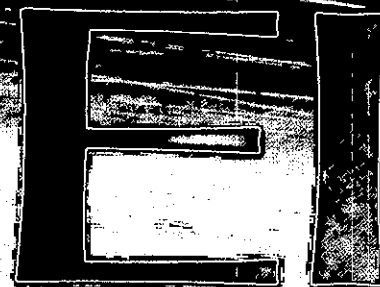
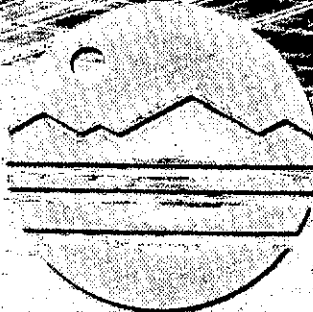
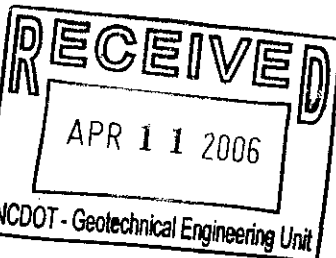


# Environmental ENVIRONMENTAL Environmental



## LIMITED PRELIMINARY SITE ASSESSMENT



**Parcel 127**  
**James David McCourry Property**  
**120 Mill Springs Road**  
**Burnsville, NC 28714**

State Project No. R-2519A  
WBS Element No. 35609.1.1  
EI Project No. ENMO060029.00

Prepared For:

Gregory A. Smith  
State of North Carolina  
Department of Transportation  
Geotechnical Unit  
GeoEnvironmental Section  
1589 Mail Service Center  
Raleigh, NC 27699-1589

Prepared by:

Environmental Investigations, Inc.  
2101 Gateway Centre Boulevard, Suite 200  
Morrisville, NC 27560  
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April 2006

## LIMITED PRELIMINARY SITE ASSESSMENT (PSA)

Conducted on

Parcel 127

James David McCourry Property  
120 Mill Springs Road  
Burnsville, NC 28714  
State Project No. R-2519A  
WBS Element No. 35609.1.1  
EI Project No. ENMO060029.00

For

Mr. Gregory A. Smith  
State of North Carolina  
Department of Transportation  
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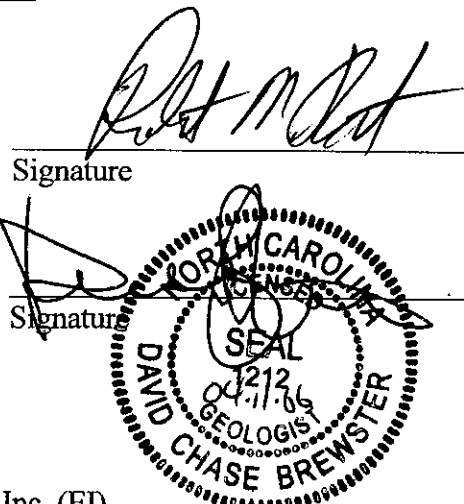
Issue Date: April 11, 2006

Robert M. Shaut  
Project Geologist/Manager

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Prepared By:

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**Appendix D: Laboratory Analytical Results Report**

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## 1.0 INTRODUCTION

Environmental Investigations, Inc. (EI) conducted a *Limited Preliminary Site Assessment* (PSA) within the existing and/or proposed North Carolina Department of Transportation (NCDOT) *right-of-way* (ROW) adjacent to a parcel (identified by the NCDOT as Parcel 127) located at 120 Mill Springs Road, Burnsville, North Carolina 28714. A residence is currently located on the adjacent parcel. The report presented herein documents the findings of the PSA that was conducted within the described ROW. For purposes of this report, the terms subject site and/or site include the existing NCDOT ROW and the proposed ROW, and/or the abutting property/parcel.

### 1.1 Report Organization

Field activities were conducted by Mr. Robert Michael Shaut, and Sterling Turner, Environmental Geologists with EI, on February 23, 2005 and March 29, 2006, respectively. The report presented herein summarizes the scope of work conducted, discusses sampling procedures, and presents our findings, conclusions and recommendations. A table entitled "Summary of Soil Analytical Results" is presented in **Table 1**, and a table entitled "Summary of Groundwater Analytical Results", is presented in **Table 2**. A "Site Location Map", a "Site Map" and "Extent of Residual Petroleum Impact (Vadose Zone) Map" are presented in **Figures 1, 2, and 3**, respectively. A compilation of "Site Photographs" are presented in **Appendix A**, the "Standard Field Operating Procedures (SOP)" are presented in **Appendix B**, "Soil Boring Logs" are included in **Appendix C**, while an "Analytical Laboratory Report" is presented in **Appendix D**.

### 1.2 Background

Mr. Eugene Tarascio, GeoEnvironmental Project Manager with the NCDOT GeoTechnical Engineering Unit submitted to EI a "*Request for Technical and Cost Proposal*" (RFP), dated February 7, 2006. The RFP solicited a technical and cost proposal to perform Limited PSAs on a total of six (6) Parcels located within a NCDOT Highway Project, identified as WBS Element #35609.1.1, State Project #R-2519A, located in Burnsville, NC. The RFP outlined site information on each of the six (6) parcels and NCDOT Figures (Plan Sheets) and applicable site photographs were attached to the RFP. Mr. Gregory A. Smith, LG, PE, GeoEnvironmental Supervisor with the NCDOT, GeoTechnical Engineering Unit, GeoEnvironmental Section authorized EI to perform the PSAs, as documented in a "Notice to Proceed" dated February 16, 2006 (verbal authorization was granted on February 10, 2006).

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State Project: R-2519A  
WBS Element: 35609.1.1120

Limited Preliminary Site Assessment  
Parcel 127 – James David McCourry Property  
Mill Springs Road, Burnsville, NC 28714

---

### 1.3 Objectives

The objective of performing the PSA was to determine if an existing residential heating oil UST has impacted the subsurface of the existing and/or proposed ROW. The study (PSA) on the referenced parcel (Parcel 127 – James David McCourry Property) included herein was performed with a reasonable effort to investigate and quantify potentially petroleum-hydrocarbon residual impacted subsurface soils. However, findings documented in the report do not constitute a guarantee that all potential source of (petroleum) environmental contamination have been assessed and subsequently analyzed.

This report is provided for the sole use of the NCDOT on the project for which it was prepared. All materials and information used for this project were obtained by EI, Inc. Use of this report by any third parties other than the NCDOT will be at such party's sole risk. EI Inc. disclaims liability for any use of or reliance on this report by third parties.

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Parcel 127 – James David McCourry Property  
Mill Springs Road, Burnsville, NC 28714

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## 2.0 SCOPE OF WORK & ENVIRONMENTAL SERVICES

### 2.1 Requested Scope of Work

Documented in the RFP, dated February 7, 2006, the NCDOT requested the following scope of work:

- Determine if contaminated soils are present around the heating oil UST;
- delineate and estimate the quantity of impacted soils and indicate the approximate area of soil contamination on a site map for the site;
- if groundwater is encountered and the project manager suspects the possibility of groundwater contamination, obtain a sample for analysis by converting one (1) of the borings to a temporary monitoring well;
- for each groundwater sample collected, also obtain a 24-hour groundwater depth; and
- prepare a report including field activities, findings, and recommendations for the site and submit in quadruplet to this office.

### 2.2 Scope of Services

To accomplish the scope-of-services, a field reconnaissance was performed to identify general site conditions, and hand augers and Direct Push Technology (DPT) were utilized to collect soil samples on the subject parcel.

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Parcel 127 – James David McCourry Property  
Mill Springs Road, Burnsville, NC 28714

To perform the requested Limited PSA, EI personnel visited the site on two (2) occasions to supervise, oversee and/or perform site reconnaissance activities and collect appropriate samples to complete the project objectives. To complete the study on the subject parcel, EI performed the following scope of services:

- Advancement of three (3) soil test borings utilizing a hand auger to a total depth of less than 2.74 meters (9.0 feet) below the land surface (bls) in the vicinity of the heating oil UST.
- Supervision, and oversight of the advancement of two (2) soil test borings utilizing DPT methods to depths of less than 3.66 meters (12.0 feet) bls in the vicinity of the heating oil UST.
- Collection and submittal of four (4) soil samples for laboratory analytical testing.
- Installation of one (1) temporary monitoring well (piezometer).
- Collection and submittal of one (1) groundwater sample for laboratory analyses
- Photo documentation of pertinent site features.
- Preparation of this *Limited PSA Report*, presenting our findings and conclusions along with our recommendations.

---

## 3.0 SITE CHARACTERIZATION

### 3.1 Site Location

A residence known as the James David McCourry property is currently located at Mill Springs Road, in Burnsville, North Carolina (**Figure 1**). The subject property is currently located immediately adjacent to the DOT ROW (**Photograph 1**) as identified in DOT's R-2519A Plan Sheet. Copies of digital site photographs are presented in **Appendix A**.

### 3.2 Physical Setting

The subject site parcel currently consists of a residence. The parcel consists of a one-story house a concrete driveway, while the house is surrounded by a fenced yard consisting of grass and/or shrubbery. Please refer to **Figure 2, Site Map** for the location of the residence.

#### 3.2.1 Number and Capacities of USTs

A heating oil UST is located on the east side of the house near the southeast house corner. A vent and fill port for the tank (**Photograph 2**) was visible and according to the NCDOT, the property owner has stated that the UST holds a capacity of 2,082 liters (550 gallons).

### 3.3 Site Topography

Site observations and review of the Burnsville, NC United States Geological Survey (USGS) Topographic Quadrangle Map (July 1, 1984) revealed that the subject site is located at an elevation of approximately 798 meters (2,620 feet) above mean sea level (msl) (**Figure 1**). Topographically, the site slopes gently to the southwest. Surface water runoff appears to flow directly southwest in the direction of Little Crabtree Creek located approximately 107 meters (350) feet from the site.

### 3.4 Land Use & Surrounding Properties

The subject property is located inside the city limits of Burlington, NC. Land use in the immediate vicinity of the site is characterized by residential properties. The site is bounded on the north by SR 1329, to the east by a residential property, to the south by an undeveloped parcel and to the west by an undeveloped parcel.

## 4.0 SUBSURFACE INVESTIGATION

### 4.1 Subsurface Soils Investigation

On March 29, 2006, an EI Geologist advanced a total of three (3) soil test borings in the vicinity of an existing residential heating oil UST with a hand-auger. Troxler Geologic Services, based in Raleigh, North Carolina, was selected and subcontracted to provide Direct Push Technology (DPT) services. On February 23, 2006, EI directed and supervised the advancement of two (2) soil test borings (GP-1 and GP-2), both in a downgradient direction approximately 10.00 meters (33 feet) from the subject UST.

In general, the borings were advanced in order to evaluate the absence/presence of potential subsurface soil (vadose zone) impact and/or subsurface groundwater (petroleum smearing) impact associated with potential petroleum releases associated with either former and/or present UST system spills and/or releases into the subsurface. The soil borings were advanced to investigative depths ranging from 0.31 meters (1.0 feet) to a depth of 3.66 meters (12.0 feet) bls.

### 4.2 Soil Test Boring Methodology

A complete descriptive explanation of EI's *Standard Field Operating Procedures* that discusses specific sampling methodology is presented in **Appendix B**.

### 4.3 Soil Sample Collection Procedures

Based on the results of site conditions (i.e., location of UST system), a total of four (4) soil samples were collected for laboratory retention from the five (5) soil test borings conducted at the property. Weathered rock and gravel were encountered approximately 10.00 meters (33 feet) downgradient from the subject UST, as both of the borings advanced in this area encountered auger refusal, approximately at a depth of 2.66 meters (12.0 feet) bls.

Soil samples retained for laboratory analysis were transferred to a representative of Paradigm Analytical Laboratory, for laboratory analytical testing. Dates and times of sample shipment may be referenced in the analytical Chain-of Custodies (COC) presented in **Appendix D**.

### 4.4 Backfill Activities

At the completion of the exploratory subsurface advancement activities, the test borings were backfilled to surface grade. A complete descriptive explanation of EI's *Standard Field Operating Procedures* that discusses backfill procedures is presented in **Appendix B**.

#### **4.5 Subsurface Soil Lithology**

During boring advancement activities, soil samples were classified in the field by an EI geologist utilizing the Unified Soil Classification System (USCS). Subsurface soils encountered in the area of study were fairly consistent. The on-site geology consists of grass with surficial topsoil from the surface to approximately 1.0-foot below grade. Layers of soil consisting of tan to light brown clayey SILT (ML) silty CLAY in some areas were encountered to the investigated depth of approximately 2.74 meters (9.0 feet) below the land surface (bls).

Detailed descriptions are presented in Soil Boring Logs included in **Appendix C**. The boring logs include an interpretation of subsurface conditions based on field samples.

#### **4.6 Groundwater Investigation**

##### **4.6.1 Temporary Monitoring Well Installation**

During the field study (February 23, 2006), soil test boring “GP-2” (located approximately 10.00 meters (33 feet) downgradient from the subject UST) was converted into a Type I (temporary) 2.54 cm (1.0 inch) diameter groundwater monitoring well (piezometer). The approximate location of the groundwater monitoring well is depicted in **Figure 2**. The well location was selected in the field by the EI Field Geologist (Robert Shaut) based on the topographic location of the boring and due to the fact that drilling equipment could not be employed adjacent to the subject UST because of site restrictions (fence). The well was advanced to the approximate depth of 3.66 meters (12.0 feet) bls.

##### **4.6.2 Monitoring Well Sampling**

On the date of installation, EI personnel collected a groundwater sample from the referenced temporary monitoring well (“GP-2”) for purposes of analytical testing. On February 23, 2006, the samples were transferred to representatives of Paradigm Analytical Laboratories for analytical laboratory testing. Groundwater sampling procedures are discussed in more detail in the *Standard Operating Procedures* presented in **Appendix C**.

The groundwater table was measured in the temporary monitoring well (“GP-2”) on February 24, 2006. Groundwater was measured at approximately 2.304 meters (7.56 feet) below the top of casing. The casing depth was level to the ground surface.

---

## 5.0 LABORATORY TESTING AND RESULTS

### 5.1 Subsurface Soil Analytical Methods

A total of four (4) soil samples (“PAR 127 GP1-8”, “HA-1”, “HA-2”, and “HA-3”) were submitted for total petroleum hydrocarbons (TPH) analyses by Method 8015B with preparation methods for the analysis of Diesel Range Organics (DRO) by GC-FID and Gasoline Range Organics (GRO) by GC-FID. The GRO method is utilized to extract volatile fuels such as gasoline, while the DRO method is utilized to extract less volatile petroleum products such as diesel fuel, No. 2 fuel oil, kerosene, and varsol.

### 5.2 Soil Laboratory Analyses Results

Laboratory analysis of soil samples collected from all four (4) soil samples collected did not detect DRO or GRO concentrations above the laboratory detection limits. The specific results of the analytical testing of the soil samples are tabulated and presented in **Table 1**. The complete laboratory results and Chain-of-Custody Records are presented in **Appendix D**.

### 5.3 Groundwater Laboratory Analytical Methods

Groundwater sample “GP-2” collected from the referenced temporary well was submitted for VOCs analysis for aromatic and halogenated volatiles by GC/PID-ELCD for EPA Method 6230D + IPE & MTBE, for semivolatile organic compounds by GC/MS for EPA Method 625 and the top ten peaks identified, for extractable petroleum hydrocarbons by GC/FID by Method MADEP EPH, and for volatile petroleum hydrocarbons by GC-PID/FID by MADEP VPH.

### 5.4 Groundwater Laboratory Analyses Results

Review of the groundwater analytical data showed concentrations of only two (2) VOC analytes, at or above the method laboratory detection limits. The detected analytes (naphthalene at 0.946 ug/L, and toluene at 1.01 ug/L) **were not reported above** the current North Carolina Groundwater Standards (15A NCAC 2L .0202). Aliphatics, aromatics and SVOCs were not reported above the laboratory method reporting limits. A summary of the analytical results is tabulated in **Table 2**.

---

## 6.0 SUMMARY OF FINDINGS

EI has reviewed information gathered during the Limited PSA study including the site reconnaissance activities, review of DOT plan sheets, review of the site investigation including soil collection activities, and review of a laboratory analyses report. Compiled below is a summarized list of the significant findings.

- Groundwater was encountered beneath the site at a location approximately 10.00 meters (33 feet) downgradient from the subject UST at a depth of 2.304 meters (7.56 feet) below the top of casing.
- Analyses of four (4) soil samples advanced within the *proposed* NCDOT ROW, collected at a depth of approximately 2.44 meters (8.0 feet) bbl, which is a depth that is beneath the bottom depth of the tank did not report concentrations of DRO or GRO above the method laboratory reporting limits or the NCDENR action limits of 10.0 mg/kg.
- Analysis of a groundwater sample collected from a temporary monitoring well installed at the site within the *proposed* NCDOT ROW did not show concentrations of all tested constituents (aliphatics, aromatics, VOCs and/or SVOCs) above the current North Carolina Groundwater Standards (15A NCAC 2L .0202).

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Limited Preliminary Site Assessment  
Parcel 127 – James David McCourry Property  
Mill Springs Road, Burnsville, NC 28714

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## 7.0 CONCLUSIONS AND RECOMMENDATIONS

EI personnel have reviewed information obtained during the *Limited PSA* at the site (Parcel 127) and present the following conclusions and recommendations.

It does not appear based on laboratory analytical data that any significant petroleum spills and/or releases have occurred in the vicinity of the heating oil UST. Based on the findings of this investigation, EI does not recommend any further assessment at this time.

*Note: This report does not constitute a guarantee that all potential sources of environmental contamination have been assessed and subsequently analyzed.*

**TABLES**

**TABLE 1**  
**SUMMARY OF SOIL ANALYTICAL RESULTS**  
 Parcel 127  
 James David McCourry Property  
 120 Mill Springs Road, Burnsville, NC 28714  
 State Project No. R-2519A  
 WBS Element No. 35609.1.1

Sample Identification	PAR 127 GP1-8	HO-1	HO-2	HO-3
Sample Depth Meters (Feet)	2.134m-2.438m (7'-8')	2.134m-2.438m (7'-8')	2.134m-2.438m (7'-8')	2.134m-2.438m (7'-8')
Sample Date	2/23/2006		3/29/2006	
Field Screening Results-PID (ppm)	0.0	0.1	0.0	0.0
<b>LABORATORY RESULTS (mg/kg)</b>				
Laboratory Analysis (Total Petroleum Hydrocarbons by GC/FID #015)	NCDENR <sup>1</sup> (Volume II) Reportable Concentration (mg/kg)			
Gasoline Range Organics	BQL	BQL	BQL	BQL
Diesel Range Organics	10	BQL	BQL	BQL

NOTE:

<sup>1</sup>mg/kg denotes parts per million

NCDENR = North Carolina Department of Environment & Natural Resources

**TABLE 2**  
**SUMMARY OF GROUNDWATER ANALYTICAL RESULTS**  
**Parcel 127 - James David McCourry Property**  
**120 Mill Springs Road**  
**Burnsville, NC 28714**  
**State Project: R-25190A**  
**WBS Element: 35609.1.1**

Sample Identification		GP-2
Groundwater Depth (feet below top of casing)		7.56
Sample Date		2/23/2006
Volatiles Method 6230D	2L Groundwater Standards (ug/L)	Laboratory Results (ug/L)
Benzene	1	BQL
sec-Butylbenzene	70	BQL
Diisopropyl ether (DIPE)	NS	BQL
Ethylbenzene	29	BQL
Isopropylbenzene	70	BQL
Methyl-tert butyl ether (MTBE)	200	BQL
Naphthalene	21	0.946
N-propylbenzene	70	BQL
Total Xylenes	530	BQL
Toluene	1,000	1.01
1,2,4-Trimethylbenzene	350	BQL
1,3,5-Trimethylbenzene	350	BQL
MTBE	200	BQL
Remaining Analytes	N/A	BQL
MADEP VPH	2L GW Standards (ug/L)	Laboratory Results (ug/L)
C5-C8 Aliphatics	420	<100
C9-C12 Aliphatics	4200	<100
C9-C10 Aliphatics	210	<100
MADEP EPH	2L GW Standards (ug/L)	Laboratory Results (ug/L)
C9-C18 Aliphatics	4200	<100
C19-C36 Aliphatics	42000	<100
C11-C22 Aromatics	210	<100
Semivolatiles - GCMS Method 625		Laboratory Results (ug/L)
Fluorene	280	BQL
Acenaphthene	80	BQL
Acenaphthylene	210	BQL
Bis (2-ethylhexyl) Phthalate	NS	BQL
Di-n-Butylphthalate	700	BQL
Diethylphthalate	5000	BQL
Dimethylphthalate	NS	BQL
Fluoranthene	280	BQL
Fluorene	280	BQL
Naphthalene	21	BQL
Phenanthrene	210	BQL
Pyrene	210	BQL
All Remaining Analytes	N/A	BQL

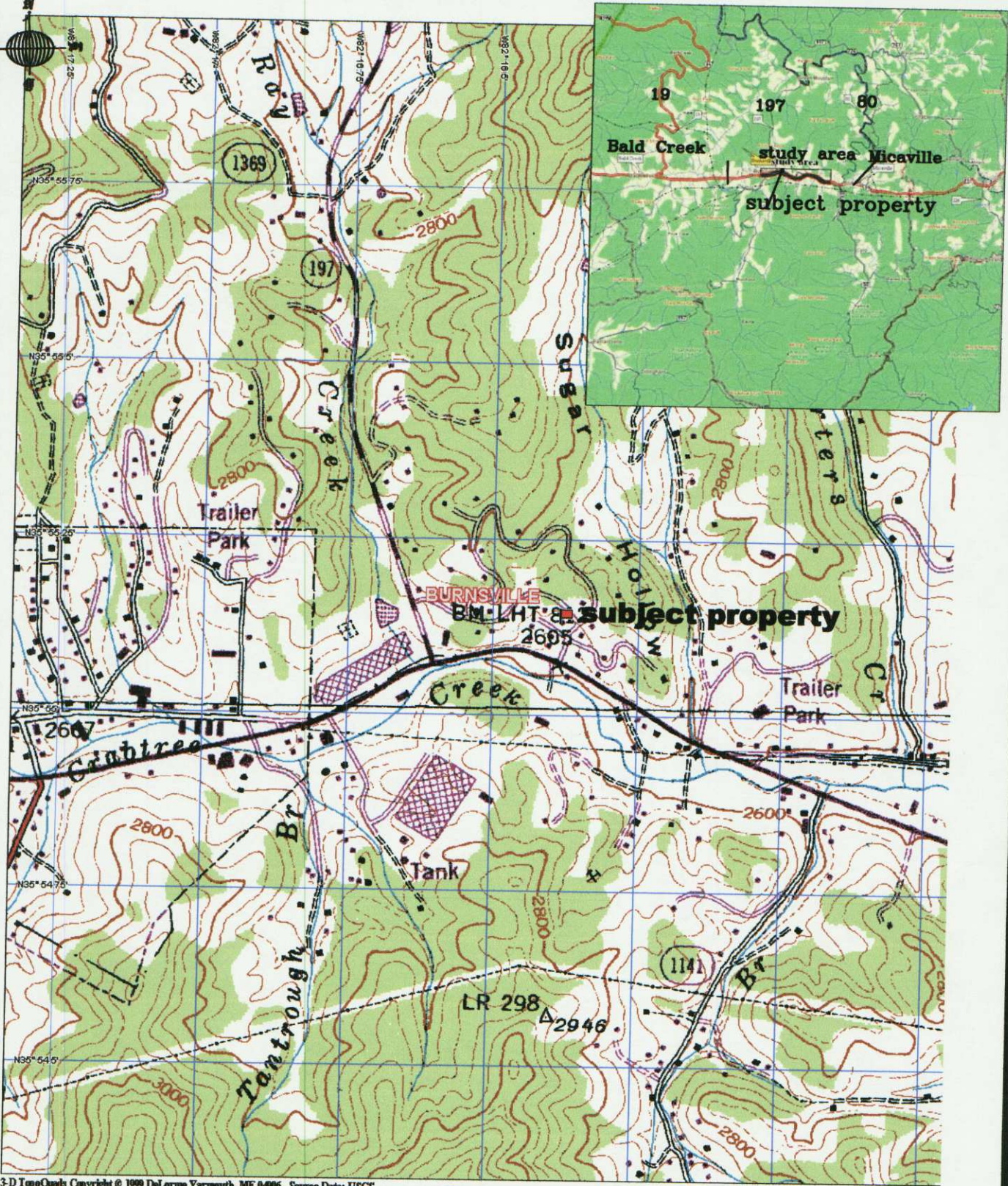
**Legend:**

BQL = Below Quantitation Limit

NA = Not Applicable

NS = No Standard

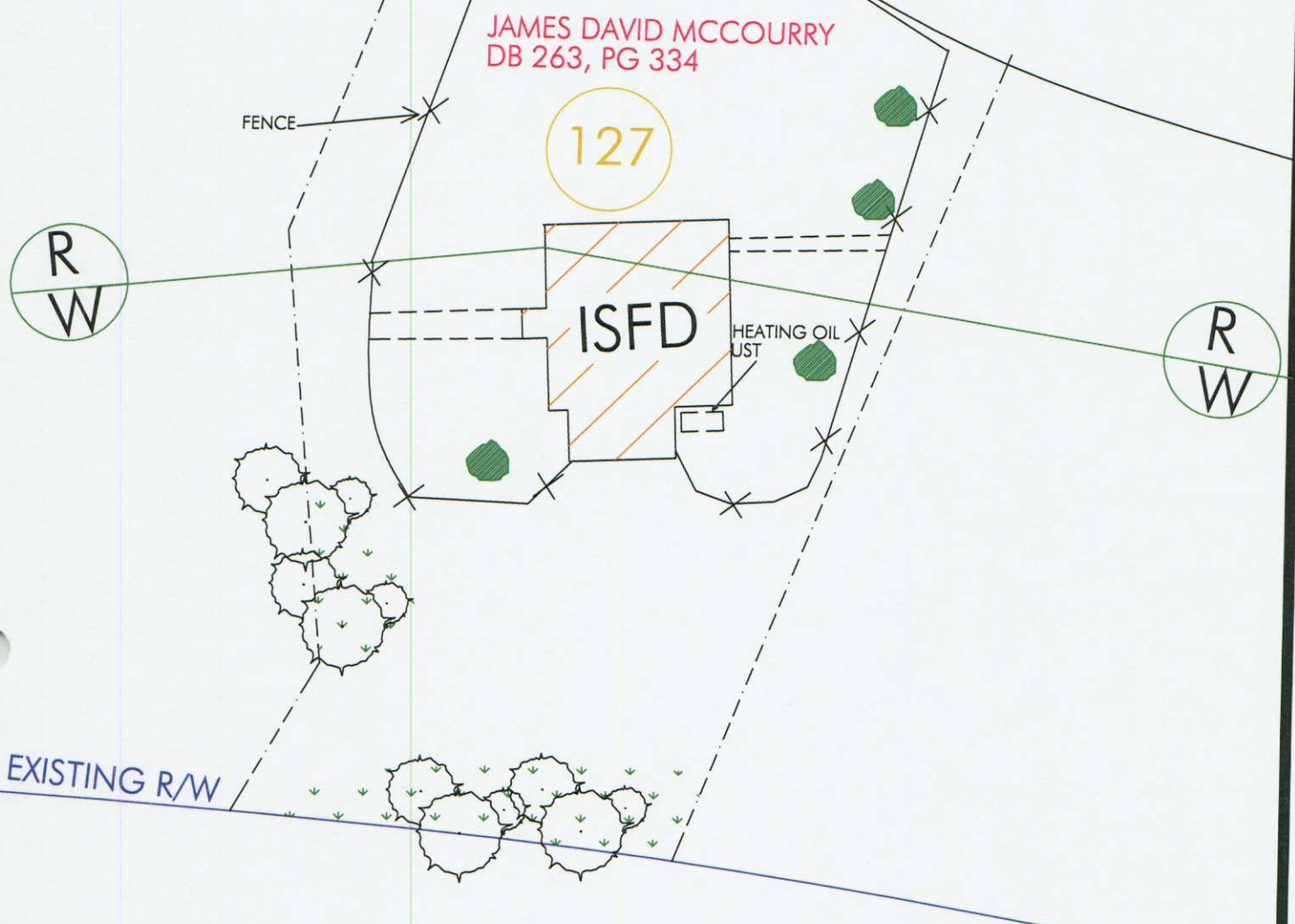
## **FIGURES**



DWN NO.	FIGURE 1
JOB NO.:	ENM0060029.00
DRAWN BY:	USGS/RMS
CHECKED BY:	DCB
DATE:	2/21/06
SCALE:	as shown

**SITE LOCATION MAP**  
**PARCEL 127**  
James David McCurry Property  
120 Mill Springs Road  
Burnsville, NC 28714  
State Project: R-2519A



**LEGEND:**

- Shrubbery
- Property Line
- Proposed Right-of-Way
- Existing Right-of-Way

5 0 10  
Scale in Meters

OWN NO.	FIGURE 2
JOB NO.:	ENM0060029.00
DRAWN BY:	RMS
CHECKED BY:	DCB
DATE:	2/21/06
SCALE:	1cm = 80m

SITE MAP  
PARCEL 127  
James David McCourry Property  
120 Mill Springs Road  
Burnsville, NC 28714  
State Project: R-2519A

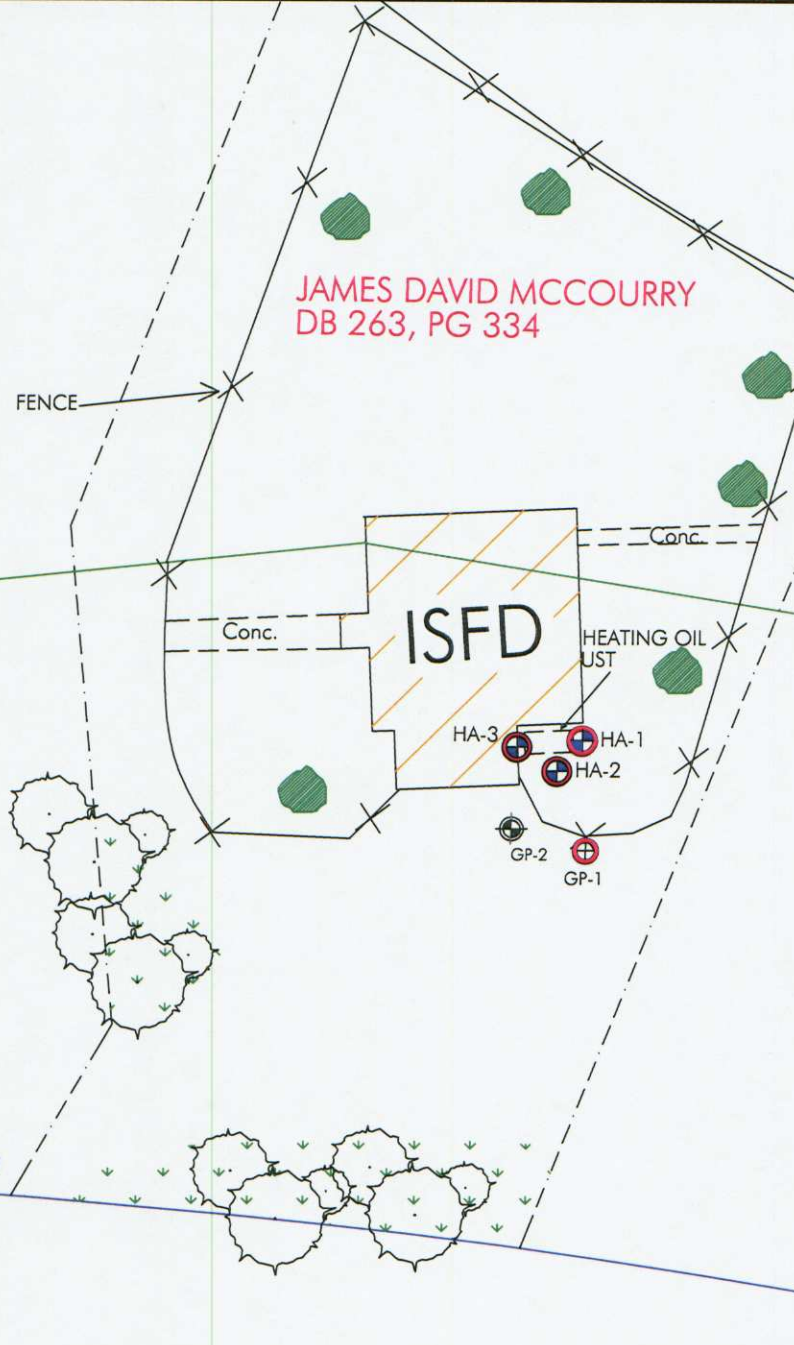




JAMES DAVID MCCOURRY  
DB 263, PG 334

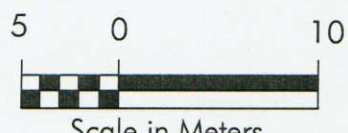


EXISTING R/W



**LEGEND:**

- Property Line
- Bushes
- Proposed Right-of-Way
- Existing Right-of-Way
- DPT Soil Test Boring
- Hand Auger Soil Test Boring
- Temporary Monitoring Well
- HA-1
- MW-2
- Estimated Extent of Residual Petroleum Impact  
TPH Concentrations Diesel  
(Vadose Zone)



OWN NO.	FIGURE 3
JOB NO.:	ENM0060029.00
DRAWN BY:	RMS
CHECKED BY:	DCB
DATE:	2/21/06
SCALE:	1cm = 40m

EXTENT OF RESIDUAL PETROLEUM IMPACT (VADOSE ZONE) MAP  
PARCEL 127  
James David McCourry Property  
120 Mill Springs Road  
Burnsville, NC 28714  
State Project: R-2519A



**EI**

ENVIRONMENTAL INVESTIGATIONS, INC.

**APPENDIX A**  
**SITE PHOTOGRAPHS**



Photograph 1: Looking west at the subject property.



Photograph 2: A closer view, note fill port and vent pipe.  
Flags outline tank boundaries.

**APPENDIX B**

**STANDARD OPERATING PROCEDURES**

**STANDARD OPERATING PROCEDURES  
Subsurface Assessment Methodology And Sampling Protocol**

**Parcel 127  
James David McCourry Property  
120 Mill Springs Road  
Burnsville, NC 28714**

**WBS Element # 35609.1.1  
State Project # R-2519A  
EI Project No. ENMO060029.00**

**Prepared For:**

**Gregory A. Smith  
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**Prepared by:**

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PH (919) 544-7500 FAX (919) 544-2199**

**March 2006**

**STANDARD OPERATING PROCEDURES**  
**(Subsurface Assessment Methodology And Sampling Protocol)**

**INTRODUCTION**

Environmental Investigations, Inc. (EI) has prepared this STANDARD OPERATING PROCEDURES - Subsurface Assessment Methodology and Sampling Protocol Plan (SPP) for a residential property owned by Paul D. Biggerstaff property located at 84 Mount View Drive, Burnsville, Yancey County, North Carolina.

The document presented herein describes the methodology and protocol that was utilized during the *Limited Preliminary Site Assessment* conducted at the above referenced project "site".

**SAMPLING DESIGN**

Prior to conducting a subsurface assessment, a sampling strategy was developed by EI based on the objectives of the investigation. After designing our soil sampling strategy, the appropriate equipment and techniques were selected to conduct the investigation. Our sampling strategy was based upon the premise of accomplishing the following performance objectives:

- collect soil samples that are representative of conditions as they exist at the study site;
- selecting the appropriate sampling device(s);
- taking measures to avoid introducing contamination as a result of poor sampling and/or poor handling techniques;
- reducing the potential of cross contamination between samples;
- defining sampling site selections and collection procedures for the appropriate individual media;
- defining the quality control assurance procedures;
- analytical requirements and limitations; and
- Data interpretation and assessment.

The sampling plan for this study was developed using the non-probabilistic (directed sampling designs) in nature. The location and frequency was based on this approach, to allow for the flexibility of the field coordinator (Geologist) to determine the number of samples collected for analysis. This approach allowed for the study objectives, properties of the matrix, resource constraints and access to sampling points to be adequately performed. Provision for access, use of sampling equipment, was also pre-determined.

The following section of the SPP discusses the sampling equipment available and collection methods which have been utilized to be technically appropriate.

## **SITE ORIENTATION**

Prior to conducting any soil sampling procedures, the EI Project Geologist/Manager reviewed and presented the Site and Safety Health Plan to all participants involved with the project which was developed based on the EI Safety and Health program. All monitoring, protective equipment (latex gloves, Tyvek® suits, etc.), potential hazards associated with the site and general health and safety standards were discussed.

### **Site Survey**

Prior to conducting specific sampling activities, EI personnel will conduct a limited site survey of the target and surrounding areas. Information discovered during the survey will be utilized to better perform the sampling activities and will provide more insight into establishment of the conclusions of this study. The site survey will consist of the following:

- General site layout (UST system layouts, overhead canopies, dispensers, etc.);
- Site access;
- Soil types and depths;
- Surface water drainage pathways;
- Existing site conditions;
- Visible staining of surface soil;
- Vegetation stress, and
- Possible offsite or non-site related sources.

## **FIELD INVESTIGATIVE PROCEDURES**

### **Sampling Objectives**

The general objective of sampling for this project was to collect a sample representative of subsurface and/or groundwater to reduce the potential bias caused by the sampling equipment used to obtain the sample.

The chosen sample locations were evaluated as discrete samples. A discrete sample is defined as "a discrete aliquot representative of a specific location at a given point in time."

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### **Areas of Environmental Concern**

The objectives of choosing the proper sampling methods to collect appropriate samples that are representative of the conditions as they exist at the site were as follows:

- Selecting the appropriate sampling device.
- Taking measures to avoid introducing contamination as a result of poor sampling and/or handling techniques.
- Reducing the potential of cross contamination between samples.

The areas of environmental concern consisted of an existing heating oil UST.

### **SOIL SAMPLING ACTIVITIES**

Manual techniques and equipment, such as hand augers, are usually used for surface or shallow, subsurface soil sampling. Power operated equipment is usually associated with collecting deep samples, but this equipment can also be used for collecting shallow samples when the auger hole begins to collapse, or when the soil is so tight that manual auguring is not practical. Based on the request of the property owner, EI mainly used hand augers and to a lesser extent we utilized Direct Push Technology (DPT). The following section discusses the DPT methods employed during the site study.

#### **Soil Sampling Collection Methods**

Soil samples were collected utilizing either Hand-auger or Direct Push Technology (DPT) methods.

#### **Direct Push Technology Methodology**

DPT refers to tools and sensors that are inserted into the subsurface without the use of drilling to remove soil and make a path for the tool. To perform the DPT activities, the contractor utilized a GeoProbe® 6600 machine. The GeoProbe® 6600 is a hydraulically-powered probing machine designed, which uses static force and a percussion hammer to advance small diameter sampling tools into the subsurface to collect soil cores, groundwater samples, and or soil gas samples. A GeoProbe relies on a relatively small amount of static (vehicle) weight combined with percussion as the energy for advancement of a tool string.

