,000	1	2700	5600 J	55 J	1400 U	31 J	92 J
	Caprolactam	ug/kg	1	:	1 3	. n 000z	19000 UJ	350 UJ	1400 U	45 J	1100 U
	Carbazole	ug/kg	10,000,000	31,000,000		310 J	19000 UJ	ر 70 ع	1400 U	33 1	230 J
	Chrysene	ug/kg	6,700,000	41,000,000	1,100,000	1900 J	730 J	460 J	680 J	400	1300

Table v-12A
Summary of Detected Chemicals in Soil
Site 23 - NS Railroad Building
ODOT Innerbelt Study
Cleveland, Ohio

PA	PARAMETER	UNITS	Commercial/	Construction Worker - Standard (2)	BUSTR Glosure Action Level	23-SS01 07/14/2006	23-SS02 07/14/2006	23-SS03 07/14/2006	23-SS04 07/14/2006	,23-SS05 07/14/2006	23-SS06 07/14/2006
	Dibenz(a,h)anthracene	ug/kg	6,700	41,000	1,100	210 J	19000 U	350 UJ	1400 U	ი 06	320 J
	Dibenzofuran	ug/kg				200 J	19000 U	350 UJ	1400 U	400 U	1100 U
	Di-n-butyl phthalate	ug/kg	100,000	100,000	•	120 J	19000 UJ	350 UJ	1400 U	400 U	1100 U
8	Fluoranthene	ug/kg	33,000,000	170,000,000		3600	£ 069	820 J	1100 J	490	2700
)C		ug/kg	120,000,000	340,000,000	1	120 J	19000 U	28 J	34 J	400 U	28 J
λS	Indeno(1,2,3-cd)pyrene	ug/kg	67,000	410,000	11,000	1100 J	19000 U	220 ל	490 J	370 J	950 J
	Naphthalene	ug/kg	530,000	1,900,000	39,800	370 J	3700 J	350 UJ	51 J	81 J	73 J
	Phenanthrene	ug/kg	1 3	1	•	ل 1800	1800 J	390 J	490 J	180 J	1200
	Phenol	ug/kg	1,000,000,000	510,000,000	1	220 J	19000 U	350 UJ	1400 U	400 U	1100 U
	Pyrene	ug/kg	25,000,000	130,000,000	1	3000	1200 J	640 J	1000 J	450	2200
ŀ	Gasoline Range Organics (C6-C12)	ug/kg	•	1	1,000,000	150 U	120 U	110 U	110 U	120 U	130 U
197	C10-C20	mg/kg		1	2,000	61 U	ל 100	110 U	45 U	12 U	51 U
	C20-C34	mg/kg			5,000	220	2000	300	96	48	93
	Arsenic	mg/kg	80	210	1	9.8	7.5	2.6	4.8	6.1	5.1
-	Barium	mg/kg	200,000	45,000	•	423	186	92.7	47	85.4	57.9
דצ	Cadmium	mg/kg	022	420	1	2.5	1.4	0.3	0.24	4.8	0.38
ΑTΞ	Chromium	mg/kg	8,900	-		37.6	30.4	12.2	11.2	364	12.2
IW	Lead	mg/kg	1		ı	761	125	19.6	14	18.9	33
	Silver	mg/kg	15,000	4,300	ı	0.50 J	0.58 U	0.53 U	0.53 U	3.0 U	0.64 U
		mg/kg	300	84	•	0.13 J	0.057 J	0.021 ט	0.020 J	0.12 U	0.051 J

<sup>-- =</sup> Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial/Industrial Land Use

<sup>(2)</sup> VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Table o-12B
Summary of Detected Chemicals in Water
Site 23 - NS Railroad Building
ODOT Innerbelt Study

Cleveland, Ohio

, c	PARAMETER	UNITS	VAP UPUS/- REUPUS <sup>(0)</sup>	Closure:Action	23-MW01 08/10/2006	23-MW02 08/10/2006	23-MW03 08/10/2006	23-MW04 08/10/2006
	1,1,1-Trichloroethane	ug/L	200		1.0 U	1.0 U	1.0 U	62
80	3 1,1-Dichloroethane	ug/L	1400		1.0 U	-	1.0 U	4.9
24	1,1-Dichloroethene	ug/L	7	-	1.0 U	1.0 U	1.0 U	0.47 J
	Toluene	ug/L	1000	1000	0.25 J	1.0 U	1.0 U	2.0 U
soo	2-Methyinaphthalene	ng/L	<ol> <li>And the Conference of the Conferenc</li></ol>	Hartenburga (Renders and the second	0.20 U	0.35	0.20 U	0.20 U
آ ءِم	Acenaphthene	ug/L	680	•	0.20 U	0.62	0.20 U	0.20 U
	Arsenic	ng/L	50	•	10.0 U	6.1 3	5.7 J	10.0 U
CT	Barium	ug/L	2000	1	57.7 J	47.4 J	51.8 J	59.5 J
A'TE	Chromium	ug/L	100	•	3.5 J	4.0 J	5.0 U	5.0 U
IAI	Lead	ng/L	15	•	3.0 U	ო	2.0 J	3.0 U
	Selenium	ng/L	50	••	5.0 U	5.0 U	5.0 U	5.3

-- = Standard not available

 $\mathsf{U}=\mathsf{The}$  analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

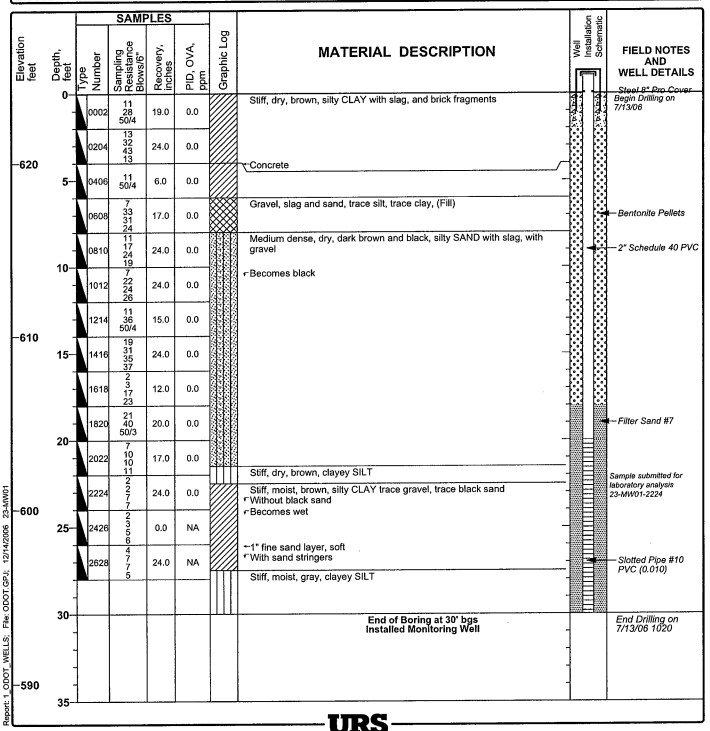
(1) VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

**BORING LOGS** 

Project Location: Site 23
Project Number: 15016633

### Log of Boring 23-MW01

Date(s) 7/13/06 Drilled 7/13/06	Logged By J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data 140# auto hammer	Total Depth of Borehole 30.0' bgs
Drill Rig CME-55 Type	Drilling Contractor HAD, Inc.	Approximate Ground Elevation <b>624'</b>
Location See Site Map	Sampling Method(s) 2" Split Spoon	Borehole Set monitoring well



Project Location: Site 23
Project Number: 15016633

# Log of Boring 23-MW02

By J. Kaminski	By M. Wolff
Hammer 140# auto hammer	Total Depth of Borehole 28.0' bgs
Drilling Contractor HAD, Inc.	Approximate Ground Elevation 630'
Sampling 2" Split Spoon	Borehole Completion Set monitoring well
	Hammer Data 140# auto hammer  Drilling Contractor HAD, Inc.  Sampling 2" Split Space

			SAME	PLES			5 ≟	
	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION  Well with the state of	WELL DETAILS
<del>-</del> 630	0-	0002	5 8 6 10	18.0	0.0		Loose, dry to moist, brown, sandy GRAVEL with slag, trace brick fragments, (Fill)	Begin Drilling on 7/13/06
:	- -	0204	8 12 12 10	10.0	0.0			
-625	5- -	0406	4 6 7 15	23.0	2.4	$\bigotimes$		- c - c - c - c - c - c - c - c - c - c
	-	0608	14 50/2	8.0	0.0		Stiff, dry to moist, brownish gray, SILT with slag, trace gravel	2" Schedule 40 PVC
<b>-620</b>	- 10-	0810	10 15 12 9	22.0	0.0		Stiff, dry to moist, brownish gray, SILT with slag, trace gravel	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
	-	1012	16 18 22	24.0	7.0		VVIIII Sanu	Bentonite Pellets
	-	1214	50/2	2.0	NA		FSolvent odor	s <i>ଲିନ୍ୟା</i> ୧୩୪ନ ।
-615	15-	1416	32 35 40	24.0	92.5		Wood fragments  Very stiff, dry, dark gray, SILT with gravel, and slag	laboratory analysis 23-MW02-1416
	_	1618	14 50/4	24.0	21.2		- Not as stiff	
-610	20-	1820	15 16 10 20	19.0	8.8			Filter Sand #7
	-	2022	4 4 5	11.0	5.5		Loose, dry, yellow, SAND trace coal, with slag	
-605	-	2224	4 6 7 6 6	12.0	NA		Loose, wet, brown, medium SAND trace fine gravel	Filter Sand #7
003	25-	2426	6 4 4 10	7.0	NA 			— Slotted Pipe #10 PVC (0.010)
	-	2628	10 11	16.0	NA		End of Boring at 28' bgs Installed Monitoring Well	End Drilling on 7/13/06 1415
-600	30						URS	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,

Project Location: Site 23
Project Number: 15016633

# Log of Boring 23-MW03

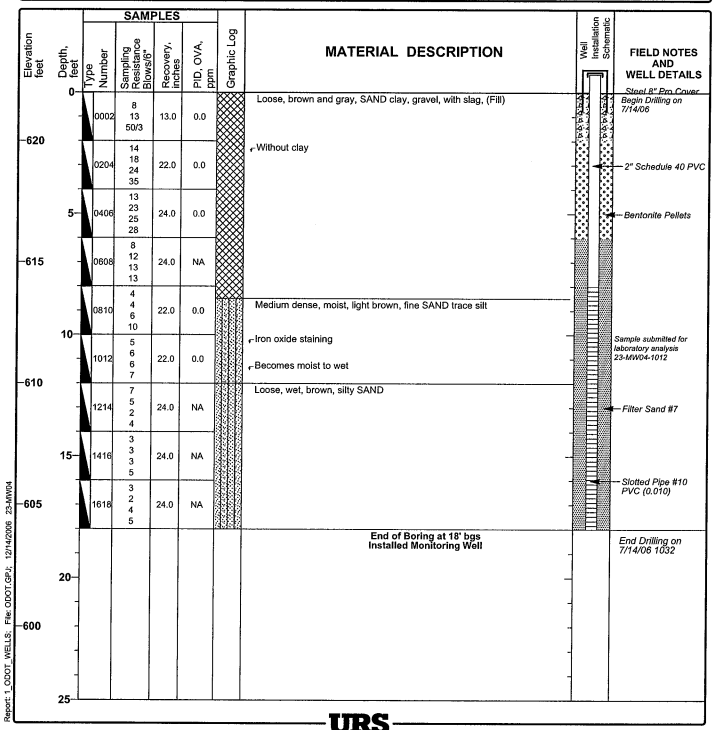
Date(s) 7/14/06 Drilled	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer 1	140# auto hammer	Total Depth of Borehole 25.0′ bgs
Drill Rig CME-55 Type	Drilling Contractor	HAD, inc.	Approximate Ground Elevation 622'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well

