Project No. 444-008E-96

PROPOSED DESIGN PLAN for a Soil Vapor Extraction (SVE) and Control System

February 1997

Prepared for

Jameco Industries 248 Wyandanch Avenue Wyandanch, NY

NYSDEC Site No. <u>1-52-006</u>

Prepared by:

GEC

 $Goldman\ Environmental\ Consultants, Inc.$

60 Brooks Drive Braintree, MA 02184 (617)-356-9140

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1.0 <u>INTRODUCTION</u>

The following Design Plan has been prepared by Goldman Environmental Consultants, Inc. (GEC), on behalf of Watts Industries, Inc. (Watts), in response to NYSDEC-approved, Interim Remedial Measure (IRM) Work Plan and NYSDEC Consent Order No. DI-0001-95-08. This Design Plan has been prepared consistent with Section 2.2 of the NYSDEC-approved Work Plan and in accordance with 6 NYCRR Part 375, for an Interim Remedial Measure (IRM). This Design Plan is recommended in order to reduce concentrations of chlorinated compounds detected beneath a portion of the site building, presumably resulting from a release of these compounds from a degreasing tank, currently in use at the site.

In the Summer of 1994, GEC oversaw a soil gas survey where soil gas samples were collected over a 111 ft X 255 ft area inside the building near the plating and degreasing operations. During this survey, concentrations of total ionizable compounds (TICs) were detected as high as 1,800 ppm and it was determined that soil gas contamination does not extend beyond the building's footprint. Chlorinated compounds are only present in the shallow portion of the water table aquifer, and there is no evidence to indicate that contamination extends to lower portions of the aquifer. Existing facility data were summarized in the "Initial Submittal", which was prepared consistent with Section I. of the Consent Order and provided to NYSDEC on May 1995.

The presence of elevated concentrations of volatile compounds in soil gas, the porous nature of the soil and volatile nature of the contamination readily lends itself to remediation by soil vapor extraction (SVE) over other remediation approaches. This August, GEC oversaw a SVE pilot test which confirmed that SVE is an effective remediation approach for chlorinated compounds in the soil. A SVE system was also viewed as a cost-effective and rapid remedy for reducing soil gas concentrations for this site.

2.0 SOIL VAPOR EXTRACTION PILOT TEST

In accordance with NYSDEC-approved, Interim Remedial Measure (IRM) Work Plan, Goldman Environmental Consultants, Inc. (GEC) in conjunction with our subcontractor GTS Environmental Technology, Inc. (GTS) conducted a Soil Vapor Extraction (SVE) pilot test on August 8, 1996, at Jameco Industries, Inc., Wyandanch, NY. The purpose of the pilot test was to determine the feasibility of remediating soils and to identify design parameters for a full scale extraction system.

The SVE pilot test included the extraction of soil vapor from existing monitoring well MW-12 (see Drawing No. L-2). This is a 2-inch diameter PVC well screened from 5 feet down to 15 feet below grade. The water table is approximately ten feet below grade. The pilot test was conducted using a Rotron Model DR505R blower driven by a five (5) horsepower (HP) motor. Two-inch and three-inch diameter PVC pipes were connected from the blower to the MW-12 well (see Figure 1). A 55 gallon water knock-out drum was installed between the extraction well and the blower to separate and collect any water extracted. The blower was driven at a constant speed of 4,500 rpm. The vacuum applied to the extraction point was adjusted by bleeding in ambient air through a bypass valve. During the pilot test, appropriate measurements were taken to determine the vacuum, flow rate, effluent concentration, and zone of influence. A 120 pound activated carbon vessel was used to control VOC air emissions prior to discharge to the atmosphere.

2.1 Soil Impedance Test

A soil impedance test was conducted to determine the flow rate as a function of applied vacuum. The following Table 1 presents the results of the soil impedance test, and Figure 1 illustrates the pilot test system configuration.

Table 1: Results of Soil Impedance Test					
Vacuum (inches of water)	Gauge Velocity (FPM)	Flow (CFM)			
6"	0	0			
10"	500	22			
15"	1,100	49			
20"	1,200	53			
25"	1,600	71			
30"	2,000	88			

CFM = Area* Velocity * 90%

The above table indicates that a 30-inch vacuum applied produced a flow of 88 CFM, indicating very permeable unsaturated zone conditions.

2.2 Zone of Influence

The zone of influence was evaluated by extracting soil vapors from test wells at a fixed rate while monitoring the subsurface pressure distribution at six soil probes installed at various distances from the test well. Soil probe locations are identified in Drawing No. L-2. At 30" vacuum applied, MW-12 produced a 3" soil vacuum at a distance of 10 feet and 1.1" soil vacuum at a distance of 40 feet. Higher vacuum applied to MW-12 would likely result in an increased zone of influence, however, during the testing, unsteady and variable vacuum and velocity readings were observed at the extraction point, presumably due to groundwater intrusions. The following Table 2 presents the zone of influence test results.

Table 2: Zone of Influence Test Results						
Vapor Probe	Distance from MW-12 (ft.)	Vacuum (inches of water)				
VP-1	6'	3" +				
VP-2	10'	3"				
VP-3	15'	2.2"				
VP-4	22'	1.3"				
VP-5	30'	0.75"				
VP-6	40'	1.1"				

The above data demonstrate that soil vacuum at all selected distances out to 40 feet can be achieved at flows in the range of 80 to 90 SCFM. An effective radius of influence significantly in excess of 40 feet appears possible.

2.3 **Groundwater Intrusion**

During the pilot test, groundwater was extracted with the VOC vapors and collected in the Knock-Out Tank. Groundwater intrusion is typical in 2inch monitoring wells with screen extension below the water table. The applied vacuum raises the water level in the well, and the high vapor flow velocity entrains water droplets. The pilot test ran for 90 minutes and groundwater accumulation was approximately 20 gallons, which is a fairly high volume.

Planned extraction wells will be set above the known high water table elevation. This will significantly reduce groundwater accumulation in the moisture separator and therefore the groundwater volume requiring treatment will be much lower.

2.4 Field Screening Results

Volatile organic compound (VOC) concentrations were measured in the field using an HNu photoionization detector. Soil vapor extracted from MW-12 was found to contain an average of 55 ppm VOCs. The following Table 3 illustrates that the pre-carbon VOC concentrations ranged from 40 to 72 ppm.

Carbon adsorption reduced VOCs in the extracted vapor by an average of more than 95%.

Table 3: VOC Concentrations in MW-12 Gas Before and After Activated Carbon Treatment					
Time (min)	Pre-Carbon Sample Concentration (ppm)	Post-Carbon Sample Concentration (ppm)	% VOC Reduction		
Start	60	5	92		
10	72				
20	70	3.5	95		
30	60	2	96.7		
40	42				
50	44	0.8	98		
60	58				
70		3	94.5		
80	44				
90	40	2	95		
Average	55 ppm	2.72 ppm	95.2%		

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VOC Concentration vs Time

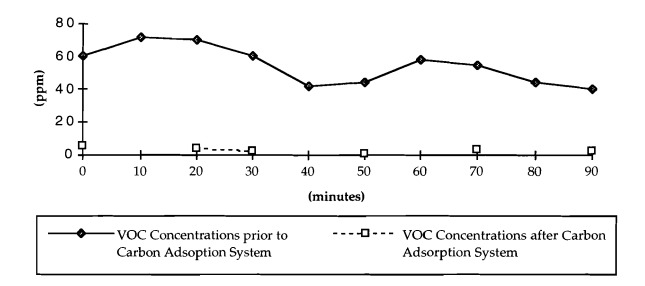


Figure 2: Results of Scrubbing Well MW-12 Gas with Activated Carbon

3.0 SAMPLING AND ANALYSIS PLAN

GEC has prepared a Sampling and Analysis (S&A) Plan which conforms with EPA and NYSDEC methods, guidance, and applicable regulations. Following are each of the specific elements required for a S&A Plan.

3.1 Quality Assurance and Quality Control Plan

The Quality Assurance and Quality Control (QA/QC) Plan for all site activities is included as Appendix A. All site QA/QC efforts are managed by Mr. Robert Fricke, GEC's Vice President and QA/QC Manager. He has data validation responsibility for site activities and over 15 years of experience managing hazardous waste site programs. Mr. Fricke's resume, along with those of all anticipated site personnel, are contained in Appendix A. All site personnel have at least 40 hours of EPA Hazardous Waste Site Worker Training and are thoroughly trained in the QA/QC Plan for the site.

The goal of the QA/QC Plan, as it relates to the S&A Plan, is to provide a framework for achieving data quality and analytical precision that is consistent with good engineering practice for the installation, and operation of a SVE and control system for the Design Plan.

3.2 Health and Safety Plan

Appendix B contains the Health and Safety (H&S) Plan that will be used for all site activities. All installation personnel have read the Plan. Daily Safety meetings will be held prior to the start of installation activities.

The Health and Safety plan addresses remediation personnel. To address concerns of local residents and Jameco employees, GEC will continuously monitor total ionizable compounds (TICs) during any invasive field activities, including the installation of the SVE system and associated drilling activities. In the event that elevated concentrations (greater than 5 parts per million) of TICs are detected, GEC will follow applicable procedures as outlined in the NYSDOH Community Air Monitoring Plan.

GEC will also implement measures to assure that fugitive dust is minimized. To minimize the volume of dust generated, GEC will assure that all work areas where the potential for dust generation exists are thoroughly wetted. GEC anticipates potential dust generation associated with the penetration of the concrete and cutting of concrete during SVE system installation. GEC does not anticipate conducting any soil excavation and will not be moving heavy equipment across unpaved areas of the site as part of the installation/operation of the SVE system. Due to the limited scope of activities in which the potential for dust generation exists, GEC does not propose to conduct particulate monitoring.

3.3 Contingency Plan

A site Contingency Plan is included with the Health and Safety Plan in Appendix B. The Contingency Plan will be used for all site activities. The Plan conforms with EPA requirements for Superfund field operations, and all site personnel are trained in Plan elements. In addition, site activities will be conducted in accordance with Jameco's Contingency Plan.

The proposed and approved S&A Plan will be thoroughly reviewed with facility personnel, who will be involved in the installation and operation of SVE system. In particular, the H&S Plan and Contingency Plan will be incorporated into and conform with facility plans that are the responsibility of Jameco's Environmental Health and Safety Manager.

After NYSDEC approves this Design Plan and the S&A Plan of this section, GEC will implement the S&A Plan and start ordering the SVE system components for installation.

4.0 SVE ENGINEERING DESIGN

The following SVE design considerations are based on information developed during the pilot test.

4.1 Design Air Flow Rate

During the pilot test, the vacuum of 30 inches of water was applied which yielded approximately 88 SCFM and an influence area of approximately 5,027 sqft. (based on 40 foot radius of influence). The estimated contaminated area is 22,231 sqft. based on the soil gas survey. The design flow for the full scale extraction well is calculated as follows:

Area of influence = $\pi \times (ROI)^2 = 3.14159 \times (40)^2 = 5,027 \text{ ft}^2$

Area of contaminated soil (oval shape)= $\pi \times (255 \text{ ft./2}) \times (111 \text{ ft/2}) = 22,231 \text{ ft}^2$

Air Flow = $(88 \text{ CFM}/5,027 \text{ ft}^2) \times (22,231 \text{ ft}^2) = 389 \text{ CFM}$

Air Flow Required for Vapor Extraction = 389 CFM

The calculated vapor extraction flow rate is 389 CFM at a vacuum of 30 inches of water. However for the full scale SVE system, the Design Air Flow Rate should be 600 SCFM at vacuum of 40 inches of water.

Design Air Flow Rate = 600 CFM

4.2 Extraction Wells

Soil vapor can be extracted either through vertical wells or horizontal trenches. Extraction wells are most appropriate when soil is fairly permeable and the water table is relatively deep. Given the prevailing conditions at the Jameco site, vertical extraction wells are appropriate.

The location and number of extraction wells that need to be located within the treatment area to provide area coverage can be calculated as follows:

Number of wells needed = $(Contaminated area (ft^2))/(Area of influence (ft^2/well)$ = $(22,231 \text{ ft}^2)/(5,027 \text{ ft}^2/\text{well}) = 4.4 \text{ wells} \approx 5 \text{ Wells}$

Number of Wells Required = 6 (see Drawing No. L-4)

Six (6) extraction wells should be installed to remediate the site. Refer to Drawing No. L-4 for proposed soil vapor extraction well locations.

4.3 SVE System Design Criteria

Based on the Design Air Flow Rate and the number of extraction wells, the SVE system should be capable of extracting VOC vapors at the rate of 600 ACFM at a vacuum of 40 inches of water and a blower discharge pressure of 0.5 psig. The SVE system should include a moisture separator with level control sensors to control the operation of the transfer pump, a blower, inlet particulate filter unit, inlet silencer, and discharge silencer. The SVE system should include provisions to continuously monitor the extraction well vacuum, air flow, and air temperature. Pressure and temperature on the discharge side of the system should also be monitored.

4.4 <u>Product Recovery Calculations</u>

During a soil gas survey, chlorinated compounds were detected within the building at concentrations as high as 1800 ppm. The chlorinated compounds were also detected at a depth of 10 feet during the installation of monitoring well MW-12. During the pilot test, the VOC concentrations were measured as high as 72 ppm and will probably drop below 50 ppm within two (2) days of system operation.

Product recovery rates and duration can be calculated as follows:

<u>Assumptions</u>

Maximum contaminant concentration of total organic vapors = 1,800 ppm

Average contaminant concentration (ACC) = 800 ppm

Estimated air flow volume (CFM) = 600 cfm

Conversion factor (CF) of TCE 1 ppm = $5.46 \text{ mg/m}^3 = 3.4 \times 10^{-7} \text{ lb VOC/ft}^3 \text{ air } @ 60^{\circ}\text{F}$

Soil Volume

Area of contaminated soil (oval shape) $= \pi \times (255 \text{ ft./2}) \times (111 \text{ ft/2}) = 22,231 \text{ ft}^2$

Thickness of contaminated soil column = 10 ft.

Total volume of contaminated soil (SV) = $22,231 \text{ ft}^2 \times 10 \text{ ft} = 222,310 \text{ ft}^3$

Total contaminant loading

800 ppm = 800×10^{-6} lb VOC/lb Soil

Assumed specfic gravity is 2

Hence specific weight $= 2 \times 62.4 \text{ lb Soil/ft}^3 \text{ Soil} = 124.8 \text{ lbs Soil/ft}^3 \text{ Soil}$

800 ppm = 800×10^{-6} lb VOC/lb Soil x 124.8 lbs Soil/ft³ Soil

= 0.1027 lb VOC/ft³ Soil

Total Contaminant loading = SV x 0.1027 lb VOC/ft³ Soil

 $= 222,310 \text{ ft}^3 \times 0.1027 \text{ lb VOC/ft}^3 \text{ Soil}$

= 22,836 lb VOC

Anticipated product recovery rate

80 ppm first two days of system startup

50 ppm (Conc.) thereafter

Estimated Product Recovery Rate at 80 ppm = (VOC Conc) x CF x CFM

 $= 80 \text{ ppm x } 3.4 \text{ X } 10^{-7} \text{x } 600 \text{ acfm}$

= 0.0163 lbs/min. x 60 min/hr

Estimated Product Recovery Rate at 80 ppm = 0.98 lb/hr

Est. removal of contaminated loading = 24 hrs/day x 0.98 lbs/hr

= 23.5 lbs/day at 80 ppm removal rate

Estimated Product Recovery Rate at 50 ppm = (VOC Conc) x CF x CFM

 $= 50 \text{ ppm x } 3.4 \text{ X } 10^{-7} \text{x } 600 \text{ acfm}$

= 0.010lbs/min. x 60 min/hr

Estimated Product Recovery Rate at 10 ppm = 0.6 lbs/hr

Anticipated product recovery Time

Time required to remove the VOC if the removal rate is 80ppm for first two days and 50 ppm thereafter:

2 days x 24 hrs/day x 0.98 lb VOC/hr + X x 0.12 lb VOC/hr = 22,836 lbs VOC

X = (22,836 lb VOC - 47 lb VOC)/0.6 lb VOC/hr = 37,982 hrs

Anticipated Prodcut Removal Time

= 4.34 years

4.5 <u>Vapor Phase Carbon Adsorption System Sizing Calculations</u>

Carbon beds are normally sized for: (1) the amount of material that is to be adsorbed, and (2) humidity associated with soil venting which causes a lengthening of the Mass Transfer Zone. The time to saturate the carbon beds is based on 7% loading and a mass balance.