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The advantages of utilizing DPT drilling methods are described as follows:

- avoids the use of drilling fluids and lubricants during drilling;
- the equipment is highly mobile;
- disturbance of geochemical conditions during installation is minimized; and
- The drilling process does not produce drill cuttings.

### **DPT Soil Sample Collection Methods**

Soil samples utilizing DPT methods were collected from the advanced DPT soil borings continuously in 4.0-foot increments using acetate liners contained in a nickel plated macro sampling tubes. Each soil-filled liner was split for field screening and soil sample collection purposes. Soil samples were collected from the liners with disposable vinyl gloves and utilized for soil vapor screening testing and/or laboratory retention. This sampling method allows for continuous soil sampling from the ground surface to the desired depth. Soil samples selected for analyses are referenced in the text section.

### **Soil Sample Collection Protocol**

The following soil sampling collection procedures were utilized during this study:

- Ensured that all equipment, samplers and tools that will come in contact with the sample media was thoroughly decontaminated.
- Informed driller of sample interval (s) for borehole and oversaw the sampling process.
- Prepared and labeled all sample containers. Samples collected for the analytes of volatiles (if applicable) were sampled first.
- Labeled the containers including the location, depth, analyte, date and time of sampling.
- Delegated the driller to prepare the sample liner by cutting the liner in half.
- Placed liners on a clean sheet of plastic.
- Cut the soil core with a clean decontaminated knife to allow of visual soil classification.
- Sniffed the soil core with a PID/FID and recorded instrument readings volatile organics (VOCs) in a logbook (discussed further below).
- Logged the soil core in a logbook, including borehole identification (ID), sample number, date, time and any pertinent data.

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- Logged soil classification including: recording percent recovery, color, description of major constituent, soil texture/structure, grading/sorting/plasticity, relative density or hardness consistency, clay, sand, silt, gravel content, grain size, moisture content, odor, staining and the Unified Soil Classification System (USCS) identifier and symbol;
- Physically collected the selected soil samples and placed these samples into laboratory prepared containers.
- Ensured the soil sample did not contain twigs, stones, and other debris from the soil.
- Packed soil samples for shipment, prepared chain-of-custody records and shipping documentation

### **Soil Vapor Screening**

An important tool in performing this study is performing the soil vapor screening or sniffing activities. Field screening is generally performed for a variety of reasons. The technique conducted during this study was used to screen soil samples for measurable levels of volatile organics. The results obtained from this procedure are not quantitative; however the results from several soil samples are relative and allowed the Field Geologist/Project Manager to select samples that are the most contaminated with the contaminated media. Generally, the presence of little or no organic vapor is possibly indicative of non-contaminated soils. Soil samples collected for purposes of soil headspace screening were tested by the following procedures:

- the field instrument was calibrated, prior to use;
- soil samples were collected directly from the DPT soil liners and placed into sealable plastic bags;
- soil samples within the bags were allowed to equilibrate for approximately five minutes;
- the headspace of each bagged sample was screened with the instrument probe for the presence of volatile organic compound (VOCs) with a Mini-RAE Photo-ionization Detector (PID);
- recording the instrument readings (VOCs) in a field logbook; and
- Verified that the FID/PID was reading background levels prior to exposing the probe into another sample.

### **Collection of Grab Soil Samples**

Soil samples may provide two (2) types of soil contamination representation including grab and composite. Samples may be generally collected in random locations from a grid pattern or selected areas believed to be contaminated as evidenced by field indicators (staining, odors and/or measurable volatile organic readings).

For this study, grab samples selected from areas showing field indicators or confirmation soil samples chosen to confirm the absence of volatile organic readings were chosen. The technical definition for a grab sample is as follows: A grab sample is a discrete aliquot representative of a specific location at a given point in time. The sample is collected at one time and at one particular sampling point and depth. Refer to the text or Chain-of-Custody in this study for soil sample selection, date, time and depths of each sample chosen for laboratory analyses.

### **Sample Handling Procedures**

The sample handling procedures were conducted as follows:

- 1) Disposable surgical latex gloves were used to avoid cross contamination of samples. Gloves were discarded in a designated "waste bag after each sample was collected.
- 2) Each confirmation sample upon collection was immediately stored in a cooler containing ice. During the sample collection process, care was taken to insure the samples were not collected in direct sunlight. In addition, during the collection process, no parts of the body without gloves touched any part of the sample.
- 3) Once placed into the cooler, each sample was protected with bubble wrap® and foam was inserted in the base, sides and top of the cooler.

### **Soil Boring Abandonment Procedures**

Due to the fact that holes in the subsurface may act as a conduit for contamination migration, proper sealing of holes is essential for ensuring that a site assessment does not contribute to the spread of contaminants. The objective of hole-sealing is to prevent preferential migration of contaminants through the bore hole. To seal the boreholes advanced during this study, the contractor utilized a method known as surface pouring. Surface pouring entails sealing the boreholes with dry products (e.g., bentonite granules, chips and/or pellets). Once the DPT drive rods have been withdrawn, dry products are physically poured into the bottom of the

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borehole and filled vertically up the column to at least two (2) feet from the base of the borehole. Once the dry products have seated into the borehole, the product is hydrated to expand the clay material. After the hydration process has been performed, the remaining portions of the boreholes are backfilled with the soil cores. Due to the nature of DPT, no soil cuttings were generated during soil boring exploration assessment work.

**GROUNDWATER INVESTIGATION**

The purpose of a monitoring well is to provide an access point for measuring groundwater levels and to collect groundwater samples representing actual in-situ groundwater conditions at that point of access. For the purpose of this investigation, based on the scope of work, EI chose to install temporary groundwater monitoring wells (Type I).

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## WELL DEVELOPMENT AND GROUNDWATER SAMPLE COLLECTION

### Water Development

The groundwater monitor well was purged with a Peristaltic™ pump. Well development allows fresh water from the formation to enter the well and the groundwater samples will more accurately represent actual groundwater conditions. The well was purged of approximately three (3) to five (5) well volumes of water or until dry prior to sampling.

### Groundwater Sampling Procedures

After well development activities were performed, groundwater samples were collected from the well(s) with the referenced pump. During the collection process, samples were poured directly from the bailer into the laboratory supplied containers which were placed into an ice chest filled with ice. Under no circumstances were any intermediate sample containers used, i.e. jar, beaker, etc., and then transferred to the sample container. In addition, water samples were not field filtered.

Prior to collecting the water sample, the containers were labeled accordingly. This procedure was performed prior to sampling because sample containers have a tendency to "sweat" when filled with groundwater; this makes it difficult to affix a label to the container after sampling. The sample label also was covered with a clear piece of tape, which was wrapped around the sample container. This procedure prevented the label from detaching from the container during sample storage and shipment.

Each sample container was labeled indicating the sample location (i.e. GP-1, or MW-1, etc.), date and time of collection, sample location, collector, project site, and analysis identification. Other pertinent information was recorded in the field book.

After the groundwater sample(s) was collected, the containers were immediately placed in a sample cooler containing ice. Upon completion, the samples were transported to Paradigm Analytical Laboratories, located in Wilmington, NC using chain-of-custody documentation.

### Soil Boring Abandonment Procedures

Due to the fact that holes in the subsurface may act as a conduit for contamination migration, proper sealing of holes is essential for ensuring that a site assessment does not contribute to the spread of contaminants. The objective of hole-sealing is to prevent preferential migration of contaminants through the bore hole. To seal the boreholes advanced during this study, the

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contractor utilized a method known as surface pouring. Surface pouring entails sealing the boreholes with dry products (e.g., bentonite granules, chips and/or pellets). Once the DPT drive rods have been withdrawn, dry products are physically poured into the bottom of the borehole and filled vertically up the column to at least two (2) feet from the base of the borehole. Once the dry products have seated into the borehole, the product is hydrated to expand the clay material. After the hydration process has been performed, the remaining portions of the boreholes are backfilled with the soil cores. Due to the nature of DPT, no soil cuttings were generated during soil boring exploration assessment work.

### **LABORATORY ANALYTICAL METHODS**

#### **Soil Analytical Methods**

Based upon verbal information provided by NCDOT personnel (Eugene Tarascio), EI selected to analyze the chosen soil samples for total petroleum hydrocarbons (TPH) analyses by Method 8015B with preparation methods for the analysis of Diesel Range Organics (DRO) by GC-FID and Gasoline Range Organics (GRO) by GC-FID. The GRO method is utilized to extract volatile fuels such as gasoline, while the DRO method is utilized to extract less volatile petroleum products such as diesel fuel, fuel oil #2, kerosene, and varsol.

One (1) soil sample from the site was analyzed for volatile organics by SW-846 Method 8260 (5035 Prep), for semi-volatiles (SVOCs) by SW-846 Method 8270, and for aliphatics and aromatics by Massachusetts Department of Environmental Protection's (MADEP) method for volatile petroleum hydrocarbons (VPH) and MADEP's method for extractable petroleum hydrocarbons (EPH), respectively.

These laboratory analytical methods were utilized as required in the *Guidelines* in order to compare results to the DWM's maximum soil contaminant concentration (MSCC) cleanup standards. The MSCC concentrations are also published in the *Guidelines*.

### **SAMPLE PACKAGING AND SHIPPING**

This section discusses the sample packaging and shipping protocol that shall be used to transport collected samples to the laboratories for analytical testing. Samples collected, prepared, preserved and stored must then be readied for packaging and shipping. It is important that the presented protocol be followed to ensure that the samples reach their destination in sound condition. In addition, the samples must be under strict COC from the time they are sampled until the analysis is

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complete.

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Samples collected for this project were classified as environmental materials samples and were not considered hazardous. In addition, the samples collected for this study were not classified as "dangerous goods".

Environmental samples collected for this field study were packed prior to shipment using the following procedures:

1. Secure drain plug on cooler with tape.
2. Place cushioned layer on bottom of cooler (vermiculite or "bubble-wrap" plastic).
3. Line cooler with large heavy duty plastic bag.
4. Place all sample containers in large plastic bag within the cooler. Be sure the lids on all bottles are tight (will not leak).
5. Cushion containers to prevent breakage.
6. Put ice that has been "double bagged" in heavy duty polyethylene bags and placed on top of and/or between the samples within the large plastic bag. Fill all remaining space between the containers with cushion materials.
7. Securely fasten the top of the large plastic bag with tape or tie.
8. Place the Chain-of-Custody Record into a plastic bag, and tape the bag to the inner side of the cooler lid.
9. Close the cooler and securely tape (preferably with fiber tape) the top of the cooler shut. Custody seals should be affixed to the top and sides of the cooler within the securing tape so that the cooler cannot be opened without breaking the seal.
10. Shipping containers (ice cooler) must be marked "THIS END UP", and arrow labels which indicate the proper upward position of the container should be affixed to the container. A label containing the name and address of the shipper should be placed on the containers exterior. Labels used in the shipment of hazardous materials (e.g., Cargo Only Air Craft, Flammable Solids, etc.) are not permitted to be on the outside of containers used to transport environmental samples.

***Shipping Note:***

"When samples are to be shipped by common carrier or sent through the United States mail, it must comply with the Department of Transportation Hazardous Materials Regulations (49 CFR 172). The person offering such material for transportation is responsible for ensuring such compliance. For the preservation requirements of 40 CFR, Part 136, Table II, the Office of Hazardous Materials, Materials Transportation Bureau, Department of Transportation has determined that the Hazardous Materials Regulations do not apply to the following materials: Hydrochloric Acid (HCL) in water solutions at concentrations of 0.04% by weight or less (pH about 1.96 or greater); Nitric acid (HN03) in water solutions at concentrations of 0.-15% by weight or less (pH about 1. 62 or greater); Sulfuric acid (H<sub>2</sub>SO<sub>4</sub>) in water solutions at concentrations of 0.35% by weight or less (pH about 1. 15 or greater); and Sodium Hydroxide (Na OH) in water solutions at concentrations of 0.08% by weight or less (pH about 12.30 or less). This footnote is wholly reproduced from 40 CFR 136.3, which is

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**Sample Transportation**

The cooler(s) containing the collected soil samples was shipped overnight via Federal Express, with COC documentation, to Prism Laboratories, Inc. in Charlotte, NC. The following protocol was used for sample handling and transportation:

- 1) The lids on all bottles were tightened to reduce the potential for leakage.
- 2) The sample identification label on each individual laboratory container was covered with a clear piece of plastic tape. Each container was then placed within an appropriately sized polyethylene bag and sealed.
- 3) The containers were placed into a bubble-wrap® lined rectangular ice chest (cooler).
- 4) Ice was placed on top and surrounding bubble-wrap® sample containers. Some of the remaining spaces between the containers were filled with bubble-wrap® and/or ice.
- 5) The cooler drain plug was secured with clear tape.
- 6) The COC's was double plastic bagged and was taped to the inner side of the cooler lid.
- 7) The cooler was closed and securely taped.
- 8) A label with adhesive tape containing the name and address of the shipper and the address of the laboratory was placed on top of the cooler.

**DECONTAMINATION PROCEDURES**

Decontamination is the process of washing, rinsing and removing contaminants from exposed surfaces of equipment. Decontamination helps prevent the spread of contamination off-site, and avoids cross-contamination to other samples. The decontamination procedures were performed as follows:

- 1) Disposable surgical latex gloves were used in lieu of decontamination procedures to collect soil samples.

The soil samples retained for laboratory analyses were placed in the appropriate clean laboratory prepared containers, labeled and subsequently delivered with chain-of-custody documentation (COC) for analysis. Dates and times of sampling may be referenced on the COC's. Specific laboratory analysis methods are referenced in the text of this Study.

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**QUALITY ASSURANCE PROTOCOL**

**Field and Laboratory Control Samples**

The purpose of this section is to describe the standard control sampling program that supported the data quality objectives for this site. These control samples will included field control Quality Assurance (QA) samples used to assess sources of error. To minimize or consider the impact these errors have on the resulting data, a combination of unique field QA/QC protocols and control samples were developed to meet the QA overall objectives.

**Field Control Samples**

The elements of the sampling and field QA/QC strategy included the following:

- (1) El developed a well thought out sampling strategy for the site. The plan adequately and sufficiently outlined the different types of environmental media and protocol to sample the media.
- (2) Sampling methodologies to obtain true representative samples.
- (3) Used decontamination procedures in order to reduce cross-contamination potential between sampling points.
- (4) Used the proper sample containers, and preservation requirements.
- (5) Used the proper storage, and shipping of samples protocol.

Techniques to verify the inclusion of the QA/QC program included scheduled field control samples consisting of field blanks (trip and temperature). The field control samples were handled similarly as the environmental samples.

**Quality Control Samples**

A trip and temperature blank were collected during this study.

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**Laboratory QA/QC Procedures**

Laboratory QA/QC procedures are implemented in order to prevent, detect, and correct potential errors during the analytical process. The reliability and credibility of analytical laboratories are corroborated by the development and performance of their respective QA/QC programs. For this project, the NCDOT contracted laboratory provided and performed their program as they see fit. Standard practices used by the selected laboratory included the following quality control sample information in their generated reports:

- (a) laboratory method blanks;
- (b) temperature blanks.

**INVESTIGATION DERIVED WASTE MANAGEMENT PROTOCOL**

The investigation derived waste (IDW) generated during the sampling activities were placed on site. These wastes include any derivative investigative soils leftover from the sampling and backfilling protocol, decontamination water (cleaning of field equipment), bailers, bailer haul-line and PPE equipment, if applicable. The management of IDW for this project complies with applicable or relevant and appropriate requirements (ARAs). The site specific ARAs were followed in consensus with the EPA Standard Operating Procedures (SOP) and Quality Assurance Manual, Region 4 and the *Guidelines For Assessment And Corrective Action*, drafted by the North Carolina Underground Storage Tank Section, effective July 1, 2001.

**APPENDIX C**  
**SOIL BORING LOGS**



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 Morrisville, North Carolina  
 919-544-7500

**SOIL BORING LOG**

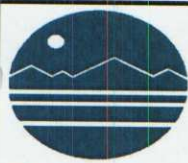
Boring No. GP-1  
 Date Drilled: 02/23/06

Client: NCDOT  
 Project Name: Parcel #127  
 Project/Site Location: 120 Mill Springs Road, Burnsville, NC  
 Project Number: ENMO060029.00

Logged By: RMS  
 Drilling Company: Troxler Geologic Services  
 Drill Device: GeoProbe 6600  
 Drill Method: DPT

Total Boring Depth: 3.66 m Weather Conditions: Cool Surface Elevation: \_\_\_\_\_  
 Boring Diameter: 10.16cm Boring Location: Downgradient of UST

Depth (Feet)	Depth (meters)	Time	Sample Analyzed	Recovery	Soil Profile	Lithological Description	Sample PID (ppm)
						Tan to light brown clayey SILT (ML), dry.	
2.00	0.61			100%			NA
4.00	1.22						NA
6.00	1.83			100%	(ML)		NA
8.00	2.44					Tan to light brown fine sandy SILT (ML), with little clay, dry to moist.	0.0
10.00	3.05	14:00	x				0.0
12.00	3.66			100%		Auger Refusal Boring terminated at 3.66 meters (12.0') bls. x denotes soil sample at 2.44 - 3.05 meters (8'-10') bls interval collected for laboratory retention.	0.0



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**SOIL BORING LOG**

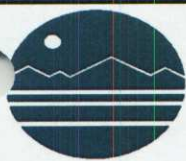
Boring No. GP-2  
 Date Drilled: 02/23/06

Client: NCDOT  
 Project Name: Parcel #127  
 Project/Site Location: 120 Mill Springs Road, Burnsville, NC  
 Project Number: ENMO060029.00

Logged By: RMS  
 Drilling Company: Troxler Geologic Services  
 Drill Device: GeoProbe 6600  
 Drill Method: DPT

Total Boring Depth: 3.66 m Weather Conditions: Cool Surface Elevation: \_\_\_\_\_  
 Boring Diameter: 10.16cm Boring Location: Downgradient of UST

Depth (Feet)	Depth (meters)	Time	Sample Analyzed	Recovery	Soil Profile	Lithological Description	Sample PID (ppm)
						Tan to light brown clayey SILT (ML), dry.	
2.00	0.61			100%			NA
4.00	1.22						NA
6.00	1.83			100%	(ML)		NA
8.00	2.44	14:00	x			Tan to light brown fine sandy SILT (ML), with little clay, dry to moist.	0.0
10.00	3.05			100%			0.0
12.00	3.66					Boring terminated at 3.66 meters (12.0') bls. x denotes soil sample at 2.44 - 3.05 meters (8'-10') bls interval collected for laboratory retention. Boring converted into a temporary monitoring well.	0.0



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## SOIL BORING LOG

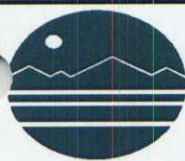
Boring No. HA-1  
 Date Drilled: 03/29/06

Client: NCDOT  
 Project Name: Parcel #127  
 Project/Site Location: 120 Mill Springs Road, Burnsville, NC  
 Project Number: ENMO060029.00

Logged By: RMS  
 Drilling Company: N/A  
 Drill Device: Hand Auger  
 Drill Method: Hand Auger

Total Boring Depth: 2.44 m Weather Conditions: Cold Surface Elevation: \_\_\_\_\_  
 Boring Diameter: 10.16cm Boring Location: Adjacent to UST

Depth (Feet)	Depth (meters)	Time	Sample Analyzed	Recovery	Soil Profile	Lithological Description	Sample PID (ppm)
2.00	0.61			100%		Reddish brown to light tan clayey SILT (ML), micaeous, dry to slightly moist	0.0
4.00	1.22						0.1
6.00	1.83			100%		Reddish brown to light tan silty CLAY (CL), very micaeous, low plasticity, dry to slightly moist	0.0
8.00	2.44	???	x			Boring terminated at 2.44m (8.0') bls. x denotes soil sample at 2.13m - 2.44m (7'-8') bls interval collected for laboratory retention.	0.3



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**SOIL BORING LOG**

Boring No. HA-2  
Date Drilled: 03/29/06

Client:	NCDOT
Project Name:	Parcel #127
Project/Site Location:	120 Mill Springs Road, Burnsville, NC
Project Number:	ENMO060029.00

Logged By:	RMS
Drilling Company:	N/A
Drill Device:	Hand Auger
Drill Method:	Hand Auger

Total Boring Depth:	2.44 m	Weather Conditions:	Cold	Surface Elevation:	
Boring Diameter:	10.16cm	Boring Location:	Proposed Drainage Piping		

Depth (Feet)	Depth (meters)	Time	Sample Analyzed	Recovery	Soil Profile	Lithological Description	Sample PID (ppm)
						Reddish brown to light tan clayey SILT (ML), micaeous, dry to slightly moist	0.0
2.00	0.61			100%	ML		0.1
4.00	1.22						
6.00	1.83			100%	CL	Reddish brown to light tan silty CLAY (CL), very micaeous, low plasticity, dry to slightly moist	0.0
8.00	2.44	???	x				0.3
						Boring terminated at 2.44m (8.0') bls. x denotes soil sample at 2.13m - 2.44m (7'-8') bls interval collected for laboratory retention.	



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## SOIL BORING LOG

Boring No. HA-3  
Date Drilled: 03/29/06

Client: NCDOT  
Project Name: Parcel #127  
Project/Site Location: 120 Mill Springs Road, Burnsville, NC  
Project Number: ENMO060029.00

Logged By: RMS  
Drilling Company: N/A  
Drill Device: Hand Auger  
Drill Method: Hand Auger

Total Boring Depth: 10.16cm Weather Conditions: Cold Surface Elevation:  
Boring Diameter: 4.0" Boring Location: Proposed Drainage Piping

Depth (Feet)	Depth (meters)	Time	Sample Analyzed	Recovery	Soil Profile	Lithological Description	Sample PID (ppm)
2.00	0.61			100%	ML	Reddish brown to light tan clayey SILT (ML), micaeous, dry to slightly moist	0.0
4.00	1.22						0.1
6.00	1.83			100%	CL	Reddish brown to light tan silty CLAY (CL), very micaeous, low plasticity, dry to slightly moist	0.0
8.00	2.44	???	x			Boring terminated at 2.44m (8.0') bls. x denotes soil sample at 2.13m - 2.44m (7'-8') bls interval collected for laboratory retention.	0.3

**APPENDIX D**  
**LABORATORY ANALYTICAL REPORT**

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Mr. Bob Shaut  
Environmental Investigations  
2101 Gateway Centre Boulevard  
Suite 200  
Morrisville NC 27560  
Report Number: G106-565

Client Project: NCDOT-Yancey

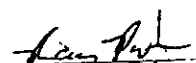
Dear Mr. Shaut:

Enclosed are the results of the analytical services performed under the referenced project. The samples are certified to meet the requirements of the National Environmental Laboratory Accreditation Conference Standards. Copies of this report and supporting data will be retained in our files for a period of five years in the event they are required for future reference. Any samples submitted to our laboratory will be retained for a maximum of thirty (30) days from the date of this report unless other arrangements are requested.

If there are any questions about the report or the services performed during this project, please call Paradigm at (910) 350-1903. We will be happy to answer any questions or concerns which you may have.

Thank you for using Paradigm Analytical Labs for your analytical services. We look forward to working with you again on any additional analytical needs which you may have.

Sincerely,  
Paradigm Analytical Laboratories, Inc.

  
Laboratory Director  
J. Patrick Weaver

3/10/2006  
Date

## CASE NARATIVE

Date: March 10, 2006

Environmental Investigations Project ID: NCDOT Yancey  
SGS-Paradigm Analytical ID: G106-565

Twenty-four samples were received at the laboratory February 24 for analysis as indicated on the chain of custody. The samples were received in good condition, within temperature and holding time limits.

All extractions and analyses were completed within holding time and without quality control exception.

Many of the 8260 sample results show low concentrations of methylene chloride below the reporting limit but above the method detection limit. This analyte is a common laboratory solvent and its detection is likely a laboratory artifact.

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 206 GP1  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-1  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 16:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 78.08

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.75	5035	1	02/25/06
Diesel Range Organics	13.8	7.51	3541	1	02/27/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 206 HA1  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-2  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 17:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 75.83

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	9.16	5035	1	02/25/06
Diesel Range Organics	BQL	7.54	3541	1	02/27/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 206 HA1A  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-3  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/23/06 9:50  
Date Received: 2/24/06  
Matrix: Soil  
Solids 76.88

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.32	5035	1	02/25/06
Diesel Range Organics	BQL	7.67	3541	1	02/27/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 206 HA2

Analyzed By: MJC  
Date Collected: 2/23/06 10:20

Client Project ID: NCDOT-Yancey

Date Received: 2/24/06

Lab Sample ID: G106-565-4

Matrix: Soil

Lab Project ID: G106-565

Solids 77.25

Report Basis: Dry Weight

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.23	5035	1	02/27/06
Diesel Range Organics	BQL	7.58	3541	1	02/27/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 206 HA3  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-5  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/23/06 11:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 70.87

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.14	5035	1	02/27/06
Diesel Range Organics	BQL	8.10	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 206 HA4  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-6  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/23/06 11:05  
Date Received: 2/24/06  
Matrix: Soil  
Solids 73.43

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.98	5035	1	02/27/06
Diesel Range Organics	109	8.47	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 127 GP1-8  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-7  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/23/06 14:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 78.88

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.89	5035	1	02/27/06
Diesel Range Organics	BQL	7.86	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 163 GP1-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-8  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 9:15  
Date Received: 2/24/06  
Matrix: Soil  
Solids 71.71

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.57	5035	1	02/27/06
Diesel Range Organics	BQL	8.14	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 163 GP2-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-9  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 9:22  
Date Received: 2/24/06  
Matrix: Soil  
Solids 77.19

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.22	5035	1	02/27/06
Diesel Range Organics	8.04	7.16	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 163 GP3-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-10  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 10:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 74.16

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.04	5035	1	02/27/06
Diesel Range Organics	BQL	8.12	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 163 GP4-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-11  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 10:45  
Date Received: 2/24/06  
Matrix: Soil  
Solids 77.80

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.54	5035	1	02/27/06
Diesel Range Organics	20.6	7.26	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 163 GP5-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-12  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 11:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 69.99

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.75	5035	1	02/27/06
Diesel Range Organics	BQL	8.31	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 199A GP1-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-13  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 12:30  
Date Received: 2/24/06  
Matrix: Soil  
Solids 86.37

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.31	5035	1	02/27/06
Diesel Range Organics	BQL	6.75	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 199A GP2-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-14  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 12:45  
Date Received: 2/24/06  
Matrix: Soil  
Solids 90.91

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.57	5035	1	02/27/06
Diesel Range Organics	BQL	6.65	3541	1	02/28/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 199A GP3-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-15  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/22/06 13:00  
Date Received: 2/24/06  
Matrix: Soil  
Solids 91.13

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.53	5035	1	02/27/06
Diesel Range Organics	BQL	6.74	3541	1	03/02/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 89 HA4-8

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/20/2006 11:20

Lab Sample ID: G106-565-16

Date Received: 2/24/2006

Lab Project ID: G106-565

Matrix: Soil

Report Basis: Dry Weight

Solids 68.16

Analyte	Result MG/KG	Report Limit MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.08	5035	1	02/24/06
Diesel Range Organics	BQL	9.13	3545	1	03/02/06

Comments:

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 89 HA6-8  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-17  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/20/06 12:45  
Date Received: 2/24/06  
Matrix: Soil  
Solids 77.20

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.69	5035	1	02/24/06
Diesel Range Organics	BQL	8.05	3541	1	03/02/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 89 HA9-9  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-18  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/21/06 12:06  
Date Received: 2/24/06  
Matrix: Soil  
Solids 69.34

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.45	5035	1	02/24/06
Diesel Range Organics	51.8	8.98	3541	1	03/02/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 89 HA10-9  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-19  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/21/2006 12:44  
Date Received: 2/24/2006  
Matrix: Soil  
Solids 61.86

Analyte	Result MG/KG	Report Limit MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.09	5035	1	02/24/06
Diesel Range Organics	BQL	9.89	3545	1	03/02/06

Comments:

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 221 GP1-8  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-20  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/21/06 15:15  
Date Received: 2/24/06  
Matrix: Soil  
Solids 75.74

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	19.0	6.65	5035	1	02/24/06
Diesel Range Organics	112	8.11	3541	1	03/02/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 221 GP2-8  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-21  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/21/06 15:28  
Date Received: 2/24/06  
Matrix: Soil  
Solids 69.37

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.50	5035	1	02/25/06
Diesel Range Organics	BQL	8.96	3541	1	03/02/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 221 GP3-8  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-22  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/21/06 15:45  
Date Received: 2/24/06  
Matrix: Soil  
Solids 72.24

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.82	5035	1	02/25/06
Diesel Range Organics	10.8	8.64	3541	1	03/02/06

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: PAR 221 GP4-10  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: G106-565-23  
Lab Project ID: G106-565  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 2/21/06 16:36  
Date Received: 2/24/06  
Matrix: Soil  
Solids 76.29

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.12	5035	1	02/25/06
Diesel Range Organics	BQL	8.10	3541	1	03/02/06

**QC Results for Total Petroleum Hydrocarbons**  
by GC/FID

Client Sample ID: Batch QC  
 Lab Sample ID: g106-565-20a  
 LCS ID: LCS4022406A / VP022406

Analyzed By: MJC  
 Matrix: Soil  
 Solids 75.74

**MS/MSD**

Analyte	Sample MG/KG	Spiked MG/KG	MS MG/KG	REC %	Spiked MG/KG	MSD MG/KG	REC % (70-130)	RPD %
GRO	17.8	44.3	62.1	100	44.3	63.2	102	1.98

**LCS**

Analyte	Spiked MG/KG	Result MG/KG	REC %	LIMITS		Lower	Upper
				Lower	Upper		
GRO	40	39	96.3	70	130		

**Comments:**

Reviewed By: \_\_\_\_\_

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: Method Blank  
Client Project ID:  
Lab Sample ID: VBLK4022406A  
Lab Project ID:  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected:  
Date Received:  
Matrix: Soil  
Solids 100.00

Analyte	Result mg/kg	RL mg/kg	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.00	5035	1	02/24/06
Surrogate Spike Results		Spike Added 50		Spike Result 49.5	Percent Recovery 99
BFB					

Comments:

Flags:

**QC Results for Total Petroleum Hydrocarbons**  
by GC/FID

Client Sample ID: Batch QC  
 Lab Sample ID: g106-565-4a  
 LCS ID: LCS4022706A / VP022706

Analyzed By: MJC  
 Matrix: Soil  
 Solids 77.25

## MS/MSD

Analyte	Sample MG/KG	Spiked MG/KG	MS MG/KG	REC %	Spiked MG/KG	MSD MG/KG	REC % (70-130)	RPD %
GRO	BQL	48.2	44.5	92.3	48.2	45.8	95	2.88

## LCS

Analyte	Spiked MG/KG	Result MG/KG	REC %	LIMITS		
				Lower	Upper	
GRO	40	38	95.3	70	130	

Comments:

Reviewed By: BY

**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: Method Blank

Analyzed By: MJC

Client Project ID:

Date Collected:

Lab Sample ID: VBLK4022706A

Date Received:

Lab Project ID:

Matrix: Soil

Report Basis: Dry Weight

Solids 100.00

Analyte	Result mg/kg	RL mg/kg	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	6.00	5035	1	02/27/06
<b>Surrogate Spike Results</b>		<b>Spike Added</b>		<b>Spike Result</b>	<b>Percent Recovery</b>
BFB		50		46.6	93.2

Comments:

Flags:

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 206 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-1A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 16:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 78.1

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	25.9	70.8	4.17	1	2/27/2006	
Benzene	BQL	7.08	4.04	1	2/27/2006	
Bromobenzene	BQL	7.08	3.49	1	2/27/2006	
Bromoform	BQL	7.08	4.14	1	2/27/2006	
Bromochloromethane	BQL	7.08	4.02	1	2/27/2006	
Bromodichloromethane	BQL	7.08	3.43	1	2/27/2006	
Bromomethane	BQL	7.08	5.94	1	2/27/2006	
2-Butanone	11.5	35.4	4.08	1	2/27/2006	J
n-Butylbenzene	BQL	7.08	4.75	1	2/27/2006	
sec-Butylbenzene	BQL	7.08	4.96	1	2/27/2006	
tert-Butylbenzene	BQL	7.08	4.92	1	2/27/2006	
Carbon disulfide	BQL	7.08	3.73	1	2/27/2006	
Carbon tetrachloride	BQL	7.08	4.90	1	2/27/2006	
Chlorobenzene	BQL	7.08	3.56	1	2/27/2006	
Chloroethane	BQL	7.08	4.45	1	2/27/2006	
Chloroform	BQL	7.08	3.56	1	2/27/2006	
Chloromethane	BQL	7.08	3.41	1	2/27/2006	
2-Chlorotoluene	BQL	7.08	4.22	1	2/27/2006	
4-Chlorotoluene	BQL	7.08	3.94	1	2/27/2006	
Dibromochloromethane	BQL	7.08	3.17	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	7.08	15.0	1	2/27/2006	
Dibromomethane	BQL	7.08	4.25	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	7.08	3.30	1	2/27/2006	
1,2-Dichlorobenzene	BQL	7.08	3.41	1	2/27/2006	
1,3-Dichlorobenzene	BQL	7.08	3.32	1	2/27/2006	
1,4-Dichlorobenzene	BQL	7.08	3.49	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	7.08	15.2	1	2/27/2006	
1,1-Dichloroethane	BQL	7.08	4.08	1	2/27/2006	
1,1-Dichloroethene	BQL	7.08	5.45	1	2/27/2006	
1,2-Dichloroethane	BQL	7.08	4.07	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	7.08	3.49	1	2/27/2006	
trans-1,2-dichloroethene	BQL	7.08	4.60	1	2/27/2006	
1,2-Dichloropropane	BQL	7.08	3.63	1	2/27/2006	
1,3-Dichloropropane	BQL	7.08	3.24	1	2/27/2006	
2,2-Dichloropropane	BQL	7.08	4.51	1	2/27/2006	
1,1-Dichloropropene	BQL	7.08	5.11	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	7.08	3.94	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	7.08	4.01	1	2/27/2006	
Dichlorodifluoromethane	BQL	7.08	5.28	1	2/27/2006	
Diisopropyl ether (Dipe)	BQL	7.08	3.36	1	2/27/2006	
Ethylbenzene	BQL	7.08	4.31	1	2/27/2006	

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 206 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-1A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 16:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 78.1

<b>Report Name Compound</b>	<b>Result UG/KG</b>	<b>Quantitation Limit UG/KG</b>	<b>MDL UG/KG</b>	<b>Dilution Factor</b>	<b>Date Analyzed</b>	<b>Flag</b>
Hexachlorobutadiene	BQL	7.08	5.60	1	2/27/2006	
2-Hexanone	BQL	7.08	3.09	1	2/27/2006	
Iodomethane	BQL	7.08	6.59	1	2/27/2006	
Isopropylbenzene	BQL	7.08	4.56	1	2/27/2006	
4-Isopropyltoluene	BQL	7.08	4.82	1	2/27/2006	
Methylene chloride	16.3	28.3	4.05	1	2/27/2006	J
4-Methyl-2-pentanone	BQL	7.08	3.27	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	7.08	3.60	1	2/27/2006	
Naphthalene	BQL	7.08	2.85	1	2/27/2006	
n-Propyl benzene	BQL	7.08	4.55	1	2/27/2006	
Styrene	BQL	7.08	5.07	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	7.08	3.85	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	7.08	3.49	1	2/27/2006	
Tetrachloroethene	BQL	7.08	4.46	1	2/27/2006	
Toluene	BQL	7.08	4.19	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	7.08	3.10	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	7.08	3.17	1	2/27/2006	
Trichloroethene	BQL	7.08	4.42	1	2/27/2006	
1,1,1-Trichloroethane	BQL	7.08	4.89	1	2/27/2006	
1,1,2-Trichloroethane	BQL	7.08	3.66	1	2/27/2006	
Trichlorofluoromethane	BQL	7.08	5.85	1	2/27/2006	
1,2,3-Trichloroproppane	BQL	7.08	3.75	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	7.08	3.97	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	7.08	4.22	1	2/27/2006	
Vinyl chloride	BQL	7.08	4.68	1	2/27/2006	
m-,p-Xylene	BQL	14.2	8.03	1	2/27/2006	
o-Xylene	BQL	7.08	3.97	1	2/27/2006	
4-Bromofluorobenzene	50	49.1		98		
1,2-Dichloroethane-d4	50	59.9		120		
Toluene-d8	50	51.4		103		

Comments:

Flags:

Reviewed By: ZYL

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 206 HA1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-2A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 17:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 75.8

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	61.2	72.0	4.23	1	2/27/2006	J
Benzene	BQL	7.20	4.10	1	2/27/2006	
Bromobenzene	BQL	7.20	3.54	1	2/27/2006	
Bromoform	BQL	7.20	4.20	1	2/27/2006	
Bromochloromethane	BQL	7.20	4.09	1	2/27/2006	
Bromodichloromethane	BQL	7.20	3.48	1	2/27/2006	
Bromomethane	BQL	7.20	6.03	1	2/27/2006	
2-Butanone	13.8	36.0	4.15	1	2/27/2006	J
n-Butylbenzene	BQL	7.20	4.82	1	2/27/2006	
sec-Butylbenzene	BQL	7.20	5.04	1	2/27/2006	
tert-Butylbenzene	BQL	7.20	5.00	1	2/27/2006	
Carbon disulfide	BQL	7.20	3.79	1	2/27/2006	
Carbon tetrachloride	BQL	7.20	4.98	1	2/27/2006	
Chlorobenzene	BQL	7.20	3.61	1	2/27/2006	
Chloroethane	BQL	7.20	4.52	1	2/27/2006	
Chloroform	BQL	7.20	3.61	1	2/27/2006	
Chloromethane	BQL	7.20	3.47	1	2/27/2006	
2-Chlorotoluene	BQL	7.20	4.29	1	2/27/2006	
4-Chlorotoluene	BQL	7.20	4.00	1	2/27/2006	
Dibromochloromethane	BQL	7.20	3.22	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	7.20	15.3	1	2/27/2006	
Dibromomethane	BQL	7.20	4.32	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	7.20	3.35	1	2/27/2006	
1,2-Dichlorobenzene	BQL	7.20	3.47	1	2/27/2006	
1,3-Dichlorobenzene	BQL	7.20	3.37	1	2/27/2006	
1,4-Dichlorobenzene	BQL	7.20	3.54	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	7.20	15.4	1	2/27/2006	
1,1-Dichloroethane	BQL	7.20	4.15	1	2/27/2006	
1,1-Dichloroethene	BQL	7.20	5.54	1	2/27/2006	
1,2-Dichloroethane	BQL	7.20	4.13	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	7.20	3.54	1	2/27/2006	
trans-1,2-dichloroethene	BQL	7.20	4.68	1	2/27/2006	
1,2-Dichloropropane	BQL	7.20	3.69	1	2/27/2006	
1,3-Dichloropropane	BQL	7.20	3.30	1	2/27/2006	
2,2-Dichloropropane	BQL	7.20	4.58	1	2/27/2006	
1,1-Dichloropropene	BQL	7.20	5.20	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	7.20	4.00	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	7.20	4.07	1	2/27/2006	
Dichlorodifluoromethane	BQL	7.20	5.37	1	2/27/2006	
Diisopropyl ether (DIPE)	BQL	7.20	3.41	1	2/27/2006	
Ethylbenzene	BQL	7.20	4.38	1	2/27/2006	