			04115	VI EO				<del></del>		1	
		-	SAME	LES		_			ation :	atic	
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	Weli	Installation		FIELD NOTES AND WELL DETAILS Steel 8" Pro Cover
-620	0	0002	11 50/4	12.0	0.0		Loose, dry, brown, silty SAND with slag, with brick fragments	प्रचक्रक		44444	Steel 8" Pro Cover Begin Drilling on 7/14/06
-020	_	0204	5 13 19 27	12.0	0.0		Stiff, wet, black, silty SAND trace slag, trace grass ←Becomes dry				
	5-	0406	40	24.0	0.0		December hypurnish group				
-615	-	0608	5 12 12 10	13.0	0.0		FBecomes brownish gray  Loose, dry, dark gray, silty SAND trace fine gravel	 			—Bentonite Pellets
	- 10	0810	4	12.0	0.0		Loose, ary, dark gray, silly SAND trace line gravel		•		— 2" Schedule 40 PVC Sample submitted for
-610	<u>-</u>	1012	4 15 19 10 WOH	17.0	0.0		சWith coal, and slag, and brick fragments				laboratory analysis 23-MW03-1012
	-	1214	1 1 1	4.0	0.0			ļ		•••	
	15— -	1416	4	3.0	0.0		+White paste material at 15'  -rWith wood fragments				
-605	-	1618	4	0.0	0.0		Soft, wet, black, silty SAND	_			
	20-	1820	7	24.0	0.0		Loose, wet, brown, medium SAND	-			
-600	-	2022	1 6 1	24.0	0.0		Augered down to 25' bgs				Filter Sand #7
-600 -595	-						, ago, ou dominio ao ogo				Slotted Pipe #10 PVC (0.010)
	25-						End of Boring at 25' bgs Installed Monitoring Well	-	<u></u> E		End Drilling on 7/14/06 1230
595	-							1			
1	30-				_			1			
							URS				

Project Location: Site 23
Project Number: 15016633

### Log of Boring 23-MW04

Date(s) 7/14/06 Drilled	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140# auto hammer	Total Depth of Borehole 18.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 622'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Groundwater Level and Date Measured 611.32 on 8/10/06			



DATA ASSESSMENT REPORT

### Data Assessment Report ODOT Innerbelt Study Site 23 – NS Railroad Building

Reviewer: P. Schuler Date: November 14, 2006

Eleven soil samples, four groundwater samples, and three trip blanks were collected at the NS Railroad Building site in Cleveland, Ohio, from July 13 through August 10, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample		Re	quested .	Analyse	es <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC	TPH	Met
A6G140115001	23-MW01-2224 <sup>(2)</sup>	07/13/2006	Soil	X	X	X	X
A6G140115002	23-MW01-2224-D <sup>(2)</sup>	07/13/2006	Soil	X	X	X	X
A6G140115003	23-MW02-1416	07/13/2006	Soil	X	X	X	X
A6G140115004	TB-071306	07/13/2006	Trip Blank	X			
A6G140397001	23-MW04-1012	07/14/2006	Soil	X	X	X	X
A6G140397002	23-SS01	07/14/2006	Soil	X	X	X	X
A6G140397003	23-SS02	07/14/2006	Soil	X	X	X	X
A6G140397004	23-SS03	07/14/2006	Soil	X	X	X	X
A6G140397005	23-SS04	07/14/2006	Soil	X	X	X	X
A6G140397006	23-SS05	07/14/2006	Soil	X	X	X	X
A6G140397007	23-SS06	07/14/2006	Soil	X	X	X	X
A6G140397008	23-MW03-1012	07/14/2006	Soil	X	X	X	X
A6G140397009	TB-071406	07/14/2006	Trip Blank	X			
A6H120107001	23-MW01	08/10/2006	Groundwater	X	X		X
A6H120107002	23-MW02	08/10/2006	Groundwater	X	X		X
A6H120107003	23-MW03	08/10/2006	Groundwater	X	X		X
A6H120107004	23-MW04	08/10/2006	Groundwater	X	X		X
A6H120107005	TB-081006	08/10/2006	Trip Blank	X			

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B]

SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

TPH = Total Petroleum Hydrocarbons (Gasoline and Diesel Range Organics) [SW-846 Method 8015A/B]

Met = RCRA Metals [SW-846 Methods 6010B/6020/7470A/7471A]

(2) Samples 23-MW01-2224 and 23-MW01-2224-D are field duplicates.

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not



include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Samples 23-MW04-1012 and 23-MW01-2224-D were analyzed for gasoline range organics four days after the 14-day holding time had expired. The nondetect results for both samples were qualified as estimated ("UJ") and may be biased low.
- Positive detections for acetone in sample 23-MW02-1416, for methylene chloride in samples 23-MW01-2224, 23-SS03, 23-SS05, 23-SS06, and 23-MW04, for bis(2-ethylhexyl)phthalate in samples 23-MW01, 23-MW02, and 23-MW03, and for gasoline range organics in samples 23-SS03, 23-SS04, 23-SS05, and 23-MW03-1012 were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks and/or trip blanks at similar concentrations.
- The semivolatile compound, bis(2-ethylhexyl)phthalate, was detected at concentrations ranging from 0.9 to 1.8 ug/L in all equipment blanks associated with project samples. Since this compound is considered a common laboratory contaminant, its presence in environmental samples at concentrations less than ten times the highest equipment blank concentration is attributed to external contamination rather than actual site conditions. Therefore, the positive detections for bis(2-ethylhexyl)phthalate in samples 23-MW01-2224-D, 23-MW04-1012, 23-SS03, 23-SS05 and 23-SS06 were qualified as nondetect ("U") at the value reported, even though it was not detected in the associated method blanks.
- One or more volatile internal standard responses were outside of the acceptance range in samples 23-MW01-2224-D, 23-SS01, 23-SS02, 23-SS06, and 23-MW03-1012. The results for all volatile analytes quantified from each of the noncompliant internal standards were qualified as estimated ("J" or "UJ").
- One or more semivolatile internal standard responses were outside of the acceptance range in samples 23-SS02 and 23-SS03. The results for all semivolatile analytes quantified from each of the noncompliant internal standards were qualified as estimated ("J" or "UJ").
- The arsenic and cadmium results for the soil samples collected 07/14/06 were qualified as estimated ("J") due to low matrix spike recoveries in sample 23-SS02.
- The lab reported some metals results with a "G" qualifier, indicating that a dilution was required due to matix interferences. The "G" qualifiers were removed in the final data set, but the sample reporting limits are elevated accordingly.
- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J" for organics, and "B" for inorganics), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, organic "J" qualifiers were retained with the numeric results. The inorganic "B"

qualifiers were changed to "J" qualifiers for the sake of consistency throughout the package.

All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-7.

# Table 2-1 Analytical Data Summary Site 23 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140115001 23-MW01-2224 07/13/2006	A6G140115002 23-MW01-2224-D 07/13/2006	A6G140115003 23-MW02-1416 07/13/2006	A6G140397001 23-MW04-1012 07/14/2006	A6G140397002 23-SS01 07/14/2006	A6G140397003 23-SS02 07/14/2006
1,1,1-Trichloroethane	ug/kg	6.1 U	6.3 U	28 U	25	7.6 U	5.8 U
1,1,2,2-Tetrachloroethane	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 UJ	5.8 UJ
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
1,1,2-Trichloroethane	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 UJ
1,1-Dichloroethane	ug/kg	6.1 U	6.3 U	28 U	1.8 J	7.6 U	5.8 U
1,1-Dichloroethene	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
1,2,4-Trichlorobenzene	ug/kg	6.1 U	6.3 UJ	6.0 J	5.8 U	7.6 UJ	5.8 UJ
1,2-Dibromo-3-chloropropane	ug/kg	12 U	13 UJ	55 U	12 U	15 UJ	12 UJ
1,2-Dibromoethane	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 UJ
1,2-Dichlorobenzene	ug/kg	6.1 U	6.3 UJ	1.6 J	5.8 U	7.6 UJ	5.8 UJ
1,2-Dichloroethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
1,2-Dichloropropane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
1,3-Dichlorobenzene	ug/kg	6.1 U	6.3 UJ	1.6 J	5.8 U	7.6 UJ	5.8 UJ
1,4-Dichlorobenzene	ug/kg	6.1 U	6.3 UJ	1.9 J	5.8 U	7.6 UJ	5.8 UJ
2-Butanone	ug/kg	1.9 J	25 U	7.1 J	1.5 J	31 U	23 U
2-Hexanone	ug/kg	25 U	25 UJ	110 U	23 U	31 U	23 UJ
4-Methyl-2-pentanone	ug/kg	25 U	25 UJ	110 U	23 U	31 UJ	23 UJ
Acetone	ug/kg	25 U	12 J	110 U	4.5 J	31 U	23 U
Benzene	ug/kg	6.1 U	1.5 J	1.4 J	5.8 U	7.6 U	0.37 J
Bromodichloromethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Bromoform	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 UJ
¹sromomethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
rbon disulfide	ug/kg	6.1 U	1.7 J	11 J	5.8 U	7.6 U	5.8 U
Carbon tetrachloride	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Chlorobenzene	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 U
Chloroethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Chloroform	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Chloromethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
cis-1,2-Dichloroethene	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
cis-1,3-Dichloropropene	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Cyclohexane	ug/kg	12 U	1.5 J	55 U	12 U	15 U	12 U
Dibromochloromethane	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 UJ
Dichlorodifluoromethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Ethylbenzene	ug/kg	6.1 U	2.5 J	11 J	5.8 U	7.6 U	5.8 UJ
Isopropylbenzene	ug/kg	6.1 U	0.80 J	15 J	5.8 U	7.6 U	5.8 UJ
Methyl acetate	ug/kg	12 U	13 U	55 U	12 U	15 U	12 U
Methyl tert-butyl ether	ug/kg	25 U	25 U	110 U	23 U	31 U	23 U
Methylcyclohexane	ug/kg	12 U	4.5 J	55 U	12 U	15 U	12 U
Methylene chloride	ug/kg	6.1 U	6.3 U	28 U	5.5 J	7.6 U	6.7
Styrene	ug/kg	6.1 U	6.3 UJ	1.6 J	5.8 U	7.6 U	5.8 UJ
Tetrachloroethene	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 UJ
Toluene	ug/kg	6.1 U	13 J	6.5 J	0.45 J	7.6 U	1.3 J
trans-1,2-Dichloroethene	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
trans-1,3-Dichloropropene	ug/kg	6.1 U	6.3 UJ	28 U	5.8 U	7.6 U	5.8 UJ
Trichloroethene	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Trichlorofluoromethane	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Vinyl chloride	ug/kg	6.1 U	6.3 U	28 U	5.8 U	7.6 U	5.8 U
Yvienes (total)	ug/kg	12 U	7.0 J	25 J	12 U	15 U	12 UJ

