

**2014 OU2 GROUNDWATER INVESTIGATION  
VPB 147  
BETHPAGE, NY**

Prepared for:



**Department of the Navy  
Naval Facilities Engineering Command, Mid-Atlantic  
9742 Maryland Ave.  
Norfolk, VA 23511-3095**

**Comprehensive Long-Term Environmental Action Navy  
Contract Number N62470-11-D-8013**

**CTO WE15**

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**February 2015**

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## List of Acronyms and Abbreviations

AOC	Area of Concern
bgs	below ground surface
DoD	Department of Defense
ELAP	Environmental Laboratory Accreditation Program
EPA	Environmental Protection Agency, United States
ft	feet
GOCO	Government-Owned Contractor-Operated
GPS	Global Positioning System
IDW	Investigation Derived Waste
IR	Installation Restoration
Katahdin	Katahdin Analytical Services, Inc
NAD	North American Datum
NAVD	North American Vertical Datum
NAVFAC	Naval Facilities Engineering Command
NG	Northrop Grumman
NWIRP	Naval Weapons Industrial Reserve Plant
NYSDEC	New York State Department of Environmental Conservation
OU	Operable Unit
PCBs	Polychlorinated Biphenyls
PCE	Tetrachloroethene
PID	Photoionization Detector
POTW	Publicly Owned Treatment Works
PPE	Personal Protective Equipment
SAP	Sampling and analysis plan
SVOC	Semivolatile Organic Compounds
TCE	Trichloroethene
TCL	Target Compound List
TCLP	Toxicity Characteristic Leaching Procedure
TOC	Total Organic Carbon
UFP	United Federal Programs
VOC	Volatile Organic Compounds
VPB	Vertical Profile Boring

## **1.0 PROJECT BACKGROUND**

Resolution Consultants has prepared this Data Summary Report for the Naval Facilities Engineering Command (NAVFAC), Mid-Atlantic under contract task order WE15 Contract N62470-11-D-8013. This report describes vertical profile boring (VPB) installation activities (specifically at the VPB 147 location) in 2014 for the Naval Weapons Industrial Reserve Plant (NWIRP) Bethpage Operable Unit (OU) 2 Site 1 offsite plume. NWIRP Bethpage is located in east-central Nassau County, Long Island, New York, approximately 30 miles east of New York City (Figure 1).

### **1.1 Scope and Objectives**

This data summary report provides information on the installation of VPB 147. The purpose of the VPB 147 investigation was to ascertain subsurface conditions and contaminant levels upgradient of the Massapequa Water District wells and to design outpost wells that will be used to provide early warning of plume migration toward the Massapequa Water District wellfield. VPB locations within the general vicinity of VPB 147 are shown in Figure 2. VPB 147 was completed to 1030 feet (ft) below ground surface (bgs).

Field tasks were conducted in 2014 in accordance with the *United Federal Programs Sampling and Analysis Plan (UFP SAP)*, Bethpage, New York and the UFP SAP Addendum Installation of Vertical Profile Borings and Monitoring Wells (Resolution Consultants, 2013). The field investigation included completing one vertical profile boring, groundwater grab samples, geophysical logging and surveying.

Documentation of these activities is included in Appendix A of this report.

### **1.2 Site History**

NWIRP Bethpage is in the Hamlet of Bethpage, Town of Oyster Bay, New York. Since its inception in 1941, the plant's primary mission was the research prototyping, testing, design, engineering, fabrication, and primary assembly of military aircraft. The facilities at NWIRP included four plants used for assembly and prototype testing, a group of quality control laboratories, two warehouse complexes (north and south), a salvage storage area, water recharge basins, the Industrial Wastewater Treatment Plant, and several smaller support buildings.

The Navy's property originally totaled 109.5 acres and was formerly a Government-Owned Contractor-Operated (GOCO) facility that was operated by Northrop Grumman (NG) until September 1998. Prior to 2002, the NWIRP property was bordered on the north, west, and south

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by current or former NG facilities, and on the east by a residential neighborhood. By March 2008, approximately 100 acres of NWIRP property were transferred to Nassau County in three separate actions. The remaining 9 acres and access easements were retained by the Navy to continue remedial efforts at Installation Restoration (IR) Site 1 – Former Drum Marshalling Area and Site 4 – Former Underground Storage Tanks (Area of Concern [AOC] 22). A parcel of land connecting the two sites was also retained. Currently, the 9-acre parcel of NWIRP is bordered on the east by the residential neighborhood and on the north, south, and west by Nassau County property. Access to the NWIRP is from South Oyster Bay Road.

### **1.3 Geology and Hydrogeology**

Overburden at the site consists of well over 1,000 ft of Cretaceous deposits overlying crystalline bedrock of the Hartland Formation. Overburden is divided into four geologic units: the upper Pleistocene deposits, the Magothy Formation, the clay member of the Raritan Formation (“Raritan Clay”) and the Lloyd Sand member of the Raritan Formation (“Lloyd Sand”) (Geraghty and Miller, 1994).

The upper Pleistocene ranges in thickness from approximately 50 to 100 ft and consists of till and outwash deposits of medium to coarse sand and gravel with lenses of fine sand, silt and clay (Smolensky and Feldman, 1990); these deposits form the Upper Glacial Aquifer. Directly underlying this unit is the Magothy Formation with a thickness of 650 to 900 ft bgs observed onsite. The Magothy is characterized by fine to medium sands and silts interbedded with zones of clays, silty sands and sandy clays. Sand and gravel lenses are found in some areas between depths of 600 and 875 ft bgs; these deposits form the Magothy Aquifer.

Investigations performed by the Navy since 2012 indicate that the bottom of the Magothy (top of the Raritan Clay) can extend to depths of 700 to greater than 1,000 ft bgs. The top of the Raritan Clay deepens to the south southeast, as evidenced by clay depths of 1,000 ft bgs (or more) in borings installed in the offsite plume to date. The Raritan Clay Unit is of continental origin and consists of clay, silty clay, clayey silt, and fine silty sand. This member acts as a confining layer over the Lloyd Sand Unit. The Lloyd Sand Unit is also of continental origin, having been deposited in a large fresh water lacustrine environment. The material consists of fine to coarse-grained sands, gravel, interbedded clay, and silty sand. These deposits form the Lloyd Aquifer.

The Upper Glacial Aquifer and the Magothy Aquifer comprise the aquifers of interest at the NWIRP. Regionally, these formations are generally considered to form a common, interconnected aquifer as

the coarse nature of each unit near their contact and the lack of any regionally confining clay unit allows for the unrestricted flow of groundwater between the formations.

The Magothy Aquifer is the major source of public water in Nassau County. The most productive water bearing zones are the discontinuous lenses of sand and gravel that occur within the siltier matrix. The major water-bearing zones are coarse sand and gravel lenses located in the lower portion of the Magothy. The Magothy Aquifer is commonly regarded to function overall as an unconfined aquifer at shallow depths and a confined aquifer at deeper depths. The drilling program at the NWIRP has revealed that clay zones beneath the facility are common but laterally discontinuous. No confining clay units of facility-wide extent have been encountered.

Groundwater is encountered at a depth of approximately 50 ft bgs at the facility. Historically, because of pumping and recharge at the facility, groundwater depths have been measured to range from 40 to 60 ft bgs. The regional groundwater flow in the area is to the south-southeast.

## **2.0 FIELD PROGRAM**

Field investigation activities at VPB 147 consisted of drilling, sampling, soil/groundwater analysis, geophysical logging, and surveying. Drilling during this investigation was performed by Delta Well and Pump Company of Ronkonkoma, New York. A description of these tasks is provided below.

### **2.1 Vertical Profile Borings**

One vertical profile boring (VPB 147) was completed during this field effort between May 29, 2014 and July 14, 2014. The total depth of VPB 147 was 1030 ft. The location is shown in Figure 2 and details are summarized in Table 1.

#### **2.1.1 Drilling**

VPB 147 was installed by drilling a 9-inch diameter hole using mud rotary drilling techniques. Drilling mud consisted of potable water and polymer-free sodium bentonite or equivalent. Drilling mud was contained and re-circulated in baffled, high capacity mud tubs. A sand separator was used intermittently to remove fines from circulation.

#### **2.1.2 Sampling**

A total of 16 split spoon samples were collected from ground surface to the bottom of the boring. A change in geology was observed by the field geologist at 1003 ft bgs and six split spoon samples were subsequently collected to confirm the presence of the Raritan Clay. Samples were logged by the field geologist and screened for Volatile Organic Compounds (VOCs) utilizing a photoionization detector (PID). A detailed boring log for VPB 147 is included in Appendix A.

Groundwater grab samples were collected every 50 ft for the first 200 ft of borehole depth. After the first 200 ft, groundwater grab samples were collected approximately every 20 ft until the boring terminated in the Raritan. Groundwater grab samples were collected with a hydropunch sampler and analyzed for VOCs using Environmental Protection Agency (EPA) Method 8260B. The groundwater grab samples were analyzed by Katahdin Analytical Services (Katahdin), a Department of Defense (DoD), Environmental Laboratory Accreditation Program (ELAP), and New York State Department of Environmental Conservation (NYSDEC)-certified laboratory. During the collection of groundwater grab samples, field parameters were measured (pH, temperature, specific conductivity, oxidation reduction potential, dissolved oxygen, and turbidity). Data validation was performed by Resolution Consultants. Groundwater grab sample logs, data validation packages, and analytical data tables are included in Appendix A.



One soil sample was collected for laboratory analysis for total organic carbon (TOC) by EPA series SW-846 method 9060A. During drilling, air sampling was conducted under a Community Air Monitoring Plan. One air sample was collected per VPB using Summa canisters and submitted for laboratory analysis by EPA Method TO-15. All analyses were performed or sub-contracted by Katahdin. Data validation of both TOC and air data was performed by Resolution Consultants. Data validation packages and analytical data tables are included in Appendix A.

### **2.1.3 Geophysics**

Borehole geophysical logs (gamma) were recorded after the borehole was drilled but prior to the removal of drill rods. A Mount Sopris Instrument model 2PGA-100 poly gamma was used. Starting at the top of the hole, the probe was advanced at a maximum rate of 12 ft per minute. A copy of the log was printed in the field for review once the probe reached the bottom of the borehole. The instrument was then raised to the top of the boring and a second log was generated and printed in the field. The down hole gamma log sheets and plots comparing the gamma log with trichloroethene (TCE) and tetrachloroethene (PCE) concentrations from hydropunch samples are included in Appendix A.

## **2.2 Decontamination and Investigation Derived Waste (IDW)**

Resolution Consultants utilized dedicated and disposable sampling equipment when possible to avoid the potential for cross-contamination of samples. The sampling equipment included dedicated plastic scoops, disposable Teflon or polyethylene tubing, disposable gloves, and laboratory supplied sample bottles. Hand held equipment, split spoons, and the hydropunch were decontaminated using Liquinox and water wash, a potable water rinse, followed by a distilled water rinse. Water was collected in 5-gallon pails or 55-gallon drums.

As part of the IDW management practices and in accordance with the SAP, the investigation waste (consisting of soil cuttings, drilling muds, IDW fluids, and personal protective equipment [PPE]) generated during the boring installation was containerized and staged at NWIRP Bethpage. IDW solids were characterized and disposed of properly. Representative samples from each roll off were submitted to Katahdin for analysis of:

- Target Compound List (TCL) VOCs
- TCL Semi-volatile Organic Compounds (SVOCs)
- Toxicity Characteristic Leaching Procedure (TCLP) Metals

- Polychlorinated Biphenyls (PCBs)
- Total petroleum hydrocarbons
- Corrosivity
- Ignitability
- Reactive Cyanide
- Reactive Sulfide
- Paint Filter

IDW water was containerized in frac tanks and stored at NWIRP Bethpage for characterization and ultimate disposal to the Publicly Owned Treatment Works (POTW), in accordance with the facilities existing discharge permit. A representative water sample was collected from each frac tank and submitted to Katahdin for analysis of VOCs via Method SW 624, pH via Method SW 9040B, PCBs via Method 8082 and Total Metals via Method SW 846 (all waters). To the extent feasible, soil and water were not mixed. All analytical criteria were met for disposal of soil and water.

### **2.3 Surveying**

A survey of the boring location was conducted at the end of fieldwork by C. T. Male, Inc., of Latham, NY, under the direct supervision of Resolution Consultants. The location was tied into the existing base map developed for this investigation. The survey elevation is referenced to the North American Vertical Datum (NAVD) 1988 and has a vertical accuracy of 0.01 foot. Vertical control is based on observations of COR Stations Queens and Central Islip. The horizontal location is referenced to the North American Datum (NAD) 1983 (2011) N.Y. Long Island Zone 3104 and has an accuracy of 0.1 foot. Local horizontal and vertical control is based on Global Positioning System (GPS) observations using the NYS Net Real Time Network.

A table of survey data (ground, latitude/longitude and northing/easting) and a survey map is included in Appendix A.

### 3.0 REFERENCES

Geraghty and Miller, Inc., 1994. *Remedial Investigation Report, Grumman Aerospace Corporation, Bethpage, New York*. Revised September 1994.

Naval Facilities Engineering Command (NAVFAC), 2003. *Record of Decision Naval Weapons Industrial Reserve Plant Bethpage, New York, Operable Unit 2 – Groundwater*, NYS Registry: 1-30-003B. April.

Resolution Consultants, 2013. *United Federal Programs Sampling and Analysis Plan, Site OU-2 Offsite TCE Groundwater Plume Investigation*, Bethpage, New York. April.

Resolution Consultants, 2013. UFP SAP Addendum, *Groundwater Sampling Using Low Stress (Low Flow) Purging and Sampling Protocol*. November.

Resolution Consultants, 2013. UFP SAP Addendum, *Installation of Vertical Profile Borings and Monitoring Wells*. December.

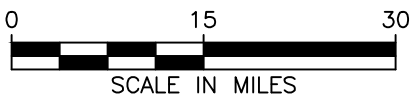
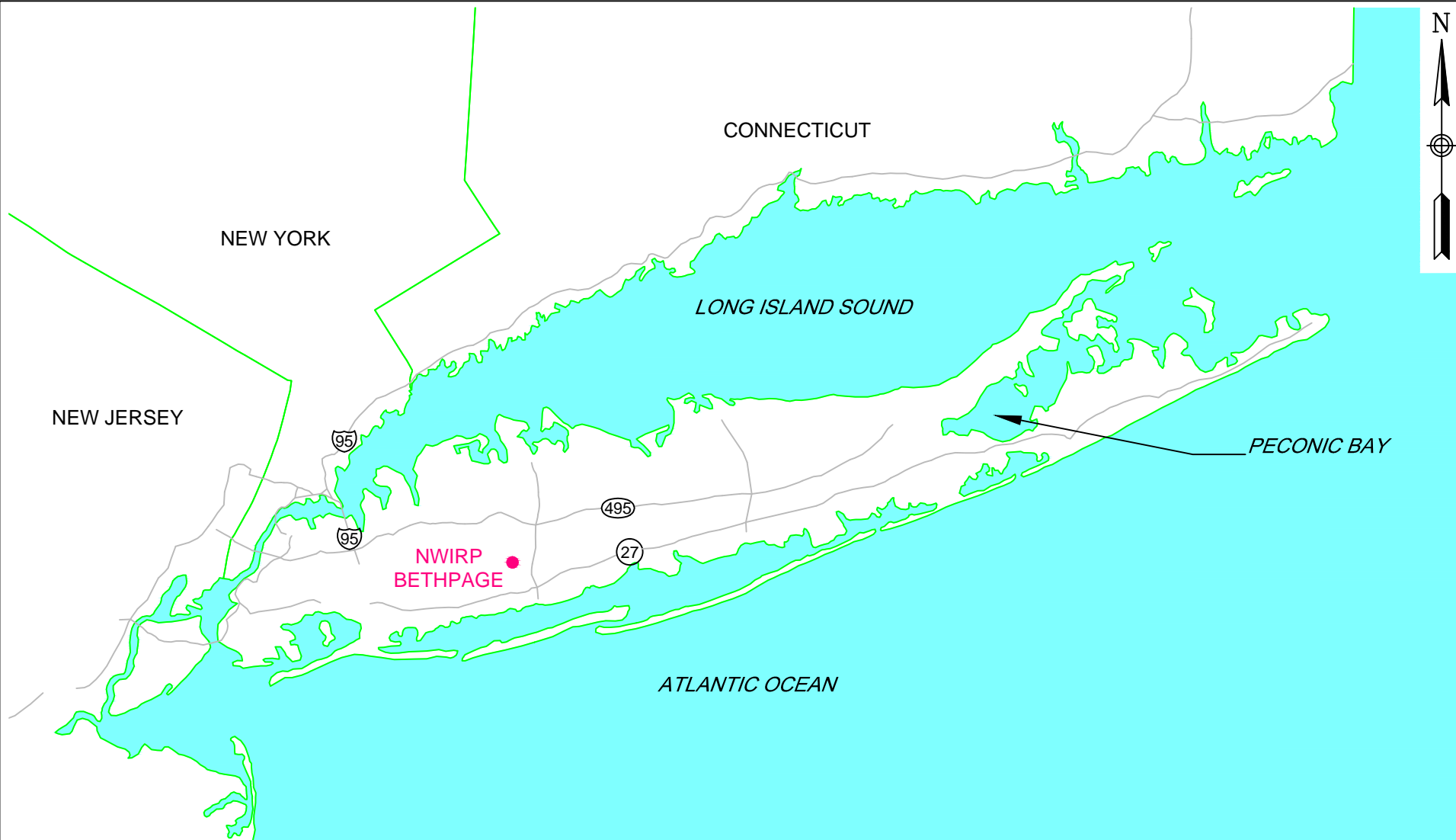
Smolensky, D., and Feldman, S., 1990. *Geohydrology of the Bethpage-Hicksville-Levittown Area, Long Island, New York*, U.S. Geological Survey Water-Resourced Investigations Report 88-4135, 25 pp.

## Tables

**TABLE 1  
VERTICAL PROFILE BORING SUMMARY  
2014 OU2 GROUNDWATER INVESTIGATION  
NWIRP BETHPAGE, NY**

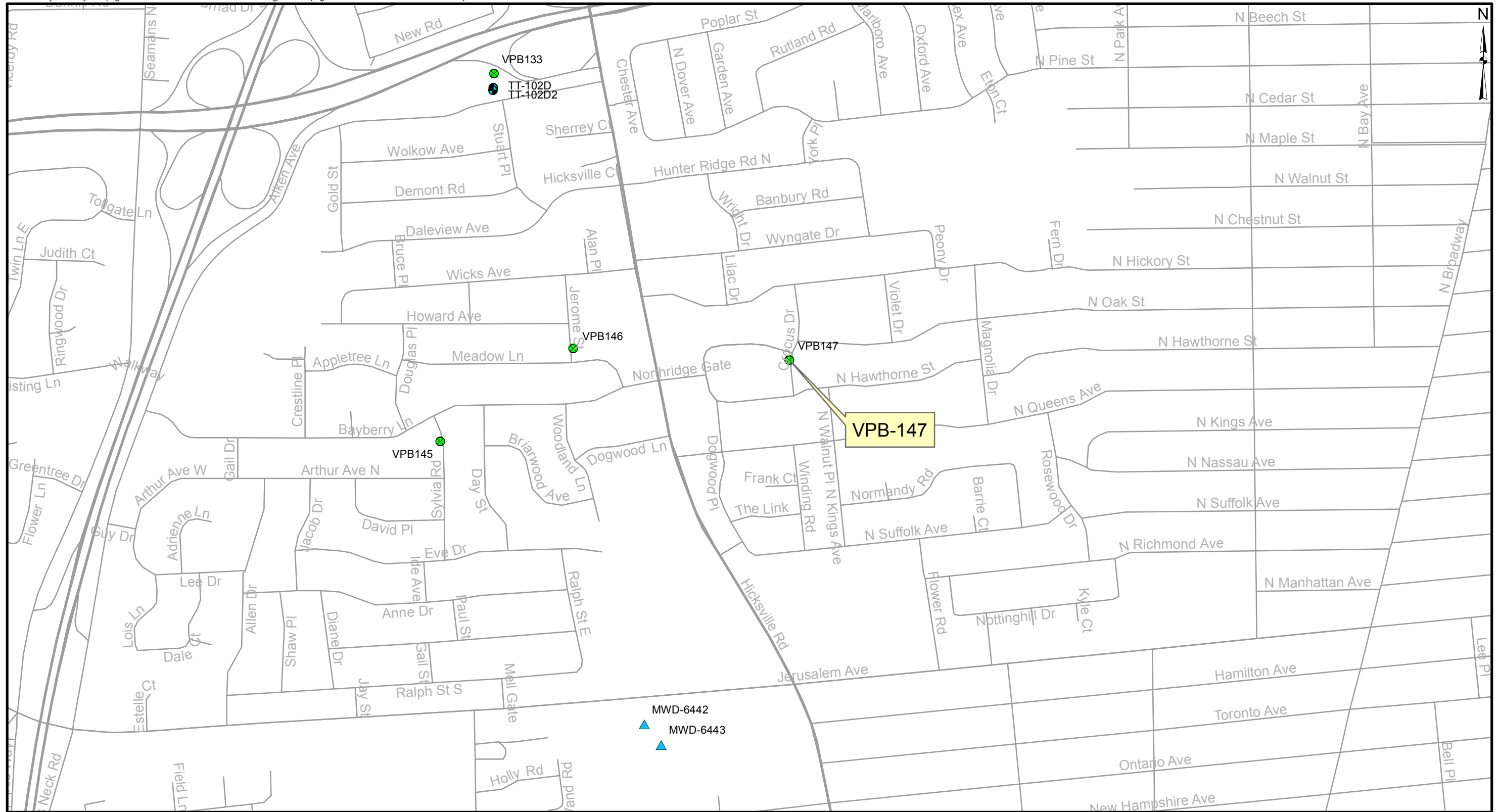
BORING	BORING START DATE	BORING COMPLETION DATE	GROUND ELEVATION (MSL)	TOTAL DEPTH (ft bgs)	SURFACE CASING SET AT (ft bgs)	NO. OF SPOON SAMPLES	GAMMA LOG (ft bgs)	NO. GW SAMPLES COLLECTED/ ATTEMPTED	TOC SAMPLES	DATE OF AIR SAMPLE	MONITORING WELLS INSTALLED AT LOCATION
VPB 147	5/29/2014	7/14/2014	44.92	1030	52.5	16	1028	40/49	1 (283 - 285 ft bgs)	6/24/2014	Pending

## Figures



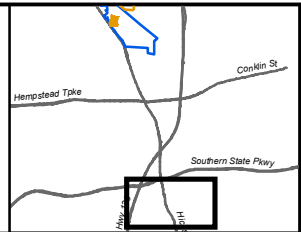
GENERAL LOCATION MAP  
NWIRP BETHPAGE  
BETHPAGE, NEW YORK

CONTRACT NUMBER N62470-11-D-8013		CTO NUMBER WE15	
APPROVED BY ---		DATE ---	
APPROVED BY ---		DATE ---	
FIGURE NO. 1			REV 0



- Vertical Profile Boring
- Monitoring Well
- Vertical Profile Boring - Proposed
- ▲ Water Supply Well
- ◆ Extraction Well

0    300    600    1,200  
Feet



**VPB 147 LOCATION MAP**  
**OPERABLE UNIT 2 (SITE 1)**  
**NAVAL WEAPONS INDUSTRIAL RESERVE PLANT**  
**BETHPAGE, NEW YORK**

CONTRACT NUMBER N62470-11-D-8013	CTO NUMBER WE15
APPROVED BY PS	DATE 12/23/2014
APPROVED BY	DATE
FIGURE NO. <b>2</b>	REV 1



## **Appendix A**

**VPB 147**

**Section 1**

**VPB 147 Boring and Gamma Logs**

<b>Client:</b> Department of the Navy, Naval Facilities Engineering Command, Mid-Atlantic			<b>Logged By:</b> V. Thayer		
<b>Location:</b> Crocus Dr. and Sheep Pasture Ln., Massapequa, NY		<b>Northing:</b> 193658.09		<b>Easting:</b> 1128979.9	
<b>Project #:</b> 60266526		<b>Ground Elevation (ft amsl):</b> 44.92		<b>Drilling Company:</b> Delta Well & Pump	
<b>Start Date:</b> 5/29/2014		<b>Drilling Method:</b> Auger (0-50' bgs) Mud Rotary (>50' bgs)		<b>Well Screen Interval (ft):</b> NA	
<b>Finish Date:</b> 7/14/2014				<b>Water Level (ft):</b> NA	
				<b>Total Depth (ft):</b> 1030.0	

Mud Rotary Drilling Note: Unless denoted by a splitspoon sample (indicated by the presence of a PID reading), boundaries between strata are approximate only and may be transitional because they are based on screened wash samples collected during mud rotary drilling at 5 ft. intervals.

DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
0					Upper Glacial			Top Soil
2						SM		Brown SILTY SAND
4						SM		
6								Brownish yellow (10 YR 6/6) well graded fine to coarse Sand, trace fine gravel
8						SW		
10								
12								
14						SM		Yellowish brown (10 YR 5/4) SILTY SAND, subrounded to subangular fine to coarse Sand, little silt, trace gravel
16								
18						SM		Yellowish brown (10 YR 5/4) SILTY SAND, fine to coarse Sand, little silt, fine to coarse gravel
20								Strong brown (7.5 YR 5/6) poorly graded SAND with Gravel, medium sand, little coarse sand, little subrounded fine to coarse gravel, trace silt
22						SP		
24								
26								
28						SM		Light brownish yellow (10 YR 6/6) SILTY SAND with Gravel, fine to coarse gravel, little silt
30								Brownish yellow (10 YR 6/6) well graded fine to coarse SAND, few subrounded fine gravel, trace silt
32						SW		
34								
36								Brownish yellow (10 YR 6/6) well graded SAND with Gravel, medium sand, some coarse sand, little subrounded gravel
38						SW		
40								
42								Brownish yellow (10 YR 6/6) well graded SAND with Gravel, medium sand, some coarse sand, little subrounded gravel
44						SW		
46								Light yellowish brown (10 YR 6/4) well graded SAND, little fine Gravel, trace subrounded coarse gravel, few silt
48						SW-SM		
50								
52						SW		Light yellowish brown (10 YR 6/4) well graded SAND, few subrounded fine Gravel
54						GP		

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DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
54	30 60 90				Upper Glacial			
56						GP		Yellow (10 YR 7/6) and very pale brown (10 YR 8/4) well graded GRAVEL, rounded fine Gravel, trace coarse gravel (continued)
58								
60			< 0.50	< 0.50		GW		Yellow (10 YR 7/6) and very pale brown (10 YR 8/4) well graded GRAVEL, subrounded fine Gravel, some subrounded to rounded coarse gravel
62								
64						GW		Yellow (10 YR 7/6) and very pale brown (10 YR 8/4) well graded GRAVEL, subrounded fine Gravel, some subrounded to rounded coarse gravel
66								
68								
70						SM		Grayish brown (10 YR 5/2) SILTY SAND, fine Sand, few coarse sand, fine gravel
72								
74					CL		Dark yellowish brown (10 YR 4/4) SANDY CLAY, Clay is dry and banding observed in nodules that came up in mud, few small gravel	
76								
78					CL		Dark yellowish brown (10 YR 4/4) SANDY CLAY with Gravel, some fine to coarse sand, little fine to coarse subrounded gravel	
80								
82					SC		Dark yellowish brown (10 YR 4/4) CLAYEY SAND	
84								
86					SC		Grayish brown (10 YR 5/2) CLAYEY SAND, Lignite, few subrounded fine gravel, one orange band of clay	
88								
90					SP-SM		Dark grayish brown (10 YR 4/2) poorly graded SAND with Silt, fine sand, trace fine gravel	
92								
94								
96								
98								
100			< 0.50	< 0.50				
102					Magothy	SM/CL		Grayish brown (10 YR 5/2) SILTY SAND, fine to coarse Sand, little silt, interbedded clay layer
104								
106					SM		Dark gray SILTY SAND with Lignite	
108								
110					SW-SM		Very dark gray (10 YR 3/1) well graded fine to medium SAND, trace coarse sand, some lignite	
112								
114					SM		Gray SILTY SAND	

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DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
116	30 60 90				Magothy			Gray SILTY SAND ( <i>continued</i> )
118						SM		Very dark grayish brown (10 YR 3/2) SILTY SAND, fine to medium Sand, trace coarse sand, lignite
120						SM		
122								
124						PT/SM		Dark gray (7.5 YR 4/1) LIGNITE microlaminated with silty fine Sand
126								
128								
130						SM		Dark gray (7.5 YR 4/1) SILTY SAND, fine to medium Sand, lignite, muscovite, some silt
132								
134						SM		Dark gray (7.5 YR 4/1) SILTY SAND, fine to medium Sand, lignite, muscovite, some silt
136								
138								
140						ML/PT		Very dark gray (Gley 1 3/1) SANDY SILT, fine to medium Sand, trace coarse sand, some lignite, nodules of lignite in friable layers
142								
144						SM/PT		Dark gray (5 Y 4/1) SILTY SAND, medium to coarse Sand, little fine sand, little silt, some black lignite
146								
148								
150			< 0.50	< 0.50		SP-SM	Very dark gray (10 YR 4/1) medium SAND, little coarse Sand, few fine sand, trace subrounded fine gravel, lignite, pyrite concretion	
152								
154						SP-SM	Very dark gray (10 YR 4/1) medium SAND, little coarse Sand, few fine sand, trace subrounded fine gravel, lignite, pyrite concretion	
156								
158								
160		0.1				SP-SM	Light brownish gray (2.5 Y 6/2) poorly graded SAND with Silt, angular medium sand, little fine sand, few silt; (2 bands of 1/4" orange stained sand)	
162								
164						SW	Dark gray (7.5 YR 4/1) well graded fine to coarse SAND, trace fine subrounded gravel, pyrite concretions	
166								
168								
170						SW	Dark gray (7.5 YR 4/1) well graded angular medium SAND, little fine sand, little coarse sand	
172								
174								
176						SM	Gray SILTY SAND	

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DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
178					Magothy			
180						SM		Gray (7.5 YR 4/1) SILTY SAND, fine to medium Sand, little silt, 1 pyrite concretion, trace angular coarse sand, lignite
182								
184						SP-SM		Gray (Gley 1 6/1) poorly graded SAND with Silt, angular medium sand, little fine sand, few silt
186								
188						SP-SM		Dark gray (7.5 YR 4/1) poorly graded SAND with Silt, angular medium sand, few silt, trace coarse sand
190								
192								
194								
196						SM		Gray SILTY SAND
198								
200			< 0.50	< 0.50		SM/ML		Gray (7.5 YR 5/1) SILTY SAND interbedded with Silt, fine sand, lignite, microlaminated clay, silt
202								
204						ML		Dark gray (Gley 1 4/1) SANDY SILT, fine Sand, little medium sand, lignite, silt
206								
208					ML		Gray (7.5 YR 5/1) SANDY SILT, fine to medium Sand, lignite, silt	
210								
212								
214					SM		Dark gray (7.5 YR 4/1) SILTY SAND, angular medium Sand, little fine sand, some silt	
216								
218								
220			< 0.50	< 0.50	SM		Very dark gray SILTY SAND, fine to coarse Sand, lignite, some silt	
222								
224					SM		Gray (Gley 1 5/1) SILTY SAND, angular medium Sand, little fine sand, little silt	
226								
228								
230					SP-SM		Dark gray (10 YR 4/1) poorly graded SAND with Silt, medium sand, few silt, lignite	
232								
234					SP-SM		Dark gray (10 YR 4/1) poorly graded SAND with Silt, medium sand, few silt, lignite, thinly laminated lignite and fine sand	
236								
238			< 0.50	< 0.50	SM			

(Continued Next Page)

DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION		
240			< 0.50	< 0.50	Magothy	SM		Dark gray (7.5 YR 4/1) SILTY SAND, angular medium Sand, little fine sand, little silt or clay, interbedded lignite, thinly laminated (continued)		
242										Dark gray (7.5 YR 4/1) SILTY SAND
244										
246							SM			
248										
250							ML		Dark gray (7.5 YR 4/1) SANDY SILT, angular medium Sand, little fine sand, trace coarse sand, silt, lignite	
252										
254							ML		Dark gray (7.5 YR 4/1) SANDY SILT, angular medium Sand, little fine sand, trace coarse sand, silt, lignite	
256										
258										
260			< 0.50	< 0.50			SP-SM		Dark gray (7.5 YR 4/1) poorly graded SAND with Silt, medium sand, trace coarse sand, lignite, few nodules of thinly laminated sand and silt	
262										
264							SM		Dark gray (7.5 YR 4/1) SILTY SAND, angular medium Sand, some silt, trace coarse sand	
266										
268							SM		Dark gray (7.5 YR 4/1) SILTY SAND, angular medium Sand, little silt, lignite	
270										
272						SM				
274										
276						ML		Dark gray (7.5 YR 4/1) SANDY SILT, fine to medium Sand, trace coarse sand, lignite.		
278										
280			< 0.50	< 0.50		SM		Gray (Gley 1) angular fine to coarse SAND, little Silt, lignite; thinly laminated lignite and fine sand		
282										
284		0				SM		Gray (Gley 1) poorly graded SAND with Silt, fine to medium sand, few silt, 2 bands of lignite, 1/2" of clay or silt		
286						SP-SM				
288										
290						SP-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, medium sand, little fine sand, silt		
292										
294						SP-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, lignite, few silt, pyrite, nodules of thinly laminated fine sand and lignite		
296										
298						SP-SM				
300			< 0.50	< 0.50		SP-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, lignite, thinly laminated fine sand, lignite		

(Continued Next Page)

DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
302					Magothy	SP-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, lignite, thinly laminated fine sand, lignite (continued)
304						SM		Gray (7.5 YR 5/1) SILTY SAND, subangular to angular medium Sand, little lignite, little silt
306						SM		Gray (Gley 1 5/1) SILTY SAND, angular fine to medium Sand, little silt, little lignite
308						SM		Dark gray (Gley 1 4/1) SILTY SAND, angular medium Sand
310						SM		Dark gray (7.5 YR 4/1) SILTY SAND, fine to medium Sand, little silt
312						SM		Very dark gray (Gley 1 4/0) well graded fine to coarse SAND, lignite, pyrite
314						SM		Very dark gray (Gley 1 4/1) well graded SAND, fine to coarse Sand, few silt, pyrite
316						SM		Dark gray SILTY SAND
318						SM		Dark gray (Gley 4/0) SILTY SAND, medium Sand, little fine sand, lignite, little silt
320						SM		SILTY SAND, subangular medium Sand, few fine sand, trace coarse sand, lignite
322						SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, angular medium sand, little fine sand, few silt, lignite
324						SW-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, angular medium sand, little fine sand, few silt, lignite
326						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
328						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
330						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
332						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
334						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
336						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
338						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
340						SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt
342	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
344	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
346	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
348	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
350	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
352	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
354	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
356	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
358	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
360	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					
362	SW-SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, pyrite, lignite, little silt					

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DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION	
364					Magothy			Dark gray (Gley 1 4/1) poorly graded SAND with Silt, angular medium sand, little fine sand, few coarse sand, lignite, few silt, pyrite	
366						SP-SM			SILTY SAND, fine to medium Sand, little silt, lignite
368									
370									
372							SM		
374									
376							SM		Gray (Gley 1 5/1) SILTY SAND, fine to medium Sand, some silt, little lignite interbedded fine sand and lignite
378									
380			< 0.50	< 0.50			SM		Gray (Gley 1 5/1) SILTY SAND, subangular fine to coarse Sand, some lignite, trace pyrite, little silt
382									
384		0.1					SP-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt, fine to medium sand, 3 bands of lignite, 1/4" thick, spaced 1.5 to 2" apart
386									
388							SM		Gray (Gley 1 5/1) SILTY SAND, fine to medium Sand, few angular coarse sand, pyrite, lignite
390									Silty Sand interbedded lignite, clay.
392							SM/CL		
394									
396							SM		Dark gray (Gley 1 4/0) SILTY SAND, fine to medium Sand, trace coarse sand, lignite, pyrite, some silt, interbedded lignite, sand, silt
398									
400			< 0.50	< 0.50					
402							ML		Gray (Gley 1 5/1) SANDY SILT, fine to medium Sand, trace coarse sand, thinly laminated lignite, sand
404									
406						SM/CL		Gray (Gley 1 5/1) fine SAND interbedded with Lignite, thinly laminated, interbedded clay or silt	
408									
410						SM/CL		Gray (Gley 5/1) SILTY SAND, angular fine to medium Sand, laminated lignite, fine sand, microlayers of silt or clay	
412									
414									
416						SM		Gray (Gley 1 5/1), SILTY SAND, fine to medium Sand, trace coarse sand, lignite, some silt	
418									
420			< 0.50	< 0.50					
422						SM/ML		Gray (Gley 1 5/1), SILTY SAND, microlaminated with Lignite and interbedded microlayers of silt	
424						SM			

(Continued Next Page)

DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
426					Magothy	SM		Dark gray (Gley 1 4/0) SILTY SAND, angular medium Sand, few coarse sand, little silt, lignite ( <i>continued</i> )
428						SP-SM		Dark gray (Gley 1 4/1) poorly graded SAND with Silt
430						SP-SM		Dark gray (Gley 1 4/1) poorly graded SAND with Silt, angular medium sand, little fine sand, few silt
432						SP-SM		Dark gray (Gley 1 4/1) SANDY SILT
434						ML		Dark gray (10 YR 4/1) SILTY SAND, subangular fine to medium Sand, muscovite flakes, 1 (1/4") layer of lignite
436						SM		Dark gray (10 YR 4/1) SILTY SAND
438						SM		Dark gray (10 YR 4/1) SILTY SAND, fine to medium Sand, some lignite
440			< 0.50	< 0.50		SM/ML		SANDY SILT
442						ML		SILTY SAND, microlaminated fine Sand, lignite, pyrite concretion
444		0				SM		Very dark gray (Gley 1 3/1) SANDY SILT, fine Sand, Lignite, microlaminated fine sand and lignite, pyrite concretion
446						ML		Dark gray (Gley 1 4/1) SANDY SILT, nodules of Clay/Silt, lignite
448						ML		Dark gray (Gley 1 4/1) SANDY SILT, nodules of Clay/Silt, lignite
450						SM		Gray (2.5 Y 5/1) SILTY SAND, fine to medium Sand, lignite, some laminated fine sand and silt
452						SM		Gray SILTY SAND, fine to medium Sand, lignite, trace coarse sand
454						SM		
456						SM		
458						SM		
460			< 0.50	< 0.50		SM		
462						SM		
464						SM		
466					SM			
468					SM			
470					SM			
472					SM			
474					SM			
476					SM			
478					SM			
480			0.45	< 0.50	SM			
482					SM			
484					SM			
486					SM			

(Continued Next Page)

DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
486	30 60 90				Magothy			
488						SM		Gray SILTY SAND, fine to medium Sand, lignite, trace coarse sand <i>(continued)</i>
490						SM		Dark gray (10 YR 4/1) SILTY SAND, fine to coarse Sand, lignite, some silt
492						SM		Dark gray SILTY SAND
494						SM		Dark gray SILTY SAND
496						SM		Dark gray SILTY SAND
498						SM		Dark gray SILTY SAND
500						SM		Gray (2.5 Y 6/1) SILTY SAND, fine Sand, lignite, thinly laminated fine sand and lignite
502						SM		Gray (2.5 Y 6/1) SILTY SAND, fine Sand, lignite, thinly laminated fine sand and lignite
504		0				SM		Gray (2.5 Y 6/1) SILTY SAND, fine Sand, 3 bands of lignite, 1/4" thick interbedded with fine sand
506						SM		Gray SILTY SAND, fine to medium Sand, little silt, lignite, trace coarse sand, pyrite concretion
508						SM		Gray SILTY SAND, fine to medium Sand, little silt, lignite, trace coarse sand, pyrite concretion
510						SM		Gray SILTY SAND, fine to medium Sand, little silt, lignite, trace coarse sand, pyrite concretion
512						CL		Dark gray (Gley 1 4/1) SANDY CLAY, some fine to medium Sand
514						CL		Dark gray (Gley 1 4/1) SANDY CLAY, some fine to medium Sand
516						CL		Dark gray (Gley 1 4/1) SANDY CLAY, some fine to medium Sand
518						SM/CL		Dark gray (Gley 1 4/1) SANDY CLAY, nodules of Silty fine Sand, lignite and clay, microlaminated
520					SM/CL		Dark gray (Gley 1 4/1) SANDY CLAY, nodules of Silty fine Sand, lignite and clay, microlaminated	
522					SM/CL		SILTY SAND, fine Sand, laminated with lignite, clay	
524					SM/CL		SILTY SAND, fine Sand, laminated with lignite, clay	
526					SM/CL		SILTY SAND, fine Sand, laminated with lignite, clay	
528					SM/CL		SILTY SAND, fine Sand, laminated with lignite, clay	
530			< 0.50	< 0.50	SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay	
532					SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay	
534					SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay, angular medium sand, little fine sand, lignite, trace fine to few silt, (nodules of laminated fine sand and lignite)	
536					SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay, angular medium sand, little fine sand, lignite, trace fine to few silt, (nodules of laminated fine sand and lignite)	
538					SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay, angular medium sand, little fine sand, lignite, trace fine to few silt, (nodules of laminated fine sand and lignite)	
540			< 0.50	< 0.50	SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay, angular medium sand, little fine sand, lignite, trace fine to few silt, (nodules of laminated fine sand and lignite)	
542					SP-SC		Dark gray (10 YR 4/1) poorly graded SAND with Clay, angular medium sand, little fine sand, lignite, trace fine to few silt, (nodules of laminated fine sand and lignite)	
544					SP-SC		Dark gray (7.5 YR 4/1) poorly graded SAND with Clay, angular medium sand, few silt	
546					SP-SC		Dark gray (7.5 YR 4/1) poorly graded SAND with Clay, angular medium sand, few silt	

(Continued Next Page)

DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
548	30 60 90				Magothy			
550						SP-SM		Dark gray (7.5 YR 4/1) poorly graded SAND with Silt, angular medium sand, lignite, trace silt
552								
554								
556						SP		Dark gray (7.5 YR 4/1) poorly graded SAND, angular medium sand, lignite, trace silt
558								
560			< 0.50	< 0.50				
562						SP		Dark gray (7.5 YR 4/1) angular medium SAND, little coarse Sand, trace silt
564								
566						SP		Gray (Gley 1 5/1) poorly graded SAND, angular medium Sand
568								
570						SM		Dark gray (Gley 1) SILTY SAND
572								
574								
576						SP-SM		Gray (Gley 1 6/1) poorly graded SAND with Silt, subangular medium sand, trace coarse sand, few silt
578								
580			< 0.50	< 0.50				
582					SP-SM		Gray (Gley 1 6/1) poorly graded SAND with Silt, subangular medium sand, trace coarse sand, few silt; microlaminated, silt, sand, lignite	
584								
586					SM		Gray (Gley 1 5/1) SILTY SAND, medium Sand, little fine sand, little silt, nodules of thinly laminated silt, sand, lignite	
588								
590								
592					SP-SC		Gray (Gley 1 6/1) poorly graded SAND with Clay, angular medium sand, few coarse sand, trace silt	
594								
596					SM		Gray (Gley 5/1) SILTY SAND, subangular medium Sand, little silt	
598								
600			< 0.50	< 0.50				
602					SP-SM		Gray (Gley 5/1) poorly graded SAND with Silt; angular medium to coarse sand, few silt, interbedded fine lignite seams	
604								
606					SP-SM		Gray (Gley 5/1) poorly graded SAND with Silt; angular medium to coarse sand, few silt, interbedded fine lignite seams	
608								
					SM		Gray (10 YR 5/1) SILTY SAND, fine Sand; some Silt	

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DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION	
610					Magothy	SM		Gray (10 YR 5/1) SILTY SAND, fine Sand; some Silt <i>(continued)</i>	
612				SM					
614				SW-SM					
616									
618									
620			< 0.50	< 0.50		SP-SM			Gray (7.5 YR 5/1) poorly graded SAND with Silt, angular medium sand, few coarse sand, few silt, thinly laminated fine sand, lignite
622									
624									Dark gray (10 YR 4/1) poorly graded SAND, angular medium Sand, trace silt
626						SP			
628									
630						SM			Gray (7.5 YR 5/1) SILTY SAND, angular medium Sand, lignite, little silt
632									
634									Gray (7.5 YR 5/1) SILTY SAND
636						SM			
638									
640			< 0.50	< 0.50		SM			Gray SILTY SAND medium to coarse Sand, little silt
642									
644						SM			Dark gray (10 YR 4/1) SILTY SAND, medium Sand, little coarse sand, few silt
646						SM			
648									
650					SC		Dark gray (10 YR 4/1) CLAYEY SAND, Pyrite		
652									
654					SM		Gray (10 YR 5/1) SILTY SAND, medium Sand, little coarse sand, little silt		
656					SM				
658									
660					SM		Dark gray (10 YR 4/1) SILTY SAND, fine to medium Sand, pyrite concretion, some silt		
662									
664		0			SM		Dark gray (10 YR 4/1) SILTY SAND, fine to medium Sand, pyrite concretion, some silt		
666									
668									
670			< 0.50	< 0.50	SM		Dark gray (10 YR 4/1) SILTY SAND, fine to coarse Sand, lignite, trace gravel, some silt		

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DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
672					Magothy	SM		Dark gray (10 YR 4/1) SILTY SAND, fine to coarse Sand, lignite, trace gravel, some silt <i>(continued)</i>
674						ML		Dark gray (10 YR 4/1) SANDY SILT, fine to medium Sand
676								
678								
680			< 0.50	< 0.50		ML		Dark gray (10 YR 4/1) SANDY SILT, fine to medium Sand
682						ML		
684						SM		Gray (7.5 YR 5/1) SILTY SAND, fine to medium Sand, some silt, lignite, interbedded fine sand and lignite, microlaminated
686						SM		
688						SM		Gray (7.5 YR 5/1) SILTY SAND, medium Sand, trace coarse sand, lignite, little to some silt
690						SM		
692						SM		Gray (7.5 YR 5/1) SILTY SAND, medium Sand, trace coarse sand, lignite, little to some silt
694						SM		
696						SM		Gray (7.5 YR 5/1) SANDY SILT
698						ML		
700						ML		Gray (7.5 YR 5/1) SANDY SILT
702						SP-SM		Gray (7.5 YR 6/1) SAND with Silt, angular medium sand, few to little silt
704						ML		
706					ML		Gray (7.5 YR 5/1) SANDY SILT, angular medium Sand, trace coarse sand, little silt, thin strip of clay or silt (nodules)	
708					ML			
710					SM		Gray (7.5 YR 5/1) SILTY SAND, medium to coarse Sand, little fine sand, little silt, trace gravel	
712					SM			
714					SM		Gray (7.5 YR 5/1) SILTY SAND, medium to coarse Sand, little fine sand, little silt, trace gravel	
716					SM			
718					SP-SM		Gray (Gley 1 5/1) poorly graded SAND with Silt; medium sand, little coarse sand, lignite	
720			< 0.50	< 0.50	SP-SM			
722					SP-SM		Gray (7.5 YR 5/1) well graded medium to coarse SAND, little fine sand, trace silt	
724					SW			
726					SW		Gray (7.5 YR 5/1) well graded medium to coarse SAND, little fine sand, few silt	
728					SW-SM			
730					SW-SM		Gray (7.5 YR 5/1) well graded medium to coarse SAND, little fine sand, few silt	
732					SW-SM			

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DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
734	30 60 90				Magothy			Gray (7.5 YR 5/1) well graded SAND with Silt, subangular medium to coarse sand, little fine sand, few silt, trace subangular gravel
736						SW-SM		
738								
740			< 0.50	< 0.50		SW-SM		Gray (7.5 YR 5/1) well graded SAND with Silt, subangular medium to coarse sand, little fine sand, few silt
742								
744						SW-SM		Gray (Gley 1 5/1) well graded SAND with Gravel and Silt, subangular to angular fine gravel, fine to coarse sand, few silt, laminated fine sand and lignite
746								
748								
750						GP-GM		Gray (7.5 YR 6/1) poorly graded GRAVEL with Silt and Sand, subangular fine gravel, fine to coarse sand, few silt
752								
754						SP		Gray (7.5 YR 6/1) poorly graded SAND, angular medium Sand, little coarse sand
756								
758								
760			< 0.50	< 0.50		SP-SM		Gray (7.5 YR 6/1) poorly Graded SAND with Silt and gravel, angular medium sand, trace coarse sand, little fine gravel, few silt
762								
764						SP-SC		Gray (7.5 YR 6/1) poorly Graded SAND with Clay, angular medium sand, trace coarse sand, few clay
766								
768								
770						SP-SM		Gray (7.5 YR 6/1) poorly graded SAND with Silt
772								
774						SW-SM		Gray (7.5 YR 6/1) well graded SAND with Silt and gravel, fine to coarse sand, little subangular fine gravel, few silt
776								
778						SW-SM		Gray (7.5 YR 6/1) well graded SAND with Silt and gravel, fine to coarse sand, little subangular fine gravel, few silt
780								
782								
784						GW-GM		Gray (7.5 YR 5/1) poorly graded GRAVEL with Sand, fine subrounded gravel, some fine to coarse sand, few silt
786								
788								
790		0.1				SM		Gray (7.5 YR 5/1) SILTY SAND, angular medium Sand, little silt
792						SM		Gray (7.5 YR 5/1) SILTY SAND
794						SM		

(Continued Next Page)

DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
796					Magothy	SM		Gray SILTY SAND with Gravel, coarse sand, little fine gravel, silt or clay ( <i>continued</i> )
798						GC		CLAYEY GRAVEL with Sand, fine subrounded gravel, some fine to coarse sand, clay
800		< 0.50	< 0.50					
802						SC		Gray (7.5 YR 6/1) CLAYEY SAND, medium to coarse Sand, trace fine gravel, little to some clay, silt
804								
806						GC		Gray (7.5 YR 6/1) CLAYEY GRAVEL, fine Gravel, little silt, clay
808								
810						SM		Light gray (10 YR 7/1) SILTY SAND and Gravel, fine to coarse sand, little subangular fine gravel, pyrite, little silt
812								
814						SP		Gray (7.5 YR 6/1) poorly graded SAND, angular medium Sand, trace coarse sand
816		< 1.0	< 1.0					
818						SM		Gray (7.5 YR 6/1) SILTY SAND, medium to coarse Sand, trace small gravel, little silt
820								
822						SC		Light gray (7.5 YR 7/1) CLAYEY SAND, medium to coarse Sand, little clay
824								
826						SM		Light gray (7.5 YR 7/1) SILTY SAND, angular medium to coarse Sand, some silt
828								
830					SM		Light gray (7.5 YR 7/1) SILTY SAND	
832		< 0.50	< 0.50					
834					SP-SM		Light gray (7.5 YR 7/1) angular medium to coarse SAND, few to little Silt	
836								
838					SM		Light gray (7.5 YR 7/1) SILTY SAND, medium to coarse Sand, little silt	
840								
842					ML		Light gray (7.5 YR 7/1) SANDY SILT, fine to coarse Sand, little white silt	
844								
846								
848								
850								
852								
854								
856								

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DEPTH (ft)	Gamma Ray 30 60 90	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
858					Magothy	ML		
860			< 0.50	< 0.50		SC		Gray (7.5 YR 5/1) CLAYEY SAND, angular fine to coarse Sand, some white silt
862								
864						SP-SM		Gray (7.5 YR 5/1) poorly graded angular medium SAND, few Silt, clay
866								
868						SM		Gray (7.5 YR 5/1) SILTY SAND, angular medium Sand, trace coarse sand, few silt or clay
870								
872						ML/CL		Gray (7.5 YR 5/1) SANDY CLAY, Silt
874								
876						ML/CL		Gray (7.5 YR 5/1) SANDY CLAY, angular medium Sand, little fine sand, little coarse sand, clay
878			< 2.0	< 2.0				
880						SC		Gray (7.5 YR 5/1) CLAYEY SAND, angular medium Sand, little coarse sand, little fine sand, little white clay, silt
882								
884						SM		Light gray (Gley 1 7/1) SILTY SAND, fine Sand
886			< 0.50	< 0.50				
888						SM/CH		Light gray (Gley 1 7/1) SILTY SAND, fine Sand, some silt, interbedded laminated gray clay
890		0						
892					SM/CH		Gray (Gley 6/1) SILTY SAND, fine to medium Sand, silty sand microlaminated with lignite	
894								
896					SM/CH		Gray (Gley 6/1) SILTY SAND, fine to medium Sand, silty sand microlaminated with lignite	
898								
900			< 1.0	< 1.0				
902					ML		Gray (Gley 6/1) SANDY SILT, some angular fine to medium Sand, trace coarse sand, silt	
904								
906					ML/CH		Gray (5 Y 6/1) SANDY SILT, fine to coarse Sand, silt interbedded with gray clay	
908								
910								
912								
914								
916								
918								

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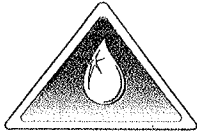
DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
918	30 60 90				Magothy			Gray (5 Y 6/1) SANDY SILT, fine to coarse Sand, silt interbedded with gray clay, trace gravel
920				ML/CH				Gray (5 Y 6/1) SANDY SILT, fine to coarse Sand, silt interbedded with gray clay, trace gravel
922				ML/CH				Gray (5 Y 6/1) SANDY SILT, medium Sand, some silt interbedded with laminated clay
924				ML/CH			Gray (10 YR 5/1) SILTY SAND, fine to medium Sand, some silt, trace coarse sand	
926				ML/CH			Gray (Gley 1 6/1) SILTY SAND, angular fine to medium Sand, trace coarse sand, little silt	
928				ML/CH			Gray (Gley 1 5/1) SILTY SAND, angular medium to coarse Sand, little silt	
930				SM			Gray (7.5 YR 5/1) SILTY SAND, fine Sand	
932				SM			Gray (Gley 1 6/1) SILTY SAND, fine Sand, micaceous, top 0.5" gray clay	
934				SM			Gray (Gley 1 6/1) SILTY SAND, angular fine to coarse Sand, trace fine gravel, little silt	
936				SM			Gray (Gley 1 5/1) SILTY SAND, angular fine to coarse Sand, little silt, lignite	
938			< 5.0	< 5.0		SM		Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
940						SM		
942						SM		
944			< 5.0	< 5.0		SM		
946						SM		
948					SM			
950					SM			
952					SM			
954					SM			
956					SM			
958					SM			
960			< 0.50	< 0.50	SM			
962					SM			
964		0			CH/SM			
966					CH/SM			
968					SM			
970					SM			
972					SM			
974					SM			
976					SM			
978					SM/CH			

(Continued Next Page)

DEPTH (ft)	Gamma Ray	PID (ppm)	TCE (ug/L)	PCE (ug/L)	Formation	USCS	GRAPHIC LOG	MATERIAL DESCRIPTION
980	30 60 90				Magothy			Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay <i>(continued)</i>
982				SM/CH				Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
984				SM/CH				Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
986				SM/CH				Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
988				SM/CH				Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
990				SM/CH				Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
992				SM/CH				Gray (Gley 1, 6/1) SILTY SAND microlaminated with Lignite, clay
994				SM				Gray (Gley 1 6/1) SILTY SAND, fine Sand, micaceous, little silt
996				SM				Gray (Gley 1 6/1) SILTY SAND, fine Sand, micaceous, little silt
998				CH				Gray fat CLAY, laminated
1000		0					Gray (Gley 1 6/1) SILTY SAND, fine Sand, micaceous, little silt. Forms sharp contact with overlying clay	
1002							Gray (Gley 1 6/1) SILTY SAND, fine Sand, micaceous, little silt. Forms sharp contact with overlying clay	
1004		0			Raritan			Gray (Gley 1 5/1) fat CLAY, laminated
1006				CH				Gray (Gley 1 5/1) fat CLAY, laminated
1008				CH				Gray (Gley 1 5/1) fat CLAY, laminated
1010		0		CH				Gray (Gley 1 5/1) fat CLAY, laminated
1012				CH				Gray (Gley 1 5/1) fat CLAY, laminated
1014		0		PT				Black (10 YR 2/1) LIGNITE and Clay, friable
1016				PT				Black (10 YR 2/1) LIGNITE and Clay, friable
1018				CH				Very dark gray (10 YR 3/1) CLAY, microlaminated
1020		0		CH				Very dark gray (10 YR 3/1) CLAY, microlaminated
1022				CH			Very dark gray (10 YR 3/1) CLAY, microlaminated	
1024		0		CH			Gray (7.5 YR 5/1) CLAY, laminated	
1026				CH			Gray (7.5 YR 5/1) CLAY, laminated	
1028				CH			Light gray (7.5 YR 7/1) with red mottling CLAY, laminated	
1030		0		CH			Light gray (7.5 YR 7/1) with red mottling CLAY, laminated	

End of boring at 1030.0 ft. bgs.

DOWN HOLE RUN



COMPANY: DELTA WELL & PUMP CO., INC.

LOCATION: NWIRP CROCUS DR.

Well: VPB-147

Depth Driller:

Depth Logger:

Date: 07/08/2014

Time:

Logged by: CMO

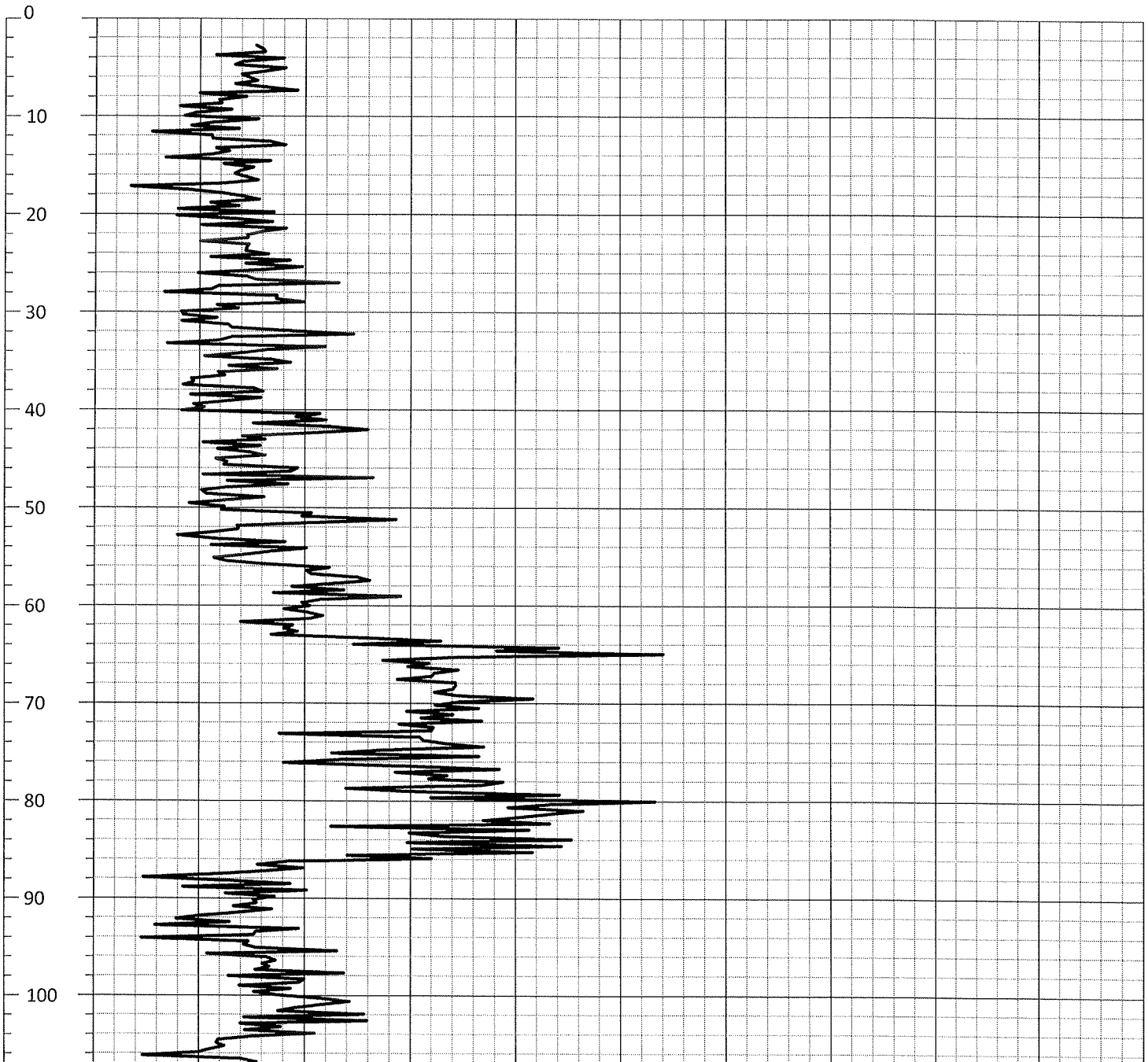
File Name: 739

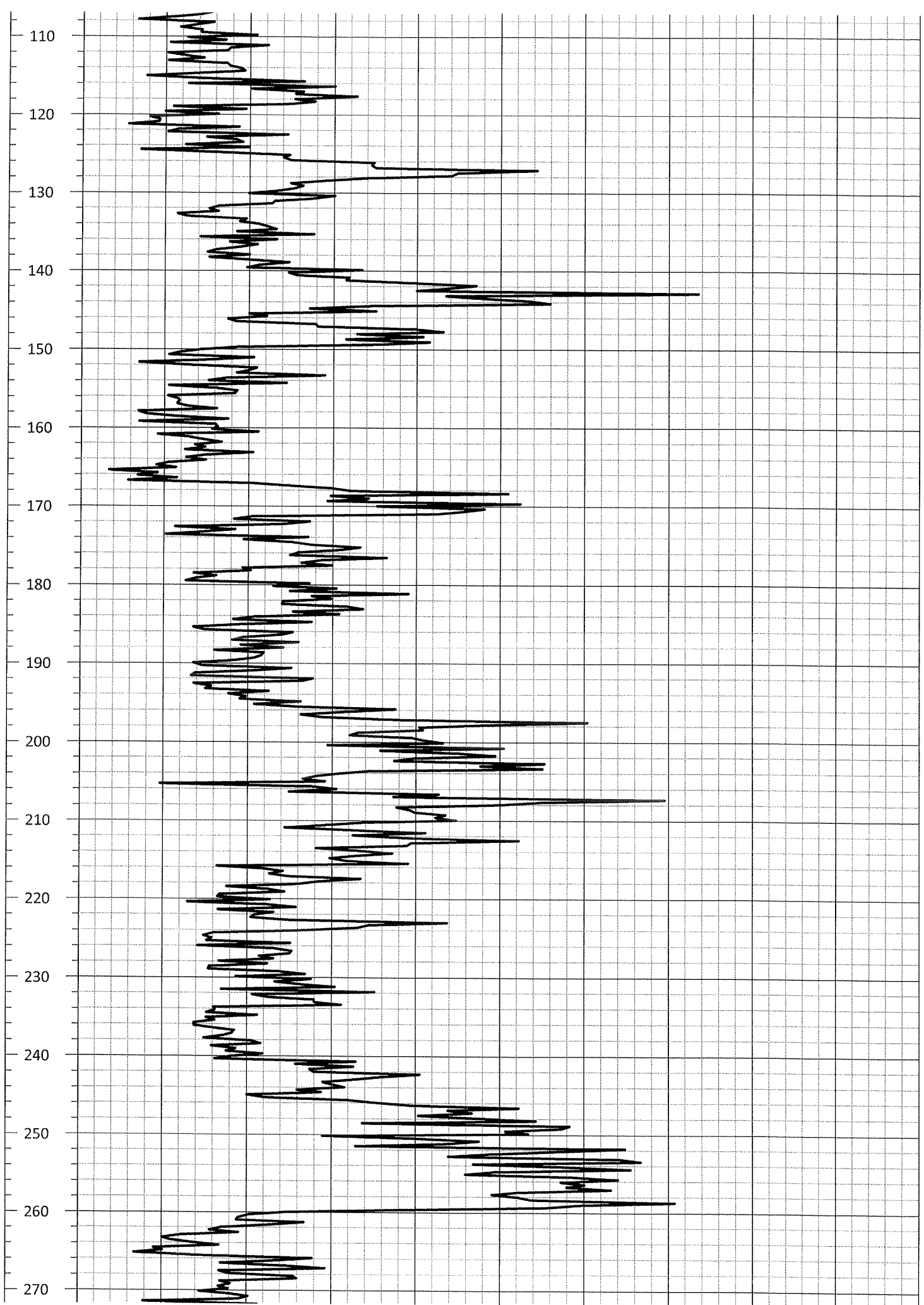
Witness: VAL

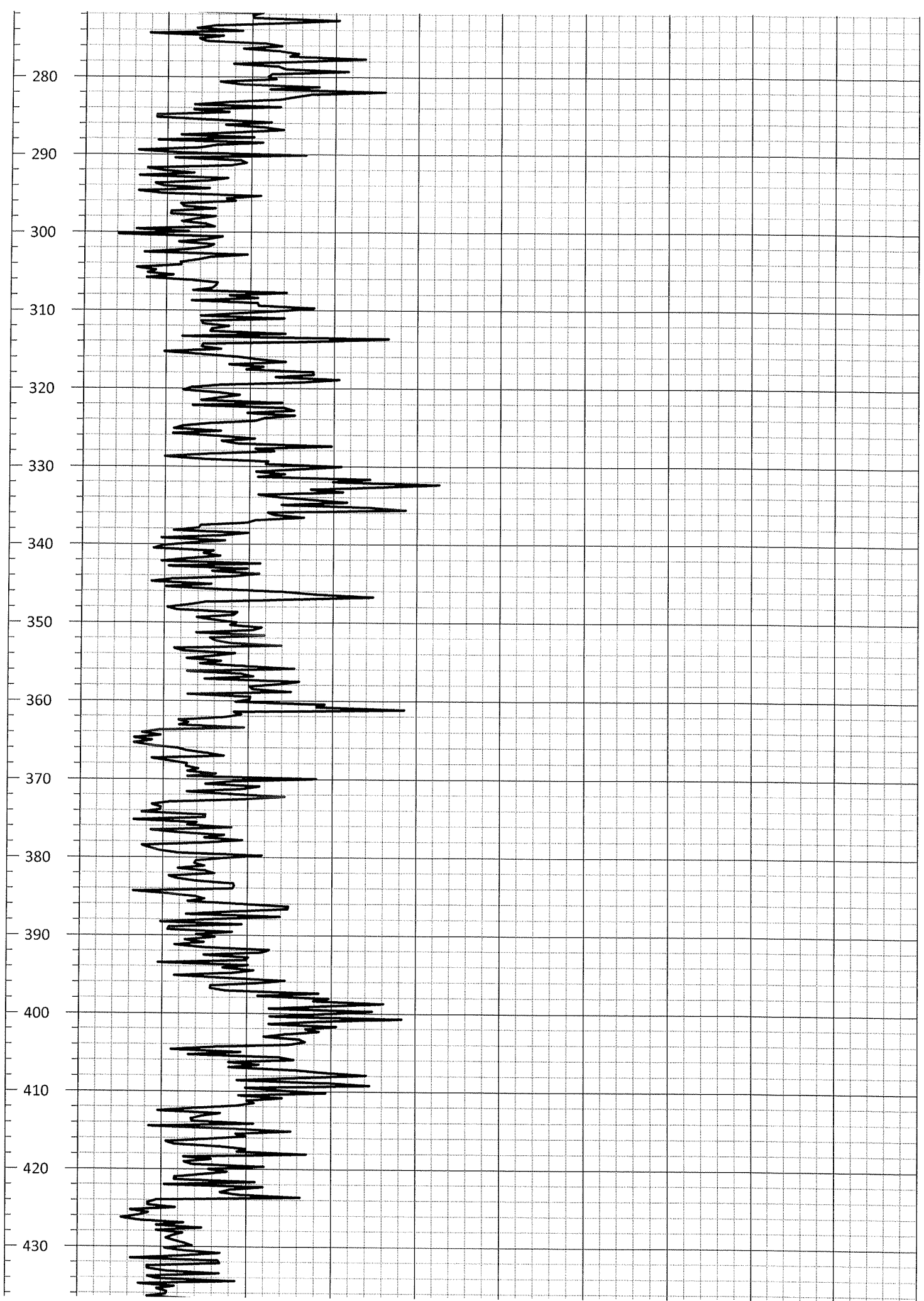
Depth (ft.) 0.0

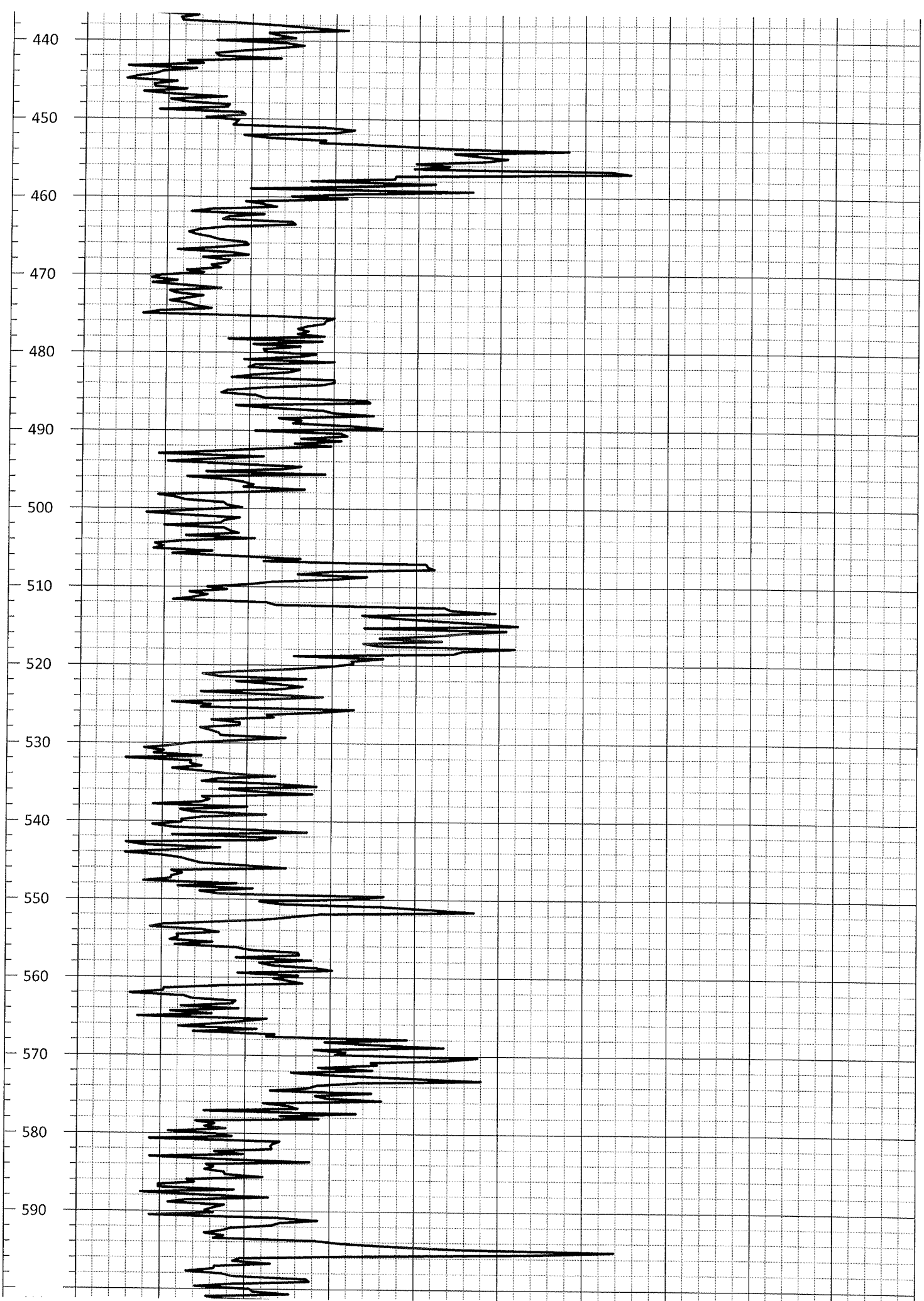
GAMMA  
(cps)