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 206 HA1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-2A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 17:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 75.8

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	7.20	5.69	1	2/27/2006	
2-Hexanone	BQL	7.20	3.14	1	2/27/2006	
Iodomethane	BQL	7.20	6.69	1	2/27/2006	
Isopropylbenzene	BQL	7.20	4.64	1	2/27/2006	
4-Isopropyltoluene	BQL	7.20	4.89	1	2/27/2006	
Methylene chloride	9.85	28.8	4.12	1	2/27/2006	J
4-Methyl-2-pentanone	BQL	7.20	3.33	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	7.20	3.66	1	2/27/2006	
Naphthalene	BQL	7.20	2.89	1	2/27/2006	
n-Propyl benzene	BQL	7.20	4.62	1	2/27/2006	
Styrene	BQL	7.20	5.15	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	7.20	3.92	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	7.20	3.54	1	2/27/2006	
Tetrachloroethene	BQL	7.20	4.53	1	2/27/2006	
Toluene	BQL	7.20	4.26	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	7.20	3.15	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	7.20	3.22	1	2/27/2006	
Trichloroethene	BQL	7.20	4.49	1	2/27/2006	
1,1,1-Trichloroethane	BQL	7.20	4.97	1	2/27/2006	
1,1,2-Trichloroethane	BQL	7.20	3.71	1	2/27/2006	
Trichlorofluoromethane	BQL	7.20	5.95	1	2/27/2006	
1,2,3-Trichloropropane	BQL	7.20	3.82	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	7.20	4.03	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	7.20	4.29	1	2/27/2006	
Vinyl chloride	BQL	7.20	4.75	1	2/27/2006	
m-,p-Xylene	BQL	14.4	8.16	1	2/27/2006	
o-Xylene	BQL	7.20	4.03	1	2/27/2006	
					Spike Recovered	
					Added	
4-Bromofluorobenzene		50	50		100	
1,2-Dichloroethane-d4		50	61.4		123	
Toluene-d8		50	52.3		105	

Comments:

Flags:

Reviewed By: Pat

**Results for Volatiles**  
by GCMS 8260-5035

Client Sample ID: PAR 206 HA1A  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-3A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 09:50  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 76.9

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	34.3	75.4	4.44	1	2/27/2006	
Benzene	BQL	7.54	4.30	1	2/27/2006	
Bromobenzene	BQL	7.54	3.71	1	2/27/2006	
Bromoform	BQL	7.54	4.41	1	2/27/2006	
Bromochloromethane	BQL	7.54	4.29	1	2/27/2006	
Bromodichloromethane	BQL	7.54	3.65	1	2/27/2006	
Bromomethane	BQL	7.54	6.32	1	2/27/2006	
2-Butanone	BQL	37.7	4.35	1	2/27/2006	
n-Butylbenzene	BQL	7.54	5.06	1	2/27/2006	
sec-Butylbenzene	BQL	7.54	5.28	1	2/27/2006	
tert-Butylbenzene	BQL	7.54	5.24	1	2/27/2006	
Carbon disulfide	BQL	7.54	3.97	1	2/27/2006	
Carbon tetrachloride	BQL	7.54	5.22	1	2/27/2006	
Chlorobenzene	BQL	7.54	3.79	1	2/27/2006	
Chloroethane	BQL	7.54	3.79	1	2/27/2006	
Chloroform	BQL	7.54	3.64	1	2/27/2006	
Chloromethane	BQL	7.54	4.50	1	2/27/2006	
2-Chlorotoluene	BQL	7.54	4.19	1	2/27/2006	
4-Chlorotoluene	BQL	7.54	3.38	1	2/27/2006	
Dibromochloromethane	BQL	7.54	16.0	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	7.54	4.53	1	2/27/2006	
Dibromomethane	BQL	7.54	3.52	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	7.54	3.64	1	2/27/2006	
1,2-Dichlorobenzene	BQL	7.54	3.53	1	2/27/2006	
1,3-Dichlorobenzene	BQL	7.54	3.71	1	2/27/2006	
1,4-Dichlorobenzene	BQL	7.54	16.1	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	7.54	4.35	1	2/27/2006	
1,1-Dichloroethane	BQL	7.54	5.81	1	2/27/2006	
1,1-Dichloroethene	BQL	7.54	4.33	1	2/27/2006	
1,2-Dichloroethane	BQL	7.54	3.71	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	7.54	4.90	1	2/27/2006	
trans-1,2-dichloroethene	BQL	7.54	3.86	1	2/27/2006	
1,2-Dichloropropane	BQL	7.54	3.46	1	2/27/2006	
1,3-Dichloropropane	BQL	7.54	4.80	1	2/27/2006	
2,2-Dichloropropane	BQL	7.54	5.45	1	2/27/2006	
1,1-Dichloropropene	BQL	7.54	4.19	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	7.54	4.27	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	7.54	5.63	1	2/27/2006	
Dichlorodifluoromethane	BQL	7.54	3.58	1	2/27/2006	
Diisopropyl ether (Dipe)	BQL	7.54	4.59	1	2/27/2006	
Ethylbenzene	BQL	7.54				

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 206 HA1A  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-3A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 09:50  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 76.9

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	7.54	5.96	1	2/27/2006	
2-Hexanone	BQL	7.54	3.29	1	2/27/2006	
Iodomethane	BQL	7.54	7.02	1	2/27/2006	
Isopropylbenzene	BQL	7.54	4.86	1	2/27/2006	
4-Isopropyltoluene	BQL	7.54	5.13	1	2/27/2006	
Methylene chloride	7.82	30.2	4.32	1	2/27/2006	J
4-Methyl-2-pentanone	BQL	7.54	3.49	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	7.54	3.83	1	2/27/2006	
Naphthalene	BQL	7.54	3.03	1	2/27/2006	
n-Propyl benzene	BQL	7.54	4.84	1	2/27/2006	
Styrene	BQL	7.54	5.40	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	7.54	4.10	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	7.54	3.71	1	2/27/2006	
Tetrachloroethene	BQL	7.54	4.75	1	2/27/2006	
Toluene	BQL	7.54	4.47	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	7.54	3.30	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	7.54	3.38	1	2/27/2006	
Trichloroethene	BQL	7.54	4.71	1	2/27/2006	
1,1,1-Trichloroethane	BQL	7.54	5.21	1	2/27/2006	
1,1,2-Trichloroethane	BQL	7.54	3.89	1	2/27/2006	
Trichlorofluoromethane	BQL	7.54	6.23	1	2/27/2006	
1,2,3-Trichloropropane	BQL	7.54	4.00	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	7.54	4.23	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	7.54	4.50	1	2/27/2006	
Vinyl chloride	BQL	7.54	4.98	1	2/27/2006	
m-,p-Xylene	BQL	15.1	8.56	1	2/27/2006	
o-Xylene	BQL	7.54	4.23	1	2/27/2006	
		Spike Added	Spike Result	Percent Recovered		
		50	48.3	97		
4-Bromofluorobenzene		50	60	120		
1,2-Dichloroethane-d4		50	50.1	100		
Toluene-d8						

Comments:

Flags:

Reviewed By: JTF

**Results for Volatiles**  
by GCMS 8260-5030

Client Sample ID: PAR 206 HA2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-4A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 10:20  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 77.3

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	BQL	60.3	3.35	1	2/27/2006	
Benzene	BQL	6.03	3.64	1	2/27/2006	
Bromobenzene	BQL	6.03	3.91	1	2/27/2006	
Bromoform	BQL	6.03	2.66	1	2/27/2006	
Bromochloromethane	BQL	6.03	3.18	1	2/27/2006	
Bromodichloromethane	BQL	6.03	2.72	1	2/27/2006	
Bromomethane	BQL	6.03	5.79	1	2/27/2006	
2-Butanone	BQL	30.1	3.87	1	2/27/2006	
n-Butylbenzene	BQL	6.03	3.30	1	2/27/2006	
sec-Butylbenzene	BQL	6.03	4.66	1	2/27/2006	
tert-Butylbenzene	BQL	6.03	4.44	1	2/27/2006	
Carbon disulfide	BQL	6.03	3.15	1	2/27/2006	
Carbon tetrachloride	BQL	6.03	4.23	1	2/27/2006	
Chlorobenzene	BQL	6.03	4.23	1	2/27/2006	
Chloroethane	BQL	6.03	5.34	1	2/27/2006	
Chloroform	BQL	6.03	3.51	1	2/27/2006	
Chloromethane	BQL	6.03	3.75	1	2/27/2006	
2-Chlorotoluene	BQL	6.03	4.17	1	2/27/2006	
4-Chlorotoluene	BQL	6.03	4.36	1	2/27/2006	
Dibromochloromethane	BQL	6.03	3.04	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	6.03	8.74	1	2/27/2006	
Dibromomethane	BQL	6.03	2.15	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	6.03	2.24	1	2/27/2006	
1,2-Dichlorobenzene	BQL	6.03	3.64	1	2/27/2006	
1,3-Dichlorobenzene	BQL	6.03	3.95	1	2/27/2006	
1,4-Dichlorobenzene	BQL	6.03	3.84	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	6.03	7.94	1	2/27/2006	
1,1-Dichloroethane	BQL	6.03	4.18	1	2/27/2006	
1,1-Dichloroethene	BQL	6.03	4.30	1	2/27/2006	
1,2-Dichloroethane	BQL	6.03	2.88	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	6.03	3.82	1	2/27/2006	
trans-1,2-dichloroethene	BQL	6.03	4.30	1	2/27/2006	
1,2-Dichloropropane	BQL	6.03	3.34	1	2/27/2006	
1,3-Dichloropropane	BQL	6.03	2.47	1	2/27/2006	
2,2-Dichloropropane	BQL	6.03	3.48	1	2/27/2006	
1,1-Dichloropropene	BQL	6.03	4.22	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	6.03	2.82	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	6.03	2.39	1	2/27/2006	
Dichlorodifluoromethane	BQL	6.03	4.34	1	2/27/2006	
Diisopropyl ether (DIPE)	BQL	6.03	3.53	1	2/27/2006	
Ethylbenzene	BQL	6.03	4.29	1	2/27/2006	

**Results for Volatiles**  
by GCMS 8260-5030

Client Sample ID: PAR 206 HA2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-4A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 10:20  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 77.3

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	6.03	4.31	1	2/27/2006	
2-Hexanone	BQL	6.03	2.15	1	2/27/2006	
Iodomethane	BQL	6.03	4.34	1	2/27/2006	
Isopropylbenzene	BQL	6.03	4.57	1	2/27/2006	
4-Isopropyltoluene	BQL	6.03	4.44	1	2/27/2006	
Methylene chloride	BQL	24.1	3.71	1	2/27/2006	
4-Methyl-2-pentanone	BQL	6.03	1.43	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	6.03	2.74	1	2/27/2006	
Naphthalene	BQL	6.03	2.87	1	2/27/2006	
n-Propyl benzene	BQL	6.03	4.48	1	2/27/2006	
Styrene	BQL	6.03	5.69	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	6.03	3.51	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	6.03	2.18	1	2/27/2006	
Tetrachloroethene	BQL	6.03	4.60	1	2/27/2006	
Toluene	BQL	6.03	3.89	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	6.03	3.17	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	6.03	2.81	1	2/27/2006	
Trichloroethene	BQL	6.03	4.19	1	2/27/2006	
1,1,1-Trichloroethane	BQL	6.03	4.22	1	2/27/2006	
1,1,2-Trichloroethane	BQL	6.03	2.42	1	2/27/2006	
Trichlorofluoromethane	BQL	6.03	4.74	1	2/27/2006	
1,2,3-Trichloropropane	BQL	6.03	2.24	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	6.03	4.36	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	6.03	4.46	1	2/27/2006	
Vinyl chloride	BQL	12.1	4.15	1	2/27/2006	
m-,p-Xylene	BQL	6.03	8.86	1	2/27/2006	
o-Xylene	BQL	6.03	4.28	1	2/27/2006	
		Spike Added	Spike Result	Percent Recovered		
4-Bromofluorobenzene	50	49.2	98			
1,2-Dichloroethane-d4	50	48.5	97			
Toluene-d8	50	49.3	99			

**Comments:****Flags:**

BQL = Below Quantitation Limits.

Reviewed By: gj

**Results for Volatiles  
by GCMS 8260-5030**

Client Sample ID: PAR 206 HA3  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-5A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 11:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 70.9

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	BQL	67.8	3.77	1	2/27/2006	
Benzene	BQL	6.78	4.10	1	2/27/2006	
Bromobenzene	BQL	6.78	4.40	1	2/27/2006	
Bromoform	BQL	6.78	3.00	1	2/27/2006	
Bromochloromethane	BQL	6.78	3.58	1	2/27/2006	
Bromodichloromethane	BQL	6.78	3.07	1	2/27/2006	
Bromomethane	BQL	6.78	6.51	1	2/27/2006	
2-Butanone	BQL	33.9	4.36	1	2/27/2006	
n-Butylbenzene	BQL	6.78	3.72	1	2/27/2006	
sec-Butylbenzene	31.1	6.78	5.25	1	2/27/2006	
tert-Butylbenzene	BQL	6.78	4.99	1	2/27/2006	
Carbon disulfide	BQL	6.78	3.54	1	2/27/2006	
Carbon tetrachloride	BQL	6.78	4.76	1	2/27/2006	
Chlorobenzene	BQL	6.78	4.76	1	2/27/2006	
Chloroethane	BQL	6.78	6.01	1	2/27/2006	
Chloroform	BQL	6.78	3.95	1	2/27/2006	
Chloromethane	BQL	6.78	4.22	1	2/27/2006	
2-Chlorotoluene	BQL	6.78	4.69	1	2/27/2006	
4-Chlorotoluene	BQL	6.78	4.91	1	2/27/2006	
Dibromochloromethane	BQL	6.78	3.42	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	6.78	9.84	1	2/27/2006	
Dibromomethane	BQL	6.78	2.42	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	6.78	2.52	1	2/27/2006	
1,2-Dichlorobenzene	BQL	6.78	4.10	1	2/27/2006	
1,3-Dichlorobenzene	BQL	6.78	4.45	1	2/27/2006	
1,4-Dichlorobenzene	BQL	6.78	4.33	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	6.78	8.94	1	2/27/2006	
1,1-Dichloroethane	BQL	6.78	4.71	1	2/27/2006	
1,1-Dichloroethene	BQL	6.78	4.84	1	2/27/2006	
1,2-Dichloroethane	BQL	6.78	3.24	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	6.78	4.30	1	2/27/2006	
trans-1,2-dichloroethene	BQL	6.78	4.84	1	2/27/2006	
1,2-Dichloropropane	BQL	6.78	3.76	1	2/27/2006	
1,3-Dichloropropane	BQL	6.78	2.78	1	2/27/2006	
2,2-Dichloropropane	BQL	6.78	3.92	1	2/27/2006	
1,1-Dichloropropene	BQL	6.78	4.75	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	6.78	3.17	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	6.78	2.69	1	2/27/2006	
Dichlorodifluoromethane	BQL	6.78	4.88	1	2/27/2006	
Diisopropyl ether (DIPE)	BQL	6.78	3.98	1	2/27/2006	
Ethylbenzene	25.8	6.78	4.83	1	2/27/2006	

**Results for Volatiles**  
by GCMS 8260-5030

Client Sample ID: PAR 206 HA3  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-5A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 11:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 70.9

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	6.78	4.86	1	2/27/2006	
2-Hexanone	BQL	6.78	2.42	1	2/27/2006	
Iodomethane	BQL	6.78	4.88	1	2/27/2006	
Isopropylbenzene	BQL	6.78	5.14	1	2/27/2006	
4-Isopropyltoluene	BQL	6.78	4.99	1	2/27/2006	
Methylene chloride	BQL	27.1	4.18	1	2/27/2006	
4-Methyl-2-pentanone	BQL	6.78	1.61	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	6.78	3.08	1	2/27/2006	
Naphthalene	15.0	6.78	3.23	1	2/27/2006	
n-Propyl benzene	7.22	6.78	5.05	1	2/27/2006	
Styrene	BQL	6.78	6.40	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	6.78	3.95	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	6.78	2.46	1	2/27/2006	
Tetrachloroethene	BQL	6.78	5.18	1	2/27/2006	
Toluene	50.5	6.78	4.38	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	6.78	3.57	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	6.78	3.16	1	2/27/2006	
Trichloroethene	BQL	6.78	4.72	1	2/27/2006	
1,1,1-Trichloroethane	BQL	6.78	4.75	1	2/27/2006	
1,1,2-Trichloroethane	BQL	6.78	2.73	1	2/27/2006	
Trichlorofluoromethane	BQL	6.78	5.33	1	2/27/2006	
1,2,3-Trichloropropane	BQL	6.78	2.52	1	2/27/2006	
1,2,4-Trimethylbenzene	50.8	6.78	4.91	1	2/27/2006	
1,3,5-Trimethylbenzene	15.6	6.78	5.02	1	2/27/2006	
Vinyl chloride	BQL	6.78	4.67	1	2/27/2006	
m-,p-Xylene	125	13.6	9.97	1	2/27/2006	
o-Xylene	57.4	6.78	4.82	1	2/27/2006	
4-Bromofluorobenzene	125.0	57.4	Spike Added	Spike Result	Percent Recovered	
1,2-Dichloroethane-d4	50	50	50	50.2	100	
Toluene-d8	182.4	50	50	45.5	91	
			50	51.2	102	

Comments:

Flags:

BQL = Below Quantitation Limits.

Reviewed By: JTF

**Results for Volatiles  
by GCMS 8260-5030**

Client Sample ID: PAR 206 HA4  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-6A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 11:05  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 73.4

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	BQL	58.2	3.24	1	2/27/2006	
Benzene	BQL	5.82	3.52	1	2/27/2006	
Bromobenzene	BQL	5.82	3.77	1	2/27/2006	
Bromoform	BQL	5.82	2.57	1	2/27/2006	
Bromochloromethane	BQL	5.82	3.07	1	2/27/2006	
Bromodichloromethane	BQL	5.82	2.63	1	2/27/2006	
Bromomethane	BQL	5.82	5.59	1	2/27/2006	
2-Butanone	BQL	29.1	3.74	1	2/27/2006	
n-Butylbenzene	BQL	5.82	3.19	1	2/27/2006	
sec-Butylbenzene	BQL	5.82	4.50	1	2/27/2006	
tert-Butylbenzene	BQL	5.82	4.28	1	2/27/2006	
Carbon disulfide	BQL	5.82	3.04	1	2/27/2006	
Carbon tetrachloride	BQL	5.82	4.09	1	2/27/2006	
Chlorobenzene	BQL	5.82	4.09	1	2/27/2006	
Chloroethane	BQL	5.82	5.16	1	2/27/2006	
Chloroform	BQL	5.82	3.39	1	2/27/2006	
Chloromethane	BQL	5.82	3.62	1	2/27/2006	
2-Chlorotoluene	BQL	5.82	4.03	1	2/27/2006	
4-Chlorotoluene	BQL	5.82	4.21	1	2/27/2006	
Dibromochloromethane	BQL	5.82	2.93	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	5.82	8.44	1	2/27/2006	
Dibromomethane	BQL	5.82	2.07	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	5.82	2.16	1	2/27/2006	
1,2-Dichlorobenzene	BQL	5.82	3.52	1	2/27/2006	
1,3-Dichlorobenzene	BQL	5.82	3.82	1	2/27/2006	
1,4-Dichlorobenzene	BQL	5.82	3.71	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	5.82	7.67	1	2/27/2006	
1,1-Dichloroethane	BQL	5.82	4.04	1	2/27/2006	
1,1-Dichloroethene	BQL	5.82	4.16	1	2/27/2006	
1,2-Dichloroethane	BQL	5.82	2.78	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	5.82	3.69	1	2/27/2006	
trans-1,2-dichloroethene	BQL	5.82	4.16	1	2/27/2006	
1,2-Dichloropropane	BQL	5.82	3.22	1	2/27/2006	
1,3-Dichloropropane	BQL	5.82	2.39	1	2/27/2006	
2,2-Dichloropropane	BQL	5.82	3.36	1	2/27/2006	
1,1-Dichloropropene	BQL	5.82	4.07	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	5.82	2.72	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	5.82	2.30	1	2/27/2006	
Dichlorodifluoromethane	BQL	5.82	4.19	1	2/27/2006	
Diisopropyl ether (Dipe)	BQL	5.82	3.41	1	2/27/2006	
Ethylbenzene	BQL	5.82	4.14	1	2/27/2006	

**Results for Volatiles  
by GCMS 8260-5030**

Client Sample ID: PAR 206 HA4  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-6A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 11:05  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 73.4

<b>Report Name Compound</b>	<b>Result UG/KG</b>	<b>Quantitation Limit UG/KG</b>	<b>MDL UG/KG</b>	<b>Dilution Factor</b>	<b>Date Analyzed</b>	<b>Flag</b>
Hexachlorobutadiene	BQL	5.82	4.17	1	2/27/2006	
2-Hexanone	BQL	5.82	2.07	1	2/27/2006	
Iodomethane	BQL	5.82	4.19	1	2/27/2006	
Isopropylbenzene	BQL	5.82	4.41	1	2/27/2006	
4-Isopropyltoluene	BQL	5.82	4.28	1	2/27/2006	
Methylene chloride	BQL	23.3	3.59	1	2/27/2006	
4-Methyl-2-pentanone	BQL	5.82	1.39	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	5.82	2.64	1	2/27/2006	
Naphthalene	BQL	5.82	2.77	1	2/27/2006	
n-Propyl benzene	BQL	5.82	4.33	1	2/27/2006	
Styrene	BQL	5.82	5.49	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	5.82	3.39	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	5.82	2.11	1	2/27/2006	
Tetrachloroethene	BQL	5.82	4.45	1	2/27/2006	
Toluene	BQL	5.82	3.76	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	5.82	3.06	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	5.82	2.71	1	2/27/2006	
Trichloroethene	BQL	5.82	4.05	1	2/27/2006	
1,1,1-Trichloroethane	BQL	5.82	4.07	1	2/27/2006	
1,1,2-Trichloroethane	BQL	5.82	2.34	1	2/27/2006	
Trichlorofluoromethane	BQL	5.82	4.57	1	2/27/2006	
1,2,3-Trichloropropane	BQL	5.82	2.16	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	5.82	4.21	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	5.82	4.31	1	2/27/2006	
Vinyl chloride	BQL	5.82	4.00	1	2/27/2006	
m-,p-Xylene	BQL	11.6	8.56	1	2/27/2006	
o-Xylene	BQL	5.82	4.13	1	2/27/2006	
		<b>Spike Added</b>	<b>Spike Result</b>	<b>Percent Recovered</b>		
4-Bromofluorobenzene		50	48.4	97		
1,2-Dichloroethane-d4		50	48.4	97		
Toluene-d8		50	52.2	104		

**Comments:****Flags:**

BQL = Below Quantitation Limits.

Reviewed By: BSW

**Results for Volatiles**  
by GCMS 8260-5035

Client Sample ID: PAR 163 GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-9A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 09:22  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 77.2

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	16.6	63.5	3.73	1	2/27/2006	
Benzene	BQL	6.35	3.62	1	2/27/2006	
Bromobenzene	BQL	6.35	3.12	1	2/27/2006	
Bromoform	BQL	6.35	3.71	1	2/27/2006	
Bromochloromethane	BQL	6.35	3.61	1	2/27/2006	
Bromodichloromethane	BQL	6.35	3.07	1	2/27/2006	
Bromomethane	BQL	6.35	5.32	1	2/27/2006	
2-Butanone	5.86	31.8	3.66	1	2/27/2006	
n-Butylbenzene	26.5	6.35	4.25	1	2/27/2006	
sec-Butylbenzene	21.2	6.35	4.45	1	2/27/2006	
tert-Butylbenzene	BQL	6.35	4.41	1	2/27/2006	
Carbon disulfide	BQL	6.35	3.34	1	2/27/2006	
Carbon tetrachloride	BQL	6.35	4.39	1	2/27/2006	
Chlorobenzene	BQL	6.35	3.19	1	2/27/2006	
Chloroethane	BQL	6.35	3.19	1	2/27/2006	
Chloroform	BQL	6.35	3.06	1	2/27/2006	
Chloromethane	BQL	6.35	3.78	1	2/27/2006	
2-Chlorotoluene	BQL	6.35	3.53	1	2/27/2006	
4-Chlorotoluene	BQL	6.35	2.85	1	2/27/2006	
Dibromochloromethane	BQL	6.35	13.5	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	6.35	3.81	1	2/27/2006	
Dibromomethane	BQL	6.35	2.96	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	6.35	3.06	1	2/27/2006	
1,2-Dichlorobenzene	BQL	6.35	2.97	1	2/27/2006	
1,3-Dichlorobenzene	BQL	6.35	3.12	1	2/27/2006	
1,4-Dichlorobenzene	BQL	6.35	13.6	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	6.35	3.66	1	2/27/2006	
1,1-Dichloroethane	BQL	6.35	4.89	1	2/27/2006	
1,1-Dichloroethene	BQL	6.35	3.65	1	2/27/2006	
1,2-Dichloroethane	BQL	6.35	3.12	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	6.35	4.13	1	2/27/2006	
trans-1,2-dichloroethene	BQL	6.35	3.25	1	2/27/2006	
1,2-Dichloropropane	BQL	6.35	2.91	1	2/27/2006	
1,3-Dichloropropane	BQL	6.35	4.04	1	2/27/2006	
2,2-Dichloropropane	BQL	6.35	4.59	1	2/27/2006	
1,1-Dichloropropene	BQL	6.35	3.53	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	6.35	3.59	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	6.35	4.74	1	2/27/2006	
Dichlorodifluoromethane	BQL	6.35	3.01	1	2/27/2006	
Diisopropyl ether (DIPE)	BQL	6.35	3.86	1	2/27/2006	
Ethylbenzene	18.8	6.35				

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 163 GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-9A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 09:22  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 77.2

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	6.35	5.02	1	2/27/2006	
2-Hexanone	BQL	6.35	2.77	1	2/27/2006	
Iodomethane	BQL	6.35	5.91	1	2/27/2006	
Isopropylbenzene	16.3	6.35	4.09	1	2/27/2006	
4-Isopropyltoluene	16.8	6.35	4.32	1	2/27/2006	
Methylene chloride	9.25	25.4	3.63	1	2/27/2006	J
4-Methyl-2-pentanone	BQL	6.35	2.93	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	6.35	3.23	1	2/27/2006	
Naphthalene	95.4	6.35	2.55	1	2/27/2006	
n-Propyl benzene	27.6	6.35	4.08	1	2/27/2006	
Styrene	BQL	6.35	4.55	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	6.35	3.45	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	6.35	3.12	1	2/27/2006	
Tetrachloroethene	BQL	6.35	4.00	1	2/27/2006	
Toluene	BQL	6.35	3.76	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	6.35	2.78	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	6.35	2.85	1	2/27/2006	
Trichloroethene	BQL	6.35	3.96	1	2/27/2006	
1,1,1-Trichloroethane	BQL	6.35	4.38	1	2/27/2006	
1,1,2-Trichloroethane	BQL	6.35	3.28	1	2/27/2006	
Trichlorofluoromethane	BQL	6.35	5.25	1	2/27/2006	
1,2,3-Trichloropropane	BQL	6.35	3.37	1	2/27/2006	
1,2,4-Trimethylbenzene	128	6.35	3.56	1	2/27/2006	
1,3,5-Trimethylbenzene	48.3	6.35	3.78	1	2/27/2006	
Vinyl chloride	BQL	6.35	4.19	1	2/27/2006	
m,p-Xylene	18.4	12.7	7.20	1	2/27/2006	
o-Xylene	34.2	6.35	3.56	1	2/27/2006	
		Spike Added	Spike Result	Percent Recovered		
4-Bromofluorobenzene		50	36.4	73		
1,2-Dichloroethane-d4		50	66.7	133		
Toluene-d8		50	50.5	101		

**Comments:****Flags:**

BQL = Below Quantitation Limits.  
 J = Detected below the quantitation limit.

Reviewed By: ✓

**Results for Volatiles**  
by GCMS 8260-5035

Client Sample ID: PAR 199A GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-14A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 12:45  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 90.9

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	3.97	47.0	2.76	1	2/27/2006	J
Benzene	BQL	4.70	2.68	1	2/27/2006	
Bromobenzene	BQL	4.70	2.31	1	2/27/2006	
Bromoform	BQL	4.70	2.75	1	2/27/2006	
Bromochloromethane	BQL	4.70	2.67	1	2/27/2006	
Bromodichloromethane	BQL	4.70	2.28	1	2/27/2006	
Bromomethane	BQL	4.70	3.94	1	2/27/2006	
2-Butanone	BQL	23.5	2.71	1	2/27/2006	
n-Butylbenzene	BQL	4.70	3.15	1	2/27/2006	
sec-Butylbenzene	BQL	4.70	3.29	1	2/27/2006	
tert-Butylbenzene	BQL	4.70	3.26	1	2/27/2006	
Carbon disulfide	BQL	4.70	2.47	1	2/27/2006	
Carbon tetrachloride	BQL	4.70	3.25	1	2/27/2006	
Chlorobenzene	BQL	4.70	2.36	1	2/27/2006	
Chloroethane	BQL	4.70	2.95	1	2/27/2006	
Chloroform	BQL	4.70	2.36	1	2/27/2006	
Chloromethane	BQL	4.70	2.27	1	2/27/2006	
2-Chlorotoluene	BQL	4.70	2.80	1	2/27/2006	
4-Chlorotoluene	BQL	4.70	2.61	1	2/27/2006	
Dibromochloromethane	BQL	4.70	2.11	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	4.70	9.97	1	2/27/2006	
Dibromomethane	BQL	4.70	2.82	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	4.70	2.19	1	2/27/2006	
1,2-Dichlorobenzene	BQL	4.70	2.27	1	2/27/2006	
1,3-Dichlorobenzene	BQL	4.70	2.20	1	2/27/2006	
1,4-Dichlorobenzene	BQL	4.70	2.31	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	4.70	10.1	1	2/27/2006	
1,1-Dichloroethane	BQL	4.70	2.71	1	2/27/2006	
1,1-Dichloroethene	BQL	4.70	3.62	1	2/27/2006	
1,2-Dichloroethane	BQL	4.70	2.70	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	4.70	2.31	1	2/27/2006	
trans-1,2-dichloroethene	BQL	4.70	3.06	1	2/27/2006	
1,2-Dichloropropane	BQL	4.70	2.41	1	2/27/2006	
1,3-Dichloropropane	BQL	4.70	2.15	1	2/27/2006	
2,2-Dichloropropane	BQL	4.70	2.99	1	2/27/2006	
1,1-Dichloropropene	BQL	4.70	3.39	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	4.70	2.61	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	4.70	2.66	1	2/27/2006	
Dichlorodifluoromethane	BQL	4.70	3.51	1	2/27/2006	
Diisopropyl ether (DIPE)	BQL	4.70	2.23	1	2/27/2006	
Ethylbenzene	BQL	4.70	2.86	1	2/27/2006	

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 199A GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-14A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-22-2006 12:45  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 90.9

<b>Report Name Compound</b>	<b>Result UG/KG</b>	<b>Quantitation Limit UG/KG</b>	<b>MDL UG/KG</b>	<b>Dilution Factor</b>	<b>Date Analyzed</b>	<b>Flag</b>
Hexachlorobutadiene	BQL	4.70	3.71	1	2/27/2006	
2-Hexanone	BQL	4.70	2.05	1	2/27/2006	
Iodomethane	BQL	4.70	4.37	1	2/27/2006	
Isopropylbenzene	BQL	4.70	3.03	1	2/27/2006	
4-Isopropyltoluene	BQL	4.70	3.20	1	2/27/2006	
Methylene chloride	<b>8.67</b>	18.8	2.69	1	2/27/2006	J
4-Methyl-2-pentanone	BQL	4.70	2.17	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	4.70	2.39	1	2/27/2006	
Naphthalene	BQL	4.70	1.89	1	2/27/2006	
n-Propyl benzene	BQL	4.70	3.02	1	2/27/2006	
Styrene	BQL	4.70	3.37	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	4.70	2.56	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	4.70	2.31	1	2/27/2006	
Tetrachloroethene	BQL	4.70	2.96	1	2/27/2006	
Toluene	BQL	4.70	2.78	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	4.70	2.06	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	4.70	2.11	1	2/27/2006	
Trichloroethene	BQL	4.70	2.93	1	2/27/2006	
1,1,1-Trichloroethane	BQL	4.70	3.24	1	2/27/2006	
1,1,2-Trichloroethane	BQL	4.70	2.43	1	2/27/2006	
Trichlorofluoromethane	BQL	4.70	3.88	1	2/27/2006	
1,2,3-Trichloropropane	BQL	4.70	2.49	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	4.70	2.63	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	4.70	2.80	1	2/27/2006	
Vinyl chloride	BQL	4.70	3.10	1	2/27/2006	
m-,p-Xylene	BQL	9.40	5.33	1	2/27/2006	
o-Xylene	BQL	4.70	2.63	1	2/27/2006	
		<b>Spike Added</b>	<b>Spike Result</b>	<b>Percent Recovered</b>		
4-Bromofluorobenzene		50	52.5	105		
1,2-Dichloroethane-d4		50	54.5	109		
Toluene-d8		50	50.6	101		

**Comments:****Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: Paul

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 89 HA6-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-17A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-20-2006 12:45  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 77.2

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	29.9	67.0	3.94	1	2/28/2006	
Benzene	BQL	6.70	3.82	1	2/28/2006	
Bromobenzene	BQL	6.70	3.30	1	2/28/2006	
Bromoform	BQL	6.70	3.92	1	2/28/2006	
Bromochloromethane	BQL	6.70	3.81	1	2/28/2006	
Bromodichloromethane	BQL	6.70	3.25	1	2/28/2006	
Bromomethane	BQL	6.70	5.62	1	2/28/2006	
2-Butanone	6.89	33.5	3.86	1	2/28/2006	J
n-Butylbenzene	BQL	6.70	4.49	1	2/28/2006	
sec-Butylbenzene	BQL	6.70	4.69	1	2/28/2006	
tert-Butylbenzene	BQL	6.70	4.65	1	2/28/2006	
Carbon disulfide	BQL	6.70	3.53	1	2/28/2006	
Carbon tetrachloride	BQL	6.70	4.64	1	2/28/2006	
Chlorobenzene	BQL	6.70	3.37	1	2/28/2006	
Chloroethane	BQL	6.70	4.21	1	2/28/2006	
Chloroform	BQL	6.70	3.37	1	2/28/2006	
Chloromethane	BQL	6.70	3.23	1	2/28/2006	
2-Chlorotoluene	BQL	6.70	4.00	1	2/28/2006	
4-Chlorotoluene	BQL	6.70	3.73	1	2/28/2006	
Dibromochloromethane	BQL	6.70	3.00	1	2/28/2006	
1,2-Dibromo-3-chloropropane	BQL	6.70	14.2	1	2/28/2006	
Dibromomethane	BQL	6.70	4.02	1	2/28/2006	
1,2-Dibromoethane (EDB)	BQL	6.70	3.12	1	2/28/2006	
1,2-Dichlorobenzene	BQL	6.70	3.23	1	2/28/2006	
1,3-Dichlorobenzene	BQL	6.70	3.14	1	2/28/2006	
1,4-Dichlorobenzene	BQL	6.70	3.30	1	2/28/2006	
trans-1,4-Dichloro-2-butene	BQL	6.70	14.3	1	2/28/2006	
1,1-Dichloroethane	BQL	6.70	3.86	1	2/28/2006	
1,1-Dichloroethene	BQL	6.70	5.16	1	2/28/2006	
1,2-Dichloroethane	BQL	6.70	3.85	1	2/28/2006	
cis-1,2-Dichloroethene	BQL	6.70	3.30	1	2/28/2006	
trans-1,2-dichloroethene	BQL	6.70	4.36	1	2/28/2006	
1,2-Dichloropropane	BQL	6.70	3.43	1	2/28/2006	
1,3-Dichloropropane	BQL	6.70	3.07	1	2/28/2006	
2,2-Dichloropropane	BQL	6.70	4.26	1	2/28/2006	
1,1-Dichloropropene	BQL	6.70	4.84	1	2/28/2006	
cis-1,3-Dichloropropene	BQL	6.70	3.73	1	2/28/2006	
trans-1,3-Dichloropropene	BQL	6.70	3.79	1	2/28/2006	
Dichlorodifluoromethane	BQL	6.70	5.00	1	2/28/2006	
Diisopropyl ether (DIPE)	BQL	6.70	3.18	1	2/28/2006	
Ethylbenzene	BQL	6.70	4.08	1	2/28/2006	

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 89 HA6-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-17A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-20-2006 12:45  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 77.2

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	6.70	5.30	1	2/28/2006	
2-Hexanone	BQL	6.70	2.92	1	2/28/2006	
Iodomethane	8.51	6.70	6.24	1	2/28/2006	
Isopropylbenzene	BQL	6.70	4.32	1	2/28/2006	
4-Isopropyltoluene	BQL	6.70	4.56	1	2/28/2006	
Methylene chloride	19.4	26.8	3.84	1	2/28/2006	J
4-Methyl-2-pentanone	BQL	6.70	3.10	1	2/28/2006	
Methyl-tert-butyl ether (MTBE)	BQL	6.70	3.41	1	2/28/2006	
Naphthalene	4.44	6.70	2.70	1	2/28/2006	J
n-Propyl benzene	BQL	6.70	4.30	1	2/28/2006	
Styrene	BQL	6.70	4.80	1	2/28/2006	
1,1,1,2-Tetrachloroethane	BQL	6.70	3.65	1	2/28/2006	
1,1,2,2-Tetrachloroethane	BQL	6.70	3.30	1	2/28/2006	
Tetrachloroethene	BQL	6.70	4.22	1	2/28/2006	
Toluene	5.20	6.70	3.97	1	2/28/2006	J
1,2,3-Trichlorobenzene	BQL	6.70	2.94	1	2/28/2006	
1,2,4-Trichlorobenzene	BQL	6.70	3.00	1	2/28/2006	
Trichloroethene	BQL	6.70	4.18	1	2/28/2006	
1,1,1-Trichloroethane	BQL	6.70	4.63	1	2/28/2006	
1,1,2-Trichloroethane	BQL	6.70	3.46	1	2/28/2006	
Trichlorofluoromethane	BQL	6.70	5.54	1	2/28/2006	
1,2,3-Trichloropropane	BQL	6.70	3.55	1	2/28/2006	
1,2,4-Trimethylbenzene	3.82	6.70	3.75	1	2/28/2006	J
1,3,5-Trimethylbenzene	BQL	6.70	4.00	1	2/28/2006	
Vinyl chloride	BQL	6.70	4.43	1	2/28/2006	
m-,p-Xylene	BQL	13.4	7.60	1	2/28/2006	
o-Xylene	BQL	6.70	3.75	1	2/28/2006	
		Spike Added	Spike Result	Percent Recovered		
4-Bromofluorobenzene		50	55.3	111		
1,2-Dichloroethane-d4		50	74.2	148		
Toluene-d8		50	52.1	104		

## Comments:

## Flags:

BQL = Below Quantitation Limits.  
 J = Detected below the quantitation limit.