For 80 ppm VOC Concentration

```
Activated Carbon (AC) Used = (lbs contaminant/min.) / 7\%
= (0.0163 \text{ lbs/min.}) / 0.07 = 0.23 \text{ lb (AC)} used per min.
= 0.23 \text{ lb/min.} \times 60 \text{ min/hr} \times 24 \text{ hr/day}
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Activated Carbon (AC) Used = 331 lbs/day for VOC concentration of 80 PPMV

For **50 ppm** VOC Concentration

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Activated Carbon (AC) Used = (lbs contaminant/min.) / 7\% = (0.01 lbs/min.) / 0.07 = 0.029 lb (AC) used per min. = 0.143lb/ min x 60 min/ hr x 24 hr/ day
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Activated Carbon (AC) Used = 206 lbs/day for VOC concentration of 50 ppmv

The size of vapor phase Carbon Adsorbers to treat off-gases from the vapor extraction system should be based on a flow of 600 cfm and an influent VOC concentration of 80 ppmv for the first two days of system startup and 10 ppmv thereafter. The carbon adsorbers sizing calculations are as follows:

For the first two days of system startup, the amount of carbon needed to adsorb VOC is 331 lbs/day and 206 lbs./day thereafter.

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1,000 lbs. AC - 662 lbs (used in first two days) = 338 lbs/ 206 lbs/day = 1.6 days
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Therefore for system startup, soil gas vapor will load one carbon canister with 7% loading capacity in 2 days + 1.6 day = 3.6 days.

Thereafter, a new 1,000 lbs carbon canister will be replaced every 5 days for the balance of time.

1,000 lbs/ 206 lbs/ day = 5 days

4.6 <u>Estimated VOC Emissions</u>

Based on the calculation in Section 4.4, an Estimated Product Recovery Rate is 0.98 lb/hr or 23.5 lbs/day for first day and thereafter is 0.6 lb/hr or 14.4 lbs/day which is based on 24 hours per day.

During pilot testing, an average %VOC reduction was 95%. The amount of VOC emission after VOC control system is calculated as follows:

The VOC emission for first two days (1st. Period) = 23.5 lbs/day X (1-95%) = 1.18 lb/day

VOC Emission for first Period = 1.18 lb/day

and thereafter
VOC Emissions after two days (2nd Period) =
14.4 lbs/day X (1-95%) = 0.72 lb/day

VOC Emission for Second Period = 0.72 lb/day

5.0 SELECTED SVE AND VAPOR CONTROL SYSTEM

Chlorinated compounds were detected during a soil gas survey, and soil gas samples were collected over a 111 ft X 255 ft area inside the building near the plating and degreasing operations. See Drawing No. L-2 for soil gas sampling locations. The soil gas samples were screened for the presence of total ionizable compounds (TICs) and total organic vapors (TOVs) using a HNu Model HW-101 photoionization detector (PID) and a Foxborough Model Century 128 organic vapor analyzer (OVA). Concentrations as high as 1800 parts per million (ppm) of TICs were detected. Soil gas contamination does not appear to extend beyond the building footprint. The groundwater samples were also collected from the monitoring wells located within the building foot-print, and analytical results indicated the presence of low concentrations of dissolved-phase chlorinated compounds. The chlorinated compounds are only present in the shallow portion of the water table aquifer, and there is no evidence to indicate that contamination extends to the lower portion of the aquifer. Facility data are summarized in a Consent Order "Initial Submittal" to NYSDEC dated May 22, 1995.

Jameco used 1,1,1-trichloroethylene (TCE) solvent in their degreasing operation and the presence of TCE in the soil indicates that the solvent tank may have leaked. According to NIOSH Pocket Guide for Chemical Hazards, TCE has a vapor pressure of 58 mm Hg, a boiling point of 189 °F and a Henry's Law Constant greater than 100 ATM. The intrinsic permeability of the soil at Jameco is greater than 10^{-8} cm².

The porous nature of the soil and volatile nature of the contamination lends itself to the SVE remediation. The information obtained during the pilot test confirms that SVE is an effective remediation approach chlorinated compounds in the soil. SVE is also viewed as a cost-effective and rapid remedy for reducing soil gas concentrations for this application.

The following selected SVE and vapor control system components are based on the pilot test study and design calculations described in Sections 2 and 4 respectively.

5.1 Water Table

The water table at the Jameco site was measured at approximately ten feet below grade and depth to water will likely to be stable throughout most of the year. However, seasonal snow melt and heavy rainfall could result in a higher water table.

5.2 Extraction Wells

Soil vapors will be extracted through six (6) vertical wells located inside the building rather than horizontal trenches because the soil is highly permeable and the water table is relatively deep. See Drawing No. L-4 for extraction well locations. Each extraction well will be installed with 4-inch diameter, Schedule 40 PVC pipe with 0.010-inch slotted screen from 4 feet to 8 feet. The well will have a bentonite seal one foot above the screen and 1.5 feet of cement seal to the sump level.

5.3 Groundwater Intrusion

Since the water table at the Jameco site is approximately 10 feet below grade, six new extraction wells will be installed 2-feet above the water table. During the pilot test, groundwater was extracted because the 2-inch monitoring well with slotted screen was extended below the water table. The groundwater intrusion probably will not occur during normal operation, but some intrusion could result from unusually high water table conditions or by applying a higher than normal vacuum to one or more of the extraction wells. Therefore, a moisture separator with liquid phase treatment system has been included to collect and treat the moisture. Estimated collected groundwater volume is less than one (1) gallon. Due to the volatile nature of the contaminant, collected groundwater will have a minimal amount of VOC. The designed moisture separator for collecting groundwater intrusion has a capacity of 120 gallons and is equipped with level sensors to operate a liquid phase treatment system. The treated groundwater will be discharged to the existing leaching pool area.

5.4 Manifold Piping

Manifold piping connects the extraction wells to the skid-mounted SVE system (see Detail A, Drawing No. PP-1). Piping and fittings will be 4-inch diameter, Schedule 80, Industrial Grade, PVC manufactured from Type 1, Grade 1 conforming to ASTM D-1784. All pipe fittings, including valves, unions and flanges will be pressure rated at 150 psi conforming to ASTM D-1875, ASTM D-2464 AND ASTM D-2467.

Each extraction well will be individually piped, valved, and connected to a manifold located upstream of the SVE system. A flow gage, vacuum gage and air flow control valve will be installed on each pipe at the extraction well. Piping will be sloped toward the extraction well so that the condensate or entrained groundwater flows back toward the well.

5.5 <u>Trenches</u>

Piping will be placed in shallow utility trenches that lead from the extraction well to the SVE system, as shown in Drawing No. UT-1. The trenches will be 12-inches or 24-inches wide and 12-inches deep. Trenches will be epoxy-coated and sloped toward the spill collection sump. Trenches will be covered with medium traffic fiberglass or steel grating panels as manufactured by Fibergrate or equivalent. The panel will be 12-inches or 24-inches wide, 48-inches long, and 1-1/2-inches thick minimum. The spill collection sump will be equipped with a sump pump to pump out any contaminated spill water into drums for disposal in an environmentally safe manner.

5.6 **Skid-Mounted SVE System**

The SVE system will be a skid-mounted VACPAC, Bisco Environmental Inc. Model URAI 59. The SVE system will consist of a moisture separator with level control sensors to control the operation of the transfer pump, a positive displacement blower, inlet particulate filter and silencer, and discharge silencer. The SVE system will include provisions to continuously monitor the extraction well vacuum, air flow, and air temperature; and pressure and temperature on the discharge side of the system will also be monitored. See Drawing No. PID-1 for more details of the

system operation and controls. The following are major components and features of the SVE system.

5.6.1 <u>Positive Displacement Blower</u> (PDB-1)

The blower will be a positive displacement blower Roots model URAI 59. The blower is capable of providing 600 CFM at a vacuum of 80 inches of water.

The blower consist of two "Figure 8" lobe impellers mounted on parallel shafts, rotating in opposite directions. As each impeller passes the blower inlet, it traps a definite volume of air and carries it around the case to the blower outlet, where the air is discharged. With constant speed operation, the displaced air volume is essentially the same regardless of pressure, temperature, or barometric pressure. Timing gears control the relative position of the impellers to each other and maintain small but definite clearances, which allows operation without lubrication being required inside the air casing.

The blower has a cast iron casing; carburized and ground alloy steel, spur timing gears that are secured to steel shafts with a taper mounting and locknut; and cast iron involute impellers.

The blower motor is a foot-mounted, NEMA A Design B, TEFC, Standard Industrial, Continuous Duty, ball bearing, variable-torque-type suitable for operation at 15 HP, 230/460 VAC, 3 phase, and 60 Hertz.

5.6.2 <u>Moisture Separator</u> (MS-1)

The moisture separator will have a liquid capacity of 120 gals and will be designed for use in the soil vapor extraction system. The separator will be capable of continuous operation with a pressure drop of less than six inches of water at the rated flow of 600 SCFM. The separator unit will be capable of operation under various inlet conditions ranging from a fine mist to slugs of water.

The separator unit will be equipped with a transfer pump and liquid level controls. The level control switches will be single polarity, reed-type, float switches. High, low, and mid-level switches will control on-off

functions for the transfer pumps and blower. Liquid levels controlled/monitored will be independently set and field adjustable with levels identified as:

"High Level" - Blower Shut-off "Medium Level" - Transfer Pump On "Low Level" - Transfer Pump Off and Blower On

5.6.3 Separator Transfer Pump (TP-1)

The transfer pump will be Burk Pump, Model GA4-3/4. The pump will be capable of providing pumping capacity of up to 30 GPM at up to 30 ft of water total head pressure. The pump body will be constructed of cast iron/bronze with a Viton mechanical seal and close-coupled to a 1/2 HP, 3450 RPM, 230/460 VAC, 3 phase, 60 Hz, TEFC motor.

5.6.4 <u>Inlet Particular Filter</u> (IPF)

The inlet particulate filter will be installed after the moisture separator and before the positive displacement blower. The inlet particulate filter will be a Solberg Model CSL-335P-400F capable of a 96-98% removal efficiency in the 2 to 4 micron size range. The filter unit will protect the blower from harmful dust and other particles that may be drawn into the blower through the air distribution system.

5.6.5 <u>Discharge Silencer</u>

The discharge silencer will be installed after the positive displacement blower and will be a Universal Silencer Model RD Series. The discharge silencer will be a heavy-duty, all welded unit constructed of carbon steel and plate. It will provide pulse control and will be equipped with an acoustically treated inlet.

5.7 Vapor Control System

During the pilot test, VOC concentrations were measured as high as 72 ppm and will probably drop below 10 ppm within two (2) days after system startup. GEC recommends carbon adsorption because of low VOC loading and relatively high air flow needs. See Section 4.5 for equipment sizing calculations.

5.7.1 <u>Vapor Phase Carbon Adsorption System</u>

Vapor phase carbon adsorption systems separate organic contaminants from an air stream by accumulation of the organics on the surface of a granular activated carbon. This process is driven by molecular attraction between the contaminant and the activated carbon surface. One gram of a typical commercial activated carbon has a surface area equivalent to 1,000 square meters. This high surface area permits the accumulation of a large number of contaminant molecules.

The specific capacity of an activated carbon to adsorb volatile organic compounds is related to: molecular surface attraction, the total surface area per unit weight of carbon, and the concentration of volatile compounds in the air stream. As a contaminated air stream passes through a confined bed of activated carbon, a dynamic condition develops which establishes a mass transfer zone. The "mass transfer zone" requires a certain carbon bed depth to reduce the contaminant concentration from the initial to the final level, at a given gas flow rate.

The activated carbon beds should be packed with 1,000 lbs of activated carbon to provide approximately 3.6 days of VOC treatment for the first cycle and 5 days thereafter. The activated carbon bed size depends on the organic contaminant concentration in the air stream. Based on the calculations described in Section 4.5, an air stream with 80 ppm of VOC contamination will use 331 lbs per day of activated carbon, and an air stream with 50 ppm of VOC contamination will use 206 lbs per day.

As the mass transfer zone moves through a carbon bed and reaches its exit boundary, contamination begins to show in the effluent. This condition is known as "breakthrough" and the amount of material adsorbed is considered the breakthrough capacity.

Carbon Adsorption is an extremely versatile technology and is the least expensive for the organic contaminated air stream that will be generated by the proposed SVE system. Activated carbon adsorption is particularly effective in treating vapor streams with low VOC concentrations. We anticipate that the mass removal rate at 80 ppm will be 0.98 lb/hr and 0.6 lb/hr at 50 ppm.

5.8 <u>Liquid Phase Carbon Adsorption System</u>

The liquid phase carbon adsorption system will collect the molecules of a dissolved compound as the molecules adhere to the surface of activated carbon. Adsorption occurs when the attractive forces at the carbon surface overcome the attractive forces of the liquid.

The carbon system will be a Bisco Model AF-55 which includes 175 pounds of activated carbon to treat the effluent from the moisture separator. The unit will be an epoxy-lined steel vessel with a 2-inch diameter inlet/outlet to treat a maximum of 10 GPM of contaminated effluent.

5.9 Control Panel

A control panel (CP-1) cabinet will mount the programmable logic controller (PLC) with remote monitoring and fax/pager notification capabilities. The control panel will be an EOS ProControl Series II which will be capable of remote control and monitoring, fax and pager reporting, and datalogging. The control panel cabinet will be a NEMA 4 weather-tight enclosure which will include: two (2) motor starters with thermal overload protection; two (2) three-position "Hand/Off/Auto" (HOA) selector switches; two (2) motor status indicator lights; three (3) alarm status indicator lights - separate and distinct lights for each alarm condition; common manual reset button; and circuitry to operate and interface a pressure transmitter with an electrically-actuated ball valve. The control panel will be designed for 230 VAC, 3 phase, 60Hz service.

6.0 SVE SYSTEM OPERATION AND MONITORING PLAN

After NYSDEC approves the SVE Design Plan, the system components will be ordered from the selected vendors; and GEC will oversee the system installation, start-up, and operation of the remediation system. During installation, GEC will inspect the system components to ensure that they are meeting the equipment specifications, and that the vendor has installed the SVE system according to the design plans. After installation is complete, GEC will implement the operation and monitoring plans. Operation and monitoring of the system are necessary to ensure that the system performance is optimized and the contaminant mass removal rate is documented.

6.1 Start-up Operation/Monitoring

The start-up phase will take 5 to 7 days and will include manifold valving adjustments. These adjustments will optimize contaminant mass removal by biasing vacuum pressure on the extraction wells that produce vapor with higher concentrations of VOC. Flow measurements and vacuum readings will be monitored continuously and vapor concentration will be monitored/recorded daily from each well, manifold, and exhaust vent.

6.2 <u>Long-Term Operations/Monitoring</u>

Long-term monitoring will consist of flow-balancing, flow and pressure measurements, and vapor concentration readings. Measurements will take place at weekly to biweekly intervals for the duration of the system's operational period.

The following Table 4 provides a brief synopsis of the system monitoring plan.

Table 4: SVE System Monitoring Recommendations						
Phase	Monitoring Frequency	WI	nat to Monitor	Wł	nere to Monitor	
Start-up (5 to 7 days)	Continuous	o	Flow	o	Extraction Wells	
	_	0	Vacuum	o	Manifold	
	Daily	o	Vapor Concentration	o Sta	Manifold & Exhaust ck	
	•					
Remedial (ongoing)	Continuous	o	Flow	o	Extraction Wells	
		o	Vacuum	0	Manifold	
	Weekly	o	Vapor Concentration	o Sta	Manifold & Exhaust ck	

SVE system performance monitoring is necessary to determine if remedial progress is proceeding at a reasonable rate. SVE system remedial progress typically exhibits asymptotic behavior with respect to VOC concentration reduction and cumulative VOC mass removal. The operator will evaluate alternatives for increasing mass removal rate by: (1) increasing flow to extraction wells with higher vapor concentration, or (2) periodic shutdown and startup operation of extraction wells to allow the subsurface environment to come to equilibrium and then start extracting vapors again.

Termination of operations may be appropriate if:

- 1. residual levels are at or below NYSDEC's limits;
- 2. asymptotic behavior is persistent for periods greater that about two months; and
- 3. the concentration rebound is sufficiently small following periods of temporary system shutdown.

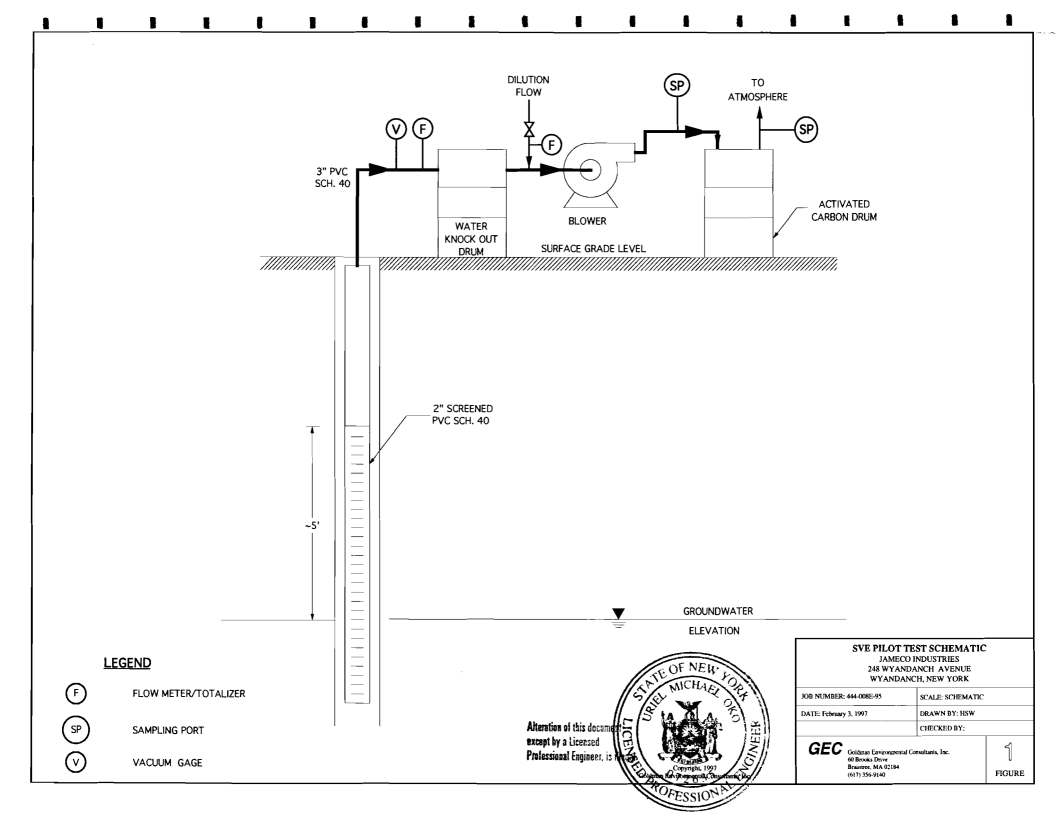
6.3 Operation and Maintenance Training

After the SVE system installation is complete, GEC will implement Operation & Maintenance (O&M) training for the continued operation of the remediation system. The O&M training will include the system components,

specifications, operations and troubleshooting, tips on how to optimize the system operation, and general routine maintenance. The training will include a sampling plan for influent and effluent air monitoring using PID, OVA and DräggerTM tubes.

The O&M manual will contain descriptions and explanations of all equipment and controls used for operation of the SVE system. Information concerning the process description and operational techniques will be included as employed during normal operations as well as in emergency situations. This manual will be have two sections: (1) for normal operation and (2) for abnormal operation, which will include emergency response procedures. This O&M manual will serve as a training manual for an inexperienced SVE system operator and will be thoroughly reviewed by all operating personnel.

The O&M manual will provide suggestions for recordkeeping, which spare parts should be stocked, and where to obtain them in the case of an emergency.



QUALITY ASSURANCE PROJECT PLAN (QAPjP) JAMECO INDUSTRIES, INC. 248 WYANDANCH, AVE WYANDANCH, NEW YORK

February 1997

Prepared For:

New York State Department of Environmental Conservation

and

Camille Gagnon Watts Industries, Inc. P.O. Box 6431 South Main Street Franklin, NH 03235

GEC

Goldman Environmental Consultants, Inc.

60 Brooks Drive Braintree, MA 02184 (617)-356-9140

QUALITY ASSURANCE PROJECT PLAN

248 WYANDANCH AVENUE WYANDANCH, NEW YORK

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1.0 Project Description

This Quality Assurance Project Plan (QAPjP) describes activities, procedures and measures that will be taken to assure quality and accuracy of analytical data relative to the Interim Remedial Measure (IRM) including the installation of a Soil Vapor Extraction (SVE) system as a means of reducing concentrations of chlorinated compounds detected beneath the Jameco facility building.

2.0 Project Organization

A Project Organization chart has been included as Figure 2 of this QAPjP. As the chart indicates, quality assurance will be the responsibility of Robert A. Fricke, GEC's Vice President of Technical Services. Mr. Fricke's resume is included with Attachment A of this Plan. GEC intends to use IEA and their Monroe, Connecticut laboratory. IEA is a New York State Department of Health (NYSDOH) Environmental Laboratory Approval Program (ELAP) certified laboratory for general and CLP analyses.

3.0 Quality Assurance Objectives (QAO) for Data Measurement

As described in the Sampling and Analysis Plan section of the attached Design Plan, sampling will be conducted during the performance of the proposed SVE system. Upon completion of the IRM, data will be collected to evaluate the completeness of the IRM and to evaluate the risk that residual contamination poses to public health safety and welfare, and the environment. All activities conducted at the site will be coordinated with a Remedial Investigation/Feasibility Study (RI/FS) that will be conducted contemporaneously with the IRM.

GEC's QA/QC Plan is attached as Attachment A to this QAPjP. In addition, IEA's Quality Assurance Plan is also included with Attachment A.

4.0 Sampling Procedure

GEC's sampling plan is described in Section 3.0 of the attached Proposed Design Plan. In addition, Standard Operating Procedures for the collection of soil and groundwater samples is included as Attachment B to this QAPjP.

5.0 Sample and Document Custody Procedures

Standard Operating Procedures for Sample and Document Custody Procedures are included at Attachment B to this QAPjP. All samples will be forwarded to IEA via overnight shipment to assure that samples are received by the laboratory within 24 hours of collection.

6.0 <u>Calibrations Procedures and Frequency</u>

Sample calibration procedures and frequency are determine by IEA with cooperation of GEC. A description of these procedures is included in Attachment A.

7.0 Analytical Procedures

Analytical procedures to be followed during the completion of this IRM are described in Section 3.0 of the Design Plan. As noted in the preceding paragraphs this IRM will be conducted in coordination with an RI/FS at the site. As such, a significant amount of information regarding site soil, groundwater and air (indoor and soil gas) will be obtained. Analytical procedures are described in IEA's documentation, attached as Attachment A to this QAPjP. Also included in Attachment A is a listing of the method reporting limits for specific analytes and laboratory analyses.

8.0 Data Reduction, Validation and Reporting

Data Reduction, Usability and Reporting procedures are described in IEA's documentation, attached as Attachment A to this QAPjP. In addition, GEC's QAO will review all data and analytical procedures to assure that data collection has been conducted in accordance with GEC's QA/QC procedures, provided in Attachment A.

9.0 <u>Internal Quality Control Checks</u>

Internal Quality Control Checks are described in IEA's documentation, included as Attachment A to this QAPjP. In addition, GEC's QAO will review all data and analytical procedures to assure that data collection has been conducted in accordance with GEC's QA/QC procedures, attached as Attachment A.

10.0 Performance and System Audits

GEC's and IEA's Performance and System Audit procedures are described in GEC's QA/QC program and in IEA's documentation, attached as Attachment A to this QAPjP. In addition, GEC's QAO audits laboratory practices and systems initially and as warranted as part of GEC's ongoing commitment to attaining data quality objectives.

11.0 Preventative Maintenance

GEC's and IEA's Preventative Maintenance procedures are described GEC's QA/QC program and in IEA's documentation, attached as Attachment A to this QAPjP. In addition, GEC's QAO audits laboratory practices and systems initially and as warranted as part of GEC's ongoing commitment to attaining data quality objectives.

12.0 Data Measurement Assessment Procedures

GEC's and IEA's Data Measurements procedures are described in GEC's for QA/QC and in IEA's documentation, attached as Attachment A to this QAPjP. In addition, GEC's QAO audits laboratory practices and systems initially and as warranted as part of GEC's ongoing commitment to attaining data quality objectives.

As part of the Data Measurement and Assessment Procedures, GEC will evaluate data suitability. Data suitability findings will be forwarded to NYSDEC prior to submission of the final IRM report.

13.0 **Corrective Action**

GEC's and IEA's Corrective Action procedures are described in GEC's for QA/QC and in IEA's documentation, attached as Attachment A to this QAPiP.

Quality Assurance Reports and Management 14.0

Upon completion of site activities, GEC documents all potential quality assurance concerns as a means of reporting and managing quality assurance. Items included in this reporting might include a statistical analysis of suspect data, presence of analytes in method or trip blanks, or the unexplained absence of known contaminants. Information of this nature is included in IRM and RI/FS Progress Reports.

Respectfully submitted, Goldman Environmental Consultants, Inc.

Prepared By:

Samuel W. Butcher

Senior Project Manager

Reviewed By:

Robert A. Fricke

Quality Assurance Officer

Vice President, Technical Services

that for Robert Fricks

<u>QAPjP</u> ATTACHMENT A

GEC QA/QC and IEA QA/QC

Quality Assurance/Quality Control (QA/QC)

I. Purpose

The purpose of the GEC QA/QC program is to generate analytical data that is of known and defensible quality. These procedures apply to all projects in which sampling is involved. QA/QC from one project is not transferable to another.

II. Decontamination

- 1) Decontamination should be performed on all reusable field sampling equipment and protective gear. Sampling equipment should be decontaminated before the collection of a sample and after sampling has been completed. Protective gear should be decontaminated after the collection of a sample.
- 2) It is necessary to use the following decontamination solutions in the field:
 - Non-phosphate detergent plus tap water wash.
 - Distilled/ deionized water rinse.
 - 10% Nitric Acid rinse.*
 - Distilled/ deionized water rinse.*
 - Methanol rinse, when sampling volatiles only.
 - Pesticide grade methanol then hexane rinse.**
 - Distilled / deionized water rinse. **
 - * Only if sample is to be analyzed for metals.
 - ** Only if sample is to be analyzed for semi-volatile organics, PCBs or pesticides.
- 3) Sample bottles and sampling equipment should not be stored near gasoline, solvents, or other potential sources of contamination. If unavoidable bottles and equipment should be sealed in containers or plastic.
- 4) Heavy equipment, including hand tools, should be cleaned by steam cleaning or manual scrubbing prior and subsequent to use in hazardous waste investigations.

III. Measures or Quality Control/Quality Assurance

1. Trip Blanks

- Trip blanks are used in order to detect additional sources of contamination that might affect analytical results. The following are potential sources of additional contamination:
 - a. Sample containers;
 - b. Contamination during shipment to and from the site;
 - c. Ambient air contact with analytical instrumentation at the laboratory during analysis; or
 - d. Laboratory reagent used in analytical procedures.
- One trip blank is required for every set of volatile organic compound samples sent to the lab, regardless of job size. If, however, VOCs are not a parameter of the sampling round, the laboratory is consulted as to which parameter should have an associated trip blank.