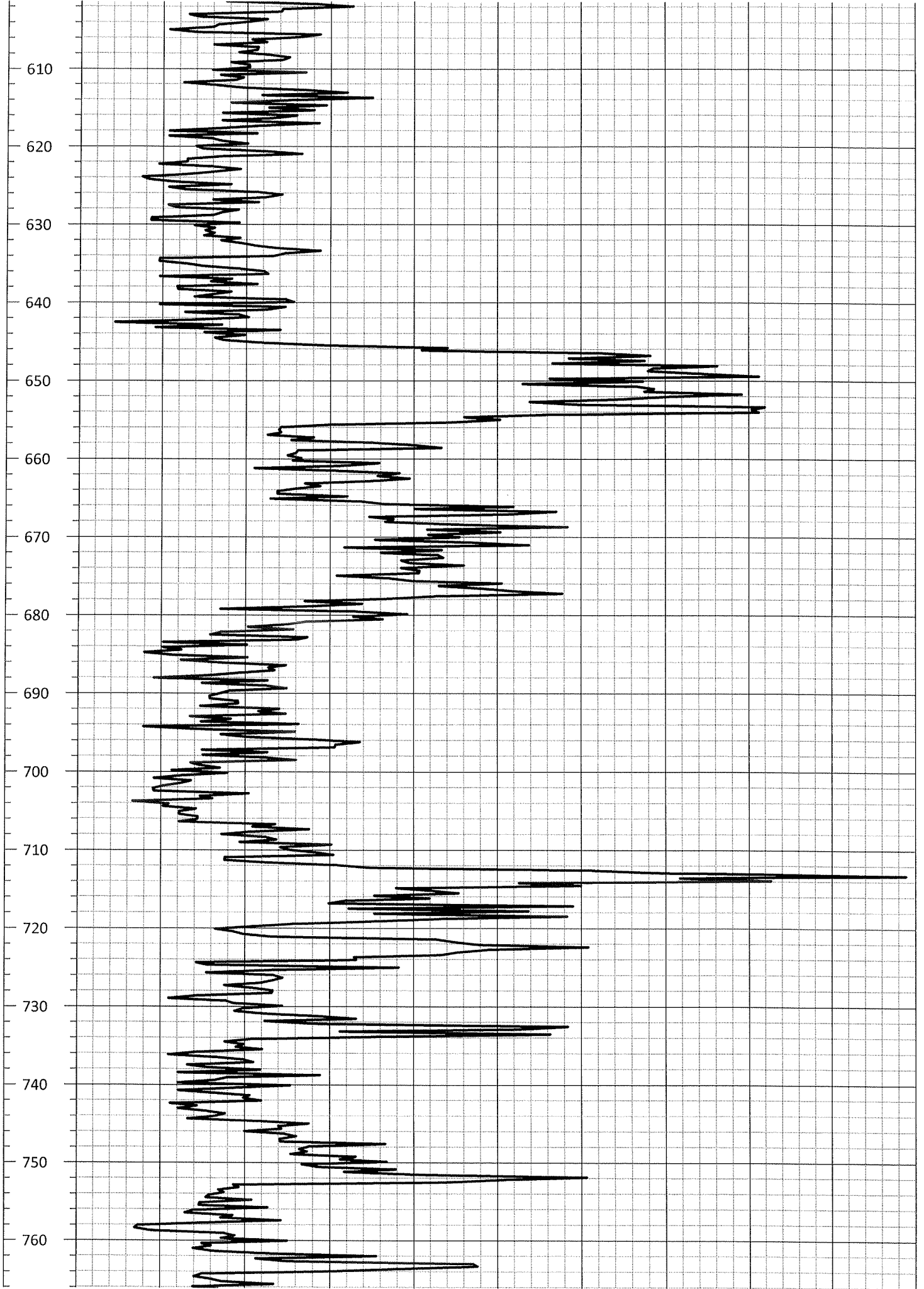
100.0



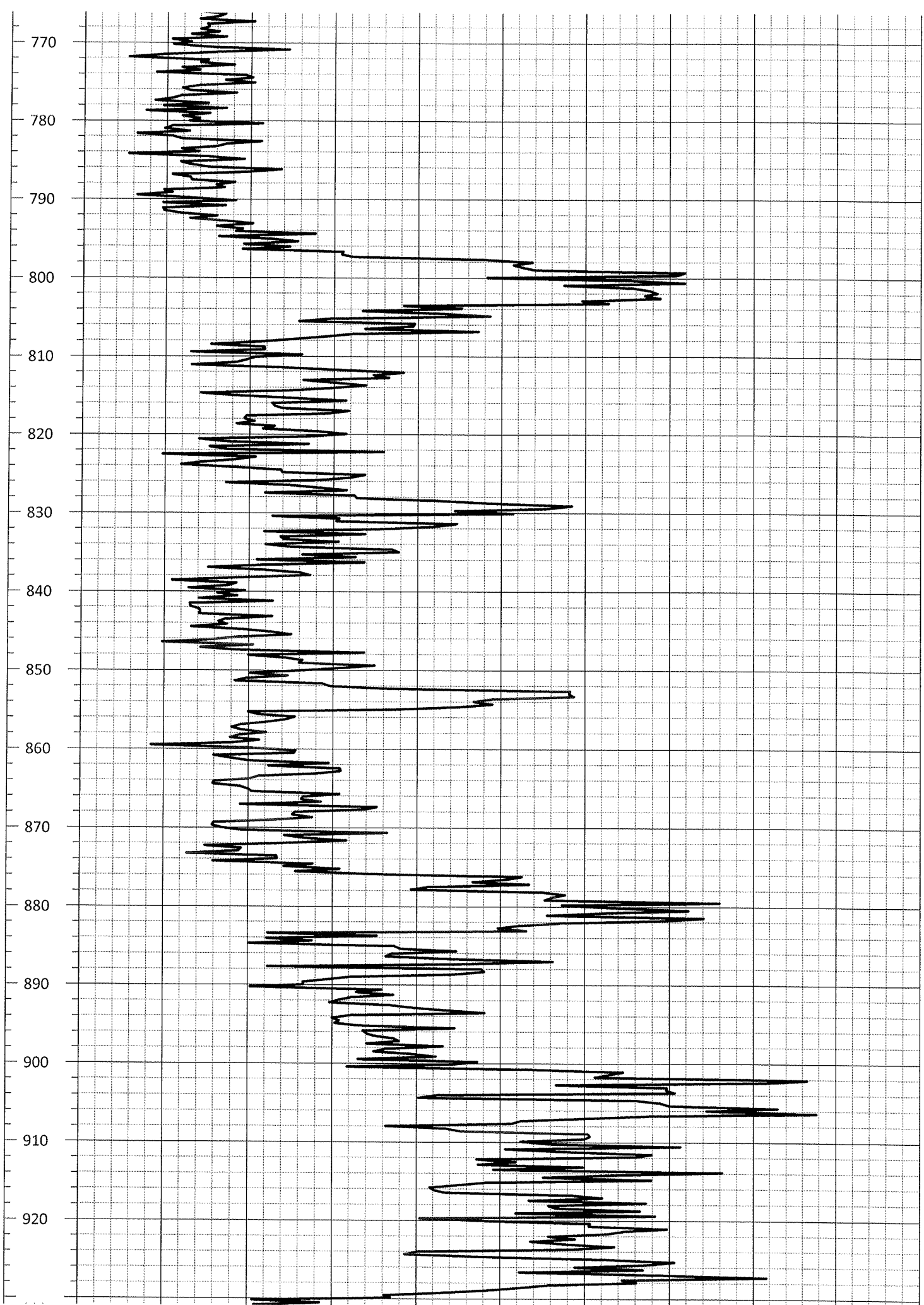


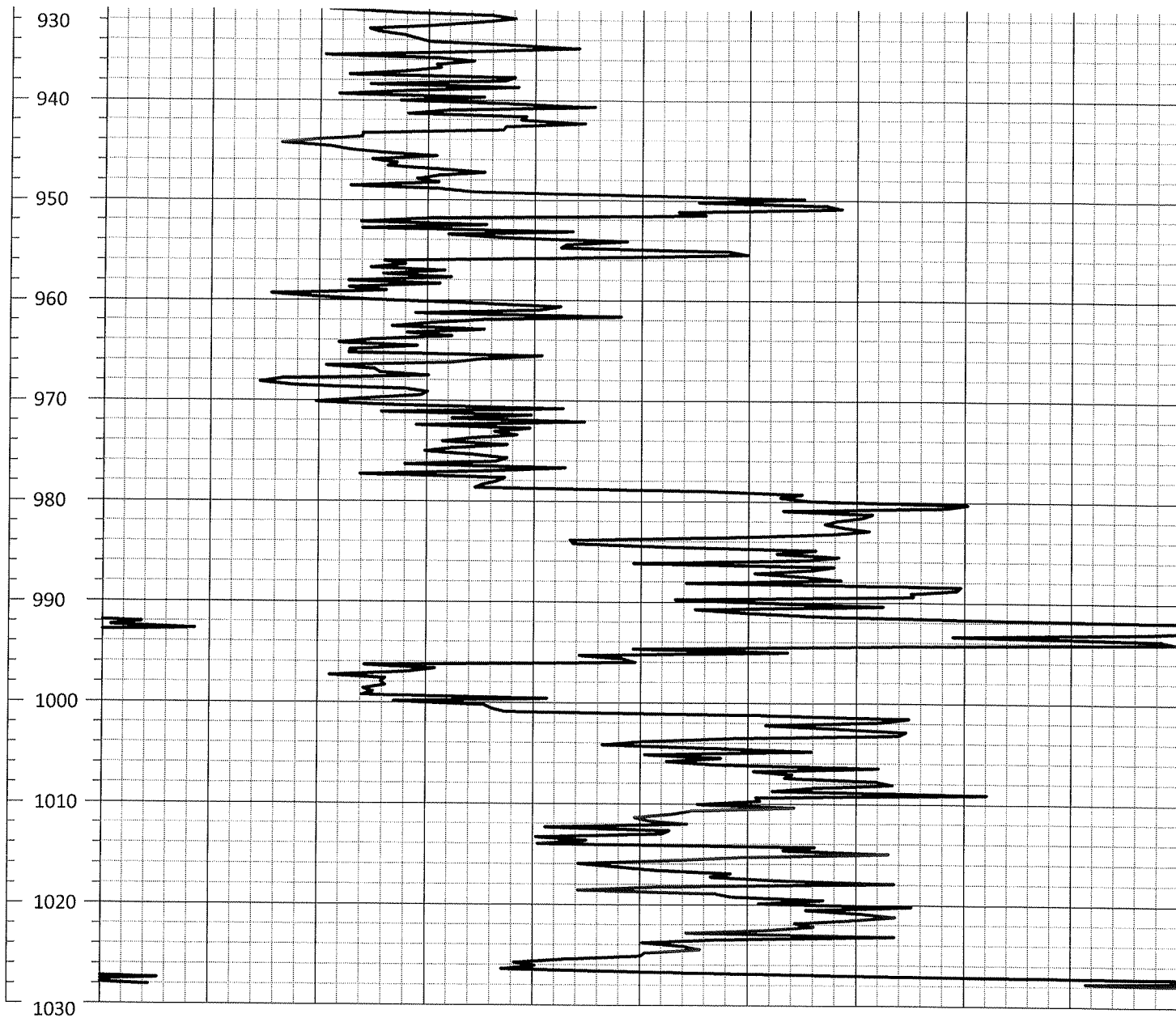










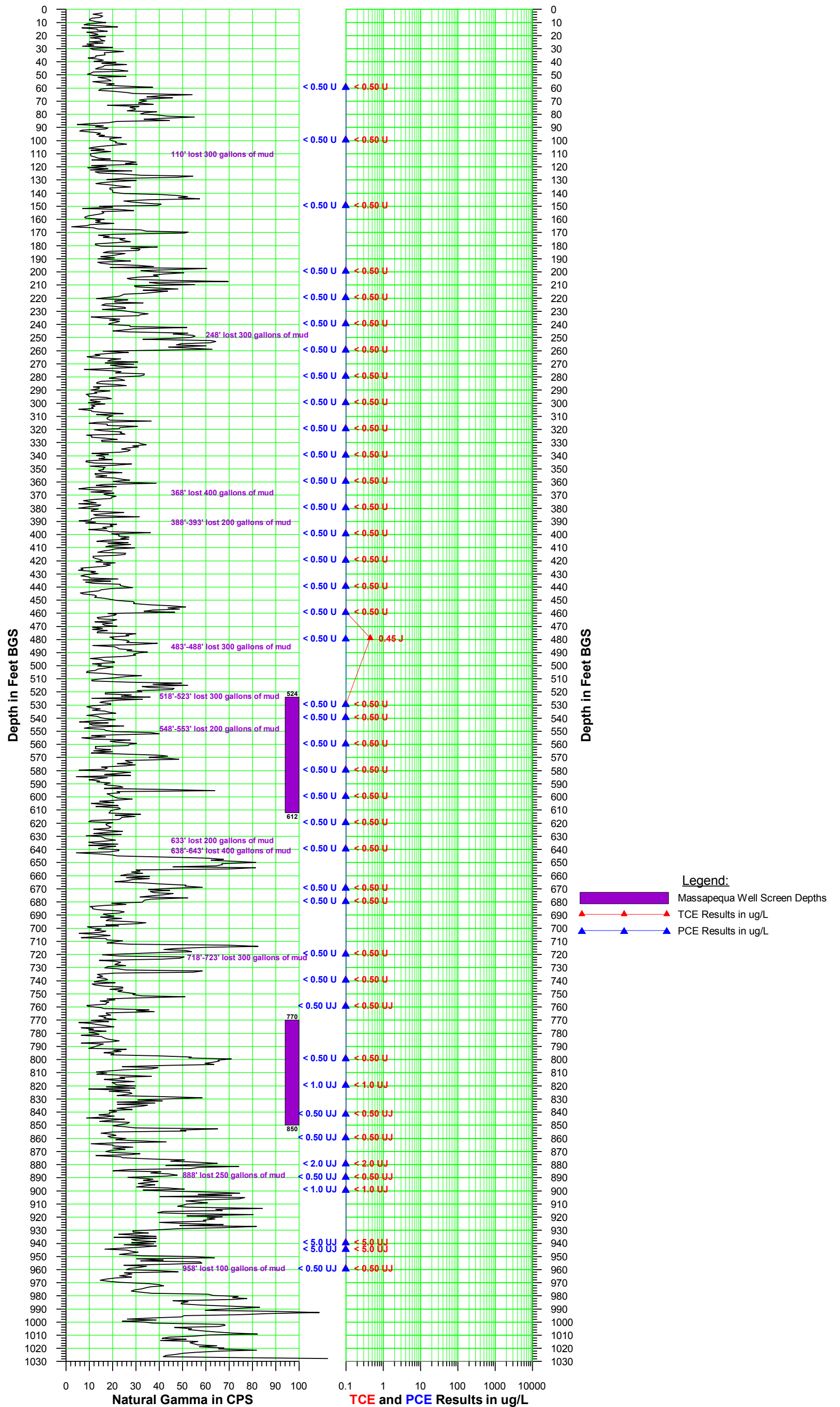


Depth (ft.)	0.0	GAMMA (cps)	100.0
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## **Section 2**

### **VPB 147 Gamma and PCE/TCE Plot**

# Vertical Profile Boring VPB-147 Downward Run - July 8, 2014 Validated Analytical Data



**Section 3**

**VPB 147 Groundwater Sample Log Sheets**

Hydropunch Sample

Client: Navy  
 Project No: 60266526  
 Site Location: intersection of Crocus Drive and sheep pasture land north Massapequa

Date: June 5 - June 10, 2014  
 VPB: 147

Collector(s): Bob Trayer

Sample Date	Time	Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Starting depth (ft)	Ending depth (ft)	Color
6/5/14	12:45	18.96	7.04	465	5.86	65.8	466	58	60	pale brown
6/5/14	15:00	19.67	6.74	414	3.39	86.0	422.8	98	100	pale gray
6/6/14	11:00	20.27	7.32	625	0.02	12.4	1000+	148	150	initially clear sample, then dark brown as pouring
6/6/14	14:45	not enough	recovery for YSI					198	200	
6/9/14	10:00	16	5.95	239	8.87	85.8	239	218	220	clear
6/9/14	12:00	17	6.00	146	7.56	82.0	136.3	238	240	clear
6/9/14	14:15	18.85	7.51	148	4.47	44.6	297.4	258	260	
6/9/14	16:00	15.27	6.90	220	6.90	55.40	73.75	278	280	clear
6/10/14	11:30	20.88	6.79	274	3.74	79.9	100.8	298	300	clear
6/10/14	14:30	21.54	7.21	310	8.64	43.2	183.6	318	320	
6/10/14	17:00	21.77	7.56	379	5.96	44.0	221.5	338	340	

Hydropunch Sample

Client:  
Project No:  
Site Location:

Navy  
60266526  
Crocus Drive &  
Sheep Pasture Lane

Date: June 11 - June 18, 2014  
VPB: 147

Collector(s): V. Thayer

Sample Date	Time	Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Starting depth (ft)	Ending depth (ft)	Color
6/11/14	10:15	19.41	6.13	229	8.27	43	211.6	358	360	clear
6/11/14	13:15	17.0	6.14	251	6.84	32.6	77.9	378	380	
6/12/14	16:30	22.5	6.12	155	5.85	47.1	116.5	398	400	
6/12/14	14:30	21.34	6.26	178	4.38	33.1	32.05	418	420	Pale gray
6/12/14	16:00	23.43	6.36	56	5.01	24.7	161.3	438	440	Pale gray
6/13/14	12:30	24.3	6.65	39	7.35	53.5	754.7	458	460	
6/13/14	14:45	23.58	6.68	107	5.00	18.0	469.9	478	480	
6/16/14	11:30	no recovery	no recovery					498	500	
6/16/14	15:00	no recovery	no recovery					518	520	Gray Silt (VT)
6/17/14	10:15	not enough recovery	not enough recovery					528	530	Brownish Gray
6/17/14	13:00	22.84	7.02	228	0.57	35.7	109.2	538	540	Gray
6/17/14	15:00	17.16	6.87	95	2.87	64.6	200.4	558	560	Gray
6/18/14	10:15	not enough recovery	not enough recovery					578	580	

not enough recovery for YSI

Hydropunch Sample

Client:  
Project No:  
Site Location:

NAVY  
00266526  
intersection of Crocers  
Drive and Sheep Pasture  
Lane, north Mkoaapogua

Date: June 18 - June 24, 2014  
VPB: 147

Collector(s): V. Thayer

Sample Date	Time	Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Starting depth (ft)	Ending depth (ft)	Color
6/18/14	13:00	23.9	6.53	76	6.14	10.4	680.7	598	600	
6/18/14	16:00	24.71	6.08	101	4.42	119.4	1000+	618	620	Gray silt
6/19/14	11:00	not enough	not enough	recovery for YSI				638	640	
6/19/14	no recovery							658	660	
6/19/14	16:00	20.73	7.24	65	4.25			668	670	
6/20/14	10:30	23.56	7.10	102	5.53	6.1	924	678	680	pale gray
6/20/14	13:45	no recovery						698	700	
6/23/14	11:15	29.8	6.22	164	4.2	74.3	812.2	718	720	
6/23/14	14:00	24.35	5.97	111	4.7	50.6	960.3	738	740	
6/24/14	11:30	23.67	6.70	147	2.37	20.4	1000+	758	760	pale gray milky color
6/24/14	13:45	no recovery						778	780	
6/24/14	16:00	no recovery						783	785	
6/25/14	12:00	not enough						798	800	



Hydropunch Sample

Navy  
 60266926  
 Crocus Avenue  
 Sheep Pasture Lane

Client:  
 Project No:  
 Site Location:

Date: June 25 - July 2, 2014  
 VPB: 1470

Collector(s): V. Trayer

Sample Date	Time	Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Starting depth (ft)	Ending depth (ft)	Color
6/25/14	14:40							818	820	Silty grey
6/26/14	11:00	24.48	7.07	288	1.51	83.5	860	840	842	Gray
6/26/14	16:00	26.17	6.61	784	0.75	9.3	1000+	858	860	Dark gray
6/27/14	10:30	21.10	7.06	775	4.62	105.1	1000+	878	880	Dark gray
6/27/14	13:45	26.88	7.08	772	0.54	1.9	1000+	888	890	Dark grey
6/30/14	12:15							898	900	
6/30/14	15:30							918	920	
7/01/14	11:30	27.26	7.58	766	1.83	18.4	1000+	938	940	GRAY
7/01/14	14:15							943	945	
7/01/14	17:00							958	960	Gray mud
7/02/14	12:00							978	980	black muck
7/02/14	14:45							988	990	

## **Section 4**

### **VPB 147 Analytical Data Validation**

- Analytical Data Sheets
- Chain of Custody Records
- Validation Letter and Table



## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	KATAHDIN ANALYTICAL	
Service Request:	SH3955	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS)	
Validation Level:	Limited	
AECOM Project Number:	60266526.SA.DV	
Prepared by:	Dawn Brule/RESCON	Completed on: 09/23/2014
Reviewed by:	Paula DiMattei/RESCON	File Name: SH3955_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 5, 2014.

Sample ID	Matrix/Sample Type
VPB147-GW-060514-58-60	Ground water
VPB147-GW-060514-98-100	Ground water
VPB147-TRIP BLANK-060514	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✗ Initial calibration/continuing calibration verification
- ✓ Laboratory blanks/trip blanks/equipment blanks
- ✗ Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicates
- ✓ Internal standards

✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2. Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks are evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

Nonconformances are summarized in Attachment A in Table A-3. Data qualification on the basis of surrogate recovery nonconformances was as follows:

Criteria	Action	
	Detected Compounds	Nondetected Compounds
%R > Upper Limit (UL)	J	No qualification
20% ≤ %R < Lower Limit (LL)	J	UJ
%R < 20%	J	R

Qualified sample results are shown in Table 1.

### **MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

## **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

## **ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-TRIP BLANK-060514	WQ	ACETONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-060514	WQ	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-060514	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-060514	WQ	CHLOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-060514	WQ	METHYL TERT-BUTYL ETHER		0.50	UG_L	UJ	c
VPB147-TRIP BLANK-060514	WQ	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c



## Attachment A

### Non Conformance Summary Tables

**Table A-1 - Initial Calibration**

Calibration Date/Time	Compound	% R	Limits
27-May-2014 10:16	chloroethane	17.2	<15%
Associated samples: VPB147-TRIP BLANK-060514			

**Table A-2 -Continuing Calibration Verification Standard**

CCV ID	Compound	% D	Limits
WG144345-4	chloromethane	21.3	<20%
	bromomethane	43.6	<20%
	chloroethane	33.7	<20%
	trichlorofluoromethane	30.2	<20%
	acetone	37.0	<20%
	Methyl tert-butyl ether	23.3	<20%
Associated sample: VPB147-TRIP BLANK-060514			

**Table A-3 - Surrogates**

Sample ID	Surrogate	% Recovery	Lower Limit	Upper Limit
VPB147-TRIP BLANK-060514	1,2-DICHLOROETHANE-D4	126	70	120

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

### Attachment C

#### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



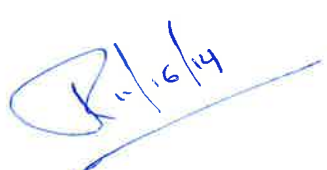
## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH3955-1  
**Client ID:** 147-TB-060514  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH3955  
**Lab File ID:** D8821.D

**Sample Date:** 05-JUN-14  
**Received Date:** 06-JUN-14  
**Extract Date:** 09-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144345

**Analysis Date:** 09-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50



*[Handwritten Signature]*  
6/16/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH3955-1  
**Client ID:** 147-TB-060514  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH3955  
**Lab File ID:** D8821.D

**Sample Date:** 05-JUN-14  
**Received Date:** 06-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144345

**Analysis Date:** 09-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene	*	74.7	%					
Toluene-d8		96.6	%					
1,2-Dichloroethane-d4	*	126.	%					
Dibromofluoromethane		102.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH3955-2RA  
**Client ID:** 147-060514-58-60  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH3955  
**Lab File ID:** D8912.D

**Sample Date:** 05-JUN-14  
**Received Date:** 06-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144658

**Analysis Date:** 13-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		12	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.69	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH3955-2RA  
**Client ID:** 147-060514-58-60  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH3955  
**Lab File ID:** D8912.D

**Sample Date:** 05-JUN-14  
**Received Date:** 06-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144658

**Analysis Date:** 13-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		102.	%					
Toluene-d8		112.	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		104.	%					



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH3955-3RA  
**Client ID:** 147-060514-98-100  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH3955  
**Lab File ID:** D8913.D

**Sample Date:** 05-JUN-14  
**Received Date:** 06-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144658

**Analysis Date:** 13-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	2.3	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH3955-3RA  
**Client ID:** 147-060514-98-100  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH3955  
**Lab File ID:** D8913.D

**Sample Date:** 05-JUN-14  
**Received Date:** 06-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144658

**Analysis Date:** 13-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		106.	%					
Toluene-d8		115.	%					
1,2-Dichloroethane-d4		118.	%					
Dibromofluoromethane		109.	%					



## Data Validation Report

Project: Regional Groundwater Investigation - NWIRP Bethpage  
Laboratory: KATAHDIN ANALYTICAL  
Service Request: SH4055  
Analyses/Method: EPA SW-846 Method 8260B for VOCs (GC/MS)  
Validation Level: Limited  
AECOM Project Number: 60266526.SA.DV  
Prepared by: Dawn Brule/RESCON Completed on: 09/24/2014  
Reviewed by: Paula DiMattei/RESCON File Name: SH4055\_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 6, 2014 and June 9, 2014.

Sample ID	Matrix/Sample Type
VPB147-GWD-060914	Field Duplicate of VPB147-GW-060914-218-220
VPB147-GW-060614-148-150	Ground water
VPB147-GW-060614-198-200	Ground water
VPB147-GW-060914-218-220	Ground water
VPB147-GW-060914-238-240	Ground water
VPB147-GW-060914-258-260	Ground water
VPB147-GW-060914-278-280	Ground water
VPB147-TRIP BLANK-060914	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✗ Initial calibration/continuing calibration verification

- ✓ Laboratory blanks/trip blanks/equipment blanks
- ✓ Surrogate spike recoveries
- ✓ Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- ✓ Field duplicates
- ✓ Internal standards
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## **RESULTS**

### **Data Completeness**

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### **Holding Times/Sample Preservation**

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2. Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

conformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus AECOM professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **MS/MSD Results**

The MS/MSD %Rs and relative percent differences (RPDs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met or qualification of the sample data was not required.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Field Duplicate Results**

Field duplicate RPDs were reviewed for conformance with the QC criterion of  $\leq 30\%$  for aqueous matrices. This criterion applies if both results were greater than five times the Limit of Quantitation (LOQ).

All QC acceptance criteria were met.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-060614-148-150	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-060614-148-150	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060614-148-150	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-060614-148-150	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060614-198-200	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-060614-198-200	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060614-198-200	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-060614-198-200	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-218-220	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-060914-218-220	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-218-220	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-218-220	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-238-240	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-060914-238-240	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-238-240	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-238-240	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-258-260	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-060914-258-260	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-258-260	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-258-260	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-278-280	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-060914-278-280	WG	BROMOMETHANE		1.0	UG_L	UJ	c



Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-060914-278-280	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-060914-278-280	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GWD-060914	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GWD-060914	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GWD-060914	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GWD-060914	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-060914	WQ	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-060914	WQ	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-060914	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-060914	WQ	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c

## Attachment A

## Non Conformance Summary Tables

Table A-1 - Initial Calibration

Calibration Date/Time	Compound	% R	Limits
09-June-2014 11:06	chloroethane	36.2	<15%
Associated samples: all samples in SDG SH4055			

Table A-2 -Continuing Calibration Verification Standard

CCV ID	Compound	% D	Limits
WG144490-4	dichlorodifluoromethane	31.3	<20%
	bromomethane	-23.9	<20%
	chloroethane	-26.3	<20%
	4-methyl-2-pentanone	25.8	<20%
Associated samples: all samples in SDG SH4055			

**Attachment B**  
**Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

### Attachment C

#### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



600 Technology Way  
 Scarborough, ME 04074  
 Tel: (207) 874-2400  
 Fax: (207) 775-4029

# CHAIN of CUSTODY

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Page \_\_\_\_ of \_\_\_\_

Client: *Resolution Consultants* Contact: *Eleanor Vivandou* Phone #: *( )* Fax #: *( )*  
 Address: *998 Cold Country Rd* City: *Plainville* State: *NY* Zip Code: *11803*  
 Purchase Order #: \_\_\_\_\_ Proj. Name / No.: *Bethpage 60266526* Katahdin Quote #: \_\_\_\_\_  
 Bill (if different than above) Address: \_\_\_\_\_

Sampler (Print / Sign) *Valerie Thayer Valerie Thayer* Copies To: *Valerie Thayer*

LAB USE ONLY WORK ORDER # *514055*  
 KATAHDIN PROJECT NUMBER \_\_\_\_\_  
 REMARKS: \_\_\_\_\_  
 SHIPPING INFO:  FED EX  UPS  CLIENT  
 AIRBILL NO: \_\_\_\_\_  
 TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

LAB USE ONLY					ANALYSIS AND CONTAINER TYPE PRESERVATIVES										
* Sample Description	Date / Time col'd	Matrix	No. of Cntrs.		Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON	Filt. OY ON
✓ VPB147-GW-060614-148-150	6/6/14 / 11:00	GW	3	X											
✓ VPB147-GW-MS/MSD-060614 148-150	6/6/14 / 11:00	GW	6	X											
✓ VPB147-GW-060614-198-200	6/6/14 / 2:45	GW	1	X											
✓ VPB147-Tripblanks - 060914	5/2/14 / 11:00	W	3	X											
✓ VPB147-GW-060914-278-280	6/9/14 / 4:00	GW	4	X											
✓ VPB147-GW-060914-258-260	6/9/14 / 14:15	GW	4	X											
✓ VPB147-GW-060914-238-240	6/9/14 / 12:00	GW	4	X											
✓ VPB147-GW-060914-218-220	6/9/14 / 10:00	GW	4	X											
✓ VPB147-GW-060914	6/9/14 / 10	GW	3	X											
Temperature	/		1												
Blank	/														

COMMENTS *VPB 147-GW-060614-198-200 - bubbles formed after sampling*

Relinquished By: (Signature) <i>Valerie Thayer</i>	Date / Time <i>6/9/14 7-</i>	Received By: (Signature) <i>[Signature]</i>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

THE TERMS AND CONDITIONS ON THE REVERSE SIDE HEREOF SHALL GOVERN SERVICES, EXCEPT WHEN A SIGNED CONTRACTUAL AGREEMENT EXISTS.