Reviewed By: Bob

**Results for Volatiles**  
by GCMS 8260-5035

Client Sample ID: PAR 221 GP1-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID G106-565-24A  
 Lab Project ID: G106-565  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 12:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 75.2

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	10.1	60.0	3.53	1	2/28/2006	J
Benzene	BQL	6.00	3.42	1	2/28/2006	
Bromobenzene	BQL	6.00	2.95	1	2/28/2006	
Bromoform	BQL	6.00	3.50	1	2/28/2006	
Bromochloromethane	BQL	6.00	3.41	1	2/28/2006	
Bromodichloromethane	BQL	6.00	2.90	1	2/28/2006	
Bromomethane	BQL	6.00	5.03	1	2/28/2006	
2-Butanone	BQL	30.0	3.46	1	2/28/2006	
n-Butylbenzene	BQL	6.00	4.02	1	2/28/2006	
sec-Butylbenzene	BQL	6.00	4.20	1	2/28/2006	
tert-Butylbenzene	BQL	6.00	4.16	1	2/28/2006	
Carbon disulfide	BQL	6.00	3.16	1	2/28/2006	
Carbon tetrachloride	BQL	6.00	4.15	1	2/28/2006	
Chlorobenzene	BQL	6.00	3.01	1	2/28/2006	
Chloroethane	BQL	6.00	3.77	1	2/28/2006	
Chloroform	BQL	6.00	3.01	1	2/28/2006	
Chloromethane	BQL	6.00	2.89	1	2/28/2006	
2-Chlorotoluene	BQL	6.00	3.58	1	2/28/2006	
4-Chlorotoluene	BQL	6.00	3.34	1	2/28/2006	
Dibromochloromethane	BQL	6.00	2.69	1	2/28/2006	
1,2-Dibromo-3-chloropropane	BQL	6.00	12.7	1	2/28/2006	
Dibromomethane	BQL	6.00	3.60	1	2/28/2006	
1,2-Dibromoethane (EDB)	BQL	6.00	2.80	1	2/28/2006	
1,2-Dichlorobenzene	BQL	6.00	2.89	1	2/28/2006	
1,3-Dichlorobenzene	BQL	6.00	2.81	1	2/28/2006	
1,4-Dichlorobenzene	BQL	6.00	2.95	1	2/28/2006	
trans-1,4-Dichloro-2-butene	BQL	6.00	12.8	1	2/28/2006	
1,1-Dichloroethane	BQL	6.00	3.46	1	2/28/2006	
1,1-Dichloroethene	BQL	6.00	4.62	1	2/28/2006	
1,2-Dichloroethane	BQL	6.00	3.44	1	2/28/2006	
cis-1,2-Dichloroethene	BQL	6.00	2.95	1	2/28/2006	
trans-1,2-dichloroethene	BQL	6.00	3.90	1	2/28/2006	
1,2-Dichloropropane	BQL	6.00	3.07	1	2/28/2006	
1,3-Dichloropropane	BQL	6.00	2.75	1	2/28/2006	
2,2-Dichloropropane	BQL	6.00	3.82	1	2/28/2006	
1,1-Dichloropropene	BQL	6.00	4.33	1	2/28/2006	
cis-1,3-Dichloropropene	BQL	6.00	3.34	1	2/28/2006	
trans-1,3-Dichloropropene	BQL	6.00	3.40	1	2/28/2006	
Dichlorodifluoromethane	BQL	6.00	4.47	1	2/28/2006	
Diisopropyl ether (Dipe)	BQL	6.00	2.84	1	2/28/2006	
Ethylbenzene	BQL	6.00	3.65	1	2/28/2006	

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: PAR 221 GP1-8

Client Project ID: NCDOT-Yancey

Lab Sample ID G106-565-24A

Lab Project ID: G106-565

Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected: 02-23-2006 12:00  
 Date Received: 2/24/2006  
 Matrix: Soil  
 %Solids: 75.2

<b>Report Name Compound</b>	<b>Result UG/KG</b>	<b>Quantitation Limit UG/KG</b>	<b>MDL UG/KG</b>	<b>Dilution Factor</b>	<b>Date Analyzed</b>	<b>Flag</b>
Hexachlorobutadiene	BQL	6.00	4.74	1	2/28/2006	
2-Hexanone	BQL	6.00	2.62	1	2/28/2006	
Iodomethane	BQL	6.00	5.58	1	2/28/2006	
Isopropylbenzene	BQL	6.00	3.86	1	2/28/2006	
4-Isopropyltoluene	BQL	6.00	4.08	1	2/28/2006	
Methylene chloride	4.55	24.0	3.43	1	2/28/2006	J
4-Methyl-2-pentanone	BQL	6.00	2.77	1	2/28/2006	
Methyl-tert-butyl ether (MTBE)	BQL	6.00	3.05	1	2/28/2006	
Naphthalene	BQL	6.00	2.41	1	2/28/2006	
n-Propyl benzene	BQL	6.00	3.85	1	2/28/2006	
Styrene	BQL	6.00	4.29	1	2/28/2006	
1,1,1,2-Tetrachloroethane	BQL	6.00	3.26	1	2/28/2006	
1,1,2,2-Tetrachloroethane	BQL	6.00	2.95	1	2/28/2006	
Tetrachloroethene	BQL	6.00	3.78	1	2/28/2006	
Toluene	BQL	6.00	3.55	1	2/28/2006	
1,2,3-Trichlorobenzene	BQL	6.00	2.63	1	2/28/2006	
1,2,4-Trichlorobenzene	BQL	6.00	2.69	1	2/28/2006	
Trichloroethene	BQL	6.00	3.74	1	2/28/2006	
1,1,1-Trichloroethane	BQL	6.00	4.14	1	2/28/2006	
1,1,2-Trichloroethane	BQL	6.00	3.10	1	2/28/2006	
Trichlorofluoromethane	BQL	6.00	4.95	1	2/28/2006	
1,2,3-Trichloropropane	BQL	6.00	3.18	1	2/28/2006	
1,2,4-Trimethylbenzene	BQL	6.00	3.36	1	2/28/2006	
1,3,5-Trimethylbenzene	BQL	6.00	3.58	1	2/28/2006	
Vinyl chloride	BQL	6.00	3.96	1	2/28/2006	
m-,p-Xylene	BQL	12.0	6.80	1	2/28/2006	
$\alpha$ -Xylene	BQL	6.00	3.36	1	2/28/2006	
4-Bromofluorobenzene	50	53.8	108			
1,2-Dichloroethane-d4	50	69.6	139			
Toluene-d8	50	51.6	103			

**Comments:****Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: \_\_\_\_\_

**Results for Volatiles**  
by GCMS 8260-5035

Client Sample ID: Method Blank  
 Client Project ID:  
 Lab Sample ID VBLK3022706B  
 Lab Project ID:  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected:  
 Date Received:  
 Matrix: Soil  
 %Solids: 100.0

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	BQL	50.0	2.94	1	2/27/2006	
Benzene	BQL	5.00	2.85	1	2/27/2006	
Bromobenzene	BQL	5.00	2.46	1	2/27/2006	
Bromoform	BQL	5.00	2.92	1	2/27/2006	
Bromochloromethane	BQL	5.00	2.84	1	2/27/2006	
Bromodichloromethane	BQL	5.00	2.42	1	2/27/2006	
Bromomethane	BQL	25.0	2.88	1	2/27/2006	
2-Butanone	BQL	5.00	3.35	1	2/27/2006	
n-Butylbenzene	BQL	5.00	3.50	1	2/27/2006	
sec-Butylbenzene	BQL	5.00	3.47	1	2/27/2006	
tert-Butylbenzene	BQL	5.00	2.63	1	2/27/2006	
Carbon disulfide	BQL	5.00	3.46	1	2/27/2006	
Carbon tetrachloride	BQL	5.00	2.51	1	2/27/2006	
Chlorobenzene	BQL	5.00	3.14	1	2/27/2006	
Chloroethane	BQL	5.00	2.51	1	2/27/2006	
Chloroform	BQL	5.00	2.41	1	2/27/2006	
Chloromethane	BQL	5.00	2.98	1	2/27/2006	
2-Chlorotoluene	BQL	5.00	2.78	1	2/27/2006	
4-Chlorotoluene	BQL	5.00	2.24	1	2/27/2006	
Dibromochloromethane	BQL	5.00	10.6	1	2/27/2006	
1,2-Dibromo-3-chloropropane	BQL	5.00	3.00	1	2/27/2006	
Dibromomethane	BQL	5.00	2.33	1	2/27/2006	
1,2-Dibromoethane (EDB)	BQL	5.00	2.41	1	2/27/2006	
1,2-Dichlorobenzene	BQL	5.00	2.34	1	2/27/2006	
1,3-Dichlorobenzene	BQL	5.00	2.46	1	2/27/2006	
1,4-Dichlorobenzene	BQL	5.00	10.7	1	2/27/2006	
trans-1,4-Dichloro-2-butene	BQL	5.00	2.88	1	2/27/2006	
1,1-Dichloroethane	BQL	5.00	3.85	1	2/27/2006	
1,1-Dichloroethene	BQL	5.00	2.87	1	2/27/2006	
1,2-Dichloroethane	BQL	5.00	2.46	1	2/27/2006	
cis-1,2-Dichloroethene	BQL	5.00	3.25	1	2/27/2006	
trans-1,2-dichloroethene	BQL	5.00	2.56	1	2/27/2006	
1,2-Dichloropropane	BQL	5.00	2.29	1	2/27/2006	
1,3-Dichloropropane	BQL	5.00	3.18	1	2/27/2006	
2,2-Dichloropropane	BQL	5.00	3.61	1	2/27/2006	
1,1-Dichloropropene	BQL	5.00	2.78	1	2/27/2006	
cis-1,3-Dichloropropene	BQL	5.00	2.83	1	2/27/2006	
trans-1,3-Dichloropropene	BQL	5.00	3.73	1	2/27/2006	
Dichlorodifluoromethane	BQL	5.00	2.37	1	2/27/2006	
Diisopropyl ether (DIPE)	BQL	5.00	3.04	1	2/27/2006	
Ethylbenzene	BQL					

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: Method Blank  
 Client Project ID:  
 Lab Sample ID VBLK3022706B  
 Lab Project ID:  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected:  
 Date Received:  
 Matrix: Soil  
 %Solids: 100.0

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	5.00	3.95	1	2/27/2006	
2-Hexanone	BQL	5.00	2.18	1	2/27/2006	
Iodomethane	BQL	5.00	4.65	1	2/27/2006	
Isopropylbenzene	BQL	5.00	3.22	1	2/27/2006	
4-Isopropyltoluene	BQL	5.00	3.40	1	2/27/2006	
Methylene chloride	BQL	20.0	2.86	1	2/27/2006	
4-Methyl-2-pentanone	BQL	5.00	2.31	1	2/27/2006	
Methyl-tert-butyl ether (MTBE)	BQL	5.00	2.54	1	2/27/2006	
Naphthalene	BQL	5.00	2.01	1	2/27/2006	
n-Propyl benzene	BQL	5.00	3.21	1	2/27/2006	
Styrene	BQL	5.00	3.58	1	2/27/2006	
1,1,1,2-Tetrachloroethane	BQL	5.00	2.72	1	2/27/2006	
1,1,2,2-Tetrachloroethane	BQL	5.00	2.46	1	2/27/2006	
Tetrachloroethene	BQL	5.00	3.15	1	2/27/2006	
Toluene	BQL	5.00	2.96	1	2/27/2006	
1,2,3-Trichlorobenzene	BQL	5.00	2.19	1	2/27/2006	
1,2,4-Trichlorobenzene	BQL	5.00	2.24	1	2/27/2006	
Trichloroethene	BQL	5.00	3.12	1	2/27/2006	
1,1,1-Trichloroethane	BQL	5.00	3.45	1	2/27/2006	
1,1,2-Trichloroethane	BQL	5.00	2.58	1	2/27/2006	
Trichlorofluoromethane	BQL	5.00	4.13	1	2/27/2006	
1,2,3-Trichloropropane	BQL	5.00	2.65	1	2/27/2006	
1,2,4-Trimethylbenzene	BQL	5.00	2.80	1	2/27/2006	
1,3,5-Trimethylbenzene	BQL	5.00	2.98	1	2/27/2006	
Vinyl chloride	BQL	5.00	3.30	1	2/27/2006	
m-,p-Xylene	BQL	10.0	5.67	1	2/27/2006	
o-Xylene	BQL	5.00	2.80	1	2/27/2006	
		Spike Added	Spike Result	Percent Recovered		
4-Bromofluorobenzene		50	50.5	101		
1,2-Dichloroethane-d4		50	46.3	93		
Toluene-d8		50	48.3	97		

**Comments:****Flags:**

BQL = Below Quantitation Limits.  
 J = Detected below the quantitation limit.

Reviewed By: RJ

**Results for Laboratory Control Spike (LCS)**  
**by GC/MS 8260/5035**

Lab Sample ID: lcs3022706c  
Analyst: JTF  
Batch ID: 3022706

Date Analyzed: 27 Feb 2006 11:13 pm

Matrix: Soil

compound	Spiked ( $\mu\text{g}/\text{Kg}$ )	Amount recovered	LCS (%)	Limits	
				Lower (%)	Upper (%)
benzene	50	48.06	96.1	77.6	122
chlorobenzene	50	47.91	95.8	75.3	125
1,1-dichloroethene	50	46.31	92.6	78.5	121
toluene	50	48.16	96.3	75.7	124
trichloroethene	50	49.42	98.8	60.8	139

**Comments:** Concentration values are on column amount.

**Flags:** \* = Out of limits.  
NA = Not applicable  
NS = Not spiked

Reviewed by: PM

## Results for MS/MSD by GC/MS 8260/5035

Client Project ID: Batch QC  
 Lab Sample ID: g122-2732-6a  
 Batch ID: 3022706

Date Analyzed: 28 Feb 2006 7:55 am  
 Matrix: Soil  
 Analyzed By: JTF

Compound	Unspiked Sample ug/L	Spike conc. ug/L	Recovered MS %	Recovered MSD %	Limits		RPD %	RPD Limit %
					Lower %	Upper %		
benzene	BQL	50	98.4	98.2	74.8	133	0.2	30
chlorobenzene	BQL	50	101.8	102.7	66.3	135	0.9	30
1,1-dichloroethene	BQL	50	96.7	97.3	72.0	135	0.6	30
toluene	BQL	50	102.2	101.3	70.5	138	0.8	30
trichloroethene	BQL	50	105.3	103.3	60.7	152	1.9	30

**Comments:**

Concentrations are on column amounts.

Concentration Units: ug/L

**Flags:**

\* = Out of limits.

NA = Not applicable

BQL = Below quantitation limit.

Reviewed By: gns

VO41.072303.1

**Results for Volatiles**  
by GCMS 8260-5035

Client Sample ID: Method Blank  
 Client Project ID:  
 Lab Sample ID VBLK3022706D  
 Lab Project ID:  
 Report Basis: Dry Weight

Analyzed By: JTF  
 Date Collected:  
 Date Received:  
 Matrix: Soil  
 %Solids: 100.0

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Acetone	BQL	50.0	2.94	1	2/28/2006	
Benzene	BQL	5.00	2.85	1	2/28/2006	
Bromobenzene	BQL	5.00	2.46	1	2/28/2006	
Bromoform	BQL	5.00	2.92	1	2/28/2006	
Bromochloromethane	BQL	5.00	2.84	1	2/28/2006	
Bromodichloromethane	BQL	5.00	2.42	1	2/28/2006	
Bromomethane	BQL	5.00	4.19	1	2/28/2006	
2-Butanone	BQL	25.0	2.88	1	2/28/2006	
n-Butylbenzene	BQL	5.00	3.35	1	2/28/2006	
sec-Butylbenzene	BQL	5.00	3.50	1	2/28/2006	
tert-Butylbenzene	BQL	5.00	3.47	1	2/28/2006	
Carbon disulfide	BQL	5.00	2.63	1	2/28/2006	
Carbon tetrachloride	BQL	5.00	3.46	1	2/28/2006	
Chlorobenzene	BQL	5.00	2.51	1	2/28/2006	
Chloroethane	BQL	5.00	3.14	1	2/28/2006	
Chloroform	BQL	5.00	2.51	1	2/28/2006	
Chloromethane	BQL	5.00	2.41	1	2/28/2006	
2-Chlorotoluene	BQL	5.00	2.98	1	2/28/2006	
4-Chlorotoluene	BQL	5.00	2.78	1	2/28/2006	
Dibromochloromethane	BQL	5.00	2.24	1	2/28/2006	
1,2-Dibromo-3-chloropropane	BQL	5.00	10.6	1	2/28/2006	
Dibromomethane	BQL	5.00	3.00	1	2/28/2006	
1,2-Dibromoethane (EDB)	BQL	5.00	2.33	1	2/28/2006	
1,2-Dichlorobenzene	BQL	5.00	2.41	1	2/28/2006	
1,3-Dichlorobenzene	BQL	5.00	2.34	1	2/28/2006	
1,4-Dichlorobenzene	BQL	5.00	2.46	1	2/28/2006	
trans-1,4-Dichloro-2-butene	BQL	5.00	10.7	1	2/28/2006	
1,1-Dichloroethane	BQL	5.00	2.88	1	2/28/2006	
1,1-Dichloroethene	BQL	5.00	3.85	1	2/28/2006	
1,2-Dichloroethane	BQL	5.00	2.87	1	2/28/2006	
cis-1,2-Dichloroethene	BQL	5.00	2.46	1	2/28/2006	
trans-1,2-dichloroethene	BQL	5.00	3.25	1	2/28/2006	
1,2-Dichloropropane	BQL	5.00	2.56	1	2/28/2006	
1,3-Dichloropropane	BQL	5.00	2.29	1	2/28/2006	
2,2-Dichloropropane	BQL	5.00	3.18	1	2/28/2006	
1,1-Dichloropropene	BQL	5.00	3.61	1	2/28/2006	
cis-1,3-Dichloropropene	BQL	5.00	2.78	1	2/28/2006	
trans-1,3-Dichloropropene	BQL	5.00	2.83	1	2/28/2006	
Dichlorodifluoromethane	BQL	5.00	3.73	1	2/28/2006	
Diisopropyl ether (DIPE)	BQL	5.00	2.37	1	2/28/2006	
Ethylbenzene	BQL	5.00	3.04	1	2/28/2006	

**Results for Volatiles  
by GCMS 8260-5035**

Client Sample ID: Method Blank

Analyzed By: JTF

Client Project ID:

Date Collected:

Lab Sample ID VBLK3022706D

Date Received:

Lab Project ID:

Matrix: Soil

Report Basis: Dry Weight

%Solids: 100.0

Report Name Compound	Result UG/KG	Quantitation Limit UG/KG	MDL UG/KG	Dilution Factor	Date Analyzed	Flag
Hexachlorobutadiene	BQL	5.00	3.95	1	2/28/2006	
2-Hexanone	BQL	5.00	2.18	1	2/28/2006	
Iodomethane	BQL	5.00	4.65	1	2/28/2006	
Isopropylbenzene	BQL	5.00	3.22	1	2/28/2006	
4-Isopropyltoluene	BQL	5.00	3.40	1	2/28/2006	
Methylene chloride	BQL	20.0	2.86	1	2/28/2006	
4-Methyl-2-pentanone	BQL	5.00	2.31	1	2/28/2006	
Methyl-tert-butyl ether (MTBE)	BQL	5.00	2.54	1	2/28/2006	
Naphthalene	BQL	5.00	2.01	1	2/28/2006	
n-Propyl benzene	BQL	5.00	3.21	1	2/28/2006	
Styrene	BQL	5.00	3.58	1	2/28/2006	
1,1,1,2-Tetrachloroethane	BQL	5.00	2.72	1	2/28/2006	
1,1,2,2-Tetrachloroethane	BQL	5.00	2.46	1	2/28/2006	
Tetrachloroethene	BQL	5.00	3.15	1	2/28/2006	
Toluene	BQL	5.00	2.96	1	2/28/2006	
1,2,3-Trichlorobenzene	BQL	5.00	2.19	1	2/28/2006	
1,2,4-Trichlorobenzene	BQL	5.00	2.24	1	2/28/2006	
Trichloroethene	BQL	5.00	3.12	1	2/28/2006	
1,1,1-Trichloroethane	BQL	5.00	3.45	1	2/28/2006	
1,1,2-Trichloroethane	BQL	5.00	2.58	1	2/28/2006	
Trichlorofluoromethane	BQL	5.00	4.13	1	2/28/2006	
1,2,3-Trichloropropane	BQL	5.00	2.65	1	2/28/2006	
1,2,4-Trimethylbenzene	BQL	5.00	2.80	1	2/28/2006	
1,3,5-Trimethylbenzene	BQL	5.00	2.98	1	2/28/2006	
Vinyl chloride	BQL	5.00	3.30	1	2/28/2006	
m-,p-Xylene	BQL	10.0	5.67	1	2/28/2006	
o-Xylene	BQL	5.00	2.80	1	2/28/2006	
		Spike Added	Spike Result	Percent Recovered		
4-Bromofluorobenzene		50	51.3	103		
1,2-Dichloroethane-d4		50	46.8	94		
Toluene-d8		50	50.2	100		

**Comments:****Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By:

**Results for MS/MSD**  
**by GC/MS 8260/5035**

Client Project ID: Batch QC  
 Lab Sample ID: g122-2735-1a  
 Batch ID: 3022706

Date Analyzed: 27 Feb 2006 8:08 pm  
 Matrix: Soil  
 Analyzed By: JTF

Compound	Unspiked Sample ug/L	Spike conc. ug/L	Recovered MS %	Recovered MSD %	Limits		RPD %	RPD Limit %
					Lower %	Upper %		
benzene	BQL	50	95.7	98.0	74.8	133	2.4	30
chlorobenzene	BQL	50	95.2	93.1	66.3	135	2.3	30
1,1-dichloroethene	BQL	50	92.8	94.5	72.0	135	1.8	30
toluene	0.5	50	94.6	95.8	70.5	138	1.3	30
trichloroethene	BQL	50	93.5	100.3	60.7	152	7.0	30

**Comments:**

Concentrations are on column amounts.

Concentration Units: ug/L

**Flags:**

\* = Out of limits.

NA = Not applicable

BQL = Below quantitation limit.

Reviewed By: JTF

VO41.072303.1

**Results for Laboratory Control Spike (LCS)**  
**by GC/MS 8260/5035**

Lab Sample ID: lcs3022706a  
Analyst: JTF  
Batch ID: 3022706

Date Analyzed: 27 Feb 2006 10:23 am  
Matrix: Soil

compound	Spiked ( $\mu\text{g}/\text{Kg}$ )	Amount recovered	LCS (%)	Limits	
				Lower (%)	Upper (%)
benzene	50	48.24	96.5	77.6	122
chlorobenzene	50	40.33	80.7	75.3	125
1,1-dichloroethene	50	43.40	86.8	78.5	121
toluene	50	45.36	90.7	75.7	124
trichloroethene	50	43.65	87.3	60.8	139

**Comments:** Concentration values are on column amount.

**Flags:** \* = Out of limits.

NA = Not applicable

NS = Not spiked

Reviewed by: \_\_\_\_\_

VO39.072303.1

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 163 GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-9K  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/22/2006 9:22  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 77.19

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	403	57.7	1	3/1/2006	
Acenaphthylene	BQL	403	53.6	1	3/1/2006	
Anthracene	BQL	403	58.5	1	3/1/2006	
Benzo[a]anthracene	BQL	403	69.8	1	3/1/2006	
Benzo[a]pyrene	BQL	403	61.7	1	3/1/2006	
Benzo[b]fluoranthene	BQL	403	70.6	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	403	110	1	3/1/2006	
Benzo[k]fluoranthene	BQL	403	77.8	1	3/1/2006	
Benzoic Acid	BQL	807	807	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	403	60.1	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	403	48.8	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	403	50.4	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	403	54.0	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	403	68.2	1	3/1/2006	
Butylbenzylphthalate	BQL	403	62.1	1	3/1/2006	
2-Chloronaphthalene	BQL	403	63.3	1	3/1/2006	
2-Chlorophenol	BQL	403	126	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	403	126	1	3/1/2006	
4-Chloroaniline	BQL	2020	307	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	403	59.3	1	3/1/2006	
Chrysene	BQL	403	43.6	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	403	113	1	3/1/2006	
Dibenzofuran	BQL	403	73.4	1	3/1/2006	
Di-n-Butylphthalate	BQL	403	48.0	1	3/1/2006	
1,2-Dichlorobenzene	BQL	403	44.8	1	3/1/2006	
1,3-Dichlorobenzene	BQL	403	44.0	1	3/1/2006	
1,4-Dichlorobenzene	BQL	403	45.6	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	807	102	1	3/1/2006	
2,4-Dichlorophenol	BQL	403	145	1	3/1/2006	
Diethylphthalate	BQL	403	52.0	1	3/1/2006	
Dimethylphthalate	BQL	403	48.8	1	3/1/2006	
2,4-Dimethylphenol	BQL	403	288	1	3/1/2006	
Di-n-octylphthalate	BQL	403	66.5	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	2020	238	1	3/1/2006	
2,4-Dinitrophenol	BQL	2020	888	1	3/1/2006	
2,4-Dinitrotoluene	BQL	403	52.4	1	3/1/2006	
2,6-Dinitrotoluene	BQL	403	73.4	1	3/1/2006	
Diphenylamine *	BQL	403	39.5	1	3/1/2006	
Fluoranthene	BQL	403	56.5	1	3/1/2006	
Fluorene	BQL	403	50.0	1	3/1/2006	
Hexachlorobenzene	BQL	403	62.1	1	3/1/2006	
Hexachlorobutadiene	BQL	403	64.5	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	807	41.5	1	3/1/2006	
Hexachloroethane	BQL	403	36.3	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	403	103	1	3/1/2006	

**Results for Semivolatiles**  
**by GCMS 8270**

Client Sample ID: PAR 163 GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-9K  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/22/2006 9:22  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 77.19

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Isophorone	BQL	403	59.3	1	3/1/2006	
2-Methylnaphthalene	BQL	403	118	1	3/1/2006	
2-Methylphenol	BQL	403	142	1	3/1/2006	
3- & 4-Methylphenol	BQL	403	137	1	3/1/2006	
Naphthalene	BQL	403	32.7	1	3/1/2006	
2-Nitroaniline	BQL	403	63.3	1	3/1/2006	
3-Nitroaniline	BQL	2020	415	1	3/1/2006	
4-Nitroaniline	BQL	2020	124	1	3/1/2006	
Nitrobenzene	BQL	403	54.4	1	3/1/2006	
2-Nitrophenol	BQL	403	125	1	3/1/2006	
4-Nitrophenol	BQL	2020	112	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	403	51.2	1	3/1/2006	
Pentachlorophenol	BQL	2020	105	1	3/1/2006	
Phenanthrene	BQL	403	46.0	1	3/1/2006	
Phenol	BQL	403	111	1	3/1/2006	
Pyrene	BQL	403	77.4	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	403	50.4	1	3/1/2006	
2,4,5-Trichlorophenol	BQL	403	156	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	403	144	1	3/1/2006	

	Spike Added	Spike Result	Percent Recovered
2-Fluorobiphenyl	10	10.1	101
2-Fluorophenol	10	11	110
Nitrobenzene-d5	10	11.2	112
Phenol-d6	10	11.2	112
2,4,6-Tribromophenol	10	10.7	107
4-Terphenyl-d14	10	10.9	109

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: ZNP

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 199A GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-14N  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/22/2006 12:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 90.91

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	343	49.1	1	3/1/2006	
Acenaphthylene	BQL	343	45.6	1	3/1/2006	
Anthracene	BQL	343	49.7	1	3/1/2006	
Benzo[a]anthracene	BQL	343	59.4	1	3/1/2006	
Benzo[a]pyrene	BQL	343	52.5	1	3/1/2006	
Benzo[b]fluoranthene	BQL	343	60.0	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	343	93.3	1	3/1/2006	
Benzo[k]fluoranthene	BQL	343	66.2	1	3/1/2006	
Benzoic Acid	BQL	686	686	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	343	51.1	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	343	41.5	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	343	42.9	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	343	46.0	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	343	58.0	1	3/1/2006	
Butylbenzylphthalate	BQL	343	52.8	1	3/1/2006	
2-Chloronaphthalene	BQL	343	53.9	1	3/1/2006	
2-Chlorophenol	BQL	343	107	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	343	107	1	3/1/2006	
4-Chloroaniline	BQL	1720	261	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	343	50.4	1	3/1/2006	
Chrysene	BQL	343	37.1	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	343	96.1	1	3/1/2006	
Dibenzofuran	BQL	343	62.4	1	3/1/2006	
Di-n-Butylphthalate	BQL	343	40.8	1	3/1/2006	
1,2-Dichlorobenzene	BQL	343	38.1	1	3/1/2006	
1,3-Dichlorobenzene	BQL	343	37.4	1	3/1/2006	
1,4-Dichlorobenzene	BQL	343	38.8	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	686	86.5	1	3/1/2006	
2,4-Dichlorophenol	BQL	343	124	1	3/1/2006	
Diethylphthalate	BQL	343	44.3	1	3/1/2006	
Dimethylphthalate	BQL	343	41.5	1	3/1/2006	
2,4-Dimethylphenol	BQL	343	245	1	3/1/2006	
Di-n-octylphthalate	BQL	343	56.6	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	1720	202	1	3/1/2006	
2,4-Dinitrophenol	BQL	1720	756	1	3/1/2006	
2,4-Dinitrotoluene	BQL	343	44.6	1	3/1/2006	
2,6-Dinitrotoluene	BQL	343	62.4	1	3/1/2006	
Diphenylamine *	BQL	343	33.6	1	3/1/2006	
Fluoranthene	BQL	343	48.0	1	3/1/2006	
Fluorene	BQL	343	42.5	1	3/1/2006	
Hexachlorobenzene	BQL	343	52.8	1	3/1/2006	
Hexachlorobutadiene	BQL	343	54.9	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	686	35.3	1	3/1/2006	
Hexachloroethane	BQL	343	30.9	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	343	87.8	1	3/1/2006	

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 199A GP2-10  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-14N  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/22/2006 12:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 90.91

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Isophorone	BQL	343	50.4	1	3/1/2006	
2-Methylnaphthalene	BQL	343	100	1	3/1/2006	
2-Methylphenol	BQL	343	121	1	3/1/2006	
3- & 4-Methylphenol	BQL	343	116	1	3/1/2006	
Naphthalene	BQL	343	27.8	1	3/1/2006	
2-Nitroaniline	BQL	343	53.9	1	3/1/2006	
3-Nitroaniline	BQL	1720	353	1	3/1/2006	
4-Nitroaniline	BQL	1720	106	1	3/1/2006	
Nitrobenzene	BQL	343	46.3	1	3/1/2006	
2-Nitrophenol	BQL	343	106	1	3/1/2006	
4-Nitrophenol	BQL	1720	95.0	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	343	43.6	1	3/1/2006	
Pentachlorophenol	BQL	1720	89.5	1	3/1/2006	
Phenanthrene	BQL	343	39.1	1	3/1/2006	
Phenol	BQL	343	94.0	1	3/1/2006	
Pyrene	BQL	343	65.9	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	343	42.9	1	3/1/2006	
2,4,5-Trichlorophenol	BQL	343	133	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	343	122	1	3/1/2006	

	Spike Added	Spike Result	Percent Recovered
2-Fluorobiphenyl	10	10.3	103
2-Fluorophenol	10	8.8	88
Nitrobenzene-d5	10	10.7	107
Phenol-d6	10	9.2	92
2,4,6-Tribromophenol	10	9	90
4-Terphenyl-d14	10	11.4	114

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: rsj

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 89 HA6-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-17L  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/20/2006 12:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 77.2

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	404	57.7	1	3/1/2006	
Acenaphthylene	BQL	404	53.7	1	3/1/2006	
Anthracene	BQL	404	58.5	1	3/1/2006	
Benzo[a]anthracene	BQL	404	69.9	1	3/1/2006	
Benzo[a]pyrene	BQL	404	61.8	1	3/1/2006	
Benzo[b]fluoranthene	BQL	404	70.7	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	404	110	1	3/1/2006	
Benzo[k]fluoranthene	BQL	404	77.9	1	3/1/2006	
Benzoic Acid	BQL	808	808	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	404	60.2	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	404	48.9	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	404	50.5	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	404	54.1	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	404	68.2	1	3/1/2006	
Butylbenzylphthalate	BQL	404	62.2	1	3/1/2006	
2-Chloronaphthalene	BQL	404	63.4	1	3/1/2006	
2-Chlorophenol	BQL	404	126	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	404	126	1	3/1/2006	
4-Chloroaniline	BQL	2020	308	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	404	59.4	1	3/1/2006	
Chrysene	BQL	404	43.6	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	404	113	1	3/1/2006	
Dibenzofuran	BQL	404	73.5	1	3/1/2006	
Di-n-Butylphthalate	BQL	404	48.1	1	3/1/2006	
1,2-Dichlorobenzene	BQL	404	44.8	1	3/1/2006	
1,3-Dichlorobenzene	BQL	404	44.0	1	3/1/2006	
1,4-Dichlorobenzene	BQL	404	45.6	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	808	102	1	3/1/2006	
2,4-Dichlorophenol	BQL	404	145	1	3/1/2006	
Diethylphthalate	BQL	404	52.1	1	3/1/2006	
Dimethylphthalate	BQL	404	48.9	1	3/1/2006	
2,4-Dimethylphenol	BQL	404	289	1	3/1/2006	
Di-n-octylphthalate	BQL	404	66.6	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	2020	238	1	3/1/2006	
2,4-Dinitrophenol	BQL	2020	889	1	3/1/2006	
2,4-Dinitrotoluene	BQL	404	52.5	1	3/1/2006	
2,6-Dinitrotoluene	BQL	404	73.5	1	3/1/2006	
Diphenylamine *	BQL	404	39.6	1	3/1/2006	
Fluoranthene	BQL	404	56.5	1	3/1/2006	
Fluorene	BQL	404	50.1	1	3/1/2006	
Hexachlorobenzene	BQL	404	62.2	1	3/1/2006	
Hexachlorobutadiene	BQL	404	64.6	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	808	41.6	1	3/1/2006	
Hexachloroethane	BQL	404	36.3	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	404	103	1	3/1/2006	

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 89 HA6-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-17L  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/20/2006 12:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 77.2

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Isophorone	BQL	404	59.4	1	3/1/2006	
2-Methylnaphthalene	BQL	404	118	1	3/1/2006	
2-Methylphenol	BQL	404	142	1	3/1/2006	
3- & 4-Methylphenol	BQL	404	137	1	3/1/2006	
Naphthalene	BQL	404	32.7	1	3/1/2006	
2-Nitroaniline	BQL	404	63.4	1	3/1/2006	
3-Nitroaniline	BQL	2020	416	1	3/1/2006	
4-Nitroaniline	BQL	2020	124	1	3/1/2006	
Nitrobenzene	BQL	404	54.5	1	3/1/2006	
2-Nitrophenol	BQL	404	125	1	3/1/2006	
4-Nitrophenol	BQL	2020	112	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	404	51.3	1	3/1/2006	
Pentachlorophenol	BQL	2020	105	1	3/1/2006	
Phenanthrene	BQL	404	46.0	1	3/1/2006	
Phenol	BQL	404	111	1	3/1/2006	
Pyrene	BQL	404	77.5	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	404	50.5	1	3/1/2006	
2,4,5-Trichlorophenol	BQL	404	156	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	404	144	1	3/1/2006	

	Spike Added	Spike Result	Percent Recovered
2-Fluorobiphenyl	10	9.9	99
2-Fluorophenol	10	8.3	83
Nitrobenzene-d5	10	10.6	106
Phenol-d6	10	9.1	91
2,4,6-Tribromophenol	10	7.5	75
4-Terphenyl-d14	10	10.7	107

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: RWJ

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 221 GP1-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-24G  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/23/2006 12:00  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 75.23

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	410	58.6	1	3/1/2006	
Acenaphthylene	BQL	410	54.5	1	3/1/2006	
Anthracene	BQL	410	59.4	1	3/1/2006	
Benzo[a]anthracene	BQL	410	70.8	1	3/1/2006	
Benzo[a]pyrene	BQL	410	62.7	1	3/1/2006	
Benzo[b]fluoranthene	BQL	410	71.7	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	410	111	1	3/1/2006	
Benzo[k]fluoranthene	BQL	410	79.0	1	3/1/2006	
Benzoic Acid	BQL	819	819	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	410	61.0	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	410	49.6	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	410	51.2	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	410	54.9	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	410	69.2	1	3/1/2006	
Butylbenzylphthalate	BQL	410	63.1	1	3/1/2006	
2-Chloronaphthalene	BQL	410	64.3	1	3/1/2006	
2-Chlorophenol	BQL	410	128	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	2050	128	1	3/1/2006	
4-Chloroaniline	BQL	410	312	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	410	60.2	1	3/1/2006	
Chrysene	BQL	410	44.2	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	410	115	1	3/1/2006	
Dibenzofuran	BQL	410	74.5	1	3/1/2006	
Di-n-Butylphthalate	BQL	410	48.7	1	3/1/2006	
1,2-Dichlorobenzene	BQL	410	45.5	1	3/1/2006	
1,3-Dichlorobenzene	BQL	410	44.6	1	3/1/2006	
1,4-Dichlorobenzene	BQL	410	46.3	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	819	103	1	3/1/2006	
2,4-Dichlorophenol	BQL	410	147	1	3/1/2006	
Diethylphthalate	BQL	410	52.8	1	3/1/2006	
Dimethylphthalate	BQL	410	49.6	1	3/1/2006	
2,4-Dimethylphenol	BQL	410	293	1	3/1/2006	
Di-n-octylphthalate	BQL	410	67.6	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	2050	241	1	3/1/2006	
2,4-Dinitrophenol	BQL	2050	902	1	3/1/2006	
2,4-Dinitrotoluene	BQL	410	53.2	1	3/1/2006	
2,6-Dinitrotoluene	BQL	410	74.5	1	3/1/2006	
Diphenylamine *	BQL	410	40.1	1	3/1/2006	
Fluoranthene	BQL	410	57.3	1	3/1/2006	
Fluorene	BQL	410	50.8	1	3/1/2006	
Hexachlorobenzene	BQL	410	63.1	1	3/1/2006	
Hexachlorobutadiene	BQL	410	65.5	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	819	42.2	1	3/1/2006	
Hexachloroethane	BQL	410	36.9	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	410	105	1	3/1/2006	

**Results for Semivolatiles**  
by GCMS 8270

Client Sample ID: PAR 221 GP1-8  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-24G  
 Lab Project ID: G106-565  
 Report Basis: Dry weight

Analyzed By: MRC  
 Date Collected: 2/23/2006 12:00  
 Date Received: 2/24/2006  
 Date Extracted: 2/28/2006  
 Matrix: Soil  
 % Solids: 75.23

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Isophorone	BQL	410	60.2	1	3/1/2006	
2-Methylnaphthalene	BQL	410	120	1	3/1/2006	
2-Methylphenol	BQL	410	144	1	3/1/2006	
3- & 4-Methylphenol	BQL	410	139	1	3/1/2006	
Naphthalene	BQL	410	33.2	1	3/1/2006	
2-Nitroaniline	BQL	410	64.3	1	3/1/2006	
3-Nitroaniline	BQL	2050	422	1	3/1/2006	
4-Nitroaniline	BQL	2050	126	1	3/1/2006	
Nitrobenzene	BQL	410	55.3	1	3/1/2006	
2-Nitrophenol	BQL	410	127	1	3/1/2006	
4-Nitrophenol	BQL	2050	113	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	410	52.0	1	3/1/2006	
Pentachlorophenol	BQL	2050	107	1	3/1/2006	
Phenanthrene	BQL	410	46.7	1	3/1/2006	
Phenol	BQL	410	112	1	3/1/2006	
Pyrene	BQL	410	78.6	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	410	51.2	1	3/1/2006	
2,4,5-Trichlorophenol	BQL	410	158	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	410	146	1	3/1/2006	

	Spike Added	Spike Result	Percent Recovered
2-Fluorobiphenyl	10	9.6	96
2-Fluorophenol	10	7.9	79
Nitrobenzene-d5	10	9.8	98
Phenol-d6	10	8.9	89
2,4,6-Tribromophenol	10	7.2	73
4-Terphenyl-d14	10	10.5	105

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.  
 J = Detected below the quantitation limit.