- Trip blanks are kept with containers used in the sampling round at all times. More specifically, they should accompany the site specific sampling containers from the time the containers leave the laboratory until they are returned for analysis.
- Obtain containers and trip blanks prepared specifically for each job from the laboratory. Return unused containers to the laboratory upon completion of a project.

2. Field Blanks

- Field blanks are used to indicate potential contamination contracted from ambient air or from sampling equipment. It also serves as a QA/QC for decontamination procedures.
- Collect one (1) set of field blanks for every 20 samples per project. It is not necessary to take a field blank for jobs in which less than 10 samples are collected.

- Procedure

- a. Collect two sets of sample containers to cover all sampling parameters. One set will be full of analyte free water (obtain extra analyte free water to fill two VOA vials). The other set is empty.
- b. Go to the most contaminated area and run the water from the full containers, through the decontaminated sampling equipment and into the associated empty containers.

- c. Send to the lab for analysis.
- Use containers and field blanks prepared specifically for job.

3. Duplicate Samples

- Duplicate samples are collected in order to serve as a laboratory check. Therefore, it is important that the lab does not know which samples are to serve for this purpose.

- Frequency

- a. Obtain one (1) duplicate sample for every 10 samples of each matrix. If less than ten samples are collected of a given matrix, a duplicate must be collected anyway.
- b. If a total of less than 10 samples are collected, collect one (1) duplicate of the majority medium.
- c. If a total of less than five (5) samples are collected, it is not necessary to collect a duplicate sample.
- * Note that the frequency as outlined here pertains to the number of samples collected per project, not per location of a given project.

- Procedures

The idea behind the duplicate sample is to collect two samples as close to identical as possible.

a. For water:

Alternately fill containers for the same parameter with equal amounts of liquid per bailer. Fill duplicate VOC vials from the same bailer of liquid.

b. For soil:

- -- VOC samples must be taken from the discreet sampling locations.
- -- For all other samples, mix the applicable soil in a decontaminated stainless steel or polyethylene bowl or tray. Then fill sample containers with the soil mix.
- -- When confronted with the option of collecting a water sample or a soil sample, choose the water sample.

- Labeling for the laboratory

a. Label the containers normally and give the duplicate samples

different reference numbers.

- b. Indicate the quantity of duplicates in the "special instructions" or "remarks" portion of the chain of custody and laboratory services sheet, however, do not indicate the reference numbers of the duplicates.
- c. Upon receipt of analytical results, contact the laboratory and convey all data pertaining to the duplicates for their QA/QC.

4. Background samples

- Background samples are taken only if it is required for comparison of site conditions to the surrounding environment. This is to be dictated by client needs on a site to site basis.
- 5. Performance Evaluation Samples
- The project manager should consider the use of the following performance evaluation samples on a periodic basis. Typically, these will be reserved for larger jobs:
 - a. Laboratory performance evaluation samples
 - -- Collect duplicate samples and send to two different laboratories for comparison. Avoid using soil samples for this procedure.
 - -- Send a sample of known quantity and quality to the laboratory in order to determine laboratory performance. Such samples can be prepared by any laboratory.
 - b. Gas chromatograph (GC) performance evaluation samples
 - -- Acquire a sample of known quantity and quality from a laboratory. Analyze the sample with the gas chromatograph in order to determine the integrity of GC results.

IV. Field Sampling QA/QC

- 1) When sampling a well, collect VOA samples first and Oil & Grease samples last.
- 2) Start sampling at the presumed least contaminated areas, proceeding to the more contaminated areas.
- 3) Preservatives

- Consult the laboratory in order to determine which sampling parameters require preservatives. The laboratory will provide sampling containers specific for each job.
- It is necessary to fill the sample container when using preserved bottles; preservative is added with this assumption
- If samples are not collected correctly, they will not pass GEC QA/QC.
- 4) A chain-of-custody must accompany each set of samples from the job site to the laboratory. Be sure to identify the presence of trip blanks on the chain-of-custody sheets.
- 5) If possible, use the numbering system outlined on the attached sheet for identifying samples.

V. Ordering Sample Containers

- 1) Pre-plan sampling strategy to determine the sample parameters, the number of sample points including QA/QC samples, and the matrix of the given sample points.
- 2) Call laboratory and tell them:
 - Sample parameters;
 - Number of samples to be collected;
 - The number of container sets needed for trip blanks, field blanks, and duplicates; and
 - The matrix of each sample to be collected.
- 3) Sample containers should be ordered specifically for each job. Any sample containers unused at the end of the job should be sent back to the laboratory.

VI. Conclusions

- 1) Pre-planning is crucial.
- 2) Keep open communication with the laboratory on all matters.
- 3) If you make a mistake in sampling collection, accept it, and retake the necessary samples.

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Lawrence M. Goldman, LSP President

Education

University of Massachusetts (Lowell) M.S. Engineering and Environmental Studies 1983

Lowell Technological Institute B.S., Textile Technology 1968

Relevant Training/Professional Registration

Princeton Course -Groundwater Pollution and Hydrology

NGWA - Risk Assessment for Environmental Professionals

Brooking Institute
Environmental Quality and
Natural Resources (80 Hours)

EPA technical course training in air pollution control, wastewater treatment, hazardous and solid waste management

Professional Registration Licensed Site Professional (MA #1509)

OSHA Hazardous Waste Site Worker Training (40 Hours)

Business/Regulatory Affiliations

Smaller Business Association of New England - Chairman, Environmental Issues Committee

Member, MDEP Licensed Site Professional Regulatory Development Committee

Member, Licensed Site Professional Association

Associated Industries of Mass Energy and Environmental Committee

American Consulting Engineering Council of N.E.

Regulatory Compliance

- Lawrence Goldman has over 22 years of experience administering and managing multi-media (air, water, and hazardous waste) environmental programs for industry and government.
- As Director of Enforcement, EPA Region I, administered a staff of 75 in support of permitting activities, compliance monitoring, and regulatory enforcement in concert with the New England States.
- Developed Region I uncontrolled sites and Superfund enforcement programs (1978-1981). Implemented cleanup programs at numerous NPL Sites throughout New England.
- Design and implementation of multi-media environmental audit programs for manufacturing/industrial clients throughout the United States.
- Expert witness for CERCLA and state cost recovery actions.
 Licensed Site Professional (MA) and knowledgeable of MGL Chapter 21E and the MCP.
- Provides technical and regulatory support to law firms, Fortune 500 companies, as well as small to medium sized manufacturers and businesses.

Hazardous Waste Management/Assessment

- Administers staff of engineers, hydrogeologists, toxicologists and support personnel in conducting walk through and subsurface investigations at an estimated 750-1,000 sites throughout the Northeast since 1985.
- Developed assessment, containment and remediation programs for numerous complex sites under MGL Chapter 21E and the MCP. Rendered LSP opinions, implemented risk reduction measures including risk characterization.
- Developed decision-making guidelines for an industrial chemical manufacturer at a nationally ranked Superfund Site, focusing upon lagoon and landfill closure and aquifer restoration.
- Developed RCRA Part B applications for storage, treatment and incineration of hazardous waste for major defense contractors.
- Developed the remedial alternatives and regulatory analysis necessary to control a 60-mile stretch of PCB-laden river sediment for a Fortune 500 client under an EPA/State enforcement order.
- Negotiated the regulatory and technical requirements with EPA Region I to stabilize a former 240 acre munitions site under the Immediate Removal Action provisions of CERCLA.

List of Publications available upon request.

Gary W. Siegel, P.E., L.S.P. Vice-President, Environmental Engineering

Education

Northeastern University M.S., Civil Engineering 1983

Specialization in Water and Wastewater Treatment

Brown University B.S., Engineering 1979

Additional Training

U.S. EPA
OSHA Hazardous Waste Site
Worker Training (40 hours)

Toxic Use Reduction Institute TUR Planners Course

Executive Enterprises
Groundwater Management and
Remediation

Certifications

Registered Professional Engineer
Massachusetts/Maine/Rhode
Island

Registered Licensed Site Professional Mass., No. 1523

Previous Employers

IBM Corporation Poughkeepsie, NY 1979 - 1980

USEPA Region I Boston, MA 1980 - 1982

W. R. Grace & Co. Lexington, MA 1982- 1988

Memberships

American Society of Civil Engineers

American Consulting Engineers Council

Environmental Engineering

- Gary Siegel has over 16 years of experience managing hazardous waste remediation projects, environmental engineering projects, and regulatory compliance issues for industry, and government.
- Manages all of GEC's remedial response actions and remediation design projects.
- Project Manager for the assessment, remediation, risk characterization and issuance of RAO's or LSP Evaluation Opinions for more than of 30 industrial, commercial and residential sites.

Major Site Assessment and Remediation Experience

- Project Manager and Design Engineer on implementing an innovative Short Term Measure to halt a continuing oil leak through a seawall in a tidal zone. Design utilized tidal action against a slurry wall to passively collect floating product.
- LSP and Project Manager for a site and risk characterization for a 200 year old mill property. Site issues involved Tier 1 Permit Application, a Method 3 risk assessment of over 50 individual contaminants and the development of a limited AUL leading to a Class B-2 RAO.
- Principal Engineer for GEC's preparation of Pollution Prevention/ Resource Conservation (P2/RC) Plans for Massport and the Massachusetts Turnpike Authority under the Clean State Initiative.
- LSP and Project manager of an RAM involving the decontamination of PCBs from two transformer vaults providing power to a large manufacturing operation.
- Project engineer for the excavation and disposal of a 32,000 cubic yard industrial waste landfill. Project involved coordinating over 1200 dumptrailers, excavation methods, site investigations and government reporting.
- LSP and Project manager for the assessment and remediation of 10 Mass. Highway Department sites located on Cape Cod, New Bedford and Martha's Vineyard.
- LSP and Project Manager for two separate disposal sites at a 100 year old manufacturing facility. Work involved utility related remediation, a limited RAM in the area of an abandoned coal bunker and developing a methodology to avoid further excavation by reclassifying the site's groundwater.
- City of Everett LSP on oversite of Monsanto's remediation and redevelopment of the closed chemical manufacturing facility in Everett. Responsible for regular site inspection, review and comment on Monsanto MCP submissions

Samuel W. Butcher Manager, Hydrogeology and Remedial Programs

Education

Brown University, M.Sc. Geology, 1988

Skidmore College B.A. Geology, 1986

Certifications

Certified Ground Water Professional (CGWP), National Ground Water Assn. (AGWSE)

Professional Registration Licensed Site Professional (MA #9185)

Additional Training

Tufts University
Post-graduate study - Hydrology

MODFLOW (USGS Modular Flow Model) for Simulation of Ground Water and Advective Transport, NGWA

Visual MODFLOW, NGWA

OSHA Hazardous Waste Site Worker Training (40 Hours)

OSHA Waste Site Supervisor Training (8 Hours)

Professional Affiliations

National Ground Water Assn.

Geological Society of America

Seminars

University of Massachusetts
Contaminated Soil Remediation

American Petroleum Institute Assessment, Control and Remediation of LNAPL Contaminated Sites

Senior Project Manager / Senior Hydrogeologist

- Samuel Butcher has over seven years of experience in environmental site investigation and assessment. He manages projects involving complex geologic and/or hydrologic conditions. Specializes in subsurface investigations associated with remedial response activities and site investigations. Also manages investigation, assessment and closure of underground and above ground storage tanks.
- Manages projects to determine subsurface soil and bedrock as well as aquifer and contaminant transport characteristics. Also responsible for ground water modeling projects.
- Conducted numerous site assessments at industrial, commercial, and military facilities throughout New England. Studies included geologic and hydrologic assessments, hazardous waste evaluations, underground storage tank removals, installations and management plans as well as geophysical investigations and emergency response activities.

Recent Work

- Initial Site Assessment, plume delineation, potential source identification and Risk Characterization of multi-media, multicontaminant releases at a former manufacturing facility. Assessment included lithologic and hydrologic characterization of substrata, decommissioning of USTs, characterization of potential off-site sources and closure of facility
- Field testing, via pump tests and slug tests, to determine hydraulic conductivity of several hazardous waste disposal sites. Incorporation of field test results into design of remedial systems and construction of 3-dimensional groundwater models.
- Initial Site Assessment, source identification and Risk Characterization of releases of chlorinated compounds to subsurface soil and ground water. Included extensive characterization and assessment of subsurface soil and hydrology to determine extent of migration and migration pathways of freephase chlorinated compounds.
- Supervision and oversight of underground storage tank and soil removal of approximately 25 tanks at military facilities in New England. Design of report format which is specifically requested in tank removal "Request for Bids".

PUBLICATIONS

- 1. Butcher S.W., 1989 "The Nickpoint Concept and Its Implications Regarding Onlap to the Stratigraphic Record" in Cross T.A. Quantitative Dynamic Stratigraphy, Prentice Hall, p. 375-385.
- 2. Butcher S.W., "Eight questions you should ask before buying a property with USTs" New England Real Estate Journal, November 24, 1989, p.22-24
- 3. Butcher S.W., 1993, "Recommended Approach for Subsurface Investigation and Model Development Case History" Northeastern Geology v. 15, No. 2, p. 176-177

Harry S. Wahra Mechanical Engineer

Education

State University of New York
B.S., Mechanical Engineering 1987

Additional Training

OSHA Hazardous Waste Site Worker Training (40 Hours)

OSHA Hazardous Waste Site Worker Refreshers (8 Hours)

OSHA Waste Site Supervisor Training (8 Hours)

Certified Novell Netware System Manager

Seminars and Classes

University of Toledo/Division of Continuing Education Industrial Wastewater Pretreatment Seminar, 1995.

University of Lowell/Division of Continuing Education Engineer in Training Seminar

Mechanical Engineer

- Eight years of experience providing a multitude of mechanical process engineering services in the environmental consulting industry.
- Designed, managed startup and installation activities of Remediation system for petroleum product recovery. Prepared Engineer's Report and Operation & Maintenance manual for the system. Conducted monthly visits to the site to ensure proper operation.
- Prepared work plan for an Interim Remedial Measure (IRM) using Soil Vapor Extraction (SVE) system for solvent recovery from soil.
 Responsible for SVE pilot test to collect data for sizing and selecting of a production scale SVE system.
- Assisted in the design of Air Stripping System to remove solvents from the groundwater and also provided assistance on system installation and preparation of O&M manual. Regularly conducted site visits to ensure the proper operation.
- Designed, supervised and managed startup and installation activities of Wastewater Treatment Systems which includes Ion Exchange Systems, Ultra filtration Systems, Reverse Osmosis Systems, and hydroxide precipitation systems for a variety of manufacturing facilities. Prepared Cost Estimates and bills of material for these systems.
- Prepared Engineer's Report, Operation and Maintenance manual, and Staffing Plan for Wastewater Treatment Systems.
- Designed and specified customized packaged HVAC, Local Exhaust Ventilation (LEV) and Air Pollution Control (APC) systems. Conducted meteorological dispersion models and BACT studies for appropriate design and specifications of APC systems. Provided management, supervision, and troubleshooting guidance for these systems during installation.
- Engaged in regulatory permitting and liaison activities. Completed various permits which include the following:
 - (1) Title V Operating Permit Application;
 - (2) DEP Air Plans Applications;
 - (3) Industrial Wastewater Discharge permit;
 - (4) DEP Sewer Connection permit;
 - (5) Air Source Registration;
 - (6) SARA/TURA Reporting; and
 - (7) DEP Cross-connection permits.
- Responsible for Water Distribution system design, planning and installation required by state and local cross-connection regulations.
- Provided classroom instructions to WWT operators regarding mechanical equipment operation and maintenance procedures as part of WWT certification course for plant operators.
- Assisted in implementing wastewater and groundwater sampling program for regulatory monitoring requirements.

Paul T. Bartlett, P.E. Vice President and Manager of Air Programs

Education

N.C. State University
B.S. Mechanical Engineering, 1966

Boston University
Courses in M.S. Administration
Program

Relevant Training/Professional Registration

Registered Professional Engineer, Connecticut

EPA technical course training in air pollution control and combustion

Clark University, COPACE, Worcester, MA- Designed and taught an introductory course in Environmental Engineering

Membership

Air and Waste Management Association

Regulatory Compliance

- Paul Bartlett has over 25 years of technical, project management, and business development experience with multi-media (air, water, and hazardous waste) environmental programs for industry and government.
- Regional Engineer for Central Region Office, Air Quality Division, North Carolina's Dept. of Natural and Economic Resources where he directed a staff of engineers, scientists, and inspectors conducting source registrations, emission audits, permit negotiations, and public relations.
- Presently managing an air emissions compliance project for a coater who is a major source of VOCs and HAPs. Tasks include preparing source registration form and a comprehensive permit application, SARA Title III and MA TURA reporting, and negotiations with MADEP.
- Managed a cogeneration feasibility study to remove a foundry and manufacturing facility from the electrical grid. The study included developing a conceptual permit application that responded to emerging NOX RACT requirements under the Clean Air Act Amendments.
- Design and implementation of multi-media environmental audit and compliance programs for manufacturing/industrial clients throughout the United States, including due diligence audits for Thermo Electron Corporation during acquisitions.
- Site investigations of 30 industries in Connecticut and Oregon to determine O&M practices and resulting air emissions from normal operations and upset conditions.

Environmental Engineering

- He is experienced in the design, installation, startup, and performance/compliance testing of industrial ventilation, air pollution control, and industrial wastewater treatment systems.
- Specialist in separations technology for pollution prevention, process recycling, toxics use and waste minimization, and site remediation applications.
- He was the Vice President of Engineering during the startup phase of a company introducing a Heat Pump Evaporator (HPE) for process recycling and wastewater treatment. No CFCs were used in the refrigeration cycle and processing was under a vacuum to eliminate emissions to the atmosphere.
- Air quality/emissions evaluation and process redesign to reduce HAPs from a mercury processing mill. Modified solids conveyance and enclosure and redesigned induced draft fans for incineration.
- Process engineering assessment to reduce HAPs from a chlorinated dry bleach plant. Work included modifying calciner's end seals and operating temperature of NH3 burner and improving O&M practices.

Robert A. Fricke Vice-President of Technical Programs

Education

Florida Institute of Technology M.S., Chemical Oceanography 1981

Specialization in laboratory analysis of organic contaminants

Bloomsburg State College B.S., Biology 1978

Additional Training

OSHA Hazardous Waste Site Worker Training (40 hours)

Tufts University
Asbestos Management Planner training (40 hours)

Tufts University
Asbestos Management Planner
refresher (8 hours)

Portable Gas Chromatograph Training Course (16 hours)

Affiliations

American Chemical Society

Current Responsibilities

 Robert Fricke is Vice President of Technical Services with 14 years of experience in environmental consulting for manufacturing clients. He currently manages a regulatory compliance group, he coordinates project activities for his group and acts as a liaison with national clients. Mr. Fricke is also responsible for developing marketing opportunities within the printing, photoprocessing, and woodfinishing industries.

Previous Superfund Experience

- Health and safety officer at private, large-scale field investigations and 13 Superfund Sites across the nation.
- Developed and taught EPA 24 and 40 hour field instrumentation training courses for two years.
- Performed numerous field investigations, involving the collection and laboratory analysis of contaminants in air, soil, surface water, and groundwater at landfills, and commercial and industrial sites.

Health and Safety Planning

 Corporate Safety Director for seven years implementing GEC's health and safety program including: coordinating a medical surveillance program; approving health and safety plans required by OSHA; selection of appropriate personal protective equipment; instruction of staff on use of field equipment.

Recent Work

- Authored and managed the preparation of six Part B applications for storage, treatment, and disposal of hazardous wastes, including the preparation and submittal of a Part B Application for an incinerator.
- Currently coordinates and directs in-house staff on environmental and workplace compliance tasks for five national printing and publishing firms with 72 printing plants in 30 states. Has conducted EPA and OSHA audits at over 70 newsprint production plants around the country.
- Has managed and conducted environmental and workplace safety compliance projects in Canada and Europe, local and international clients. The projects ranged from responding to basic compliance issues, regulatory research, identification of local permit and enforcement policies, and regulatory applicability to new industrial equipment. European target countries included England and Germany.
- Conducted over 50 regulatory audits for clients with the following industries: electroplating, wood products, specialty chemicals/plastics; ink and paint manufacturing; biotechnology inorganic chemical processing; general manufacturing; information storage media; and defense equipment manufacturing. The audits included environmental and/or workplace safety issues, employee training, and relevant compliance documentations.

Robert A. Fricke Vice-President of Technical Programs

PUBLICATIONS & PRESENTATIONS (partial listing)

- "A Novel Approach to Reducing the Hazards of Buried Tank and Drum Excavation: Case History", Fricke, R., and D. Dahlstrom, American Chemical Society National Meeting, September, 1982.
- 2. "Sampling and Preparation of Soil Homogenates for Interlaboratory Comparison Of Analysis of 2,3,7,8-TCDD and Surrogate Compounds", Spatola, J., Beckett, W., Schoengold, D., Fricke, R., and J. Schwartz, EPA research publication.
- 3. "Strategies for Maintaining EPA/OSHA Compliance", presented at the 1991 America East Newspaper Operations Conference, Harrisburg, Pennsylvania.
- 4. "Strategies for Maintaining OSHA Compliance", presentation at the 1991 Newspaper Operations of America Conference, Newport, Rhode Island.
- 5. "Evaluation of Chemical Products in the Newsprint Industry", presented at the 1991 America East Newspaper Operations Conference, Harrisburg, Pennsylvania.
- 6. "Liability Issues Related to Printing Plant Audits", 1992 Printing Industries of New England Environmental Expo, Milford, Massachusetts.
- 7. "Negotiating with Regulatory Agencies", presentation at the 1992 Newspaper Association of America, St. Petersburg, Florida.
- 8. "EPA/OSHA Regulatory Review", presentation at the 1993 Newspaper Operations Association, Danvers, Massachusetts.
- 9. "Personal Protective Equipment", presentation at the 1994 Printing Industries of New England Environmental Expo, Marlboro, Massachusetts.
- 10. "Green Technology in the Printing Industry", presentation at the 1994 EcoExpo, World Trade Center, Boston, Massachusetts.

Eileen A. Furlong Manager, Site Assessments

Education

Michigan State University
M.S., Limnology Option,
Department of Fisheries and
Wildlife, 1982

Southeastern Massachusetts University, B.S., Marine Biology, 1978

Additional Training

Agency for Toxic Substances and Disease Registry Health Assessment Training Workshop

Harvard Educational and Resource Center, Hazardous Substance Training Program, Risk Communication and the News Media Workshop

Harvard School of Public Health, Harvard Center for Risk Analysis, Analyzing Risk: Science, Assessment and Management

General Sciences Corporation, Soil and Groundwater Modeling for Risk Assessment and Soil Clean-up Level Evaluation

EPA Hazardous Waste Site Worker Training (40 hours)

OSHA Hazardous Waste Site Supervisor Training Course

Memberships

National Society for Risk Analysis

Society for Risk Analysis - New England Chapter

Boston Bar Association, Environmental Law, Adjunct Member

Community

Randolph Conservation Commission

Manager, Site Assessments

- Eileen Furlong is the Senior Toxicologist and Manager of the Site Assessment Group. She has 5 years of experience characterizing human health risks from multi-media contaminants at hazardous waste disposal sites. She has extensive experience conducting health assessments (environmental fate, human exposure, and toxicology) for Superfund sites.
- Manages and directs personnel involved in Preliminary Site Assessments, Release Notification, assessments for LSP Evaluation Opinions, Numerical Ranking Systems Scoring, Tier classification, Tier 1 Permit Application, Remedial Abatement Measure, Immediate Response Action, Phase I and Phase II subsurface investigations, relative to MCP and MGL Chapter 21 E.
- Prepares Scope of Work and Cost Estimates for site investigations. Prepares and reviews final MCP response action documents.

Previous Work

- Massachusetts Department of Public Health. Conducted health assessments for Superfund sites per a cooperative agreement with the Agency for Toxic Substances and Disease Registry. Technical advisor to the Woburn Environment and Birth Study; represented the DPH at interagency and community meetings pertaining to Superfund sites. Responded to citizen concerns regarding environmental health issues.