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## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-1  
**Client ID:** 147-060614-148-150  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7582.D

**Sample Date:** 06-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>UMM</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	<del>UM</del> U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J	0.35	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		13	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	<del>UMM</del> U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	<del>UM</del> U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R. 1/29/15*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4055-1  
 Client ID: 147-060614-148-150  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4055  
 Lab File ID: C7582.D

Sample Date: 06-JUN-14  
 Received Date: 10-JUN-14  
 Extract Date: 11-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144490

Analysis Date: 11-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	<del>UMM</del> U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	<del>UMM</del> U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	<del>UMM</del> U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		94.2	%					
Toluene-d8		97.0	%					
1,2-Dichloroethane-d4		109.	%					
Dibromofluoromethane		97.9	%					

*Handwritten signature and date: J. 129/15*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-2  
**Client ID:** 147-060614-198-200  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7583.D

**Sample Date:** 06-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J	0.29	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		6.9	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 12/2/17



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-2  
**Client ID:** 147-060614-198-200  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7583.D

**Sample Date:** 06-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		88.2	%					
Toluene-d8		91.8	%					
1,2-Dichloroethane-d4		103.	%					
Dibromofluoromethane		94.1	%					

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4055-7  
 Client ID: 147-060914-218-220  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4055  
 Lab File ID: C7587.D

Sample Date: 09-JUN-14  
 Received Date: 10-JUN-14  
 Extract Date: 11-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144490

Analysis Date: 11-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*REC 12/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-7  
**Client ID:** 147-060914-218-220  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7587.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.1	%					
Toluene-d8		92.3	%					
1,2-Dichloroethane-d4		106.	%					
Dibromofluoromethane		96.4	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-6  
**Client ID:** 147-060914-238-240  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7586.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	3.3	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Handwritten signature and date: 12/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-6  
**Client ID:** 147-060914-238-240  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7586.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		90.8	%					
Toluene-d8		94.1	%					
1,2-Dichloroethane-d4		105.	%					
Dibromofluoromethane		96.0	%					

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4055-5  
 Client ID: 147-060914-258-260  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4055  
 Lab File ID: C7585.D

Sample Date: 09-JUN-14  
 Received Date: 10-JUN-14  
 Extract Date: 11-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144490

Analysis Date: 11-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	⊕ UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	⊕ UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	⊕ UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	⊕ UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Handwritten signature/initials*  
 6/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-5  
**Client ID:** 147-060914-258-260  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7585.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.0	%					
Toluene-d8		93.3	%					
1,2-Dichloroethane-d4		107.	%					
Dibromofluoromethane		94.6	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-4  
**Client ID:** 147-060914-278-280  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7584.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> U5	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> U5	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> U5	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> U5	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-4  
**Client ID:** 147-060914-278-280  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7584.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		94.3	%					
Toluene-d8		96.6	%					
1,2-Dichloroethane-d4		111.	%					
Dibromofluoromethane		98.9	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-8  
**Client ID:** VPB147-GWD-060914  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7588.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50


 REC 12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-8  
**Client ID:** VPB147-GWD-060914  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7588.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		90.4	%					
Toluene-d8		93.6	%					
1,2-Dichloroethane-d4		106.	%					
Dibromofluoromethane		95.4	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-3  
**Client ID:** VPB147-TB-060914  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7579.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*REC 12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4055-3  
**Client ID:** VPB147-TB-060914  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4055  
**Lab File ID:** C7579.D

**Sample Date:** 09-JUN-14  
**Received Date:** 10-JUN-14  
**Extract Date:** 11-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144490

**Analysis Date:** 11-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.0	%					
Toluene-d8		93.6	%					
1,2-Dichloroethane-d4		101.	%					
Dibromofluoromethane		93.1	%					



## Data Validation Report

Project: Regional Groundwater Investigation - NWIRP Bethpage  
Laboratory: KATAHDIN ANALYTICAL  
Service Request: SH4157  
Analyses/Method: EPA SW-846 Method 8260B for VOCs (GC/MS) and EPA SW-846 Method 9060 for TOC (Carbonaceous analyzer, IR or FID)  
Validation Level: Limited  
AECOM Project Number: 60266526.SA.DV  
Prepared by: Dawn Brule/RESCON Completed on: 09/29/2014  
Reviewed by: Paula DiMattei/RESCON File Name: SH4157\_8260B\_9060

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 10, 2014 and June 11, 2014.

Sample ID	Matrix/Sample Type
VPB147-EB-061114	Equipment blank
VPB147-SOIL-D-061014	Field Duplicate of VPB147-SOIL-061014-283-285
VPB147-GW-061014-298-300	Ground water
VPB147-GW-061014-318-320	Ground water
VPB147-GW-061014-338-340	Ground water
VPB147-GW-061114-358-360	Ground water
VPB147-GW-061114-378-380	Ground water
VPB147-SOIL-061014-283-285	Soil
VPB147-TB-061114	Trip Blank

The samples were analyzed in accordance with *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846* (USEPA, 1996), specifically:

- Method 8260B, *Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry*
- Method 9060A, *Total Organic Carbon*

Data validation activities were conducted with reference to these methods, *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), *USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review* (January 2010), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010) where applicable. In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification
- X Laboratory blanks/trip blanks/equipment blanks
- ✓ Surrogate spike recoveries
- ✓ Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- X Field duplicates
- ✓ Internal standards
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated or negated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.



**GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

**Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

**ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus AECOM professional judgment was used		

**CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

**Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL).

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.



Method, equipment rinsate and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

Nonconformances are summarized in Attachment A in Tables A-3 . Sample results were qualified as follows:

Blank type	Blank result	Sample result	Action for samples
Method, Storage, Field, Trip, or Instrument*	Detects	Not detected	No qualification
	≤ LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and ≤ 2x LOQ	Report the sample result with a U**
		≥ 2x the LOQ	No qualifications
	> LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and < blank contamination	Report the sample result with a U or reject the sample result as unusable R
		≥ LOQ and ≥ blank contamination	If the result is ≤2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
	TIC detected	Detects	If the result is ≤2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
* Qualifications based on instrument blank results affect only the sample analyzed immediately after the sample that has target compounds that exceed the calibration range or non-target compounds that exceed 100 g/L.			
**Based on professional judgment.			

LOQ - Limit of Quantitation. Qualified sample results are shown in Table 1.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **MS/MSD Results**

The MS/MSD %Rs and relative percent differences (RPDs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Field Duplicate Results**

Field duplicate RPDs were reviewed for conformance with the QC criterion of  $\leq 30\%$  for soil matrices. This criterion applies if both results were less than five times the Limit of Quantitation (LOQ).

Nonconformances are summarized in Attachment A in Table A-4.

Data qualification to the analytes associated with the specific field duplicate RPDs was as follows:

**Actions:** professional judgment was used

Criteria	RPD	Action	
		Detected	Nondetected
Sample and duplicate are nondetect results	Not calculable (NC)	No qualification	No qualification
Sample and duplicate results $\geq 5x$ LOQ	$>30$ (soil)	J	Not Applicable
Sample and duplicate results $<5x$ LOQ	$>60$ (soil)	J	Not Applicable
If sample or duplicate result is $>5x$ LOQ and the other is not detected	NC	J	UJ
If sample or duplicate result is $<5x$ LOQ and the other is not detected	NC	No qualification	No qualification

Qualified sample results are shown in Table 1.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-EB-061114	WQ	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-EB-061114	WQ	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-EB-061114	WQ	CARBON DISULFIDE		1.0	UG_L	U	bl
VPB147-EB-061114	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-EB-061114	WQ	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061014-298-300	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061014-298-300	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061014-298-300	WG	CARBON DISULFIDE		1.0	UG_L	U	bl
VPB147-GW-061014-298-300	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061014-298-300	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061014-318-320	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061014-318-320	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061014-318-320	WG	CARBON DISULFIDE		1.0	UG_L	U	bl
VPB147-GW-061014-318-320	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061014-318-320	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061014-338-340	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061014-338-340	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061014-338-340	WG	CARBON DISULFIDE		1.0	UG_L	U	bl
VPB147-GW-061014-338-340	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061014-338-340	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061114-358-360	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061114-358-360	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-061114-358-360	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061114-358-360	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061114-378-380	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061114-378-380	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061114-378-380	WG	CARBON DISULFIDE		1.0	UG_L	U	bl
VPB147-GW-061114-378-380	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061114-378-380	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-SOIL-061014-283-285	SO	TOTAL ORGANIC CARBON	1500	490	UG_G	J	fd
VPB147-SOIL-D-061014	SO	TOTAL ORGANIC CARBON	3600	500	UG_G	J	fd
VPB147-TB-061114	WQ	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-TB-061114	WQ	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-TB-061114	WQ	CARBON DISULFIDE		1.0	UG_L	U	bl
VPB147-TB-061114	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-TB-061114	WQ	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c

## Attachment A

## Non Conformance Summary Tables

Table A-1 - Initial Calibration

Calibration Date/Time	Compound	% R	Limits
09-June-2014 11:06	chloroethane	36	<15%
Associated samples: VPB147-EB-061114, VPB147-GW-061014-298-300, VPB147-GW-061014-318-320, VPB147-GW-061014-338-340, VPB147-GW-061114-378-380, VPB147-TB-061114			

Table A-2 -Continuing Calibration Verification Standard

CCV ID	Compound	% R	Limits
WG144656-4	dichlorodifluoromethane	45	<20%
	chloroethane	-26	<20%
	4-methyl-2-pentanone	30	<20%
	2-hexanone	21	<20%
Associated samples: VPB147-EB-061114, VPB147-GW-061014-298-300, VPB147-GW-061014-318-320, VPB147-GW-061014-338-340, VPB147-GW-061114-378-380, VPB147-TB-061114			

Table A-3 - Lab Blanks

Blank ID	Compound	Result	QL	Units	Associated Samples
WG144656-2	CARBON DISULFIDE	0.33	0.50	UG_L	VPB147-EB-061114 VPB147-GW-061014-298-300 VPB147-GW-061014-318-320 VPB147-GW-061014-338-340 VPB147-GW-061114-378-380 VPB147-TB-061114

Table A-4 - Field Duplicates

Sample ID	Duplicate ID	Analyte	Sample Result	Qual	Duplicate Result	Qual	LOQ	Units	RPD
VPB147-SOIL-061014-283-285	VPB147-SOIL-D-061014	TOC	1500		3600		490	ug/g	82.4

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

### Attachment C

#### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance





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# CHAIN of CUSTODY

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Page \_\_\_\_ of \_\_\_\_

Client: Resolution Consultants Contact: Eleanor Vukobrodic Phone #: ( ) Fax #: ( )  
 Address: 998 C Old Country Rd City: Plainville State: NY Zip Code: 11803  
 Purchase Order #: \_\_\_\_\_ Proj. Name / No.: \_\_\_\_\_ Katahdin Quote #: \_\_\_\_\_

Bill (if different than above) Address: \_\_\_\_\_

Sampler (Print / Sign): Valerie Thayer / Valerie Thayer Copies To: Valerie Thayer

LAB USE ONLY WORK ORDER # / 5141570  
 KATAHDIN PROJECT NUMBER \_\_\_\_\_  
 REMARKS: \_\_\_\_\_  
 SHIPPING INFO:  FED EX  UPS  CLIENT  
 AIRBILL NO: \_\_\_\_\_  
 TEMP °C  TEMP BLANK  INTACT  NOT INTACT

ANALYSIS AND CONTAINER TYPE PRESERVATIVES

Fit.	Fit.	Fit.	Fit.	Fit.	Fit.	Fit.	Fit.	Fit.	Fit.
OY	ON	OY	ON	OY	ON	OY	ON	OY	ON

*	Sample Description	Date / Time coll'd	Matrix	No. of Cntrs.															
✓	VPB147-Trip blanks - 061114	5/21/14 / 10:00	W	3	X														
	Trip blank	/	W	1															
✓	VPB147-GW-061014-298-300	06102014 / 11:30	GW	4	X														
	VPB147-GW-061014-318-320	06102014 / 14:30	GW	3	X														
✓	VPB147-GW-061014-338-340	06102014 / 5:00 PM	GW	4	X														
* ✓	VPB147-Soil-061014-283-285	06102014 / 8:45 AM	Soil	1	X	X													
✓	VPB147-Soil/D-061014-VI	06102014 / 8:45	Soil	1		X													
✓	VPB147-GW-061114-358-360	06112014 / 10:15	GW	3	X														
✓	VPB147-GW-061114-378-380	06112014 / 13:15	GW	4	X														
* ✓	VPB147-GW-Equipment Rinse Blank - 061114	06112014 / 14:20	W	3	X														

COMMENTS: \* Please use 8oz jar for TOC sample & MS/MSD  
 \* Equipment Rinse Blank using hydro punch & distilled water (A-B)

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>06/11/14 7-</u>	Received By: (Signature) <u>[Signature]</u>	Date / Time <u>6-12-14 0800</u>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Date / Time	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

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000002 ORIGINAL

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-9  
**Client ID:** VPB147-EB-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7630.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U U	<del>0.32</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

REC 12/12/14

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-9  
**Client ID:** VPB147-EB-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7630.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.0	%					
Toluene-d8		95.4	%					
1,2-Dichloroethane-d4		103.	%					
Dibromofluoromethane		92.3	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-2  
**Client ID:** 147-061014-298-300  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7632.D

**Sample Date:** 10-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U U	0.31 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

REC 12/12/14

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-2  
**Client ID:** 147-061014-298-300  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7632.D

**Sample Date:** 10-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		93.5	%					
Toluene-d8		95.8	%					
1,2-Dichloroethane-d4		107.	%					
Dibromofluoromethane		95.1	%					

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4157-3  
 Client ID: 147-061014-318-320  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4157  
 Lab File ID: C7633.D

Sample Date: 10-JUN-14  
 Received Date: 12-JUN-14  
 Extract Date: 13-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144656

Analysis Date: 13-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U U	<del>0.29</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	4.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-3  
**Client ID:** 147-061014-318-320  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7633.D

**Sample Date:** 10-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.0	%					
Toluene-d8		96.3	%					
1,2-Dichloroethane-d4		108.	%					
Dibromofluoromethane		97.4	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-4  
**Client ID:** 147-061014-338-340  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7634.D

**Sample Date:** 10-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U U	<del>0.39</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone		12	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*REC 12/12/14*



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-4  
**Client ID:** 147-061014-338-340  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7634.D

**Sample Date:** 10-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		98.7	%					
Toluene-d8		99.5	%					
1,2-Dichloroethane-d4		111.	%					
Dibromofluoromethane		101.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-7  
**Client ID:** 147-061114-358-360  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7635.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone		5.2	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

UJ 12/12/14

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-7  
**Client ID:** 147-061114-358-360  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7635.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		94.3	%					
Toluene-d8		95.3	%					
1,2-Dichloroethane-d4		106.	%					
Dibromofluoromethane		95.1	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-8  
**Client ID:** 147-061114-378-380  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7636.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	U U	<del>0.31</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		6.4	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-8  
**Client ID:** 147-061114-378-380  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7636.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		96.1	%					
Toluene-d8		96.8	%					
1,2-Dichloroethane-d4		107.	%					
Dibromofluoromethane		96.9	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-1  
**Client ID:** VPB147-TB-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7629.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	U U	<del>0.28</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4157-1  
**Client ID:** VPB147-TB-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4157  
**Lab File ID:** C7629.D

**Sample Date:** 11-JUN-14  
**Received Date:** 12-JUN-14  
**Extract Date:** 13-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144656

**Analysis Date:** 13-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 14-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		94.4	%					
Toluene-d8		95.4	%					
1,2-Dichloroethane-d4		104.	%					
Dibromofluoromethane		93.2	%					

## Report of Analytical Results

**Client:** Rick Purdy  
AECOM  
701 Edgewater Drive  
Wakefield, MA 01880

**Lab Sample ID:** SH4157-5  
**Report Date:** 01-JUL-14  
**Client PO:** 16518  
**Project:** Navy Clean WE15-03-0  
**SDG:** SH4157

Sample Description  
147-061014-283-285

Matrix      Date Sampled      Date Received  
SL              10-JUN-14              12-JUN-14

Parameter	Result	Adj LOQ	Adj MDL	Adj LOD	Anal. Method	QC.Batch	Anal. Date	Prep. Method	Prep. Date	Footnotes
TOC In Soil	1500 ug/gdrywt	490	100	N/A	SW846 9060A Mod.	WG145000	17-JUN-14 13:38:35	N/A	N/A	
Total Solids	81. %	1	N/A	N/A	SM2540G	WG144717	16-JUN-14 09:29:58	SM2540G	14-JUN-14	

SE 12/12/14



## Report of Analytical Results

Client: Rick Purdy  
 AECOM  
 701 Edgewater Drive  
 Wakefield, MA 01880

Lab Sample ID: SH4157-6  
 Report Date: 01-JUL-14  
 Client PO: 16518  
 Project: Navy Clean WE15-03-0  
 SDG: SH4157

Sample Description  
 147-SOIL-D-061014

Matrix      Date Sampled      Date Received  
 SL              10-JUN-14              12-JUN-14

Parameter	Result	Adj LOQ	Adj MDL	Adj LOD	Anal. Method	QC.Batch	Anal. Date	Prep. Method	Prep. Date	Footnotes
TOC In Soil	3600 ug/gdrywt	500	100	N/A	SW846 9060A Mod.	WG145000	17-JUN-14 14:17:12	N/A	N/A	
Total Solids	80. %	1		N/A	SM2540G	WG144717	16-JUN-14 09:30:09	SM2540G	14-JUN-14	

*R. 12/12/14*



Resolution Consultants  
250 Apollo Drive  
Chelmsford, MA 01824

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## Data Validation Report

Project: Regional Groundwater Investigation - NWIRP Bethpage

Laboratory: Katahdin Analytical

Service Request: SH4221

Analyses/Method: EPA SW-846 Method 8260B for VOCs (GC/MS) and Standard Method 5310 for Total Organic Carbon by High-Temperature Combustion

Validation Level: 3

AECOM Project Number: 60266526.SA.DV

Prepared by: Dawn Brule/RESCON Completed on: 09/25/2014

Reviewed by: Lori Herberich/RESCON File Name: SH4221\_5310B and 8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 12, 2014 and May 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-EB-061214	Equipment blank
VPB147-GWD-061214	Field Duplicate of VPB147-GW-061214-398-400
VPB147-GW-061214-398-400	Groundwater
VPB147-GW-061214-418-420	Groundwater
VPB147-GW-061214-438-440	Groundwater
VPB147-TRIP BLANK-061214	Trip Blank

The samples were analyzed in accordance with:

*Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW-846, Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (USEPA, 1996).*

*Standard Methods for the Examination of Water and Wastewater, Method SM5310B, Total Organic Carbon by High-Temperature Combustion*

Data validation activities were conducted with reference to these methods, *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), *USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review* (January 2010), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2 (DoD, October 2010)* where applicable. In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

## REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- ✓ Data completeness (chain-of-custody [COC])/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✗ Initial calibration/continuing calibration verification
- ✗ Laboratory blanks/equipment blanks/trip blanks
- ✓ Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- ✓ Field duplicate results
- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (✗) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated or negated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness (COC)/Sample Integrity

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times and Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria. The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met. The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and/or
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICV and CCV was as follows:

#### **ICV Recovery Nonconformances:**

Nonconformance	Actions	
	Detected Compounds	Nondetected Compounds
%R > 120%	J	No qualification
20% < %R < 80%	J	UJ
%R < 20% (see note)	J	R*

Notes: Based on NFG 2008 VOC guidance, professional judgment is used to reject (R) nondetects in all associated samples for any analyte with < 20% recovery. Also, professional judgment is used to estimate (UJ) rather the reject (R) sample results previously negated (U) on the basis of blank contamination.

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift > 20%	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL).

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method, equipment rinsate and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

Nonconformances are summarized in Attachment A in Table A-3.

TOC sample results were qualified as follows:

Blank Type	Blank Result	Sample Result	Action for Samples
ICB/CCB (Positive)	≥DL but ≤ LOQ	Nondetect	No action
		≥DL but <LOQ	Qualify as nondetect (U) at the LOQ
		> LOQ	Use professional judgment (see below [1])
	>LOQ	≥DL but <LOQ	Qualify as nondetect (U) at the LOQ
		> LOQ but < ICB/CCB Result	Qualify at level of Blank Result with a "U" or Qualify result as unusable
		>ICB/CCB but <10x the ICB/CCB result	Qualify as estimated (J)
		≥10x ICB/CCB	No action is taken based on professional judgment
PB / EB/ FB (Positive)	> LOQ	≥DL but ≤ LOQ	Qualify as nondetect (U) at the LOQ
		>LOQ but < 10x Blank Result	Qualify results as unusable
		≥10x Blank Result	No action
	≥DL but ≤LOQ	Nondetect	No action
		≥DL but <LOQ	Qualify as nondetect (U) at the LOQ
		> LOQ	Use professional judgment (see below [1])

[1] Establish an action level (AL) at 5x the blank contamination. If sample result is <AL, qualify the reported result with a U.

LOQ - Limit of Quantitation.

Qualified sample results are shown in Table 1.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

**Field Duplicate Results**

Field duplicate RPDs were reviewed for conformance with the QC criterion of  $\leq 60\%$  for aqueous matrices. This criterion applies if both results were less than five times the Limit of Quantitation (LOQ). All QC acceptance criteria were met.

**Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

**Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

**QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-EB-061214	WQ	TOTAL ORGANIC CARBON		1.0*	MG/L	U	bl
VPB147-GW-061214-398-400	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	c
VPB147-GW-061214-398-400	WG	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-GW-061214-398-400	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-GW-061214-398-400	WG	ACETONE	7.8	2.5	UG/L	J	c
VPB147-GW-061214-418-420	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	c
VPB147-GW-061214-418-420	WG	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-GW-061214-418-420	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-GW-061214-438-440	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	c
VPB147-GW-061214-438-440	WG	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-GW-061214-438-440	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-GW-061214-438-440	WG	ACETONE	7.3	2.5	UG/L	J	c
VPB147-GWD-061214	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	c
VPB147-GWD-061214	WG	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-GWD-061214	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-GWD-061214	WG	ACETONE	11	2.5	UG/L	J	c
VPB147-TRIP BLANK-061214	WQ	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	c
VPB147-TRIP BLANK-061214	WQ	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-TRIP BLANK-061214	WQ	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c

\*LOQ

**Attachment A****Nonconformance Summary Tables****Table A-1 - Initial Calibration Verification Standard**

ICV ID	Compound	% R	Limits
WG144515-7	acetone	125	80-120%
Associated samples: all samples in SDG SH4221			

**Table A-2 -Continuing Calibration Verification Standard**

CCV ID	Compound	% D	Limits
WG144718-4	4-methyl-2-pentanone	21	≤20%
	2-hexanone	28	≤20%
	1,2-dibromo-3-chloropropane	22	≤20%
Associated samples: all samples in SDG SH4221			

**Table A-3 - Lab Blanks**

Blank ID	Compound	Result	LOD	Units	Associated Samples
WG145514-1	TOTAL ORGANIC CARBON	0.21	0.50	MG/L	VPB147-EB-061214



**Attachment B****Qualifier Codes and Explanations**

Qualifier	Explanation
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

## Attachment C

### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance





600 Technology Way  
 Scarborough, ME 04074  
 Tel: (207) 874-2400  
 Fax: (207) 775-4029

# CHAIN of CUSTODY

PLEASE BEAR DOWN AND  
 PRINT LEGIBLY IN PEN

Client: Resolution Consultants Contact: Eleonor Vidouzak Phone # ( ) Fax # ( )

Address: 100 Red Schoolhouse Rd Chestnut Ridge NY State NY Zip Code

Purchase Order #  Proj. Name / No. Bethpage Katahdin Quote #

Bill (if different than above) Address

Sampler (Print / Sign) Valerie Thayer VAL Thayer Copies To: Valerie Thayer

**LAB USE ONLY**

WORK ORDER #: SH4221

KATAHDIN PROJECT NUMBER

**ANALYSIS AND CONTAINER TYPE PRESERVATIVES**

REMARKS:

Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.
<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N

SHIPPING INFO:  FED EX  UPS  CLIENT

AIRBILL NO:

TEMP °C   TEMP BLANK  INTACT  NOT INTACT

VOCs 8240

TOC 112504

* Sample Description	Date / Time coll'd	Matrix	No. of Cntrs.
<u>VPB147-GW-061214-438-490</u>	<u>6/12/14 14:00</u>	<u>GW</u>	<u>4</u>
<u>VPB147-GW-061214-418-420</u>	<u>6/12/14 12:30</u>	<u>GW</u>	<u>4</u>
<u>VPB147-EB-061214</u>	<u>6/12/14 10:15</u>	<u>W</u>	<u>3</u>
<u>VPB147-GW-061214-398-400</u>	<u>6/12/14 9:45</u>	<u>GW</u>	<u>4</u>
<u>VPB147-GWD-061214</u>	<u>9:45</u>	<u>GW</u>	<u>3</u>
<u>Temperature Blank</u>	<u></u>	<u>W</u>	<u>1</u>
<u>VPB147-Trip blanks 061214</u>	<u>5/21/10:00 2014</u>	<u>W</u>	<u>3</u>
<u></u>	<u>/</u>	<u></u>	<u></u>
<u></u>	<u>/</u>	<u></u>	<u></u>
<u></u>	<u>/</u>	<u></u>	<u></u>
<u></u>	<u>/</u>	<u></u>	<u></u>

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COMMENTS

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>06/12/14 7:00</u>	Received By: (Signature) <u>[Signature]</u> <u>6-13-14 09:00</u>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-4  
**Client ID:** 147-061214-398-400  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8942.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	7.8	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U J5	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U J5	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 12/12/14

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4221-4  
 Client ID: 147-061214-398-400  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4221  
 Lab File ID: D8942.D

Sample Date: 12-JUN-14  
 Received Date: 13-JUN-14  
 Extract Date: 14-JUN-14  
 Extracted By: DJP  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144718

Analysis Date: 14-JUN-14  
 Analyst: DJP  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		101.	%					
Toluene-d8		111.	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		108.	%					

12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-2  
**Client ID:** 147-061214-418-420  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8941.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	# UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	# UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 12/12/14



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-2  
**Client ID:** 147-061214-418-420  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8941.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U 45	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		103.	%					
Toluene-d8		113.	%					
1,2-Dichloroethane-d4		111.	%					
Dibromofluoromethane		108.	%					

R12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-1  
**Client ID:** 147-061214-438-440  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8940.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	<b>J</b>	7.3	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> <b>UJ</b>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> <b>UJ</b>	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-1  
**Client ID:** 147-061214-438-440  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8940.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	<del>U</del> UJ	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		104.	%					
Toluene-d8		112.	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		104.	%					

R. 12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-5  
**Client ID:** VPB147-GWD-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8943.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	11	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U US	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U US	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/12/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4221-5  
 Client ID: VPB147-GWD-061214  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4221  
 Lab File ID: D8943.D

Sample Date: 12-JUN-14  
 Received Date: 13-JUN-14  
 Extract Date: 14-JUN-14  
 Extracted By: DJP  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144718

Analysis Date: 14-JUN-14  
 Analyst: DJP  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylencs (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	<del>U</del> UJ	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		103.	%					
Toluene-d8		110.	%					
1,2-Dichloroethane-d4		111.	%					
Dibromofluoromethane		108.	%					

R12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4221-6  
**Client ID:** VPB147-TB-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4221  
**Lab File ID:** D8933.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144718

**Analysis Date:** 14-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> U5	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> U5	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4221-6  
 Client ID: VPB147-TB-061214  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4221  
 Lab File ID: D8933.D

Sample Date: 12-JUN-14  
 Received Date: 13-JUN-14  
 Extract Date: 14-JUN-14  
 Extracted By: DJP  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144718

Analysis Date: 14-JUN-14  
 Analyst: DJP  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 16-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	<del>U</del> <b>U</b>	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		99.5	%					
Toluene-d8		110.	%					
1,2-Dichloroethane-d4		110.	%					
Dibromofluoromethane		104.	%					

## Report of Analytical Results

**Client:** Rick Purdy  
 AECOM  
 701 Edgewater Drive  
 Wakefield, MA 01880

**Lab Sample ID:** SH4221-3  
**Report Date:** 01-JUL-14  
**Client PO:** 16518  
**Project:** Navy Clean WE15-03-0  
**SDG:** SH4221

**Sample Description**  
 VPB147-EB-061214

**Matrix** AQ  
**Date Sampled** 12-JUN-14  
**Date Received** 13-JUN-14

Parameter	Result	Adj LOQ	Adj MDL	Adj LOD	Anal. Method	QC Batch	Anal. Date	Prep. Method	Prep. Date	Footnotes
Total Organic Carbon	<del>10.23</del> mg/L 1.00	1.0	0.10	.5	SM5310B	WG145514	26-JUN-14 19:13:05	N/A	N/A	

6/12/14







Resolution Consultants  
250 Apollo Drive  
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## Data Validation Report

Project: Regional Groundwater Investigation - NWIRP Bethpage

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Laboratory: Katahdin Analytical

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Service Request: SH4349

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Analyses/Method: EPA SW-846 Method 8260B for VOCs (GC/MS) and Standard Method 5310 for Total Organic Carbon by High-Temperature Combustion

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Validation Level: 3

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AECOM Project Number: 60266526.SA.DV

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Prepared by: Dawn Brule/RESCON Completed on: 09/30/2014

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Reviewed by: Lori Herberich/RESCON File Name: SH4349\_5310B and 8260B

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### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 13 and 16, 2014 and May 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-FB-061614	Field blank
VPB147-GW-061314-458-460	Groundwater
VPB147-GW-061314-478-480	Groundwater
VPB147-TRIP BLANKS-061614	Trip Blank

The samples were analyzed in accordance with:

- *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (USEPA, 1996).*
- *Standard Methods for the Examination of Water and Wastewater, Method SM5310B, Total Organic Carbon by High-Temperature Combustion*

Data validation activities were conducted with reference to these methods, *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), *USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review* (January 2010), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2 (DoD, October 2010)* where applicable. In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

## REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- ✓ Data completeness (chain-of-custody [COC])/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification
- X Laboratory blanks/equipment blanks/trip blanks
- ✓ Surrogate spike recoveries
- ✓ Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- X Laboratory control sample (LCS) results
- NA Field duplicate results
- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated or negated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness (COC)/Sample Integrity

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times and Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria. The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met. The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and/or
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift >20%	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method, field, and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

Nonconformances are summarized in Attachment A in Table A-3.

Sample results were qualified as follows:

Blank type	Blank result	Sample result	Action for samples
Method, Storage, Field, Trip, or Instrument*	Detects	Not detected	No qualification
	≤ LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and ≤ 2x LOQ	Report the sample result with a U**
		≥ 2x the LOQ	No qualifications
	> LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and < blank contamination	Report the sample result with a U or reject the sample result as unusable R
		≥ LOQ and ≥ blank contamination	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
	TIC detected	Detects	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
* Qualifications based on instrument blank results affect only the sample analyzed immediately after the sample that has target compounds that exceed the calibration range or non-target compounds that exceed 100 g/L.			
**Based on professional judgment.			

LOQ - Limit of Quantitation.

Qualified sample results are shown in Table 1.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **MS/MSD Results**

The MS/MSD %Rs and relative percent differences (RPDs) were reviewed for conformance with the QC acceptance criteria.

Several 8260B compounds exceeded the QC limits for %R in the MSD and/or the MS/MSD RPD for sample VPB147-GW-061314-458-460. The sample results were nondetect for these compounds and were accepted without qualification.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

Nonconformances are summarized in Attachment A in Table A-4.

Data qualification to the analytes associated with the specific LCS %Rs or RPDs was as follows:

Nonconformances <sup>1</sup>	Action	
	Detected Compounds	Nondetected Compounds
%R or RPD > UL	J	No qualification
%R < LL	J	UJ
%R < 20% (see note 1) (LL = lower limit, UL = upper limit)	J	R
Notes:		
1. Based on NFG 2008 VOC guidance, professional judgment is used to reject (R) nondetects in all associated samples for any analyte with < 20% recovery. Also, professional judgment is used to estimate (UJ) rather the reject sample results previously negated (U) on the basis of blank contamination.		