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

# Table 2-1 Analytical Data Summary Site 23 Soil Volatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140397004 23-SS03 07/14/2006	A6G140397005 23-SS04 07/14/2006	A6G140397006 23-SS05 07/14/2006	A6G140397007 23-SS06 07/14/2006	A6G140397008 23-MW03-1012 07/14/2006
1.1.1-Trichloroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
1,1,2,2-Tetrachloroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 UJ	5.9 UJ
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
1,1,2-Trichloroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
1,1-Dichloroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
1,1-Dichloroethene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
1,2,4-Trichlorobenzene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 UJ	5.9 UJ
1,2-Dibromo-3-chloropropane	ug/kg	11 U	11 U	12 U	13 UJ	12 UJ
1,2-Dibromoethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
1,2-Dichlorobenzene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 UJ	5.9 UJ
1,2-Dichloroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
1,2-Dichloropropane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
1,3-Dichlorobenzene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 UJ	5.9 UJ
1,4-Dichlorobenzene	ug/kg ug/kg	5.3 U	5.3 U	6.0 U	6.4 UJ	5.9 UJ
2-Butanone	ug/kg	21 U	3.3 U 21 U	24 U	1.7 J	8.8 J
2-Hexanone	ug/kg ug/kg	21 U	21 U	24 U	26 U	23 UJ
4-Methyl-2-pentanone	ug/kg ug/kg	21 U	21 U	24 U	26 UJ	23 UJ
Acetone		21 U	21 U	24 U	26 U	45
Benzene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	0.86 J
The first the same as the same of the same	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Bromodichloromethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
Bromoform	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	the second second
Bromomethane	ug/kg	A CONTROL OF THE PARTY OF THE P	A A COLOR OF THE STATE OF THE S			5.9 U
Carbon disulfide	ug/kg	5.3 U	5.3 U	6.0 U	0.57 J	2.4 J
Carbon tetrachloride	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Chlorobenzene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
Chloroethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Chloroform	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Chloromethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
cis-1,2-Dichloroethene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
cis-1,3-Dichloropropene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Cyclohexane	ug/kg	11 U	11 U	12 U	13 U	0.86 J
Dibromochloromethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
Dichlorodifluoromethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Ethylbenzene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	1.6 J
Isopropylbenzene	ug/kg	5.3 U	5.3 U	6.0 U	1.0 J	7.5 J
Methyl acetate	ug/kg	11 U	11 U	12 U	13 U	12 U
Methyl tert-butyl ether	ug/kg	21 U	21 U	24 U	26 U	23 U
Methylcyclohexane	ug/kg	11 U	11 U	12 U	13 U	2.7 J
Methylene chloride	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	7.9
Styrene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
Tetrachloroethene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
Toluene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	3.6 J
trans-1,2-Dichloroethene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
trans-1,3-Dichloropropene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 UJ
Trichloroethene	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Trichlorofluoromethane	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Vinyl chloride	ug/kg	5.3 U	5.3 U	6.0 U	6.4 U	5.9 U
Xylenes (total)	ug/kg	11 U	11 U	12 U	13 U	4.5 J

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

Table 2-1
Analytical Data Summary
Site 23 Soil Volatiles
ODOT Innerbelt Study

PARAMETER	UNITS	A6G140115004 TB-071306 07/13/2006	A6G140397009 TB-071406 07/14/2006
1,1,1-Trichloroethane	ug/L	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L ug/L	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L ug/L	2.0 U	2.0 U
The second secon	ug/L	1.0 U	1.0 U
1,2-Dibromoethane		1.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L		1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U
2-Butanone	ug/L	10 U	1
2-Hexanone	ug/L	10 U	10 U 10 U
4-Methyl-2-pentanone	ug/L	10 U	
Acetone	ug/L	10 U	10 U
Benzene	ug/L	1.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U
Chloroethane	ug/L	1.0 U	1.0 U
Chloroform	ug/L	1.0 U	1.0 U
Chloromethane	ug/L	1.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	1.0 U
Dibromochloromethane	ug/L	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U
Methylene chloride	ug/L	0.52 J	0.39 J
Styrene	ug/L	1.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U
Toluene	ug/L	1.0 U	1.0 U
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	2.0 U
Aylelles (total)		The analyte was anal	

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-2 Analytical Data Summary Site 23 Soil Semivolatiles ODOT Innerbelt Study

		A6G140115001 23-MW01-2224	A6G140115002 23-MW01-2224-D	A6G140115003 23-MW02-1416	A6G140397001 23-MW04-1012	A6G140397002 23-SS01	A6G140397003 23-SS02
PARAMETER	UNITS	07/13/2006	07/13/2006	07/13/2006	07/14/2006	07/14/2006	07/14/2006
1,1'-Biphenyl	ug/kg	810 U	40 J	2800 J	380 U	2000 U	19000 U
2,2'-oxybis(1-Chloropropane)	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
2,4,5-Trichlorophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2,4,6-Trichlorophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2,4-Dichlorophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
2,4-Dimethylphenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
2,4-Dinitrophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2,4-Dinitrotoluene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2,6-Dinitrotoluene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2-Chloronaphthalene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2-Chlorophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
2-Methylnaphthalene	ug/kg	820	560	11000 J	41 J	480 J	6800 J
2-Methylphenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
2-Nitroaniline	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
2-Nitrophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
3.3'-Dichlorobenzidine	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
3-Nitroaniline	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
4,6-Dinitro-2-methylphenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
4-Bromophenyl phenyl ether	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
4-Chloro-3-methylphenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
4-Chloroaniline	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Chlorophenyl phenyl ether	ug/kg ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
ethylphenol / 3-Methylphenol	ug/kg ug/kg	810 U	72 J	18000 U	380 U	2000 U	19000 UJ
4-Nitroaniline	ug/kg ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
4-Nitrophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
Acenaphthene	ug/kg	810 U	43 J	21000	12 J	120 J	19000 U
Acenaphthylene	ug/kg	810 U	35 J	18000 U	380 U	140 J	19000 U
	ug/kg ug/kg	160 U	84 U	3700 U	77 U	380 J	3900 UJ
Acetophenone Anthracene	ug/kg ug/kg	260 J	120 J	16000 J	18 J	350 J	19000 UJ
		810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Atrazine	ug/kg ug/kg	810 U	410 U	18000 U	380 U	2900	19000 UJ
Benzaldehyde		890	460	11000 J	59 J	1600 J	600 J
Benzo(a)anthracene	ug/kg	810	420	5300 J	57 J	1500 J	520 J
Benzo(a)pyrene	ug/kg	1000	680	8000 J	84 J	2800	1100 J
Benzo(b)fluoranthene	ug/kg	460 J	280 J	1900 J	49 J	1200 J	830 J
Benzo(ghi)perylene	ug/kg	440 J	210 J	3200 J	30 J	970 J	19000 U
Benzo(k)fluoranthene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
bis(2-Chloroethoxy)methane	ug/kg		410 U	18000 U	380 U	2000 U	19000 UJ
bis(2-Chloroethyl) ether	ug/kg	810 U	the contract of the contract of	the contract of the contract o	the state of the s	2700	
bis(2-Ethylhexyl) phthalate	ug/kg	810 U	410 U	18000 U	380 U		5600 J
Butyl benzyl phthalate	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
Caprolactam	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Carbazole	ug/kg	120 J	61 J	10000 J	380 U	310 J	19000 UJ
Chrysene	ug/kg	900	460	9500 J	69 J	1900 J	730 J
Dibenz(a,h)anthracene	ug/kg	150 J	76 J	18000 U	13 J	210 J	19000 U
Dibenzofuran	ug/kg	180 J	130 J	13000 J	380 U	200 J	19000 U
Diethyl phthalate	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
Dimethyl phthalate	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
n-butyl phthalate	ug/kg	810 U	410 U	18000 U	380 U	120 J	19000 UJ
∫ı-octyl phthalate	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
Fluoranthene	ug/kg	1800	880	46000	110 J	3600	690 J
Fluorene	ug/kg	99 J	52 J	22000	380 U	120 J	19000 U

# Table 2-2 Analytical Data Summary Site 23 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140115001 23-MW01-2224 07/13/2006	A6G140115002 23-MW01-2224-D 07/13/2006	A6G140115003 23-MW02-1416 07/13/2006	A6G140397001 23-MW04-1012 07/14/2006	A6G140397002 23-SS01 07/14/2006	A6G140397003 23-SS02 07/14/2006
Hexachlorobenzene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Hexachlorobutadiene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Hexachlorocyclopentadiene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 U
Hexachloroethane	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Indeno(1,2,3-cd)pyrene	ug/kg	440 J	260 J	1800 J	38 J	1100 J	19000 U
Isophorone	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Naphthalene	ug/kg	550 J	400 J	31000	16 J	370 J	3700 J
Nitrobenzene	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
N-Nitrosodi-n-propylamine	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
N-Nitrosodiphenylamine	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Pentachlorophenol	ug/kg	810 U	410 U	18000 U	380 U	2000 U	19000 UJ
Phenanthrene	ug/kg	1100	590	74000	76 J	1800 J	1800 J
Phenol	ug/kg	810 U	410 U	18000 U	380 U	220 J	19000 U
Pyrene	ug/kg	1500	750	34000	100 J	3000	1200 J