- Department of Cancer Pharmacology, Dana-Farber Cancer Institute. Performed pharmacokinetic studies of alkylating agents in humans and animals; developed microanalytical techniques involving gas chromatography and high pressure liquid chromatography for alkylating agents and DNA damage products.

Recent Work

- Managed and implemented a Phase II-Comprehensive Site Assessment under MCP, as revised, for a petroleum release site. Phase II investigation included a Method 2 Risk Characterization consisting of 25 contaminants, a potentially productive aquifer, and a Method 3 Stage I Environmental Screening.
- Managed MCP response actions for an industrial site. Site had 52 contaminants and is located in an Interim Wellhead Protection Area and Area of Critical Environmental Concern. Conducted Numerical Rank System scoring for the site. Planned and directed the field investigation to support the Risk Characterization. Directed a Method 3 Risk Characterization, and conducted a Method 3 Stage I Environmental Screening. Site qualified for a Class B-2 Response Action Outcome, with an Activity and Use Limitation.

- Directed tank removal and MCP response action activities for an industrial site. Response actions included: 72-hour notification of release with verbal approval for an Immediate Response Action; implementation of IRA and preparation of IRA Completion Report; subsurface investigation; documentation of coal/coal ash as source of background; risk characterization; and preparation of a Class A-2 RAO.
- Provided litigation support for a lawsuit involving contamination at a former circuit board research and development site.
- Investigated the industrial toxicology of propylene glycol ethers and methylene chloride used in the printing facilities of a national newspaper syndicate.

PUBLICATIONS (partial listing)

- 1. "Consideration of Non-Routinely Monitored Hazardous Substances for the Health Assessment", Proceedings of the HMCRI Superfund 90 Conference (pp. 144-147). Furlong, E.A., Barry, T.A., and Condon, S.K., 1990.
- "Health Assessment for Baird and McGuire, Norfolk County, Holbrook, Massachusetts", Agency for Toxic Substances and Disease Registry and Massachusetts Department of Public Health. Furlong, E.A., Barry, T.A., and Condon, S.K., 1990.
- 3. "Role of Environmental Fate and Transport Data in Health Assessments: A Case Study", Proceedings of the HMCRI Superfund 90 Conference (pp. 128-132). Ulirsch, G., Hayes, L., and Furlong, E.A., 1990.
- 4. "Health Assessment for Wells G and H, Woburn, Massachusetts", Agency for Toxic Substances and Disease Registry and Massachusetts Department of Public Health. Furlong, E.A., Condon, S.K., DiSirio, M., and House, L., 1989.
- 5. "Addendum to Health Assessment for Sullivan's Ledge, New Bedford, Massachusetts", Agency to Toxic Substances and Disease Registry and Massachusetts Department of Public Health. Furlong, E.A., Condon, S.K., DiSirio, M., House, L., 1989.
- 6. "Health Assessment for Sullivan's Ledge, New Bedford, Massachusetts", 1 bid., 1988.
- 7. "Production of dihydrothymidine stereoisomers in DNA by gamma irradiation", Biochemistry 25:4344-4349, Furlong, E.A., Jorgensen, T.J., Henner, W.D., 1986.
- 8. "Trihalomethane levels in chlorinated Michigan drinking water", Ecological Modelling 32:215-225, Furlong, E.A., D'Itri, F.M., 1986.

Daniel N. Kirichok Chemical Engineer

Education

Northeastern University (Boston) B.S. in Chemical Engineering 1994

Certifications

Fundamentals of Engineering (Formerly E.I.T. Exam) 1994

Buisness/Regulatory Affiliations

Associate, American Institute of Chemical Engineers

Member: Order of the Engineer

Air & Waste Management Association

Member: AESF

 Daniel Kirichok is a chemical engineer with GEC, specializing in air regulatory, permitting and control issues.

Regulatory Compliance

- Over 3 years of experience with assessing regulatory requirements and environmental impacts for clients in general industry.
- Strong working knowledge of EPA, MDEP air regulations and state interpretation of these regulations.
- Completed several air permit applications for industry, particularly within the Printing and Metal Products Industry.
- Developed Title V operating permits for industrial clients

Materials Use Reduction/Waste Treatment

- Conducted review of feasible treatment technologies for removal of mercury and other metals from wastewater.
- Conducted preliminary design of wastewater piping system for client laboratory system. Currently evaluating treatment options for removal of mercury.
- Conducted Massachusetts Toxics Use Reduction option investigations for clients including two metals finishing operations, utilities companies, and fabric coating operations.
- Design of Air Pollution Collection/PMID removal system
- Designed wastewater handling system
- Helped to develop/select treatment strategies for wastewater mercury reduction/removal

Health & Safety Regulations

- Assisted in creating hazard communications plan and SOP's for major computer component manufacturer.
- Historical file search for facility site assesments

Current Responsibilities

 Develop and implement plans, reports and permit applications for industrial clients:

NPDES General Discharge; Sewer Connection/Extension; SARA Title III & Right-to-Know; Air Emission Permits; BACT evaluations; source registration; TURA reports and evaluations; OSHA hazard communications.

Education

Northeastern University (Boston) B.S. in Civil Engineering 1995

Certifications

OSHA Hazardous Waste Site Worker Training (40 hours)

Business/Regulatory Affiliations

Member, American Society of Civil Engineers

Member, Order of the Engineer

Jason Mauro is an environmental engineer with GEC with a background in the petroleum industry and strong experience in remediation of petroleum product releases.

Remediation Experience

- Strong understanding of remedial systems, including, but not limited to, soil vapor extraction, groundwater pump and treat, and sparging systems.
- Involved in preliminary design of Soil Vapor Extraction and Groundwater Pump & Treat systems for the remediation of petroleum impacted media. Provided field management during the installation and start up of these systems, as well as, assistance in the construction of these systems. Construction activities included well installation, connection to wells, trenching, installation of necessary piping, remediation equipment installation (air strippers, blowers, carbon vessels, catalytic oxidizers), shed construction, and associated electrical work.
- Developed Operation and Maintenance manuals for SVE and Groundwater Pump & Treat systems.
- Conducted monthly site visits to perform system maintenance including troubleshooting, making necessary adjustments to the system in order to maintain efficiency, and repair system when necessary. Repairs included replacing valves, rerouting of air stacks, cleaning out iron clogged air strippers, repairing faulty pumps, repairing damaged wells, and any associated electrical work.

Environmental Engineering

- Developed spreadsheets to track SVE system efficiency, including mass removal(s), and fuel consumption.
- Field experience in groundwater and soil sampling and well monitoring.
- Field engineer for the removal and installation of underground storage tanks. Designed and oversaw the installation of dewatering activities for projects taking place in coastal or high water table areas. Dewatering activities included the installation of tongue and groove sheet piles and whalers, pumps to remove the water, temporary treatment systems for contaminated water, permitting required to reinject water back into the ground via reinjection wells or to dispose of water via municipal storm sewer.

Field engineer for EPA retrofits to existing underground storage tanks. Retrofits included the installation of spill containment measures around fill pipes, sump pumps, and product dispensers. Also included was the installation and start up of modern leak detection equipment.

Project Manager for the demolition and rebuilding of a gasoline service station. Demolition activities included asbestos removal, the removal of USTs, associated piping, dispensers and hydraulic lifts in repair area. Responsible for the coordination of contaminated soil removal. Oversaw dewatering activities and temporary pump and treat system. Field engineer for installation of new building footings and pre-fabricated building. Responsible for the installation of new USTs, piping, and product dispensers. Field Engineer for the start up of UST leak detection system and construction punch list items.

Current Responsibilities

- Responsibilities include conducting environmental site assessments in support of MGL Chapter 21E and subsurface investigations.
- Performing field work, including groundwater and soil sampling for laboratory analysis, well monitoring, and site remediation.

Parrish C. Smolcha Environmental Toxicologist

Education

Union Collate B.S., Biology 1994

SUNY College of Environmental Science and Forestry Continuing Ed., Organic Chemistry

Additional Training

OSHA Hazardous Waste Site Worker Training [Title 29 CFR 1910.120(e)(8)]

Hazard Communication Training [Title 29 CFR 1910.1200]

American Red Cross - Adult CPR; Standard First Aid

Memberships

Sigma Xi National Research Honor Society, Associate Member

Environmental Toxicologist

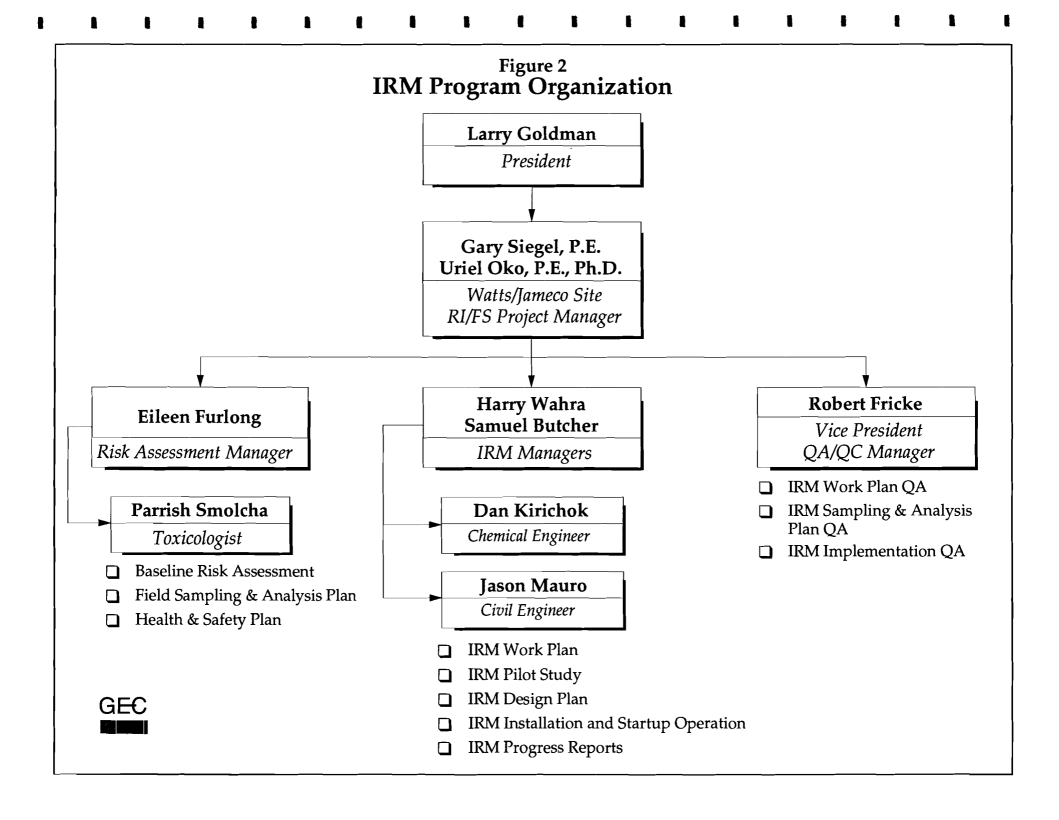
 Parrish Smolcha has over 2 years of experience performing Risk Assessments (RAs) under the Massachusetts Contingency Plan (MCP) and in conjunction with Remedial Investigations/Feasibility Studies (RI/FS) under CERCLA/SARA. These RAs include data evaluation; contamination assessment; exposure assessment for soil, groundwater, and air; environmental fate and transport assessment; toxicity assessment; and risk characterization.

Recent Work

- At a former industrial facility in eastern Massachusetts, assisted in the preparation of a Method 2 risk characterization under the MCP. Identified human and environmental receptors, exposure points and pathways, exposure point concentrations, and site soil and groundwater categories. Developed Method 2 soil and groundwater standards for several site-related constituents using MADEP protocol.
- At a junior high school in southeastern Massachusetts, conducted a Method 1 risk characterization under the MCP resulting in a Class B-2 Response Action Outcome Statement.
- At a newspaper printing and publishing facility in southeastern Massachusetts, conducted a Method 1 risk characterization under the MCP resulting in a Class A-2 Response Action Outcome Statement.
- At a Superfund site in eastern Massachusetts, assisted in the preparation of a Method 2 risk characterization under the MCP. Identified potential human and environmental receptors and developed Method 2 soil standards for several site-related constituents.

Previous Work

- At an industrial site in eastern Massachusetts, assisted in the preparation of a combined Method 2 / Method 3 risk characterization resulting in a Class A-2 Response Action Outcome Statement.
- At a RCRA facility in western Massachusetts, assisted in the development of a background sampling plan for soil and groundwater. The plan was consistent with MCP and Hazardous and Solid Waste Amendments (1984) protocols.
- Following MCP guidance, prepared a target constituent analysis for sediment and floodplain soil along a major river system in western Massachusetts. The analysis was used to identify "target" constituents present at concentrations that warranted further downstream sampling.
- Prepared a baseline RA for a major New York State utility in support of the conversion of a warehouse into a child daycare facility. Developed hypothetical future exposure scenarios for daycare children and adult workers and recommended remedial measures to mitigate potential exposure to lead-based paint.



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IEA-CT Laboratory Quality Assurance Program

prepared by

Marsha K. Culik Quality Assurance Manager

for

IEA Corporation
Monroe, Connecticut

This document has been prepared by IEA Corporation and will be updated annually. The material contained herein is not to be disclosed to, or made available to any third party without the prior expressed written approval of the Corporate Quality Assurance Director.

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1.0 QUALITY ASSURANCE PROGRAM-IDENTIFICATION FORM

Document Title:

IEA-CONNECTICUT QUALITY ASSURANCE PROGRAM PLAN

Corporate Address:

IEA Connecticut 200 Monroe Turnpike Monroe, Connecticut 06468

Company Official:

Title: Telephone: Mr. Michael V. Bonomo Director of Operations

(203) 261-4458

Company Official:

Title:

Mr. Jeffrey C. Curran Laboratory Manager

Company Official:

Title:

Ms. Marsha K. Culik

Quality Assurance Manager

Plan Coverage:

IEA-Connecticut Laboratory including the following functions:

Administration Sample Receipt GC Laboratories Quality Assurance

Data Entry Report Production Computer Systems Inorganics Laboratories GC/MS Laboratories Facilities and Safety

Sample Preparation Laboratories

Concurrences:

Name: Title:

Mr. Michael Bonomo Director of Operations Signature:

Date:

Name:

Mr. Jeffrey Curran

Signature:

Title:

Laboratory Manager

Date:

Name: Title:

Ms. Marsha Culik

Quality Assurance Manager

Signature:

Date:

Name:

Mr. David Houle President - IEA

Signature:

Date:

12/4/95

Title: Location:

Cary, North Carolina

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2.0 INTRODUCTION

2.1 Background

Industrial & Environmental Analyst's, Inc. (IEA) is a full-service environmental organization specializing in laboratory analytical services and field support services.

The IEA organization is a network of seven (6) integrated environmental laboratories located throughout the Eastern United States with over 300 employees, making it one of the top ten environmental testing companies in the United States. The corporation serves a broad range of industries including environmental consulting and engineering firms, state and federal agencies, pharmaceutical, petroleum, and electronic component manufacturers. In support of these activities the corporation presently maintains environmental laboratory certifications in over twenty five state programs. IEA Corporate headquarters are located in Cary, North Carolina.

IEA is a wholly-owned subsidiary of the AQUARION Company, headquartered in Bridgeport, Connecticut. AQUARION is listed on the New York Stock Exchange and has annual revenues exceeding 100 million. It is also the largest investor-owned water utility in the country.

The IEA laboratories are located as follows:

IEA/ConnecticutMonroeIEA/IllinoisSchaumburgIEA/North CarolinaCaryIEA/NC-RadiologicalMorrisvilleIEA/MassachusettsN. BillericaIEA/New JerseyWhippany

Detailed information such as mailing addresses and telephone numbers for each of the laboratories is presented in Table 2.2.1.

HISTORY OF IEA

IEA was founded in 1977, in Burlington, Vermont, as a water resources testing facility in support of IBM's facility in Essex Junction, Vermont. IEA served the IBM site exclusively for three years performing ultrapure water analysis, wastewater treatment and pollution control. In 1982, IEA opened a second facility in Research Triangle Park (RTP), North Carolina in order to provide desired services from the IBM facility in RTP. In 1984 IEA expanded its market and began serving the developing environmental testing market. By 1985 IEA had expanded to a full service laboratory offering complete soil and water analysis, field sampling, groundwater analysis and evaluation of hazardous waste. The North Carolina laboratory, which serves as IEA's corporate headquarters, is located in Cary, North Carolina.

In the fall of 1988, IEA positioned itself as one of the leading laboratories in the country by qualifying for the USEPA Contract Laboratory Program (CLP). This development created a favorable position for winning major consulting engineering contracts. As such, IEA grew rapidly and expanded its commercial client base considerably. Due to the rapid increase in demand for environmental services IEA sought potential buyers in 1989 in order to provide resources for future expansion. As a result, IEA was purchased by The Aquarion Corporation, based in Bridgeport, Connecticut in 1989. Aquarion is a New York Stock Exchange-listed corporation that traces its roots to 1857. It has the distinction of being the largest investor-owned water utility in the nation. Annual revenues of Aquarion exceed 100 million.

Since the initial purchase, IEA has acquired several existing environmental laboratories which were operated in strategic locations along the Eastern United States. As a result, IEA now offers very comprehensive environmental testing services including mixed waste radiological testing and a full range of chemical testing performed in support of DOD, DOE, RCRA, CERCLA, NPDES, TSCA and SDWA regulations.

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This plan is intended to describe the quality assurance program of the IEA-Connecticut facility located at 200 Monroe Turnpike, Monroe, Connecticut. IEA operates a corporate wide quality assurance program (Doc.# QAQ00102.NET) and this facility QA program complies with the requirements set forth in the corporate program. In some cases, the requirements in the facility QA program may be more stringent than the corporate program, but in no case can they be less stringent.

TABLE 2.1.1 IEA NETWORK LOCATIONS

North Carolina Corporate Headquarters 3000 Weston Parkway Cary, NC 27513 (919) 677-0090 (919) 677-0427 (Fax)

(800) 444-9919

North Carolina Radiological Laboratory 120 South Center Court Suite 300 Morrisville, NC 27560 (919) 460-8505 (919) 469-2646 (Fax)

Massachusetts 149 Rangeway Road N. Billerica, MA 01862 (617) 272-5212 (508) 667-7871 (Fax) (800) 950-5212 Connecticut 200 Monroe Turnpike Monroe, CT 06468 (203) 261-4458

(203) 268-5346 (Fax)

New Jersey 628 Route 10 Whippany, NJ 07981 (201) 428-8181 (201) 428-5222 (Fax)

Illinois 126 West Center Court Schaumburg, IL 60195 (708) 705-0740 (708) 705-1567 (Fax) (800) 933-2580

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2.2 Definition of Terms

A number of terms are used within this document to describe the corporate QA program in effect at IEA laboratories. To ensure effective communication, the following terms are being defined:

Accuracy

- the degree of agreement of a measurement with an accepted reference or true value. Accuracy is usually expressed as the difference between the measurement and the true value. It is a measurement of the bias in a system.

Analytical Report Turnaround Time - in order to ensure proper communication is maintained, IEA has defined analytical report turnaround times to be always based upon calendar days, not business days. Analytical holding times are also based on calendar days.

Audit

- a systematic check to determine the quality of some function or activity. Audits may be of two basic types, performance audits or system audits. Performance audits involve a quantitative comparison of the labs results to that of a proficiency sample containing known concentrations of analytes. A system audit is a qualitative evaluation that normally consists of an on-site review of a laboratory's quality assurance system and physical facilities.

Batch

- the basic unit for analytical quality control. It is defined as a group of samples which are analyzed together with the same method sequence and the same lots of reagents and with the manipulations common to each sample within the same time period or in continuous sequential time periods. Samples in each batch should be of similar composition (matrix). At IEA laboratories, the maximum batch size has been set at 20 samples. At IEA's smaller laboratories where the number of samples received daily may be low, samples received in a given week may be combined into one analytical batch. Due to holding time constraints, individual samples may be extracted on different days as compared to other samples in the batch. If this is the case, a method blank must be performed daily with every sample extraction. The other QC samples such as MS and MSD are only performed for the total analytical batch.

Comparability

- a measure of the confidence with which one data set can be compared to another.

Completeness

- a measure of the amount of valid data obtained from a measurement system compared to the amount that was expected to be obtained under routine operating conditions.

Data Quality Objectives

- during the planning phase of a project requiring laboratory support, the data user must establish the quality of data required from the investigation. Such statements of data quality are known as data quality objectives (DQOs). The DQOs are qualitative and quantitative statements of the quality of data required to support specific decisions or regulatory actions.

Data Validation

- a systematic effort to review data to identify any outliers or errors and thereby cause deletion or flagging of suspect values to assure the validity of the data to the user. This process may be done by manual or computer methods.

Field Blank

- contaminant free water, or appropriate matrix, used during sampling activities to determine if there is any potential for sample contamination associated with the field sampling or equipment.

Library Search

- a technique used by which a mass spectrum of an unknown compound is compared to the mass spectrum of compounds contained in a computer library in an effort to identify

4 4

(MDL)

Quality Control (QC)

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unknown compounds.	Compounds i	dentified	in this	manner	аге	referred to	as	"tentatively
identified compounds"	(TICs).							•

Matrix Spike - the process of adding a known amount of analyte to a sample and analyzing the sample.

The amount of analyte recovered is calculated as a percent recovery. This technique is

used to assess accuracy of analysis.

Matrix Spike Duplicate - a second matrix spike is compared to the results of the matrix spike to assess precision of

the analysis.

Method Blank - contaminant free water, or appropriate matrix, taken through the entire analytical process to determine if there is any contamination associated with the analytical procedures.

Method Detection Limit - the minimum concentration of a substance that can be

measured and reported with 99% confidence that the analyte concentration is greater than

Practical Quantitation

Limit (PQL)

- is the lowest level that can be reliably achieved within specified limits of precision and accuracy during routine operating conditions.

Precision - a measure of mutual agreement among individual measurements of the same property, usually under prescribed similar conditions. Precision is usually expressed in terms of

standard deviation.

. ..

Quality Assurance (QA) - the total integrated program put in place to assure the reliability of data generated in the laboratory.

- the routine application of specific, well-defined procedures which ensure the generation

of data which fulfill the objectives of the QA program.

Quality Assurance - a written assembly of management policies, objectives,

Program Plan (QAPP) - a written assembly of management policies, objectives,

principles and general procedures which outline how the laboratory intends to generate

data of known and accepted quality.

Quality Assurance - a written document, which presents, in specific terms,

Project Plan (QAPjP) the policies, organization, objectives, functional activities and specific QA/QC activities designed to achieve the data quality objectives of a specific project. There are 16 essential elements which EPA has mandated to be addressed in a project plan.

Relative Percent - relative percent difference (RPD) is used as the measure of precision between Difference (RPD) sample duplicates. The formula utilized to calculate RPD is as follows:

Relative Percent Difference (RPD)

RPD = (Sample Result - Duplicate Result) x 100
Mean of Sample and Duplicate Results

Note: RPD is expressed as the absolute value obtained from the above formula.

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Representativeness

- the degree to which data accurately and precisely represents a characteristic of a population, parameter variations at a sampling point, or an environmental condition.

Standard Operating Procedure (SOP)

- a detailed, written description of how a laboratory executes a particular procedure or method. It is intended to standardize the performance of the procedure.

Surrogates

- generally, organic compounds which are not target analytes, that are added to samples to assess analytical performance of a method. These compounds are spiked into all blanks, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

Trip Blank

- contaminant free water, or appropriate matrix, which accompanies bottles and samples during shipment to assess the potential for sample contamination during shipment. Trip blanks are not opened in the field.

Tuning

- a technique used in GC/MS procedures to verify that the instrument is properly calibrated to produce reliable mass spectral information.

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2.3 Purpose

The IEA-Connecticut quality assurance program serves as an operational charter for the organization. It defines the purpose, organizational structure, and operating principles of the laboratory and presents an overview of the key elements of the quality assurance program. This quality assurance program will be reviewed and modified as necessary on an annual basis. Any deviation from this program must be approved in writing by the facility QA manager and copied to the President.

This quality assurance program has been prepared according to guidelines presented in the USEPA document entitled "Guidelines and Specifications for Preparing Quality Assurance Program Plans", Office of Monitoring Systems and Quality Assurance, Office of Research and Development, USEPA, (QAMS-004/80), EPA-600/8-83-024, June, 1983.

2.4 Scope

This QA program applies to the generation of analytical data at the IEA-Connecticut lab location. Since the vast majority of environmental client needs are driven by various federal and state regulations, the program has been designed to meet the requirements of the following programs:

Clean Water Act (CWA)
Clean Air Act (CAA)
Safe Drinking Water Act (SDWA)
Resource Conservation and Recovery Act (RCRA)
Comprehensive Environmental Response, Compensation and Liability Act (CERCLA)

This Quality Assurance Program Plan (QAPmP) covers laboratory operation at IEA-Connecticut. The purpose of this QAPmP is to provide information on laboratory operations as required for specific Quality Assurance Project Plans (QAPjPs), and to provide the basis for the Quality Assurance Program at IEA-Connecticut. This program is based on the IEA Corporate Quality Assurance Program Plan (Doc# QAQ00102.NET).

This QA program applies to the generation of analytical data utilized for environmental monitoring and assessment programs. The major types of laboratory support for government regulations are as follows:

- Analysis and characterization of environmental (soil, sediment, water and air) and waste samples per the Resource Conservation and Recovery Act (RCRA) for either compliance, disposal or delisting purposes.
- Analysis of drinking water samples in support of the Safe Drinking Water Act (SDWA).
- Analysis of environmental samples in accordance with contracts with the USEPA CLP program and various state agencies (CERCLA and NYSDEC).
- Analysis of environmental samples (soil, sediment, water and air) for contaminants such as those compounds
 found on the EPA priority pollutant list, target compound list, etc. for site assessment purposes.
- Analysis of waste stream samples in accordance with NPDES requirements.

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3.0 QUALITY ASSURANCE POLICY STATEMENT

It is the intention of IEA corporation to consistently produce analytical data of known and documented quality at all network laboratories which fully meet clients' data quality objectives.

The contents of the QA program describe the activities which are utilized in order to ensure this commitment is maintained.

IEA Quality Policy