Qualified sample results are shown in Table 1.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-FB-061614	WQ	TOTAL ORGANIC CARBON		1.0*	MG/L	U	bl
VPB147-FB-061614	WQ	2-BUTANONE		2.5	UG/L	UJ	c
VPB147-FB-061614	WQ	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-FB-061614	WQ	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-FB-061614	WQ	BROMODICHLOROMETHANE	0.70	0.50	UG/L	J	l
VPB147-FB-061614	WQ	CHLOROETHANE		1.0	UG/L	UJ	c
VPB147-FB-061614	WQ	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c
VPB147-GW-061314-458-460	WG	2-BUTANONE		2.5	UG/L	UJ	c
VPB147-GW-061314-458-460	WG	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-GW-061314-458-460	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-GW-061314-458-460	WG	CARBON DISULFIDE		1.0*	UG/L	U	bl
VPB147-GW-061314-458-460	WG	CHLOROETHANE		1.0	UG/L	UJ	c
VPB147-GW-061314-458-460	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c
VPB147-GW-061314-478-480	WG	2-BUTANONE		2.5	UG/L	UJ	c
VPB147-GW-061314-478-480	WG	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-GW-061314-478-480	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
VPB147-GW-061314-478-480	WG	CARBON DISULFIDE		1.0*	UG/L	U	bl
VPB147-GW-061314-478-480	WG	CHLOROETHANE		1.0	UG/L	UJ	c
VPB147-GW-061314-478-480	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c
VPB147-TRIP BLANKS-061614	WQ	2-BUTANONE		2.5	UG/L	UJ	c
VPB147-TRIP BLANKS-061614	WQ	2-HEXANONE		2.5	UG/L	UJ	c
VPB147-TRIP BLANKS-061614	WQ	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c



Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-TRIP BLANKS-061614	WQ	CHLOROETHANE		1.0	UG/L	UJ	c
VPB147-TRIP BLANKS-061614	WQ	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c

\*LOQ value

## Attachment A

### Nonconformance Summary Tables

**Table A-1 - Initial Calibration**

Calibration Date/Time	Compound	% RSD	Limits
09-June-2014	chloroethane	36	≤15%
Associated samples: all samples in SDG SH4349			

**Table A-2 -Continuing Calibration Verification Standard**

CCV ID	Compound	% D	Limits
WG145023-4	dichlorodifluoromethane	55	≤20%
	2-butanone	29	≤20%
	4-methyl-2-pentanone	37	≤20%
	2-hexanone	21	≤20%
Associated samples: all samples in SDG SH4349			

**Table A-3 - Lab Blanks**

Blank ID	Compound	Result	LOD	Units	Associated Samples
WG145514-1	TOTAL ORGANIC CARBON	0.21	0.50	MG/L	VPB147-FB-061614
WG145023-2	CARBON DISULFIDE	0.32	0.50	UG/L	VPB147-GW-061314-458-460, VPB147-GW-061314-478-480

**Table A-4 - Lab Control Samples**

LCS ID	Compound	LCS % Recovery	Lower Limit	Upper Limit	Associated Samples
WG145023-1	XYLENES, TOTAL	117	89	116	SDG SH4349
WG145023-1	BROMODICHLOROMETHANE	126	75	120	SDG SH4349
WG145023-1	DICHLORODIFLUOROMETHANE	166	30	155	SDG SH4349
WG145023-1	O-XYLENE	121	80	120	SDG SH4349

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

### Attachment C

#### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance





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Fax: (207) 775-4029

# CHAIN of CUSTODY

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Page \_\_\_ of \_\_\_

Client Resolution Consultants Contact Eleanore Vivadou Phone # ( ) Fax # ( )

Address 100 Reel Schoolhouse Road City Chestnut Ridge State NY Zip Code \_\_\_\_\_

Purchase Order # \_\_\_\_\_ Proj. Name / No. Beth page 60266526 Katahdin Quote # \_\_\_\_\_

Bill (if different than above) \_\_\_\_\_ Address \_\_\_\_\_

Sampler (Print / Sign) Valerie Thayer, Valerie Thayer Copies To: V. Thayer

LAB USE ONLY WORK ORDER #: 5H4349  
KATAHDIN PROJECT NUMBER \_\_\_\_\_

### ANALYSIS AND CONTAINER TYPE PRESERVATIVES

REMARKS: \_\_\_\_\_

SHIPPING INFO:  FED EX  UPS  CLIENT

AIRBILL NO: \_\_\_\_\_

TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

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<u>VOLs</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>8260 HCL</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>TDC</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>Small sulfuric acid</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>VPB147-GW-061314-478-480</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>VPB147-GW-061314-458-460</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>VPB147-GW-ms/msd-061314-458-460</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>Temp blank</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>VPB147-Tripblanks-061614</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>VPB147-FB-061614</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>
<u>VPB147-FB-061614</u>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>

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Relinquished By: (Signature) _____	Date / Time _____	Received By: (Signature) _____	Date / Time _____	Relinquished By: (Signature) _____	Date / Time _____	Received By: (Signature) _____

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061614 2



### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4349-4  
 Client ID: VPB147-FB-061614  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4349  
 Lab File ID: C7752.D

Sample Date: 16-JUN-14  
 Received Date: 18-JUN-14  
 Extract Date: 19-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145023

Analysis Date: 19-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.36	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Bromodichloromethane</b>	<del>U</del> J	0.70	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
<b>Dibromochloromethane</b>	J	0.85	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 12/12/14



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4349-4  
**Client ID:** VPB147-FB-061614  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4349  
**Lab File ID:** C7752.D

**Sample Date:** 16-JUN-14  
**Received Date:** 18-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145023

**Analysis Date:** 19-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	UL	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	<del>UL</del> U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	<del>UL</del> U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.0	%					
Toluene-d8		95.3	%					
1,2-Dichloroethane-d4		116.	%					
Dibromofluoromethane		101.	%					

R12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4349-2  
**Client ID:** 147-061314-458-460  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4349  
**Lab File ID:** C7760.D

**Sample Date:** 13-JUN-14  
**Received Date:** 18-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145023

**Analysis Date:** 19-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>UL</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	<del>J</del> U	<del>0.26</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	J	4.9	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	UL	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4349-2  
**Client ID:** 147-061314-458-460  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4349  
**Lab File ID:** C7760.D

**Sample Date:** 13-JUN-14  
**Received Date:** 18-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145023

**Analysis Date:** 19-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	UL	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	<del>UL</del> U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	<del>UL</del> U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		92.9	%					
Toluene-d8		94.7	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		98.3	%					

*R 12/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4349-1  
**Client ID:** 147-061314-478-480  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4349  
**Lab File ID:** C7759.D

**Sample Date:** 13-JUN-14  
**Received Date:** 18-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145023

**Analysis Date:** 19-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>UL</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	<del>J</del> U	<del>0.38</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	J	4.2	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>	J	0.45	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	UL	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 12/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4349-1  
**Client ID:** 147-061314-478-480  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4349  
**Lab File ID:** C7759.D

**Sample Date:** 13-JUN-14  
**Received Date:** 18-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145023

**Analysis Date:** 19-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	UL	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	UL U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	UL U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		96.0	%					
Toluene-d8		96.9	%					
1,2-Dichloroethane-d4		116.	%					
Dibromofluoromethane		103.	%					

R 12/14/12

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4349-3  
 Client ID: VPB147-TB-061614  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4349  
 Lab File ID: C7751.D

Sample Date: 16-JUN-14  
 Received Date: 18-JUN-14  
 Extract Date: 19-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145023

Analysis Date: 19-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	<del>U</del> U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4349-3  
**Client ID:** VPB147-TB-061614  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4349  
**Lab File ID:** C7751.D

**Sample Date:** 16-JUN-14  
**Received Date:** 18-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145023

**Analysis Date:** 19-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 20-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	UL	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	<del>UL</del> U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	<del>UL</del> U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		94.3	%					
Toluene-d8		95.6	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		98.0	%					

*6/21/14*



ANALYTICAL SERVICES



Cert No E87604

### Report of Analytical Results

**Client:** Rick Purdy  
AECOM  
701 Edgewater Drive  
Wakefield, MA 01880

**Lab Sample ID:** SH4349-4  
**Report Date:** 01-JUL-14  
**Client PO:** 16518  
**Project:** Navy Clean WE15-03-0  
**SDG:** SH4349

**Sample Description**  
VPB147-FB-061614

**Matrix** AQ  
**Date Sampled** 16-JUN-14  
**Date Received** 18-JUN-14

Parameter	Result	Adj LOQ	Adj MDL	Adj LOD	Anal. Method	QC Batch	Anal. Date	Prep. Method	Prep. Date	Footnotes
Total Organic Carbon	10.28 mg/L <b>1.0 U</b>	1.0	0.10	.5	SM5310B	WG145514	26-JUN-14 20:14:04	N/A	N/A	N/A

12/12/14







## Data Validation Report

Project: Regional Groundwater Investigation - NWIRP Bethpage

Laboratory: KATAHDIN ANALYTICAL

Service Request: SH4407

Analyses/Method: EPA SW-846 Method 8260B for VOCs (GC/MS)

Validation Level: Limited

AECOM Project Number: 60266526.SA.DV

Prepared by: Dawn Brule/RESCON Completed on: 09/24/2014

Reviewed by: Paula DiMattei/RESCON File Name: SH4407\_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 17, 2014, June 18, 2014 and May 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-GW-061714-528-530	Ground water
VPB147-GW-061714-538-540	Ground water
VPB147-GW-061714-558-560	Ground water
VPB147-GW-061814-578-580	Ground water
VPB147-GW-061814-598-600	Ground water
VPB147-GW-061814-618-620	Ground water
VPB147-TRIP BLANK-061814	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✗ Initial calibration/continuing calibration verification
- ✗ Laboratory blanks/trip blanks/equipment blanks
- ✓ Surrogate spike recoveries

- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicates
- ✓ Internal standards
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated or negated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

Nonconformances are summarized in Attachment A in Table A-3.

Blank type	Blank result	Sample result	Action for samples
Method, Storage, Field, Trip, or Instrument*	Detects	Not detected	No qualification
	≤ LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and ≤ 2x LOQ	Report the sample result with a U**
		≥ 2x the LOQ	No qualifications
	> LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and < blank contamination	Report the sample result with a U or reject the sample result as unusable R
		≥ LOQ and ≥ blank contamination	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
	TIC detected	Detects	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
* Qualifications based on instrument blank results affect only the sample analyzed immediately after the sample that has target compounds that exceed the calibration range or non-target compounds that exceed 100 g/L.			
**Based on professional judgment.			

LOQ - Limit of Quantitation.

Qualified sample results are shown in Table 1.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

**Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

**Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

**QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-061714-528-530	WG	2-BUTANONE	5.0	2.5	UG_L	J	c
VPB147-GW-061714-528-530	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-528-530	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-528-530	WG	CARBON DISULFIDE		1.0	UG_L	U	bt
VPB147-GW-061714-528-530	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061714-528-530	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061714-538-540	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-538-540	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-538-540	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-538-540	WG	CARBON DISULFIDE		1.0	UG_L	U	bt
VPB147-GW-061714-538-540	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061714-538-540	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061714-558-560	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-558-560	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-558-560	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061714-558-560	WG	CARBON DISULFIDE		1.0	UG_L	U	bt
VPB147-GW-061714-558-560	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061714-558-560	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061814-578-580	WG	2-BUTANONE	1.9	2.5	UG_L	J	c
VPB147-GW-061814-578-580	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-578-580	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-578-580	WG	CARBON DISULFIDE		1.0	UG_L	U	bt

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-061814-578-580	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061814-578-580	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061814-598-600	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-598-600	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-598-600	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-598-600	WG	CARBON DISULFIDE		1.0	UG_L	U	bt
VPB147-GW-061814-598-600	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061814-598-600	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061814-618-620	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-618-620	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-618-620	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061814-618-620	WG	CARBON DISULFIDE		1.0	UG_L	U	bt
VPB147-GW-061814-618-620	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061814-618-620	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-061814	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-061814	WQ	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-061814	WQ	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-061814	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-061814	WQ	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c



## Attachment A

## Non Conformance Summary Tables

Table A-1 - Initial Calibration

Calibration Date/Time	Compound	% R	Limits
09-June-2014 11:06	chloroethane	36	<15%
Associated samples: all samples in SDG SH4407			

Table A-2 -Continuing Calibration Verification Standard

CCV ID	Compound	% D	Limits
WG144656-4	dichlorodifluoromethane	61	<20%
	2-butanone	24	<20%
	4-methyl-2-pentanone	40	<20%
	2-hexanone	21	<20%
Associated samples: all samples in SDG SH4407			

Table A-3 - Field Blanks

Blank ID	Compound	Result	LOD	Units	Associated Samples
VPB147-TRIP BLANK-061814	CARBON DISULFIDE	0.25	0.50	UG_L	All samples in SDG SH4407

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

### Attachment C

#### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bt	Trip blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



600 Technology Way  
 Scarborough, ME 04074  
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 Fax: (207) 775-4029

# CHAIN of CUSTODY

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Page \_\_\_\_ of \_\_\_\_

Client <i>Resolution Consultants</i>	Contact <i>Eleanor Vivandout</i>	Phone # ( )	Fax # ( )
Address <i>100 Reed Schoolhouse Rd</i> City <i>Chestnut Ridge</i> State <i>NY</i>		Zip Code	
Purchase Order #	Proj. Name / No. <i>60266526 Bethpage</i>	Katahdin Quote #	
Bill (if different than above)		Address	
Sampler (Print / Sign) <i>Valerie Thayer Valerie Thayer</i>		Copies To: <i>Val Thayer</i>	

**LAB USE ONLY** WORK ORDER #: *SH4407*  
 KATAHDIN PROJECT NUMBER \_\_\_\_\_

REMARKS: \_\_\_\_\_

SHIPPING INFO:  FED EX  UPS  CLIENT

AIRBILL NO: \_\_\_\_\_

TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

					ANALYSIS AND CONTAINER TYPE PRESERVATIVES												
					Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	
					OY	ON	OY	ON	OY	ON	OY	ON	OY	ON	OY	ON	
					<i>VOCs</i>												
	<i>VPB147-GW-061714-528-530</i>	<i>06/17/2014</i>	<i>10:15</i>	<i>GW</i>	<i>1</i>	<i>X</i>											
	<i>VPB147-GW-061714-538-540</i>	<i>06/17/2014</i>	<i>13:00</i>	<i>GW</i>	<i>4</i>	<i>X</i>											
	<i>VPB147-GW-061814-548-600</i>	<i>6/18/2014</i>	<i>13:00</i>	<i>GW</i>	<i>4</i>	<i>X</i>											
	<i>VPB147-GW-061814-578-580</i>	<i>6/18/2014</i>	<i>10:15</i>	<i>GW</i>	<i>3</i>	<i>X</i>											
	<i>VPB147-GW-061714-558-560</i>	<i>6/17/2014</i>	<i>15:00</i>	<i>GW</i>	<i>4</i>	<i>X</i>											
	<i>VPB147-GW-061814-618-620</i>	<i>6/18/2014</i>	<i>13:30</i>	<i>GW</i>	<i>6</i>	<i>X</i>											
	<i>Temp blank</i>	<i>06/18/2014</i>	<i>11:00</i>	<i>W</i>	<i>1</i>	<i>X</i>											
	<i>VPB147-Trip blanks-061814</i>	<i>6/18/2014</i>	<i>11:00</i>	<i>W</i>	<i>3</i>	<i>X</i>											

COMMENTS

Relinquished By: (Signature) <i>Valerie Thayer</i>	Date / Time <i>6/18/2014 7-</i>	Received By: (Signature) <i>[Signature]</i>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

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## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-1  
**Client ID:** 147-061714-528-530  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7781.D

**Sample Date:** 17-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
<b>Chloromethane</b>	J	0.56	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J U	<del>0.32</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		29	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>2-Butanone</b>	J	5.0	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/18/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-1  
**Client ID:** 147-061714-528-530  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7781.D

**Sample Date:** 17-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		100.	%					
Toluene-d8		98.1	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		101.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-2  
**Client ID:** 147-061714-538-540  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7782.D

**Sample Date:** 17-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	U U	<del>0.66</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		6.3	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R. 12/18/14*



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-2  
**Client ID:** 147-061714-538-540  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7782.D

**Sample Date:** 17-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.2	%					
Toluene-d8		93.4	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		100.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-5  
**Client ID:** 147-061714-558-560  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7785.D

**Sample Date:** 17-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U W	<del>0.30</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone		19	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-5  
**Client ID:** 147-061714-558-560  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7785.D

**Sample Date:** 17-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		99.6	%					
Toluene-d8		97.4	%					
1,2-Dichloroethane-d4		115.	%					
Dibromofluoromethane		99.8	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-4  
**Client ID:** 147-061814-578-580  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7784.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
<b>Chloromethane</b>	J	0.43	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J U	<del>0.65</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		14	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>2-Butanone</b>	J J	1.9	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Rec 6/21/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4407-4  
 Client ID: 147-061814-578-580  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4407  
 Lab File ID: C7784.D

Sample Date: 18-JUN-14  
 Received Date: 19-JUN-14  
 Extract Date: 20-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145087

Analysis Date: 20-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		96.4	%					
Toluene-d8		96.0	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		101.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-3  
**Client ID:** 147-061814-598-600  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7783.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U U	<del>0.36</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	3.7	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-3  
**Client ID:** 147-061814-598-600  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7783.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		94.5	%					
Toluene-d8		95.0	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		97.4	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-6  
**Client ID:** 147-061814-618-620  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7786.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	<del>U</del> U	<del>0.34</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		7.2	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-6  
**Client ID:** 147-061814-618-620  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7786.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		96.9	%					
Toluene-d8		97.3	%					
1,2-Dichloroethane-d4		116.	%					
Dibromofluoromethane		100.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-7  
**Client ID:** VPB147-TB-061814  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7775.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J	0.25	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/18/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4407-7  
**Client ID:** VPB147-TB-061814  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4407  
**Lab File ID:** C7775.D

**Sample Date:** 18-JUN-14  
**Received Date:** 19-JUN-14  
**Extract Date:** 20-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145087

**Analysis Date:** 20-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 21-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.4	%					
Toluene-d8		96.9	%					
1,2-Dichloroethane-d4		111.	%					
Dibromofluoromethane		98.9	%					



## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	KATAHDIN ANALYTICAL	
Service Request:	SH4476	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS)	
Validation Level:	Limited	
AECOM Project Number:	60266526.SA.DV	
Prepared by:	Dawn Brule/RESCON	Completed on: 09/25/2014
Reviewed by:	Paula DiMattei/RESCON	File Name: SH4476_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 19, 2014 and May 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-GW-061914-638-640	Ground water
VPB147-GW-061914-668-670	Ground water
VPB147-TRIP BLANKS-061914	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✗ Initial calibration/continuing calibration verification
- ✓ Laboratory blanks/trip blanks/equipment blanks
- ✓ Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicates
- ✓ Internal standards

✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus AECOM professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met or qualification of the sample data was not required..

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

## **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

## **ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-061914-638-640	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-061914-638-640	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061914-638-640	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061914-638-640	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061914-638-640	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061914-638-640	WG	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061914-668-670	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-061914-668-670	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-061914-668-670	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-061914-668-670	WG	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-GW-061914-668-670	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-061914-668-670	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANKS-061914	WQ	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANKS-061914	WQ	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANKS-061914	WQ	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANKS-061914	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANKS-061914	WQ	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANKS-061914	WQ	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c



## Attachment A

## Non Conformance Summary Tables

Table A-1 - Initial Calibration

Calibration Date/Time	Compound	% R	Limits
09-June-2014 11:06	chloroethane	36	<15%
Associated samples: all samples within SDG SH4476			

Table A-2 – Continuing Calibration Verification Standard

CCV ID	Compound	% D	Limits
WG145161-4	dichlorodifluoromethane	65	<20%
	Trichlorofluoromethane	21	<20%
	2-butanone	32	<20%
	4-methyl-2-pentanone	56	<20%
	2-hexanone	35	<20%
Associated samples: VPB147-TRIP BLANK-061914,VPB147-GW-061914-638-640			
WG145199-4	dichlorodifluoromethane	36	<20%
	bromomethane	-35	<20%
	chloroethane	-21	<20%
	2-butanone	32	<20%
	4-methyl-2-pentanone	41	<20%
	2-hexanone	26	<20%
Associated sample: VPB147-GW-061914-668-670			

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

## Attachment C

## Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



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# CHAIN of CUSTODY

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Page \_\_\_\_ of \_\_\_\_

Client: Resolution Consultants Contact: Eleanor Vianaclou Phone #: ( ) Fax #: ( )  
 Address: 100 Reel Schoolhouse Rd City: Chestnut Ridge State: NY Zip Code: \_\_\_\_\_  
 Purchase Order #: \_\_\_\_\_ Proj. Name / No.: Bethpage, 100266526 Katahdin Quote #: \_\_\_\_\_  
 Bill (if different than above) Address: \_\_\_\_\_  
 Sampler (Print / Sign): Valerie Thayer Valerie Thayer Copies To: V Thayer

LAB USE ONLY WORK ORDER #: 544476  
 KATAHDIN PROJECT NUMBER: \_\_\_\_\_  
 REMARKS: \_\_\_\_\_  
 SHIPPING INFO:  FED EX  UPS  CLIENT  
 AIRBILL NO: \_\_\_\_\_  
 TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

LAB USE ONLY					ANALYSIS AND CONTAINER TYPE PRESERVATIVES											
					Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.
					YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON	YOY ON
*	Sample Description	Date / Time coll'd	Matrix	No. of Cntrs.	VOCs 8260											
	VPB147-Trip blanks - 061914	5/21/14 11:00	GW	3	X											
	VPB147-GW-061914-668-670	4/9/14 4:30 pm	GW	4	X											
	VPB147-GW-061914-638-640	6/19/14 11:00	GW	6	X											
	Temp blank	/	W	1												

COMMENTS

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>6/19/14</u>	Received By: (Signature) <u>[Signature]</u>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

THE TERMS AND CONDITIONS ON THE REVERSE SIDE HEREOF SHALL GOVERN SERVICES, EXCEPT WHEN A SIGNED CONTRACTUAL AGREEMENT EXISTS.

ORIGINAL



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4476-3  
**Client ID:** 147-061914-638-640  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4476  
**Lab File ID:** C7816.D

**Sample Date:** 19-JUN-14  
**Received Date:** 20-JUN-14  
**Extract Date:** 21-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145161

**Analysis Date:** 21-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		8.8	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/18/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4476-3  
**Client ID:** 147-061914-638-640  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4476  
**Lab File ID:** C7816.D

**Sample Date:** 19-JUN-14  
**Received Date:** 20-JUN-14  
**Extract Date:** 21-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145161

**Analysis Date:** 21-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		100.	%					
Toluene-d8		99.3	%					
1,2-Dichloroethane-d4	*	120.	%					
Dibromofluoromethane		106.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4476-2RA  
**Client ID:** 147-061914-668-670  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4476  
**Lab File ID:** C7827.D

**Sample Date:** 19-JUN-14  
**Received Date:** 20-JUN-14  
**Extract Date:** 22-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145199

**Analysis Date:** 22-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Handwritten signature/initials: G. 2/29/15*



### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4476-2RA  
**Client ID:** 147-061914-668-670  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4476  
**Lab File ID:** C7827.D

**Sample Date:** 19-JUN-14  
**Received Date:** 20-JUN-14  
**Extract Date:** 22-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145199

**Analysis Date:** 22-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	UL U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		99.5	%					
Toluene-d8		98.4	%					
1,2-Dichloroethane-d4		118.	%					
Dibromofluoromethane		101.	%					

*Riz/10/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4476-1  
**Client ID:** VPB147-TB-061914  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4476  
**Lab File ID:** C7801.D

**Sample Date:** 19-JUN-14  
**Received Date:** 20-JUN-14  
**Extract Date:** 21-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145161

**Analysis Date:** 21-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R12/18/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4476-1  
 Client ID: VPB147-TB-061914  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4476  
 Lab File ID: C7801.D

Sample Date: 19-JUN-14  
 Received Date: 20-JUN-14  
 Extract Date: 21-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145161

Analysis Date: 21-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.7	%					
Toluene-d8		95.8	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		99.4	%					



## Data Validation Report

Project: Regional Groundwater Investigation - NWIRP Bethpage  
Laboratory: KATAHDIN ANALYTICAL  
Service Request: SH4576  
Analyses/Method: EPA SW-846 Method 8260B for VOCs (GC/MS)  
Validation Level: Limited  
AECOM Project 60266526.SA.DV  
Number:  
Prepared by: Dawn Brule/RESCON Completed on: 09/25/2014  
Reviewed by: Paula DiMattei/RESCON File Name: SH4576\_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 20, 2014, June 23, 2014 and May 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-GW-062014-678-680	Ground water
VPB147-GW-062314-718-720	Ground water
VPB147-GW-062314-738-740	Ground water
VPB147-TRIP BLANK-062314	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✗ Initial calibration/continuing calibration verification
- ✓ Laboratory blanks/trip blanks/equipment blanks
- ✓ Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicates

- ✓ Internal standards
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## **RESULTS**

### **Data Completeness**

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### **Holding Times/Sample Preservation**

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

**Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

**MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

**LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

**Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

**Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

**Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

**QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062014-678-680	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-062014-678-680	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-062014-678-680	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-062014-678-680	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-062014-678-680	WG	CYCLOHEXANE		0.50	UG_L	UJ	c
VPB147-GW-062014-678-680	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-062014-678-680	WG	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	CYCLOHEXANE		0.50	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-062314-718-720	WG	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	CYCLOHEXANE		0.50	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-GW-062314-738-740	WG	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-062314	WQ	2-BUTANONE		2.5	UG_L	UJ	c



Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-TRIP BLANK-062314	WQ	2-HEXANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-062314	WQ	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-062314	WQ	CHLOROETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-062314	WQ	CYCLOHEXANE		0.50	UG_L	UJ	c
VPB147-TRIP BLANK-062314	WQ	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-062314	WQ	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	c

## Attachment A

### Non Conformance Summary Tables

**Table A-1 - Initial Calibration**

Calibration Date/Time	Compound	% R	Limits
09-June-2014 11:06	chloroethane	36	<15%
Associated samples: all samples in SDG SH4576			

**Table A-2 -Continuing Calibration Verification Standard**

CCV ID	Compound	% R	Limits
WG145394-4	dichlorodifluoromethane	38	<20%
	trichlorofluoromethane	33	<20%
	2-butanone	40	<20%
	cyclohexane	21	<20%
	4-methyl-2-pentanone	46	<20%
	2-hexanone	28	<20%
Associated samples: all samples in SDG SH4576			

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

## Attachment C

### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



600 Technology Way  
 Scarborough, ME 04074  
 Tel: (207) 874-2400  
 Fax: (207) 775-4029

# CHAIN of CUSTODY

PLEASE BEAR DOWN AND  
 PRINT LEGIBLY IN PEN

Page \_\_\_\_ of \_\_\_\_

Client: Resolution Consultants Contact: Eleanor Vivandou Phone #: ( ) Fax #: ( )  
 Address: 100 red schoolhouse Rd City: Chestnut Ridge State: NY Zip Code: \_\_\_\_\_  
 Purchase Order #: \_\_\_\_\_ Proj. Name / No.: Bethpage 60266526 Katahdin Quote #: \_\_\_\_\_  
 Bill (if different than above): \_\_\_\_\_ Address: \_\_\_\_\_

Sampler (Print / Sign): Valerie Thayer, Valerie Thayer Copies To: Valerie Thayer

LAB USE ONLY WORK ORDER #: 514576  
 KATAHDIN PROJECT NUMBER: \_\_\_\_\_

ANALYSIS AND CONTAINER TYPE PRESERVATIVES										
Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.
OY	ON	OY	ON	OY	ON	OY	ON	OY	ON	OY

REMARKS: \_\_\_\_\_  
 SHIPPING INFO:  FED EX  UPS  CLIENT  
 AIRBILL NO: \_\_\_\_\_  
 TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

* Sample Description	Date / Time coll'd	Matrix	No. of Cntrs.
<u>Temp</u>	<u>/</u>	<u>W</u>	<u>1</u>
<u>VPB147-GW-062014-678-680</u>	<u>6/20/10:30</u>	<u>GW</u>	<u>2</u>
<u>VPB147-Trip Blank-062314</u>	<u>05/2/2014 11:00</u>	<u>W</u>	<u>3</u>
<u>VPB147-GW-062314-718-720</u>	<u>06/22/2014 11:15</u>	<u>GW</u>	<u>4</u>
<u>VPB147-GW-062314-738-740</u>	<u>6/23/2014 14:00</u>	<u>GW</u>	<u>4</u>

<u>VPB 8260</u>										

COMMENTS

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>6/23/14 7</u>	Received By: (Signature) <u>[Signature]</u>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-1  
**Client ID:** 147-062014-678-680  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7908.D

**Sample Date:** 20-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone		7.2	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U UJ	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*6/26/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-1  
**Client ID:** 147-062014-678-680  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7908.D

**Sample Date:** 20-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		88.9	%					
Toluene-d8		90.7	%					
1,2-Dichloroethane-d4		118.	%					
Dibromofluoromethane		99.7	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-3  
**Client ID:** 147-062314-718-720  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7909.D

**Sample Date:** 23-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone		15	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R. 2/18/14*



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-3  
**Client ID:** 147-062314-718-720  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7909.D

**Sample Date:** 23-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		90.5	%					
Toluene-d8		90.4	%					
1,2-Dichloroethane-d4		116.	%					
Dibromofluoromethane		98.9	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-4  
**Client ID:** 147-062314-738-740  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7910.D

**Sample Date:** 23-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>		8.0	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Riz 1/8/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-4  
**Client ID:** 147-062314-738-740  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7910.D

**Sample Date:** 23-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.4	%					
Toluene-d8		92.4	%					
1,2-Dichloroethane-d4		118.	%					
Dibromofluoromethane		102.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-2  
**Client ID:** 147-TB-062314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7906.D

**Sample Date:** 23-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	⊕ UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	⊕ UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	⊕ UJ	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	⊕ UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	⊕ UJ	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	⊕ UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	⊕ UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

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## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4576-2  
**Client ID:** 147-TB-062314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4576  
**Lab File ID:** C7906.D

**Sample Date:** 23-JUN-14  
**Received Date:** 24-JUN-14  
**Extract Date:** 25-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145394

**Analysis Date:** 25-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 26-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		88.4	%					
Toluene-d8		91.3	%					
1,2-Dichloroethane-d4		119.	%					
Dibromofluoromethane		99.7	%					



## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	Katahdin Analytical	
Service Request:	SH4749	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS) and Standard Method 5310 for Total Organic Carbon by High-Temperature Combustion	
Validation Level:	3	
AECOM Project Number:	60266526.SA.DV	
Prepared by:	Dawn Brule/RESCON	Completed on: 10/02/2014
Reviewed by:	Lori Herberich/RESCON	File Name: SH4749_5310B and 8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 24 - 26, 2014 and May 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-EB-062514	Equipment blank (TOC)
VPB147-EB-062514	Equipment blank (VOC)
VPB147-GW-062414-758-760	Groundwater
VPB147-GW-062514-798-800	Groundwater
VPB147-GW-062514-818-820	Groundwater
VPB147-GW-062614-840-842	Groundwater
VPB147-GW-062614-858-860	Groundwater
VPB147-TRIPBLANKS-062614	Trip Blank

The samples were analyzed in accordance with:

- *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW-846, Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (USEPA, 1996).*
- *Standard Methods for the Examination of Water and Wastewater, Method SM5310B, Total Organic Carbon by High-Temperature Combustion*

Data validation activities were conducted with reference to these methods, *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008)*, *USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review (January 2010)*, and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2 (DoD, October 2010)* where applicable. In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- X Data completeness (chain-of-custody [COC])/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification
- X Laboratory blanks/equipment blanks/trip blanks
- ✓ Surrogate spike recoveries
- ✓ Matrix spike (MS) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicate results
- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated or negated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### **Data Completeness (COC)/Sample Integrity**

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Selected samples were mostly soil and had very little standing water.

- For samples VPB147-GW-062614-840-842, VPB147-GW-062414-758-760, and VPB147-GW-062614-858-860, the laboratory decanted the liquid from one vial each prior to analysis.
- For sample VPB147-GW-062514-818-820 the laboratory decanted the water from two individual vials into one vial as a composite. The sample was analyzed at a dilution because of insufficient volume.

Positive and nondetect results for these sample were qualified as estimated (J and UJ) respectively, due to possible loss of sample integrity during the decanting procedure. Qualified sample results are shown in Table 1.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### **Holding Times and Sample Preservation**

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria. The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met. The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and/or
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1, A-2, and A-3.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

#### **ICV Recovery Nonconformances:**

Nonconformance	Actions	
	Detected Compounds	Nondetected Compounds
%R > 120%	J	No qualification
20% < %R < 80%	J	UJ
%R < 20% (see note)	J	R*

Notes: Based on NFG 2008 VOC guidance, professional judgment is used to reject (R) nondetects in all associated samples for any analyte with < 20% recovery. Also, professional judgment is used to estimate (UJ) rather the reject (R) sample results previously negated (U) on the basis of blank contamination.



**CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift >20%	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

**Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL).

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method, equipment rinsate and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

Nonconformances are summarized in Attachment A in Tables A-4 and A-5.