Reviewed By: sw

**Results for Semivolatiles**  
**by GCMS 8270**

Client Sample ID: Method Blank  
 Client Project ID:  
 Lab Sample ID: PB4597  
 Lab Project ID:  
 Report Basis: Dry Weight

Analyzed By: MRC  
 Date Collected:  
 Date Received:  
 Date Extracted: 2/28/2006  
 Matrix: SOIL  
 % Solids: 100

Compound	Result ug/Kg	RL ug/Kg	MDL ug/Kg	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	313	44.7	1	3/1/2006	
Acenaphthylene	BQL	313	41.6	1	3/1/2006	
Anthracene	BQL	313	45.3	1	3/1/2006	
Benzo[a]anthracene	BQL	313	54.1	1	3/1/2006	
Benzo[a]pyrene	BQL	313	47.8	1	3/1/2006	
Benzo[b]fluoranthene	BQL	313	54.7	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	313	85.0	1	3/1/2006	
Benzo[k]fluoranthene	BQL	313	60.3	1	3/1/2006	
Benzoic Acid	BQL	625	625	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	313	46.6	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	313	37.8	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	313	39.1	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	313	41.9	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	313	52.8	1	3/1/2006	
Butylbenzylphthalate	BQL	313	48.1	1	3/1/2006	
2-Chloronaphthalene	BQL	313	49.1	1	3/1/2006	
2-Chlorophenol	BQL	313	97.8	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	313	97.5	1	3/1/2006	
4-Chloroaniline	BQL	1560	238	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	313	45.9	1	3/1/2006	
Chrysene	BQL	313	33.8	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	313	87.5	1	3/1/2006	
Dibenzofuran	BQL	313	56.9	1	3/1/2006	
Di-n-Butylphthalate	BQL	313	37.2	1	3/1/2006	
1,2-Dichlorobenzene	BQL	313	34.7	1	3/1/2006	
1,3-Dichlorobenzene	BQL	313	34.1	1	3/1/2006	
1,4-Dichlorobenzene	BQL	313	35.3	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	625	78.8	1	3/1/2006	
2,4-Dichlorophenol	BQL	313	113	1	3/1/2006	
Diethylphthalate	BQL	313	40.3	1	3/1/2006	
Dimethylphthalate	BQL	313	37.8	1	3/1/2006	
2,4-Dimethylphenol	BQL	313	223	1	3/1/2006	
Di-n-octylphthalate	BQL	313	51.6	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	1560	184	1	3/1/2006	
2,4-Dinitrophenol	BQL	1560	688	1	3/1/2006	
2,4-Dinitrotoluene	BQL	313	40.6	1	3/1/2006	
2,6-Dinitrotoluene	BQL	313	56.9	1	3/1/2006	
Diphenylamine *	BQL	313	30.6	1	3/1/2006	
Fluoranthene	BQL	313	43.7	1	3/1/2006	
Fluorene	BQL	313	38.8	1	3/1/2006	
Hexachlorobenzene	BQL	313	48.1	1	3/1/2006	
Hexachlorobutadiene	BQL	313	50.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	625	32.2	1	3/1/2006	
Hexachloroethane	BQL	313	28.1	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	313	80.0	1	3/1/2006	

**Results For Matrix Spike / Matrix Spike Duplicate and Laboratory Control Standard (MS/MSD/LCS)**  
by GCMS

Client Sample ID: Batch QC

Date Collected:

Client Project ID:

Date Received:

Lab Sample ID: Batch-4597-MS/MSD/LCS

Date Extracted: 02/28/06

Lab Project ID:

Date Analyzed: 03/02/06

Matrix: SOIL

Analyzed By: MRC

Prep Method: 3540

Dilution: 1

	Sample	MS	MS	MS	MSD	MSD	MSD	RPD	QC Limits	
	Amount ( $\mu\text{g/kg}$ )	Spike ( $\mu\text{g/kg}$ )	Conc. ( $\mu\text{g/kg}$ )	% Rec.	Spike ( $\mu\text{g/kg}$ )	Conc. ( $\mu\text{g/kg}$ )	% Rec.		RPD	% Rec.
Acenaphthylene	BQL	3360	4160	124	3350	3880	116.0	6.59	30	73.0-140
4-Chloro-3-methylphenol	BQL	3360	3550	106	3350	3320	99.1	6.45	30	80.0-115
2-Chlorophenol	BQL	3360	3380	101	3350	3140	93.8	7.00	30	77.1-111
1,4-Dichlorobenzene	BQL	3360	3180	94.8	3350	2990	89.3	5.98	30	70.6-117
2,4-Dinitrotoluene	BQL	3360	3500	104	3350	3270	97.6	6.45	30	67.6-136
N-Nitrosodi-n-propylamine	BQL	3360	3240	96.6	3350	2960	88.4	8.86	30	74.3-133
4-Nitrophenol	BQL	3360	3780	113	3350	3450	103.0	9.00	30	56.8-133
Pentachlorophenol	BQL	3360	3410	101	3350	3190	95.2	6.31	30	29.2-108
Phenol	BQL	3360	3430	102	3350	3240	96.8	5.33	30	71.2-120
Pyrene	BQL	3360	3360	100	3350	3190	95.4	4.81	30	68.5-140
1,2,4-Trichlorobenzene	BQL	3360	3200	95.4	3350	3010	89.9	5.94	30	68.9-119

	Spiked Amount ( $\mu\text{g/kg}$ )	LCS Conc. ( $\mu\text{g/kg}$ )	LCS Spike %	QC Limits % Rec.
Acenaphthylene	3125	3770	121	80.9-143
4-Chloro-3-methylphenol	3125	3200	102	83.9-124
2-Chlorophenol	3125	3080	98.7	80.3-119
1,4-Dichlorobenzene	3125	2940	94.0	76.3-118
2,4-Dinitrotoluene	3125	3210	103	80.6-126
N-Nitrosodi-n-propylamine	3125	2910	93.0	80.3-131
4-Nitrophenol	3125	3370	108	60.0-145
Pentachlorophenol	3125	3080	98.4	36.4-114
Phenol	3125	3100	99.3	74.3-117
Pyrene	3125	3120	99.7	74.7-141
1,2,4-Trichlorobenzene	3125	2950	94.4	74.1-120

## Comments:

Concentrations reflect the spiked sample amounts.

## Flags:

\* = Out of limits.

NA = Not applicable.

Reviewed By: RL

**Results for Oil and Grease**

Client Sample ID: PAR 206 GP1  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: g106-565-1p  
Lab Project ID: G106-565  
Matrix: Soil

Date Analyzed: 3/2/2006  
Analyzed By: nio  
Date Collected: 2/22/2006 16:00  
Date Received: 2/24/2006  
Solids: 78.08

Parameter	Method	RL	Result
		MG/KG	MG/KG
Oil & Grease	9071	38.5	BQL

**Comments:**

BQL = Below Quantitation Limit  
All soils are corrected for percent solids.

Reviewed By: apl  
9071\_LIMS\_v1.35

**Results for Oil and Grease**

Client Sample ID: PAR 206 HA1  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: g106-565-2L  
Lab Project ID: G106-565  
Matrix: Soil

Date Analyzed: 3/2/2006  
Analyzed By: nio  
Date Collected: 2/22/2006 17:00  
Date Received: 2/24/2006  
Solids: 75.83

Parameter	Method	RL	Result
		MG/KG	MG/KG
Oil & Grease	9071	40.7	BQL

**Comments:**

BQL = Below Quantitation Limit  
All soils are corrected for percent solids.

Reviewed By: pmf  
9071\_LIMS\_v1.35

**Results for Oil and Grease**

Client Sample ID: PAR 206 HA1A  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: g106-565-3L  
Lab Project ID: G106-565  
Matrix: Soil

Date Analyzed: 3/2/2006  
Analyzed By: nio  
Date Collected: 2/23/2006 9:50  
Date Received: 2/24/2006  
Solids: 76.88

Parameter	Method	RL	Result
		MG/KG	MG/KG
Oil & Grease	9071	40.3	BQL

**Comments:**

BQL = Below Quantitation Limit  
All soils are corrected for percent solids.

Reviewed By: ZV  
9071\_LIMS\_v1.35

**Results for Oil and Grease**

Client Sample ID: PAR 206 HA2  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: g106-565-41  
Lab Project ID: G106-565  
Matrix: Soil

Date Analyzed: 3/2/2006  
Analyzed By: nio  
Date Collected: 2/23/2006 10:20  
Date Received: 2/24/2006  
Solids: 77.25

Parameter	Method	RL	Result
		MG/KG	MG/KG
Oil & Grease	9071	40	BQL

**Comments:**

BQL = Below Quantitation Limit  
All soils are corrected for percent solids.

Reviewed By: Bob  
9071\_LIMS\_v1.35

**Results for Oil and Grease**

Client Sample ID: PAR 206 HA3.  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: g106-565-51  
Lab Project ID: G106-565  
Matrix: Soil

Date Analyzed: 3/2/2006  
Analyzed By: nio  
Date Collected: 2/23/2006 11:00  
Date Received: 2/24/2006  
Solids: 70.87

Parameter	Method	RL	Result
		MG/KG	MG/KG
Oil & Grease	9071	42.3	BQL

**Comments:**

BQL = Below Quantitation Limit  
All soils are corrected for percent solids.

Reviewed By: psj  
9071\_LIMS\_v1.35

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**Results for Oil and Grease**

Client Sample ID: PAR 206 HA4  
Client Project ID: NCDOT-Yancey  
Lab Sample ID: g106-565-6J  
Lab Project ID: G106-565  
Matrix: Soil

Date Analyzed: 3/2/2006  
Analyzed By: nio  
Date Collected: 2/23/2006 11:05  
Date Received: 2/24/2006  
Solids: 73.43

Parameter	Method	RL	Result
		MG/KG	MG/KG
Oil & Grease	9071	39.7	<b>373</b>

**Comments:**

BQL = Below Quantitation Limit  
All soils are corrected for percent solids.

Reviewed By: PL  
9071\_LIMS\_v1.35

Client Sample ID: Batch QC  
 Lab Sample ID: G106-565-6J  
 Batch ID: 4612

Analyzed by: nio  
 Matrix: Soil  
 Solids: 73.43

## MS/MSD

Analyte	Sample MG/KG	Spiked MG/KG	MS MG/KG	REC %	Spiked MG/KG	MSD MG/KG	REC %	RPD %
Oil & Grease	373	421	282	N/A	415	365	N/A	166

## LCS

Analyte		Spiked MG/KG	Result MG/KG	REC %	Limits	
					Lower	Upper
Oil & Grease		312	228	73	70	130

## Prep Blank

Analyte	Sample MG/KG
Oil & Grease	BQL

**VPH (Aliphatics/Aromatics) Laboratory Reporting Form**Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 163 GP2-10
Sample Matrix	Soil
Collection Option (for Soil)*	2
Date Collected	02/22/06
Date Received	02/24/06
Date Extracted	02/22/06
Date Analyzed	02/27/06
Dry Weight	77
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	12 (mg/Kg)
C <sub>9</sub> -C <sub>10</sub> Aromatics**	10 (mg/Kg)
Surrogate % Recovery - PID	100
Surrogate % Recovery - FID	100

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-565-9d

Reviewed By: Bvd

**VPH (Aliphatics/Aromatics) Laboratory Reporting Form**Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 199A GP2-10
Sample Matrix	Soil
Collection Option (for Soil)*	2
Date Collected	02/22/06
Date Received	02/24/06
Date Extracted	02/22/06
Date Analyzed	02/27/06
Dry Weight	91
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>10</sub> Aromatics**	< 10 (mg/Kg)
Surrogate % Recovery - PID	96
Surrogate % Recovery - FID	94

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-565-14d

Reviewed By: RW

**VPH (Aliphatics/Aromatics) Laboratory Reporting Form**Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 89 HA6-8
Sample Matrix	Soil
Collection Option (for Soil)*	2
Date Collected	02/20/06
Date Received	02/24/06
Date Extracted	02/20/06
Date Analyzed	02/24/06
Dry Weight	77
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>10</sub> Aromatics**	< 10 (mg/Kg)
Surrogate % Recovery - PID	96
Surrogate % Recovery - FID	93

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-565-17g

Reviewed By: RvP

**VPH (Aliphatics/Aromatics) Laboratory Reporting Form**Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 221 GP1-8
Sample Matrix	Soil
Collection Option (for Soil)*	2
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/23/06
Date Analyzed	02/27/06
Dry Weight	75
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	< 10 (mg/Kg)
C <sub>9</sub> -C <sub>10</sub> Aromatics**	< 10 (mg/Kg)
Surrogate % Recovery - PID	98
Surrogate % Recovery - FID	95

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-565-24c

Reviewed By: DR

Attachment 2  
VPH Laboratory Reporting Form

<b>Calibration and QA/QC Information</b>
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FID Initial Calibration Date:	<u>02/11/06</u>	PID Initial Calibration Date:	<u>02/11/06</u>
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**Calibration Ranges and Limits**

Range	MDL (07/15/2004) ( $\mu\text{g/L}$ )	ML ( $\mu\text{g/L}$ )	RL ( $\mu\text{g/L}$ )	RL (mg/Kg)
C <sub>5</sub> -C <sub>8</sub> Aliphatics	4.4	14	100	10
C <sub>9</sub> -C <sub>12</sub> Aliphatics	3.4	11	100	10
C <sub>9</sub> -C <sub>10</sub> Aromatics	0.13	0.41	100	10

**Calibration Concentration Levels**

Range	Levels ( $\mu\text{g/L}$ )	%RSD or CCC	Method of Quantitation
C <sub>5</sub> -C <sub>8</sub> Aliphatics	40	10.8	Calibration Factor
	1000		
	2000		
	3000		
	4000		
C <sub>9</sub> -C <sub>12</sub> Aliphatics	10	0.99	Linear Regression
	250		
	500		
	750		
	1000		
C <sub>9</sub> -C <sub>10</sub> Aromatics	10	19.30	Calibration Factor
	250		
	500		
	750		
	1000		

Calibration Check Date: 02/24/06

**Calibration Check**

Range	Levels ( $\mu\text{g/L}$ )	RPD
C <sub>5</sub> -C <sub>8</sub> Aliphatics	2000      200	0.4
C <sub>9</sub> -C <sub>12</sub> Aliphatics	500      50	-9.2
C <sub>9</sub> -C <sub>10</sub> Aromatics	500      50	-8.2

MDL = Method Detection Limit

RPD = Relative Percent Difference

ML = Minimum Limit

%RSD = Percent Relative Standard Deviation

RL = Reportable Limit

CCC = Correlation Coefficient of Curve

Attachment 2  
VPH Laboratory Reporting Form

<b>Calibration and QA/QC Information</b>
--

FID Initial Calibration Date: 02/11/06PID Initial Calibration Date: 02/11/06**Calibration Ranges and Limits**

Range	MDL (07/15/2004) ( $\mu\text{g/L}$ )	ML ( $\mu\text{g/L}$ )	RL ( $\mu\text{g/L}$ )	RL (mg/Kg)
C <sub>5</sub> -C <sub>8</sub> Aliphatics	4.4	14	100	10
C <sub>9</sub> -C <sub>12</sub> Aliphatics	3.4	11	100	10
C <sub>9</sub> -C <sub>10</sub> Aromatics	0.13	0.41	100	10

**Calibration Concentration Levels**

Range	Levels ( $\mu\text{g/L}$ )	%RSD or CCC	Method of Quantitation
C <sub>5</sub> -C <sub>8</sub> Aliphatics	40	10.8	Calibration Factor
	1000		
	2000		
	3000		
	4000		
C <sub>9</sub> -C <sub>12</sub> Aliphatics	10	0.99	Linear Regression
	250		
	500		
	750		
	1000		
C <sub>9</sub> -C <sub>10</sub> Aromatics	10	19.30	Calibration Factor
	250		
	500		
	750		
	1000		

Calibration Check Date: 02/27/06**Calibration Check**

Range	Levels ( $\mu\text{g/L}$ )	RPD
C <sub>5</sub> -C <sub>8</sub> Aliphatics	2000 200	3.5
C <sub>9</sub> -C <sub>12</sub> Aliphatics	500 50	-3.4
C <sub>9</sub> -C <sub>10</sub> Aromatics	500 50	-2.9

MDL = Method Detection Limit

RPD = Relative Percent Difference

ML = Minimum Limit

%RSD = Percent Relative Standard Deviation

RL = Reportable Limit

CCC = Correlation Coefficient of Curve

**EPH (Aliphatics/Aromatics) Results**

by MDEP-EPH

Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 163 GP2-10
Sample Matrix	Soil
Date Collected	02/22/06
Date Received	02/24/06
Date Extracted	02/27/06
Date Analyzed	03/06/06
Dry Weight	77.2
Dilution Factor	1:1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	160 (mg/Kg)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	20 (mg/Kg)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	33 (mg/Kg)
Aliphatic Surrogate % Recovery	88
Aromatic Surrogate % Recovery	85
Fractionation Surrogate 1 % Recovery	85

**Comments:**

\* = Excludes any surrogates or internal standards.

Lab info: G106-565-9J

Reviewed By: Paul

**EPH (Aliphatics/Aromatics) Results**

by MDEP-EPH

Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 199A GP2-10
Sample Matrix	Soil
Date Collected	02/22/06
Date Received	02/24/06
Date Extracted	02/27/06
Date Analyzed	02/28/06
Dry Weight	90.9
Dilution Factor	1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	< 10 (mg/Kg)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	< 10 (mg/Kg)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	< 10 (mg/Kg)
Aliphatic Surrogate % Recovery	96
Aromatic Surrogate % Recovery	96

**Comments:**

\* = Excludes any surrogates or internal standards.

Sample did not require fractionation.

Lab info: G106-565-14M

Reviewed By: 12-1

**EPH (Aliphatics/Aromatics) Results**

by MDEP-EPH

Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 89 HA6-8
Sample Matrix	Soil
Date Collected	02/20/06
Date Received	02/24/06
Date Extracted	02/27/06
Date Analyzed	02/28/06
Dry Weight	77.2
Dilution Factor	1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	< 10 (mg/Kg)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	< 10 (mg/Kg)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	< 10 (mg/Kg)
Aliphatic Surrogate % Recovery	95
Aromatic Surrogate % Recovery	92

**Comments:**

\* = Excludes any surrogates or internal standards.

Sample did not require fractionation.

Lab info: G106-565-17J

Reviewed By: PV

**EPH (Aliphatics/Aromatics) Results**

by MDEP-EPH

Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 221 GP1-8
Sample Matrix	Soil
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/27/06
Date Analyzed	02/28/06
Dry Weight	75.2
Dilution Factor	1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	< 10 (mg/Kg)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	< 10 (mg/Kg)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	< 10 (mg/Kg)
Aliphatic Surrogate % Recovery	94
Aromatic Surrogate % Recovery	92

**Comments:**

\* = Excludes any surrogates or internal standards.

Sample did not require fractionation.

Lab info: G106-565-24F

Reviewed By: DWJ

## Attachment 3

## EPH Laboratory Reporting Form

## Calibration and QA/QC Information

Initial Calibration Date: 12/28/05

## Calibration Ranges and Limits

Range	MDL (2/2004) ( $\mu\text{g}/\text{L}$ )	ML ( $\mu\text{g}/\text{L}$ )	RL ( $\mu\text{g}/\text{L}$ )	RL (mg/Kg)
C <sub>9</sub> -C <sub>18</sub> Aliphatics	3.84	12.2	100	10
C <sub>19</sub> -C <sub>36</sub> Aliphatics	0.57	1.8	100	10
C <sub>11</sub> -C <sub>22</sub> Aromatics	4.54	14.4	100	10

## Calibration Concentration Levels

Range	Levels ( $\mu\text{g}/\text{mL}$ )	%RSD or CCC	Method of Quantitation
C <sub>9</sub> -C <sub>18</sub> Aliphatics	6	24.90	Calibration Factor
	30		
	60		
	120		
	240		
C <sub>19</sub> -C <sub>36</sub> Aliphatics	8	15.4	Calibration Factor
	40		
	80		
	160		
	320		
C <sub>11</sub> -C <sub>22</sub> Aromatics	17	9.8	Calibration Factor
	85		
	170		
	340		
	680		

Calibration Check Date: 03/06/06

## Calibration Check

Range	Levels ( $\mu\text{g}/\text{mL}$ )	RPD
C <sub>9</sub> -C <sub>18</sub> Aliphatics	120	12.7
C <sub>19</sub> -C <sub>36</sub> Aliphatics	160	6.7
C <sub>11</sub> -C <sub>22</sub> Aromatics	340	12.9

MDL = Method Detection Limit

RPD = Relative Percent Difference

ML = Minimum Limit

%RSD = Percent Relative Standard Deviation

RL = Reportable Limit

CCC = Correlation Coefficient of Curve

Attachment 3

## EPH Laboratory Reporting Form

## Calibration and QA/QC Information

Initial Calibration Date: 12/28/05

## Calibration Ranges and Limits

Range	MDL (2/2004) ( $\mu\text{g}/\text{L}$ )	ML ( $\mu\text{g}/\text{L}$ )	RL ( $\mu\text{g}/\text{L}$ )	RL ( $\text{mg}/\text{Kg}$ )
C <sub>9</sub> -C <sub>18</sub> Aliphatics	3.84	12.2	100	10
C <sub>19</sub> -C <sub>36</sub> Aliphatics	0.57	1.8	100	10
C <sub>11</sub> -C <sub>22</sub> Aromatics	4.54	14.4	100	10

## Calibration Concentration Levels

Range	Levels ( $\mu\text{g}/\text{mL}$ )	%RSD or CCC	Method of Quantitation
C <sub>9</sub> -C <sub>18</sub> Aliphatics	6	24.90	Calibration Factor
	30		
	60		
	120		
	240		
C <sub>19</sub> -C <sub>36</sub> Aliphatics	8	15.4	Calibration Factor
	40		
	80		
	160		
	320		
C <sub>11</sub> -C <sub>22</sub> Aromatics	17	9.8	Calibration Factor
	85		
	170		
	340		
	680		

Calibration Check Date: 02/28/06

## Calibration Check

Range	Levels ( $\mu\text{g}/\text{mL}$ )	RPD
C <sub>9</sub> -C <sub>18</sub> Aliphatics	120	17.0
C <sub>19</sub> -C <sub>36</sub> Aliphatics	160	10.3
C <sub>11</sub> -C <sub>22</sub> Aromatics	340	10.7

MDL = Method Detection Limit

ML = Minimum Limit

RL = Reportable Limit

RPD = Relative Percent Difference

%RSD = Percent Relative Standard Deviation

CCC = Correlation Coefficient of Curve

**Results for Metals**

Client Sample ID: PAR 206 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-1  
 Lab Project ID: G106-565  
 Batch ID: 4586 4592  
 Report Basis: Dry

Analyzed By: PSW  
 Date Collected: 2/22/2006 16:00  
 Date Received: 2/24/2006  
 Matrix: SOIL  
 Solids 78.08

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	BQL	7.68	0.553	1	MG/KG	6010B	2/28/2006	B
Arsenic	3.72	1.28	0.541	1	MG/KG	6010B	2/28/2006	
Beryllium	1.41	1.28	0.0152	1	MG/KG	6010B	2/28/2006	
Cadmium	0.756	1.28	0.0215	1	MG/KG	6010B	2/28/2006	
Chromium	53.8	1.28	0.0608	1	MG/KG	6010B	2/28/2006	
Copper	45.1	2.56	0.0569	1	MG/KG	6010B	2/28/2006	B
Lead	53.6	1.28	0.150	1	MG/KG	6010B	2/28/2006	B
Mercury	0.0291	0.0235	0.00411	1	MG/KG	7471	2/28/2006	
Nickel	23.2	5.12	0.0383	1	MG/KG	6010B	2/28/2006	B
Selenium	BQL	2.56	0.566	1	MG/KG	6010B	2/28/2006	
Silver	BQL	1.28	0.0512	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.28	0.585	1	MG/KG	6010B	2/28/2006	
Zinc	77.0	2.56	0.223	1	MG/KG	6010B	2/28/2006	B

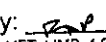
**Comments**

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; MDL

Reviewed By:   
MET\_LIMS\_40

## Results for Metals

Client Sample ID: PAR 206 HA1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-2  
 Lab Project ID: G106-565  
 Batch ID: 4586 4592  
 Report Basis: Dry

Analyzed By: PSW  
 Date Collected: 2/22/2006 17:00  
 Date Received: 2/24/2006  
 Matrix: SOIL  
 Solids 75.83

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	BQL	7.33	0.527	1	MG/KG	6010B	2/28/2006	B
Arsenic	3.89	1.22	0.515	1	MG/KG	6010B	2/28/2006	
Beryllium	1.83	1.22	0.0145	1	MG/KG	6010B	2/28/2006	
Cadmium	0.884	1.22	0.0205	1	MG/KG	6010B	2/28/2006	JB
Chromium	58.4	1.22	0.0580	1	MG/KG	6010B	2/28/2006	
Copper	57.5	2.44	0.0542	1	MG/KG	6010B	2/28/2006	B
Lead	32.0	1.22	0.143	1	MG/KG	6010B	2/28/2006	B
Mercury	0.0829	0.0225	0.00394	1	MG/KG	7471	2/28/2006	
Nickel	20.9	4.88	0.0365	1	MG/KG	6010B	2/28/2006	B
Selenium	1.57	2.44	0.540	1	MG/KG	6010B	2/28/2006	
Silver	BQL	1.22	0.0488	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.22	0.558	1	MG/KG	6010B	2/28/2006	
Zinc	158	2.44	0.212	1	MG/KG	6010B	2/28/2006	B

## Comments

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; MDL

Reviewed By: PW  
MET LIMS 4.0

**Results for Metals**

Client Sample ID: PAR 206 HA1A  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-3  
 Lab Project ID: G106-565  
 Batch ID: 4586 4592  
 Report Basis: Dry

Analyzed By: PSW  
 Date Collected: 2/23/2006 09:50  
 Date Received: 2/24/2006  
 Matrix: SOIL  
 Solids 76.88

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	BQL	7.36	0.530	1	MG/KG	6010B	2/28/2006	B
Arsenic	4.95	1.23	0.518	1	MG/KG	6010B	2/28/2006	
Beryllium	1.09	1.23	0.0146	1	MG/KG	6010B	2/28/2006	J
Cadmium	0.216	1.23	0.0206	1	MG/KG	6010B	2/28/2006	JB
Chromium	52.4	1.23	0.0583	1	MG/KG	6010B	2/28/2006	
Copper	59.3	2.45	0.0545	1	MG/KG	6010B	2/28/2006	B
Lead	22.7	1.23	0.144	1	MG/KG	6010B	2/28/2006	B
Mercury	0.0206	0.0248	0.00434	1	MG/KG	7471	2/28/2006	J
Nickel	23.1	4.91	0.0367	1	MG/KG	6010B	2/28/2006	B
Selenium	BQL	2.45	0.542	1	MG/KG	6010B	2/28/2006	
Silver	BQL	1.23	0.0491	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.23	0.561	1	MG/KG	6010B	2/28/2006	
Zinc	75.2	2.45	0.213	1	MG/KG	6010B	2/28/2006	B

**Comments**

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; MDL

Reviewed By: *[Signature]*  
MET\_LIMS 4.0

**Results for Metals**

Client Sample ID:	PAR 206 HA2	Analyzed By:	PSW
Client Project ID:	NCDOT-Yancey	Date Collected:	2/23/2006 10:20
Lab Sample ID:	G106-565-4	Date Received:	2/24/2006
Lab Project ID:	G106-565	Matrix:	SOIL
Batch ID:	4586 4592	Solids	77.25
Report Basis:	Dry		

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	BQL	6.58	0.474	1	MG/KG	6010B	2/28/2006	B
Arsenic	3.51	1.10	0.463	1	MG/KG	6010B	2/28/2006	
Beryllium	1.80	1.10	0.0131	1	MG/KG	6010B	2/28/2006	
Cadmium	2.22	1.10	0.0184	1	MG/KG	6010B	2/28/2006	
Chromium	53.3	1.10	0.0521	1	MG/KG	6010B	2/28/2006	
Copper	29.8	2.19	0.0487	1	MG/KG	6010B	2/28/2006	
Lead	24.6	1.10	0.128	1	MG/KG	6010B	2/28/2006	
Mercury	0.123	0.0247	0.00431	1	MG/KG	7471	2/28/2006	
Nickel	22.6	4.39	0.0328	1	MG/KG	6010B	2/28/2006	
Selenium	BQL	2.19	0.485	1	MG/KG	6010B	2/28/2006	
Silver	BQL	1.10	0.0439	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.10	0.501	1	MG/KG	6010B	2/28/2006	
Zinc	150	2.19	0.191	1	MG/KG	6010B	2/28/2006	

**Comments**

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; MDL

Reviewed By: *[Signature]*  
MET\_LIMS\_4.0

**Results for Metals**

Client Sample ID:	PAR 206 HA3	Analyzed By:	PSW
Client Project ID:	NCDOT-Yancey	Date Collected:	2/23/2006 11:00
Lab Sample ID:	G106-565-5	Date Received:	2/24/2006
Lab Project ID:	G106-565	Matrix:	SOIL
Batch ID:	4586 4592	Solids	70.87
Report Basis:	Dry		

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	BQL	8.30	0.597	1	MG/KG	6010B	2/28/2006	B
Arsenic	4.54	1.38	0.584	1	MG/KG	6010B	2/28/2006	
Beryllium	2.57	1.38	0.0165	1	MG/KG	6010B	2/28/2006	
Cadmium	0.609	1.38	0.0232	1	MG/KG	6010B	2/28/2006	JB
Chromium	92.8	1.38	0.0657	1	MG/KG	6010B	2/28/2006	
Copper	48.9	2.77	0.0614	1	MG/KG	6010B	2/28/2006	B
Lead	12.2	1.38	0.162	1	MG/KG	6010B	2/28/2006	B
Mercury	0.0469	0.0264	0.00462	1	MG/KG	7471	2/28/2006	
Nickel	36.2	5.53	0.0414	1	MG/KG	6010B	2/28/2006	B
Selenium	BQL	2.77	0.611	1	MG/KG	6010B	2/28/2006	B
Silver	BQL	1.38	0.0553	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.38	0.632	1	MG/KG	6010B	2/28/2006	
Zinc	160	2.77	0.240	1	MG/KG	6010B	2/28/2006	B

**Comments**

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; MDL

Reviewed By: PM  
MET\_LIMS\_4.0

**Results for Metals**

Client Sample ID: PAR 206 HA4  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-565-6  
 Lab Project ID: G106-565  
 Batch ID: 4586 4592  
 Report Basis: Dry

Analyzed By: PSW  
 Date Collected: 2/23/2006 11:05  
 Date Received: 2/24/2006  
 Matrix: SOIL  
 Solids: 73.43

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	BQL	6.92	0.498	1	MG/KG	6010B	2/28/2006	B
Arsenic	3.84	1.15	0.487	1	MG/KG	6010B	2/28/2006	
Beryllium	1.41	1.15	0.0137	1	MG/KG	6010B	2/28/2006	
Cadmium	1.68	1.15	0.0194	1	MG/KG	6010B	2/28/2006	
Chromium	53.4	1.15	0.0548	1	MG/KG	6010B	2/28/2006	
Copper	61.6	2.31	0.0512	1	MG/KG	6010B	2/28/2006	B
Lead	44.3	1.15	0.135	1	MG/KG	6010B	2/28/2006	
Mercury	0.102	0.0255	0.00445	1	MG/KG	7471	2/28/2006	
Nickel	23.2	4.62	0.0345	1	MG/KG	6010B	2/28/2006	
Selenium	0.945	2.31	0.510	1	MG/KG	6010B	2/28/2006	
Silver	BQL	1.15	0.0462	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.15	0.528	1	MG/KG	6010B	2/28/2006	
Zinc	171	2.31	0.201	1	MG/KG	6010B	2/28/2006	B

**Comments**

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; MDL

Reviewed By: EML  
MET\_LIMS\_4.0

**Results for Metals**

Client Sample ID:	Lab Blank	Analyzed By:	PSW
Client Project ID:		Date Collected:	
Lab Sample ID:	pb4586	Date Received:	
Lab Project ID:		Matrix:	SOIL
Batch ID:	4586	Solids	100.00
Report Basis:	Dry		

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Antimony	0.747	6.00	0.432	1	MG/KG	6010B	2/28/2006	JB
Arsenic	BQL	1.00	0.422	1	MG/KG	6010B	2/28/2006	
Beryllium	BQL	1.00	0.0119	1	MG/KG	6010B	2/28/2006	
Cadmium	0.0350	1.00	0.0168	1	MG/KG	6010B	2/28/2006	JB
Chromium	BQL	1.00	0.0475	1	MG/KG	6010B	2/28/2006	
Copper	0.287	2.00	0.0444	1	MG/KG	6010B	2/28/2006	JB
Lead	0.317	1.00	0.117	1	MG/KG	6010B	2/28/2006	JB
Nickel	0.349	4.00	0.0299	1	MG/KG	6010B	2/28/2006	JB
Selenium	0.648	2.00	0.442	1	MG/KG	6010B	2/28/2006	JB
Silver	BQL	1.00	0.0400	1	MG/KG	6010B	2/28/2006	
Thallium	BQL	1.00	0.457	1	MG/KG	6010B	2/28/2006	
Zinc	0.225	2.00	0.174	1	MG/KG	6010B	2/28/2006	JB

**Comments**

BQL = Below Quantitation Limits

DF = Dilution Factor

J = Between MDL and RL

B= Amount in Prep Blank &gt; RL

Reviewed By: P.W.  
PrepBlank

**Results for Metals**

Client Sample ID: Lab Blank                          Analyzed By: PSW  
Client Project ID:                                      Date Collected:  
Lab Sample ID: pb4592                                Date Received:  
Lab Project ID:                                         Matrix: SOIL  
Batch ID: 4592                                         Solids 100.00  
Report Basis: Dry

Metals	Result	RL	MDL	DF	Units	Method	Date Analyzed	Flags
Mercury	BQL	0.02	0.0035	1	MG/KG	7471	2/28/2006	

**Comments**

BQL = Below Quantitation Limits  
DF = Dilution Factor  
J = Between MDL and RL  
B= Amount in Prep Blank > RL

Reviewed By:   
PrepBlank

## METALS Results for LCS/LCD

ICP Batch: 4586

Matrix: SOIL

HG Batch: 4592

Units: MG/KG

Other:

Analyte	TRUE Value	LCS	LCS %REC	LCD	LCD %REC	Limit		RPD	RPD Limit
						Lower	Upper		
Antimony	40.0	38.1	95.3	39.2	98.0	80	120	2.85	20
Arsenic	40.0	35.9	89.8	38.1	95.3	80	120	5.95	20
Beryllium	40.0	36.1	90.2	38.3	95.8	80	120	5.91	20
Cadmium	40.0	35	87.5	36.3	90.7	80	120	3.65	20
Chromium	40.0	36.7	91.8	38.3	95.8	80	120	4.27	20
Copper	40.0	38	95.0	40.3	101	80	120	5.87	20
Lead	40.0	36.5	91.2	37.6	94.0	80	120	2.97	20
Mercury	0.500	0.483	96.6	0.484	96.8	80	120	0.207	20
Nickel	40.0	37.1	92.8	38.4	96.0	80	120	3.44	20
Selenium	40.0	32.8	82.0	34.5	86.2	80	120	5.05	20
Silver	40.0	35	87.5	36.0	90.0	80	120	2.82	20
Thallium	40.0	34	85.0	35.7	89.3	80	120	4.88	20
Zinc	40.0	35.4	88.5	36.6	91.5	80	120	3.33	20

Reviewed By: BW

## MS/MSD Results for METALS

Lab ID: G106-565-1

MS Lab ID: G106-565-1

MSD Lab ID: G106-565-1

ICP Batch: 4586

HG Batch: 4592

Other:

Analyzed By: PSW

Matrix: Soil

Units: MG/KG

Solids: 78.08

Analyte	Sample Result	SA MS	MS Result	MS %REC	SA MSD	MSD Result	MSD %REC	Limit		RPD	RPD Limit
								Lower	Upper		
Antimony	BQL	43.4	8.54	19.7	*	46.6	8.55	18.3	*	75	125
Arsenic	3.72	43.4	40.5	84.7		46.6	44.0	86.4		75	125
Beryllium	1.41	43.4	41.6	92.6		46.6	44.4	92.3		75	125
Cadmium	BQL	43.4	38.2	88.0		46.6	41.8	89.7		75	125
Chromium	53.8	43.4	103	113		46.6	102	103		75	125
Copper	45.1	43.4	94.8	114		46.6	92.6	102		75	125
Lead	53.6	43.4	90.1	84.1		46.6	137	179	*	75	125
Mercury	0.0291	0.572	0.661	111		0.562	0.593	100		75	125
Nickel	23.2	43.4	65.1	96.5		46.6	67.8	95.7		75	125
Selenium	BQL	43.4	30.9	71.2	*	46.6	33.7	72.3	*	75	125
Silver	BQL	43.4	36.8	84.8		46.6	40.6	87.1		75	125
Thallium	BQL	43.4	31.0	71.4	*	46.6	36.0	77.3		75	125
Zinc	77	43.4	120	99.0		46.6	131	116		75	125

## Comments

\*=Out of Limits

NA = Not applicable, due to sample concentration greater than three times spike concentration

Reviewed By: Rae

**List of Reporting Abbreviations  
and Data Qualifiers**

B = Compound also detected in batch blank

BQL = Below Quantitation Limit (RL or MDL)

DF = Dilution Factor

Dup = Duplicate

D = Detected, but RPD is > 40% between results in dual column method.