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

Table 2-2 Analytical Data Summary Site 23 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140397004 23-SS03 07/14/2006	A6G140397005 23-SS04 07/14/2006	A6G140397006 23-SS05 07/14/2006	A6G140397007 23-SS06 07/14/2006	A6G140397008 23-MW03-1012 07/14/2006
1,1'-Biphenyl	ug/kg	350 UJ	1400 U	400 U	1100 U	290 J
· ' '	ug/kg ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2,2'-oxybis(1-Chloropropane)	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2,4,5-Trichlorophenol	ug/kg ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2,4,6-Trichlorophenol		350 UJ	1400 U	400 U	1100 U	1500 U
2,4-Dichlorophenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2,4-Dimethylphenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2,4-Dinitrophenol	ug/kg	and the state of t	1400 U	400 U	1100 U	1500 U
2,4-Dinitrotoluene	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2,6-Dinitrotoluene	ug/kg	350 UJ		400 U	1100 U	1500 U
2-Chloronaphthalene	ug/kg	350 UJ	1400 U	the second of the second	1100 U	1500 U
2-Chlorophenol	ug/kg	350 UJ	1400 U	400 U	100 J	5200
2-Methylnaphthalene	ug/kg	350 UJ	68 J	67 J	. 7.7-	1500 U
2-Methylphenol	ug/kg	350 UJ	1400 U	400 U	1100 U	
2-Nitroaniline	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
2-Nitrophenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
3,3'-Dichlorobenzidine	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
3-Nitroaniline	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4,6-Dinitro-2-methylphenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Bromophenyl phenyl ether	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Chloro-3-methylphenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Chloroaniline	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Chlorophenyl phenyl ether	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Methylphenol / 3-Methylphenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Nitroaniline	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
4-Nitrophenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Acenaphthene	ug/kg	27 J	28 J	14 J	57 J	1500 U
Acenaphthylene	ug/kg	24 J	37 J	400 U	54 J	1500 U
Acetophenone	ug/kg	70 UJ	280 U	80 U	210 U	310 U
Anthracene	ug/kg	96 J	130 J	51 J	260 J	190 J
Atrazine	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Benzaldehyde	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Benzo(a)anthracene	ug/kg	370 J	560 J	330 J	1300	560 J
Benzo(a)pyrene	ug/kg	380 J	710 J	540	1400	480 J
Benzo(b)fluoranthene	ug/kg	620 J	980 J	680	2100	680 J
Benzo(ghi)perylene	ug/kg	300 J	570 J	450	1000 J	310 J
Benzo(k)fluoranthene	ug/kg	230 J	340 J	330 J	760 J	230 J
bis(2-Chloroethoxy)methane	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
bis(2-Chloroethyl) ether	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
bis(2-Ethylhexyl) phthalate	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Butyl benzyl phthalate	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Caprolactam	ug/kg	350 UJ	1400 U	45 J	1100 U	1500 U
Carbazole	ug/kg	70 J	1400 U	33 J	230 J	150 J
Chrysene	ug/kg	460 J	680 J	400	1300	640 J
Dibenz(a,h)anthracene	ug/kg	350 UJ	1400 U	90 J	320 J	1500 U
Dibenzofuran	ug/kg	350 UJ	1400 U	400 U	1100 U	950 J
Diethyl phthalate	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Dimethyl phthalate	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Di-n-butyl phthalate	ug/kg ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Di-n-octyl phthalate	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
		820 J	1100 J	490	2700	840 J
Fluoranthene Fluorene	ug/kg ug/kg	820 J 28 J	34 J	400 U	28 J	120 J

#### Table 2-2 Analytical Data Summary Site 23 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G140397004 23-SS03 07/14/2006	A6G140397005 23-SS04 07/14/2006	A6G140397006 23-SS05 07/14/2006	A6G140397007 23-SS06 07/14/2006	A6G140397008 23-MW03-1012 07/14/2006
Hexachlorobenzene	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Hexachlorobutadiene	ug/kg	350 ÚJ	1400 U	400 U	1100 U	1500 U
Hexachlorocyclopentadiene	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Hexachloroethane	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Indeno(1,2,3-cd)pyrene	ug/kg	220 J	490 J	370 J	950 J	260 J
Isophorone	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Naphthalene	ug/kg	350 UJ	51 J	81 J	73 J	3000
Nitrobenzene	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
N-Nitrosodi-n-propylamine	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
N-Nitrosodiphenylamine	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Pentachlorophenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Phenanthrene	ug/kg	390 J	490 J	180 J	1200	1500
Phenol	ug/kg	350 UJ	1400 U	400 U	1100 U	1500 U
Pyrene	ug/kg	640 J	1000 J	450	2200	870 J

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.

# Table 2-3 Analytical Data Summary Site 23 Soil TPH ODOT Innerbelt Study

PARAMETER	UNITS	A6G140115001 23-MW01-2224 07/13/2006	A6G140115002 23-MW01-2224-D 07/13/2006	A6G140115003 23-MW02-1416 07/13/2006	A6G140397001 23-MW04-1012 07/14/2006	A6G140397002 23-SS01 07/14/2006
Gasoline Range Organics (C6-C12)	ug/kg	120 U	88 J	5600	120 U	150 U
C10-C20	mg/kg	25	12 J	88	7.2	61 U
C20-C34	mg/kg	130	52	130	16	220

PARAMETER	UNITS	A6G140397003 23-SS02 07/14/2006	A6G140397004 23-SS03 07/14/2006	A6G140397005 23-SS04 07/14/2006	A6G140397006 23-SS05 07/14/2006	A6G140397007 23-SS06 07/14/2006
Gasoline Range Organics (C6-C12)	ug/kg	120 U	110 U	110 U	120 U	130 U
C10-C20	mg/kg	100 J	110 U	42 U	12 U	51 U
C20-C34	mg/kg	2000	300	96	48	93

PARAMETER	UNITS	A6G140397008 23-MW03-1012 07/14/2006
Gasoline Range Organics (C6-C12)	ug/kg	140 U
C10-C20	mg/kg	49
C20-C34	mg/kg	310

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-4
Analytical Data Summary
Site 23 Soil Metals
ODOT Innerbelt Study

PARAMETER	UNITS	A6G140115001 23-MW01-2224 07/13/2006	A6G140115002 23-MW01-2224-D 07/13/2006	A6G140115003 23-MW02-1416 07/13/2006	A6G140397001 23-MW04-1012 07/14/2006	A6G140397002 23-SS01 07/14/2006	A6G140397003 23-SS02 07/14/2006
Percent Solids	%	81.5	79.6	90.4	86.1	65.6	86.1
Arsenic	mg/kg	18.6 J	15.6 J	5.5 J	4.6	9.8	7.5
Barium	mg/kg	160	133	45.7	13.8 J	423	186
Cadmium	mg/kg	1.7 J	1.2 J	0.27 J	0.20 J	2.5	1.4
Chromium	mg/kg	20.8	17.3	5.5	6.7	37.6	30.4
Lead	mg/kg	221	212	16.2	6.6	761	125
Selenium	mg/kg	0.61 U	0.63 U	0.55 U	0.58 U	0.76 U	0.58 U
Silver	mg/kg	0.61 U	0.63 ∪	0.55 U	0.58 U	0.50 J	0.58 U
Mercury	mg/kg	0.16	0.073 J	0.031 J	0.12 U	0.13 J	0.057 J

PARAMETER	UNITS	A6G140397004 23-SS03 07/14/2006	A6G140397005 23-SS04 07/14/2006	A6G140397006 23-SS05 07/14/2006	A6G140397007 23-SS06 07/14/2006	A6G140397008 23-MW03-1012 07/14/2006
Percent Solids	%	94.5	94.2	83.5	77.9	85.3
Arsenic	mg/kg	2.6	4.8	6.1	5.1	14.4
Barium	mg/kg	92.7	47	85.4	57.9	311
Cadmium	mg/kg	0.3	0.24	1.8	0.38	1.6
Chromium	mg/kg	12.2	11.2	364	12.2	45.7
Lead	mg/kg	19.6	14	18.9	33	128
Selenium	mg/kg	0.53 U	0.53 U	3.0 U	0.64 U	2.9 U
Silver	mg/kg	0.53 U	0.53 U	3.0 U	0.64 U	0.59 U
Mercury	mg/kg	0.021 J	0.020 J	0.12 U	0.051 J	0.089 J

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-5
Analytical Data Summary
Site 23 Water Volatiles
ODOT Innerbelt Study

		A6H120107001 23-MW01	A6H120107002 23-MW02	A6H120107003 23-MW03	A6H120107004 23-MW04	A6H120107005 TB-081006
PARAMETER	UNITS	08/10/2006	08/10/2006	08/10/2006	08/10/2006	08/10/2006
1,1,1-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U	62	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,1-Dichloroethane	ug/L	1.0 U	1.1	1.0 U	4.9	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U	0.47 J	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U	2.0 U	4.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,2-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,3-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
1,4-Dichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
2-Butanone	ug/L	10 U	10 U	10 U	20 U	10 U
2-Hexanone	ug/L	10 U	10 U	10 U	20 U	10 U
4-Methyl-2-pentanone	ug/L	10 U	10 U	10 U	20 U	10 U
Acetone	ug/L	10 U	10 U	10 U	20 U	10 U
Benzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Chloroethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Chloroform	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Chloromethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
and the first of t	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
cis-1,3-Dichloropropene	Carlotte and the second second	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Cyclohexane	ug/L	1.0 U				
Dibromochloromethane	ug/L	ere and the second of the second	1.0 U	1.0 U	· 2.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U	10 U	20 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U	5.0 U	10 U	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Methylene chloride	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Styrene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Toluene	ug/L	0.25 J	1.0 U	1.0 U	2.0 U	1.0 U
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Vinyl chloride	ug/L	1.0 U	1.0 U	1.0 U	2.0 U	1.0 U
Xylenes (total)	ug/L	2.0 U	2.0 U	2.0 U	4.0 U	2.0 U

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.



Table 2-6 Analytical Data Summary Site 23 Water Semivolatiles ODOT Innerbelt Study

Augusta (n. 1905) Raid			A6H120107001 23-MW01	A6H120107002 23-MW02	A6H120107003 23-MW03	A6H120107004 23-MW04
PARAMETER		UNITS	08/10/2006	08/10/2006	08/10/2006	08/10/2006
1,1'-Biphenyl		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2,2'-oxybis(1-Chloropropane)		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2,4,5-Trichlorophenol		ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,4,6-Trichlorophenol		ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,4-Dichlorophenol	• •	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
2,4-Dimethylphenol		ug/L	2.0 U	2.0 U	2.0 U	2.0 U
2,4-Dinitrophenol	•	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2.4-Dinitrotoluene		ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2,6-Dinitrotoluene		ug/L	5.0 U	5.0 U	5.0 U	5.0 U
2-Chloronaphthalene		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2-Chlorophenol		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
2-Methylnaphthalene		ug/L ug/L	0.20 U	0.35	0.20 U	0.20 U
2-Methylphenol		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
		ug/∟ ug/L	2.0 U	2.0 U	2.0 U	2.0 U
2-Nitroaniline 2-Nitrophenol		ug/L ug/L	2.0 U	2.0 U	2.0 U	2.0 U
3,3'-Dichlorobenzidine		ug/L ug/L	5.0 U	5.0 U	5.0 U	5.0 U
The second secon		ug/L	2.0 U	2.0 U	2.0 U	2.0 U
3-Nitroaniline		_	5.0 U	5.0 U	5.0 U	5.0 U
4,6-Dinitro-2-methylphenol	a - 5 - 178-1	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Bromophenyl phenyl ether		ug/L				
4-Chloro-3-methylphenol		ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Chloroaniline		ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Chlorophenyl phenyl ether		ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Methylphenol		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
4-Nitroaniline	,	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
4-Nitrophenol		ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Acenaphthene	*	ug/L	0.20 U	0.62	0.20 U	0.20 U
Acenaphthylene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Acetophenone		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Anthracene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Atrazine		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Benzaldehyde		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Benzo(a)anthracene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Benzo(a)pyrene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Benzo(b)fluoranthene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Benzo(ghi)perylene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Benzo(k)fluoranthene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
bis(2-Chloroethoxy)methane	*. * *	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
bis(2-Chloroethyl) ether		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
bis(2-Ethylhexyl) phthalate		ug/L	8.0 U	1.5 U	1.1 U	1.0 U
Butyl benzyl phthalate		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Caprolactam	·	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Carbazole		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Chrysene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Dibenz(a,h)anthracene		ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Dibenzofuran		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Diethyl phthalate		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Dimethyl phthalate		ug/L ug/L	1.0 U	1.0 U	1.0 U	1.0 U
		ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Di-n-butyl phthalate			1.0 U	1.0 U	1.0 U	1.0 U
Di-n-octyl phthalate		ug/L	0.20 U	0.48	0.20 U	0.20 U
Fluoranthene Fluorene		ug/L ug/L	0.20 U	0.43	0.20 U	0.20 U