Sample results were qualified as follows:

For common lab contaminants (methylene chloride, acetone, 2-butanone):

Blank type	Blank result	Sample result	Action for samples
Method, Storage, Field, Trip, or Instrument*	Detects	Not detected	No qualification
		< 2x LOQ	Report sample LOQ value with a U
		≥ 2x LOQ and ≤ 4x LOQ	Report the sample result with a U**
	> 2x LOQ	≥ 4x LOQ	No qualifications
		< 2x LOQ	Report sample LOQ value with a U
		≥ 2x LOQ and < blank contamination	Report the sample result with a U
	≥ 2x LOQ and ≥ blank contamination	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**	
* Qualifications based on instrument blank results affect only the sample analyzed immediately after the sample that has target compounds that exceed the calibration range or non-target compounds that exceed 100 g/L.			
**Based on professional judgment			

For TOC analysis:

Blank Type	Blank Result	Sample Result	Action for Samples
ICB/CCB (Positive)	≥DL but ≤ LOQ	Nondetect	No action
		≥DL but ≤LOQ	Qualify as nondetect (U) at the LOQ

Blank Type	Blank Result	Sample Result	Action for Samples
	>LOQ	> LOQ	Use professional judgment (see below [1])
		$\geq$ DL but $\leq$ LOQ	Qualify as nondetect (U) at the LOQ
		> LOQ but < ICB/CCB Result	Qualify at level of Blank Result with a "U" or Qualify result as unusable
		>ICB/CCB but <10x the ICB/CCB result	Qualify as estimated (J)
		$\geq$ 10x ICB/CCB	No action is taken based on professional judgment
PB / EB/ FB (Positive)	> LOQ	$\geq$ DL but $\leq$ LOQ	Qualify as nondetect (U) at the LOQ
		>LOQ but < 10x Blank Result	Qualify results as unusable
		$\geq$ 10x Blank Result	No action
	$\geq$ DL but $\leq$ LOQ	Nondetect	No action
		$\geq$ DL but $\leq$ LOQ	Qualify as nondetect (U) at the LOQ
		> LOQ	Use professional judgment (see below [1])

[1] Establish an action level (AL) at 5x the blank contamination. If sample result is <AL, qualify the reported result with a U.  
LOQ - Limit of Quantitation

Qualified sample results are shown in Table 1.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **MS Results**

The MS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

### **ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-EB-062514	WQ	TOTAL ORGANIC CARBON		1.0*	MG/L	U	bl
VPB147-EB-062514-2	WQ	1,1-DICHLOROETHENE		0.50	UG/L	UJ	c
VPB147-EB-062514-2	WQ	BROMOMETHANE		1.0	UG/L	UJ	c
VPB147-GW-062414-758-760	WG	1,1,1-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,1,2,2-TETRACHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,1,2-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,1-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,1-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2,4-TRICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2-DIBROMOETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2-DICHLOROETHENE, TOTAL		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,2-DICHLOROPROPANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,3-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	1,4-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	2-BUTANONE		2.5	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	2-HEXANONE		2.5	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	ACETONE	24	2.5	UG/L	J	mc,c

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062414-758-760	WG	BENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	BROMODICHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	BROMOFORM		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	BROMOMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CARBON DISULFIDE	0.58	0.50	UG/L	J	mc
VPB147-GW-062414-758-760	WG	CARBON TETRACHLORIDE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CHLOROFORM		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CHLOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CIS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CIS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	DIBROMOCHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	ETHYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	ISOPROPYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	M- AND P-XYLENE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	METHYL ACETATE		0.75	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	METHYL CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	METHYL TERT-BUTYL ETHER		0.50	UG/L	UJ	mc,c
VPB147-GW-062414-758-760	WG	METHYLENE CHLORIDE		2.5	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062414-758-760	WG	O-XYLENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	STYRENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	TETRACHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	TOLUENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	TRANS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	TRANS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	TRICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	TRICHLOROFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	VINYL CHLORIDE		1.0	UG/L	UJ	mc
VPB147-GW-062414-758-760	WG	XYLENES, TOTAL		1.5	UG/L	UJ	mc
VPB147-GW-062514-798-800	WG	1,1-DICHLOROETHENE		0.50	UG/L	UJ	c
VPB147-GW-062514-798-800	WG	BROMOMETHANE		1.0	UG/L	UJ	c
VPB147-GW-062514-818-820	WG	1,1,1-TRICHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,1,2,2-TETRACHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,1,2-TRICHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,1-DICHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,1-DICHLOROETHENE		1.0	UG/L	UJ	mc,c
VPB147-GW-062514-818-820	WG	1,2,4-TRICHLOROBENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,2-DIBROMO-3-CHLOROPROPANE		1.5	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,2-DIBROMOETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,2-DICHLOROBENZENE		1.0	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062514-818-820	WG	1,2-DICHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,2-DICHLOROETHENE, TOTAL		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,2-DICHLOROPROPANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,3-DICHLOROBENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	1,4-DICHLOROBENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	2-BUTANONE		5.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	2-HEXANONE		5.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	4-METHYL-2-PENTANONE		5.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	ACETONE		34**	UG/L	UJ	mc,be
VPB147-GW-062514-818-820	WG	BENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	BROMODICHLOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	BROMOFORM		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	BROMOMETHANE		2.0	UG/L	UJ	mc,c
VPB147-GW-062514-818-820	WG	CARBON DISULFIDE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CARBON TETRACHLORIDE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CHLOROBENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CHLOROETHANE		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CHLOROFORM		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CHLOROMETHANE		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CIS-1,2-DICHLOROETHENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CIS-1,3-DICHLOROPROPENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	CYCLOHEXANE		1.0	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062514-818-820	WG	DIBROMOCHLOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	DICHLORODIFLUOROMETHANE		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	ETHYLBENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	ISOPROPYLBENZENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	M- AND P-XYLENE		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	METHYL ACETATE		1.5	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	METHYL CYCLOHEXANE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	METHYL TERT-BUTYL ETHER		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	METHYLENE CHLORIDE		5.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	O-XYLENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	STYRENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	TETRACHLOROETHENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	TOLUENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	TRANS-1,2-DICHLOROETHENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	TRANS-1,3-DICHLOROPROPENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	TRICHLOROETHENE		1.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	TRICHLOROFLUOROMETHANE		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	VINYL CHLORIDE		2.0	UG/L	UJ	mc
VPB147-GW-062514-818-820	WG	XYLENES, TOTAL		3.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,1,1-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,1,2,2-TETRACHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		0.50	UG/L	UJ	mc



Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062614-840-842	WG	1,1,2-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,1-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,1-DICHLOROETHENE		0.50	UG/L	UJ	mc,c
VPB147-GW-062614-840-842	WG	1,2,4-TRICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,2-DIBROMOETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,2-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,2-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,2-DICHLOROETHENE, TOTAL		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,2-DICHLOROPROPANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,3-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	1,4-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	2-BUTANONE		2.5	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	2-HEXANONE		2.5	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	ACETONE		5.0*	UG/L	UJ	mc,be
VPB147-GW-062614-840-842	WG	BENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	BROMODICHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	BROMOFORM		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	BROMOMETHANE		1.0	UG/L	UJ	mc,c
VPB147-GW-062614-840-842	WG	CARBON DISULFIDE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CARBON TETRACHLORIDE		0.50	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062614-840-842	WG	CHLORO BENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CHLOROFORM		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CHLOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CIS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CIS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	DIBROMOCHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	ETHYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	ISOPROPYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	M- AND P-XYLENE		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	METHYL ACETATE		0.75	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	METHYL CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	METHYL TERT-BUTYL ETHER		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	METHYLENE CHLORIDE		2.5	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	O-XYLENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	STYRENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	TETRACHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	TOLUENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	TRANS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	TRANS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062614-840-842	WG	TRICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	TRICHLOROFUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	VINYL CHLORIDE		1.0	UG/L	UJ	mc
VPB147-GW-062614-840-842	WG	XYLENES, TOTAL		1.5	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,1,1-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,1,2,2-TETRACHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,1,2-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,1-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,1-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2,4-TRICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2-DIBROMOETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2-DICHLOROETHENE, TOTAL		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,2-DICHLOROPROPANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,3-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	1,4-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	2-BUTANONE		2.5	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	2-HEXANONE		2.5	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062614-858-860	WG	ACETONE		12**	UG/L	UJ	mc,be
VPB147-GW-062614-858-860	WG	BENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	BROMODICHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	BROMOFORM		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	BROMOMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CARBON DISULFIDE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CARBON TETRACHLORIDE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CHLOROETHANE		1.0	UG/L	UJ	mc,c
VPB147-GW-062614-858-860	WG	CHLOROFORM		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CHLOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CIS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CIS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	DIBROMOCHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	ETHYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	ISOPROPYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	M- AND P-XYLENE		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	METHYL ACETATE		0.75	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	METHYL CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	METHYL TERT-BUTYL ETHER		0.50	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062614-858-860	WG	METHYLENE CHLORIDE		2.5	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	O-XYLENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	STYRENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	TETRACHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	TOLUENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	TRANS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	TRANS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	TRICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	TRICHLOROFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	VINYL CHLORIDE		1.0	UG/L	UJ	mc
VPB147-GW-062614-858-860	WG	XYLENES, TOTAL		1.5	UG/L	UJ	mc
VPB147-TRIPBLANKS-062614	WQ	1,1-DICHLOROETHENE		0.50	UG/L	UJ	c
VPB147-TRIPBLANKS-062614	WQ	BROMOMETHANE		1.0	UG/L	UJ	c

\*LOQ

\*\*sample value

## Attachment A

## Nonconformance Summary Tables

Table A-1 - Initial Calibration

Calibration Date/Time	Compound	% RSD	Limits
01-July-2014 09:21	chloroethane	16	≤15%
Associated sample: VPB147-GW-062614-858-860			

Table A-2 - Initial Calibration Verification Standard

ICV ID	Compound	% R	Limits
WG144515-7	acetone	125	80-120%
Associated samples: VPB147-EB-062514,VPB147-TRIP BLANKS-062614,VPB147-GW-062614-840-842,VPB147-GW-062514-798-800,VPB147-GW-062514-818-820,VPB147-GW-062414-758-760			

Table A-3 -Continuing Calibration Verification Standard

CCV ID	Compound	% D	Limits
WG145586-4	bromomethane	-24	≤20%
	1,1-dichloroethene	-21	≤20%
Associated samples: VPB147-EB-062514,VPB147-TRIP BLANKS-062614,VPB147-GW-062614-840-842,VPB147-GW-062514-798-800,VPB147-GW-062514-818-820			
WG145700-4	methyl tert-butyl ether	-21	≤20%
Associated sample: VPB147-GW-062414-758-760			

Table A-4 - Lab Blanks

Blank ID	Compound	Result	QL	Units	Associated Samples
WG146111-1	TOTAL ORGANIC CARBON	0.26	0.50	MG/L	VPB147-EB-062514

Table A-5 - Field Blanks

Blank ID	Compound	Result	QL	Units	Associated Samples
VPB147-EB-062514-2	ACETONE	3.5	2.5	UG/L	VPB147-GW-062614-840-842, VPB147-GW-062514-798-800, VPB147-GW-062514-818-820, VPB147-GW-062414-758-760, VPB147-GW-062614-858-860

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

## Attachment C

### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



Client: Resolution Consultants Contact: Eleanor Vitvudae Phone #: ( ) Fax #: ( )  
 Address: 100 Red Schoolhouse Rd City: Chestnut Ridge State: NY Zip Code: \_\_\_\_\_  
 Purchase Order #: \_\_\_\_\_ Proj. Name / No.: \_\_\_\_\_ Katahdin Quote #: \_\_\_\_\_

Bill (if different than above) Address: \_\_\_\_\_  
 Sampler (Print / Sign): Valerie Thayer Valerie Thayer Copies To: Valerie Thayer

LAB USE ONLY WORK ORDER #: 514749  
 KATAHDIN PROJECT NUMBER \_\_\_\_\_

ANALYSIS AND CONTAINER TYPE PRESERVATIVES

REMARKS: \_\_\_\_\_

SHIPPING INFO:  FED EX  UPS  CLIENT  
 AIRBILL NO: \_\_\_\_\_  
 TEMP °C  TEMP BLANK  INTACT  NOT INTACT

	Fill.	Fill.	Fill.	Fill.	Fill.	Fill.	Fill.	Fill.	Fill.	Fill.	Fill.
	OY ON	OY ON	OY ON	OY ON	OY ON	OY ON	OY ON	OY ON	OY ON	OY ON	OY ON
✓ VPB147-EB-062514											
VPB147-EB-062514											
✓ VPB147-Tripblanks - 062614											
✓ VPB147-GW-062614-840											
VPB147-GW-062514-798-800											
VPB147-GW-062414-758-760											
✓ VPB147-GW-062514-818-820											
Temp blank											
VPB147-GW-062614-858-860											

COMMENTS: EB = Equipment Rinse Blank  
\* mostly used

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>6/26/14 7-</u>	Received By: (Signature) <u>[Signature]</u> <u>6-27-14 09:15</u>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-2  
**Client ID:** VPB147-EB-062514  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9214.D

**Sample Date:** 25-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	J	3.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Handwritten signature/initials*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-2  
**Client ID:** VPB147-EB-062514  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9214.D

**Sample Date:** 25-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.3	%					
Toluene-d8		112.	%					
1,2-Dichloroethane-d4	*	120.	%					
Dibromofluoromethane	*	116.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-6RA  
**Client ID:** 147-062414-758-760  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9236.D

**Sample Date:** 24-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 30-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145700

**Analysis Date:** 30-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J	0.58	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	J	24	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Rizkiy*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-6RA  
**Client ID:** 147-062414-758-760  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9236.D

**Sample Date:** 24-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 30-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145700

**Analysis Date:** 30-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U <b>UJ</b>	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		96.4	%					
Toluene-d8		109.	%					
1,2-Dichloroethane-d4		101.	%					
Dibromofluoromethane		100.	%					

*Riz/18/17*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-5  
**Client ID:** 147-062514-798-800  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9220.D

**Sample Date:** 25-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R 12/18/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-5  
**Client ID:** 147-062514-798-800  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9220.D

**Sample Date:** 25-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.8	%					
Toluene-d8		108.	%					
1,2-Dichloroethane-d4	*	122.	%					
Dibromofluoromethane		114.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-7DL  
**Client ID:** 147-062514-818-820  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9216.D

**Sample Date:** 25-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	2.0	ug/L	2	2	4.0	0.48	2.0
Chloromethane	U	2.0	ug/L	2	2	4.0	0.72	2.0
Vinyl Chloride	U	2.0	ug/L	2	2	4.0	0.50	2.0
Bromomethane	U	2.0	ug/L	2	2	4.0	0.98	2.0
Chloroethane	U	2.0	ug/L	2	2	4.0	1.1	2.0
Trichlorofluoromethane	U	2.0	ug/L	2	2	4.0	0.48	2.0
1,1-Dichloroethene	U	1.0	ug/L	2	1	2.0	0.70	1.0
Carbon Disulfide	U	1.0	ug/L	2	1	2.0	0.50	1.0
Freon-113	U	1.0	ug/L	2	1	2.0	0.62	1.0
Methylene Chloride	U	5.0	ug/L	2	5	10.	2.3	5.0
<b>Acetone</b>		34	ug/L	2	5	10.	4.4	5.0
trans-1,2-Dichloroethene	U	1.0	ug/L	2	1	2.0	0.50	1.0
Methyl tert-butyl Ether	U	1.0	ug/L	2	1	2.0	0.72	1.0
1,1-Dichloroethane	U	1.0	ug/L	2	1	2.0	0.42	1.0
cis-1,2-Dichloroethene	U	1.0	ug/L	2	1	2.0	0.42	1.0
Chloroform	U	1.0	ug/L	2	1	2.0	0.64	1.0
1,1,1-Trichloroethane	U	1.0	ug/L	2	1	2.0	0.40	1.0
2-Butanone	U	5.0	ug/L	2	5	10.	2.6	5.0
Cyclohexane	U	1.0	ug/L	2	1	2.0	0.62	1.0
Carbon Tetrachloride	U	1.0	ug/L	2	1	2.0	0.44	1.0
Benzene	U	1.0	ug/L	2	1	2.0	0.52	1.0
1,2-Dichloroethane	U	1.0	ug/L	2	1	2.0	0.40	1.0
Trichloroethene	U	1.0	ug/L	2	1	2.0	0.56	1.0
1,2-Dichloropropane	U	1.0	ug/L	2	1	2.0	0.50	1.0
Bromodichloromethane	U	1.0	ug/L	2	1	2.0	0.66	1.0
cis-1,3-Dichloropropene	U	1.0	ug/L	2	1	2.0	0.38	1.0
Toluene	U	1.0	ug/L	2	1	2.0	0.54	1.0
4-Methyl-2-Pentanone	U	5.0	ug/L	2	5	10.	2.6	5.0
trans-1,3-Dichloropropene	U	1.0	ug/L	2	1	2.0	0.40	1.0
1,1,2-Trichloroethane	U	1.0	ug/L	2	1	2.0	0.66	1.0
Tetrachloroethene	U	1.0	ug/L	2	1	2.0	0.80	1.0
Dibromochloromethane	U	1.0	ug/L	2	1	2.0	0.60	1.0
2-Hexanone	U	5.0	ug/L	2	5	10.	3.4	5.0
Chlorobenzene	U	1.0	ug/L	2	1	2.0	0.44	1.0
Ethylbenzene	U	1.0	ug/L	2	1	2.0	0.42	1.0

*12/18/14*



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-7DL  
**Client ID:** 147-062514-818-820  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9216.D

**Sample Date:** 25-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U UJ	3.0	ug/L	2	3	6.0	0.50	3.0
Styrene	U	1.0	ug/L	2	1	2.0	0.46	1.0
Bromoform	U	1.0	ug/L	2	1	2.0	0.46	1.0
Isopropylbenzene	U	1.0	ug/L	2	1	2.0	0.46	1.0
1,1,2,2-Tetrachloroethane	U	1.0	ug/L	2	1	2.0	0.76	1.0
1,3-Dichlorobenzene	U	1.0	ug/L	2	1	2.0	0.52	1.0
1,4-Dichlorobenzene	U	1.0	ug/L	2	1	2.0	0.48	1.0
1,2-Dichlorobenzene	U	1.0	ug/L	2	1	2.0	0.30	1.0
1,2,4-Trichlorobenzene	U	1.0	ug/L	2	1	2.0	0.74	1.0
Methyl Acetate	U	1.5	ug/L	2	1	2.0	1.1	1.5
Methylcyclohexane	U	1.0	ug/L	2	1	2.0	0.60	1.0
o-Xylene	U	1.0	ug/L	2	1	2.0	0.50	1.0
M+P-Xylenes	U	2.0	ug/L	2	2	4.0	1.2	2.0
1,2-Dichloroethylene (Total)	U	2.0	ug/L	2	2	4.0	0.42	2.0
1,2-Dibromoethane	U	1.0	ug/L	2	1	2.0	0.44	1.0
1,2-Dibromo-3-Chloropropane	U	1.5	ug/L	2	1	2.0	1.0	1.5
P-Bromofluorobenzene		92.0	%					
Toluene-d8		108.	%					
1,2-Dichloroethane-d4		117.	%					
Dibromofluoromethane		110.	%					

*Riz/10/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-4  
**Client ID:** 147-062614-840-842  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9219.D

**Sample Date:** 26-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone		<del>7.3</del> 5.0	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-4  
**Client ID:** 147-062614-840-842  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9219.D

**Sample Date:** 26-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U UJ	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.0	%					
Toluene-d8		112.	%					
1,2-Dichloroethane-d4		118.	%					
Dibromofluoromethane		110.	%					

*Rizley*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-9  
**Client ID:** 147-062614-858-860  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9266.D

**Sample Date:** 26-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 01-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145769

**Analysis Date:** 01-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 02-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	L	12	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*G. 2/15/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-9  
**Client ID:** 147-062614-858-860  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9266.D

**Sample Date:** 26-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 01-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145769

**Analysis Date:** 01-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 02-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U JS	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		84.9	%					
Toluene-d8		111.	%					
1,2-Dichloroethane-d4		117.	%					
Dibromofluoromethane		105.	%					

*Riz/ks/ku*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-3  
**Client ID:** VPB174-TB-062614  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9215.D

**Sample Date:** 26-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> <i>U</i>	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	<del>U</del> <i>U</i>	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R. 12/18/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4749-3  
**Client ID:** VPB174-TB-062614  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4749  
**Lab File ID:** D9215.D

**Sample Date:** 26-JUN-14  
**Received Date:** 27-JUN-14  
**Extract Date:** 28-JUN-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145586

**Analysis Date:** 28-JUN-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 01-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		89.6	%					
Toluene-d8		106.	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		107.	%					

**Report of Analytical Results**

**Client:** Rick Purdy  
AECOM  
701 Edgewater Drive  
Wakefield, MA 01880

**Lab Sample ID:** SH4749-1  
**Report Date:** 11-JUL-14  
**Client PO:** 16518  
**Project:** Navy Clean WE15-03-0  
**SDG:** SH4749

**Sample Description:**  
VPB147-EB-062514

**Matrix:** AQ  
**Date Sampled:** 25-JUN-14  
**Date Received:** 27-JUN-14

Parameter	Result	Adj LOQ	Adj MDL	Adj LOD	Anal. Method	QC Batch	Anal. Date	Prep. Method	Prep. Date	Footnotes
Total Organic Carbon	10.22 mg/L <i>1.00</i>	1.0	0.10	.5	SM5310B	WG146111	30-JUN-14 19:10:03	N/A	N/A	

*6/12/14*





## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	Test America, South Burlington, Vermont	
Service Request:	200-22957	
Analyses/Method:	EPA Method TO-15, VOCs Collected in Canisters - GC/MS	
Validation Level:	3	
AECOM Project Number:	60266526.SA.DV	
Prepared by:	Dawn Brule/RESCON	Completed on: 10/06/2014
Reviewed by:	Lori Heberich/RESCON	File Name: 200-22957_TO-15

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 24, 2014.

Sample ID	Matrix/Sample Type
VPB147-AIR-062414	Ambient air

Data validation activities were conducted with reference to *Determination of Volatile Organic Compounds (VOCs) In Air Collected In Specially-Prepared Canisters and Analyzed By Gas Chromatography/Mass Spectrometry (GC/MS)* (USEPA, Method TO-15) and the *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- ✓ Data completeness (chain-of-custody [COC])/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- ✓ Initial calibration/continuing calibration verification
- ✓ Laboratory blanks
- NA Matrix duplicate (MD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicates
- ✓ Internal standards
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this

validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. There were no data points qualified or rejected on the basis of this data review.

## **RESULTS**

### **Data Completeness**

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

### **Holding Times/Sample Preservation**

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria. The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met. The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

The QC acceptance criteria were met.

### **Laboratory Blanks**

Laboratory method blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). Blank results were reviewed for conformance with the QC acceptance criteria. Data validation qualifications for individual samples are based on the maximum

contaminant concentration detected in all associated blanks. The QC acceptance criteria were met and qualification of the sample results was not required.

### **MD Results**

MD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as undetected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

No sample results were qualified as a result of this data review.

### **ATTACHMENTS**

Attachment A: Qualifier Codes and Explanations

**Attachment A****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

TestAmerica Burlington  
30 Community Drive  
Suite 11

South Burlington, VT 05403

phone 802-660-1990 fax 802-660-1919

## Canister Samples Chain of Custody Record

TestAmerica Analytical Testing Corp. assumes no liability with respect to the collection and shipment of these samples.

Client Contact Information		Samples Collected By: <i>Valerie Thayer</i>			of COCs																																								
Company: <i>Resolution Consultants</i>		Project Manager: <i>Eleanor Vivaudou</i>																																											
Address: <i>999 S. Duxbury Road</i>		Phone: <i>845-425-4980</i>																																											
City/State/Zip: <i>Bethel, NH</i>		Email: <i>Eleanor.vivaudou@resolution.com</i>																																											
Phone: <i>207-650-6572</i>		Site Contact: <i>Valerie Thayer</i>																																											
FAX:		TA Contact:																																											
Project Name: <i>Both Age</i>		Analysis Turnaround Time																																											
Site: <i>VP0147</i>		Standard (Specify)																																											
PO #		Rush (Specify)																																											
Sample Identification		Sample Date(s)	Time Start	Time Stop	Canister Vacuum In Field, Hg (Start)	Canister Vacuum In Field, Hg (Stop)	Flow Controller ID	Canister ID	MA-APH	TO-15	EPA 3C	EPA 25C	ASTM D-1946	Other (Please specify in notes section)	Landfill Gas	Soil Gas	Ambient Air	Indoor Air	Other (Please specify in notes section)																										
<i>VP0147-AIC-062-114</i>		<i>6/24/14</i>	<i>9:00 am</i>	<i>4:20 pm</i>	<i>31</i>	<i>70</i>	<i>2811</i>	<i>ESV #28597</i>																																					
<div style="display: flex; justify-content: space-between; align-items: center;"> <div style="text-align: left;"> <table border="1" style="border-collapse: collapse; width: 100%;"> <tr> <td style="width: 50%; padding: 5px;">Temperature (Fahrenheit)</td> <td style="width: 50%;"></td> </tr> <tr> <td style="padding: 5px;">Interior</td> <td></td> </tr> <tr> <td style="padding: 5px;">Ambient</td> <td></td> </tr> <tr> <td style="padding: 5px;">Start</td> <td></td> </tr> <tr> <td style="padding: 5px;">Stop</td> <td></td> </tr> <tr> <td style="padding: 5px;">Interior</td> <td></td> </tr> <tr> <td style="padding: 5px;">Ambient</td> <td></td> </tr> <tr> <td style="padding: 5px;">Start</td> <td></td> </tr> <tr> <td style="padding: 5px;">Stop</td> <td></td> </tr> <tr> <td style="padding: 5px;">Pressure (Inches of Hg)</td> <td></td> </tr> <tr> <td style="padding: 5px;">Ambient</td> <td></td> </tr> <tr> <td style="padding: 5px;">Start</td> <td></td> </tr> <tr> <td style="padding: 5px;">Stop</td> <td></td> </tr> </table> </div> <div style="text-align: center;"> </div> <div style="text-align: right;">                 200-22957 COC             </div> </div>																				Temperature (Fahrenheit)		Interior		Ambient		Start		Stop		Interior		Ambient		Start		Stop		Pressure (Inches of Hg)		Ambient		Start		Stop	
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Pressure (Inches of Hg)																																													
Ambient																																													
Start																																													
Stop																																													

Special Instructions/QC Requirements & Comments:

*Summa canisters placed downwind of drilling*

Sample collected by <i>Valerie Thayer</i>	Samples Received by: <i>Valerie Thayer</i>	Received by: (Signature) Received by: (Signature)
Date/Time: <i>June 24/2014 7:00</i>	Date/Time: <i>6/25/14 1030</i>	Date/Time:
Collected by:	Opened by:	Condition:
Shipper Name:	Shipped by:	Condition:

Analytical Data

Client: Katahdin Analytical Services

Job Number: 200-22957-1

Sdg Number: 200-22957

Client Sample ID: VPB147-AIR-062414

Lab Sample ID: 200-22957-1

Date Sampled: 06/24/2014 1620

Client Matrix: Air

Date Received: 06/25/2014 1030

TO-15 Volatile Organic Compounds in Ambient Air

Analysis Method:	TO-15	Analysis Batch:	200-74232	Instrument ID:	CHW.i
Prep Method:	Summa Canister	Prep Batch:	N/A	Lab File ID:	8298_011.d
Dilution:	1.0			Initial Weight/Volume:	200 mL
Analysis Date:	06/26/2014 2100			Final Weight/Volume:	200 mL
Prep Date:	06/26/2014 2100			Injection Volume:	200 mL

Analyte	Result (ppb v/v)	Qualifier	DL	LOQ
1,1,1-Trichloroethane	0.080	U	0.20	0.20
1,1,2,2-Tetrachloroethane	0.030	U	0.20	0.20
1,1,2-Trichloro-1,2,2-trifluoroethane	0.030	U	0.20	0.20
1,1,2-Trichloroethane	0.030	U	0.20	0.20
1,1-Dichloroethane	0.080	U	0.20	0.20
1,1-Dichloroethene	0.080	U	0.20	0.20
1,2,4-Trichlorobenzene	0.080	U	0.50	0.50
1,2-Dibromoethane (EDB)	0.080	U	0.20	0.20
1,2-Dichlorobenzene	0.030	U	0.20	0.20
1,2-Dichloroethane	0.030	U	0.20	0.20
1,2-Dichloropropane	0.080	U	0.20	0.20
Acetone	2.5	U	5.0	5.0
1,3-Dichlorobenzene	0.030	U	0.20	0.20
1,4-Dichlorobenzene	0.030	U	0.20	0.20
2-Butanone (MEK)	0.69		0.50	0.50
2-Hexanone	0.20	U	0.50	0.50
4-Methyl-2-pentanone	0.080	U	0.50	0.50
Benzene	0.030	U	0.20	0.20
Bromoform	0.030	U	0.20	0.20
Bromomethane	0.080	U	0.20	0.20
Carbon disulfide	0.20	U	0.50	0.50
Carbon tetrachloride	0.080	U	0.20	0.20
Chlorobenzene	0.030	U	0.20	0.20
Dibromochloromethane	0.030	U	0.20	0.20
Chloroethane	0.080	U	0.50	0.50
Chloroform	0.080	U	0.20	0.20
Chloromethane	0.55		0.50	0.50
cis-1,2-Dichloroethene	0.080	U	0.20	0.20
cis-1,3-Dichloropropene	0.080	U	0.20	0.20
Cyclohexane	0.080	UM	0.20	0.20
Bromodichloromethane	0.030	U	0.20	0.20
Dichlorodifluoromethane	0.54		0.50	0.50
Ethylbenzene	0.030	U	0.20	0.20
Isopropylbenzene	0.030	U	0.20	0.20
Methyl tert-butyl ether	0.080	U	0.20	0.20
Methylene Chloride	0.20	UM	0.50	0.50
m,p-Xylene	0.080	U	0.50	0.50
Xylene, o-	0.030	U	0.20	0.20
Styrene	0.030	U	0.20	0.20
Tetrachloroethene	0.030	U	0.20	0.20
Toluene	0.030	U	0.20	0.20
trans-1,2-Dichloroethene	0.080	U	0.20	0.20
trans-1,3-Dichloropropene	0.080	U	0.20	0.20
Trichloroethene	0.080	U	0.20	0.20
Trichlorofluoromethane	0.24		0.20	0.20
Vinyl chloride	0.080	U	0.20	0.20

*Rizoblu*

Client: Katahdin Analytical Services

Job Number: 200-22957-1

Sdg Number: 200-22957

Client Sample ID: VPB147-AIR-062414

Lab Sample ID: 200-22957-1

Date Sampled: 06/24/2014 1620

Client Matrix: Air

Date Received: 06/25/2014 1030

TO-15 Volatile Organic Compounds in Ambient Air

Analysis Method:	TO-15	Analysis Batch:	200-74232	Instrument ID:	CHW.i
Prep Method:	Summa Canister	Prep Batch:	N/A	Lab File ID:	8298_011.d
Dilution:	1.0			Initial Weight/Volume:	200 mL
Analysis Date:	06/26/2014 2100			Final Weight/Volume:	200 mL
Prep Date:	06/26/2014 2100			Injection Volume:	200 mL

Analyte	Result (ppb v/v)	Qualifier	DL	LOQ
Xylene (total)	0.080	U	0.20	0.20

Analyte	Result (ug/m3)	Qualifier	DL	LOQ
1,1,1-Trichloroethane	0.44	U	1.1	1.1
1,1,2,2-Tetrachloroethane	0.21	U	1.4	1.4
1,1,2-Trichloro-1,2,2-trifluoroethane	0.23	U	1.5	1.5
1,1,2-Trichloroethane	0.16	U	1.1	1.1
1,1-Dichloroethane	0.32	U	0.81	0.81
1,1-Dichloroethene	0.32	U	0.79	0.79
1,2,4-Trichlorobenzene	0.59	U	3.7	3.7
1,2-Dibromoethane (EDB)	0.61	U	1.5	1.5
1,2-Dichlorobenzene	0.18	U	1.2	1.2
1,2-Dichloroethane	0.12	U	0.81	0.81
1,2-Dichloropropane	0.37	U	0.92	0.92
Acetone	5.9	U	12	12
1,3-Dichlorobenzene	0.18	U	1.2	1.2
1,4-Dichlorobenzene	0.18	U	1.2	1.2
2-Butanone (MEK)	2.0	U	1.5	1.5
2-Hexanone	0.82	U	2.0	2.0
4-Methyl-2-pentanone	0.33	U	2.0	2.0
Benzene	0.096	U	0.64	0.64
Bromoform	0.31	U	2.1	2.1
Bromomethane	0.31	U	0.78	0.78
Carbon disulfide	0.62	U	1.6	1.6
Carbon tetrachloride	0.50	U	1.3	1.3
Chlorobenzene	0.14	U	0.92	0.92
Dibromochloromethane	0.26	U	1.7	1.7
Chloroethane	0.21	U	1.3	1.3
Chloroform	0.39	U	0.98	0.98
Chloromethane	1.1	U	1.0	1.0
cis-1,2-Dichloroethene	0.32	U	0.79	0.79
cis-1,3-Dichloropropene	0.36	U	0.91	0.91
Cyclohexane	0.28	U M	0.69	0.69
Bromodichloromethane	0.20	U	1.3	1.3
Dichlorodifluoromethane	2.7	U	2.5	2.5
Ethylbenzene	0.13	U	0.87	0.87
Isopropylbenzene	0.15	U	0.98	0.98
Methyl tert-butyl ether	0.29	U	0.72	0.72
Methylene Chloride	0.69	U M	1.7	1.7
m,p-Xylene	0.35	U	2.2	2.2
Xylene, o-	0.13	U	0.87	0.87
Styrene	0.13	U	0.85	0.85
Tetrachloroethene	0.20	U	1.4	1.4
Toluene	0.11	U	0.75	0.75
trans-1,2-Dichloroethene	0.32	U	0.79	0.79
trans-1,3-Dichloropropene	0.36	U	0.91	0.91

*Handwritten signature/initials in blue ink.*

**Analytical Data**

Client: Katahdin Analytical Services

Job Number: 200-22957-1

Sdg Number: 200-22957

Client Sample ID: VPB147-AIR-062414

Lab Sample ID: 200-22957-1

Date Sampled: 06/24/2014 1620

Client Matrix: Air

Date Received: 06/25/2014 1030

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**TO-15 Volatile Organic Compounds in Ambient Air**

Analysis Method:	TO-15	Analysis Batch:	200-74232	Instrument ID:	CHW.i
Prep Method:	Summa Canister	Prep Batch:	N/A	Lab File ID:	8298_011.d
Dilution:	1.0			Initial Weight/Volume:	200 mL
Analysis Date:	06/26/2014 2100			Final Weight/Volume:	200 mL
Prep Date:	06/26/2014 2100			Injection Volume:	200 mL

Analyte	Result (ug/m3)	Qualifier	DL	LOQ
Trichloroethene	0.43	U	1.1	1.1
Trichlorofluoromethane	1.3		1.1	1.1
Vinyl chloride	0.20	U	0.51	0.51
Xylene (total)	0.35	U	0.87	0.87







## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	KATAHDIN ANALYTICAL	
Service Request:	SH4830	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS)	
Validation Level:	Limited	
AECOM Project Number:	60266526.SA.DV	
Prepared by:	Dawn Brule/RESCON	Completed on: 09/26/2014
Reviewed by:	Paula DiMattei/RESCON	File Name: SH4830_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on June 27, 2014 and June 30, 2014.