"Management and staff are committed to maintaining a carefully controlled analytical environment in order to ensure the consistent generation of accurate data which meets or exceeds the data quality objectives of our clientele."

IEA recognizes that maintaining a proper ethical standard is an important element of an effective quality assurance program. In order to ensure that all personnel understand the importance the company places on maintaining high ethical standards at all times, IEA has established an "Ethics Policy" and it is presented for your information. This policy is used to set the standard within the organization for day-to-day performance. Each employee is requested to sign the ethics policy, signifying agreed compliance with it's stated purpose. Copies of all signed ethics policy statements are maintained in personnel files.

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IEA ETHICS POLICY

The management of IEA corporation recognizes our responsibility to clients and fellow employees to ensure that fair and ethical business practices are followed at all facilities.

Our clients have placed their trust in our organization to continually provide high quality data which is valid, defensible and represents sound professional judgement at all times. In order to meet this responsibility it is imperative that high ethical standards be maintained at all times by all employees.

The management and staff are committed to maintaining a carefully controlled analytical environment which assures the consistent generation of accurate data which meets the data quality objectives of our clientele.

The following represents the IEA ethics policy which has been adopted to clearly identify the corporate position on ethical practices. Failure to comply with this policy cannot and will not be tolerated.

The Company and all its Employees will:

- Fully comply with all applicable federal, state, and local laws and regulations.
- Produce analytical products that are accurate, defensible and which represent sound professional judgement at all times.
- Provide employees with guidance and an understanding of the ethical and quality standards required in
 the environmental industry. In this regard, all employees should feel free to identify any ethical
 misconduct without fear of retribution. Any employee involved in any form of ethical misconduct will be
 subject to immediate disciplinary action including potential termination of employment.
- Present services to clients in a confidential, honest and forthright manner and strive to deliver quality products at a fair price.
- Treat employees equitably by compensating them fairly, acknowledging their scientific contributions, and
 providing them opportunities for professional growth and development.
- Offer employment opportunities to qualified candidates regardless of their race, creed, color, sex or age.
- Be a responsible corporate citizen of the community by operating in an environmentally sound manner at all times.
- Maintain all facilities in a safe and professional manner through maintenance of a safety awareness
 program and provide the necessary safety equipment and training to protect all employees from
 preventable injury and chemical exposure.

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4.0 QUALITY ASSURANCE MANAGEMENT

4.1 Introduction

The management of IEA-Connecticut is committed to the execution of the quality assurance program described in this document. The officers of IEA as well as lab directors and lab managers are required to comply with the program's stated goals, requirements and responsibilities.

In addition, each staff member has a responsibility to ensure compliance at all times with the QA program.

4.2 Assignment of Responsibilities

The primary objective of the network quality assurance program is to ensure that systems are in place such that all network laboratories consistently generate high quality analytical data.

Additionally, the QA program provides a mechanism to identify and implement policies to improve the quality of products and services. Records must also be maintained to document the laboratory's performance.

Quality assurance at IEA is monitored at both the corporate and laboratory levels. IEA's network quality assurance program is led by the president of IEA. The QA program at each network lab is directed by the QA manager at that facility, who reports directly to the laboratory's director and indirectly to the president. Figure 4.2.1 presents the organizational structure of network quality assurance functions and Figure 4.2.2 illustrates the overall general management of the corporation.

The following provides a listing of responsibilities and authority of key managerial personnel. Section 5 of the Appendix presents the organizational structure of the IEA-Connecticut facility.

Director of Operations

Responsibility:

All corporate directors and managers comply with the quality assurance program and require similar compliance by all staff personnel.

Ensure that all laboratory operations under their control are active participants in attaining the network quality assurance objectives.

Ensure compliance with methods and procedures as written.

Timely compliance with any corrective action requirements.

Ensure that instrument tunings and calibrations are performed at the required frequency and that instrument maintenance and logbooks are maintained in an orderly manner.

Authority:

Maintain the authority to suspend or terminate employees for dishonesty, or non-compliance with established QA policies and procedures.

The directors' or managers' authority is granted from a vice president of IEA, to whom they report.

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Laboratory Manager

Responsibility:

Ensure compliance with methods and procedures as written.

Ensure that analytical procedures are performed in accordance with the requested method and SOPs.

Oversee preparation of analytical reports and data review.

Authority:

Maintain the authority to suspend or terminate employees for dishonesty, or non-compliance with established QA policies and procedures.

Authority is granted from the Director of Operations, to whom they report.

Laboratory Quality Assurance Manager

Responsibility:

Responsible for recommending pertinent additions to the network QA program.

Responsible for monitoring and assessing compliance of the laboratory with the requirements contained in the QA program.

Function as a liaison between the corporate QA director and laboratory staff at their facility.

Represent the laboratory during all external audits conducted by clients or regulatory agencies.

Conduct semi-annual audits and inspections to assess compliance with established methods, policies and procedures. Results of these audits are reported to the network QA director and the laboratory director.

Maintain a document control system containing current policies and procedures utilized by the laboratory.

Maintain various certification programs for the laboratory.

Review laboratory performance on various QC proficiency samples submitted to laboratories by state and federal agencies.

Inform local and corporate management of the status of the QA program at the particular facility through a monthly QA report.

Investigate all inquiries relative to data quality issues and follow up on corrective action if necessary.

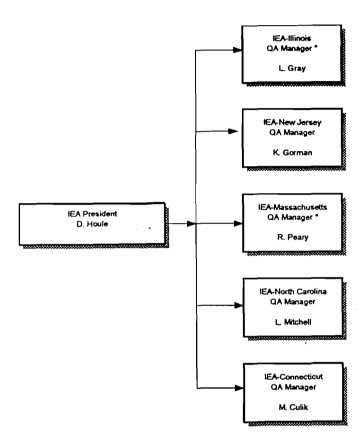
Authority:

The quality assurance staff has the authority to stop or change any analytical procedure in order to assure that data quality is maintained.

The authority of the QA staff is granted by the director of the facility.

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FIGURE 4.2.1 NETWORK QUALITY ASSURANCE ORGANIZATIONAL CHART

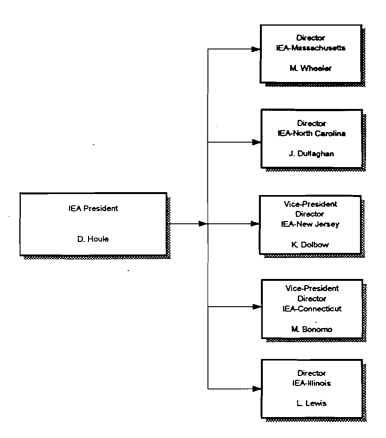


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Above positions are part-time QA positions. These individuals also have operational responsibilities.

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FIGURE 4.2.2 NETWORK ORGANIZATIONAL CHART



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4.3 Communications

The quality assurance department communicates internally and externally through various means. Communication can take place via telephone, memoranda or take the form of audit reports. At the present time, the quality assurance department participates in a weekly conference call to discuss relevant issues and disseminate information.

In addition, various quality assurance reports are routinely generated as discussed in section 4.5.

4.4 Document Control

A system of document control is essential to provide the framework necessary to ensure that methods and procedures are followed in a consistent manner.

IEA has developed a centralized document control system which is maintained for the entire network and is administered by the corporate staff located at the Cary, North Carolina facility. The document control system provides for the following:

- A unique document control number for each document
- A central location for all documents
- A systematic method for distribution of approved documents
- A tracking system for existing documents
- Identification of document revisions
- A mechanism for periodic review of documents
- · Archival of outdated material
- A focal point for information exchange
- Facilitates the establishment of standardized methods and procedures

A detailed description of the document control system is contained in IEA document number QAS00101.NET. This document is available for inspection and review during a site visit. The Quality Assurance Manager is responsible for ensuring that the document control system is properly managed. Any new or revised document must be submitted to the QA Manager for review and distribution.

It is the responsibility of all members of the laboratory to maintain complete records of all operations performed. All records shall be neat and organized. All laboratory records are the property of the laboratory and shall not be removed from the premises without permission from supervisors. All records are considered confidential and must be safeguarded. Unauthorized changes, loss or destruction of records can be grounds for dismissal from the laboratory. Consult the IEA, Inc. Ethics Policy regarding integrity of data and employee conduct.

Measurement records must be recorded in pre-printed record logs or pre-printed measurement logs. This policy will facilitate the organization and archival of all laboratory data for future reference.

All injection forms, instrumentation forms, sample prep forms, QC forms, etc. which are used to process samples and measurement results are described and attached to each analytical SOP. The SOP specifies where these records and forms are cataloged and stored.

All measurement data is recorded in logbooks or on pre-printed log sheets in permanent ink. Transcriptions will be avoided whenever possible. The record will reflect the measurement performed and all appropriate details for conclusions related to the measurement. The record must be initialed and dated by the individual performing the measurement on the day the measurement is performed. Corrections shall be made by drawing a single line through the error, initialing and dating the error. All forms will be reviewed by the QA Manager annually. If it is found that the document does not meet the requirements of the SOP, the discrepancy is forwarded to the group/section leader through the corrective action process (reference SOP on Corrective Action Reports -QAS00501.CT). Further detail on laboratory document control is found in the SOP on Document Control - QAS00301.CT.

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4.5 QA Program Assessment

The quality assurance program can only accomplish its objectives if management and staff are committed to adherence to the program. In order to assess continued compliance and to identify strong and weak points of the program, the network QA director conducts annual assessments at each location.

Each quality assurance manager conducts an annual audit of the particular laboratory. A copy of the audit along with any proficiency test results obtained are submitted to the president.

A written status report is prepared monthly by each of the facility QA managers. A copy of this report is issued to the facility laboratory director as well as the corporate president. The corporate staff provides a summary of these reports each month to upper management. A typical status report would include such information as:

- · Changes in the quality assurance program
- Summary of proficiency results at each network lab
- · Summary of on-time report issuance
- Changes in certification status
- Summary of system audits conducted at each network lab
- Significant QA concerns and recommendations for resolution
- Accomplishments since the previous report

4.6 Additional Lab Policies to Achieve QA Objectives

In addition to policies and procedures specified in other sections of this document there are numerous policies and standard procedures which have been implemented to ensure that data of known quality is continually generated by all network laboratories. Examples and a brief description of a few of these additional policies are presented below:

4.6.1 Participation in EPA Water Supply and Water Pollution Proficiencies

The USEPA currently operates a Water Supply (WS) and a Water Pollution (WP) proficiency program. Each program consists of the issuance of proficiency samples twice in a calendar year. Analysis of proficiency samples on the second set of samples in a year are only required by EPA for those parameters which the laboratory failed during the first round in a given year. As part of IEA's QA program, full participation and analysis of all appropriate parameters is required of all IEA labs regardless of past performance. This serves as an important indicator on the continuing quality of data being generated at each facility.

The laboratory also participates in the NYSDOH proficiency testing program for Potable Water, Hazardous Waste and CLP. The lab currently analyzes quarterly organic PE samples from EPA for the CLP program.

4.6.2 Corporate Laboratory Performance Evaluation Program

In addition to participating in various agency sponsored performance evaluation programs such as Water Supply (WS) and Water Pollution (WP) studies, the corporate quality assurance office conducts additional performance evaluation studies.

Periodically, performance evaluation samples are submitted to each laboratory for parameters which are not addressed in other performance evaluation programs (ie. TCLP testing). In this type of testing, the laboratory is aware the samples are performance check samples but the "true" concentration values are unknown. The results are submitted to corporate QA for evaluation and a report is issued on the findings. Corrective actions are taken if required, as a result of these test findings.

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4.6.3 Routine Use of QC Check Samples

One of the most important goals of a strong quality assurance program is to ensure that data of known quality is consistently generated during day-to-day operations. IEA accomplishes this through the routine inclusion of a QC check sample in every inorganic analytical batch which includes metals and wet chemistries. For organic testing including GC and GC/MS a QC check sample is analyzed at the frequency required in the particular method. Section 8 in the Appendix provides QC check sample requirements for selected methods. A QC check sample is an artificially prepared sample which contains the analytes of interest. The source of the standards used for preparation of the check sample must be independent (either another vendor or a different lot from the same vendor) from those used to prepare a calibration curve. The QC check sample is an important mechanism to confirm the method is being executed properly during routine analysis. The QC check also serves as a useful tool in identifying possible problems such as matrix interference, degraded analytical standards, and inaccurate standard preparation.

In certain cases, reliable QC check samples are not available for a particular procedure. In such cases, the QA manager has the authority to waive this requirement for that particular test. The QA manager must document this waiver in writing.

4.6.4 Central Solvent Monitoring Program

IEA has established a central monitoring program for commonly used solvents within the corporation. Prior to use, a specific lot number of these solvents is provided to the laboratory for testing. The solvents are concentrated and tested for the presence of interfering substances relative to their intended use. If the particular lot of solvent passes the defined acceptance criteria, the vendor is notified and the solvent lot is reserved for use by the entire corporation. The approved lot numbers are provided to all laboratories and only approved solvents can be employed. IEA Document # QAS00400.NET describes the details of the solvent approval program and is available for review during a site audit.

4.6.5 Quality Assurance Final Report Review

An integral portion of the overall quality assurance program is the consistent monitoring of final reports as they leave IEA facilities. Each QA manager is responsible for reviewing 5 percent of the final data reports issued each month. The reports to be reviewed are picked at random. The reports are reviewed for typographical errors, technical clarity and overall presentation.

4.6.6 Lateness of Data Reports

IEA recognizes that one cannot overlook the timeliness of data generation when assessing the quality of our services from our client's perspective. High quality data, when delivered several weeks late is not acceptable. In recognition of this, IEA monitors the lateness of all reports on a monthly basis from each of its laboratory operations. The actual report shipment date is compared to the date originally projected to the client. This information is gathered monthly through the QA department and a monthly report is issued to each laboratory director and to corporate management. This monitoring program serves to identify service trends, and to ensure that corrective action will be taken before problems occur.

4.6.7 Method Detection Limit Verification

Each laboratory is required to perform a method detection limit study for all commonly performed test methods. The study must be performed during the initial setup and verification of the particular method. In addition, the MDL study must be conducted in the event of a major change in the technique or instrumentation. The results of the MDL studies must be fully documented and available for review upon request. The quality assurance manager is responsible for maintaining such records. Specific state

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certification programs may require MDLs to be determined annually. If this is the case, the laboratory will comply with this requirement.

4.6.8 Establishment of IEA Good Laboratory Practices

In order to ensure that various procedures are executed in a consistent and comprehensive manner, IEA has developed a series of procedures which fall into the category of "Good Laboratory Practices". These practices have been endorsed by the corporation for routine use at each laboratory facility and are defined in various standard operating procedures throughout the organization. Examples of a few of these "Good Lab Practices" are presented below for the reader's information:

A. Standardized logbook requirements (Doc# QAS01201.NET)

Preprinted pages
Prenumbered pages
Dedicated logbooks per test method
Bound logbooks
Use of black ink only
Document controlling of logbooks
Archival of old logbooks
Acceptance criteria in logbook
Making corrections
Secondary review of logbook entries

B. Balance calibration (Doc# QAS01002.NET)

Unique identifier for each balance
Balance must be checked daily with use and documented
Acceptance ranges are established for each balance
Balance must be checked in the weight range normally used
All balances must be professionally serviced and calibrated annually

C. Temperature monitoring requirements for lab apparatus (Doc# QAS00801.NET)

Refrigerators, freezers and lab ovens are checked each work day
Unique identifier assigned for each unit
Acceptance ranges are established for each unit
Thermometers used in monitoring must be calibrated to a NIST traceable thermometer annually, at a minimum. State certification requirements may require more frequent calibration
All thermometers are immersed in appropriate media to avoid temperature fluctuations during measurement

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D. Correcting data and general laboratory records (Doc# QAS01300.NET)

All entries must be entered in black ink.

"White Out" is not to be used at any time within the laboratory for alteration or correction of lab documents

Corrections are made using a one-line strikeout

All corrections are initialed and dated by the data editor

E. Handling reagents and analytical standards (including the following)

Recording receipt and expiration dates

Documenting preparation of reagents and standards

Labelling requirements

Disposal

F. Cleaning procedures for sample containers and laboratory glassware (Doc# QAS01400.NET) (including the following)

Cleaning sample containers Cleaning inorganic glassware Procedures for cleaning organic glassware

G. Requirements for general lab calibration curves (including the following)

In cases where the referenced analytical method does not provide specific guidance or requirements for development of initial or continuing calibration curves, the following procedure is to be utilized by the laboratory.

All standard calibration curves must consist of a minimum of three points. Any deviation from this must be approved in writing by the facility QA manager.

All calibration points must be recalculated using the generated curve and all calibration points must be within 10% of the expected value for the curve to be considered acceptable.

Concentration of compounds or analytes must fall within the calibration range of the curve to be acceptable for quantitation for inorganic and organic methodology.

H. Method blank subtraction

Subtraction of method blanks from sample results is not permitted unless specifically authorized by the laboratory QA manager.

4.6.9 Quality Control Charts

Maintaining quality control charts is currently not mandatory under IEA's corporate quality assurance program, however, many state certification programs require them. As a result, laboratories are required to comply with such state certification requirements.

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5.0 PERSONNEL QUALIFICATIONS

5.1 Introduction

IEA's management is very proud of its highly qualified and professional staff. The IEA-CT staff consists of over 50 professionals and support personnel which include:

Analytical Chemists
Quality Assurance Specialists
Computer Systems Analysts
Environmental Technicians
Customer service Staff
Account Executives

5.2 Education and Experience

In order to ensure that employees have sufficient education and experience to perform a particular task, requirements have been defined for each laboratory position.

The personnel who are responsible for operations of sample analyses and data validation are outlined in Section 5 of the Appendix. Section 1 of the appendix presents professional profiles of key personnel within the IEA-Connecticut organization. Profiles of additional IEA staff members are available for review during a facility visit or are available upon special request.

Throughout the years, IEA has performed sophisticated environmental analysis for a significant number of large corporations. Examples of relevant experience are available upon request.

5.3 Training

IEA is committed to furthering the technical and interpersonal skills of employees at all levels. Technical training is accomplished within each laboratory by management to ensure method comprehension. It is at these training sessions that staff is updated on all current technical advances. It is IEA policy that all new personnel must demonstrate competency in performing a particular method through the analysis of QC check samples prior to the analyst conducting analysis independently on client samples. New analysts may conduct analysis on client samples along with another experienced analyst prior to the completion of the training period. All laboratory personnel are required to acknowledge through signature that they have read and understood the SOP's that are appropriate for their particular area.

All laboratory personnel must have adequate education, training, and experience to carry out their responsibilities. The QA Manager and the Laboratory Management will periodically review the training needs of the staff and make recommendations for any additional training. Each department within the laboratory is responsible for personnel training. Training sessions are scheduled on a monthly basis. Each training session, whether it be individual or group training must be documented utilizing the forms attached to the corporate SOP for Employee Training QAS01600.NET. The completed forms must be submitted to the Human Resource department for placement into the employee training files. Included in the training process is analyst proficiency testing. A successful QC check sample must be analyzed and documented for each analyst. This information is on file with the QA Manager.

5.4 Certifications

Table 5.4.1 presents the state certifications held by the IEA-Connecticut laboratory. Many states certify laboratories for specific parameters or tests within a category (i.e. method 325.2 for wastewater). The information in the following table indicates the lab is certified in a general category of testing such as drinking water or wastewater analysis. The laboratory should be contacted directly if parameter-specific certification information is required.

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IEA-CT currently participates in the USEPA Superfund Contract Laboratory Program (CLP). The lab is also approved to perform work for the Army Corps of Engineers which validates laboratories on a project-by-project basis.

This document is updated annually; therefore, it is likely that additional certifications, beyond those listed, may be currently available. This information can be obtained easily by calling the specific laboratory (See Table 2.2.1 for phone) and asking for the QA manager.

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TABLE 5.4.1

STATE CERTIFICATIONS

In some instances it may be necessary for environmental data to be reported to a regulatory authority with reference to a certified laboratory. For your convenience, the laboratory identification numbers for the IEA-Connecticut laboratory are provided in the following table. Many states certify laboratories for specific parameters or tests within a category (i.e. method 325.2 for wastewater). The information in the following table indicates the lab is certified in a general category of testing such as drinking water or wastewater analysis. The laboratory should be contacted directly if parameter-specific certification information is required.

IEA-Connecticut Certification Summary (as of June 1993)

State	Responsible Agency	Certification	Lab Number
Connecticut	Department of Health Services	Drinking Water, Wastewater	PH-0497
Kansas	Department of Health and Environmental Services	Drinking Water, Wastewater/Solid, Hazardous Waste	E-210/E-1185
Massachusetts	Department of Environmental Protection	Potable/Non-Potable Water	CT023
New Hampshire	Department of Environmental Services	Drinking Water, Wastewater	252891
New Jersey	Department of Environmental Protection	Drinking Water, Wastewater	46410
New York	Department of Health	CLP, Drinking Water, Wastewater, Solid/ Hazardous Waste	10602
North Carolina	Division of Environmental Management	Wastewater	388
Rhode Island	Department of Health	ChemistryNon- Potable Water and Wastewater	A43
California	Department of Health Services	Hazardous Waste	1778

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6.0 FACILITIES, EQUIPMENT AND SERVICES