Table 2-6 Analytical Data Summary Site 23 Water Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6H120107001 23-MW01 08/10/2006	A6H120107002 23-MW02 08/10/2006	A6H120107003 23-MW03 08/10/2006	A6H120107004 23-MW04 08/10/2006
Hexachlorobenzene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Hexachlorobutadiene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U	10 U	10 U	10 U
Hexachloroethane	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U
Isophorone	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Naphthalene	ug/L	0.20 U	1.8	0.20 U	0.20 U
Nitrobenzene	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Pentachlorophenol	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Phenanthrene	ug/L	0.20 U	1.1	0.20 U	0.20 U
Phenoi	ug/L	1.0 U	1.0 U	1.0 U	1.0 U
Pyrene	ug/L	0.20 U	0.20 U	0.20 U	0.20 U

 $<sup>\</sup>ensuremath{\mathsf{U}}$  = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

Table 2-7
Analytical Data Summary
Site 23 Water Metals
ODOT Innerbelt Study

PARAMETER	UNITS	A6H120107001 23-MW01 08/10/2006	A6H120107002 23-MW02 08/10/2006	A6H120107003 23-MW03 08/10/2006	A6H120107004 23-MW04 08/10/2006
Arsenic	ug/L	10.0 U	6.1 J	5.7 J	10.0 U
Barium	ug/L	57.7 J	47.4 J	51.8 J	59.5 J
Cadmium	ug/L	2.0 U	2.0 U	2.0 U	2.0 U
Chromium	ug/L	3.5 J	4.0 J	5.0 U	5.0 U
Lead	ug/L	3.0 U	3.0	2.0 J	3.0 U
Selenium	ug/L	5.0 U	5.0 U	5.0 U	5.3
Silver	ug/L	5.0 U	5.0 U	5.0 U	5.0 U
Mercury	ug/L	0.20 U	0.20 U	0.20 U	0.20 U

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

### 6.13 SITE 24 – JAMES VINCENT

According to the Cleveland City Directories, the site was listed as the Pump and Ice Company from 1954 through 1999.

According to the Cleveland Fire Prevention Bureau files, the site was occupied by the American Vinegar Company in 1941. In 1964, the site was occupied by the Pump & Ice Machine Company Inc.

At the time of the URS Phase I reconnaissance, the site was occupied by the Pump Ice Machine Company. No surface staining, stressed vegetation and/or the handling and storage of hazardous materials were observed on the site.

#### 6.13.1 Field Activities

A total of three monitoring wells were proposed for Site 24. Groundwater was encountered in three soil borings (MW01, MW02, and MW03) during field activities and monitoring wells were installed. Three soil and groundwater samples were collected and analyzed for VOCs and SVOCs. A Sample Location Map is included as **Figure 6-13A**.

## 6.13.2 Site-Specific Geology/Hydrogeology

Soils at Site 24 consisted primarily of sand, silty sand, and silty clay with minor amounts of slag and gravel. Bedrock was not encountered in any of the soil borings, which were advanced to 16 feet bgs.

Groundwater elevations were measured at Site 24 on August 11, 2006. Localized groundwater flow across Site 24 is likely influenced by both natural features and urban development, including paved surfaces, buildings, and underground utilities. The general flow direction at Site 24 is to the south-southwest, towards the Cuyahoga River.

# 6.13.3 Soil Analytical Results

A total of three VOCs were detected in two soil samples submitted from Site 24. The chemicals detected in the samples submitted were 2-butanone (2.4 ug/kg to 23 ug/kg), acetone (66 ug/kg), and carbon disulfide (3.3 ug/kg). All other VOCs were below the detection limits.

A total of 20 SVOCs were detected in two samples submitted from Site 24. Concentrations of 2-methylnaphthalene (1,200 ug/kg and 7,900 ug/kg), 4-methylphenol / 3-methylphenol (220 ug/kg), acenaphthene (6,000 ug/kg), anthracene (120 ug/kg and 19,000 ug/kg), benzaldahyde (210 ug/kg), benzo(a)anthracene (320 ug/kg and 36,000 ug/kg), benzo(a)pyrene (220 ug/kg and 30,000 ug/kg), benzo(b)fluoranthene (260 ug/kg and 36,000 ug/kg), benzo(g,h,i)perylene (190 ug/kg and 18,000 ug/kg), benzo(k)fluoranthene (130 ug/kg and 15,000 ug/kg), carbazole (6,100 ug/kg), chrysene (300 ug/kg and 32,000 ug/kg), dibenz(a,h)anthracene (4,100 ug/kg), dibenzofuran (240 ug/kg and 6,400 ug/kg), fluoranthene (460 ug/kg and 85,000 ug/kg), fluorene (8,500 ug/kg), indeno(1,2,3)pyrene (120 ug/kg and 15,000 ug/kg), naphthalene (660 ug/kg and

**Phase II Findings** 

6,900 ug/kg), phenanthrene (630 ug/kg and 68,000 ug/kg), and pyrene (490 ug/kg and 70,000 ug/kg) were detected in the soil samples submitted.

The analytical results are presented in Table 6-13A.

## 6.13.4 Groundwater Analytical Results

Toluene (0.20 ug/L) was detected in a groundwater sample collected from Site 24. All other VOCs were below the detection limits.

No SVOCs were detected in the groundwater samples submitted from Site 24.

The analytical results are presented in Table 6-13B.

### 6.13.5 Comparison Standards

The analytical results were compared to the OEPAs VAP Generic Direct-Contact Standards for Commercial and Industrial Land Use (VAP, 2002a), the Construction and Excavation Worker Activities Category (VAP, 2002b), the Generic Unrestricted Potable Use Standards Based on MCLs or Other Regulatory Established Criteria (VAP, 2002c), the Risk-Derived Generic Unrestricted Potable Use Standards (VAP, 2002d), and the BUSTR Closure Action Levels. The VAP and BUSTR standards are included on **Tables 6-13A** and **6-13B**.

There were no VOCs detected in any of the soil samples from Site 24, which exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

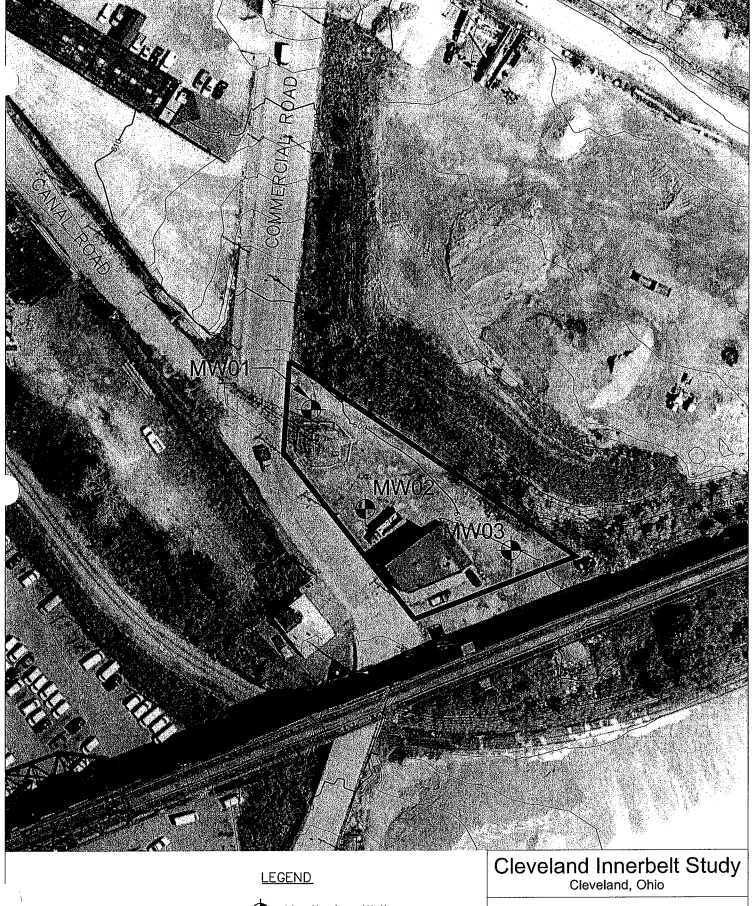
The concentrations of benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, dibenz(a,h)anthracene, and indeno(1,2,3-cd)pyrene detected in sample 24-MW030204 exceeded the OEPA VAP standards for commercial and industrial land use, the construction and excavation worker activities category, or the BUSTR closure action levels.

No VOCs detected in the groundwater samples from Site 24 exceeded the OEPA VAP standard for generic unrestricted potable use standards or BUSTR closure action levels.

### 6.13.6 Conclusions

Based on the analytical results, the soil at Site 24 may require special disposal and/or worker protection protocols (plan note) during construction activities.