Sample ID	Matrix/Sample Type
VPB147-GW-062714-878-880	Ground water
VPB147-GW-062714-888-890	Ground water
VPB147-GW-063014-898-900	Ground water
VPB147-TRIP BLANK-063014	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following parameters (where applicable to the method):

- X Data completeness (chain-of-custody (COC))/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification
- ✓ Laboratory blanks/trip blanks/equipment blanks
- X Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicates

- ✓ Internal standards
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Selected samples were mostly soil and had very little standing water.

- For sample VPB147-GW-063014-898-900, the laboratory decanted the liquid from one vial prior to analysis.
- For samples VPB147-GW-062714-888-890 and VPB147-GW-062714-878-880, the laboratory decanted the water from individual vials into one vial as a composite per sample.
- Due to limited volume, samples VPB147-GW-063014-898-900 and VPB147-GW-062714-878-880 were analyzed at dilutions and their detection limits are elevated accordingly.

Positive and non-detect results for these sample were qualified as estimated (J and UJ) respectively, due to possible loss of sample integrity during the decanting procedure.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Table A-1.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

Nonconformances are summarized in Attachment A in Table A-2.

Data qualification on the basis of surrogate recovery nonconformances was as follows:

Criteria	Action	
	Detected Compounds	Nondetected Compounds
%R > Upper Limit (UL)	J	No qualification
20% ≤ %R < Lower Limit (LL)	J	UJ
%R < 20%	J	R

Qualified sample results are shown in Table 1.

### **MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062714-878-880	WG	1,1,1-TRICHLOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,1,2,2-TETRACHLOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,1,2-TRICHLOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,1-DICHLOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,1-DICHLOROETHENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2,4-TRICHLOROBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2-DIBROMO-3-CHLOROPROPANE		3.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2-DIBROMOETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2-DICHLOROBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2-DICHLOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2-DICHLOROETHENE, TOTAL		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,2-DICHLOROPROPANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,3-DICHLOROBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	1,4-DICHLOROBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	2-BUTANONE		10	UG_L	UJ	mc,c
VPB147-GW-062714-878-880	WG	2-HEXANONE		10	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	4-METHYL-2-PENTANONE		10	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	ACETONE	24	10.	UG_L	J	mc,s
VPB147-GW-062714-878-880	WG	BENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	BROMODICHLOROMETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	BROMOFORM		2.0	UG_L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062714-878-880	WG	BROMOMETHANE		4.0	UG_L	UJ	mc,c
VPB147-GW-062714-878-880	WG	CARBON DISULFIDE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CARBON TETRACHLORIDE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CHLOROBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CHLOROETHANE		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CHLOROFORM		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CHLOROMETHANE		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CIS-1,2-DICHLOROETHENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CIS-1,3-DICHLOROPROPENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	CYCLOHEXANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	DIBROMOCHLOROMETHANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	DICHLORODIFLUOROMETHANE		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	ETHYLBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	ISOPROPYLBENZENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	M- AND P-XYLENE		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	METHYL ACETATE		3.0	UG_L	UJ	mc,c
VPB147-GW-062714-878-880	WG	METHYL CYCLOHEXANE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	METHYL TERT-BUTYL ETHER		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	METHYLENE CHLORIDE	16	10	UG_L	J	mc,s
VPB147-GW-062714-878-880	WG	O-XYLENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	STYRENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	TETRACHLOROETHENE		2.0	UG_L	UJ	mc
VPB147-GW-	WG	TOLUENE		2.0	UG_L	UJ	mc



Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
062714-878-880							
VPB147-GW-062714-878-880	WG	TRANS-1,2-DICHLOROETHENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	TRANS-1,3-DICHLOROPROPENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	TRICHLOROETHENE		2.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	TRICHLOROFLUOROMETHANE		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	VINYL CHLORIDE		4.0	UG_L	UJ	mc
VPB147-GW-062714-878-880	WG	XYLENES, TOTAL		6.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,1,1-TRICHLOROETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,1,2,2-TETRACHLOROETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,1,2-TRICHLOROETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,1-DICHLOROETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,1-DICHLOROETHENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2,4-TRICHLOROBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2-DIBROMOETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2-DICHLOROBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2-DICHLOROETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2-DICHLOROETHENE, TOTAL		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,2-DICHLOROPROPANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,3-DICHLOROBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	1,4-DICHLOROBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	2-BUTANONE	13	2.5	UG_L	J	mc,c,s

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-062714-888-890	WG	2-HEXANONE		2.5	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	4-METHYL-2-PENTANONE		2.5	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	ACETONE	85	2.5	UG_L	J	mc,s
VPB147-GW-062714-888-890	WG	BENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	BROMODICHLOROMETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	BROMOFORM		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	BROMOMETHANE		1.0	UG_L	UJ	mc,c
VPB147-GW-062714-888-890	WG	CARBON DISULFIDE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CARBON TETRACHLORIDE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CHLOROBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CHLOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CHLOROFORM		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CHLOROMETHANE		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CIS-1,2-DICHLOROETHENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CIS-1,3-DICHLOROPROPENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	CYCLOHEXANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	DIBROMOCHLOROMETHANE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	DICHLORODIFLUOROMETHANE		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	ETHYLBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	ISOPROPYLBENZENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	M- AND P-XYLENE		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	METHYL ACETATE		0.75	UG_L	UJ	mc,c
VPB147-GW-	WG	METHYL CYCLOHEXANE		0.50	UG_L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
062714-888-890							
VPB147-GW-062714-888-890	WG	METHYL TERT-BUTYL ETHER		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	METHYLENE CHLORIDE		2.5	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	O-XYLENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	STYRENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	TETRACHLOROETHENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	TOLUENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	TRANS-1,2-DICHLOROETHENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	TRANS-1,3-DICHLOROPROPENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	TRICHLOROETHENE		0.50	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	TRICHLOROFLUOROMETHANE		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	VINYL CHLORIDE		1.0	UG_L	UJ	mc
VPB147-GW-062714-888-890	WG	XYLENES, TOTAL		1.5	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,1,1-TRICHLOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,1,2,2-TETRACHLOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,1,2-TRICHLOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,1-DICHLOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,1-DICHLOROETHENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,2,4-TRICHLOROBENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,2-DIBROMO-3-CHLOROPROPANE		1.5	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,2-DIBROMOETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,2-DICHLOROBENZENE		1.0	UG_L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-063014-898-900	WG	1,2-DICHLOROETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,2-DICHLOROETHENE, TOTAL		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,2-DICHLOROPROPANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,3-DICHLOROBENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	1,4-DICHLOROBENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	2-BUTANONE		5.0	UG_L	UJ	mc,c
VPB147-GW-063014-898-900	WG	2-HEXANONE		5.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	4-METHYL-2-PENTANONE		5.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	ACETONE	56	5.0	UG_L	J	mc,s
VPB147-GW-063014-898-900	WG	BENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	BROMODICHLOROMETHANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	BROMOFORM		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	BROMOMETHANE		2.0	UG_L	UJ	mc,c
VPB147-GW-063014-898-900	WG	CARBON DISULFIDE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CARBON TETRACHLORIDE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CHLOROBENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CHLOROETHANE		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CHLOROFORM		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CHLOROMETHANE		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CIS-1,2-DICHLOROETHENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CIS-1,3-DICHLOROPROPENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	CYCLOHEXANE		1.0	UG_L	UJ	mc
VPB147-GW-	WG	DIBROMOCHLOROMETHANE		1.0	UG_L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
063014-898-900							
VPB147-GW-063014-898-900	WG	DICHLORODIFLUOROMETHANE		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	ETHYLBENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	ISOPROPYLBENZENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	M- AND P-XYLENE		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	METHYL ACETATE		1.5	UG_L	UJ	mc,c
VPB147-GW-063014-898-900	WG	METHYL CYCLOHEXANE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	METHYL TERT-BUTYL ETHER		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	METHYLENE CHLORIDE	3.8	5.0	UG_L	J	mc,s
VPB147-GW-063014-898-900	WG	O-XYLENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	STYRENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	TETRACHLOROETHENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	TOLUENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	TRANS-1,2-DICHLOROETHENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	TRANS-1,3-DICHLOROPROPENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	TRICHLOROETHENE		1.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	TRICHLOROFLUOROMETHANE		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	VINYL CHLORIDE		2.0	UG_L	UJ	mc
VPB147-GW-063014-898-900	WG	XYLENES, TOTAL		3.0	UG_L	UJ	mc
VPB147-TRIP BLANK-063014	WQ	2-BUTANONE		2.5	UG_L	UJ	c
VPB147-TRIP BLANK-063014	WQ	BROMOMETHANE		1.0	UG_L	UJ	c
VPB147-TRIP BLANK-063014	WQ	METHYL ACETATE		0.75	UG_L	UJ	c

## Attachment A

## Non Conformance Summary Tables

Table A-1 - Continuing Calibration Verification Standard

CCV ID	Compound	% R	Limits
WG145865-4	BROMOMETHANE	21	<20%
	2-BUTANONE	25	<20%
	METHYL ACETATE	22	<20%
Associated samples: all samples in SDG SH4830			

Table A-2 - Surrogates

Sample ID	Surrogate	% Recovery	Lower Limit	Upper Limit
VPB147-GW-062714-878-880	1,2-DICHLOROETHANE-D4	136	70	120
VPB147-GW-062714-878-880	DIBROMOFLUOROMETHANE	116	85	115
VPB147-GW-062714-888-890	1,2-DICHLOROETHANE-D4	131	70	120
VPB147-GW-063014-898-900	1,2-DICHLOROETHANE-D4	131	70	120
VPB147-TRIP BLANK-063014	1,2-DICHLOROETHANE-D4	138	70	120

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

### Attachment C

#### Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance





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 Scarborough, ME 04074  
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 Fax: (207) 775-4029

# CHAIN of CUSTODY

PLEASE BEAR DOWN AND  
 PRINT LEGIBLY IN PEN

Page \_\_\_\_ of \_\_\_\_

Client: Resolution Consultants Contact: Eleanor Vivandell Phone # \_\_\_\_\_ Fax # \_\_\_\_\_  
 Address \_\_\_\_\_ City \_\_\_\_\_ State \_\_\_\_\_ Zip Code \_\_\_\_\_

Purchase Order # \_\_\_\_\_ Proj. Name / No. Bethpage 60266526 Katahdin Quote # \_\_\_\_\_  
 Bill (if different than above) Address \_\_\_\_\_

Sampler (Print / Sign) Valerie Thayer Valerie Thayer Copies To: Val Thayer

LAB USE ONLY WORK ORDER #: SH 4830  
 KATAHDIN PROJECT NUMBER \_\_\_\_\_

ANALYSIS AND CONTAINER TYPE PRESERVATIVES									
Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.
Y	N	Y	N	Y	N	Y	N	Y	N

REMARKS: \_\_\_\_\_  
 SHIPPING INFO:  FED EX  UPS  CLIENT  
 AIRBILL NO: \_\_\_\_\_  
 TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

* Sample Description	Date / Time coll'd	Matrix	No. of Cntrs.											
VFB 147-GW-063014-048-900	6/30/14 12:15	GW	1	X										
IDWGW-063014-White frac TANK	6/30/14 14:30	W	1		X									
VFB 147-GW-062714-888-870	6/27/14 13:45	GW	4	X										
Temp Blank	/ /	W	1											
VFB 147-GW-062714-878-880	6/27/14 10:30	GW	6	X										
VFB 147-Tripblanks-063014	5/2/14 10:00	W	3	X										

COMMENTS: IDWGW - sample from frac tank - white

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>6/30/14 7</u>	Received By: (Signature) <u>[Signature]</u>	Relinquished By: (Signature) _____	Date / Time _____	Received By: (Signature) _____
Relinquished By: (Signature) _____	Date / Time _____	Received By: (Signature) _____	Relinquished By: (Signature) _____	Date / Time _____	Received By: (Signature) _____

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4830-4DL  
**Client ID:** 147-062714-878-880  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4830  
**Lab File ID:** D9288.D

**Sample Date:** 27-JUN-14  
**Received Date:** 01-JUL-14  
**Extract Date:** 02-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145865

**Analysis Date:** 02-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	4.0	ug/L	4	2	8.0	0.96	4.0
Chloromethane	U	4.0	ug/L	4	2	8.0	1.4	4.0
Vinyl Chloride	U	4.0	ug/L	4	2	8.0	1.0	4.0
Bromomethane	U	4.0	ug/L	4	2	8.0	2.0	4.0
Chloroethane	U	4.0	ug/L	4	2	8.0	2.2	4.0
Trichlorofluoromethane	U	4.0	ug/L	4	2	8.0	0.96	4.0
1,1-Dichloroethene	U	2.0	ug/L	4	1	4.0	1.4	2.0
Carbon Disulfide	U	2.0	ug/L	4	1	4.0	1.0	2.0
Freon-113	U	2.0	ug/L	4	1	4.0	1.2	2.0
<b>Methylene Chloride</b>	J	16	ug/L	4	5	20.	4.5	10.
<b>Acetone</b>	J	24	ug/L	4	5	20.	8.8	10.
trans-1,2-Dichloroethene	U	2.0	ug/L	4	1	4.0	1.0	2.0
Methyl tert-butyl Ether	U	2.0	ug/L	4	1	4.0	1.4	2.0
1,1-Dichloroethane	U	2.0	ug/L	4	1	4.0	0.84	2.0
cis-1,2-Dichloroethene	U	2.0	ug/L	4	1	4.0	0.84	2.0
Chloroform	U	2.0	ug/L	4	1	4.0	1.3	2.0
1,1,1-Trichloroethane	U	2.0	ug/L	4	1	4.0	0.80	2.0
2-Butanone	U	10	ug/L	4	5	20.	5.2	10.
Cyclohexane	U	2.0	ug/L	4	1	4.0	1.2	2.0
Carbon Tetrachloride	U	2.0	ug/L	4	1	4.0	0.88	2.0
Benzene	U	2.0	ug/L	4	1	4.0	1.0	2.0
1,2-Dichloroethane	U	2.0	ug/L	4	1	4.0	0.80	2.0
Trichloroethene	U	2.0	ug/L	4	1	4.0	1.1	2.0
1,2-Dichloropropane	U	2.0	ug/L	4	1	4.0	1.0	2.0
Bromodichloromethane	U	2.0	ug/L	4	1	4.0	1.3	2.0
cis-1,3-Dichloropropene	U	2.0	ug/L	4	1	4.0	0.76	2.0
Toluene	U	2.0	ug/L	4	1	4.0	1.1	2.0
4-Methyl-2-Pentanone	U	10	ug/L	4	5	20.	5.3	10.
trans-1,3-Dichloropropene	U	2.0	ug/L	4	1	4.0	0.80	2.0
1,1,2-Trichloroethane	U	2.0	ug/L	4	1	4.0	1.3	2.0
Tetrachloroethene	U	2.0	ug/L	4	1	4.0	1.6	2.0
Dibromochloromethane	U	2.0	ug/L	4	1	4.0	1.2	2.0
2-Hexanone	U	10	ug/L	4	5	20.	6.8	10.
Chlorobenzene	U	2.0	ug/L	4	1	4.0	0.88	2.0
Ethylbenzene	U	2.0	ug/L	4	1	4.0	0.84	2.0

*Gizella*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4830-4DL  
**Client ID:** 147-062714-878-880  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4830  
**Lab File ID:** D9288.D

**Sample Date:** 27-JUN-14  
**Received Date:** 01-JUL-14  
**Extract Date:** 02-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145865

**Analysis Date:** 02-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U <b>UJ</b>	6.0	ug/L	4	3	12.	1.0	6.0
Styrene	U	2.0	ug/L	4	1	4.0	0.92	2.0
Bromoform	U	2.0	ug/L	4	1	4.0	0.92	2.0
Isopropylbenzene	U	2.0	ug/L	4	1	4.0	0.92	2.0
1,1,2,2-Tetrachloroethane	U	2.0	ug/L	4	1	4.0	1.5	2.0
1,3-Dichlorobenzene	U	2.0	ug/L	4	1	4.0	1.0	2.0
1,4-Dichlorobenzene	U	2.0	ug/L	4	1	4.0	0.96	2.0
1,2-Dichlorobenzene	U	2.0	ug/L	4	1	4.0	0.60	2.0
1,2,4-Trichlorobenzene	U	2.0	ug/L	4	1	4.0	1.5	2.0
Methyl Acetate	U	3.0	ug/L	4	1	4.0	2.1	3.0
Methylcyclohexane	U	2.0	ug/L	4	1	4.0	1.2	2.0
o-Xylene	U	2.0	ug/L	4	1	4.0	1.0	2.0
M+P-Xylenes	U	4.0	ug/L	4	2	8.0	2.4	4.0
1,2-Dichloroethylene (Total)	U	4.0	ug/L	4	2	8.0	0.84	4.0
1,2-Dibromoethane	U	2.0	ug/L	4	1	4.0	0.88	2.0
1,2-Dibromo-3-Chloropropane	U	3.0	ug/L	4	1	4.0	2.0	3.0
P-Bromofluorobenzene		84.6	%					
Toluene-d8		114.	%					
1,2-Dichloroethane-d4	*	136.	%					
Dibromofluoromethane	*	116.	%					

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4830-3  
**Client ID:** 147-062714-888-890  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4830  
**Lab File ID:** D9285.D

**Sample Date:** 27-JUN-14  
**Received Date:** 01-JUL-14  
**Extract Date:** 02-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145865

**Analysis Date:** 02-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	J	85	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U UJ	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>2-Butanone</b>	J	13	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U UJ	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Rizki/17*

## Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4830-3  
 Client ID: 147-062714-888-890  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4830  
 Lab File ID: D9285.D

Sample Date: 27-JUN-14  
 Received Date: 01-JUL-14  
 Extract Date: 02-JUL-14  
 Extracted By: DJP  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145865

Analysis Date: 02-JUL-14  
 Analyst: DJP  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U <i>UJ</i>	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		78.0	%					
Toluene-d8	*	85.0	%					
1,2-Dichloroethane-d4	*	131.	%					
Dibromofluoromethane		101.	%					

*DJP 7/18/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4830-1DL  
**Client ID:** 147-063014-898-900  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4830  
**Lab File ID:** D9286.D

**Sample Date:** 30-JUN-14  
**Received Date:** 01-JUL-14  
**Extract Date:** 02-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145865

**Analysis Date:** 02-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	2.0	ug/L	2	2	4.0	0.48	2.0
Chloromethane	U	2.0	ug/L	2	2	4.0	0.72	2.0
Vinyl Chloride	U	2.0	ug/L	2	2	4.0	0.50	2.0
Bromomethane	U	2.0	ug/L	2	2	4.0	0.98	2.0
Chloroethane	U	2.0	ug/L	2	2	4.0	1.1	2.0
Trichlorofluoromethane	U	2.0	ug/L	2	2	4.0	0.48	2.0
1,1-Dichloroethene	U	1.0	ug/L	2	1	2.0	0.70	1.0
Carbon Disulfide	U	1.0	ug/L	2	1	2.0	0.50	1.0
Freon-113	U	1.0	ug/L	2	1	2.0	0.62	1.0
<b>Methylene Chloride</b>	J	3.8	ug/L	2	5	10.	2.3	5.0
<b>Acetone</b>	J	56	ug/L	2	5	10.	4.4	5.0
trans-1,2-Dichloroethene	U	1.0	ug/L	2	1	2.0	0.50	1.0
Methyl tert-butyl Ether	U	1.0	ug/L	2	1	2.0	0.72	1.0
1,1-Dichloroethane	U	1.0	ug/L	2	1	2.0	0.42	1.0
cis-1,2-Dichloroethene	U	1.0	ug/L	2	1	2.0	0.42	1.0
Chloroform	U	1.0	ug/L	2	1	2.0	0.64	1.0
1,1,1-Trichloroethane	U	1.0	ug/L	2	1	2.0	0.40	1.0
2-Butanone	U	5.0	ug/L	2	5	10.	2.6	5.0
Cyclohexane	U	1.0	ug/L	2	1	2.0	0.62	1.0
Carbon Tetrachloride	U	1.0	ug/L	2	1	2.0	0.44	1.0
Benzene	U	1.0	ug/L	2	1	2.0	0.52	1.0
1,2-Dichloroethane	U	1.0	ug/L	2	1	2.0	0.40	1.0
Trichloroethene	U	1.0	ug/L	2	1	2.0	0.56	1.0
1,2-Dichloropropane	U	1.0	ug/L	2	1	2.0	0.50	1.0
Bromodichloromethane	U	1.0	ug/L	2	1	2.0	0.66	1.0
cis-1,3-Dichloropropene	U	1.0	ug/L	2	1	2.0	0.38	1.0
Toluene	U	1.0	ug/L	2	1	2.0	0.54	1.0
4-Methyl-2-Pentanone	U	5.0	ug/L	2	5	10.	2.6	5.0
trans-1,3-Dichloropropene	U	1.0	ug/L	2	1	2.0	0.40	1.0
1,1,2-Trichloroethane	U	1.0	ug/L	2	1	2.0	0.66	1.0
Tetrachloroethene	U	1.0	ug/L	2	1	2.0	0.80	1.0
Dibromochloromethane	U	1.0	ug/L	2	1	2.0	0.60	1.0
2-Hexanone	U	5.0	ug/L	2	5	10.	3.4	5.0
Chlorobenzene	U	1.0	ug/L	2	1	2.0	0.44	1.0
Ethylbenzene	U	1.0	ug/L	2	1	2.0	0.42	1.0

*Riz/2/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4830-1DL  
 Client ID: 147-063014-898-900  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4830  
 Lab File ID: D9286.D

Sample Date: 30-JUN-14  
 Received Date: 01-JUL-14  
 Extract Date: 02-JUL-14  
 Extracted By: DJP  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145865

Analysis Date: 02-JUL-14  
 Analyst: DJP  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U UJ	3.0	ug/L	2	3	6.0	0.50	3.0
Styrene	U	1.0	ug/L	2	1	2.0	0.46	1.0
Bromoform	U	1.0	ug/L	2	1	2.0	0.46	1.0
Isopropylbenzene	U	1.0	ug/L	2	1	2.0	0.46	1.0
1,1,2,2-Tetrachloroethane	U	1.0	ug/L	2	1	2.0	0.76	1.0
1,3-Dichlorobenzene	U	1.0	ug/L	2	1	2.0	0.52	1.0
1,4-Dichlorobenzene	U	1.0	ug/L	2	1	2.0	0.48	1.0
1,2-Dichlorobenzene	U	1.0	ug/L	2	1	2.0	0.30	1.0
1,2,4-Trichlorobenzene	U	1.0	ug/L	2	1	2.0	0.74	1.0
Methyl Acetate	U	1.5	ug/L	2	1	2.0	1.1	1.5
Methylcyclohexane	U	1.0	ug/L	2	1	2.0	0.60	1.0
o-Xylene	U	1.0	ug/L	2	1	2.0	0.50	1.0
M+P-Xylenes	U	2.0	ug/L	2	2	4.0	1.2	2.0
1,2-Dichloroethylene (Total)	U	2.0	ug/L	2	2	4.0	0.42	2.0
1,2-Dibromoethane	U	1.0	ug/L	2	1	2.0	0.44	1.0
1,2-Dibromo-3-Chloropropane	U	1.5	ug/L	2	1	2.0	1.0	1.5
P-Bromofluorobenzene		85.9	%					
Toluene-d8		115.	%					
1,2-Dichloroethane-d4	*	131.	%					
Dibromofluoromethane		114.	%					

*Handwritten signature/initials*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4830-5  
**Client ID:** VPB147-TB-063014  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4830  
**Lab File ID:** D9283.D

**Sample Date:** 30-JUN-14  
**Received Date:** 01-JUL-14  
**Extract Date:** 02-JUL-14  
**Extracted By:** DJP  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG145865

**Analysis Date:** 02-JUL-14  
**Analyst:** DJP  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	<del>U</del> UJ	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

Riz 1/8/14



### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4830-5  
 Client ID: VPB147-TB-063014  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4830  
 Lab File ID: D9283.D

Sample Date: 30-JUN-14  
 Received Date: 01-JUL-14  
 Extract Date: 02-JUL-14  
 Extracted By: DJP  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG145865

Analysis Date: 02-JUL-14  
 Analyst: DJP  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 03-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	<del>U</del> <i>U5</i>	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		86.0	%					
Toluene-d8		117.	%					
1,2-Dichloroethane-d4	*	138.	%					
Dibromofluoromethane		114.	%					

*Riz/10/14*



## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	Katahdin Analytical	
Service Request:	SH4962	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS)	
Validation Level:	3	
AECOM Project Number:	60266526.SA.DV	
Prepared by:	Dawn Brule/RESCON	Completed on: 12/8/2014
Reviewed by:	Lori Herberich/RESCON	File Name: SH4962_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants from the Regional Groundwater Investigation - NWIRP Bethpage site on July 1 and 2, 2014.

Sample ID	Matrix/Sample Type
VPB147-GW-070114-938-940	Groundwater
VPB147-GW-070114-943-945	Groundwater
VPB147-GW-070114-958-960	Groundwater
VPB147-TRIPBLANKS-070214	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW-46, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- X Data completeness (chain-of-custody [COC])/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification
- ✓ Laboratory blanks/equipment blanks/trip blanks
- ✓ Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicate results

- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### **Data Completeness (COC)/Sample Integrity**

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Selected samples were mostly soil and had very little standing water.

- For samples VPB147-GW-070114-958-960, VPB147-GW-070114-943-945, and VPB147-GW-070114-938-940, the laboratory decanted the water from individual vials into one vial as one composite per sample.
- Due to limited volume, samples VPB147-GW-070114-943-945 and VPB147-GW-070114-938-940 were analyzed at dilutions and their detection limits are elevated accordingly.

Positive and nondetect results for these sample were qualified as estimated (J and UJ, respectively), due to possible loss of sample integrity during the decanting procedure.

Due to limitations in the reporting system, the laboratory omitted the "VPB-" prefix and the "GW" from the sample ID, and truncated the ID for the Trip Blank in the report. The submitted EDD file reflects the full sample ID. Qualified sample results are shown in Table 1.

### **Holding Times and Sample Preservation**

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria. The QC acceptance criteria were met.

### **GC/MS Performance Checks**

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met. The QC acceptance criteria were met.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and/or
- the retention time method acceptance criteria were met.

Nonconformances are summarized in Attachment A in Tables A-1 and A-2.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift >20%	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1.

### **Laboratory Blanks/Equipment Blanks/Trip Blanks**

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL). An equipment blank was not submitted with the samples in this data set.

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required. The QC acceptance criteria were met and/or qualification of the sample results was not required.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **MS/MSD Results**

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as not detected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

## **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

**ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-938-940	WG	1,1,1-TRICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,1,2,2-TETRACHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,1,2-TRICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,1-DICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,1-DICHLOROETHENE		5.0	UG/L	UJ	mc,c
VPB147-GW-070114-938-940	WG	1,2,4-TRICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,2-DIBROMO-3-CHLOROPROPANE		7.5	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,2-DIBROMOETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,2-DICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,2-DICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,2-DICHLOROETHENE, TOTAL		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,2-DICHLOROPROPANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,3-DICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	1,4-DICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	2-BUTANONE		25	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	2-HEXANONE		25	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	4-METHYL-2-PENTANONE		25	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	ACETONE		25	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	BENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	BROMODICHLOROMETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	BROMOFORM		5.0	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-938-940	WG	BROMOMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CARBON DISULFIDE		5.0	UG/L	UJ	mc,c
VPB147-GW-070114-938-940	WG	CARBON TETRACHLORIDE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CHLOROETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CHLOROFORM		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CHLOROMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CIS-1,2-DICHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CIS-1,3-DICHLOROPROPENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	CYCLOHEXANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	DIBROMOCHLOROMETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	DICHLORODIFLUOROMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	ETHYLBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	ISOPROPYLBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	M- AND P-XYLENE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	METHYL ACETATE		7.5	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	METHYL CYCLOHEXANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	METHYL TERT-BUTYL ETHER		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	METHYLENE CHLORIDE		25	UG/L	UJ	mc,c
VPB147-GW-070114-938-940	WG	O-XYLENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	STYRENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	TETRACHLOROETHENE		5.0	UG/L	UJ	mc



Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-938-940	WG	TOLUENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	TRANS-1,2-DICHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	TRANS-1,3-DICHLOROPROPENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	TRICHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	TRICHLOROFLUOROMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	VINYL CHLORIDE		10	UG/L	UJ	mc
VPB147-GW-070114-938-940	WG	XYLENES, TOTAL		15	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,1,1-TRICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,1,2,2-TETRACHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,1,2-TRICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,1-DICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,1-DICHLOROETHENE		5.0	UG/L	UJ	mc,c
VPB147-GW-070114-943-945	WG	1,2,4-TRICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,2-DIBROMO-3-CHLOROPROPANE		7.5	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,2-DIBROMOETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,2-DICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,2-DICHLOROETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,2-DICHLOROETHENE, TOTAL		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,2-DICHLOROPROPANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,3-DICHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	1,4-DICHLOROBENZENE		5.0	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-943-945	WG	2-BUTANONE		25	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	2-HEXANONE		25	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	4-METHYL-2-PENTANONE		25	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	ACETONE	38	25	UG/L	J	mc,c
VPB147-GW-070114-943-945	WG	BENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	BROMODICHLOROMETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	BROMOFORM		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	BROMOMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CARBON DISULFIDE		5.0	UG/L	UJ	mc,c
VPB147-GW-070114-943-945	WG	CARBON TETRACHLORIDE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CHLOROBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CHLOROETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CHLOROFORM		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CHLOROMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CIS-1,2-DICHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CIS-1,3-DICHLOROPROPENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	CYCLOHEXANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	DIBROMOCHLOROMETHANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	DICHLORODIFLUOROMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	ETHYLBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	ISOPROPYLBENZENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	M- AND P-XYLENE		10	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-943-945	WG	METHYL ACETATE		7.5	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	METHYL CYCLOHEXANE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	METHYL TERT-BUTYL ETHER		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	METHYLENE CHLORIDE		25	UG/L	UJ	mc,c
VPB147-GW-070114-943-945	WG	O-XYLENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	STYRENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	TETRACHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	TOLUENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	TRANS-1,2-DICHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	TRANS-1,3-DICHLOROPROPENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	TRICHLOROETHENE		5.0	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	TRICHLOROFLUOROMETHANE		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	VINYL CHLORIDE		10	UG/L	UJ	mc
VPB147-GW-070114-943-945	WG	XYLENES, TOTAL		15	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,1,1-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,1,2,2-TETRACHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,1,2-TRICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,1-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,1-DICHLOROETHENE		0.50	UG/L	UJ	mc,c
VPB147-GW-070114-958-960	WG	1,2,4-TRICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,2-DIBROMO-3-CHLOROPROPANE		0.75	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-958-960	WG	1,2-DIBROMOETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,2-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,2-DICHLOROETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,2-DICHLOROETHENE, TOTAL		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,2-DICHLOROPROPANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,3-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	1,4-DICHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	2-BUTANONE	5.2	2.5	UG/L	J	mc,c
VPB147-GW-070114-958-960	WG	2-HEXANONE		2.5	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	ACETONE	24	2.5	UG/L	J	mc,c
VPB147-GW-070114-958-960	WG	BENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	BROMODICHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	BROMOFORM		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	BROMOMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CARBON DISULFIDE		0.50	UG/L	UJ	mc,c
VPB147-GW-070114-958-960	WG	CARBON TETRACHLORIDE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CHLOROBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CHLOROETHANE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CHLOROFORM		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CHLOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CIS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-GW-070114-958-960	WG	CIS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	DIBROMOCHLOROMETHANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	ETHYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	ISOPROPYLBENZENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	M- AND P-XYLENE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	METHYL ACETATE		0.75	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	METHYL CYCLOHEXANE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	METHYL TERT-BUTYL ETHER		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	METHYLENE CHLORIDE		2.5	UG/L	UJ	mc,c
VPB147-GW-070114-958-960	WG	O-XYLENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	STYRENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	TETRACHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	TOLUENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	TRANS-1,2-DICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	TRANS-1,3-DICHLOROPROPENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	TRICHLOROETHENE		0.50	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	TRICHLOROFLUOROMETHANE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	VINYL CHLORIDE		1.0	UG/L	UJ	mc
VPB147-GW-070114-958-960	WG	XYLENES, TOTAL		1.5	UG/L	UJ	mc
VPB147-TRIPBLANKS-070214	WQ	1,1-DICHLOROETHENE		0.50	UG/L	UJ	c

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
VPB147-TRIPBLANKS-070214	WQ	CARBON DISULFIDE		0.50	UG/L	UJ	c
VPB147-TRIPBLANKS-070214	WQ	METHYLENE CHLORIDE		2.5	UG/L	UJ	c

## Attachment A

## Nonconformance Summary Tables

Table A-1 - Initial Calibration Verification Standard

ICV ID	Compound	% R	Limits
WG146050-7	CARBON DISULFIDE	141	80-120%
	ACETONE	163	80-120%
	2-BUTANONE	122	80-120%
Associated samples: all samples in SDG SH4962			

Table A-2 -Continuing Calibration Verification Standard

CCV ID	Compound	% D	Limits
WG146127-4	1,1-DICHLOROETHENE	22	≤20%
	CARBON DISULFIDE	27	≤20%
	1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	39	≤20%
	METHYLENE CHLORIDE	24	≤20%
	METHYLCYCLOHEXANE	27	≤20%
Associated samples: all samples in SDG SH4962			

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.