6.1 Introduction

The following describes the physical facility of the IEA-Connecticut laboratory.

6.2 Facilities

IEA-Connecticut

The laboratory currently maintains a staff of approximately 50 environmental professionals and occupies a facility of approximately 13,000 sq. ft. Separate laboratory areas are dedicated to GC instrumentation, GC/MS instrumentation, extractions for organic parameters, sample preparation for metals analysis, metals analysis and wet chemistries.

The volatiles analysis laboratory containing GC/MS instrumentation has a separate air handling system which is maintained at a positive pressure at all times. The organic sample preparation laboratory has a separate HVAC system that creates negative pressure in the area. This design results in a contaminant-free environment for trace-level volatiles analysis.

Critical instrumentation such as GC/MS units, ICP's, AA's, data systems and gas chromatographs are tied into an uninterruptable power supply system (UPS) to minimize instrument downtime and damage for short duration power interruptions.

The floor plan of the analytical laboratory is included in Section 4 of the Appendix.

Security of Facilities

The laboratory is secured by a card key access system. Only authorized IEA-CT personnel have access to the facility. All visitors must sign in with the receptionist and must be accompanied by an IEA-CT employee.

The sample receipt and storage area is under the responsibility of the sample custodian. This area is a locked, secure area opened by the sample control department each day. A walk-in refrigeration unit and 10 locked commercial refrigerator units are used to house samples waiting for analysis. Samples for volatile analysis are stored in separate units. Locked laboratory refrigerators, located throughout the laboratory, are used to maintain sample extracts or laboratory reagents. Each laboratory refrigerator is dedicated to sample, sample extract, or reagent storage.

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6.3 Equipment

The following is a summary listing of equipment utilized at the IEA-CT facility. A more detailed listing is presented in Table 6.3.1.

Analytical instrumentation at IEA-Connecticut includes:

- 9 Gas Chromatographs/Mass Spectrometers (GC/MS)
- 5 Gas Chromatographs (GC)
- 3 Atomic Absorption Spectrometers (Graphite Furnace/AA)
- 2 Inductively Coupled Argon Plasma (ICP) Emission Spectrometer
- 2 Mercury analyzers
- 2 Gel Permeation Chromatographs
- 2 Infrared Spectrometer (IR)
- 1 Total Organic Halide (TOX) Analyzer
- 1 Total Organic Carbon Analyzer
- 1 Automated Analyzer for Wet Chemistries
- 1 LIMS (Laboratory Information System)
 Automated Data Acquisition Management System (ADAM)

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Table 6.3.1-Laboratory Equipment Listing

WET CHEMISTRY

Equipment Name	Manufacturer	Model Number	Serial Number
Centirfuge	DYNAC	0101	16846
Spectrophotometer, UV-VIS	Perkin-Elmer	35	34630
IR-Spectrophotometer	Perkin-Elmer	1310	134423
Turbidimeter	Hach Company	2100A	851017142
TOC Analyzer	Xertex-Dohrmann	DC-80	HF2029
TOX Analyzer	Xertex-Dohrmann	мсз а,в	MF 2106
Fluorometer	Sequoia-Turner Corp.	112-003	D 01491
pH/ISE Meter	Orion	SA 720	SR45A
pH/ISE Meter	Beckman	12	0232578
Conductivity Meter	Cole-Parmer Instrument	1484-20	1421
Flash Point Apparatus	Precision Scientific	Pensky-Martin	10 Au-12
Oven	Fisher Scientific	SSG	291
Oven	VWR	1320	0701090
Incubator	Blue M Electric	100 A	IN1-1362
Bio Refrigerator	Frost Queen	R20/L	00029
BOD Incubator (2)	Precision Scientific	FU199JRW2/FU178RRW2	FLC02662
Midi Distillation Setup (2)	Andrews Glass Co.	110-10-R	A4W0309/0209
D.O. Meter	YSI	51A	0241
Autoclave	Market Forge	STM-E	034200
COD Reactor	НАСН	45600	920300006892
Muffle Furnace	Thermolyne	-	_
TKN block digestor	Scientific Instruments	AD-4020	8915049
Digital Hot Plate/Stirrer	PMC	730	0298E
Digital Hot Plate/Stirrer	PMC	730	0299E
Semiautomated Analyzer	LACHAT	Quikchem	125360

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Table 6.3.1-Laboratory Equipment Listing

METALS

Equipment Name	Manufacturer	Model Number	Serial Number
Mercury Analyzer	Spectro-Products	HG4	4708
Mercury Analyzer	Jarrell-Ash	QS1	1210031
ICP-Trace	Jarell-Ash	JA61T	349490
ICP-Simultaneous	Jarrell-Ash	JA61	67782
Furnace AA	Perkin-Elmer	Z3030	3131
Furnace AA	Perkin-Elmer	Z5100	130911
Furnace AA	Perkin-Elmer	Z5100 PC	135141

ORGANIC EXTRACTIONS

Equipment Name	Manufacturer	Model Number	Serial Number
Gas Chromatograph	Perkin-Elmer	8320	83N546502
Gel Permeation Chromatograph	ABC	1002B	7323
Gel Permeation Chromatograph	ABC	AP1000	9228
Refrigerator	ww	4EF	F3978U
Oven	ASP	D 1142	144011
Oven	ASP	D 1162	149010
Sonicator	Sonics & Materials	SM500	6892
Sonicator	Sonics & Materials	VCX-400	00030C
Sonicator	Tekmar	TM500	7264
Auto Sampler	Perkin-Elmer	AS100	95234
Rotary Evaporator	BUCH I	R-114	-
Seporatory Funnal Shaker	Glas-Col	Series 100	F715-10-B5J

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Table 6.3.1-Laboratory Equipment Listing

GC/MS VOLATILES

Equipment Name	Manufacturer	Model Number	Serial Number
Purge & Trap	Tekmar	LSC 2000	91318021
Purge & Trap	Tekmar	ALS 2016	91322002
Purge & Trap	Tekmar	LSC 2000	91203019
Purge & Trap	Tekmar	ALS 2016	91232007
Tube Desorber	Envirochem	810TD	268153
Tube ALS	Envirochem	MTD	MT-1005
Purge & Trap	Tekmar	LSC 4000	254
Purge & Trap	Tekmar	ALS	372
Purge & Trap	Tekmar	LSC-2	1824
Computer/Data System	Hewlett Packard	425T	3048T147545
Data System-Enviroquant	Hewlett-Packard	Vectra XM2	-
Data System -Enviroquant	Hewlett-Packard	Vectra	-
Terminal	Hewlett Packard	X-window	3048T18725
Terminal	Hewlett Packard	X-window	3048T18726
Disc Drive	Hewlett Packard	7914	-
Disc Drive	Hewlett Packard	7914	
P&T	Tekmar	LSC-2	227
P&T	Tekmar	4000	192
P&T	Tekmar	4000	398
P&T	Tekmar	LSC-2	1324
P&T	Tekmar	ALS	679
P&T	Tekmar	ALS	494
P&T	Tekmar	ALS	1068
GC/MS	Hewlett Packard	5995B	2217A00358
GC/MS	Hewlett Packard	5995C	2413A00659
GC/MS	Hewlett Packard	5995C	2413A00430
GC/MS	Hewlett-Packard	5890 Series II/5972 MSD	-
Equipment Name	Manufacturer	Model Number	Serial Number
GC/MS	Hewlett-Packard	5890 Scrics II/5972 MSD	-
Terminal	Hewlett Packard	45849A	2530A13541
Terminal	Hewlett Packard	35751	2643A07666

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Table 6.3.1-Laboratory Equipment Listing

CRT	Hewlett Packard	35731A	8633K26810
Printers (partial list)	Hewlett Packard	2934A	2635A32940
Printers	Hewlett Packard	2934A	2715A43948
Printers	Hewlett Packard	2225A	2512830379
Printers	Hewlett Packard	2225A	2510832359
Terminal	Hewlett Packard	35751	2630A06622
CRT	Hewlett Packard	35731A	8610K20516
Magnetic Tape Unit	Hewlett Packard	7970E	N/A
Scanning Interface	Hewlett Packard	59824A	N/A
Scanning Interface	Hewlett Packard	59824A	N/A
Cart. Tape Unit	Hewlett Packard	7914	N/A
5010 Auto Desorber	Tekmar	14-2150-000	133-GT
Cart. Tape Unit	Hewlett Packard	7914	N/A

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Table 6.3.1-Laboratory Equipment Listing

GC/MS SEMI-VOLATILE

Equipment Name	Manufacturer	Model Number	Serial Number
Gas Chromatograph	Hewlett Packard	5890	7518A05422
Gas Chromatograph	Hewlett Packard	5890	2728A14615
Auto Sampler	Hewlett Packard	76732A	2441 A03468
Mass Selective Detector	Hewlett Packard	5970	2513A00923
Mass Selective Detector	Hewlett Packard	5970	2716A10638
Computer Terminal	Hewlett Packard	150 П	2720Y05798
Computer Terminal	Hewlett Packard	150 11	2720Y03266
Computer Terminal	Hewlett Packard	150 П	2530A13540
Scanning Interface (2)	Hewlett Packard	59824A	_
Tape Drive	Hewlett Packard	9144	
Disc Drive	Hewlett Packard	7958	
9 Track Magnetic Tape	Hewlett Packard	7970E	_
9 Track Magnetic Tape	Hewlett Packard	7970E	
Computer	Hewlett Packard	НР1000А	_
Computer	Hewlett Packard	HP1000	_
Printer	Hewlett Packard	2934A	2524A19296
Printer	Hewlett Packard	2235A	2814A11816
Printer	Hewlett Packard	2225A	2618S30681
Printer	Hewlett Packard	2934A	2643A35608
Autosampler	Hewlett-Packard	7673A	2546A01489
GC	Hewlett-Packard	5890A	
MSD	Hewlett-Packard	5971 A	3040A01426
Autosampler	Hewlett-Packard	7673	3120A28431
Computer	Gateway	386/25 DX	365201578025
Terminal	Hewlett Packard	36731A	8635K28238
GC/MS/MSD	Hewlett-Packard	5890 SeriesII/5972MSD	
Data System -Enviroquant	Hewlett-Packard	Vectra XM2	-

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Table 6.3.1-Laboratory Equipment Listing

GAS CHROMATOGRAPHY

Equipment Name	Manufacturer	Model Number	Serial Number
GC	Hewlett-Packard	5890	2541A06301
GC	Hewlett-Packard	5890	2750A14840
Autosampler	Hewlett-Packard	7673A	2546A00709
Autosampler	Hewlett-Packard	7673A	3123A25128
Autosampler	Hewlett-Packard	7673 A	2718A0653A
Integrator	Hewlett-Packard	3396A	2804A01106
Integrator	Hewlett-Packard	3393A	2332A00D80
GC	Hewlett-Packard	5890 Series II	3121A35826
GC	Hewlett-Packard	5890 Series II	3235A44989
Data System	Hewlett-Packard	HP1000A	3020A05230
Terminals (3)	Hewlett-Packard	35741A	
Printers (3)	Hewlett-Packard	35741A	
Tape Drive	Hewlett Packard	9144	2724E13732
Data System-Enviroquant	Hewlett-Packard	Vectra XM2	-

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6.4 Instrument Maintenance

Where it is economically feasible, the IEA-CT laboratory has service contracts for major instruments. These contracts provide routine preventive maintenance according to the manufacturer's requirements. Additionally the laboratory maintains an inventory of expendable parts and supplies to minimize downtime and to allow laboratory personnel to make minor repairs if necessary.

Each analytical measurement SOP lists the preventive maintenance schedule for each instrument which is to be followed by in-house and extramural repair contractors. In addition, each measurement group must maintain a log of all in-house and extramural preventive maintenance activities. Table 6.4.1 presents examples of general measures which are performed throughout the laboratory.

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Table 6.4.1 Laboratory Preventative Maintenance

GC/MS SYSTEMS			
EQUIPMENT	ACTION PERFORMED	FREQUENCY	
Hewlett-Packard 5995 GC/MS	Check oil level in mechanical pumps	Weekly	
	Check water level and operating condition in the Neslab cooling units	Weekly	
	Check compressed air gas supply	Daily	
	Check helium gas supply	Daily	
	Check carbon dioxide gas supply	Daily	
	Change the oil in the mechanical pumps	Every 6 months	
	Inspect the pump hoses and replace if required	Every 6 months	
	Change oil in the diffusion pump	Every 6 months	
	Change foreline and exhaust trap absorbent	Every 6 months	
	Inspect and refill the calibration sample vial with PFTBA	Every 6 months	
	Vacuum fan grills and filters	Every 6 months	
	Check fore and separator pump pressures	Weekly	
	Ion source cleaning and filament replacement	As needed	
	Column replacement and conditioning	As needed	
	Column cutting and reinstallation	As needed	
	Manual tuning	As needed	
	Change compressed air gas supply	As needed	
	Change helium gas supply	As needed	
	Change carbon dioxide gas supply	As needed	
	Recharge Neslab cooling units	As needed	
	Replace electron multiplier	As needed	
	Remove and clean or replace jet separator	As needed	

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Table 6.4.1 Laboratory Preventative Maintenance

EQUIPMENT	ACTION PERFORMED	FREQUENCY
Hewlett-Packard 5970 MSD / 5971 MSD/5972 MSD	Check oil level in mechanical pumps	Weekly
	Change the oil in the mechanical pumps	Every 6 months
	Inspect the pump hoses and replace if required	Every 6 months
	Change oil in the turbo pump	Every 6 months
	Change exhaust trap absorbent	Every 6 months
	Inspect and refill the calibration sample vial with PFTBA	Every 6 months
	Vacuum fan grills and filters	Every 6 months
	Ion source cleaning and filament replacement	As needed
	Manual tuning	As needed
	Replace electron multiplier	As needed
	Clean out transfer line to GC	After every column removal
Hewlett-Packard 5890 GC	Check helium gas supply	Daily
	Change split vent trap	Every 3 months
	Column replacement and conditioning	As needed
	Column cutting and reinstallation	Daily or as needed
	Change helium gas cylinder	As needed
	Change liner and septum	Daily or as needed
	Clean injection port	As needed

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Table 6.4.1 Laboratory Preventative Maintenance

EQUIPMENT	ACTION PERFORMED	FREQUENCY
Hewlett-Packard 7672A Autosampler	Inspect and correct injector alignment	After reseating
	Inspect syringe	Daily
	Check compressed air gas supply	Daily
	Inspect and adjust tension on sample tray	Daily
	Change rinse vials	Daily
	Change waste vials	Weekly
	Replace syringe	As needed
	Sand injector post	As needed
	Realign autosampler on brackets	As needed
	Change compressed air cylinder	As needed
Hewlett-Packard 7673 A Autosampler	Inspect syringe	Daily
	Inspect seating of injector	Daily
	Change rinse vials	Daily
•	Change waste vials	Weekly
	Replace syringe	As needed
	Reset control box #	As needed
Tekmar Purge and Trap Sample Concentrators and Autosamplers	Inspect spargers and fittings	Daily
	Check purge flow	Daily
	Inspect line and valve temperatures	Daily
	Change and condition trap	As needed
	Adjust purge flow	As needed
	Rinse or clean sparging vessels	As needed
	Rinse sample lines	As needed
	Bake out trap	After each analysis, extend as needed
	Replace lines and fittings	As needed
	Adjust line and valve temperatures	As needed

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Table 6.4.1 Laboratory Preventative Maintenance

EQUIPMENT	ACTION PERFORMED	FREQUENCY
Envirochem Air Sample Concentrator and Autosampler	Inspect fittings	Daily
	Check flows	Daily
	Inspect line and valve temperatures	Daily
	Change and condition internal traps	As needed
	Adjust flow	As needed
	Bake out trap	After each analysis, extend as needed
	Replace lines and fittings	As needed
	Adjust line and valve temperatures	As needed

	GC SYSTEMS					
EQUIPMENT	ACTION PERFORMED	FREQUENCY				
Hewlett-Packard 5890A GC (GC-1,4,5 Dual ECD)	Check gas supply	Daily				
	Check breakdown criteria	As required by run sequence				
	Vacuum filters and grills	Quarterly				
	Column replacement and conditioning	As needed				
	Column cutting and reinstallation	As needed				
	Change gas cylinders	As needed				
	Change liner and septum	As needed				
	Replace guard column	As needed				
	Clean injection port	As needed				
	Recondition ECD	As needed				
	Change ECD vent absorbent traps	Quarterly				

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Table 6.4.1 Laboratory Preventative Maintenance

EQUIPMENT	ACTION PERFORMED	FREQUENCY	
Hewlett-Packard 5890A GC (GC-3 FID/NPD)	Check gas supply	Daily	
	Vacuum filters and grills	Quarterly	
	Column replacement and conditioning	As needed	
	Column cutting and reinstallation	As needed	
	Change gas cylinders	As needed	
	Change liner and septum	As needed	
	Clean injection port	As needed	
	Replace or reactivate the NPD collector	As needed	
Hewlett-Packard 7673 A Autosampler	Inspect syringe	Daily	
	Inspect seating of injector	Daily	
	Inspect rinse and waste vials	. Daily	
	Vacuum filters and grills	Quarterly	
	Replace syringe	As needed	
	Change rinse and waste vials	As needed	

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Table 6.4.1 Laboratory Preventative Maintenance

EQUIPMENT	ACTION PERFORMED	FREQUENCY
Perkin-Elmer AS-100B Autosampler	Inspect syringe	Daily
	Inspect rinse and waste vials	Daily
	Check flushing efficiency	Daily
	Clean or replace syringe	As needed
	Change rinse and waste vials	As needed
	Change diverter valve septum	As needed

METALS SYSTEMS				
Graphite Furnace	Clean contact rings, furnace housing and quartz windows	Daily		
	Inspect, clean or replace graphite tubes	As needed		
	Replenish matrix modifiers	Daily		
	Check lamp alignments and energies	Daily		
	Clean mirrors for the optical sensors	Weekly		
	Clean windows on furnace housing	Weekly		
	Inspect contact rings for excessive wear	Monthly		
Inductively Coupled Plasma	Change capillary and pump tubing	Twice weekly		
	Replace liquid argon tank	As required		
	Reprofile via slit micrometer	Per manual		
	Replace and realign plasma torch	As needed		
	Clean nebulizer and spray chamber	As needed		
	Check primary imaging mirror	Weekly		
Mercury Analyzer	Clean sample cell and tubing	Monthly		
	Check sparger condition	Daily		
	Check level of mercury scrubber solution	Daily		
	Replace lamps	As required		

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WET CHEMISTRY SYSTEMS						
EQUIPMENT	ACTION PERFORMED	FREQUENCY				
pH Meters	Clean electrode if calibration has deteriorated	As needed				
	Store pH electrodes in pH 7.0 buffer	Daily				
	Check ISE electrodes and meter	Per manual				
Analytical Balances	Surfaces cleaned and covered	Daily				
	Calibrated and cleaned by manufacturer	Semi-annually				
	Accuracy checked by class "S" weights	Prior to use				
Conductivity Meters	Instrument surfaces inspected and cleaned	Daily				
	Calibrated using 0.01M potassium chloride	Daily				
	Spare cells on inventory	As needed				
Spectrophotometers	Instrument cleaned	Daily use				
Total Organic Halogen Analyzer (TOX)	Instrument cleaned	Daily use				
	Perform cell performance checks	Daily				
	Flush cells and check heated tapes	Daily				
	Inspect sample boats, inlet and exit tubes, o-rings and seals	Daily				
Autoanalyzer Systems	Clean all components and flush system	Daily use				
	Inspect all pump tubes and sample lines	Daily use				
	Inspect line coils, heating baths and filters	Weekly				
	Inspect all colorimeter filters	Weekly				
	Inspect and clean chemical manifolds	Monthly				

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7.0 DATA GENERATION

7.1 Introduction

There are numerous policies and standard procedures which have been implemented to ensure that data of known quality is continually generated by the IEA-CT laboratory. The IEA Corporate and Laboratory Facility Quality Assurance Plans are examples of documents which are generated. Guidelines for the facility QA plans are detailed in section 7.2.1 of the Corporate Quality Assurance Program Plan Doc#QAQ00102.NET.

7.2 Quality Assurance Project Plans

Quality Assurance Project Plans (QAPjP) are developed to meet contract and agency requirements on a project specific basis. These plans discuss specific terms, policies, objectives and QA activities designed to achieve the data quality objectives of the project.

All QA project plans are written in accordance with the following USEPA Document: <u>USEPA Guidelines and Specification for Preparing Quality Assurance Project Plans.</u> QAMS-005/80, Washington DC: USEPA, Quality Assurance Management Staff, October 17, 1980.

Guidelines for preparing QA project plans are also detailed in the Corporate Quality Assurance Program Plan Doc#QAQ00102.NET.

7.3 Methods

IEA-CT utilizes a wide variety of analytical methods. A listing of general analytical capabilities is presented in Table 7.3.1. Section 8 of the Appendix lists the analytical method and detection limits associated with various analytical procedures.

Each department is required to have a written standard operating procedure (SOP) in use which describes how the requirements of the method are met. All SOPs must be prepared in accordance with IEA Doc.#QAS00200.NET.

Analytical methodologies and quality assurance protocols in use are based on the following guidelines:

"Methods of Organic Chemical Analysis of Municipal and Industrial Wastewater", Federal Register Vol. 49, No. 209, October 26, 1984;

"Test Methods for Evaluating Solid Waste", SW-846 Third Edition, September 1986, USEPA, plus updates;

"Standard Methods for the Examination of Water and Wastewater" 1985, 14th, 15th and 16th Edition;

"Methods for Chemical Analysis of Water and Wastes" March 1983, EMSL, EPA;

"Manual of Analytical Methods for the Analysis of Pesticides in Humans and Environmental Samples", EPA 600/8-80-038, June 1980;

Organic Analysis: Multi-media, Multi-Concentration-IFB-CLP, January 1991, Document Number OLM01.9 (plus revisions);

Organic Analysis: Multi-media, Multi-Concentration-IFB-CLP, Document Number OLM03.1

Inorganic Analysis: Multi-media, Multi-Concentration-IFB-CLP, Document Number ILM03.0, ILM04.0;

"Handbook for Analytical Quality Control in Water and Wastewater Laboratories", EPA-600/4-79-019, March 1979;

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National Enforcement Investigation Center Policies and Procedures Manual, EPA-330/9/78/001-R, Revised May 1986

"Manual for the Certification of Laboratories Analyzing Drinking Water", April 1990, EPA/570/9-90/008.

"EML Procedures Manual", HASL-300, November 1990, 27th Edition.

"Prescribed Procedures for Measurement of Radioactivity in Drinking Water", EPA-600/4-80-032, August 1980.

"Health and Environmental Chemistry: Analytical Techniques, Data Management, and Quality Assurance" LA-10300-M, Vol. 1-3 Manual, Los Alamos National Laboratory, April 1992.

"Radiochemical Analytical Procedures for Analysis of Environmental Samples," EMSL-LV 0539-17.

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TABLE 7.3.1

IEA-CT ANALYTICAL CAPABILITIES

1. ORGANICS-GC/MS

Volatile Organics-524.2 Volatile Organics-CLP Volatile Organics-8240 Volatile Organics-8260 Volatile Organics-T01/T02 Volatile Organics-Appendix IX Acid & Base/Neutrals-8270 Acid & Base/Neutrals-CLP Acid & Base/Neutrals-Appendix IX Volatile Organics-624 Acid & Base/Neutrals-625

III. INORGANIC METALS

ICP Metals Furnace Metals **CLP Metals**

V. INORGANIC WET CHEMISTRY*

Acidity Alkalinity Ammonia Bicarbonate

Biochemical Oxygen Demand (BOD)

Bromide Chloride

Chlorine Demand Chlorine Residual

Chemical Oxygen Demand

Color Conductivity Chromium (VI) Cyanide - Amenable Cyanide - Total Cyanide (CLP) Dissolved Oxygen Flashpoint Fluoride Grain Size

Hydrocarbon analysis

MBAS Nitrate Nitrite Odor Oil and Grease Paint Filter Test ρН Phenols

II. ORGANICS-GC

Misc. DAI - 8015 Organohalide Pesticides & PCBs-608 Organohalide Pesticides & PCBs-8080 Organohalide Pesticides & PCBs-CLP Organophosphate Pesticides-8140 Organohalide Pesticides & PCBs-Appendix IX

Chlorinated Herbicides-8150

Chlorinated Herbicides-Appendix IX

Appendix IX Metals TCLP Metals Drinking Water Metals

Phosphate **Phosphorus** Settleable Solids Silica

Specific Gravity Sulfate Sulfide

Sulfite Sludge Volume Index

Tannins and Lignins Total Dissolved Solids Total Kjeldahl Nitrogen Total Organic Carbon Total Organic Halides **Total Solids**

Total Suspended Solids

Turbidity

Volatile Solids

Corrosivity Characteristics Ignitability Characteristics

EPTOX TCLP SPLP

Extactable Organic Halides

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			F QC REQUIREMENTS NIC ANALYSIS METH		
Requirement	Drinking Water Analysis Method 524	Water and Wastewater Analysis Method 624	RCRA Solid Waste Analysis Methods 8240/8260	Superfund Hazardous Waste Analysis CLP SOW OLM01.9	NYSDEC
Tuning Frequency Criteria	25 ng BFB 8 hrs See following page	50 ng BFB Daily See following page	50 ng BFB 12 hrs See following page	50 ng BFB 12 hrs See following page	50 ng BFB 12 hrs See following pa
Initial Calibration Maximum % RSD Minimum RRF	3-5 standards <20% NS	3 standards <35% NS	5 standards CCC <30%* SPCC > 0.250- 0.300*	5 standards <20.5%* 0.01-0.500*	5 standards <20.5%* 0.01-0.500* 10 compounds wi max 100% RSD
Continuing Calibration Frequency Maximum %D Minimum RRF IS Area	8 hrs ±30% NS ±30% of last CC or ±50% of IC	Daily QC Limits NS NS	12 hrs CCC ±25%* SPCC >0.250- 0.300* -50 to +100% of last CC	12 hrs ±25.0%* 0.01-0.500* NS	12 hrs ±25.0%* 0.01-0.500* NS 10 compounds wi max 100% RSD
QC Check Sample/LCS Frequency Criteria	Quarterly QC Limits*	Daily QC Limits*	Each batch or if MS % recovery not in QC limits QC Limits*	NS QC Limits*	Each cat. B SDO
Method Blank Frequency Criteria	Daily <mdl< td=""><td>Daily In control</td><td>12 hrs In control</td><td>12 hrs <crql*< td=""><td>12 hrs <crql*< td=""></crql*<></td></crql*<></td></mdl<>	Daily In control	12 hrs In control	12 hrs <crql*< td=""><td>12 hrs <crql*< td=""></crql*<></td></crql*<>	12 hrs <crql*< td=""></crql*<>
Spikes Frequency % Recovery	Blank spike Daily or 5% 80-120%	Matrix spike 5% QC Limits*	Matrix spike 5% QC Limits*	Matrix spike 5% or 1/SDG QC Limits*	Matrix spike 5% or 1/SDG QC Limits*
Duplicates Frequency Precision	BS duplicate Quarterly <20% RSD	Field duplicate NS NS	MS duplicate or sample duplicate 5% SD Limits*	MS duplicate 5% or 1/SDG RPD Limits*	MS duplicate MSB required* 5% or 1/SDG RPD Limits*
Sample Analysis Holding time Internal standards Criteria	14 days 1 @ 2-10 ug/L NS	14 days 3 @ 30 ug/L NS	14 days 3-4 @ 50 ug/L NS	10 days from receipt 3 @ 50 ug/L Area -50 to +100T RT ±30 sec	7 days from rece 3 @ 50 ug/L Area -50 to +10 RT ±30 sec
Surrogate Criteria Analyte ID	2 @ 5 ug/L 80-120% RT ±3x SD window 3 ions ±20%	3 @ 30 ug/L NS RT ±30 sec 3 ions ±20%	3 @ 50 ug/L See following page RRT ±0.06 Ions >10% ±20%	3 @ 50 ug/L See following page RRT ±0.06 Ions >10% ±20%	3 @ 50 ug/L See following pa RRT ±0.06 lons >10% ±20

^{*}For complete information refer to method or protocol

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SUMMARY OF VOLATILE SURROGATE RECOVERY LIMITS								
Compound	Method 524 (%)	Method 624 (%)	Method 8240 Water (%)	Method 8240 Soil (%)	Method 8260 Water (%)	Method 8260 Soil (%)	CLP SOW Water (%)	CLP SOW Soil (%)
4-Bromofluorobenzene	80-120	NS	86 <u>-1</u> 15	74-121	86-115	72-121	86-115	59-113
1,2-Dichloroethane-d4	80-120	NS	76-114	70-121	NS	NS	76-114	70-121
Toluene-d8	NS	NS	86-110	81-117	88-110	81-117	88-110	84-138
Dibromofluoromethane	NS	NS	NS	NS	86-118	80-120	NS	NS

SUMMARY OF VOLATILE SPIKE RECOVERY LIMITS					
Method Compound	Method 524 (%)	Method 624 (%)	Method 8240 (%)	CLP SOW Soil (%)	CLP SOW Water (%)
Benzene	80-120	37-151	37-151	66-142	66-142
Chlorobenzene	80-120	37-160	37-160	75-130	60-133
1,1-Dichloroethane	80-120	59-155	59-155	61-145	59-172
Toluene	80-120	47-150	47-150	76-125	59-139
Trichloroethene	80-120	71-157	71-157	71-120	62-137

SUMMARY OF INSTRUMENT TUNING REQUIREMENTS					
BFB Ion Abundance Criteria	NYSDEC 1/91	Method 524 (%)	Method 624 (%)	Methods 8240/8260 (%)	CLP SOW OLM01.8 (%)
50 - % of mass 95	15-40	15-40	15-40	15-40	8.0-40.0
75 - % of mass 95	30-60	30-80	30-60	30-60	33.0-66.0
95	100	100	100	100	100.0
96 - % of mass 95	5-9	5-9	5-9	5-9	5.0-9.0
173 - % of mass 174	_<2	<2	<2	<2	<2.0
174 - % of mass 95	>50	>50	>50	<50	50.0-120.0
175 - % of mass 174	5-9	5-9	5-9	5-9	4.0-9.0
176 - % of mass 174	95-101	95-101	95-101	95-101	95.0-101.0
177 - % of mass 176	5-9	5-9	5-9	5-9	5.0-9.0

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TABLE 7.3.2 SUMMARY OF QC REQUIREMENTS FOR EPA SEMI-VOLATILE ORGANIC ANALYSIS METHODS

<u></u>	Era Jenu-VO	LATILE ORGANIC ANAL	1313 METHODS	
Requirement	Water and Wastewater Analysis Method 625	RCRA Solid Waste Analysis Method 8270	Superfund Hazardous Waste Analysis CLP SOW OLM01.9	NYSDEC
Tuning Frequency Criteria	50 ng DFTPP Daily See following page	50 ng DFTPP 12 hrs See following page	50 ng DFTPP 12 hrs See following page	50 ng DFTPP 12 hrs See following page
Initial Calibration Maximum %RSD Minimum RRF	3 standards <35% NS	5 standards CCC <30%* SPCC >0.050*	5 standards <20.5%* 0.01-1.300*	5 standards <20.5%* 0.01-1.300* 20 compounds Max 100% RSD
Continuing Calibration Frequency Maximum %D Minimum RRF IS Area	Daily ±20% NS NS	12 hrs CCC ±30%* SPCC > 0.050* -50 to +100% of last CC	12 hrs ±25%* 0.01-1.300* NS	12 hrs ±25%* 0.01-1.300* NS 20 compounds Max 100% RSD
QC Check Sample/LCS Frequency Criteria	≤5% QC Limits*	If MS % recovery not in QC limits QC Limits*	Each SDG QC Limits*	Each SDG with cat. B QC Limits*
Method Blank Frequency Criteria	1 per batch In control	1 per batch In control	1 per batch <crql*< td=""><td>1 per batch < CRQL*</td></crql*<>	1 per batch < CRQL*
Spikes Frequency % Recovery	Matrix spike 5% QC Limits*	Matrix spike 5% QC Limits*	Matrix spike 5% or I/SDG QC Limits*	Matrix spike 5% or 1/SDG QC Limits*
Duplicates Frequency Precision	Field duplicates NS NS	MS duplicate or sample duplicate 5% SD Limits*	MS duplicate 5% or 1/SDG RPD Limits*	MS duplicate MS blank 5% or 1/SDG RPD Limits*
Sample Analysis Holding Time Water extraction	7 days	7 days	5 days from receipt	completed within 5 days
Soil extraction	NA	14 days	10 days from receipt	from receipt completed within 5 days from receipt
Analysis Internal standards Criteria	40 days from extraction 3 NS	40 days from extraction 6 @ 40 ug/L NS	35 days from extraction 6 @ 20 ug/L Area -50/+100% RT ±30 sec	35 days from extraction 6 @ 20 ug/L Area -50/+100% RT ±30 sec