E = Estimated concentration, exceeds calibration range.

J = Estimated concentration, below calibration range and above MDL

LCS(D) = Laboratory Control Spike (Duplicate)

MDL = Method Detection Limit

MS(D) = Matrix Spike (Duplicate)

PQL = Practical Quantitation Limit

RL = Reporting Limit

RPD = Relative Percent Difference

mg/kg = milligram per kilogram, ppm, parts per million

ug/kg = micrograms per kilogram, ppb, parts per billion

mg/L = milligram per liter, ppm, parts per million

ug/L = micrograms per liter, ppb, parts per billion

% Rec = Percent Recovery

% soilds = Percent Solids

**Special Notes:**

- 1) Metals and mercury samples are digested with a hot block, see the standard operating procedure document for details.
- 2) Uncertainty for all reported data is less than or equal to 30 percent.



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PARADIGM ANALYTICAL LABORATORIES, INC.

<b>1</b> CLIENT: <u>BOB SHATI</u> CONTACT: <u>EL</u> PHONE NO.: <u>(919) 544-7500</u> PROJECT: <u>DOT - Hwy</u> SITE/PWSID#: <u>R-2519A</u> REPORTS TO: <u>BOB SHATI</u> INVOICE TO: <u>NC DOT</u> FAX NO.: <u>(919) 544-2199</u> QUOTE #: <u>NBS LT 35609.1.1</u> P.O. NUMBER					SGS Reference: <u>G106-565</u> PAGE <u>1</u> OF <u>1</u>													
<b>2</b>	LAB NO.	SAMPLE IDENTIFICATION	DATE	TIME	MATRIX	No C N O C T E R E N A G G	SAMPLE TYPE	Pesticides Used Analysts Required	REMARKS									
									<b>3</b>	<b>TPH - ORO</b>	<b>TPH - ORO</b>	<b>9071-00-0000</b>	<b>9071-00-0000</b>	<b>VOCs - 0260</b>	<b>VOCs - 0260</b>	<b>PP METALS</b>	<b>PP METALS</b>	<b>PP METALS</b>
<b>4</b>	Collected By: <u>J. D. Shat</u> Relinquished By: <u>J. D. Shat</u> Date: <u>7/23/96</u> Time: <u>1100</u> Received By: <u>John Shat</u> Shipping Carrier: <u></u> Shipping Ticket No: <u></u> Special Deliverable Requirements: <u></u> Chain of Custody Seal: (Circle) <u>INTACT</u> <u>BROKEN</u> <u>ABSENT</u> Requested Turnaround Time and Special Instructions: <u></u>					Samples Received Cold? (Circle) YES <u>NO</u> Temperature (C): <u>2, 3, 4, 7, 10</u>												
<b>5</b>	Relinquished By: (3) Date: <u>7/24/96</u> Time: <u>1000</u> Received By: <u>John Shat</u> Relinquished By: (4) Date: <u></u> Time: <u></u> Received By: <u></u>																	

N.C. CERTIFICATION #481

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PARADIGM ANALYTICAL LABORATORIES, INC.

1	CLIENT: <u>EZ</u>	PHONE NO.: <u>919) 544-7500</u>										
CONTACT: <u>BOB SHAW</u>	SITE/PSID#: <u>R-259A</u>											
PROJECT: <u>NC DOT - VANCEY</u>	REPORT TO: <u>BOB SHAW - EZ</u>	FAX NO.: <u>919) 544-2199</u>										
INVOICE TO: <u>NC DOT</u>	QUOTE #: <u>NBS# 25609.1.1</u>											
P.O. NUMBER												
LAB NO.	SAMPLE IDENTIFICATION		DATE	TIME	MATRIX	C O N T E R S	SAMPLE TYPE	Preservatives Used	Analysis Required	G C O M P A G R U P	REMARKS	
	<u>PAR 89 HA4-B</u>		<u>2-20-06</u>	<u>1120</u>	<u>Soil</u>		X	X	X		<u>TPH GRO</u>	
	<u>PAR 89 HA6-B</u>		<u>2-20-06</u>	<u>1245</u>			X	X	X		<u>TPH DRO</u>	
	<u>PAR 89 HA9-B</u>		<u>2-21-06</u>	<u>1206</u>			X	X	X		<u>VOCs-8260</u>	
	<u>PAR 89 HA10-B</u>		<u>2-21-06</u>	<u>1244</u>			X	X	X		<u>VOCs-8270</u>	
	<u>PAR 221 GP1-B</u>		<u>2-21-06</u>	<u>1515</u>			X	X	X		<u>MAAP VPH</u>	
	<u>PAR 221 GP2-B</u>		<u>2-21-06</u>	<u>1528</u>			X	X	X		<u>MAAP EPH</u>	
	<u>PAR 221 GP3-B</u>		<u>2-21-06</u>	<u>1545</u>								
	<u>PAR 221 GP4-B</u>		<u>2-24-06</u>	<u>1636</u>			X	X	X			
	<u>PAR 221 GP1-B</u>		<u>2-23-06</u>	<u>1200</u>								

5	Collected/Relinquished By: <u>(1)</u>	Date <u>1-23-06</u>	Time <u>1000</u>	Received By: <u>John Shaver</u>	Shipping Carrier:	Samples Received Cold? (Circle YES) <u>NO</u>
	Relinquished By: <u>(2)</u>				Shipping Ticket No:	Temperature (C): <u>2, 3°C</u> <u>2, 7°C</u>
	Relinquished By: <u>(3)</u>				Special Deliverable Requirements:	Chain of Custody Seal: (Circle)  <u>INTACT</u> <u>BROKEN</u> <u>ABSENT</u>
	Relinquished By: <u>(4)</u>				Requested Turnaround Time and Special Instructions:	

N.C. CERTIFICATION #481

□ 200 W. Price Ave., Anchorage, AK 99518 Tel: (907) 342-2313 Fax: (907) 544-5301  
 □ 6500 Balsam St., Suite 100, Wilmington, NC 28405 Tel: (910) 360-1603 Fax: (910) 360-1567

Yellow - Handwritten by Lab  
 Pink - Printed with Report  
 Red - Handwritten by Sampler

□ 1258 Greenbrier Street, Charleston, WV 25311 Tel: (304) 344-0725 Fax: (304) 344-0781



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PARADIGM ANALYTICAL LABORATORIES, INC.

						PAGE _____ OF _____
1	CLIENT: <u>ET</u>					SGS Reference: <u>G106 -565</u>
CONTACT:	<u>Bob Shatto</u>					PHONE NO.: <u>(919) 544-7500</u>
PROJECT:	<u>MCOT - VNCY</u>					SITE/PWSID: <u>R - 2519A</u>
REPORTS TO:	<u>Bob Shatt - ET</u>					FAX NO.: <u>(919) 544-2199</u>
INVOICE TO:						QUOTE # <u>NBS# 35609.1.1</u>
2	P.O. NUMBER	SAMPLE IDENTIFICATION	DATE	TIME	MATRIX	No
3	C= COMP	Preservative Used	C= GRAB	Analysis Required	REMARKS	SGS
4	Shipping Carrier:	Samples Received Cold? (Circle) <input checked="" type="checkbox"/> YES <input type="checkbox"/> NO	Temperature (C): <u>2-32</u>	Chain of Custody Seal: (Circle)	INTACT <input checked="" type="checkbox"/>	BROKEN <input type="checkbox"/>
5	Collected/Relinquished By: <u>J. H. H.</u>	Date <u>2/23/06</u>	Time <u>1005</u>	Received By: <u>John H. H.</u>	Special Deliverable Requirements:	ABSENT <input checked="" type="checkbox"/>
Relinquished By: (2)	Date <u>2/24/06</u>	Time <u>1005</u>	Received By: <u>John H. H.</u>	Requested Turnaround Time and Special Instructions:		
Relinquished By: (3)	Date	Time	Received By:			
Relinquished By: (4)	Date	Time	Received By:			

**SGS**

Mr. Bob Shaut  
Environmental Investigations  
2101 Gateway Centre Boulevard  
Suite 200  
Morrisville NC 27560  
Report Number: G106-566  
Client Project: NCDOT-Yancey

Dear Mr. Shaut:

Enclosed are the results of the analytical services performed under the referenced project. The samples are certified to meet the requirements of the National Environmental Laboratory Accreditation Conference Standards. Copies of this report and supporting data will be retained in our files for a period of five years in the event they are required for future reference. Any samples submitted to our laboratory will be retained for a maximum of thirty (30) days from the date of this report unless other arrangements are requested.

If there are any questions about the report or the services performed during this project, please call Paradigm at (910) 350-1903. We will be happy to answer any questions or concerns which you may have.

Thank you for using Paradigm Analytical Labs for your analytical services. We look forward to working with you again on any additional analytical needs which you may have.

Sincerely,  
Paradigm Analytical Laboratories, Inc.

*Jay P. Weaver*  
Laboratory Director  
J. Patrick Weaver

3/10/2006  
Date



## CASE NARRATIVE

Date: March 6, 2006

EI Project ID: NCDOT-Yancey  
SGS-Paradigm Analytical ID: G106-566

Five water samples were received at the laboratory February 24 for analysis as indicated on the chain of custody. The samples were received in good condition, within temperature and holding time limits.

All extractions and analyses were completed within holding.

All laboratory control spikes and matrix spikes recovered within control limits.

All blanks except one of the 6230D blanks were free of target compounds. Blank VBLK3022806b contained a detection of the target compound 1,2,3-trichlorobenzene at 0.738 ug/L. This detection was due to carry-over contamination from a preceding quality control check. The batch was accepted as the target was not detected in any sample.

**Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 199A GP2

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/22/2006 14:45

Lab Sample ID: G106-566-1A

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	BQL	0.500	0.161	1	3/1/2006	
Bromobenzene	BQL	0.500	0.929	1	3/1/2006	
Bromochloromethane	BQL	0.500	0.171	1	3/1/2006	
Bromodichloromethane	BQL	0.500	0.213	1	3/1/2006	
Bromoform	BQL	0.500	0.197	1	3/1/2006	
Bromomethane	BQL	0.500	0.405	1	3/1/2006	
n-Butylbenzene	BQL	0.500	0.376	1	3/1/2006	
sec-Butylbenzene	BQL	0.500	0.185	1	3/1/2006	
tert-Butylbenzene	BQL	0.500	0.175	1	3/1/2006	
Carbon tetrachloride	BQL	0.500	0.41	1	3/1/2006	
Chlorobenzene	BQL	0.500	0.177	1	3/1/2006	
Chloroethane	BQL	0.500	0.384	1	3/1/2006	
Chloroform	BQL	0.500	0.179	1	3/1/2006	
Chloromethane	BQL	0.500	0.38	1	3/1/2006	
2-Chlorotoluene	BQL	0.500	0.207	1	3/1/2006	
4-Chlorotoluene	BQL	0.500	0.377	1	3/1/2006	
Dibromochloromethane	BQL	0.500	0.177	1	3/1/2006	
1,2-Dibromo-3-chloropropane	BQL	0.500	0.233	1	3/1/2006	
1,2-Dibromoethane (EDB)	BQL	0.500	0.208	1	3/1/2006	
Dibromomethane	BQL	0.500	0.207	1	3/1/2006	
1,2-Dichlorobenzene	BQL	0.500	0.195	1	3/1/2006	
1,3-Dichlorobenzene	BQL	0.500	0.218	1	3/1/2006	
1,4-Dichlorobenzene	BQL	0.500	0.211	1	3/1/2006	
Dichlorodifluoromethane	BQL	0.500	0.407	1	3/1/2006	
1,1-Dichloroethane	BQL	0.500	0.177	1	3/1/2006	
1,2-Dichloroethane	BQL	0.500	0.19	1	3/1/2006	
1,1-Dichloroethene	BQL	0.500	0.14	1	3/1/2006	
cis-1,2-Dichloroethene	BQL	0.500	0.423	1	3/1/2006	
trans-1,2-Dichloroethene	BQL	0.500	0.186	1	3/1/2006	
1,2-Dichloropropane	BQL	0.500	0.187	1	3/1/2006	
2,2-Dichloropropane	BQL	0.500	0.423	1	3/1/2006	
cis-1,3-Dichloropropene	BQL	0.500	0.21	1	3/1/2006	
trans-1,3-Dichloropropene	BQL	0.500	0.205	1	3/1/2006	
Diisopropyl ether (DIPE)	BQL	0.500	0.176	1	3/1/2006	
Ethylbenzene	BQL	0.500	0.166	1	3/1/2006	
Hexachlorobutadiene	BQL	0.500	0.188	1	3/1/2006	
Isopropylbenzene	BQL	0.500	0.18	1	3/1/2006	
p-Isopropyltoluene	BQL	0.500	0.383	1	3/1/2006	
Methyl-tert butyl ether (MTBE)	BQL	0.500	0.347	1	3/1/2006	
Methylene Chloride	BQL	5.00	0.464	1	3/1/2006	

Reviewed By: MM  
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N.C. CERTIFICATION #481

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## Results for Volatiles

by GC 6230D

Client Sample ID: PAR 199A GP2

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/22/2006 14:45

Lab Sample ID: G106-566-1A

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	BQL	0.500	0.2	1	3/1/2006	
n-Propylbenzene	BQL	0.500	0.176	1	3/1/2006	
Styrene	BQL	1.00	0.348	1	3/1/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	3/1/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	3/1/2006	
Tetrachloroethene	BQL	0.500	0.445	1	3/1/2006	
Toluene	BQL	0.500	0.187	1	3/1/2006	
1,2,3-Trichlorobenzene	BQL	0.500	0.237	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	3/1/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	3/1/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	3/1/2006	
Trichloroethene	BQL	0.500	0.266	1	3/1/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	3/1/2006	
1,2,3-Trichloropropane	BQL	0.500	0.929	1	3/1/2006	
1,2,4-Trimethylbenzene	BQL	0.500	0.174	1	3/1/2006	
1,3,5-Trimethylbenzene	BQL	0.500	0.374	1	3/1/2006	
Vinyl Chloride	BQL	1.00	0.424	1	3/1/2006	
m/p-Xylene	BQL	1.00	0.35	1	3/1/2006	
o-Xylene	BQL	1.00	0.348	1	3/1/2006	
Surrogate Spike Recoveries		Spike Added		Spike Result	Percent Recovery	
Trifluorotoluene		40		39.9	99.7	
1,4-Dichlorobutane		40		38.1	95.2	

## Comments:

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: ony  
GC\_LIMS\_v2.0

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**Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 206 GP1

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 9:30

Lab Sample ID: G106-566-2A

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	BQL	0.500	0.161	1	3/1/2006	
Bromobenzene	BQL	0.500	0.929	1	3/1/2006	
Bromochloromethane	BQL	0.500	0.171	1	3/1/2006	
Bromodichloromethane	BQL	0.500	0.213	1	3/1/2006	
Bromoform	BQL	0.500	0.197	1	3/1/2006	
Bromomethane	BQL	0.500	0.405	1	3/1/2006	
n-Butylbenzene	BQL	0.500	0.376	1	3/1/2006	
sec-Butylbenzene	BQL	0.500	0.185	1	3/1/2006	
tert-Butylbenzene	BQL	0.500	0.175	1	3/1/2006	
Carbon tetrachloride	BQL	0.500	0.41	1	3/1/2006	
Chlorobenzene	BQL	0.500	0.177	1	3/1/2006	
Chloroethane	BQL	0.500	0.384	1	3/1/2006	
Chloroform	BQL	0.500	0.179	1	3/1/2006	
Chloromethane	BQL	0.500	0.38	1	3/1/2006	
2-Chlorotoluene	BQL	0.500	0.207	1	3/1/2006	
4-Chlorotoluene	BQL	0.500	0.377	1	3/1/2006	
Dibromochloromethane	BQL	0.500	0.177	1	3/1/2006	
1,2-Dibromo-3-chloropropane	BQL	0.500	0.233	1	3/1/2006	
1,2-Dibromoethane (EDB)	BQL	0.500	0.208	1	3/1/2006	
Dibromomethane	BQL	0.500	0.207	1	3/1/2006	
1,2-Dichlorobenzene	BQL	0.500	0.195	1	3/1/2006	
1,3-Dichlorobenzene	BQL	0.500	0.218	1	3/1/2006	
1,4-Dichlorobenzene	BQL	0.500	0.211	1	3/1/2006	
Dichlorodifluoromethane	BQL	0.500	0.407	1	3/1/2006	
1,1-Dichloroethane	BQL	0.500	0.177	1	3/1/2006	
1,2-Dichloroethane	BQL	0.500	0.19	1	3/1/2006	
1,1-Dichloroethene	BQL	0.500	0.14	1	3/1/2006	
cis-1,2-Dichloroethene	BQL	0.500	0.423	1	3/1/2006	
trans-1,2-Dichloroethene	BQL	0.500	0.186	1	3/1/2006	
1,2-Dichloropropane	BQL	0.500	0.187	1	3/1/2006	
2,2-Dichloropropane	BQL	0.500	0.423	1	3/1/2006	
cis-1,3-Dichloropropene	BQL	0.500	0.21	1	3/1/2006	
trans-1,3-Dichloropropene	BQL	0.500	0.205	1	3/1/2006	
Diisopropyl ether (DIPE)	BQL	0.500	0.176	1	3/1/2006	
Ethylbenzene	BQL	0.500	0.166	1	3/1/2006	
Hexachlorobutadiene	BQL	0.500	0.188	1	3/1/2006	
Isopropylbenzene	BQL	0.500	0.18	1	3/1/2006	
p-Isopropyltoluene	BQL	0.500	0.383	1	3/1/2006	
Methyl-tert butyl ether (MTBE)	BQL	0.500	0.347	1	3/1/2006	
Methylene Chloride	BQL	5.00	0.464	1	3/1/2006	

Reviewed By: 34  
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**Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 206 GP1

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 9:30

Lab Sample ID: G106-566-2A

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	BQL	0.500	0.2	1	3/1/2006	
n-Propylbenzene	BQL	0.500	0.176	1	3/1/2006	
Styrene	BQL	1.00	0.348	1	3/1/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	3/1/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	3/1/2006	
Tetrachloroethene	BQL	0.500	0.445	1	3/1/2006	
Toluene	BQL	0.500	0.187	1	3/1/2006	
1,2,3-Trichlorobenzene	BQL	0.500	0.237	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	3/1/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	3/1/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	3/1/2006	
Trichloroethene	BQL	0.500	0.266	1	3/1/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	3/1/2006	
1,2,3-Trichloroproppane	BQL	0.500	0.929	1	3/1/2006	
1,2,4-Trimethylbenzene	BQL	0.500	0.174	1	3/1/2006	
1,3,5-Trimethylbenzene	BQL	0.500	0.374	1	3/1/2006	
Vinyl Chloride	BQL	0.500	0.424	1	3/1/2006	
m/p-Xylene	BQL	1.00	0.35	1	3/1/2006	
o-Xylene	BQL	1.00	0.348	1	3/1/2006	
<b>Surrogate Spike Recoveries</b>		Spike Added	Spike Result	Percent Recovery		
Trifluorotoluene		40	39.9	99.7		
1,4-Dichlorobutane		40	38.5	96.1		

**Comments:**

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: pmf  
GC\_LIMS\_V2.0

**Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 221 GP1

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 11:30

Lab Sample ID: G106-566-3A

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	3.32	2.00	0.644	4	3/2/2006	
Bromobenzene	BQL	0.500	0.929	1	3/1/2006	
Bromo-chloromethane	BQL	0.500	0.171	1	3/1/2006	
Bromo-dichloromethane	BQL	0.500	0.213	1	3/1/2006	
Bromoform	BQL	0.500	0.197	1	3/1/2006	
Bromo-methane	BQL	0.500	0.405	1	3/1/2006	
n-Butylbenzene	BQL	2.00	1.5	4	3/2/2006	
sec-Butylbenzene	6.17	2.00	0.741	4	3/2/2006	
tert-Butylbenzene	BQL	2.00	0.7	4	3/2/2006	
Carbon tetrachloride	BQL	0.500	0.41	1	3/1/2006	
Chloro-benzene	BQL	0.500	0.177	1	3/1/2006	
Chloro-ethane	BQL	0.500	0.384	1	3/1/2006	
Chloroform	BQL	0.500	0.179	1	3/1/2006	
Chloro-methane	BQL	0.500	0.38	1	3/1/2006	
2-Chloro-toluene	BQL	0.500	0.207	1	3/1/2006	
4-Chloro-toluene	BQL	0.500	0.377	1	3/1/2006	
Dibromo-chloro-methane	BQL	0.500	0.177	1	3/1/2006	
1,2-Dibromo-3-chloro-propane	BQL	0.500	0.233	1	3/1/2006	
1,2-Dibromo-ethane (EDB)	BQL	0.500	0.208	1	3/1/2006	
Dibromo-methane	BQL	0.500	0.207	1	3/1/2006	
1,2-Dichloro-benzene	BQL	0.500	0.195	1	3/1/2006	
1,3-Dichloro-benzene	BQL	0.500	0.218	1	3/1/2006	
1,4-Dichloro-benzene	BQL	0.500	0.211	1	3/1/2006	
Dichloro-di-fluoro-methane	BQL	0.500	0.407	1	3/1/2006	
1,1-Dichloro-ethane	BQL	0.500	0.177	1	3/1/2006	
1,2-Dichloro-ethane	BQL	0.500	0.19	1	3/1/2006	
1,1-Dichloro-ethene	BQL	0.500	0.14	1	3/1/2006	
cis-1,2-Dichloro-ethene	BQL	0.500	0.423	1	3/1/2006	
trans-1,2-Dichloro-ethene	BQL	0.500	0.186	1	3/1/2006	
1,2-Dichloro-propane	BQL	0.500	0.187	1	3/1/2006	
2,2-Dichloro-propane	BQL	0.500	0.423	1	3/1/2006	
cis-1,3-Dichloro-propene	BQL	0.500	0.21	1	3/1/2006	
trans-1,3-Dichloro-propene	BQL	0.500	0.205	1	3/1/2006	
Di-isopropyl ether (DIPE)	BQL	2.00	0.703	4	3/2/2006	
Ethyl-benzene	11.4	2.00	0.662	4	3/2/2006	
Hexa-chloro-butadiene	BQL	0.500	0.188	1	3/1/2006	
Isopropyl-benzene	5.27	2.00	0.718	4	3/2/2006	
p-Isopropyl-toluene	BQL	2.00	1.53	4	3/2/2006	
Methyl-tert-butyl-ether (MTBE)	BQL	2.00	1.39	4	3/2/2006	
Methylene Chloride	BQL	5.00	0.464	1	3/1/2006	

Reviewed By: P.J.S.  
GC\_LIMS\_V2.0

**SGS****Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 221 GP1

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 11:30

Lab Sample ID: G106-566-3A

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	40.7	2.00	0.8	4	3/2/2006	
n-Propylbenzene	6.10	2.00	0.702	4	3/2/2006	
Styrene	BQL	4.00	1.39	4	3/2/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	3/1/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	3/1/2006	
Tetrachloroethene	BQL	0.500	0.445	1	3/1/2006	
Toluene	13.7	2.00	0.748	4	3/2/2006	
1,2,3-Trichlorobenzene	BQL	0.500	0.237	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	3/1/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	3/1/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	3/1/2006	
Trichloroethene	BQL	0.500	0.266	1	3/1/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	3/1/2006	
1,2,3-Trichloropropane	BQL	0.500	0.929	1	3/1/2006	
1,2,4-Trimethylbenzene	47.5	2.00	0.695	4	3/2/2006	
1,3,5-Trimethylbenzene	6.50	2.00	1.5	4	3/2/2006	
Vinyl Chloride	BQL	0.500	0.424	1	3/1/2006	
m/p-Xylene	30.1	4.00	1.4	4	3/2/2006	
o-Xylene	24.3	4.00	1.39	4	3/2/2006	
<b>Surrogate Spike Recoveries</b>		<b>Spike Added</b>	<b>Spike Result</b>	<b>Percent Recovery</b>		
Trifluorotoluene		40	40.9	102		
1,4-Dichlorobutane		40	39.6	99		

**Comments:**

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: PJW  
 GC\_LIMS v2.0  
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**Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 163 GP2

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 9:00

Lab Sample ID: G106-566-4B

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	BQL	0.500	0.161	1	3/2/2006	
Bromobenzene	BQL	0.500	0.929	1	3/2/2006	
Bromochloromethane	BQL	0.500	0.171	1	3/2/2006	
Bromodichloromethane	BQL	0.500	0.213	1	3/2/2006	
Bromoform	BQL	0.500	0.197	1	3/2/2006	
Bromomethane	BQL	0.500	0.405	1	3/2/2006	
n-Butylbenzene	BQL	0.500	0.376	1	3/2/2006	
sec-Butylbenzene	0.322	0.500	0.185	1	3/2/2006	J
tert-Butylbenzene	BQL	0.500	0.175	1	3/2/2006	
Carbon tetrachloride	BQL	0.500	0.41	1	3/2/2006	
Chlorobenzene	BQL	0.500	0.177	1	3/2/2006	
Chloroethane	BQL	0.500	0.384	1	3/2/2006	
Chloroform	BQL	0.500	0.179	1	3/2/2006	
Chloromethane	BQL	0.500	0.38	1	3/2/2006	
2-Chlorotoluene	BQL	0.500	0.207	1	3/2/2006	
4-Chlorotoluene	BQL	0.500	0.377	1	3/2/2006	
Dibromochloromethane	BQL	0.500	0.177	1	3/2/2006	
1,2-Dibromo-3-chloropropane	BQL	0.500	0.233	1	3/2/2006	
1,2-Dibromoethane (EDB)	BQL	0.500	0.208	1	3/2/2006	
1,2-Dibromomethane	BQL	0.500	0.207	1	3/2/2006	
1,2-Dichlorobenzene	BQL	0.500	0.195	1	3/2/2006	
1,3-Dichlorobenzene	BQL	0.500	0.218	1	3/2/2006	
1,4-Dichlorobenzene	BQL	0.500	0.211	1	3/2/2006	
Dichlorodifluoromethane	BQL	0.500	0.407	1	3/2/2006	
1,1-Dichloroethane	BQL	0.500	0.177	1	3/2/2006	
1,2-Dichloroethane	BQL	0.500	0.19	1	3/2/2006	
1,1-Dichloroethene	BQL	0.500	0.14	1	3/2/2006	
cis-1,2-Dichloroethene	BQL	0.500	0.423	1	3/2/2006	
trans-1,2-Dichloroethene	BQL	0.500	0.186	1	3/2/2006	
1,2-Dichloropropane	BQL	0.500	0.187	1	3/2/2006	
2,2-Dichloropropane	BQL	0.500	0.423	1	3/2/2006	
cis-1,3-Dichloropropene	BQL	0.500	0.21	1	3/2/2006	
trans-1,3-Dichloropropene	BQL	0.500	0.205	1	3/2/2006	
Diisopropyl ether (DIPE)	BQL	0.500	0.176	1	3/2/2006	
Ethylbenzene	BQL	0.500	0.166	1	3/2/2006	
Hexachlorobutadiene	BQL	0.500	0.188	1	3/2/2006	
Isopropylbenzene	BQL	0.500	0.18	1	3/2/2006	
p-Isopropyltoluene	BQL	0.500	0.383	1	3/2/2006	
Methyl-tert butyl ether (MTBE)	BQL	0.500	0.347	1	3/2/2006	
Methylene Chloride	BQL	5.00	0.464	1	3/2/2006	

Reviewed By: *[Signature]*  
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**Results for Volatiles**

by GC 6230D

Client Sample ID: PAR 163 GP2

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 9:00

Lab Sample ID: G106-566-4B

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	2.17	0.500	0.2	1	3/2/2006	
n-Propylbenzene	0.197	0.500	0.176	1	3/2/2006	J
Styrene	BQL	1.00	0.348	1	3/2/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	3/2/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	3/2/2006	
Tetrachloroethene	BQL	0.500	0.445	1	3/2/2006	
Toluene	BQL	0.500	0.187	1	3/2/2006	
1,2,3-Trichlorobenzene	BQL	0.500	0.237	1	3/2/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	3/2/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	3/2/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	3/2/2006	
Trichloroethene	BQL	0.500	0.266	1	3/2/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	3/2/2006	
1,2,3-Trichloropropane	BQL	0.500	0.929	1	3/2/2006	
1,2,4-Trimethylbenzene	1.24	0.500	0.174	1	3/2/2006	
1,3,5-Trimethylbenzene	0.380	0.500	0.374	1	3/2/2006	J
Vinyl Chloride	BQL	0.500	0.424	1	3/2/2006	
m/p-Xylene	BQL	1.00	0.35	1	3/2/2006	
o-Xylene	0.547	1.00	0.348	1	3/2/2006	J
<b>Surrogate Spike Recoveries</b>		Spike Added		Spike Result	Percent Recovery	
Trifluorotoluene		40		40.3	101	
1,4-Dichlorobutane		40		42.9	107	

**Comments:**

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: RJ  
GC\_LIMS\_v2.0

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## Results for Volatiles

by GC 6230D

Client Sample ID: PAR 127 GP1

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 15:30

Lab Sample ID: G106-566-5B

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	BQL	0.500	0.161	1	3/2/2006	
Bromobenzene	BQL	0.500	0.929	1	3/2/2006	
Bromoform	BQL	0.500	0.171	1	3/2/2006	
Bromochloromethane	BQL	0.500	0.213	1	3/2/2006	
Bromodichloromethane	BQL	0.500	0.197	1	3/2/2006	
Bromoform	BQL	0.500	0.405	1	3/2/2006	
Bromomethane	BQL	0.500	0.376	1	3/2/2006	
n-Butylbenzene	BQL	0.500	0.185	1	3/2/2006	
sec-Butylbenzene	BQL	0.500	0.175	1	3/2/2006	
tert-Butylbenzene	BQL	0.500	0.41	1	3/2/2006	
Carbon tetrachloride	BQL	0.500	0.177	1	3/2/2006	
Chlorobenzene	BQL	0.500	0.384	1	3/2/2006	
Chloroethane	BQL	0.500	0.179	1	3/2/2006	
Chloroform	BQL	0.500	0.38	1	3/2/2006	
Chloromethane	BQL	0.500	0.207	1	3/2/2006	
2-Chlorotoluene	BQL	0.500	0.377	1	3/2/2006	
4-Chlorotoluene	BQL	0.500	0.177	1	3/2/2006	
Dibromochloromethane	BQL	0.500	0.233	1	3/2/2006	
1,2-Dibromo-3-chloropropane	BQL	0.500	0.208	1	3/2/2006	
1,2-Dibromoethane (EDB)	BQL	0.500	0.207	1	3/2/2006	
Dibromomethane	BQL	0.500	0.195	1	3/2/2006	
1,2-Dichlorobenzene	BQL	0.500	0.218	1	3/2/2006	
1,3-Dichlorobenzene	BQL	0.500	0.211	1	3/2/2006	
1,4-Dichlorobenzene	BQL	0.500	0.407	1	3/2/2006	
Dichlorodifluoromethane	BQL	0.500	0.407	1	3/2/2006	
1,1-Dichloroethane	BQL	0.500	0.177	1	3/2/2006	
1,2-Dichloroethane	BQL	0.500	0.19	1	3/2/2006	
1,1-Dichloroethene	BQL	0.500	0.14	1	3/2/2006	
cis-1,2-Dichloroethene	BQL	0.500	0.423	1	3/2/2006	
trans-1,2-Dichloroethene	BQL	0.500	0.186	1	3/2/2006	
1,2-Dichloropropane	BQL	0.500	0.187	1	3/2/2006	
2,2-Dichloropropane	BQL	0.500	0.423	1	3/2/2006	
cis-1,3-Dichloropropene	BQL	0.500	0.21	1	3/2/2006	
trans-1,3-Dichloropropene	BQL	0.500	0.205	1	3/2/2006	
Diisopropyl ether (DIPE)	BQL	0.500	0.176	1	3/2/2006	
Ethylbenzene	BQL	0.500	0.166	1	3/2/2006	
Hexachlorobutadiene	BQL	0.500	0.188	1	3/2/2006	
Isopropylbenzene	BQL	0.500	0.18	1	3/2/2006	
p-Isopropyltoluene	BQL	0.500	0.383	1	3/2/2006	
Methyl-tert butyl ether (MTBE)	BQL	0.500	0.347	1	3/2/2006	
Methylene Chloride	BQL	5.00	0.464	1	3/2/2006	

Reviewed By: TJW  
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## Results for Volatiles

by GC 6230D

Client Sample ID: PAR 127 GP1

Analyzed By: MJC

Client Project ID: NCDOT-Yancey

Date Collected: 2/23/2006 15:30

Lab Sample ID: G106-566-5B

Date Received: 2/24/2006

Lab Project ID: G106-566

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	0.946	0.500	0.2	1	3/2/2006	
n-Propylbenzene	BQL	0.500	0.176	1	3/2/2006	
Styrene	BQL	1.00	0.348	1	3/2/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	3/2/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	3/2/2006	
Tetrachloroethene	BQL	0.500	0.445	1	3/2/2006	
Toluene	1.01	0.500	0.187	1	3/2/2006	
1,2,3-Trichlorobenzene	BQL	0.500	0.237	1	3/2/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	3/2/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	3/2/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	3/2/2006	
Trichloroethene	BQL	0.500	0.266	1	3/2/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	3/2/2006	
1,2,3-Trichloropropane	BQL	0.500	0.929	1	3/2/2006	
1,2,4-Trimethylbenzene	BQL	0.500	0.174	1	3/2/2006	
1,3,5-Trimethylbenzene	BQL	0.500	0.374	1	3/2/2006	
Vinyl Chloride	BQL	0.500	0.424	1	3/2/2006	
m/p-Xylene	BQL	1.00	0.35	1	3/2/2006	
o-Xylene	BQL	1.00	0.348	1	3/2/2006	
Surrogate Spike Recoveries		Spike Added		Spike Result	Percent Recovery	
Trifluorotoluene		40		40.1	100	
1,4-Dichlorobutane		40		42.7	107	

## Comments:

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: RW  
GC\_LIMS\_V2.0

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## Results for Volatiles

by GC 6230D

Client Sample ID: Method Blank

Analyzed By: MJC

Client Project ID:

Date Collected:

Lab Sample ID: VBLK3030206A

Date Received:

Lab Project ID:

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	BQL	0.500	0.161	1	3/2/2006	
Bromobenzene	BQL	0.500	0.929	1	3/2/2006	
Bromochloromethane	BQL	0.500	0.171	1	3/2/2006	
Bromodichloromethane	BQL	0.500	0.213	1	3/2/2006	
Bromoform	BQL	0.500	0.197	1	3/2/2006	
Bromomethane	BQL	0.500	0.405	1	3/2/2006	
n-Butylbenzene	BQL	0.500	0.376	1	3/2/2006	
sec-Butylbenzene	BQL	0.500	0.185	1	3/2/2006	
tert-Butylbenzene	BQL	0.500	0.41	1	3/2/2006	
Carbon tetrachloride	BQL	0.500	0.177	1	3/2/2006	
Chlorobenzene	BQL	0.500	0.384	1	3/2/2006	
Chloroethane	BQL	0.500	0.179	1	3/2/2006	
Chloroform	BQL	0.500	0.38	1	3/2/2006	
Chloromethane	BQL	0.500	0.207	1	3/2/2006	
2-Chlorotoluene	BQL	0.500	0.377	1	3/2/2006	
4-Chlorotoluene	BQL	0.500	0.177	1	3/2/2006	
Dibromochloromethane	BQL	0.500	0.233	1	3/2/2006	
1,2-Dibromo-3-chloropropane	BQL	0.500	0.208	1	3/2/2006	
1,2-Dibromoethane (EDB)	BQL	0.500	0.207	1	3/2/2006	
Dibromomethane	BQL	0.500	0.195	1	3/2/2006	
1,2-Dichlorobenzene	BQL	0.500	0.218	1	3/2/2006	
1,3-Dichlorobenzene	BQL	0.500	0.211	1	3/2/2006	
1,4-Dichlorobenzene	BQL	0.500	0.407	1	3/2/2006	
Dichlorodifluoromethane	BQL	0.500	0.177	1	3/2/2006	
1,1-Dichloroethane	BQL	0.500	0.19	1	3/2/2006	
1,2-Dichloroethane	BQL	0.500	0.14	1	3/2/2006	
1,1-Dichloroethylene	BQL	0.500	0.423	1	3/2/2006	
cis-1,2-Dichloroethylene	BQL	0.500	0.186	1	3/2/2006	
trans-1,2-Dichloroethylene	BQL	0.500	0.187	1	3/2/2006	
1,2-Dichloropropane	BQL	0.500	0.423	1	3/2/2006	
2,2-Dichloropropane	BQL	0.500	0.21	1	3/2/2006	
cis-1,3-Dichloropropene	BQL	0.500	0.205	1	3/2/2006	
trans-1,3-Dichloropropene	BQL	0.500	0.176	1	3/2/2006	
Diisopropyl ether (DIPE)	BQL	0.500	0.166	1	3/2/2006	
Ethylbenzene	BQL	0.500	0.188	1	3/2/2006	
Hexachlorobutadiene	BQL	0.500	0.18	1	3/2/2006	
Isopropylbenzene	BQL	0.500	0.383	1	3/2/2006	
p-Isopropyltoluene	BQL	0.500	0.347	1	3/2/2006	
Methyl-tert butyl ether (MTBE)	BQL	0.500	0.464	1	3/2/2006	
Methylene Chloride		5.00				

Reviewed By: *DMP*  
GC\_LIMS\_v2.0

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**Results for Volatiles**

by GC 6230D

Client Sample ID: Method Blank

Analyzed By: MJC

Client Project ID:

Date Collected:

Lab Sample ID: VBLK3030206A

Date Received:

Lab Project ID:

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	BQL	0.500	0.2	1	3/2/2006	
n-Propylbenzene	BQL	0.500	0.176	1	3/2/2006	
Styrene	BQL	1.00	0.348	1	3/2/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	3/2/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	3/2/2006	
Tetrachloroethene	BQL	0.500	0.445	1	3/2/2006	
Toluene	BQL	0.500	0.187	1	3/2/2006	
1,2,3-Trichlorobenzene	BQL	0.500	0.237	1	3/2/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	3/2/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	3/2/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	3/2/2006	
Trichloroethene	BQL	0.500	0.266	1	3/2/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	3/2/2006	
1,2,3-Trichloropropane	BQL	0.500	0.929	1	3/2/2006	
1,2,4-Trimethylbenzene	BQL	0.500	0.174	1	3/2/2006	
1,3,5-Trimethylbenzene	BQL	0.500	0.374	1	3/2/2006	
Vinyl Chloride	BQL	1.00	0.424	1	3/2/2006	
m/p-Xylene	BQL	1.00	0.35	1	3/2/2006	
o-Xylene	BQL	1.00	0.348	1	3/2/2006	
<b>Surrogate Spike Recoveries</b>		Spike Added	Spike Result	Percent Recovery		
Trifluorotoluene		40	40.2	101		
1,4-Dichlorobutane		40	39.9	99.8		

**Comments:**

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: R.M.  
GC\_LIMS\_V2.0

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## Control Limits for QC Check / Laboratory Control Spike

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Filename : 030206\006f0101.txt  
030206\006r0101.txt

Compound	ppb	Q / L (%)	Limits	
			Lower	Upper
Benzene	10.3	103.0	60.0	140.0
Bromobenzene	33.0	110.0	60.0	140.0
Bromochloromethane	10.5	105.3	60.0	140.0
Bromodichloromethane	10.8	107.6	60.0	140.0
Bromoform	10.3	102.7	60.0	140.0
Bromomethane	9.8	98.6	60.0	140.0
n-Butylbenzene	21.1	106.8	60.0	140.0
sec-Butylbenzene	10.4	104.1	60.0	140.0
tert-Butylbenzene	10.4	104.0	60.0	140.0
Carbon tetrachloride	21.6	107.8	60.0	140.0
Chlorobenzene	10.4	104.4	60.0	140.0
Chloroethane	10.4	104.1	60.0	140.0
Chloroform	10.6	105.5	60.0	140.0
Chloromethane	10.6	104.7	60.0	140.0
2-Chlorotoluene	10.4	104.0	60.0	140.0
4-Chlorotoluene	10.6	105.8	60.0	140.0
Dibromochloromethane	10.6	106.3	60.0	140.0
1,2-Dibromo-3-chloropropane	9.3	92.6	60.0	140.0
1,2-Dibromoethane (EDB)	10.7	107.2	60.0	140.0
Dibromomethane	10.6	106.0	60.0	140.0
1,2-Dichlorobenzene	10.7	107.1	60.0	140.0
1,3-Dichlorobenzene	10.6	106.0	60.0	140.0
1,4-Dichlorobenzene	10.7	107.1	60.0	140.0
Dichlorodifluoromethane	10.8	108.2	60.0	140.0
1,1-Dichloroethane	10.3	103.3	60.0	140.0
1,2-Dichloroethane	10.8	107.6	60.0	140.0
1,1-Dichloroethene	9.8	97.9	60.0	140.0
cis-1,2-Dichloroethene	21.2	106.2	60.0	140.0
trans-1,2-Dichloroethene	10.6	105.7	60.0	140.0
1,2-Dichloropropane	10.7	107.4	60.0	140.0
2,2-Dichloropropane	21.2	106.2	60.0	140.0
cis-1,3-Dichloropropene	10.8	108.1	60.0	140.0
trans-1,3-Dichloropropene	10.5	104.8	60.0	140.0
Diisopropyl ether	10.3	102.9	60.0	140.0
Ethylbenzene	10.6	106.1	60.0	140.0
Hexachlorobutadiene	11.0	110.3	60.0	140.0
Isopropylbenzene	10.6	105.4	60.0	140.0
p-Isopropyltoluene	21.6	107.5	60.0	140.0
Methyl-tert butyl ether	20.8	104.1	60.0	140.0
Methylene Chloride	10.8	107.8	60.0	140.0
Naphthalene	12.1	121.2	60.0	140.0
n-Propylbenzene	10.4	103.9	60.0	140.0
Styrene	21.1	105.3	60.0	140.0
1,1,1,2-Tetrachloroethane	10.7	106.7	60.0	140.0
1,1,2,2-Tetrachloroethane	33.0	110.0	60.0	140.0
Tetrachloroethene	21.3	106.6	60.0	140.0
Toluene	10.3	102.7	60.0	140.0
1,2,3-Trichlorobenzene	11.4	114.4	60.0	140.0
1,2,4-Trichlorobenzene	11.1	111.2	60.0	140.0
1,1,1-Trichloroethane	10.6	106.0	60.0	140.0

Reviewed by: PYL



## Control Limits for QC Check / Laboratory Control Spike

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Filename : 030206\006f0101.txt  
030206\006r0101.txt

Compound	ppb	Q / L (%)	Limits	
			Lower	Upper
1,1,2-Trichloroethane	10.6	106.4	60.0	140.0
Trichloroethene	10.7	107.4	60.0	140.0
Trichlorofluromethane	10.6	106.3	60.0	140.0
1,2,3-Trichloropropane	33.0	110.0	60.0	140.0
1,2,4-Trimethylbenzene	10.4	104.3	60.0	140.0
1,3,5-Trimethylbenzene	21.1	105.7	60.0	140.0
Vinyl Chloride	10.8	107.5	60.0	140.0
m/p-Xylene	21.2	106.8	60.0	140.0
o-Xylene	21.1	105.3	60.0	140.0

## Flags :

- + = out of QC limits.
- = lab generated limits.
- D = Detected

Reviewed by: PJY



## Control Limits for MS-MSD

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Sample : 030206\022f0101.txt  
 MS : 030206\023f0101.txt  
 MSD : 030206\024f0101.txt

030206\022r0101.txt  
 030206\023r0101.txt  
 030206\024r0101.txt

Compound	µg/L			P(%)		P Limits	
	Sam.	MS	MSD	MS	MSD	Lower	Upper
Benzene	ND	11.4	10.9	108	101	51	132
Bromobenzene	ND	21.8	30.6	73	102	51	132
Bromoform	ND	10.8	10.1	108	101	51	132
Bromochloromethane	ND	11.1	10.6	111	108	42	172
Bromodichloromethane	ND	10.0	9.2	100	92	13	168
Bromomethane	ND	8.8	7.8	82	72	1	144
n-Butylbenzene	3.54	23.9	23.0	102	97	51	132
sec-Butylbenzene	1.54	12.0	11.7	105	101	51	132
tert-Butylbenzene	ND	10.6	10.0	105	100	51	132
Carbon tetrachloride	ND	22.4	21.2	112	106	43	143
Chlorobenzene	ND	10.6	10.2	106	102	38	160
Chloroethane	ND	10.9	10.7	109	107	46	137
Chloroform	ND	11.2	10.7	112	107	49	133
Chloromethane	ND	10.8	10.7	108	107	1	193
2-Chlorotoluene	ND	11.1	10.4	111	104	51	132
4-Chlorotoluene	ND	11.7	10.5	117	105	51	132
Dibromochloromethane	ND	10.6	9.9	106	99	24	191
1,2-Dibromo-3-chloropropane	ND	10.7	9.4	100	87	51	132
1,2-Dibromomethane (EDB)	ND	11.9	10.2	119	102	51	132
Dibromomethane	ND	9.8	8.3	98	93	51	132
1,2-Dichlorobenzene	ND	11.1	10.2	111	102	1	208
1,3-Dichlorobenzene	ND	10.7	10.1	107	101	7	187
1,4-Dichlorobenzene	ND	11.2	10.3	112	103	42	143
Dichlorodifluoromethane	ND	10.7	10.7	107	107	51	132
1,1-Dichloroethane	ND	11.3	10.5	113	106	47	132
1,2-Dichloroethane	ND	11.1	10.4	111	104	51	147
1,1-Dichloroethene	ND	11.2	10.7	112	107	28	167
cis-1,2-Dichloroethene	ND	21.8	20.6	109	102	51	132
trans-1,2-Dichloroethene	ND	11.3	10.6	113	106	38	155
1,2-Dichloropropane	ND	11.4	10.8	114	108	44	156
2,2-Dichloropropane	ND	21.8	20.5	109	102	51	132
cis-1,3-Dichloropropene	ND	11.0	10.1	110	101	22	178
trans-1,3-Dichloropropene	ND	10.1	9.1	101	91	22	178
Diisopropyl ether	ND	10.2	9.6	102	96	51	132
Ethylbenzene	ND	10.2	9.9	102	99	51	132
Hexachlorobutadiene	ND	10.4	11.1	104	111	51	132
Isopropylbenzene	1.32	12.0	11.6	107	103	51	132
p-Isopropyltoluene	8.26	19.5	18.6	66	61	51	132
Methyl-tert butyl ether	ND	20.9	19.9	104	100	51	132
Methylene Chloride	ND	11.1	10.7	111	107	25	162
Naphthalene	10.16	20.1	21.6	89	113	51	132
n-Propylbenzene	1.63	12.1	11.6	106	100	51	132
Styrene	6.06	27.3	27.6	106	107	51	132
1,1,1,2-Tetrachloroethane	ND	12.0	11.1	120	111	51	132
1,1,2,2-Tetrachloroethane	ND	21.8	30.6	73	102	2	184
Tetrachloroethene	1.83	13.8	12.9	60	65	26	162
Toluene	3.44	13.9	13.6	104	101	51	132
1,2,3-Trichlorobenzene	ND	21.8	30.6	73	102	51	132
1,2,4-Trichlorobenzene	ND	10.1	10.5	101	105	51	132

Reviewed by: P.Y.



## Control Limits for MS-MSD

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Sample : 030206\022f0101.txt

030206\022r0101.txt

MS : 030206\023f0101.txt

030206\023r0101.txt

MSD : 030206\024f0101.txt

030206\024r0101.txt

Compound	µg/L			P(%)		P Limits	
	Sam.	MS	MSD	MS	MSD	Lower	Upper
1,1,1-Trichloroethane	ND	10.9	10.6	109	106	41	138
1,1,2-Trichloroethane	ND	11.0	10.4	110	104	39	136
Trichloroethene	ND	11.0	10.6	110	106	36	146
Trichlorofluoromethane	ND	10.8	10.8	108	108	21	156
1,2,3-Trichloropropane	ND	21.8	30.6	73	102	51	132
1,2,4-Trimethylbenzene	11.87	22.3	21.3	104	96	51	132
1,3,5-Trimethylbenzene	1.62	23.1	22.0	107	102	51	132
Vinyl Chloride	ND	11.4	11.3	114	113	28	163
m/p-Xylene	7.63	29.2	27.8	108	101	51	132
o-Xylene	6.06	27.3	27.5	106	107	51	132

## Flags :

- + = out of QC limits.
- = lab generated limits.
- D = Detected
- ND = None Detected

Reviewed by: Vol

**Results for Volatiles**

by GC 6230D

Client Sample ID: Method Blank

Analyzed By: MJC

Client Project ID:

Date Collected:

Lab Sample ID: VBLK3022806B

Date Received:

Lab Project ID:

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Benzene	BQL	0.500	0.161	1	2/28/2006	
Bromobenzene	BQL	0.500	0.929	1	2/28/2006	
Bromochloromethane	BQL	0.500	0.171	1	2/28/2006	
Bromodichloromethane	BQL	0.500	0.213	1	2/28/2006	
Bromoform	BQL	0.500	0.197	1	2/28/2006	
Bromomethane	BQL	0.500	0.405	1	2/28/2006	
n-Butylbenzene	BQL	0.500	0.376	1	2/28/2006	
sec-Butylbenzene	BQL	0.500	0.185	1	2/28/2006	
tert-Butylbenzene	BQL	0.500	0.175	1	2/28/2006	
Carbon tetrachloride	BQL	0.500	0.41	1	2/28/2006	
Chlorobenzene	BQL	0.500	0.177	1	2/28/2006	
Chloroethane	BQL	0.500	0.384	1	2/28/2006	
Chloroform	BQL	0.500	0.179	1	2/28/2006	
Chloromethane	BQL	0.500	0.38	1	2/28/2006	
2-Chlorotoluene	BQL	0.500	0.207	1	2/28/2006	
4-Chlorotoluene	BQL	0.500	0.377	1	2/28/2006	
Dibromochloromethane	BQL	0.500	0.177	1	2/28/2006	
1,2-Dibromo-3-chloropropane	BQL	0.500	0.233	1	2/28/2006	
1,2-Dibromoethane (EDB)	BQL	0.500	0.208	1	2/28/2006	
Dibromomethane	BQL	0.500	0.207	1	2/28/2006	
1,2-Dichlorobenzene	BQL	0.500	0.195	1	2/28/2006	
1,3-Dichlorobenzene	BQL	0.500	0.218	1	2/28/2006	
1,4-Dichlorobenzene	BQL	0.500	0.211	1	2/28/2006	
Dichlorodifluoromethane	BQL	0.500	0.407	1	2/28/2006	
1,1-Dichloroethane	BQL	0.500	0.177	1	2/28/2006	
1,2-Dichloroethane	BQL	0.500	0.19	1	2/28/2006	
1,1-Dichloroethene	BQL	0.500	0.14	1	2/28/2006	
cis-1,2-Dichloroethene	BQL	0.500	0.423	1	2/28/2006	
trans-1,2-Dichloroethene	BQL	0.500	0.186	1	2/28/2006	
1,2-Dichloropropane	BQL	0.500	0.187	1	2/28/2006	
2,2-Dichloropropane	BQL	0.500	0.423	1	2/28/2006	
cis-1,3-Dichloropropene	BQL	0.500	0.21	1	2/28/2006	
trans-1,3-Dichloropropene	BQL	0.500	0.205	1	2/28/2006	
Diisopropyl ether (DIPE)	BQL	0.500	0.176	1	2/28/2006	
Ethylbenzene	BQL	0.500	0.166	1	2/28/2006	
Hexachlorobutadiene	BQL	0.500	0.188	1	2/28/2006	
Isopropylbenzene	BQL	0.500	0.18	1	2/28/2006	
p-Isopropyltoluene	BQL	0.500	0.383	1	2/28/2006	
Methyl-tert butyl ether (MTBE)	BQL	0.500	0.347	1	2/28/2006	
Methylene Chloride	BQL	5.00	0.464	1	2/28/2006	

Reviewed By: DM  
GC\_LIMS\_v2.0

**Results for Volatiles**

by GC 6230D

Client Sample ID: Method Blank

Analyzed By: MJC

Client Project ID:

Date Collected:

Lab Sample ID: VBLK3022806B

Date Received:

Lab Project ID:

Matrix: Water

Analyte	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flags
Naphthalene	BQL	0.500	0.2	1	2/28/2006	
n-Propylbenzene	BQL	0.500	0.176	1	2/28/2006	
Styrene	BQL	1.00	0.348	1	2/28/2006	
1,1,1,2-Tetrachloroethane	BQL	0.500	0.198	1	2/28/2006	
1,1,2,2-Tetrachloroethane	BQL	0.500	0.929	1	2/28/2006	
Tetrachloroethene	BQL	0.500	0.445	1	2/28/2006	
Toluene	BQL	0.500	0.187	1	2/28/2006	
1,2,3-Trichlorobenzene	0.738	0.500	0.237	1	2/28/2006	
1,2,4-Trichlorobenzene	BQL	0.500	0.2	1	2/28/2006	
1,1,1-Trichloroethane	BQL	0.500	0.177	1	2/28/2006	
1,1,2-Trichloroethane	BQL	0.500	0.252	1	2/28/2006	
Trichloroethene	BQL	0.500	0.266	1	2/28/2006	
Trichlorofluoromethane	BQL	0.500	0.402	1	2/28/2006	
1,2,3-Trichloropropane	BQL	0.500	0.929	1	2/28/2006	
1,2,4-Trimethylbenzene	BQL	0.500	0.174	1	2/28/2006	
1,3,5-Trimethylbenzene	BQL	0.500	0.374	1	2/28/2006	
Vinyl Chloride	BQL	0.500	0.424	1	2/28/2006	
m/p-Xylene	BQL	1.00	0.35	1	2/28/2006	
o-Xylene	BQL	1.00	0.348	1	2/28/2006	
<b>Surrogate Spike Recoveries</b>		Spike Added	Spike Result	Percent Recovery		
Trifluorotoluene		40	40.1	100		
1,4-Dichlorobutane		40	38.8	97.1		

**Comments:**

All values corrected for dilution.

BQL = Below quantitation limit.

Reviewed By: DMV  
GC\_LIMS v2.0  
20 of 55



## Control Limits for QC Check / Laboratory Control Spike

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Filename : 022806\025f0101.txt  
022806\025r0101.txt

Compound	ppb	Q / L (%)	Limits	
			Lower	Upper
Benzene	10.9	109.0	60.0	140.0
Bromobenzene	33.1	110.3	60.0	140.0
Bromoform	11.2	111.6	60.0	140.0
Bromochloromethane	11.2	111.5	60.0	140.0
Bromodichloromethane	10.4	103.8	60.0	140.0
Bromoform	9.0	90.0	60.0	140.0
Bromomethane	21.8	108.9	60.0	140.0
n-Butylbenzene	10.6	106.6	60.0	140.0
sec-Butylbenzene	10.7	106.7	60.0	140.0
Carbon tetrachloride	22.1	110.3	60.0	140.0
Chlorobenzene	9.9	99.3	60.0	140.0
Chloroethane	9.8	98.4	60.0	140.0
Chloroform	10.7	107.0	60.0	140.0
Chloromethane	10.6	105.0	60.0	140.0
2-Chlorotoluene	10.6	105.6	60.0	140.0
4-Chlorotoluene	10.8	107.8	60.0	140.0
Dibromochloromethane	11.0	109.8	60.0	140.0
1,2-Dibromo-3-chloropropane	12.1	121.3	60.0	140.0
1,2-Dibromoethane (EDB)	11.1	110.8	60.0	140.0
Dibromomethane	10.5	104.5	60.0	140.0
1,2-Dichlorobenzene	11.1	111.4	60.0	140.0
1,3-Dichlorobenzene	10.6	104.7	60.0	140.0
1,4-Dichlorobenzene	11.1	110.7	60.0	140.0
Dichlorodifluoromethane	9.8	97.7	60.0	140.0
1,1-Dichloroethane	10.3	103.4	60.0	140.0
1,2-Dichloroethane	11.2	112.6	60.0	140.0
1,1-Dichloroethene	9.2	92.4	60.0	140.0
cis-1,2-Dichloroethene	20.6	103.0	60.0	140.0
trans-1,2-Dichloroethene	10.5	104.8	60.0	140.0
1,2-Dichloropropane	11.4	114.2	60.0	140.0
2,2-Dichloropropane	20.6	103.0	60.0	140.0
cis-1,3-Dichloropropene	11.1	111.1	60.0	140.0
trans-1,3-Dichloropropene	10.8	108.4	60.0	140.0
Diisopropyl ether	11.6	115.0	60.0	140.0
Ethylbenzene	11.0	109.7	60.0	140.0
Hexachlorobutadiene	10.9	108.7	60.0	140.0
Isopropylbenzene	10.9	108.7	60.0	140.0
p-Isopropyltoluene	22.3	111.6	60.0	140.0
Methyl-tert butyl ether	21.7	108.7	60.0	140.0
Methylene Chloride	10.6	106.4	60.0	140.0
Naphthalene	14.6	144.7	60.0	140.0
n-Propylbenzene	10.6	106.2	60.0	140.0
Styrene	21.6	107.8	60.0	140.0
1,1,1,2-Tetrachloroethane	12.1	120.7	60.0	140.0
1,1,2,2-Tetrachloroethane	33.1	110.3	60.0	140.0
Tetrachloroethene	21.9	108.6	60.0	140.0
Toluene	10.8	107.8	60.0	140.0
1,2,3-Trichlorobenzene	13.4	133.9	60.0	140.0
1,2,4-Trichlorobenzene	11.7	117.1	60.0	140.0
1,1,1-Trichloroethane	10.6	105.8	60.0	140.0

Reviewed by: VJF



## Control Limits for QC Check / Laboratory Control Spike

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Filename : 022806\025f0101.txt

022806\025r0101.txt

Compound	ppb	Q / L (%)	Limits	
			Lower	Upper
1,1,2-Trichloroethane	11.4	114.2	60.0	140.0
Trichloroethene	10.0	109.2	60.0	140.0
Trichlorofluoromethane	9.7	97.0	60.0	140.0
1,2,3-Trichloropropane	33.1	110.3	60.0	140.0
1,2,4-Trimethylbenzene	10.8	107.8	60.0	140.0
1,3,5-Trimethylbenzene	21.9	108.6	60.0	140.0
Vinyl Chloride	9.9	98.8	60.0	140.0
m/p-Xylene	22.0	110.0	60.0	140.0
o-Xylene	21.6	107.8	60.0	140.0

## Flags :

- + = out of QC limits.
- = lab generated limits.
- D = Detected

Reviewed by: Tom



## Control Limits for MS-MSD

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Sample : 022806\041f0101.txt 022806\041r0101.txt

MS : 022806\042f0101.txt 022806\042r0101.txt

MSD : 022806\043f0101.txt 022806\043r0101.txt

Compound	µg/L			P(%)		P Limits	
	Sam.	MS	MSD	MS	MSD	Lower	Upper
Benzene	3.16	14.7	14.6	115	115	51	132
Bromobenzene	ND	21.0	21.2	70	71	51	132
Bromochloromethane	ND	11.4	10.9	114	109	51	132
Bromodichloromethane	ND	11.6	11.1	115	111	42	172
Bromoform	ND	10.9	10.7	109	107	13	159
Bromomethane	ND	7.3	7.9	73	79	1	144
n-Butylbenzene	ND	20.7	20.6	104	103	51	132
sec-Butylbenzene	ND	10.5	10.3	105	103	51	132
tert-Butylbenzene	ND	10.6	10.4	105	103	51	132
• Carbon tetrachloride	ND	23.0	22.3	115	111	43	143
Chlorobenzene	ND	10.2	10.1	102	101	38	150
Chloroethane	ND	9.6	10.3	86	103	46	137
Chloroform	ND	11.1	10.9	111	109	49	133
Chloromethane	1.00	10.2	10.8	82	88	1	193
2-Chlorotoluene	ND	10.6	10.4	106	104	51	132
4-Chlorotoluene	ND	10.6	10.3	106	103	51	132
• Dibromochloromethane	ND	11.6	11.1	116	111	24	191
1,2-Dibromo-3-chloropropane	ND	11.3	10.4	113	104	51	132
1,2-Dibromomethane (EDB)	ND	11.2	11.0	112	110	51	132
Dibromomethane	ND	10.8	10.4	108	104	51	132
1,2-Dichlorobenzene	ND	10.6	10.6	106	106	1	208
1,3-Dichlorobenzene	ND	10.2	10.0	102	100	7	187
1,4-Dichlorobenzene	ND	10.7	10.6	107	106	42	143
Dichlorodifluoromethane	ND	10.0	10.7	100	107	51	132
1,1-Dichloroethane	ND	10.9	10.6	109	106	47	132
1,2-Dichloroethane	ND	11.5	11.3	115	113	51	147
1,1-Dichloroethene	ND	10.6	10.3	106	103	28	167
cis-1,2-Dichloroethene	ND	20.7	19.9	104	100	51	132
trans-1,2-Dichloroethene	ND	11.1	10.8	111	108	38	156
1,2-Dichloropropane	ND	11.6	11.4	116	114	44	156
2,2-Dichloropropane	ND	20.7	19.9	104	100	51	132
cis-1,3-Dichloropropene	ND	11.2	10.7	112	107	22	178
trans-1,3-Dichloropropene	ND	10.7	10.3	106	101	22	178
Diisopropyl ether	ND	12.6	12.4	122	121	51	132
Ethylbenzene	ND	10.8	10.6	108	105	51	132
Hexachlorobutadiene	ND	9.6	9.8	86	98	51	132
Isopropylbenzene	ND	11.0	10.8	110	108	51	132
p-Isopropyltoluene	ND	23.1	20.9	113	102	51	132
Methyl-tert butyl ether	ND	22.6	22.3	113	111	51	132
Methylene Chloride	ND	11.2	10.8	112	108	26	162
Naphthalene	2.36	10.2	12.8	78	104	51	132
n-Propylbenzene	ND	10.8	10.6	108	106	51	132
• Styrene	ND	25.2	26.0	126	124	51	132
1,1,1,2-Tetrachloroethane	ND	12.1	11.9	121	119	51	132
• 1,1,2,2-Tetrachloroethane	ND	21.0	21.2	70	71	8	184
Tetrachloroethene	ND	13.8	13.6	69	67	26	162
• Toluene	1.20	12.8	12.4	114	112	51	132
• 1,2,3-Trichlorobenzene	ND	21.0	21.2	70	71	51	132
1,2,4-Trichlorobenzene	ND	9.6	10.3	84	102	51	132

Reviewed by: DRW

**SGS****Control Limits for MS-MSD**

Method : 6230D Spike[ppb] : 10

Instrument : gc3

Sample : 022806\041f0101.txt      022806\041r0101.txt  
 MS : 022806\042f0101.txt      022806\042r0101.txt  
 MSD : 022806\043f0101.txt      022806\043r0101.txt

Compound	µg/L			P(%)		P Limits	
	Sam.	MS	MSD	MS	MSD	Lower	Upper
1,1,1-Trichloroethane	ND	11.1	10.8	111	108	41	138
1,1,2-Trichloroethane	ND	11.7	11.3	117	113	39	136
Trichloroethene	ND	10.9	10.6	109	106	36	146
Trichlorofluoromethane	ND	9.4	10.1	94	101	21	166
1,2,3-Trichloropropane	ND	21.0	21.2	70	71	51	132
1,2,4-Trimethylbenzene	1.38	12.5	12.3	111	110	51	132
1,3,5-Trimethylbenzene	ND	22.0	21.6	110	108	51	132
Vinyl Chloride	ND	9.7	10.6	97	106	28	163
m/p-Xylene	6.33	28.2	28.3	114	116	51	132
o-Xylene	ND	26.2	25.0	125	124	51	132

## Flags :

- + = out of QC limits.
- ♦ = lab generated limits.
- D = Detected
- ND = None Detected

Reviewed by: DRW



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: PAR 199A GP2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-11  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/22/2006 14:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	10.0	1.22	1	3/1/2006	
Acenaphthylene	BQL	10.0	1.12	1	3/1/2006	
Anthracene	BQL	10.0	1.75	1	3/1/2006	
Benzo[a]anthracene	BQL	10.0	1.36	1	3/1/2006	
Benzo[a]pyrene	BQL	10.0	1.27	1	3/1/2006	
Benzo[b]fluoranthene	BQL	10.0	1.43	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	10.0	4.57	1	3/1/2006	
Benzo[k]fluoranthene	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	10.0	1.11	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	10.0	1.57	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	10.0	1.33	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	10.0	1.99	1	3/1/2006	
Butylbenzylphthalate	BQL	10.0	1.53	1	3/1/2006	
2-Chloronaphthalene	BQL	10.0	1.25	1	3/1/2006	
2-Chlorophenol	BQL	10.0	4.22	1	3/1/2006	
2-Chlorophenol	BQL	10.0	3.26	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	10.0	1.42	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	10.0	1.11	1	3/1/2006	
Chrysene	BQL	10.0	4.87	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	10.0	1.65	1	3/1/2006	
Di-n-Butylphthalate	BQL	10.0	1.25	1	3/1/2006	
1,2-Dichlorobenzene	BQL	10.0	1.24	1	3/1/2006	
1,3-Dichlorobenzene	BQL	10.0	1.20	1	3/1/2006	
1,4-Dichlorobenzene	BQL	10.0	4.10	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	20.0	3.75	1	3/1/2006	
2,4-Dichlorophenol	BQL	10.0	1.48	1	3/1/2006	
Diethylphthalate	11.2	10.0	1.04	1	3/1/2006	
Dimethylphthalate	BQL	10.0	9.25	1	3/1/2006	
2,4-Dimethylphenol	BQL	10.0	1.16	1	3/1/2006	
Di-n-octylphthalate	BQL	10.0	3.71	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	50.0	4.20	1	3/1/2006	
2,4-Dinitrophenol	BQL	50.0	1.52	1	3/1/2006	
2,4-Dinitrotoluene	BQL	10.0	1.41	1	3/1/2006	
2,6-Dinitrotoluene	BQL	10.0	1.53	1	3/1/2006	
Diphenylamine *	BQL	10.0	1.41	1	3/1/2006	
Fluoranthene	BQL	10.0	1.22	1	3/1/2006	
Fluorene	BQL	10.0	1.22	1	3/1/2006	
Hexachlorobenzene	BQL	10.0	1.58	1	3/1/2006	
Hexachlorobutadiene	BQL	20.0	20.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	10.0	1.58	1	3/1/2006	
Hexachloroethane	BQL	10.0	4.57	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	10.0	1.27	1	3/1/2006	
Isophorone	BQL	10.0	1.08	1	3/1/2006	
Naphthalene	BQL	10.0	1.32	1	3/1/2006	
Nitrobenzene	BQL					



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: PAR 199A GP2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-11  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/22/2006 14:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
2-Nitrophenol	BQL	10.0	3.52	1	3/1/2006	
4-Nitrophenol	BQL	50.0	3.17	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	10.0	1.87	1	3/1/2006	
Pentachlorophenol	BQL	50.0	2.83	1	3/1/2006	
Phenanthrene	BQL	10.0	1.38	1	3/1/2006	
Phenol	BQL	10.0	3.38	1	3/1/2006	
Pyrene	BQL	10.0	2.08	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	10.0	1.33	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	10.0	2.92	1	3/1/2006	
		Spike Added	Spike Result	Percent Recovered		
2-Fluorobiphenyl		10	8.9	89		
2-Fluorophenol		10	8.5	85		
Nitrobenzene-d5		10	9.7	97		
Phenol-d6		10	8.9	89		
2,4,6-Tribromophenol		10	9.8	98		
4-Terphenyl-d14		10	10.3	103		

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: DRW



**Results of Library Search for Semivolatile Compounds  
by GCMS**

Client Sample ID: PAR 199A GP2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-1I  
 Lab Project ID: G106-566  
 Sample Wt/Vol: 500 ML  
 Dilution: 1

Analyzed By: MRC  
 Date Collected: 2/22/2006 14:45  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Date Analyzed: 3/1/2006  
 Matrix: Water

No.	Compound	Retention Time	CAS#	Match Probability	Result (ug/L)
1	No library search compounds detected.				
2					
3					
4					
5					
6					
7					
8					
9					
10					

**Comment:**

Tentatively Identified Compound (TIC) refers to substances which are not present in the list of target compounds. Therefore, not all TICs are identified and quantitated using individual standards. TIC listings are prepared utilizing a computerized library search of electron impact mass spectral data and evaluation of the relevant data by a mass spectral data specialist.

Quantitation is accomplished by relative peak area of the compound compared to that of the nearest internal standard from the total ion chromatogram. TICs are identified and quantitated only if the peak area is equal to or greater than 10% of that of the nearest internal standard. Quantitation provided is an estimate.

Reviewed by: D.W.



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: PAR 206 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-2E  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 9:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	10.0	1.22	1	3/1/2006	
Acenaphthylene	BQL	10.0	1.12	1	3/1/2006	
Anthracene	BQL	10.0	1.75	1	3/1/2006	
Benzo[a]anthracene	BQL	10.0	1.36	1	3/1/2006	
Benzo[a]pyrene	BQL	10.0	1.27	1	3/1/2006	
Benzo[b]fluoranthene	BQL	10.0	4.57	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	10.0	1.09	1	3/1/2006	
Benzo[k]fluoranthene	BQL	10.0	1.11	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	10.0	1.57	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	10.0	1.33	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	10.0	1.99	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	10.0	1.53	1	3/1/2006	
Butylbenzylphthalate	BQL	10.0	1.25	1	3/1/2006	
2-Chloronaphthalene	BQL	10.0	4.22	1	3/1/2006	
2-Chlorophenol	BQL	10.0	3.26	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	10.0	1.42	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	10.0	1.11	1	3/1/2006	
Chrysene	BQL	10.0	4.87	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	10.0	1.65	1	3/1/2006	
Di-n-Butylphthalate	BQL	10.0	1.25	1	3/1/2006	
1,2-Dichlorobenzene	BQL	10.0	1.24	1	3/1/2006	
1,3-Dichlorobenzene	BQL	10.0	1.20	1	3/1/2006	
1,4-Dichlorobenzene	BQL	20.0	4.10	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	10.0	3.75	1	3/1/2006	
2,4-Dichlorophenol	BQL	10.0	1.48	1	3/1/2006	
Diethylphthalate	BQL	10.0	1.04	1	3/1/2006	
Dimethylphthalate	BQL	10.0	9.25	1	3/1/2006	
2,4-Dimethylphenol	BQL	10.0	1.16	1	3/1/2006	
Di-n-octylphthalate	BQL	50.0	3.71	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	50.0	4.20	1	3/1/2006	
2,4-Dinitrophenol	BQL	10.0	1.52	1	3/1/2006	
2,4-Dinitrotoluene	BQL	10.0	1.41	1	3/1/2006	
2,6-Dinitrotoluene	BQL	10.0	1.53	1	3/1/2006	
Diphenylamine *	BQL	10.0	1.41	1	3/1/2006	
Fluoranthene	BQL	10.0	1.22	1	3/1/2006	
Fluorene	BQL	10.0	1.22	1	3/1/2006	
Hexachlorobenzene	BQL	10.0	1.58	1	3/1/2006	
Hexachlorobutadiene	BQL	20.0	20.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	10.0	1.58	1	3/1/2006	
Hexachloroethane	BQL	10.0	4.57	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	10.0	1.27	1	3/1/2006	
Isophorone	BQL	10.0	1.08	1	3/1/2006	
Naphthalene	BQL	10.0	1.32	1	3/1/2006	
Nitrobenzene	BQL					

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**Results for Semivolatiles  
by GCMS 625**

Client Sample ID: PAR 206 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-2E  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 9:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
2-Nitrophenol	BQL	10.0	3.52	1	3/1/2006	
4-Nitrophenol	BQL	50.0	3.17	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	10.0	1.87	1	3/1/2006	
Pentachlorophenol	BQL	50.0	2.83	1	3/1/2006	
Phenanthrene	BQL	10.0	1.38	1	3/1/2006	
Phenol	BQL	10.0	3.38	1	3/1/2006	
Pyrene	BQL	10.0	2.08	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	10.0	1.33	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	10.0	2.92	1	3/1/2006	
	Spike Added		Spike Result	Percent Recovered		
2-Fluorobiphenyl	10		9.1	91		
2-Fluorophenol	10		7.9	79		
Nitrobenzene-d5	10		9.5	95		
Phenol-d6	10		8.4	84		
2,4,6-Tribromophenol	10		9.5	95		
4-Terphenyl-d14	10		10.6	106		

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: PSL



**Results of Library Search for Semivolatile Compounds**  
by GCMS

Client Sample ID: PAR 206 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-2E  
 Lab Project ID: G106-566  
 Sample Wt/Vol: 500 ML  
 Dilution: 1

Analyzed By: MRC  
 Date Collected: 2/23/2006 9:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Date Analyzed: 3/1/2006  
 Matrix: Water

No.	Compound	Retention Time	CAS#	Match Probability	Result (ug/L)
1	No library search compounds detected.				
2					
3					
4					
5					
6					
7					
8					
9					
10					

**Comment:**

Tentatively Identified Compound (TIC) refers to substances which are not present in the list of target compounds. Therefore, not all TICs are identified and quantitated using individual standards. TIC listings are prepared utilizing a computerized library search of electron impact mass spectral data and evaluation of the relevant data by a mass spectral data specialist.

Quantitation is accomplished by relative peak area of the compound compared to that of the nearest internal standard from the total ion chromatogram. TICs are identified and quantitated only if the peak area is equal to or greater than 10% of that of the nearest internal standard. Quantitation provided is an estimate.

Reviewed by: mtw



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: PAR 221 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-3I  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 11:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed:	Flag
Acenaphthene	8.30	10.0	1.22	1	3/1/2006	J
Acenaphthylene	2.70	10.0	1.12	1	3/1/2006	
Anthracene	BQL	10.0	1.75	1	3/1/2006	
Benzo[a]anthracene	BQL	10.0	1.36	1	3/1/2006	
Benzo[a]pyrene	BQL	10.0	1.27	1	3/1/2006	
Benzo[b]fluoranthene	BQL	10.0	1.43	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	10.0	4.57	1	3/1/2006	
Benzo[k]fluoranthene	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	10.0	1.11	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	10.0	1.57	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	10.0	1.33	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	10.0	1.99	1	3/1/2006	
Butylbenzylphthalate	BQL	10.0	1.53	1	3/1/2006	
2-Choronaphthalene	BQL	10.0	1.25	1	3/1/2006	
2-Chlorophenol	BQL	10.0	4.22	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	10.0	3.26	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	10.0	1.42	1	3/1/2006	
Chrysene	BQL	10.0	1.11	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	10.0	4.87	1	3/1/2006	
Di-n-Butylphthalate	BQL	10.0	1.65	1	3/1/2006	
1,2-Dichlorobenzene	BQL	10.0	1.25	1	3/1/2006	
1,3-Dichlorobenzene	BQL	10.0	1.24	1	3/1/2006	
1,4-Dichlorobenzene	BQL	10.0	1.20	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	20.0	4.10	1	3/1/2006	
2,4-Dichlorophenol	BQL	10.0	3.75	1	3/1/2006	
Diethylphthalate	BQL	10.0	1.48	1	3/1/2006	
Dimethylphthalate	BQL	10.0	1.04	1	3/1/2006	
2,4-Dimethylphenol	BQL	10.0	9.25	1	3/1/2006	
Di-n-octylphthalate	BQL	10.0	1.16	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	50.0	3.71	1	3/1/2006	
2,4-Dinitrophenol	BQL	50.0	4.20	1	3/1/2006	
2,4-Dinitrotoluene	BQL	10.0	1.53	1	3/1/2006	
2,6-Dinitrotoluene	BQL	10.0	1.52	1	3/1/2006	
Diphenylamine *	1.60	10.0	1.41	1	3/1/2006	
Fluoranthene	10.7	10.0	1.22	1	3/1/2006	
Fluorene	BQL	10.0	1.22	1	3/1/2006	
Hexachlorobenzene	BQL	10.0	1.58	1	3/1/2006	
Hexachlorobutadiene	BQL	20.0	20.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	10.0	1.58	1	3/1/2006	
Hexachloroethane	BQL	10.0	4.57	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	10.0	1.27	1	3/1/2006	
Isophorone	61.8	10.0	1.08	1	3/1/2006	
Naphthalene	BQL	10.0	1.32	1	3/1/2006	
Nitrobenzene						

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**Results for Semivolatiles  
by GCMS 625**

Client Sample ID: PAR 221 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-3I  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 11:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
2-Nitrophenol	BQL	10.0	3.52	1	3/1/2006	
4-Nitrophenol	BQL	50.0	3.17	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	10.0	1.87	1	3/1/2006	
Pentachlorophenol	BQL	50.0	2.83	1	3/1/2006	
Phenanthrene	30.7	10.0	1.38	1	3/1/2006	
Phenol	BQL	10.0	3.38	1	3/1/2006	J
Pyrene	7.10	10.0	2.08	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	10.0	1.33	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	10.0	2.92	1	3/1/2006	
	Spike Added	Spike Result	Percent Recovered			
2-Fluorobiphenyl	10	7.9	79			
2-Fluorophenol	10	7.7	77			
Nitrobenzene-d5	10	9.7	97			
Phenol-d6	10	8	80			
2,4,6-Tribromophenol	10	7.9	79			
4-Terphenyl-d14	10	7.8	78			

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: pd



**Results of Library Search for Semivolatile Compounds**  
*by GCMS*

Client Sample ID: PAR 221 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-3I  
 Lab Project ID: G106-566  
 Sample Wt/Vol: 500 ML  
 Dilution: 1

Analyzed By: MRC  
 Date Collected: 2/23/2006 11:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Date Analyzed: 3/1/2006  
 Matrix: Water

No.	Compound	Retention Time	CAS#	Match Probability	Result (ug/L)
1	Alkane, Unknown	7.91			175
2	2-Methylnaphthalene	10.03	000091-57-6	97	141
3	Alkane, Unknown	6.76			118
4	Alkane, Unknown	9.94			105
5	1-Methylnaphthalene	10.18	000090-12-0	99	84.3
6	Alkane, Unknown	16.61			72.8
7	Alkane, Unknown	13.30			71
8	Unknown	7.57			66.5
9	Ethenyldimethylbenzene, Isomer of	8.50			59.5
10	Alkane, Unknown	9.66			56.8

**Comment:**

Tentatively Identified Compound (TIC) refers to substances which are not present in the list of target compounds. Therefore, not all TICs are identified and quantitated using individual standards. TIC listings are prepared utilizing a computerized library search of electron impact mass spectral data and evaluation of the relevant data by a mass spectral data specialist.