## Attachment C

## Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance



600 Technology Way  
 Scarborough, ME 04074  
 Tel: (207) 874-2400  
 Fax: (207) 775-4029

**CHAIN of CUSTODY**

**PLEASE BEAR DOWN AND  
 PRINT LEGIBLY IN PEN**

Client Resolution Consultants Contact Eleanor Vivaudou Phone # ( ) Fax # ( )  
 Address 100 Red Schoolhouse Rd City Chestnut Ridge State NY Zip Code \_\_\_\_\_  
 Purchase Order # \_\_\_\_\_ Proj. Name / No. Bothpac 160266526 Katahdin Quote # \_\_\_\_\_  
 Bill (if different than above) Address \_\_\_\_\_

Sampler (Print / Sign) Valerie Thayer VAL THAYER Copies To: Valerie Thayer

LAB USE ONLY WORK ORDER #: 5H4962  
 KATAHDIN PROJECT NUMBER \_\_\_\_\_

ANALYSIS AND CONTAINER TYPE PRESERVATIVES

Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.	Filt.
<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="checkbox"/> Y <input type="checkbox"/> N

REMARKS: \_\_\_\_\_

SHIPPING INFO:  FED EX  UPS  CLIENT

AIRBILL NO: \_\_\_\_\_

TEMP °C \_\_\_\_\_  TEMP BLANK  INTACT  NOT INTACT

* Sample Description	Date / Time coll'd	Matrix	No. of Cntrs.											
<u>temp blank</u>	<u>1</u>	<u>W</u>	<u>1</u>											
<u>VPB147-tripblanks-07014</u>	<u>7/21/14 10:00</u>	<u>W</u>	<u>3</u>	<u>X</u>										
<u>VPB147-GW-070114-958-960</u>	<u>7/1/14 16:45</u>	<u>GW</u>	<u>2</u>	<u>X</u>										
<u>VPB147-GW-070114-943-945</u>	<u>7/1/14 14:00</u>	<u>GW</u>	<u>2</u>	<u>X</u>										
<u>VPB147-GW-070114-938-940</u>	<u>7/1/14 11:30</u>	<u>GW</u>	<u>5</u>	<u>X</u>										

COMMENTS \_\_\_\_\_

Relinquished By: (Signature) <u>Valerie Thayer</u>	Date / Time <u>7/4/14 7-</u>	Received By: (Signature) <u>Ed M...</u>	Relinquished By: (Signature)	Date / Time	Received By: (Signature)
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Relinquished By: (Signature)	Date / Time	Received By: (Signature)

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4962-4DL  
**Client ID:** 147-070114-938-940  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4962  
**Lab File ID:** C8040.D

**Sample Date:** 01-JUL-14  
**Received Date:** 05-JUL-14  
**Extract Date:** 08-JUL-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG146127

**Analysis Date:** 08-JUL-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260C  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UJ	10	ug/L	10	2	20.	2.4	10.
Chloromethane	U	10	ug/L	10	2	20.	3.6	10.
Vinyl Chloride	U	10	ug/L	10	2	20.	2.5	10.
Bromomethane	U	10	ug/L	10	2	20.	4.9	10.
Chloroethane	U	10	ug/L	10	2	20.	5.5	10.
Trichlorofluoromethane	U	10	ug/L	10	2	20.	2.4	10.
1,1-Dichloroethene	U	5.0	ug/L	10	1	10.	3.5	5.0
Carbon Disulfide	U	5.0	ug/L	10	1	10.	2.5	5.0
Freon-113	U	5.0	ug/L	10	1	10.	3.1	5.0
Methylene Chloride	U	25	ug/L	10	5	50.	11.	25.
Acetone	U	25	ug/L	10	5	50.	22.	25.
trans-1,2-Dichloroethene	U	5.0	ug/L	10	1	10.	2.5	5.0
Methyl tert-butyl Ether	U	5.0	ug/L	10	1	10.	3.6	5.0
1,1-Dichloroethane	U	5.0	ug/L	10	1	10.	2.1	5.0
cis-1,2-Dichloroethene	U	5.0	ug/L	10	1	10.	2.1	5.0
Chloroform	U	5.0	ug/L	10	1	10.	3.2	5.0
1,1,1-Trichloroethane	U	5.0	ug/L	10	1	10.	2.0	5.0
2-Butanone	U	25	ug/L	10	5	50.	13.	25.
Cyclohexane	U	5.0	ug/L	10	1	10.	3.1	5.0
Carbon Tetrachloride	U	5.0	ug/L	10	1	10.	2.2	5.0
Benzene	U	5.0	ug/L	10	1	10.	2.6	5.0
1,2-Dichloroethane	U	5.0	ug/L	10	1	10.	2.0	5.0
Trichloroethene	U	5.0	ug/L	10	1	10.	2.8	5.0
1,2-Dichloropropane	U	5.0	ug/L	10	1	10.	2.5	5.0
Bromodichloromethane	U	5.0	ug/L	10	1	10.	3.3	5.0
cis-1,3-Dichloropropene	U	5.0	ug/L	10	1	10.	1.9	5.0
Toluene	U	5.0	ug/L	10	1	10.	2.7	5.0
4-Methyl-2-Pentanone	U	25	ug/L	10	5	50.	13.	25.
trans-1,3-Dichloropropene	U	5.0	ug/L	10	1	10.	2.0	5.0
1,1,2-Trichloroethane	U	5.0	ug/L	10	1	10.	3.3	5.0
Tetrachloroethene	U	5.0	ug/L	10	1	10.	4.0	5.0
Dibromochloromethane	U	5.0	ug/L	10	1	10.	3.0	5.0
2-Hexanone	U	25	ug/L	10	5	50.	17.	25.
Chlorobenzene	U	5.0	ug/L	10	1	10.	2.2	5.0
Ethylbenzene	U	5.0	ug/L	10	1	10.	2.1	5.0

*REC/12/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4962-4DL  
 Client ID: 147-070114-938-940  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4962  
 Lab File ID: C8040.D

Sample Date: 01-JUL-14  
 Received Date: 05-JUL-14  
 Extract Date: 08-JUL-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG146127

Analysis Date: 08-JUL-14  
 Analyst: REC  
 Analysis Method: SW846 8260C  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U <i>U5</i>	15	ug/L	10	3	30.	2.5	15.
Styrene	U	5.0	ug/L	10	1	10.	2.3	5.0
Bromoform	U	5.0	ug/L	10	1	10.	2.3	5.0
Isopropylbenzene	U	5.0	ug/L	10	1	10.	2.3	5.0
1,1,2,2-Tetrachloroethane	U	5.0	ug/L	10	1	10.	3.8	5.0
1,3-Dichlorobenzene	U	5.0	ug/L	10	1	10.	2.6	5.0
1,4-Dichlorobenzene	U	5.0	ug/L	10	1	10.	2.4	5.0
1,2-Dichlorobenzene	U	5.0	ug/L	10	1	10.	1.5	5.0
1,2,4-Trichlorobenzene	U	5.0	ug/L	10	1	10.	3.7	5.0
Methyl Acetate	U	7.5	ug/L	10	1	10.	5.3	7.5
Methylcyclohexane	U	5.0	ug/L	10	1	10.	3.0	5.0
o-Xylene	U	5.0	ug/L	10	1	10.	2.5	5.0
M+P-Xylenes	U	10	ug/L	10	2	20.	5.9	10.
1,2-Dichloroethylene (Total)	U	10	ug/L	10	2	20.	2.1	10.
1,2-Dibromoethane	U	5.0	ug/L	10	1	10.	2.2	5.0
1,2-Dibromo-3-Chloropropane	U	7.5	ug/L	10	1	10.	5.0	7.5
P-Bromofluorobenzene		94.1	%					
Toluene-d8		103.	%					
1,2-Dichloroethane-d4		104.	%					
Dibromofluoromethane		98.2	%					

*Rizhik*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4962-3DL  
**Client ID:** 147-070114-943-945  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4962  
**Lab File ID:** C8039.D

**Sample Date:** 01-JUL-14  
**Received Date:** 05-JUL-14  
**Extract Date:** 08-JUL-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG146127

**Analysis Date:** 08-JUL-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260C  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	10	ug/L	10	2	20.	2.4	10.
Chloromethane	U	10	ug/L	10	2	20.	3.6	10.
Vinyl Chloride	U	10	ug/L	10	2	20.	2.5	10.
Bromomethane	U	10	ug/L	10	2	20.	4.9	10.
Chloroethane	U	10	ug/L	10	2	20.	5.5	10.
Trichlorofluoromethane	U	10	ug/L	10	2	20.	2.4	10.
1,1-Dichloroethene	U	5.0	ug/L	10	1	10.	3.5	5.0
Carbon Disulfide	U	5.0	ug/L	10	1	10.	2.5	5.0
Freon-113	U	5.0	ug/L	10	1	10.	3.1	5.0
Methylene Chloride	U	25	ug/L	10	5	50.	11.	25.
<b>Acetone</b>	J	38	ug/L	10	5	50.	22.	25.
trans-1,2-Dichloroethene	U	5.0	ug/L	10	1	10.	2.5	5.0
Methyl tert-butyl Ether	U	5.0	ug/L	10	1	10.	3.6	5.0
1,1-Dichloroethane	U	5.0	ug/L	10	1	10.	2.1	5.0
cis-1,2-Dichloroethene	U	5.0	ug/L	10	1	10.	2.1	5.0
Chloroform	U	5.0	ug/L	10	1	10.	3.2	5.0
1,1,1-Trichloroethane	U	5.0	ug/L	10	1	10.	2.0	5.0
2-Butanone	U	25	ug/L	10	5	50.	13.	25.
Cyclohexane	U	5.0	ug/L	10	1	10.	3.1	5.0
Carbon Tetrachloride	U	5.0	ug/L	10	1	10.	2.2	5.0
Benzene	U	5.0	ug/L	10	1	10.	2.6	5.0
1,2-Dichloroethane	U	5.0	ug/L	10	1	10.	2.0	5.0
Trichloroethene	U	5.0	ug/L	10	1	10.	2.8	5.0
1,2-Dichloropropane	U	5.0	ug/L	10	1	10.	2.5	5.0
Bromodichloromethane	U	5.0	ug/L	10	1	10.	3.3	5.0
cis-1,3-Dichloropropene	U	5.0	ug/L	10	1	10.	1.9	5.0
Toluene	U	5.0	ug/L	10	1	10.	2.7	5.0
4-Methyl-2-Pentanone	U	25	ug/L	10	5	50.	13.	25.
trans-1,3-Dichloropropene	U	5.0	ug/L	10	1	10.	2.0	5.0
1,1,2-Trichloroethane	U	5.0	ug/L	10	1	10.	3.3	5.0
Tetrachloroethene	U	5.0	ug/L	10	1	10.	4.0	5.0
Dibromochloromethane	U	5.0	ug/L	10	1	10.	3.0	5.0
2-Hexanone	U	25	ug/L	10	5	50.	17.	25.
Chlorobenzene	U	5.0	ug/L	10	1	10.	2.2	5.0
Ethylbenzene	U	5.0	ug/L	10	1	10.	2.1	5.0

*Rizholu*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4962-3DL  
**Client ID:** 147-070114-943-945  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4962  
**Lab File ID:** C8039.D

**Sample Date:** 01-JUL-14  
**Received Date:** 05-JUL-14  
**Extract Date:** 08-JUL-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG146127

**Analysis Date:** 08-JUL-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260C  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U WJ	15	ug/L	10	3	30.	2.5	15.
Styrene	U	5.0	ug/L	10	1	10.	2.3	5.0
Bromoform	U	5.0	ug/L	10	1	10.	2.3	5.0
Isopropylbenzene	U	5.0	ug/L	10	1	10.	2.3	5.0
1,1,2,2-Tetrachloroethane	U	5.0	ug/L	10	1	10.	3.8	5.0
1,3-Dichlorobenzene	U	5.0	ug/L	10	1	10.	2.6	5.0
1,4-Dichlorobenzene	U	5.0	ug/L	10	1	10.	2.4	5.0
1,2-Dichlorobenzene	U	5.0	ug/L	10	1	10.	1.5	5.0
1,2,4-Trichlorobenzene	U	5.0	ug/L	10	1	10.	3.7	5.0
Methyl Acetate	U	7.5	ug/L	10	1	10.	5.3	7.5
Methylcyclohexane	U	5.0	ug/L	10	1	10.	3.0	5.0
o-Xylene	U	5.0	ug/L	10	1	10.	2.5	5.0
M+P-Xylenes	U	10	ug/L	10	2	20.	5.9	10.
1,2-Dichloroethylene (Total)	U	10	ug/L	10	2	20.	2.1	10.
1,2-Dibromoethane	U	5.0	ug/L	10	1	10.	2.2	5.0
1,2-Dibromo-3-Chloropropane	U	7.5	ug/L	10	1	10.	5.0	7.5
P-Bromofluorobenzene		93.2	%					
Toluene-d8		101.	%					
1,2-Dichloroethane-d4		102.	%					
Dibromofluoromethane		96.8	%					

*R 12/14/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4962-2  
**Client ID:** 147-070114-958-960  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4962  
**Lab File ID:** C8038.D

**Sample Date:** 01-JUL-14  
**Received Date:** 05-JUL-14  
**Extract Date:** 08-JUL-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG146127

**Analysis Date:** 08-JUL-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260C  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U JJ	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
<b>Acetone</b>	U JJ	24	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U JJ	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>2-Butanone</b>	U JJ	5.2	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U JJ	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Rizkelly*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4962-2  
 Client ID: 147-070114-958-960  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4962  
 Lab File ID: C8038.D

Sample Date: 01-JUL-14  
 Received Date: 05-JUL-14  
 Extract Date: 08-JUL-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG146127

Analysis Date: 08-JUL-14  
 Analyst: REC  
 Analysis Method: SW846 8260C  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U UJ	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		88.7	%					
Toluene-d8		96.0	%					
1,2-Dichloroethane-d4		100.	%					
Dibromofluoromethane		94.5	%					

R 12/18/14



## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4962-1  
**Client ID:** 147-TB-070214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4962  
**Lab File ID:** C8037.D

**Sample Date:** 02-JUL-14  
**Received Date:** 05-JUL-14  
**Extract Date:** 08-JUL-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG146127

**Analysis Date:** 08-JUL-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260C  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	<del>U</del> UJ	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	<del>U</del> UJ	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*R. 12/14/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4962-1  
**Client ID:** 147-TB-070214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4962  
**Lab File ID:** C8037.D

**Sample Date:** 02-JUL-14  
**Received Date:** 05-JUL-14  
**Extract Date:** 08-JUL-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG146127

**Analysis Date:** 08-JUL-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260C  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 09-JUL-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		89.9	%					
Toluene-d8		102.	%					
1,2-Dichloroethane-d4		101.	%					
Dibromofluoromethane		94.6	%					

**Section 5**

**VPB 147 Analytical Data Table**

Location		VPB147	VPB147	VPB147	VPB147
Sample Date	NYSDEC	6/5/2014	6/5/2014	6/6/2014	6/6/2014
Sample ID	Groundwater Guidance or Standard Value	VPB147-GW-060514- 58-60	VPB147-GW-060514- 98-100	VPB147-GW-060614- 148-150	VPB147-GW-060614- 198-200
Sample Interval	(Note 1)	58 - 60 ft	98 - 100 ft	148 - 150 ft	198 - 200 ft
Sample type code		N	N	N	N
<b>VOC 8260B (ug/L)</b>					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< <b>0.75 U</b>	< <b>0.75 U</b>	< <b>0.75 U</b>	< <b>0.75 U</b>
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
2-HEXANONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
4-METHYL-2-PENTANONE	NL	< 2.5 U	< 2.5 U	< 2.5 UJ	< 2.5 UJ
ACETONE	50	<b>12</b>	<b>2.3</b>	<b>13</b>	<b>6.9</b>
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 UJ	< 1.0 UJ
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	<b>0.35 J</b>	<b>0.29 J</b>
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 U	< 1.0 U	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	<b>0.69 J</b>	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Location		VPB147	VPB147	VPB147	VPB147
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/9/2014	6/9/2014	6/9/2014	6/9/2014
Sample ID		VPB147-GW-060914- 218-220	VPB147-GWD-060914	VPB147-GW-060914- 238-240	VPB147-GW-060914- 258-260
Sample Interval		218 - 220 ft	218 - 220 ft	238 - 240 ft	258 - 260 ft
Sample type code		N	FD	N	N
VOC 8260B (ug/L)					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
2-HEXANONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 2.5 U	3.3 J	< 2.5 U
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Location	VPB147	VPB147	VPB147	VPB147	
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/9/2014	6/10/2014	6/10/2014	6/10/2014
Sample ID		VPB147-GW-060914- 278-280	VPB147-GW-061014- 298-300	VPB147-GW-061014- 318-320	VPB147-GW-061014- 338-340
Sample Interval		278 - 280 ft	298 - 300 ft	318 - 320 ft	338 - 340 ft
Sample type code		N	N	N	N
<b>VOC 8260B (ug/L)</b>					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< <b>0.75 U</b>	< <b>0.75 U</b>	< <b>0.75 U</b>	< <b>0.75 U</b>
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
2-HEXANONE	50	< 2.5 U	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 2.5 U	<b>4.5 J</b>	<b>12</b>
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 UJ	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Location	VPB147	VPB147	VPB147	VPB147
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/11/2014	6/11/2014	6/12/2014
Sample ID	VPB147-GW-061114- 358-360	VPB147-GW-061114- 378-380	VPB147-GW-061214- 398-400	VPB147-GWD-061214
Sample Interval	358 - 360 ft	378 - 380 ft	398 - 400 ft	398 - 400 ft
Sample type code	N	N	N	FD
VOC 8260B (ug/L)				
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 UJ
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 U	< 2.5 U	< 2.5 U
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	5.2	6.4	7.8 J
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 1.0 U	< 0.50 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 U
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 U
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U

Location	VPB147	VPB147	VPB147	VPB147	
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/12/2014	6/12/2014	6/13/2014	6/13/2014
Sample ID		VPB147-GW-061214- 418-420	VPB147-GW-061214- 438-440	VPB147-GW-061314- 458-460	VPB147-GW-061314- 478-480
Sample Interval		418 - 420 ft	438 - 440 ft	458 - 460 ft	478 - 480 ft
Sample type code		N	N	N	N
<b>VOC 8260B (ug/L)</b>					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 UJ	< 0.75 UJ	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 U	< 2.5 U	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	7.3 J	4.9 J	4.2 J
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	< 1.0 U	< 1.0 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 U	< 1.0 U	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	0.45 J
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U



Location	VPB147	VPB147	VPB147	VPB147	
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/17/2014	6/17/2014	6/17/2014	6/18/2014
Sample ID		VPB147-GW-061714- 528-530	VPB147-GW-061714- 538-540	VPB147-GW-061714- 558-560	VPB147-GW-061814- 578-580
Sample Interval		528 - 530 ft	538 - 540 ft	558 - 560 ft	578 - 580 ft
Sample type code		N	N	N	N
VOC 8260B (ug/L)					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	5.0 J	< 2.5 UJ	< 2.5 UJ	1.9 J
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	29	6.3	19	14
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	0.56 J	< 1.0 U	< 1.0 U	0.43 J
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Location	VPB147	VPB147	VPB147	VPB147	
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/18/2014	6/18/2014	6/19/2014	6/19/2014
Sample ID		VPB147-GW-061814- 598-600	VPB147-GW-061814- 618-620	VPB147-GW-061914- 638-640	VPB147-GW-061914- 668-670
Sample Interval		598 - 600 ft	618 - 620 ft	638 - 640 ft	668 - 670 ft
Sample type code		N	N	N	N
VOC 8260B (ug/L)					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	3.7 J	7.2	8.8	< 2.5 U
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 UJ
CARBON DISULFIDE	60	< 1.0 U	< 1.0 U	< 0.50 U	< 0.50 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 UJ	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Location	VPB147	VPB147	VPB147	VPB147
Sample Date	6/20/2014	6/23/2014	6/23/2014	6/24/2014
Sample ID	NYSDEC Groundwater Guidance or Standard Value (Note 1)	VPB147-GW-062014- 678-680	VPB147-GW-062314- 718-720	VPB147-GW-062314- 738-740
Sample Interval	678 - 680 ft	718 - 720 ft	738 - 740 ft	758 - 760 ft
Sample type code	N	N	N	N
<b>VOC 8260B (ug/L)</b>				
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< <b>0.75 U</b>	< <b>0.75 U</b>	< <b>0.75 U</b>
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U
2-BUTANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	<b>7.2</b>	<b>15</b>	<b>8.0</b>
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	<b>0.58 J</b>
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>
CYCLOHEXANE	NL	< 0.50 UJ	< 0.50 UJ	< 0.50 UJ
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< <b>0.50 U</b>	< <b>0.50 U</b>	< <b>0.50 U</b>
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U

Location	VPB147	VPB147	VPB147	VPB147	
Sample Date	NYSDEC Groundwater Guidance or Standard Value (Note 1)	6/25/2014	6/25/2014	6/26/2014	6/26/2014
Sample ID		VPB147-GW-062514- 798-800	VPB147-GW-062514- 818-820	VPB147-GW-062614- 840-842	VPB147-GW-062614- 858-860
Sample Interval		798 - 800 ft	818 - 820 ft	840 - 842 ft	858 - 860 ft
Sample type code		N	N	N	N
VOC 8260B (ug/L)					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,1-DICHLOROETHANE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,1-DICHLOROETHENE	5	< 0.50 UJ	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 1.5 UJ	< 0.75 UJ	< 0.75 UJ
1,2-DIBROMOETHANE	NL	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,2-DICHLOROBENZENE	3	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,2-DICHLOROETHANE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
1,2-DICHLOROPROPANE	1	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,3-DICHLOROBENZENE	3	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
1,4-DICHLOROBENZENE	3	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
2-BUTANONE	50	< 2.5 U	< 5.0 UJ	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 U	< 5.0 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 U	< 5.0 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 34 UJ	< 5.0 UJ	< 12 UJ
BENZENE	1	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
BROMODICHLOROMETHANE	50	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
BROMOFORM	50	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
BROMOMETHANE	5	< 1.0 UJ	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
CARBON DISULFIDE	60	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
CARBON TETRACHLORIDE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
CHLOROBENZENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
CHLOROETHANE	5	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
CHLOROMETHANE	5	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
CYCLOHEXANE	NL	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
DICHLORODIFLUOROMETHANE	5	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
ISOPROPYLBENZENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
M- AND P-XYLENE	NL	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
METHYL ACETATE	NL	< 0.75 U	< 1.5 UJ	< 0.75 UJ	< 0.75 UJ
METHYL CYCLOHEXANE	NL	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
METHYLENE CHLORIDE	5	< 2.5 U	< 5.0 UJ	< 2.5 UJ	< 2.5 UJ
O-XYLENE	NL	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
STYRENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
TETRACHLOROETHENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
TOLUENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
TRICHLOROETHENE	5	< 0.50 U	< 1.0 UJ	< 0.50 UJ	< 0.50 UJ
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
VINYL CHLORIDE	2	< 1.0 U	< 2.0 UJ	< 1.0 UJ	< 1.0 UJ
XYLENES, TOTAL	5	< 1.5 U	< 3.0 UJ	< 1.5 UJ	< 1.5 UJ

Location		VPB147	VPB147
Sample Date	NYSDEC	6/27/2014	6/27/2014
Sample ID	Groundwater Guidance or Standard Value	VPB147-GW-062714- 878-880	VPB147-GW-062714- 888-890
Sample Interval	(Note 1)	878 - 880 ft	888 - 890 ft
Sample type code		N	N
<b>VOC 8260B (ug/L)</b>			
1,1,1-TRICHLOROETHANE	5	< 2.0 UJ	< 0.50 UJ
1,1,2,2-TETRACHLOROETHANE	5	< 2.0 UJ	< 0.50 UJ
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 2.0 UJ	< 0.50 UJ
1,1,2-TRICHLOROETHANE	1	< 2.0 UJ	< 0.50 UJ
1,1-DICHLOROETHANE	5	< 2.0 UJ	< 0.50 UJ
1,1-DICHLOROETHENE	5	< 2.0 UJ	< 0.50 UJ
1,2,4-TRICHLOROBENZENE	5	< 2.0 UJ	< 0.50 UJ
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 3.0 UJ	< 0.75 UJ
1,2-DIBROMOETHANE	NL	< 2.0 UJ	< 0.50 UJ
1,2-DICHLOROBENZENE	3	< 2.0 UJ	< 0.50 UJ
1,2-DICHLOROETHANE	5	< 2.0 UJ	< 0.50 UJ
1,2-DICHLOROETHENE, TOTAL	5	< 4.0 UJ	< 1.0 UJ
1,2-DICHLOROPROPANE	1	< 2.0 UJ	< 0.50 UJ
1,3-DICHLOROBENZENE	3	< 2.0 UJ	< 0.50 UJ
1,4-DICHLOROBENZENE	3	< 2.0 UJ	< 0.50 UJ
2-BUTANONE	50	< 10 UJ	13 J
2-HEXANONE	50	< 10 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 10 UJ	< 2.5 UJ
ACETONE	50	24 J	85 J
BENZENE	1	< 2.0 UJ	< 0.50 UJ
BROMODICHLOROMETHANE	50	< 2.0 UJ	< 0.50 UJ
BROMOFORM	50	< 2.0 UJ	< 0.50 UJ
BROMOMETHANE	5	< 4.0 UJ	< 1.0 UJ
CARBON DISULFIDE	60	< 2.0 UJ	< 0.50 UJ
CARBON TETRACHLORIDE	5	< 2.0 UJ	< 0.50 UJ
CHLOROBENZENE	5	< 2.0 UJ	< 0.50 UJ
CHLOROETHANE	5	< 4.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 2.0 UJ	< 0.50 UJ
CHLOROMETHANE	5	< 4.0 UJ	< 1.0 UJ
CIS-1,2-DICHLOROETHENE	5	< 2.0 UJ	< 0.50 UJ
CIS-1,3-DICHLOROPROPENE	0.4	< 2.0 UJ	< 0.50 UJ
CYCLOHEXANE	NL	< 2.0 UJ	< 0.50 UJ
DIBROMOCHLOROMETHANE	5	< 2.0 UJ	< 0.50 UJ
DICHLORODIFLUOROMETHANE	5	< 4.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 2.0 UJ	< 0.50 UJ
ISOPROPYLBENZENE	5	< 2.0 UJ	< 0.50 UJ
M- AND P-XYLENE	NL	< 4.0 UJ	< 1.0 UJ
METHYL ACETATE	NL	< 3.0 UJ	< 0.75 UJ
METHYL CYCLOHEXANE	NL	< 2.0 UJ	< 0.50 UJ
METHYL TERT-BUTYL ETHER	10	< 2.0 UJ	< 0.50 UJ
METHYLENE CHLORIDE	5	16 J	< 2.5 UJ
O-XYLENE	NL	< 2.0 UJ	< 0.50 UJ
STYRENE	5	< 2.0 UJ	< 0.50 UJ
TETRACHLOROETHENE	5	< 2.0 UJ	< 0.50 UJ
TOLUENE	5	< 2.0 UJ	< 0.50 UJ
TRANS-1,2-DICHLOROETHENE	5	< 2.0 UJ	< 0.50 UJ
TRANS-1,3-DICHLOROPROPENE	0.4	< 2.0 UJ	< 0.50 UJ
TRICHLOROETHENE	5	< 2.0 UJ	< 0.50 UJ
TRICHLOROFLUOROMETHANE	5	< 4.0 UJ	< 1.0 UJ
VINYL CHLORIDE	2	< 4.0 UJ	< 1.0 UJ
XYLENES, TOTAL	5	< 6.0 UJ	< 1.5 UJ

**Notes:**

1 New York State Department of Environmental Conservation Division of Water Technical and Operation Guidance series  
(6 NYCRR 700-706, Part 703.5 summarized in TOGS 1.1.1)

Ambient water quality standards and groundwater effluent limitations, class GA; NL = Not Listed

**Bold** = Detected; ***Bold and Italics*** = Detection limit exceeds NYS Groundwater Standards or guidance value

**Yellow** highlighted values exceed Groundwater Standards or guidance value

Sample type codes: N - normal environmental sample, FD - field duplicate

U = Nondetected result. The analyte was analyzed for, but was not detected above the reported sample quantitation limit.

UJ = The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte.

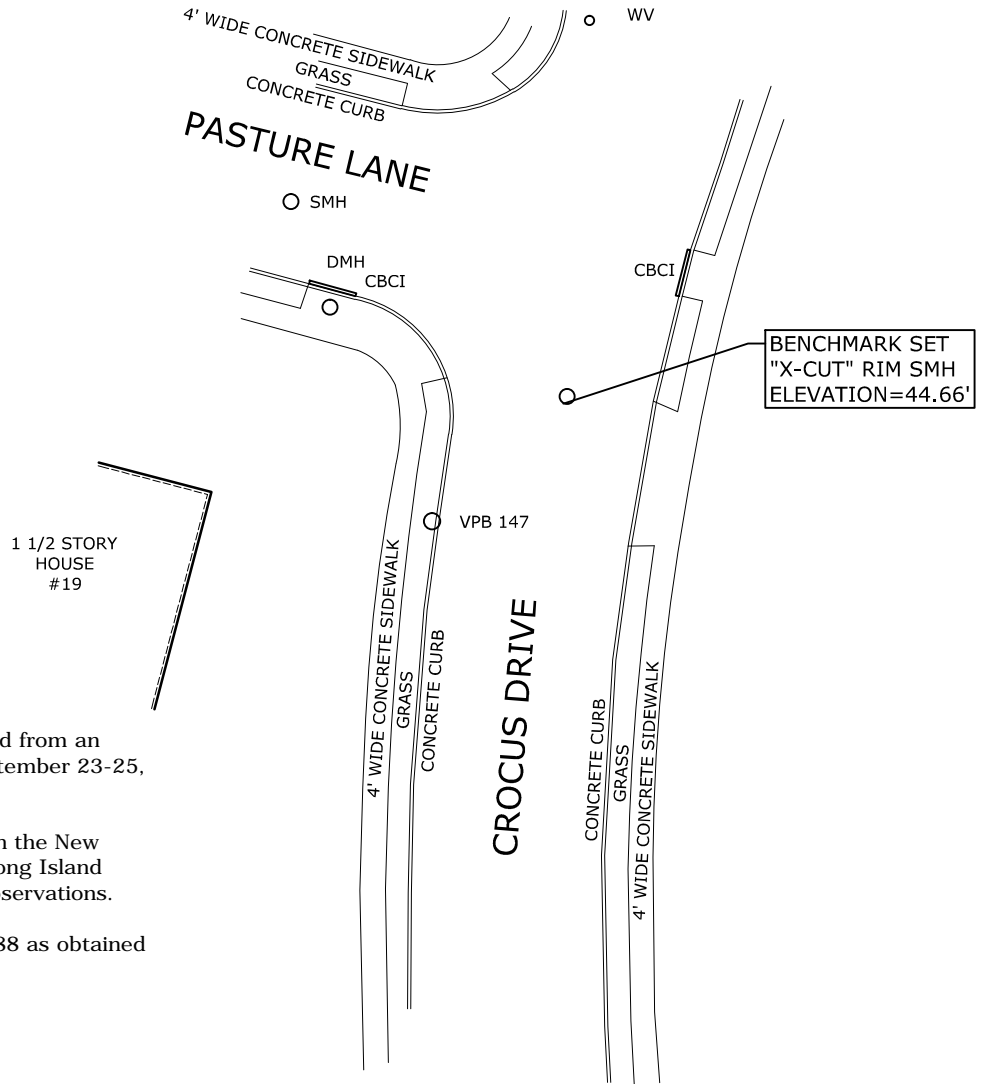
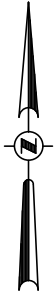
J = The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.

## **Section 6**

### **Survey**

UNAUTHORIZED ALTERATION OR ADDITION TO THIS DOCUMENT IS A VIOLATION OF SECTION 7209 SUBDIVISION 2 OF THE NEW YORK STATE EDUCATION LAW.

Description	Northing	Easting	Latitude	Longitude	Ground	Rim	PVC
VPB 147	193658.09	1128979.90	N40-41-49.34	W73-28-41.01	44.92	NA	NA

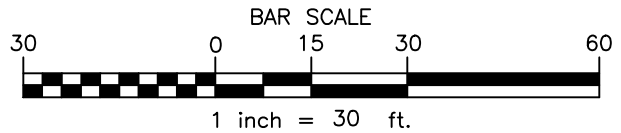


Map Notes

- Information shown hereon was compiled from an actual field survey conducted from September 23-25, 2014.
- North orientation is Grid North based on the New York State Plane Coordinate System, Long Island Zone, NAD 83 as obtained from GPS observations.
- Vertical datum shown hereon is NAVD 88 as obtained from GPS observations.

Legend

- CIBC Catch Basin Curb Inlet
- DMH Drainage Manhole
- SMH Sanitary Manhole
- VPB Vertical Profile Boring
- WV Water Valve



DWG NO. 14-501

Date	RECORD OF WORK	Appr.	VERTICAL PROFILE BORING 147 SURVEY LOCATION 19 CROCUS DRIVE	
			TOWN OF MASSAPEQUA	NASSAU COUNTY, NEW YORK
			<b>C.T. MALE ASSOCIATES</b> Engineering, Surveying, Architecture & Landscape Architecture, D.P.C.	
			50 CENTURY HILL DRIVE, LATHAM, NY 12110 518.786.7400 * FAX 518.786.7299	
			SCALE: 1"=30'	
			DATE: SEPT. 24, 2014	
Drafter: LMK		Checker: JFC		
Appr. by: JFC		Proj. No. 14.4121		