Surrogate Criteria Analyte ID	3 @ 100 ug/L NS RT ±30 sec 3 ions ±20%	6 @ 100-200 ug/L See following page RRT ±0.06 Ions >10% ±20%	8 @ 100-150 ug/L See following page RRT ±0.06 Ions > 10% ±20%	8 @ 100-150 ug/L See following page RRT ±0.06 Ions > 10% ±20%

^{*}For complete information refer to method or protocol

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SUMMARY OF SEMI-VOLATILE SURROGATE RECOVERY LIMITS					
Compound	Method 625	Method 8270 Water (%)	Method 8270 Soil (%)	NYSDEC '91 ASP CLP SOW Water (%)	NYSDEC '91 ASP CLP SOW Soil (%)
Nitrobenzene-d5	NS	35-114	23-120	34-114	23-120
2-Fluorobiphenyl	NS	43-116	30-115	43-116	30-115
p-Terphenyl-d14	NS	33-141	18-137	33-141	18-137
Phenol-d6	NS	10-94	24-113	10-110	24-113
2-Fluorophenol	NS	21-100	25-121	21-110	25-121
2,4,6-Tribromophenol	NS	10-123	19-122	10-123	19-122
1,2-Dichlorobenzene-d4	NS	NS	NA NA	16-110*	20-130*
2-Chlorophenol-d4	NS	NS	NA	33-110*	20-130*
Perylene-d12	NS	NS	NA	NA	NA

SUMMARY OF SEMI-VOLATILE SPIKE RECOVERY LIMITS						
Compound	Method 625 (%)	Method 8270 (%)	NYSDEC '91 ASP' CLP SOW Water (%)	NYSDEC '91 ASP CLP SOW Soil (%)		
Acenaphthene	47-145	47-145	46-118	31-137		
1,4-Dichlorobenzene	20-124	20-124	36-97	28-104		
2,4-Dinitrotoluene	D-112	D-112	24-96	28-89		
N-Nitroso-di-n-propylamine	D-230	D-230	41-116	41-126		
Pyrene	52-115	52-115	26-127	35-142		
1,2,4-Trichlorobenzene	44-142	44-142	39-98	38-107		
4-Chloro-3-methylphenol	22-147	22-147	23-97	26-103		
2-Chlorophenol	23-134	23-134	27-123	25-102		
4-Nitrophenol	D-132	D-132	10-80	11-114		
Pentachlorophenol	14-176	14-176	9-103	17-109		
Phenol	5-112	5-112	12-110	26-90		

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SUMMARY OF GC/MS INSTRUMENT TUNING REQUIREMENTS					
DFTPP Ion Abundance Criteria	NYSDEC	Method 525 (%)	Method 625 (%)	Method 8270 (%)	CLP SOW (%)
51 - % of mass 198	30-60	10-80_	30-60	30-60	30.0-80.0
68 - % of mass 69	<2	<2	<2	<2	<2.0
70 - % of mass 69	<2	<2	<2	<2	Present
127 - % of mass 198	40-60	10-80	40-60	40-60	25.0-75.0
197 - % of mass 198	<1	<2	<1	<1	<1.0
198	100	100	100	100	100
199 - % of mass 198	5-9	5-9	5-9	5-9	5.0-9.0
275 - % of mass 198	10-30	10-60	10-30	10-30	10.0-30.0
365 - % of mass 198	>1	>1	>1	>1	>0.75
441	< mass 443	< mass 443	< mass 443	< mass 443	< mass 443
442 - % of mass 198	40-110	> 50	>40	>40	40.0-110.0
443 - % of mass 442	17-23	15-24	17-23	17-23	15.0-24.0

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TABLE 7.3,2 SUMMARY OF QC REQUIREMENTS FOR PESTICIDE/PCB ANALYSIS METHODS					
Requirement	Water and Wastewater Analysis Method 608	RCRA SW-846 Solid Waste Analysis Method 8080	Superfund Hazardous Waste Analysis CLP SOW OLM01.8	NYSDEC	
Initial Calibration	3 standards	5 standards	3 standards (1 for multicomponent)	3 standards (1 for multicomponent)	
Maximum % RSD DDT/Endrin Breakdown Resolution	<10% NS NS	<20% <20% NS	<10.0-15.0%* <20.0% 90-110%*	<10.0-15.0%* <20.0% 90-110%*	
Continuing Calibration Frequency Maximum %D RT Criteria	Mid-level standard Daily ±15% NS	Mid-level standard Daily ±15% NS	Mid-level standard 12 hrs ±25.0% ±0.05-0.07 min of mean RT	Mid-level standard 12 hrs ±25.0% ±0.05-0.07 min of mean RT	
QC Check Sample/LCS Frequency Criteria	≤10% QC Limits*	If MS % recovery not in QC limits QC Limits*	NS QC Limits*	Each cat. B SDG QC Limits*	
Method Blank Frequency Criteria	1/batch In control	1/batch In control	1/batch < CRQL	1/batch < CRQL	
Spikes Frequency % Recovery	Matrix spike 10% QC Limits*	Matrix spike 5% QC Limits*	Matrix spike 5% or 1/SDG QC Limits*	Matrix spike 5% or 1/SDG QC Limits*	
Duplicates Frequency Precision	Field duplicate NS NS	MSD or sample duplicate 5% SD Limits*	MSD 5% or 1/SDG RPD Limits*	MSD/MS Blank 5% or 1/SDG RPD Limits*	
Sample Analysis Holding Time Water extraction	7 days	7 days	5 days VTSR	Completed within 5	
Soil extraction	NA	14 days	10 days VTSR	days VTSR Completed within 5 days VTSR	
Analysis Analyte ID	40 days RT within 3x SD of std. RT window	40 days RT within 3x SD of std. RT window	35 days RT ±0.05-0.07 min of std. RT on both columns; Conc. ±25.0%	35 days RT ±0.05-0.07 min of std. RT on both columns;	
Confirmation	2nd column for unknown samples	2nd column for positive ID	2 column required; GC/MS if > 10 ng/uL	Conc. ±25.0% 2 column required; GC/MS if >10 ng/uL	

^{*}For complete information refer to method or protocol

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SUMMARY OF PESTICIDE SURROGATE RECOVERY LIMITS					
Compound	Method 508 (%)	Method 608	Method 8080	CLP SOW (%)	NYSDE C
Tetrachloro-m-xylene	NS	NS	Lab limits	60-150	60-150
Decachlorobiphenyl	NS	NS	Lab limits	60-150	60-150
Dibutylchlorendate	NS	NS	Lab limits	NS	

SUMMARY OF PESTICIDE SPIKE RECOVERY LIMITS					
Compound	Method 608 (%)	Method 8080 (%)	CLP SOW Water (%)	CLP SOW Soil (%)	NYSDEC
gamma-BHC (Lindane)	19-140	19-140	56-123	46-127	46-127
Aldrin	42-122	42-122	40-120	34-132	34-132
Dieldrin	36-146	36-146	52-126	. 31-134	31-134
4,4'-DDT	25-160	25-160	38-127	23-134	23-134
Endrin	30-147	30-147	56-121	42-139	42-139
Heptachlor	34-111	34-111	40-131	35-130	35-130

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TABLE 7.3.2 SUMMARY OF QC REQUIREMENTS FOR EPA METALS ANALYSIS METHODS USING ATOMIC ABSORPTION (AA) SPECTROSCOPY

Requirement	Water and Wastewater Analysis Method 200.0	RCRA Solid Waste Analysis Method 7000	Superfund Hazardous Waste Analysis CLP SOW ILM03.0
Initial Calibration Frequency Criteria	3 standards and a blank Daily r ≥0.995	3 standards and a blank Daily r ≥0.995	3 standards and a blank Daily or every 24 hrs r ≥0.995
Calibration Verification Frequency Criteria	A standard at or near MCL After initial calibration and every 20 samples 90-110% recovery	Mid-range standard Every 10 samples ICV: 90-110% recovery CCV: 80-120% recovery	Mid-range standard Beginning, end, and every 10 samples or every 2 hrs 90-110% recovery Hg: 80-120% recovery
Detection Limits Standard Frequency Criteria	NS NS NS	NS NS NS	Standard at the CRDL or IDL Beginning of each sample run EPA QC limits
Calibration Blanks Frequency Criteria	After each calibration	After each calibration	Beginning, end, and every 10 samples or every 2 hrs All analytes ≤ CRDL
Preparation Blanks Frequency Criteria	Each digestion batch NS	Each digestion batch NS	1 per SDG or digestion batch All analytes ≤ CRDL
QC Check Sample/LCS .:Frequency - Criteria	NS NS	1 per batch NS	1 per matrix per SDG or digestion batch 80-120% recovery
Matrix Spike Samples Frequency Criteria	10% or 1 per batch NS	5% or 1 per batch NS	5% or 1 per SDG per matrix per level (predigestion) 75-125% recovery
Duplicate Samples Frequency Criteria	10% or 1 per batch NS	5% or 1 per batch NS	5% or 1 per SDG per matrix per level (predigestion) ≤20% RPD for values ≥5 x CRDL ±1 x CRDL for values <5 x CRDL
Furnace Quality Control Frequency Criteria	MSA as needed	MSA as needed Serial dilution: 1 per batch per matrix MSA: r≥0.995 5¥ dilution within ±10%	Duplicate injections on all; Post digestion spikes on all samples, blanks, and LCS; MSA as needed Duplicate injections: ≤20% RSD/CV Spikes: 85-115% recovery MSA: r≥0.995

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TABLE 7.3.2 SUMMARY OF QC REQUIREMENTS FOR EPA METALS ANALYSIS METHODS USING INDUCTIVELY COUPLED PLASMA (ICP) SPECTROSCOPY					
Requirement	Water and Wastewater Method 200.7	RCRA Solid Waste Analysis Method 6010	Superfund Hazardous Waste Analysis CLP SOW ILM03,0		
Initial Calibration Frequency	1 standard and a blank Daily	1 standard and a blank Daily	1 standard and a blank Daily or every 24 hrs		
Calibration Verification Frequency Criteria	Mid-range standard Every 10 samples 95-105% recovery	Mid-range standard Every 10 samples and at end 90-110% recovery	Mid-range standard Beginning, end, and every 10 samples or every 2 hours 90-110% recovery		
Other Standards Frequency	Highest mixed standard Before sample analyses	Highest mixed standard Before sample analyses	Standard at 2 x CRDL or IDL Beginning and end of each run or 2 every 8 hrs		
Criteria	95-105% recovery	95-105% recovery	EPA QC Limits		
Interference Check Sample Frequency	Beginning, end, and periodic intervals	Beginning and end of each run or every 8 hours 80-120% recovery	Beginning and end of each run or 2 every 8 hrs 80-120% recovery		
Criteria	±1.5 x SD of mean value				
Calibration Blanks Frequency	Every 10 samples	Every 10 samples and at end	Beginning, end, and 10% of samples		
Criteria	±2 x SD of mean value	±3 x SD of mean value	or every 2 hrs All analytes ≤ CRDL		
Preparation Blanks Frequency Criteria	1 per batch NS	1 per batch NS	1 per SDG or digestion batch All analytes ≤CRDL		
QC Check Sample/LCS Frequency Criteria	Each IC and weekly 95-105% recovery	Each IC and weekly 90-110% recovery	1 per SDG or digestion batch for each matrix 80-120% recovery		
Matrix Spike Samples Frequency Criteria	1 every new sample matrix 90-110% recovery	5% or 1 per batch 75-125% recovery	5% or 1 per SDG per matrix per level (predigestion) 75-125% recovery		
Duplicate Samples Frequency Criteria	NS NS	5% or 1 per batch ≤20% RPD for values >10 x	5% or 1 per SDG per matrix per level (predigestion) ≤20% RPD for values ≥5 x CRDL		
		IDL	±1 x CRDL for values <5 x CRDL		
Serial Dilution Frequency Criteria	1 every new sample matrix Dilution within ±5%	1 every new sample matrix 4 x dilution within ±10%	1 per SDG per matrix per level 5 x dilution within ±10%		

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TABLE 7.3.2 SUMMARY OF QC REQUIREMENTS FOR EPA MERCURY ANALYSIS METHODS USING COLD VAPOR ATOMIC ABSORPTION (AA) SPECTROSCOPY						
Requirement	Water and Wastewater Analysis Method 245.1/245.5	RCRA Solid Waste Analysis Method 7470/7471	Superfund Hazardous Waste Analysis CLP SOW ILM03.0			
Method Detection Limit	0.2 ug/L	0.2 ug/L	CRDL: 0.2 ug/L			
Holding Time	28 days	28 days	26 days			
Initial Calibration Frequency Criteria	6: blank and 5 standards Daily r ≥0.995	6: blank and 5 standards Daily and every hour of analysis r ≥0.995	5: blank and 4 standards Daily or every 24 hours r ≥0.995			
Calibration Verification Frequency	A standard at or near MCL After initial calibration and every 20 samples	Mid-range standard Every 10 samples	Independent standard CCV: diff. conc. than ICV, or at near the mid-range ICV: After initial calibration CCV: 10% or every 2 hours			
Criteria	90-110% Recovery	80-120% Recovery	80-120% Recovery			
Calibration Blanks Frequency Criteria	After each calibration	After each calibration	Beginning, end, and every 10 samples or every 2 hours CRDL			
Preparation Blanks						
Frequency Criteria	1 per digestion batch NS	1 per digestion batch NS	1/SDG/digestion batch ≤CRDL			
QC Check Sample/LCS Frequency	Blind performance sample 1 per year (Optional: 1 per quarter).	Independent standard Every 15 samples	EPA standard 1/SDG/batch (solid samples only)			
Criteria	EPA control limits	80-120% Recovery	80-120% Recovery			
Matrix Spike Samples Frequency Criteria	NS NS	5% or 1 per batch NS	5% or 1/SDG/matrix/level 75-125% Recovery			
Duplicate Samples Frequency Criteria	10% or 1 per batch EPA control limits	5% or 1 per batch NS	5% or 1/SDG/matrix/level ≤20% RPD			
Other Method Criteria Frequency Criteria	Method of standard addition As needed NS	Method of standard addition As needed NS	Standard at the CRDL or IDL Beginning of each sample run EPA control limits			

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TABLE 7.3.2 SUMMARY OF QC REQUIREMENTS FOR EPA CYANIDE ANALYSIS METHODS						
Requirement	Water and Wastewater Analysis Method 335.4	RCRA Solid Waste Analysis Method 9012	Superfund Hazardous Waste Analysis CLP SOW ILM03.0			
Method Detection Limit	Titration: 1 mg/L Colorimetric: 0.02 mg/L	Titration: 0.1 mg/L Colorimetric: 0.02 mg/L	CRDL: 10 ug/L			
Holding Time	14 days (24 hours when sulfide is present)	14 days	12 days from sample receipt			
Initial Calibration ⁽¹⁾	6 standards and a blank	6 standards and a blank	3 standards and a blank (one standard at the CRDL)			
Frequency	Daily_	Daily	Daily			
Calibration Verification ⁽¹⁾ Frequency	NS NS	Mid-range standard Every 15 samples 85-115% Recovery	CCV: Mid-range standard Beginning, end, and every 10 samples or 2 hours 85-115% Recovery			
Criteria	143	83-113% Recovery	85-115 % Recovery			
Other Standards (Distilled) Frequency Criteria	High and low standard 1 each per batch 90-110% Recovery	High and low standard 1 each per batch 90-110% Recovery	Mid-level standard 1 per batch 85-115% Recovery			
Calibration Blanks Frequency Criteria	Colorimetric: 1 per batch Use in initial calibration	Colorimetric: 1 per batch Use in initial calibration	Colorimetric: Beginning, end and every 10 samples or 2 hours			
Preparation Blanks Frequency Criteria	Titration: 1 per batch Colorimetric: Not specified Titration: Use in calculation Colorimetric: Not specified	Titration: 1 per batch Colorimetric: Not specified Titration: Use in calculation Colorimetric: Not specified	Titration: 1 per batch Colorimetric: 1 per batch Titration: Use in calculation Colorimetric: CRDL			
Laboratory Control Standard Frequency Criteria	NS NS NS	Independent check standard 1 per batch 85-115% Recovery	Distilled independent standard (ICV) 1 per batch 85-115% Recovery			
Matrix Spike Samples Frequency Criteria	1 per batch to check distillation efficiency NS	Matrix spike and matrix spike duplicate per batch NS	1 per matrix per concentration level per batch 75-125% Recovery			
Duplicate Samples Frequency Criteria	NS NS	1 matrix spike duplicate per batch NS	1 per matrix per concentration level per batch ≤20% RPD for values >5 x CRDL			
Other Method Criteria	Verify sample pH ≥12; Check for oxidizing agents and sulfides	Verify sample pH ≥12; Check for oxidizing agents and sulfides	Verify sample pH ≥12; Check for oxidizing agents and sulfides			

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KEY TO CHART				
BS	Blank Spike			
cc	Continuing Calibration			
ccc	Calibration Check Compounds			
CCV	Continuing Calibration Verification			
CRDL	Contract-Required Detection Limit			
CRQL	Contract-Required Quantitation Limit			
CV	Coefficient of Variation			
D	Detected			
IC	Initial Calibration			
ICV	Initial Calibration Verification			
IDL	Instrument Detection Limit			
IS	Internal Standard			
LCS	Laboratory Control Sample			
MCL	Maximum Contaminant Level			
MDL	Method Detection Limit			

KEY TO CHART			
MS	Matrix Spike		
MSA	Method of Standard Additions		
NA	Not Applicable		
NS	Not Specified		
%D	Percent Difference		
%Rec.	Percent Recovery		
PQL	Practical Quantitation Limit		
r	Correlation Coefficient		
RF	Response Factor		
RPD	Relative Percent Difference		
RRT	Relative Retention Time		
RSD	Relative Standard Deviation		
RT	Retention Time		
SD	Standard Deviation		
SDG	Sample Delivery Group		
SPCC	System Performance Check Compounds		

NOTES

- (1) Calibration standards must be distilled for EPA Methods 335.4 and 9012 when sulfides are present in the samples.
- ⁽²⁾ CLP SOW specifies that the initial calibration verification standard (ICV) be distilled and analyzed as the laboratory control standard (LSC).

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7.4 Standard Operating Procedures

All laboratory activities, from sample receipt to analysis to final report generation, must adhere to the laboratory Standard Operating Procedures (SOPs) which have been developed to provide quality environmental data with adequate documentation to be of known quality and hence of maximum use to our clients. All SOPs provide complete documentation as to how each sample is measured for each parameter. Reference corporate document QAS00200.NET for the IEA corporate format for generating SOPs. Each SOP shall have a unique code in accordance with the IEA corporate document control procedure as outlined in the corporate SOP on document control.

On a regular basis the QA Manager will review data to check for compliance to SOPs. Additionally the QA Manager will review SOPs to ensure they meet the requirements of the methodologies and applicable regulations. If it is found that the document does not meet the requirements, the discrepancy is forwarded to the group/section leader through the corrective action process. (reference SOP on Corrective Action Reports -QAS00501.CT).

In addition to method SOPs, at minimum the laboratory is required to have on file SOPs for the following operations. Many of these SOPs have been generated by the IEA corporate QA department.

Sample Receipt, Logging and Disposal
Chain-of-Custody Procedures
Sample Storage
Security of Samples and Laboratory Facility
Purity of Standards and Standards Preparation Documentation
Maintaining Laboratory Records and Logbooks
Sample Analysis and Data Control Systems
Sample Bottle and Glassware Cleaning Procedures
Laboratory Waste Disposal

An example listing of laboratory SOPs is presented in Section 7 of the Appendix. A complete list of all laboratory SOPs is available upon request.

7.5 Chain-of-Custody

Samples are physical evidence and are handled at IEA according to certain procedural safeguards. For the purposes of legal proceedings, a demonstration to the court that the laboratory is a secure area may be all that is required for the analyzed evidence to be admitted. However, in some cases, the court may require a presentation of the hand-to-hand custody of the samples while they were at the laboratory. In the event that a client requires such a comprehensive chain-of-custody demonstration, upon special request, IEA is capable of producing documentation that traces the in-house custody of the samples from the time of receipt to completion of analysis.

The National Enforcement Investigations Center (NEIC) of EPA defines custody of evidence in the following ways:

- It is in your actual possession; or
- It is in your view, after being in your physical possession; or
- It was in your possession and then you locked or sealed it up to prevent tampering; or it
 is in a secure area

At IEA-CT, chain of custody begins with shipment of the sample bottles and coolers. IEA-CT has a printed external chain-of-custody form that accompanies each sample shipment. An example of this form is found in Section 2 of the appendix.

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Upon receipt of the samples in the laboratory the sample custodian and the sample control group are responsible for obtaining all necessary shipping documentation and verification of all data entered into the laboratory sample custody records. The internal chain of custody form is generated at this point.

All samples and projects entering the laboratory are identified with a job/project number. Individual samples are then identified using the job number and sample counter. The samples are then stored according to the requirements of the analytical protocols (refrigeration).

Preliminary sample receipt notifications are distributed to each department to notify department of sample arrival and facilitate the analysis of parameters with short holding times. Each department has a system of tracking sample analysis throughout their respective departments.

All documentation received with samples is reviewed by the sample custodian at the time of receipt. The project manager then reviews the paperwork again at the time of log-in to the LIMS computer system. If there are any discrepancies noted by the sample custodian, a corrective action report is filled out and submitted to the project manager. The client is then contacted for resolution.

The specific procedures and requirements for receiving samples are specified in the SOP for sample control - "Sample Processing Methods Performed at Sample Arrival" (Doc# SMS00401.CT). IEA's chain-of-custody record is designed to meet the legal requirements of federal, state and local government agencies and the courts of law. The record covers:

- Labeling of sample bottles, packing the shipping container and transferring the shipping container under seal to the custody of a shipper;
- · Outgoing shipping manifests;
- The chain-of-custody form completed by the person(s) breaking the shipping container seal, taking the sample, resealing the shipping container and transferring custody to a shipper;
- Incoming shipping manifests;
- · Breaking the shipping container's reseal;
- Storing each labeled sample bottle in a secured area;
- Disposition of each sample to an analyst or technician; and
- The use of the sample in each bottle in a testing procedure appropriate to the intended purpose of the sample.

For each link in this process the records indicate the following:

- The person with custody; and
- The time and date each person accepted or relinquished custody.

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IEA has implemented the following standard operating procedures with regard to laboratory chain-of-custody:

- Samples are stored in a secure area:
- Non-employee access to the laboratories are controlled through the use of limited access
 points at each facility. Outside personnel can access the facility either through the front
 receptionist or the sample receipt area. Other access doors to the laboratory are
 maintained in a secure manner at all times;
- All visitors to each facility are required to sign-in at the reception area and must be
 escorted by an IEA representative at all times while in the laboratory;
- · Refrigerators, freezers, and other sample storage areas are kept locked, when not in use;
- The designated sample custodian and supervisory personnel control access to the sample storage area(s); and
- Samples remain in secured sample storage until removed for sample preparation or analysis; and
- Upon special request, all transfers of samples into and out of storage are documented through an internal chain-of-custody procedure. This procedure is not normally employed in daily operations but is available upon special request by the client.

7.6 Analytical Calibration Standards

The calibration standards used for instruments and equipment are described in the specific analytical methods, or instrument manufacturers' operational guides. All standard preparations are recorded in a bound "Standards Preparation Log Book" with the lot number, method of preparation, date and analyst's initials. This log provides the internal documentation which traces the internal working standards to primary and secondary (purchased) stocks.

The stock solutions are all kept in a daily monitored 40 C refrigerator with the exception of the organic stock solutions which are kept in a 00 C freezer. Stock calibration standards are coded in the "Prep Log" mentioned above with the analyte, concentration, date prepared, initials, and referenced to the book and page where a description of the preparation can be found and traced. No samples are maintained in the same areas as the standards.

Records on the traceability of the standards are maintained in the office of the Quality Assurance Manager. These records include sources, dates of receipt, lot numbers (if Applicable) and expiration dates (if applicable).

Table 7.6.1 provides an overview of the standard sources, types and preparation by instrument group.

Metals Calibration Standards

Commercially available at 1000 ppm levels from Inorganic Ventures and prepared from primary standard material traceable to EPA A2LA standards. Stock standards solutions are prepared every six months or when needed as multi-element stocks.

Inorganic Calibration Standards

Most calibration standards described in the methodology used ACS Reagent Grade materials. Some reference materials are available from NIST to standardize titrating solutions. Stock solutions are prepared every three months while diluted working standards are prepared daily at the time of analysis. Spike solution preparation is also documented in the solution/standard log book.

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Organic Calibration Standards

Pure compounds for organic calibration materials are available through EPA EMSL in Cincinnati, EPA in Research Triangle Park, EPA Las Vegas, Supelco, Inc., Restek, Inc. and Accustandard, Inc. Organic stocks are prepared every six months and diluted working standards are prepared weekly. Stock non-volatile solutions can be prepared every six months and diluted working standards are prepared weekly. Stock non-volatile solutions can be prepared every six months with working standards made weekly. Organic spike solutions are prepared from neat solutions and documented.

pH Calibration Standards

Calibration materials which are certified by the manufacturer to be standardized against NIST Standards are commercially available and are used by the laboratory. Three standards - 4,7, and 10 are used daily to calibrate the pH meters.

Weighing Calibration Standards

Analytical balances are certified annually. Calibration is performed on a weekly or daily basis using class "S" weights (0.50, 5.00, and 50g).

Oven Calibration Standards

Daily calibration by monitoring oven temperature with a thermometer calibrated annually with a NIST Certified Thermometer.

Conductivity Calibration Standard

Conductivity solutions are described in Standard Methods, 15th edition, Section 502.

Turbidity Standards

Formazin solution prepared from CMS neat standard according to EPA Method 180.1-2. Four standards are used to prepare a calibration curve and are made fresh daily. The stock formazin standard is prepared every three months and kept under refrigeration.

Photometer Calibration Standard

Spectronic Standards - Catalog #331-31-50 (wavelength calibration).

Refrigerators

All refrigerators are checked daily for temperature stability. Yearly, the refrigerator thermometers are calibrated against an NIST thermometer. Daily readings are recoreded in a bound logbook.

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TABLE 7.6.1 STANDARD SOURCES AND PREPARATION						
Source	Form Received	Storage	Preparation from Source	Laboratory Stock Storage	Preparation Frequency	
Restek, Inc. EPA Supelco	Neat Solutions >	Frozen	Primary stocks are prepared from source stocks	Frozen	Semi-annual	
Accustandard	F F		Intermediate stocks are prepared from primary or source stocks	Refrigerator	Weekly	
			Working stocks are prepared from intermediates	N/A	Weekly	
Restek, Inc. EPA RTP	Neat Solutions	Frozen	Primary stocks are prepared from source stocks	Frozen	Semi-annual	
Accustandard	2 Tood ppiii	l rozen	Intermediate stocks are prepared from primary or source stocks	Refrigerator	Semi-annually	
			Working stocks are prepared from intermediates	N/A	Semi-annually	
Inorganic Ventures	Solutions of 1000ppm	Room temp.	Primary stocks (1 - 10 ppm) are prepared from source	O.15% HNO, at room temperature	Annually	
		li I	Intermediate stocks (1ppb - 1 ppm)	0.15% HNO ₃ at room temperature	Semi-annually or as needed	
			Working stocks	0.15% HNO ₃ at room temperature	Daily	
	Restek, Inc. EPA Supelco Accustandard Restek, Inc. EPA RTP Supelco Accustandard	Restek, Inc. EPA Supelco Accustandard Restek, Inc. EPA RTP Supelco Accustandard Neat Solutions > 1000 ppm Neat Solutions > 1000 ppm	Restek, Inc. EPA Supelco Accustandard Restek, Inc. EPA RTP Supelco Accustandard Restek, Inc. EPA RTP Supelco Accustandard Neat Solutions > 1000 ppm Frozen Frozen Frozen Frozen Frozen Frozen Solutions > 1000 ppm Frozen Frozen Restek, Inc. EPA RTP Supelco Accustandard Neat Solutions > 1000 ppm Frozen Frozen	Restek, Inc. EPA Supelco Accustandard Restek, Inc. EPA Solutions > 1000 ppm Restek, Inc. EPA RTP Supelco Accustandard Restek, Inc. EPA RTP Supelco Accustandard Restek Solutions > 1000 ppm Restek, Inc. EPA RTP Supelco Accustandard Restek Solutions > 1000 ppm Restek Solutions S	Restek, Inc. EPA Solutions > 1000 ppm Frozen Frozen Intermediate stocks are prepared from source stocks N/A	

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7.7 Instrument Calibration Procedures

The proper calibration of instrumentation and equipment is a key element in the quality of the analysis done by the laboratory. Each type of instrumentation and each EPA approved method has specific requirements for the calibration procedures, depending on the analytes of interest and the medium of the sample.

- Tables 7.7.1 list in tabular form the procedures which are followed by IEA Connecticut. The calibration protocols meet or exceed the minimum method criteria requirements. If a method calibration requirement is more stringent than those listed in the Quality Assurance Plan, the more stringent will be followed in each case.