- Monitoring Well

## Figure 6-13A

Sample Location Map, Site #24 2515 Canal Road

URS

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Dec 18, 2006 - 4:54pm

**TABLES** 

Table 0=13A
Summary of Detected Chemicals in Soil
Site 24 - James Vincent
ODOT Innerbelt Study
Cleveland, Ohio

<u>a</u>	PARAMETER	SLIND	Commercial E Industrial Standard	Construction  Worker Standard <sup>21</sup>	EUSTR Cosure Action Lavel	24-MW01-0608 07/18/2006	24-MW02-1416 07/19/2006	24-MW03-0204 07/19/2006
s	2-Butanone	ug/kg	71,600,000	80,000,000	1	23 J	2.4 J	24 U
၁၀	Acetone	ug/kg	100,000,000	100,000,000		99	25 U	24 U
۸	Carbon disulfide	ug/kg	720,000	720,000		3.3 J	6.2 U	6.0 U
	2-Methylnaphthalene	ug/kg	Ī		•	1200 J	410 U	ቦ 0062
	4-Methylphenol / 3-Methylphenol	ug/kg	1			ב 220	410 U	40000 U
	Acenaphthene	ug/kg	180,000,000	530,000,000		2000 U	410 U	۲ 0009
	Anthracene	ug/kg	880,000,000	1,000,000,000	The state of the s	120 J	410 U	ر 19000
	Benzaldehyde	ug/kg	•	•	1	210 J	410 U	40000 U
	Benzo(a)anthracene	ug/kg	63,000	810,000	11,000	320 J	410 U	36000 J
	Benzo(a)pyrene	ug/kg	6,300	81,000	1,100	220 J	410 U	30000
	Benzo(b)fluoranthene	ug/kg	63,000	810,000	11,000	260 J	410 U	36000 J
	Benzo(ghi)perylene	ug/kg		1	1	190 J	410 U	ر 18000 ع
300	3	ug/kg	630,000	8,100,000	110,000	130 J	410 U	15000 J
λS	: :	ug/kg	10,000,000	31,000,000	1 :	2000 U	410 U	6100 J
		ug/kg	6,700,000	41,000,000	1,100,000	300 J	410 U	32000 J
	Dibenz(a,h)anthracene	ug/kg	6,700	41,000	1,100	2000 U	410 U	4100 J
	Dibenzofuran	ug/kg				240 J	410 U	6400 J
	Fluoranthene	ug/kg	33,000,000	170,000,000	•	460 J	410 U	85000
	Fluorene	ug/kg	120,000,000	340,000,000	•	2000 U	410 U	8500 J
	Indeno(1,2,3-cd)pyrene	ug/kg	67,000	410,000	11,000	120 J	410 U	15000 J
	Naphthalene	ug/kg	530,000	1,900,000	39,800	r 099	410 U	r 0069
	Phenanthrene	ug/kg	•		The state of the s	630 J	410 U	00089
	Pyrene	ug/kg	25,000,000	130,000,000		490 J	410 U	20000

<sup>-- =</sup> Standard not available

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

<sup>(1)</sup> VAP Generic Direct Contact Soil Standard, Commercial/ Industrial Land Use

<sup>(2)</sup> VAP Generic Direct Contact Soil Standard, Construction and Excavation Activities

Table v-13B

## Summary of Detected Chemicals in Water Site 24 - James Vincent ODOT Innerbelt Study Cleveland, Ohio

24-MW03 08/11/2006	1.0 U
24-MW02 08/11/2006	0.20 ئ
24-MW01 08/10/2006	1.0 U
SBUSTRIC Closure Action	1000
VAPUPUS' RDUPUS <sup>(0)</sup>	1000
UNITS	ug/L
PARAMETER	Toluene
	VOCs

-- = Standard not available

U = The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

(1) VAP Unrestricted Potable Use and Risk-Derived Unrestricted Potable Use Standards

**BORING LOGS** 

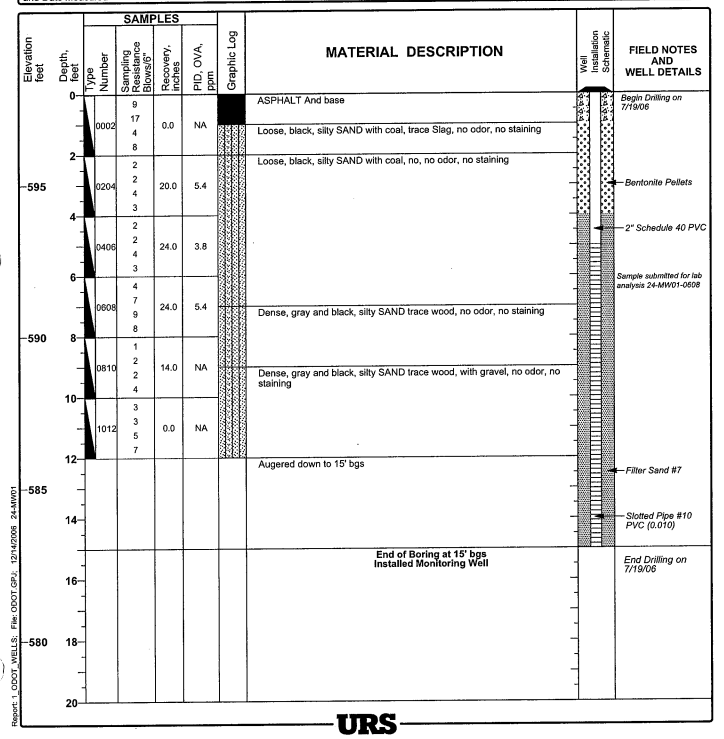
Project: ODOT - Innerbelt Corridor

Project Location: Site 24
Project Number: 15016633

## Log of Boring 24-MW01

Sheet 1 of 1

Logged By	J. Kaminski	Checked M. Wolff
Hammer Data	140 lb Hammer 30 inches	Total Depth of Borehole 15.0' bgs
Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 598'
Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
	Hammer Data Drilling Contractor Sampling	By 3. Kammer  Hammer Data Drilling Contractor Sampling 2" Split Speen



**Project: ODOT - Innerbelt Corridor** 

Project Location: Site 24
Project Number: 15016633

## Log of Boring 24-MW02

Sheet 1 of 1

Date(s) 7/19/06 Drilled	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140 lb Hammer 30 inches	Total Depth of Borehole 30.0' bgs
Drill Rig Type CME-55	Drilling Contractor	HAD, Inc.	Approximate Ground Elevation 599'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well

			SAME	LES		Ī		Т		Т	
Elevation feet	Depth, feet	Type Number	Sampling Resistance Blows/6"	Recovery, inches	PID, OVA, ppm	Graphic Log	MATERIAL DESCRIPTION	Well	Installation	Schematic	FIELD NOTES AND WELL DETAILS
	0	0002	25 15 11 10	18.0	1.5		Loose, dry, dark brown and black, medium to coarse SAND with gravel, with Slag, trace coal refuse	4.4.		4.4	Begin Drilling on 7/19/06
	-	0204	3 4 4 2	24.0	2.4		Loose, dry, dark brown and black, medium to coarse SAND with gravel, with Slag, trace Brick		•		– 2" Schedule 40 PVC – Bentonite Pellets
	5— -	0406	4	18.0	0.7		Dense, moist, gray and black, fine clayey SAND				
	-	0608	5 3	24.0	0.1		Dense, moist, greenish gray, fine to medium SAND				
-590	10-	0810	7	24.0	0.0		Hard, dry, gray and black sandy CLAY, with brick				
	-	1012	6	20.0	0.0		Medium stiff, moist, greennish gray clayey SILT				
	- 15–	1416	10 5 5	24.0	4.0		Stiff, moist, brown SILT, trace sand				Sample submitted for lab analysis 24-MW02-1416
	-	1618	7 2	24.0	3.2		r-With clay				
-580	-	1820	7 4	24.0	0.9		Hard, dry, reddish brown silty CLAY ←Becomes gray		<b>V</b> 60000000		
	20-	2022	9 5 5 3	0.0	NA		-	-			
	-	2224	3 1 7 9 10	0.0	1.8		Dense, moist, gray silty SAND, trace clay				
7 0007	25-	2426	4	24.0	1.1		rNo clay				— Filter Sand #7
2007 1 21 10 20 1 1 1 1 1 1 1 1 1 1 1 1 1 1	-	2628	8	24.0	0.9			-			Slotted Pipe #10
<b>570</b>	30–	2830	2 3 5 5	24.0	0.9		End of Boring at 30' bgs	-			PVC (0.010)  End Drilling on
,		-					Installed Monitoring Well	1			7/19/06
	-							-			
r poor	35-	l		L	I	J	URS				

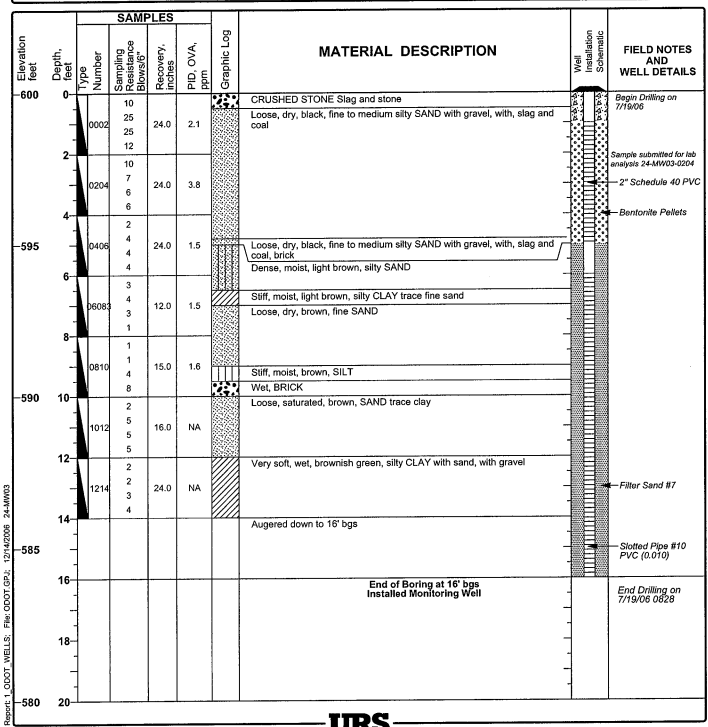
Project: ODOT - Innerbelt Corridor

Project Location: Site 24
Project Number: 15016633

### Log of Boring 24-MW03

Sheet 1 of 1

Date(s) 7/19/06 Drilled 7/19/06	Logged By	J. Kaminski	Checked M. Wolff
Drilling Method & Hollow Stem Auger Drill Bit size/type 4-1/4" ID HSA	Hammer Data	140 lb Hammer 30 inches	Total Depth of Borehole 16.0' bgs
Drill Rig CME-55 Type	Drilling Contractor	HAD, Inc.	Approximate 600'
Location See Site Map	Sampling Method(s)	2" Split Spoon	Borehole Completion Set monitoring well
Location See Site Map  Groundwater Level 591.20 on 8/11/06 and Date Measured	Method(s)	2 Split Spoon	



DATA ASSESSMENT REPORT

#### Data Assessment Report ODOT Innerbelt Study Site 24 – James Vincent

Reviewer: P. Schuler Date: November 14, 2006

Three soil samples and three groundwater samples were collected at the James Vincent site at 2515 Canal Road in Cleveland, Ohio, from July 18 through August 11, 2006. The samples were submitted to Severn Trent Laboratories, Inc. in North Canton, Ohio, for analysis of the parameters listed in Table 1.