Quantitation is accomplished by relative peak area of the compound compared to that of the nearest internal standard from the total ion chromatogram. TICs are identified and quantitated only if the peak area is equal to or greater than 10% of that of the nearest internal standard. Quantitation provided is an estimate.

Reviewed by: RW



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: PAR 163 GP2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-4I  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 9:00  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	10.0	1.22	1	3/1/2006	
Acenaphthylene	BQL	10.0	1.12	1	3/1/2006	
Anthracene	BQL	10.0	1.75	1	3/1/2006	
Benzo[a]anthracene	BQL	10.0	1.36	1	3/1/2006	
Benzo[a]pyrene	BQL	10.0	1.27	1	3/1/2006	
Benzo[b]fluoranthene	BQL	10.0	4.57	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	10.0	1.09	1	3/1/2006	
Benzo[k]fluoranthene	BQL	10.0	1.11	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	10.0	1.57	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	10.0	1.33	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	10.0	1.99	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	10.0	1.53	1	3/1/2006	
Butylbenzylphthalate	BQL	10.0	1.25	1	3/1/2006	
2-Chloronaphthalene	BQL	10.0	4.22	1	3/1/2006	
2-Chlorophenol	BQL	10.0	3.26	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	10.0	1.42	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	10.0	1.11	1	3/1/2006	
Chrysene	BQL	10.0	4.87	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	10.0	1.65	1	3/1/2006	
Di-n-Butylphthalate	BQL	10.0	1.25	1	3/1/2006	
1,2-Dichlorobenzene	BQL	10.0	1.24	1	3/1/2006	
1,3-Dichlorobenzene	BQL	10.0	1.20	1	3/1/2006	
1,4-Dichlorobenzene	BQL	20.0	4.10	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	10.0	3.75	1	3/1/2006	
2,4-Dichlorophenol	BQL	10.0	1.48	1	3/1/2006	
Diethylphthalate	BQL	10.0	1.04	1	3/1/2006	
Dimethylphthalate	BQL	10.0	9.25	1	3/1/2006	
2,4-Dimethylphenol	BQL	10.0	1.16	1	3/1/2006	
Di-n-octylphthalate	BQL	50.0	3.71	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	50.0	4.20	1	3/1/2006	
2,4-Dinitrophenol	BQL	10.0	1.52	1	3/1/2006	
2,4-Dinitrotoluene	BQL	10.0	1.41	1	3/1/2006	
2,6-Dinitrotoluene	BQL	10.0	1.53	1	3/1/2006	
Diphenylamine *	BQL	10.0	1.41	1	3/1/2006	
Fluoranthene	BQL	10.0	1.22	1	3/1/2006	
Fluorene	BQL	10.0	1.22	1	3/1/2006	
Hexachlorobenzene	BQL	10.0	1.58	1	3/1/2006	
Hexachlorobutadiene	BQL	20.0	20.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	10.0	1.58	1	3/1/2006	
Hexachloroethane	BQL	10.0	4.57	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	10.0	1.27	1	3/1/2006	
Isophorone	3.10	10.0	1.08	1	3/1/2006	
Naphthalene	BQL	10.0	1.32	1	3/1/2006	
Nitrobenzene						

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**Results for Semivolatiles  
by GCMS 625**

Client Sample ID: PAR 163 GP2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-4I  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 9:00  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
2-Nitrophenol	BQL	10.0	3.52	1	3/1/2006	
4-Nitrophenol	BQL	50.0	3.17	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	10.0	1.87	1	3/1/2006	
Pentachlorophenol	BQL	50.0	2.83	1	3/1/2006	
Phenanthrene	BQL	10.0	1.38	1	3/1/2006	
Phenol	BQL	10.0	3.38	1	3/1/2006	
Pyrene	BQL	10.0	2.08	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	10.0	1.33	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	10.0	2.92	1	3/1/2006	
	Spike Added	Spike Result	Percent Recovered			
2-Fluorobiphenyl	10	8.7	87			
2-Fluorophenol	10	7.7	77			
Nitrobenzene-d5	10	9.3	93			
Phenol-d6	10	8.2	82			
2,4,6-Tribromophenol	10	9.5	95			
4-Terphenyl-d14	10	10.3	103			

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: PNP



**Results of Library Search for Semivolatile Compounds**  
*by GCMS*

Client Sample ID: PAR 163 GP2  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-4I  
 Lab Project ID: G106-566  
 Sample Wt/Vol: 500 ML  
 Dilution: 1

Analyzed By: MRC  
 Date Collected: 2/23/2006 9:00  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Date Analyzed: 3/1/2006  
 Matrix: Water

No.	Compound	Retention Time	CAS#	Match Probability	Result (ug/L)
1	No library search compounds detected.				
2					
3					
4					
5					
6					
7					
8					
9					
10					

**Comment:**

Tentatively Identified Compound (TIC) refers to substances which are not present in the list of target compounds. Therefore, not all TICs are identified and quantitated using individual standards. TIC listings are prepared utilizing a computerized library search of electron impact mass spectral data and evaluation of the relevant data by a mass spectral data specialist.

Quantitation is accomplished by relative peak area of the compound compared to that of the nearest internal standard from the total ion chromatogram. TICs are identified and quantitated only if the peak area is equal to or greater than 10% of that of the nearest internal standard. Quantitation provided is an estimate.

Reviewed by: Phl



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: PAR 127 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-5I  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 15:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	10.0	1.22	1	3/1/2006	
Acenaphthylene	BQL	10.0	1.12	1	3/1/2006	
Anthracene	BQL	10.0	1.75	1	3/1/2006	
Benzo[a]anthracene	BQL	10.0	1.36	1	3/1/2006	
Benzo[a]pyrene	BQL	10.0	1.27	1	3/1/2006	
Benzo[b]fluoranthene	BQL	10.0	1.43	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	10.0	4.57	1	3/1/2006	
Benzo[k]fluoranthene	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	10.0	1.11	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	10.0	1.57	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	10.0	1.33	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	10.0	1.99	1	3/1/2006	
Butylbenzylphthalate	BQL	10.0	1.53	1	3/1/2006	
2-Chloronaphthalene	BQL	10.0	1.25	1	3/1/2006	
2-Chlorophenol	BQL	10.0	4.22	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	10.0	3.26	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	10.0	1.42	1	3/1/2006	
Chrysene	BQL	10.0	1.11	1	3/1/2006	
Dibenzo[a,h]anthracene	BQL	10.0	4.87	1	3/1/2006	
Di-n-Butylphthalate	BQL	10.0	1.65	1	3/1/2006	
1,2-Dichlorobenzene	BQL	10.0	1.25	1	3/1/2006	
1,3-Dichlorobenzene	BQL	10.0	1.24	1	3/1/2006	
1,4-Dichlorobenzene	BQL	10.0	1.20	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	20.0	4.10	1	3/1/2006	
2,4-Dichlorophenol	BQL	10.0	3.75	1	3/1/2006	
Diethylphthalate	BQL	10.0	1.48	1	3/1/2006	
Dimethylphthalate	BQL	10.0	1.04	1	3/1/2006	
2,4-Dimethylphenol	BQL	10.0	9.25	1	3/1/2006	
Di-n-octylphthalate	BQL	10.0	1.16	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	50.0	3.71	1	3/1/2006	
2,4-Dinitrophenol	BQL	50.0	4.20	1	3/1/2006	
2,4-Dinitrotoluene	BQL	10.0	1.52	1	3/1/2006	
2,6-Dinitrotoluene	BQL	10.0	1.41	1	3/1/2006	
Diphenylamine *	BQL	10.0	1.53	1	3/1/2006	
Diphenylamine *	BQL	10.0	1.41	1	3/1/2006	
Fluoranthene	BQL	10.0	1.22	1	3/1/2006	
Fluorene	BQL	10.0	1.22	1	3/1/2006	
Hexachlorobenzene	BQL	10.0	1.58	1	3/1/2006	
Hexachlorobutadiene	BQL	10.0	20.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	20.0	1.58	1	3/1/2006	
Hexachloroethane	BQL	10.0	4.57	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	10.0	1.27	1	3/1/2006	
Isophorone	BQL	10.0	1.08	1	3/1/2006	
Naphthalene	BQL	10.0	1.32	1	3/1/2006	
Nitrobenzene	BQL					



**Results for Semivolatiles  
by GCMS 625**

Client Sample ID: PAR 127 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-5I  
 Lab Project ID: G106-566

Analyzed By: MRC  
 Date Collected: 2/23/2006 15:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Matrix: Water

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
2-Nitrophenol	BQL	10.0	3.52	1	3/1/2006	
4-Nitrophenol	BQL	50.0	3.17	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	10.0	1.87	1	3/1/2006	
Pentachlorophenol	BQL	50.0	2.83	1	3/1/2006	
Phenanthrene	BQL	10.0	1.38	1	3/1/2006	
Phenol	BQL	10.0	3.38	1	3/1/2006	
Pyrene	BQL	10.0	2.08	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	10.0	1.33	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	10.0	2.92	1	3/1/2006	
	Spike Added	Spike Result	Percent Recovered			
2-Fluorobiphenyl	10	9.2	92			
2-Fluorophenol	10	8.8	88			
Nitrobenzene-d5	10	10.1	101			
Phenol-d6	10	9.1	91			
2,4,6-Tribromophenol	10	10.4	104			
4-Terphenyl-d14	10	7.2	72			

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: PL



**Results of Library Search for Semivolatile Compounds**  
*by GCMS*

Client Sample ID: PAR 127 GP1  
 Client Project ID: NCDOT-Yancey  
 Lab Sample ID: G106-566-51  
 Lab Project ID: G106-566  
 Sample Wt/Vol: 500 ML  
 Dilution: 1

Analyzed By: MRC  
 Date Collected: 2/23/2006 15:30  
 Date Received: 2/24/2006  
 Date Extracted: 2/27/2006  
 Date Analyzed: 3/1/2006  
 Matrix: Water

No.	Compound	Retention Time	CAS#	Match Probability	Result (ug/L)
1	Unknown	7.22			16.4
2	Unknown	12.61			8.24
3	Unknown	12.66			5.22
4	Unknown	7.60			4.52
5	Unknown	10.44			4.51
6					
7					
8					
9					
10					

**Comment:**

Tentatively Identified Compound (TIC) refers to substances which are not present in the list of target compounds. Therefore, not all TICs are identified and quantitated using individual standards. TIC listings are prepared utilizing a computerized library search of electron impact mass spectral data and evaluation of the relevant data by a mass spectral data specialist.

Quantitation is accomplished by relative peak area of the compound compared to that of the nearest internal standard from the total ion chromatogram. TICs are identified and quantitated only if the peak area is equal to or greater than 10% of that of the nearest internal standard. Quantitation provided is an estimate.

Reviewed by: znd



**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: Method Blank  
 Client Project ID:  
 Lab Sample ID: PB4588  
 Lab Project ID:

Analyzed By: MRC  
 Date Collected:  
 Date Received:  
 Date Extracted: 2/27/2006  
 Matrix: WATER

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
Acenaphthene	BQL	10.0	1.22	1	3/1/2006	
Acenaphthylene	BQL	10.0	1.12	1	3/1/2006	
Anthracene	BQL	10.0	1.75	1	3/1/2006	
Benzo[a]anthracene	BQL	10.0	1.36	1	3/1/2006	
Benzo[a]pyrene	BQL	10.0	1.27	1	3/1/2006	
Benzo[b]fluoranthene	BQL	10.0	4.57	1	3/1/2006	
Benzo[g,h,i]perylene	BQL	10.0	1.09	1	3/1/2006	
Benzo[k]fluoranthene	BQL	10.0	1.11	1	3/1/2006	
Bis(2-chloroethoxy)methane	BQL	10.0	1.09	1	3/1/2006	
Bis(2-chloroethyl)ether	BQL	10.0	1.57	1	3/1/2006	
Bis(2-chloroisopropyl)ether	BQL	10.0	1.33	1	3/1/2006	
Bis(2-ethylhexyl)phthalate	BQL	10.0	1.99	1	3/1/2006	
4-bromophenyl phenyl ether	BQL	10.0	1.53	1	3/1/2006	
Butylbenzylphthalate	BQL	10.0	1.25	1	3/1/2006	
2-Chloronaphthalene	BQL	10.0	4.22	1	3/1/2006	
2-Chlorophenol	BQL	10.0	3.26	1	3/1/2006	
4-Chloro-3-methylphenol	BQL	10.0	1.42	1	3/1/2006	
4-Chlorophenyl phenyl ether	BQL	10.0	1.11	1	3/1/2006	
Chrysene	BQL	10.0	4.87	1	3/1/2006	
Dibenz[a,h]anthracene	BQL	10.0	1.65	1	3/1/2006	
Di-n-Butylphthalate	BQL	10.0	1.25	1	3/1/2006	
1,2-Dichlorobenzene	BQL	10.0	1.24	1	3/1/2006	
1,3-Dichlorobenzene	BQL	10.0	1.20	1	3/1/2006	
1,4-Dichlorobenzene	BQL	20.0	4.10	1	3/1/2006	
3,3'-Dichlorobenzidine	BQL	10.0	3.75	1	3/1/2006	
2,4-Dichlorophenol	BQL	10.0	1.48	1	3/1/2006	
Diethylphthalate	BQL	10.0	1.04	1	3/1/2006	
Dimethylphthalate	BQL	10.0	9.25	1	3/1/2006	
2,4-Dimethylphenol	BQL	10.0	1.16	1	3/1/2006	
Di-n-octylphthalate	BQL	50.0	3.71	1	3/1/2006	
4,6-Dinitro-2-methylphenol	BQL	50.0	4.20	1	3/1/2006	
2,4-Dinitrophenol	BQL	10.0	1.52	1	3/1/2006	
2,4-Dinitrotoluene	BQL	10.0	1.41	1	3/1/2006	
2,6-Dinitrotoluene	BQL	10.0	1.53	1	3/1/2006	
Diphenylamine *	BQL	10.0	1.41	1	3/1/2006	
Fluoranthene	BQL	10.0	1.22	1	3/1/2006	
Fluorene	BQL	10.0	1.22	1	3/1/2006	
Hexachlorobenzene	BQL	10.0	1.58	1	3/1/2006	
Hexachlorobutadiene	BQL	20.0	20.0	1	3/1/2006	
Hexachlorocyclopentadiene	BQL	10.0	1.58	1	3/1/2006	
Hexachloroethane	BQL	10.0	4.57	1	3/1/2006	
Indeno(1,2,3-c,d)pyrene	BQL	10.0	1.27	1	3/1/2006	
Isophorone	BQL	10.0	1.08	1	3/1/2006	
Naphthalene	BQL	10.0	1.32	1	3/1/2006	
Nitrobenzene	BQL					

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**Results for Semivolatiles**  
by GCMS 625

Client Sample ID: Method Blank  
 Client Project ID:  
 Lab Sample ID: PB4588  
 Lab Project ID:

Analyzed By: MRC  
 Date Collected:  
 Date Received:  
 Date Extracted: 2/27/2006  
 Matrix: WATER

Compound	Result ug/L	RL ug/L	MDL ug/L	Dilution Factor	Date Analyzed	Flag
2-Nitrophenol	BQL	10.0	3.52	1	3/1/2006	
4-Nitrophenol	BQL	50.0	3.17	1	3/1/2006	
N-Nitrosodi-n-propylamine	BQL	10.0	1.87	1	3/1/2006	
Pentachlorophenol	BQL	50.0	2.83	1	3/1/2006	
Phenanthrene	BQL	10.0	1.38	1	3/1/2006	
Phenol	BQL	10.0	3.38	1	3/1/2006	
Pyrene	BQL	10.0	2.08	1	3/1/2006	
1,2,4-Trichlorobenzene	BQL	10.0	1.33	1	3/1/2006	
2,4,6-Trichlorophenol	BQL	10.0	2.92	1	3/1/2006	
		Spike Added	Spike Result	Percent Recovered		
2-Fluorobiphenyl		10	8.8	88		
2-Fluorophenol		10	8.5	85		
Nitrobenzene-d5		10	9.7	97		
Phenol-d6		10	8.9	88		
2,4,6-Tribromophenol		10	9.7	97		
4-Terphenyl-d14		10	9.6	96		

**Comments:**

\* N-Nitrosodiphenylamine is reported as the breakdown product Diphenylamine.

**Flags:**

BQL = Below Quantitation Limits.

J = Detected below the quantitation limit.

Reviewed By: PY



**Results For Matrix Spike / Matrix Spike Duplicate and Laboratory Control Standard (MS/MSD/LCS)**  
by GCMS

Client Sample ID: Batch QC

Date Collected:

Client Project ID:

Date Received:

Lab Sample ID: Batch-4588-MS/MSD/LCS

Date Extracted: 02/27/06

Lab Project ID:

Date Analyzed: 03/01/06

Matrix: WATER

Analyzed By: MRC

Prep Method: 3520

Dilution: 1

	Sample	MS	MS	MS	MSD	MSD	MSD	RPD	QC Limits	
	Amount ( $\mu\text{g}/\text{L}$ )	Spike ( $\mu\text{g}/\text{L}$ )	Conc. ( $\mu\text{g}/\text{L}$ )	% Rec.	Spike ( $\mu\text{g}/\text{L}$ )	Conc. ( $\mu\text{g}/\text{L}$ )	% Rec.		RPD	% Rec.
Acenaphthylene	BQL	222	202	91.1	222	213	95.8	5.03	30	62.0-119
4-Chloro-3-methylphenol	BQL	222	197	88.7	222	193	86.7	2.28	30	67.0-109
2-Chlorophenol	BQL	222	182	81.8	222	181	81.4	0.490	30	59.0-95.0
1,4-Dichlorobenzene	BQL	222	125	56.3	222	122	54.8	2.70	30	29.0-86.0
2,4-Dinitrotoluene	BQL	222	195	87.8	222	195	87.7	0.114	30	63.0-103
N-Nitrosodi-n-propylamine	BQL	222	176	79.4	222	174	78.1	1.65	30	67.0-107
4-Nitrophenol	BQL	222	241	108	222	228	103.0	5.31	30	49.0-146
Pentachlorophenol	BQL	222	205	92.2	222	203	91.4	0.871	30	43.0-106
Phenol	BQL	222	180	81.0	222	176	79.4	2.00	30	61.0-100
Pyrene	BQL	222	172	77.2	222	179	80.5	4.19	30	41.0-123
1,2,4-Trichlorobenzene	BQL	222	154	69.1	222	156	70.2	1.58	30	41.0-96.0

	Spiked	LCS	LCS	QC Limits % Rec.
	Amount ( $\mu\text{g}/\text{L}$ )	Conc. ( $\mu\text{g}/\text{L}$ )	Spike %	
Acenaphthylene	100	96.2	96.2	72.9-127
4-Chloro-3-methylphenol	100	87.8	87.8	61.6-113
2-Chlorophenol	100	80.3	80.3	52.3-104
1,4-Dichlorobenzene	100	47.9	47.9	27.3-85.0
2,4-Dinitrotoluene	100	88.8	88.8	66.5-117
N-Nitrosodi-n-propylamine	100	81.3	81.3	54.4-119
4-Nitrophenol	100	102	102	32.4-150
Pentachlorophenol	100	91.3	91.3	23.9-115
Phenol	100	79.9	79.9	54.5-106
Pyrene	100	85.9	85.9	63.5-126
1,2,4-Trichlorobenzene	100	65.3	65.3	49.1-98.5

**Comments:**

Concentrations reflect the spiked sample amounts.

**Flags:**

\* = Out of limits.

NA = Not applicable.

Reviewed By: RNP



## VPH (Aliphatics/Aromatics) Laboratory Reporting Form

Client Name: Environmental Investigations

Project Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 199A GP2
Sample Matrix	Water
Collection Option (for Soil)*	
Date Collected	02/22/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	02/28/06
Dry Weight	
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 100 ( $\mu\text{g/L}$ )
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	< 100 ( $\mu\text{g/L}$ )
C <sub>9</sub> -C <sub>10</sub> Aromatics**	< 100 ( $\mu\text{g/L}$ )
Surrogate % Recovery - PID	100
Surrogate % Recovery - FID	99

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-566-1d

Reviewed By: RNP



## VPH (Aliphatics/Aromatics) Laboratory Reporting Form

Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

<b>Sample Information and Analytical Results</b>	
Sample Identification	PAR 221 GP1
Sample Matrix	Water
Collection Option (for Soil)*	
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	02/28/06
Dry Weight	
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 100 ( $\mu\text{g}/\text{L}$ )
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	270 ( $\mu\text{g}/\text{L}$ )
C <sub>9</sub> -C <sub>10</sub> Aromatics**	220 ( $\mu\text{g}/\text{L}$ )
Surrogate % Recovery - PID	100
Surrogate % Recovery - FID	110

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-566-3d

Reviewed By: TKJ



## VPH (Aliphatics/Aromatics) Laboratory Reporting Form

Client Name: Environmental InvestigationsProject Name: NCDOT-Yancey

<b>Sample Information and Analytical Results</b>	
Sample Identification	PAR 163 GP2
Sample Matrix	Water
Collection Option (for Soil)*	
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	02/28/06
Dry Weight	
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 100 (µg/L)
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	< 100 (µg/L)
C <sub>9</sub> -C <sub>10</sub> Aromatics**	< 100 (µg/L)
Surrogate % Recovery - PID	100
Surrogate % Recovery - FID	98

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-566-4d

Reviewed By: EN



### VPH (Aliphatics/Aromatics) Laboratory Reporting Form

Client Name: Environmental Investigations

Project Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 127 GP1
Sample Matrix	Water
Collection Option (for Soil)*	
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	02/28/06
Dry Weight	
Dilution Factor	1
C <sub>5</sub> -C <sub>8</sub> Aliphatics**	< 100 ( $\mu\text{g/L}$ )
C <sub>9</sub> -C <sub>12</sub> Aliphatics**	< 100 ( $\mu\text{g/L}$ )
C <sub>9</sub> -C <sub>10</sub> Aromatics**	< 100 ( $\mu\text{g/L}$ )
Surrogate % Recovery - PID	100
Surrogate % Recovery - FID	100

\* = Option 1 = Established fill line on vial, Option 2 = Sampling Device/Brand, or Option 3 = Field weight of soil.

\*\* = Excludes any surrogates or internal standards.

Lab Info: g106-566-5d

Reviewed By: Paul

**SGS**

## Attachment 2

## VPH Laboratory Reporting Form

## Calibration and QA/QC Information

FID Initial Calibration Date:	<u>02/11/06</u>	PID Initial Calibration Date:	<u>02/11/06</u>
-------------------------------	-----------------	-------------------------------	-----------------

## Calibration Ranges and Limits

Range	MDL (07/15/2004) ( $\mu\text{g/L}$ )	ML ( $\mu\text{g/L}$ )	RL ( $\mu\text{g/L}$ )	(mg/Kg)
C <sub>5</sub> -C <sub>8</sub> Aliphatics	4.4	14	100	10
C <sub>9</sub> -C <sub>12</sub> Aliphatics	3.4	11	100	10
C <sub>9</sub> -C <sub>10</sub> Aromatics	0.13	0.41	100	10

## Calibration Concentration Levels

Range	Levels ( $\mu\text{g/L}$ )	%RSD or CCC	Method of Quantitation
C <sub>5</sub> -C <sub>8</sub> Aliphatics	40	10.8	Calibration Factor
	1000		
	2000		
	3000		
	4000		
C <sub>9</sub> -C <sub>12</sub> Aliphatics	10	0.99	Linear Regression
	250		
	500		
	750		
	1000		
C <sub>9</sub> -C <sub>10</sub> Aromatics	10	19.30	Calibration Factor
	250		
	500		
	750		
	1000		

Calibration Check Date: 02/28/06

## Calibration Check

Range	Levels ( $\mu\text{g/L}$ )	RPD
C <sub>5</sub> -C <sub>8</sub> Aliphatics	2000 200	2.0
C <sub>9</sub> -C <sub>12</sub> Aliphatics	500 50	-5.9
C <sub>9</sub> -C <sub>10</sub> Aromatics	500 50	-4.9

MDL = Method Detection Limit

RPD = Relative Percent Difference

ML = Minimum Limit

%RSD = Percent Relative Standard Deviation

RL = Reportable Limit

CCC = Correlation Coefficient of Curve



**EPH (Aliphatics/Aromatics) Results**  
by MDEP-EPH

Client Name: Environmental Investigations

Project Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 199A GP2
Sample Matrix	Water
Date Collected	02/22/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	02/28/06
Dry Weight	
Dilution Factor	1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	< 100 (ug/L)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	< 100 (ug/L)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	< 100 (ug/L)
Aliphatic Surrogate % Recovery	80
Aromatic Surrogate % Recovery	85

**Comments:**

\* = Excludes any surrogates or internal standards.

Sample did not require fractionation.

Lab info: G106-566-1J

Reviewed By: erf



## EPH (Aliphatics/Aromatics) Results

by MDEP-EPH

Client Name: Environmental Investigations

Project Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 221 GP1
Sample Matrix	Water
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	03/06/06
Dry Weight	
Dilution Factor	1:1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	1600 (ug/L)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	180 (ug/L)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	840 (ug/L)
Aliphatic Surrogate % Recovery	65
Aromatic Surrogate % Recovery	64
Fractionation Surrogate 1 % Recovery	93

## Comments:

\* = Excludes any surrogates or internal standards.

Lab Info: G106-566-3J

Reviewed By: RNP



## EPH (Aliphatics/Aromatics) Results

by MDEP-EPH

Client Name: Environmental Investigations

Project Name: NCDOT-Yancey

<b>Sample Information and Analytical Results</b>	
Sample Identification	PAR 163 GP2
Sample Matrix	Water
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	02/28/06
Dry Weight	
Dilution Factor	1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	< 100 (ug/L)
C <sub>19</sub> -C <sub>36</sub> Aliphatics*	< 100 (ug/L)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	< 100 (ug/L)
Aliphatic Surrogate % Recovery	58
Aromatic Surrogate % Recovery	88

**Comments:**

\* = Excludes any surrogates or internal standards.

Sample did not require fractionation.

Lab info: G106-566-4J

Reviewed By: ren

**SGS****EPH (Aliphatics/Aromatics) Results**

by MDEP-EPH

Client Name: Environmental Investigations

Project Name: NCDOT-Yancey

Sample Information and Analytical Results	
Sample Identification	PAR 127 GP1
Sample Matrix	Water
Date Collected	02/23/06
Date Received	02/24/06
Date Extracted	02/28/06
Date Analyzed	03/06/06
Dry Weight	
Dilution Factor	1:1
C <sub>9</sub> -C <sub>18</sub> Aliphatics*	< 100 (ug/L)
C <sub>18</sub> -C <sub>36</sub> Aliphatics*	< 100 (ug/L)
C <sub>11</sub> -C <sub>22</sub> Aromatics*	< 100 (ug/L)
Aliphatic Surrogate % Recovery	39
Aromatic Surrogate % Recovery	42
Fractionation Surrogate 1 % Recovery	93

**Comments:**

\* = Excludes any surrogates or internal standards.

Duplicate analysis confirms low surrogates.

Lab info: G106-566-5L

Reviewed By: EY



Attachment 3

## EPH Laboratory Reporting Form

## Calibration and QA/QC Information

Initial Calibration Date: 12/28/05

## Calibration Ranges and Limits

Range	MDL (2/2004) ( $\mu\text{g}/\text{L}$ )	ML ( $\mu\text{g}/\text{L}$ )	( $\mu\text{g}/\text{L}$ )	RL (mg/Kg)
C <sub>9</sub> -C <sub>18</sub> Aliphatics	3.84	12.2	100	10
C <sub>19</sub> -C <sub>36</sub> Aliphatics	0.57	1.8	100	10
C <sub>11</sub> -C <sub>22</sub> Aromatics	4.54	14.4	100	10

## Calibration Concentration Levels

Range	Levels ( $\mu\text{g}/\text{mL}$ )	%RSD or CCC	Method of Quantitation
C <sub>9</sub> -C <sub>18</sub> Aliphatics	6	24.90	Calibration Factor
	30		
	60		
	120		
	240		
C <sub>19</sub> -C <sub>36</sub> Aliphatics	8	15.4	Calibration Factor
	40		
	80		
	160		
	320		
C <sub>11</sub> -C <sub>22</sub> Aromatics	17	9.8	Calibration Factor
	85		
	170		
	340		
	680		

Calibration Check Date: 03/06/06

## Calibration Check

Range	Levels ( $\mu\text{g}/\text{mL}$ )	RPD
C <sub>9</sub> -C <sub>18</sub> Aliphatics	120	12.7
C <sub>19</sub> -C <sub>36</sub> Aliphatics	160	6.7
C <sub>11</sub> -C <sub>22</sub> Aromatics	340	12.9

MDL = Method Detection Limit

ML = Minimum Limit

RL = Reportable Limit

RPD = Relative Percent Difference

%RSD = Percent Relative Standard Deviation

CCC = Correlation Coefficient of Curve



Attachment 3  
EPH Laboratory Reporting Form

<b>Calibration and QA/QC Information</b>
--

Initial Calibration Date: 12/28/05

**Calibration Ranges and Limits**

Range	MDL (2/2004) ( $\mu\text{g}/\text{L}$ )	ML ( $\mu\text{g}/\text{L}$ )	RL ( $\mu\text{g}/\text{L}$ )	RL (mg/Kg)
C <sub>9</sub> -C <sub>18</sub> Aliphatics	3.84	12.2	100	10
C <sub>19</sub> -C <sub>36</sub> Aliphatics	0.57	1.8	100	10
C <sub>11</sub> -C <sub>22</sub> Aromatics	4.54	14.4	100	10

**Calibration Concentration Levels**

Range	Levels ( $\mu\text{g}/\text{mL}$ )	%RSD or CCC	Method of Quantitation
C <sub>9</sub> -C <sub>18</sub> Aliphatics	6	24.90	Calibration Factor
	30		
	60		
	120		
	240		
C <sub>19</sub> -C <sub>36</sub> Aliphatics	8	15.4	Calibration Factor
	40		
	80		
	160		
	320		
C <sub>11</sub> -C <sub>22</sub> Aromatics	17	9.8	Calibration Factor
	85		
	170		
	340		
	680		

Calibration Check Date: 02/28/06

**Calibration Check**

Range	Levels ( $\mu\text{g}/\text{mL}$ )	RPD
C <sub>9</sub> -C <sub>18</sub> Aliphatics	120	17.0
C <sub>19</sub> -C <sub>36</sub> Aliphatics	160	10.3
C <sub>11</sub> -C <sub>22</sub> Aromatics	340	10.7

MDL = Method Detection Limit

ML = Minimum Limit

RL = Reportable Limit

RPD = Relative Percent Difference

%RSD = Percent Relative Standard Deviation

CCC = Correlation Coefficient of Curve



List of Reporting Abbreviations  
and Data Qualifiers

B = Compound also detected in batch blank

BQL = Below Quantitation Limit (RL or MDL)

DF = Dilution Factor

Dup = Duplicate

D = Detected, but RPD is > 40% between results in dual column method.

E = Estimated concentration, exceeds calibration range.

J = Estimated concentration, below calibration range and above MDL

LCS(D) = Laboratory Control Spike (Duplicate)

MDL = Method Detection Limit

MS(D) = Matrix Spike (Duplicate)

PQL = Practical Quantitation Limit

RL = Reporting Limit

RPD = Relative Percent Difference

mg/kg = milligram per kilogram, ppm, parts per million

ug/kg = micrograms per kilogram, ppb, parts per billion

mg/L = milligram per liter, ppm, parts per million

ug/L = micrograms per liter, ppb, parts per billion

% Rec = Percent Recovery

% solids = Percent Solids

Special Notes:

1) Metals and mercury samples are digested with a hot block, see the standard operating procedure document for details.

2) Uncertainty for all reported data is less than or equal to 30 percent.

MI34.030606.3

**SGS**

Mr. Bob Shaut  
Environmental Investigations  
2101 Gateway Centre Boulevard  
Suite 200  
Morrisville NC 27560  
Report Number: G106-583

Client Project: Yancy DOT

Dear Mr. Shaut:

Enclosed are the results of the analytical services performed under the referenced project. The samples are certified to meet the requirements of the National Environmental Laboratory Accreditation Conference Standards. Copies of this report and supporting data will be retained in our files for a period of five years in the event they are required for future reference. Any samples submitted to our laboratory will be retained for a maximum of thirty (30) days from the date of this report unless other arrangements are requested.

If there are any questions about the report or the services performed during this project, please call Paradigm at (910) 350-1903. We will be happy to answer any questions or concerns which you may have.

Thank you for using Paradigm Analytical Labs for your analytical services. We look forward to working with you again on any additional analytical needs which you may have.

Sincerely,  
Paradigm Analytical Laboratories, Inc.

Laboratory Director  
J. Patrick Weaver

4/10/06

Date

C

C

C



**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: HO-1  
Client Project ID: Yancy DOT  
Lab Sample ID: G106-583-1  
Lab Project ID: G106-583  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 3/29/2006 17:00  
Date Received: 4/5/2006  
Matrix: Soil  
Solids 82.85

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	8.01	5035	1	04/05/06
Diesel Range Organics	BQL	7.45	3541	1	04/06/06

Reviewed By: MJC  
TPH\_DMS\_V2.0  
2 of 6



**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: HO-2  
Client Project ID: Yancy DOT  
Lab Sample ID: G106-583-2  
Lab Project ID: G106-583  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 3/29/2006 17:20  
Date Received: 4/5/2006  
Matrix: Soil  
Solids 80.00

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	7.89	5035	1	04/05/06
Diesel Range Organics	BQL	7.79	3541	1	04/06/06

Reviewed By:

3 of 6



**Results for Total Petroleum Hydrocarbons**  
by GC/FID 8015

Client Sample ID: HO-3  
Client Project ID: Yancy DOT  
Lab Sample ID: G106-583-3  
Lab Project ID: G106-583  
Report Basis: Dry Weight

Analyzed By: MJC  
Date Collected: 3/29/2006 17:40  
Date Received: 4/5/2006  
Matrix: Soil  
Solids 78.48

Analyte	Result MG/KG	RL MG/KG	Prep Method	Dilution Factor	Date Analyzed
Gasoline Range Organics	BQL	9.46	5035	1	04/05/06
Diesel Range Organics	BQL	7.81	3541	1	04/06/06



List of Reporting Abbreviations  
and Data Qualifiers

B = Compound also detected in batch blank

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ug/kg = micrograms per kilogram, ppb, parts per billion

mg/L = milligram per liter, ppm, parts per million

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% soilds = Percent Solids

Special Notes:

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- 2) Uncertainty for all reported data is less than or equal to 30 percent.

MI34.030606.3

# SGS

## CHAIN OF CUSTODY RECORD

**SGS Environmental Services Inc.**

Locations Nationwide  
 • Alaska      • Hawaii  
 • Louisiana    • Maryland  
 • New Jersey   • North Carolina  
[www.us.sgs.com](http://www.us.sgs.com)

056740

SGS Reference:

G106-583

PAGE 1 OF 1

<b>1</b> CLIENT: <b>EJ, Inc.</b> CONTACT: <b>BOB SHAW</b> PHONE NO: <b>(919) 624-5569</b> PROJECT: <b>YANKEE DOT</b> SITE/PWSID: <b>MCGUIGER RES.</b> REPORTS TO:  INVOICE TO: <b>NC DOT</b> FAX NO.: <b>( )</b> QUOTE #: <b>                </b> P.O. NUMBER: <b>                </b>						SGS Reference: <b>G106-583</b>  <b>PAGE 1 OF 1</b>											
<b>2</b>	LAB NO.	SAMPLE IDENTIFICATION	DATE	TIME	MATRIX	No	SAMPLE TYPE	Preservative Used	R	Analysis Required	C <sup>c</sup> COMP	C <sup>c</sup> GRAS	G <sup>c</sup> GRAS	G <sup>c</sup> DRO	TPH GRO	TPH DRO	REMARKS
<b>3</b>	Collected/Relinquished By: (1)	Date <b>4/4/06</b>	Time <b>1700</b>	Received By: <b>J. J. H.</b>	<b>4</b>	Shipping Carrier:	Samples Received Cold? (Circle) <b>YES</b> NO	Temperature (C): <b>55°C</b>	Chain of Custody Seal: (Circle)	<b>INTACT</b>	<b>BROKEN</b>	<b>ABSENT</b>					
<b>5</b>	Relinquished By: (2)	Date <b>4/5/06</b>	Time <b>1030</b>	Received By: <b>J. J. H.</b>	<b>6</b>	Special Deliverable Requirements:											
<b>7</b>	Relinquished By: (3)	Date	Time	Received By:	<b>7</b>	Requested Turnaround Time and Special Instructions:	<b>JUST 24 HR TAT</b>										
<b>8</b>	Published By: (4)	Date	Time	Received By:	<b>8</b>												
<b>9</b>	<b>SGS</b>				<b>9</b>												