Table 1
Sample and Analysis Summary

		Sample	2 - 1	-	ested yses <sup>(1)</sup>
Laboratory ID	Sample ID	Date	Matrix	VOC	SVOC
A6G190360001	24-MW01-0608	07/18/2006	Soil	X	X
A6G190360002	24-MW03-0204	07/19/2006	Soil	X	X
A6G190360003	24-MW02-1416	07/19/2006	Soil	X	X
A6H120112001	24-MW01	08/10/2006	Groundwater	X	X
A6H120112002	24-MW02	08/11/2006	Groundwater	X	X
A6H120112003	24-MW03	08/11/2006	Groundwater	X	X

(1) VOC = Volatile Organic Compounds [SW-846 Method 8260B] SVOC = Semivolatile Organic Compounds [SW-846 Method 8270C]

A standard review for analytical data quality was performed by URS Corporation (URS) for the above referenced samples. A standard review includes assessment of supporting quality control (QC) parameters such as associated laboratory control sample (LCS) recoveries, laboratory and field blank results, surrogate recoveries, internal standard responses, matrix spike/matrix spike duplicate recoveries, field duplicate results, detection limits, and holding times. A standard review does not include examination of the raw data or reconstruction of the analytical results. The significant findings (findings that resulted in qualification of data or otherwise affected data quality) were as follows:

- Positive detections for acetone in samples 24-MW03-0204 and 24-MW02-1416, for bis(2-ethylhexyl)phthalate in samples 24-MW02-1416, 24-MW01, and 24-MW03, and for caprolactam in sample 24-MW02 were qualified as nondetect ("U") due to the presence of these analytes in the associated method blanks and/or trip blanks at similar concentrations.
- One volatile internal standard response was outside of the acceptance range in sample 24-MW03-0204. The results for all volatile analytes quantified from each of the noncompliant internal standards were qualified as estimated ("J" or "UJ").

- The lab reported results below their reporting limit but above the method detection limit (MDL) with a qualifier ("J"), in accordance with USEPA Contract Laboratory Program (CLP) conventions. During the data assessment, the "J" qualifiers were retained with the numeric results.
- All soil analytical results were reported on a dry weight (moisture-corrected) basis.

No other significant findings were identified and all data are considered usable. The analytical results, with qualifiers, are summarized in Tables 2-1 through 2-4.

# Table 2-1 Analytical Data Summary Site 24 Soil Volatiles ODOT Innerbelt Study

		A6G190360001 24-MW01-0608 07/18/2006	A6G190360002 24-MW03-0204 07/19/2006	A6G190360003 24-MW02-1416 07/19/2006
PARAMETER	UNITS		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	07/19/2006
Percent Solids	%	81.5	82.7	80.9
1,1,1-Trichloroethane	ug/kg	6.1 U	6.0 U	6.2 U
1,1,2,2-Tetrachloroethane	ug/kg	6.1 U	6.0 UJ	6.2 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/kg	6.1 U	6.0 U	6.2 U
1,1,2-Trichloroethane	ug/kg	6.1 U	6.0 U	6.2 U
1,1-Dichloroethane	ug/kg	6.1 U	6.0 U	6.2 U
1,1-Dichloroethene	ug/kg	6.1 U	6.0 U	6.2 U
1,2,4-Trichlorobenzene	ug/kg	6.1 U	6.0 UJ	6.2 U
1,2-Dibromo-3-chloropropane	ug/kg	12 U	12 UJ	12 U
1,2-Dibromoethane	ug/kg	6.1 U	6.0 U	6.2 U
1,2-Dichlorobenzene	ug/kg	6.1 U	6.0 UJ	6.2 U
1,2-Dichloroethane	ug/kg	6.1 U	6.0 U	6.2 U
1,2-Dichloropropane	ug/kg	6.1 U	6.0 U	6.2 U
1,3-Dichlorobenzene	ug/kg	6.1 U	6.0 UJ	6.2 U
1,4-Dichlorobenzene	ug/kg	6.1 U	6.0 UJ	6.2 U
2-Butanone	ug/kg	23 J	24 U	2.4 J
2-Hexanone	ug/kg	25 U	24 U	25 U
4-Methyl-2-pentanone	ug/kg	25 U	24 UJ	25 U
Acetone	ug/kg	66	24 U	25 U
Benzene	ug/kg	6.1 U	6.0 U	6.2 U
Bromodichloromethane	ug/kg	6.1 U	6.0 U	6.2 U
Bromoform	ug/kg	6.1 U	6.0 U	6.2 U
Bromomethane	ug/kg	6.1 U	6.0 U	6.2 U
Carbon disulfide	ug/kg	3.3 J	6.0 U	6.2 U
Carbon tetrachloride	ug/kg	6.1 U	6.0 U	6.2 U
Chlorobenzene	ug/kg	6.1 U	6.0 U	6.2 U
Chloroethane	ug/kg	6.1 U	6.0 U	6.2 U
Chloroform	ug/kg	6.1 U	6.0 U	6.2 U
Chloromethane	ug/kg	6.1 U	6.0 U	6.2 U
cis-1,2-Dichloroethene	ug/kg	6.1 U	6.0 U	6.2 U
cis-1,3-Dichloropropene	ug/kg	6.1 U	6.0 U	6.2 U
Cyclohexane	ug/kg	12 U	12 U	12 U
Dibromochloromethane	ug/kg	6.1 U	6.0 U	6.2 U
Dichlorodifluoromethane	ug/kg	6.1 U	6.0 U	6.2 U
Ethylbenzene	ug/kg	6.1 U	6.0 U	6.2 U
Isopropylbenzene	ug/kg	6.1 U	6.0 U	6.2 U
Methyl acetate	ug/kg	12 U	12 U	12 U
Methyl tert-butyl ether	ug/kg	25 U	24 U	25 U
Methylcyclohexane	ug/kg	12 U	12 U	12 U
Methylene chloride	ug/kg	6.1 U	6.0 U	6.2 U
Styrene	ug/kg	6.1 U	6.0 U	6.2 U
Tetrachloroethene	ug/kg	6.1 U	6.0 U	6.2 U
Toluene	ug/kg	6.1 U	6.0 U	6.2 U
trans-1,2-Dichloroethene	ug/kg	6.1 U	6.0 U	6.2 U
trans-1,3-Dichloropropene	ug/kg	6.1 U	6.0 U	6.2 U
Trichloroethene	ug/kg	6.1 U	6.0 U	6.2 U
Trichlorofluoromethane	ug/kg	6.1 U	6.0 U	6.2 U
Vinyl chloride	ug/kg	6.1 U	6.0 U	6.2 U
Xylenes (total)	ug/kg	12 U	12 U	12 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

UJ = The analyte was not detected at or above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.



J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-2 Analytical Data Summary Site 24 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G190360001 24-MW01-0608 07/18/2006	A6G190360002 24-MW03-0204 07/19/2006	A6G190360003 24-MW02-1416 07/19/2006
1,1'-Biphenyl	ug/kg	2000 U	40000 U	410 U
2,2'-oxybis(1-Chloropropane)	ug/kg	2000 U	40000 U	410 U
		2000 U	40000 U	410 U
2,4,5-Trichlorophenol	ug/kg	2000 U	40000 U	410 U
2,4,6-Trichlorophenol	ug/kg	2000 U	40000 U	410 U
2,4-Dichlorophenol	ug/kg			410 U
2,4-Dimethylphenol	ug/kg	2000 U	40000 U	5 1
2,4-Dinitrophenol	ug/kg	2000 U	40000 U	410 U
2,4-Dinitrotoluene	ug/kg	2000 U	40000 U	410 U
2,6-Dinitrotoluene	ug/kg	2000 U	40000 U	410 U
2-Chioronaphthalene	ug/kg	2000 U	40000 U	410 U
2-Chlorophenol	ug/kg	2000 U	40000 U	410 U
2-Methylnaphthalene	ug/kg	1200 J	7900 J	410 U
2-Methylphenol	ug/kg	2000 U	40000 U	410 U
2-Nitroaniline	ug/kg	2000 U	40000 U	410 U
2-Nitrophenol	ug/kg	2000 U	40000 U	410 U
3,3'-Dichlorobenzidine	ug/kg	2000 U	40000 U	410 U
3-Nitroaniline	ug/kg	2000 U	40000 U	410 U
4,6-Dinitro-2-methylphenol	ug/kg	2000 U	40000 U	410 U
4-Bromophenyl phenyl ether	ug/kg	2000 U	40000 U	410 U
4-Chloro-3-methylphenol	ug/kg	2000 U	40000 U	410 U
4-Chloroaniline	ug/kg	2000 U	40000 U	410 U
4-Chlorophenyl phenyl ether	ug/kg	2000 U	40000 U	410 U
4-Methylphenol / 3-Methylphenol	ug/kg	220 J	40000 U	410 U
4-Nitroaniline	ug/kg	2000 U	40000 U	410 U
4-Nitrophenol	ug/kg	` 2000 U	40000 U	410 U
Acenaphthene	ug/kg	2000 U	6000 J	410 U
Acenaphthylene	ug/kg	2000 U	40000 U	410 U
Acetophenone	ug/kg	410 U	8100 U	82 U
Anthracene	ug/kg	120 J	19000 J	410 U
Atrazine	ug/kg	2000 U	40000 U	410 U
Benzaldehyde	ug/kg	210 J	40000 U	410 U
Benzo(a)anthracene	ug/kg	320 J	36000 J	410 U
Benzo(a)pyrene	ug/kg	220 J	30000 J	410 U
Benzo(b)fluoranthene	ug/kg	260 J	36000 J	410 U
Benzo(ghi)perylene	ug/kg	190 J	18000 J	410 U
Benzo(k)fluoranthene	ug/kg	130 J	15000 J	410 U
bis(2-Chloroethoxy)methane	ug/kg	2000 U	40000 U	410 U
bis(2-Chloroethyl) ether	ug/kg	2000 U	40000 U	410 U
bis(2-Ethylhexyl) phthalate	ug/kg	2000 U	40000 U	410 U
Butyl benzyl phthalate	ug/kg	2000 U	40000 U	410 U
Caprolactam	ug/kg	2000 U	40000 U	410 U
Carbazole	ug/kg	2000 U	6100 J	410 U
Chrysene	ug/kg	300 J	32000 J	410 U
Dibenz(a,h)anthracene	ug/kg	2000 U	4100 J	410 U
Dibenzofuran	ug/kg	240 J	6400 J	410 U
Diethyl phthalate	ug/kg	2000 U	40000 U	410 U
Dimethyl phthalate	ug/kg	2000 U	40000 U	410 U
Di-n-butyl phthalate	ug/kg	2000 U	40000 U	410 U
Di-n-octyl phthalate	ug/kg ug/kg	2000 U	40000 U	410 U
Fluoranthene	ug/kg ug/kg	460 J	85000	410 U
Control of the Contro	ug/kg ug/kg	2000 U	8500 J	410 U
Fluorene	ug/kg ug/kg	2000 U	40000 U	410 U

Table 2-2 Analytical Data Summary Site 24 Soil Semivolatiles ODOT Innerbelt Study

PARAMETER	UNITS	A6G190360001 24-MW01-0608 07/18/2006	A6G190360002 24-MW03-0204 07/19/2006	A6G190360003 24-MW02-1416 07/19/2006
Hexachlorobutadiene	ug/kg	2000 U	40000 U	410 U
Hexachlorocyclopentadiene	ug/kg	2000 U	40000 U	410 U
Hexachioroethane	ug/kg	2000 U	40000 U	410 U
Indeno(1,2,3-cd)pyrene	ug/kg	120 J	15000 J	410 U
Isophorone	ug/kg	2000 U	40000 U	410 U
Naphthalene	ug/kg	660 J	6900 J	410 U
Nitrobenzene	ug/kg	2000 U	40000 U	410 U
N-Nitrosodi-n-propylamine	ug/kg	2000 U	40000 U	410 U
N-Nitrosodiphenylamine	ug/kg	2000 U	40000 U	410 U
Pentachlorophenol	ug/kg	2000 U	40000 U	410 U
Phenanthrene	ug/kg	630 J	68000	410 U
Phenol	ug/kg	2000 U	40000 U	410 U
Pyrene	ug/kg	490 J	70000	410 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-3
Analytical Data Summary
Site 24 Water Volatiles
ODOT Innerbelt Study

PARAMETER	UNITS	A6H120112001 24-MW01 08/10/2006	A6H120112002 24-MW02 08/11/2006	A6H120112003 24-MW03 08/11/2006
1.1.1-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U
1,1,2,2-Tetrachloroethane	ug/L	1.0 U	1.0 U	1.0 U
1,1,2-Trichloro-1,2,2-trifluoroethane	ug/L	1.0 U	1.0 U	1.0 U
1,1,2-Trichloroethane	ug/L	1.0 U	1.0 U	1.0 U
1.1-Dichloroethane	ug/L	1.0 U	1.0 U	1.0 U
1,1-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U
1,2,4-Trichlorobenzene	ug/L	1.0 U	1.0 U	1.0 U
1,2-Dibromo-3-chloropropane	ug/L	2.0 U	2.0 U	2.0 U
1,2-Dibromoethane	ug/L	1.0 U	1.0 U	1.0 U
1,2-Diolinoemane	ug/L	1.0 U	1.0 U	1.0 U
1,2-Dichloroethane	ug/L	1.0 U	1.0 U	1.0 U
		1.0 U	1.0 U	1.0 U
1,2-Dichloropropane	ug/L	1.0 U	1.0 U	1.0 U
1,3-Dichlorobenzene	ug/L ug/L	1.0 U	1.0 U	1.0 U
1,4-Dichlorobenzene	1 1 1 1 1 1 1 1	1.0 U	1.0 U	1.0 U
2-Butanone	ug/L	10 U	10 U	10 U
2-Hexanone	ug/L	4 44 A	10 U	10 U
4-Methyl-2-pentanone	ug/L	10 U		•
Acetone	ug/L	10 U	10 U	10 U
Benzene	ug/L	1.0 U	1.0 U	1.0 U
Bromodichloromethane	ug/L	1.0 U	1.0 U	1.0 U
Bromoform	ug/L	1.0 U	1.0 U	1.0 U
Bromomethane	ug/L	1.0 U	1.0 U	1.0 U
Carbon disulfide	ug/L	1.0 U	1.0 U	1.0 U
Carbon tetrachloride	ug/L	1.0 U	1.0 U	1.0 U
Chlorobenzene	ug/L	1.0 U	1.0 U	1.0 U
Chloroethane	ug/L	1.0 U	1.0 U	1.0 U
Chloroform	ug/L	1.0 U	1.0 U	1.0 U
Chloromethane	ug/L	1.0 U	1.0 U	1.0 U
cis-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U
cis-1,3-Dichloropropene	ug/L	1.0 U	1.0 U	1.0 U
Cyclohexane	ug/L	1.0 U	1.0 U	1.0 U
Dibromochloromethane	ug/L	1.0 U	1.0 U	1.0 U
Dichlorodifluoromethane	ug/L	1.0 U	1.0 U	1.0 U
Ethylbenzene	ug/L	1.0 U	1.0 U	1.0 U
Isopropylbenzene	ug/L	1.0 U	1.0 U	1.0 U
Methyl acetate	ug/L	10 U	10 U	10 U
Methyl tert-butyl ether	ug/L	5.0 U	5.0 U	5.0 U
Methylcyclohexane	ug/L	1.0 U	1.0 U	1.0 U
Methylene chloride	ug/L	1.0 U	1.0 U	1.0 U
Styrene	ug/L	1.0 U	1.0 U	1.0 U
Tetrachloroethene	ug/L	1.0 U	1.0 U	1.0 U
Toluene	ug/L	1.0 U	0.20 J	1.0 U
trans-1,2-Dichloroethene	ug/L	1.0 U	1.0 U	1.0 U
trans-1,3-Dichloropropene	ug/L	1.0 U	1.0 U	1.0 U
Trichloroethene	ug/L	1.0 U	1.0 U	1.0 U
Trichlorofluoromethane	ug/L	1.0 U	1.0 U	1.0 U
A CHARLES AND A STATE OF THE ST	1.1 1.35 July 1	1.0 U	1.0 U	1.0 U
Vinyl chloride Xylenes (total)	ug/L ug/L	2.0 U	2.0 U	2.0 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.

J = Estimated concentration because the result was below the sample reporting limit or quality control criteria were not met.

Table 2-4
Analytical Data Summary
Site 24 Water Semivolatiles
ODOT Innerbelt Study

		A6H120112001 24-MW01 08/10/2006	A6H120112002 24-MW02 08/11/2006	A6H120112003 24-MW03 08/11/2006
PARAMETER	UNITS			
1,1'-Biphenyl	ug/L	1.0 U	1.0 U	1.0 U
2,2'-oxybis(1-Chloropropane)	ug/L	1.0 U	1.0 U	1.0 U
2,4,5-Trichlorophenol	ug/L	5.0 U	5.0 U	5.0 U
2,4,6-Trichlorophenol	ug/L	5.0 U	5.0 U	5.0 U
2,4-Dichlorophenol	ug/L	2.0 U	2.0 U	2.0 U
2,4-Dimethylphenol	ug/L	2.0 U	2.0 U	2.0 U
2,4-Dinitrophenol	ug/L	5.0 U	5.0 U	5.0 U
2,4-Dinitrotoluene	ug/L	5.0 U	5.0 U	5.0 U
2,6-Dinitrotoluene	ug/L	5.0 U	5.0 U	5.0 U
2-Chloronaphthalene	ug/L	1.0 U	1.0 U	1.0 U
2-Chlorophenol	ug/L	1.0 U	1.0 U	1.0 U
2-Methylnaphthalene	ug/L	0.20 U	0.20 U	0.20 U
2-Methylphenol	ug/L	1.0 U	1.0 U	1.0 U
2-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U
2-Nitrophenol	ug/L	2.0 U	2.0 U	2.0 U
3,3'-Dichlorobenzidine	ug/L	5.0 U	5.0 U	5.0 U
3-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U
4,6-Dinitro-2-methylphenol	ug/L	5.0 U	5.0 U	5.0 U
4-Bromophenyl phenyl ether	ug/L	2.0 U	2.0 U	2.0 U
4-Chloro-3-methylphenol	ug/L	2.0 U	2.0 U	2.0 U
4-Chloroaniline	ug/L	2.0 U	2.0 U	2.0 U
4-Chlorophenyl phenyl ether	ug/L	2.0 U	2.0 U	2.0 U
4-Methylphenol	ug/L	1.0 U	1.0 U	1.0 U
4-Nitroaniline	ug/L	2.0 U	2.0 U	2.0 U
4-Nitrophenol	ug/L	5.0 U	5.0 U	5.0 U
Acenaphthene	ug/L	0.20 U	0.20 U	0.20 U
Acenaphthylene	ug/L	0.20 U	0.20 U	0.20 U
Acetophenone	ug/L	1.0 U	1.0 U	1.0 U
Anthracene	ug/L	0.20 U	0.20 U	0.20 U
Atrazine	ug/L	1.0 U	1.0 U	1.0 U
Benzaldehyde	ug/L	1.0 U	1.0 U	1.0 U
Benzo(a)anthracene	ug/L	0.20 U	0.20 U	0.20 U
Benzo(a)pyrene	ug/L	0.20 U	0.20 U	0.20 U
Benzo(b)fluoranthene	ug/L	0.20 U	0.20 U	0.20 U
Benzo(ghi)perylene	ug/L	0.20 U	0.20 U	0.20 U
Benzo(k)fluoranthene	ug/L	0.20 U	0.20 U	0.20 U
bis(2-Chloroethoxy)methane	ug/L	1.0 U	1.0 U	1.0 U
bis(2-Chloroethyl) ether	ug/L	1.0 U	1.0 U	1.0 U
bis(2-Ethylhexyl) phthalate	ug/L	1.0 U	1.0 U	2.4 U
Butyl benzyl phthalate	ug/L	1.0 U	1.0 U	1.0 U
Caprolactam	ug/L	5.0 U	5.0 U	5.0 U
Carbazole	ug/L	1.0 U	1.0 U	1.0 U
Chrysene	ug/L	0.20 U	0.20 U	0.20 U
Dibenz(a,h)anthracene	ug/L	0.20 U	0.20 U	0.20 U
Dibenzofuran	ug/L	1.0 U	1.0 U	1.0 U
Diethyl phthalate	ug/L	1.0 U	1.0 U	1.0 U
Dimethyl phthalate	ug/L	1.0 U	1.0 U	1.0 U
Di-n-butyl phthalate	ug/L	1.0 U	1.0 U	1.0 U
Di-n-octyl phthalate	ug/L	1.0 U	1.0 U	1.0 U
Fluoranthene	ug/L	0.20 U	0.20 U	0.20 U
Fluorene	ug/L	0.20 U	0.20 U	0.20 U
Hexachlorobenzene	ug/L	0.20 U	0.20 U	0.20 U

Table 2-4
Analytical Data Summary
Site 24 Water Semivolatiles
ODOT Innerbelt Study

PARAMETER	UNITS	A6H120112001 24-MW01 08/10/2006	A6H120112002 24-MW02 08/11/2006	A6H120112003 24-MW03 08/11/2006
Hexachlorobutadiene	ug/L	1.0 U	1.0 U	1.0 U
Hexachlorocyclopentadiene	ug/L	10 U	10 U	10 U
Hexachloroethane	ug/L	1.0 U	1.0 U	1.0 U
Indeno(1,2,3-cd)pyrene	ug/L	0.20 U	0.20 U	0.20 U
Isophorone	ug/L	1.0 U	1.0 U	1.0 U
Naphthalene	ug/L	0.20 U	0.20 U	0.20 U
Nitrobenzene	ug/L	1.0 U	1.0 U	1.0 U
N-Nitrosodi-n-propylamine	ug/L	1.0 U	1.0 U	1.0 U
N-Nitrosodiphenylamine	ug/L	1.0 U	1.0 U	1.0 U
Pentachlorophenol	ug/L	5.0 U	5.0 U	5.0 U
Phenanthrene	ug/L	0.20 U	0.20 U	0.20 U
Phenol	ug/L	1.0 U	1.0 U	1.0 U
Pyrene	ug/L	0.20 U	0.20 U	0.20 U

U =The analyte was analyzed for, but was not detected. Value shown is the reporting limit.