- Documentation and records on calibrations are maintained in instrument logs and also with the data sets of the samples which are analyzed and related to them. In addition, laboratory department managers monitor the results of the calibration program to ensure the proper implementation at the analyst level.

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Analysis Col #Standards Type of curve Acceptance/rejection Fe						
Analysis	Cal. Type	# Standards	Type of curve	Acceptance/rejection criteria	Frequency	
GC Pesticides Herbicides OP pesticides	Initial	5 concentration levels	Lincar	≤ 20% RSD	continuing calibration fails	
Or positions	Continuing	1 standard (mid)		+/- 15% Difference	Daily and every 10 samples	
GC/MS quadrupole	Initial	5 concentration levels; tuning with BFB/DFTPP	Linear; tuned to manufacturer's specifications	≤ 20% RSD	continuing calibration failure	
	Every 12 hours	1 standard; tuning with BFB/DFTPP		+/- 15% Diff	Daily	
AAS Graphite	Initially .	5 concentration levels	Linear	> .995 coefficient of variation	continuing calibration failure	
	Continuing	1 standard 		+/- 95% of value	Every 10 samples	
ICP	Initially Daily	5 concentration levels 2 levels	Linear	According to instrument manufactures's instructions	Quarterly	
23331	Continuing	1 standard			Every 10 samples	
Lachat Analysis	Initially, Daily	5 concentration levels	Linear	<.995 coefficient of variation	continuing calibration failure	
	Continuing	1 standard			Every 10 samples	
pH Meters	Initially and daily	2 standards (pH 7 and 4 or 10)	Linear	+/- 95% of value	Daily	
	Continuing	1 standard			Every 10 samples	

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TABLE 7.7.1 INSTRUMENT CALIBRATION SUMMARY						
Analysis	Cal. Type	# Standards	Type of curve	Acceptance/rejection criteria	Frequency	
Spectrophoto- meter	Initially and daily	5 concentration levels plus set %T with no cuvette in holder	Linear	<.995 coefficient of variation	Daily	
	Continuing	1 standard		+/- 95% of value	Every 10 samples	
Infrared Spectrophoto- meter	Initially and monthly	5 concentration levels	Linear	<.995 coefficient of variation	Daily	
	Continuing	1 level		+/- 95% of value	Every 10 samples	
Conductivity meter	Daily	3 concentration levels	Linear	<.995 coefficient of variation	Daily	
	Continuing	3 concentration levels		+/- 95% of value	Every 10 samples	
Turbidimeter	Daily	3 concentration levels	Linear	<.995 coefficient of variation	Daily	
	Continuing	3 concentration levels		+/- 95% of value	Every 10 samples	
Balance	Daily	3 levels Class "S" weights	Point		Check single weight upon use	

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8.0 DATA PROCESSING

8.1 Introduction

Data processing is defined as the mechanisms employed for collecting, reviewing, transcribing, reporting and storing of analytical data and related information.

Because of the critical relationship between instrument calibration, the accuracy of the analytical data generated, and specific method protocols that determine data quality, IEA maintains strict controls on the calibration procedures for the various types of analytical equipment. Each type of instrumentation is calibrated prior to sample analysis according to method criteria. Specific criteria for the instrument calibrations must be met before samples may be processed. Corrective action must be taken to remedy any out of control situations.

8.2 Collection

Data in the environmental laboratory make take several forms. Some are manually generated, while others are automated computer outputs. Some examples of typical data are:

Field measurements or observations made on-site during the sample collection effort as part of a monitoring program.

Information provided on chain-of-custody forms such as sampler, sampling date, sample location, sample identification, weather observations and custody transfer information.

Recordkeeping information such as instrument run logs, standards traceability, sample preparation logbooks and balance calibrations which represent information not normally required for inclusion in client reports.

Analytical data produced by various instrumentation such as GC/MS units, gas chromatographs, atomic absorption spectrophotometers, and automated analyzers. This includes various associated outputs such as chromatograms, strip chart recordings and computer tape readouts.

Records of standard calibration curves as well as associated quality control data such as method blanks, matrix spikes, matrix spike duplicate, replicate and QC check samples.

Consistent data collection is achieved through the existence and use of standard operating procedures at each facility. For example, chain-of-custody forms are routinely checked for completeness and if omissions occur, the sampler is contacted for the missing information.

Laboratory data sheets or logbooks have a standard format to ensure that all pertinent information is recorded consistently. These items are regularly monitored to ensure compliance with established requirements.

Outputs from all instruments are monitored for readability and consistency. If clarity is less than desired, corrective actions are undertaken to rectify the output based on instrument manufacturers' recommendations.

The following sections will describe the general procedures which are employed at the IEA-CT laboratory. More specific detail can be found in the standard operating procedures.

Gas Chromatography

Data from the Gas Chromatographs is collected through interfaces and processed by a Hewlett Packard computer system (HP-1000) with RTE-A operating system and 3550A LAS software or and HP Chemstation with Enviroquant software. Data is reviewed at the bench level by the analyst. If all required QC is met then the data is reviewed for chromatographic scaling and dilutions. If necessary reintegrations and rescalings

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are done using the LAS system or Enviroquant software. The binary result files are then converted to ASCII report files for transfer to the Seedpak system for data report forms generation.

GC/Mass Spectrometry

GC/MS data is collected utilizing Hewlett Packard 1000 RTE, RTA or DOS chemstation computer systems with Aquarius or Environquant software. This software allows for the comparison of sample non-target spectrum against reference library spectra. The most recent NIST/EPA mass spectral library supported by the system must be used. Data is reviewed by the analyst. If the data meets QC requirements, then binary data files are then converted to ASCII report files for transfer to the Seedpak II computer system via the network for data report forms generation.

Atomic Absorption

ICAP metals are analyzed by a Thermo-Jarrel Ash 61 or 61E. The data collected is transferred via a network system to the Seedpak system. Furnace data analyzed by the Perkin Elmer 5100s are collected on PCs and also transferred to the network to the Seedpak system for forms generation. Mercury data is analyzed on the TJA mercury analyzer and entered into Seedpak.

Classical Chemistry

Routine wet chemistry analyses have pre-printed logbooks, such as distillation logs and digestion logs. The less frequent analyses are recorded in analysts' notebooks. Raw data is then entered into the LIMS computer for data calculation. This includes the calibration curve data which may have been previously entered. Semi-automated analyses performed on the Lachat produce calculated final results. These results are then entered into LIMS. Any raw data produced is stored in a central file. Quality control data is manually calculated. Results data is reported off LIMS in the required format.

8.3 Review

Data review can be defined as the process whereby data is accepted or rejected based on specific criteria in order to ensure that the data are adequate for the intended purpose. In most cases, the criteria is defined by the particular analytical method.

Data review is performed prior to release of the data to the client. It is performed as soon as possible after data acquisition in order to provide sufficient time for corrective action if required.

In general, the procedure presented in Figure 8.3.1 is utilized by laboratory personnel throughout the network for data review purposes.

There are numerous policies and standard procedures which have been implemented to ensure that data of known quality is continually generated by the IEA-CT laboratory.

Each analytical SOP details the type and frequency of quality control checks. This includes such items as analysis of client reference standards, matrix spikes, blanks, the use of internal standards and surrogate spikes, etc. All calibrations are checked before sample analysis can begin. If the analytical system does not pass the initial QC limits, then the system is determined to be "out of control", and the cause of the problem must be determined and corrected before measurements can continue. Once the problem is corrected, QC measurements are repeated to verify the calibration. If the system is still out of control, the system is re-examined until the problem is corrected. General requirements are listed below:

Organics

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- . A minimum of one method blank is analyzed per 20 samples (or batch) per matrix, per concentration level or extraction procedure. A method blank is required every 12 hours for volatile analysis. Blanks and samples are analyzed on the same instrumentation. Pesticides/PCB's also require instrument blanks.
- Holding blanks are placed in volatile refrigerators on a weekly basis. For EPA CLP SOW volatile analysis, holding blanks are analyzed once per SDG.
- . A matrix spike/matrix spike duplicate is analyzed at a frequency of one per 20 samples per matrix, per concentration level or per SDG, whichever is more frequent.
- . Prior to sample processing, surrogates are added to all samples and method blanks. GC/MS analyses also require the use of internal standards.
- . Multi-level initial calibration curves are performed with continuing calibration standards analyzed every 12 hours. Recalibration is required if criteria cannot be met.
- . GC/MS system tuning is verified every 12 hours.

Inorganics

- . Multi-level calibration is performed on required instrumentation and verified as required.
- . Calibration and prep blanks are analyzed at required frequencies.
- . A matrix spike and sample duplicate are analyzed every 20 samples/SDG per matrix type.
- . A Laboratory Control Sample is analyzed every 20 samples or per batch.
- . Multi-level calibrations are performed for all manual and semi-automated wet chemistry methods and verified as required (if applicable).
- . Method blanks are analyzed at required frequencies.

The precision and accuracy control limits employed by IEA are based primarily on limits contained in the published methods or required by the U.S. Environmental Protection Agency's Contract Laboratory Program (CLP). When warranted by IEA's historical data, more restrictive control limits are set than those cited by the method or the CLP.

When the CLP protocol is not applicable to analysis of samples, the precision and accuracy requirements for each analytical method are included in the individual laboratory Standard Operating Procedure (SOPs). Examples of data acceptance criteria is detailed in table 7.3.2.

At a minimum, all data will be subject to supervisory review. Sensitive data requires higher level review and release. All releases must be in writing. Oral or Faxed preliminary releases are prohibited unless prior permission of the appropriate supervisor(s) is granted.

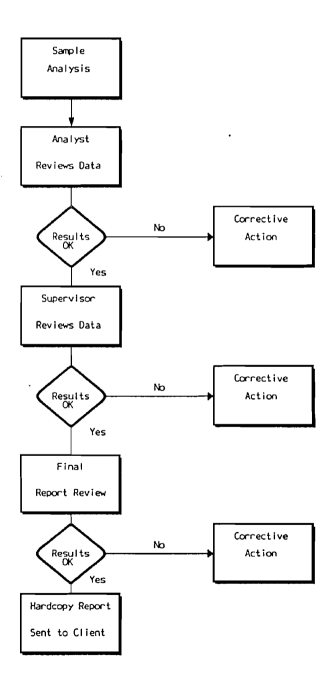
Each analytical group in the laboratory is responsible for generating the data for all analyses the group performs. In general the data must first meet all the specific QA/QC associated with the SOP that was used for the analysis prior to any release of the data. The analytical group leader (supervisor) is responsible for the final verification of the data from the analysis.

The laboratory employs a system of QA sign-off sheets called QC Batch Approval Forms and Quality Control Approval Reports (QCAR's), where each analyst must sign off that their respective part of the analysis is complete and meets the QA/QC requirements of the governing SOP. Both the Volatile and semi-volatile RTE computer systems produce batch-specific QC summary reports to check various analytical parameters. Analysis QCAR's are filled with the analysis batches while the final deliverable QCAR's are signed and placed in each job folder along with any Corrective Action Forms (CAF) which details any problems which were encountered in the measurement of samples. Any deviations from SOPs are noted on CAF's and explained in the SDG narrative which is incorporated into the final report. The group leader has final sign-off responsibility on the QCAR and is responsible for assuring the overall quality of the data.

The laboratory Quality Assurance Manager periodically examines data packages at random to ensure that all QCAR's are present and to ascertain that the data package meets the requirements as stated in the SOP. These findings are transmitted to laboratory management via progress reports.

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FIGURE 8.3.1 NETWORK DATA REVIEW PROCESS (GENERAL)



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8.4 Data and Report Storage

Unless specified otherwise by the client, all analytical data and associated information is stored for a minimum period of three years. Local state data storage requirements may vary from the corporate requirement and must be met by the laboratory if they are more stringent.

Stored information may consist of hardcopy or electronic data stored on a magnetic media.

All hardcopy information is stored at the laboratory that generated the data or off-site at a commercial document storage facility equipped with a professional security system.

All electronic data is stored on-site at the laboratory that generated the data or off-site at a commercial document storage facility equipped with a professional security system and a controlled environment suitable for storage of magnetic media.

Access to archived information is controlled by the appropriate data management custodian or facility manager.

At IEA-Connecticut, reports for the current year are filed in the data management area in filing cabinets. If the report has a larger data package, such as "CLP like" deliverables, it is then stored in numbered boxes. The number of the box is recorded into the cross reference logs and then stored in the locked storage area in the basement. All jobs must be signed out if being taken from the data management area.

8.5 Transcription

Whenever possible, manual data transcription is avoided through the use of electronic data transfer within the laboratories. In cases where manual transcription is employed, information is checked and verified by the supervisor or designee within the department.

It may not be possible to totally eliminate transcription related errors, however, section 4.6.9, paragraphs A and D, list procedures which are designed to minimize their occurrence and impact on data quality.

8.6 Data Reduction

Data reduction includes all processes that change either the form of expression (i.e., the units of measure) or the quantity of data values (rounding). It often involves statistical and mathematical analysis of data and usually results in a reduced subset of the original data set. Data reduction is performed either manually by the analyst or by computer systems interfaced to the analytical instruments. Whenever such procedures are employed within the laboratory network, mathematical procedures have been verified for accuracy of computation.

An example of this would be for CLP data packages, the data is transferred directly onto the Seedpak II system computer software from the Metals, GC and GC/MS systems via the network. The data is further processed and stored in the database. Other relevant data is transferred via the network at this point such as client ID's, etc. All calculations and final results are performed by the Seedpak II software. Many of these calculations are also done at the instrumentation level as a secondary review. Data in the database is sorted by client delivery group for easy retrieval. CLP forms are generated after all data is entered and reviewed. The forms and raw data are compiled into a data package. Tabular results are also generated at this point for level I reports reducing the occurance of typographic errors.

The data associated with each analysis is hardcopied for permanent storage either through the printing of computer files or through hand entry into bound laboratory notebooks. All notebook entries are dated and signed by the analyst.

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Job packages which include 20 samples or samples received by the laboratory during a one week time frame will comprise an "SDG". All organic parameter results will be reported in ug/L for aqueous samples and ug/Kg dry weight for soil/sediment samples. Inorganic result units vary according to the methodology.

It is laboratory policy that any and all problems related to client samples and the measurement of client samples be documented in the SDG narrative of the final laboratory report which goes to the client. The mechanism for documenting problems which shall be included in the SDG narrative is described in Section 10.0. It is the responsibility of the data management group to see that information on CAR's is included in the final SDG narrative.

After final review by the department manager, the data is placed in sample control for tracking on the project status sheet. If possible the data is placed into the job folder. When all parameters are complete the folder is removed by the data management department. It is the responsibility of the data management group to make sure that all the data is present and deliverable requirements are complete. This may include chain of custody forms, special instructions, and case narratives. The data is then compiled and sent to the report production group for word processing.

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9.0 DATA QUALITY ASSESSMENT

Data quality is assessed based on five main characteristics:

Precision
Accuracy
Completeness
Representativeness
Comparability

Each of these characteristics have been previously defined in section 2.2 of this document.

Laboratory Quality Assurance Objectives

Precision:

The objective of the network laboratories concerning precision is to equal or exceed the precision demonstrated in the published analytical method on similar samples. Relative Percent Difference (RPD) is used as the measure of precision sample duplicates. The formula utilized to calculate RPD is as follows:

Relative Percent Difference (RPD)

RPD = (Sample Result - Duplicate Result) x 100 Mean of Sample and Duplicate Results

Note: RPD is expressed as the absolute value obtained from the above formula.

Accuracy:

The objective of the network laboratories concerning accuracy is to equal or exceed the accuracy demonstrated in the published analytical method on similar samples. Accuracy is determined on matrix spikes and/or blank spikes and is calculated as follows:

Percent Recovery = (Observed-Sample) Concentration x 100 Spiked Concentration

Completeness:

IEA's objective for completeness is to be able to provide analytical data for 100 % of samples received intact and have sufficient sample volume for conducting re-analysis if initial analysis does not meet QC acceptance criteria.

Representativeness:

Representativeness of the analytical data is primarily a function of the sampling procedures and techniques employed in the field. As such, the sampling plan must be designed to provide representative samples to the laboratory. Once received at the laboratory, samples are homogenized in an effort to yield representative data on the sample submitted for analysis.

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Comparability:

IEA's objective for comparability is that all data be fully comparable with data from other network laboratories. This is accomplished through use of the following practices:

- Demonstrate traceability of standards to NIST or EPA sources
- Use of standard and approved methodologies
- Standardized units of measure
- Standardized QC acceptance criteria
- Participation in interlaboratory studies to demonstrate laboratory performance

9.1 Content of Analytical Reports

Laboratory customers have a wide variety of analytical needs. In order to meet these varied requirements, the laboratory offer several levels of data reporting options ranging from very simple format to an extreme level of documentation. Table 9.1.1 presents the contents of various levels of reports offered by the laboratory. Custom reporting beyond those listed is usually available but may require additional cost. The information provided in Table 9.1.1 is a summary only. In some cases, individual methods may not include the indicated items. For example, in metals graphite furnace analysis an ICP interference check would not be included since it is inappropriate for that method.

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Table 9.1.1 Report Content Options

	Data Reporting Options			
Wet Chemistry	Level 1	Level 2	Level 3	Level 4 (CLP)
Case narrative	Yes	Yes		Yes
Sample Results	Tabular	Tabular		Form I
Method Blank	Yes	Yes	_	Yes
External Chain of Custody	Yes	Yes		Yes
Internal Chain of Custody	Yes	Yes	<u>-</u>	Yes
Duplicate	•	Yes		Yes
Matrix Spike	-	Yes		Yes
Initial Calibration Verification (ICV)	•			Yes
Continuing Calibration Verification (CCV)	-	-		Yes
Laboratory Control Sample (LCS)	-	-		Yes
EPA Forms 1-14	-	-		Yes
Metals	Level 1	Level 2	Level 3	Level 4 (CLP)
Case Narrative	Yes	Yes	_	Yes
Sample Results	Tabular	Tabular		Form I
Method Blank	Yes	Yes	-	Yes
External Chain of Custody	Yes	Yes		Yes
Internal Chain of Custody	Yes	Yes		Yes
Duplicate	-	Yes	-	Yes
Matrix Spike	-	Yes		Yes
Initial Calibration Verification (ICV)	-			Yes
Continuing Calibration Verification (CCV)	-		_	Yes
Laboratory Control Sample (LCS)	-		-	Yes
ICP Interference Check	-	-	,	Yes
ICP Linear Range	-	-		Yes
ICP Post Spike	-	-	· 	Yes
EPA Forms 1-14	-	-	<u> </u>	Yes
Organics	Level 1	Level ?	Level 3	Level 4 (CLP)
Case Narrative	Yes	Yes		Yes
Sample Results	Tabular	TAbular	-	Form I
Method Blank	Yes	Yes		Yes
External Chain of Custody	Yes	Yes	_	Yes
Internal Chain of Custody	Yes	Yes		Yes
Matrix Spike	-	Yes		Yes
Matrix Spike Duplicate	-	Yes		Yes
Laboratory Control Sample (LCS)	-	-		as needed
Surrogate Recovery Information	-	Yes		Yes
Tuning Data (GC/MS only)	-	•		Yea
Initial Calibration Information	•			Yes
Continuing Calibration Information	-			Yes
Run Sequence Logs	-	-		EPA only
Sample Preparation Logs	<u> </u>	-		Yes
Chromatograms and Mass Spectra	•	-		Yes
EPA Forms 1-8		-		Yes

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10.0 CORRECTIVE ACTION

10.1 Introduction

The Corrective action form (CAF), presented in Section 7 of the Appendix, provides a routine written communication vehicle to describe most types of problems which may occur throughout the laboratory or as a result of a client inquiry. Problems described in SDG narratives should be supported by a CAF.

Corrective actions can be initiated at several operational levels; however they must always involve the QA Manager. Corrective actions are reviewed, documented and distributed to the appropriate personnel through the QA department. Responses are returned to QA for review and redistributed in a specified time frame.

Examples of three types of corrective actions which may be initiated are as follows:

Sample problems

Individual samples or matrix problems may cause documented corrective actions such as re-extraction, reanalysis, cleanups or dilutions.

OC problems

Corrective action may occur on entire batches of samples when QC criteria cannot be achieved.

Systematic problems

Specific project issues and procedural issues may require corrective actions. These are handled by laboratory management and the QA department.

The QA Manager will monitor and log the progress of CAF's and will report in the QA Progress Report the status of major corrective actions taken in the past month. It is the QA Manager's responsibility to see that laboratory problems are documented and solved in a timely manner. This system is outlined in the SOP for Corrective Action Reports - QAS00501.CT.

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10.2 System Audit

A system audit is an inspection and review of the entire data generation and support system of a laboratory. Activities related to the established requirements in the quality assurance program are reviewed for compliance. A typical system audit includes an evaluation of the following:

- Assessment of degree of compliance with the quality assurance program
- Continuing compliance with corrective actions identified in a previous audit of the facility
- Calibration procedures and documentation
- Sample handling procedures including chain-of-custody
- · Experience of laboratory personnel
- Existence and routine use of standard operating procedures
- Analytical data review and validation procedures
- · Data storage and recordkeeping

A system audit is performed by the on-site quality assurance manager at each facility annually. In addition to the above, a system audit is also conducted at the corporate level at each laboratory annually. The audits are staggered so that each facility is audited semi-annually, either by the local QA manager or corporate.

As previously indicated, all system audits are conducted utilizing a comprehensive standardized checklist (IEA Doc.# QAS00300.NET). Copies of the system audits conducted by the QA managers are submitted to the appropriate laboratory director/manager and president for review.

The auditor will identify any deficiencies in the audit report which is to be generated within a week of the actual audit. The laboratory director/manager is required to respond, in writing, no later than 30 days from issuance of the audit report. The response must address each of the items contained in the audit. If corrective action cannot be taken immediately, the anticipated date of compliance must be presented. If the auditor identifies issues which are significant (in their opinion), a follow-up audit can be conducted prior to the regularly scheduled audit.

A summary of the audit report findings is included in the quality assurance status report provided to management by the corporate quality assurance director.

10.3 Performance Audits

A performance audit is a quantitative check on the accuracy and/or precision of analytical data.

IEA network laboratories participate in a number of contracts and certification programs (see Table 5.4.1). Many of the certification programs employ rigorous performance evaluations which take the form of proficiency samples submitted to the laboratories on a regular basis. The following represents typical examples of routine proficiency programs.

All network laboratories are active participants in EPA Water Pollution (WP) and Water Supply (WS) proficiency programs which issue performance check samples on a semi-annual frequency.

IEA-CT participates in a number of contracts and certification programs. Many of these programs employ performance evaluations which take the form of proficiency samples submitted to the laboratory on a regular basis.

Bi-annually, the laboratory participates in the USEPA Water Supply (WS) and Water Pollution (WP) proficiency programs. IEA-CT also participates in the NYSDOH proficiency testing program for Potable Water, Hazardous Waste and CLP. The lab currently analyzes quarterly organic PE samples from EPA for the CLP program.

On a semi-annual basis, the QA Manager will submit QC samples supplied from an external source to the laboratory. The purpose of this is to check the accuracy of results, assess data quality, documentation and completeness of data

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reporting. The QA Manager shall submit an data review report to the laboratory. A written response must be submitted to QA within two weeks addressing the unacceptable findings. Corrective actions shall be put in place and monitored.

A copy of all analytical results associated with any proficiency samples is submitted to the operations director and president by each laboratory. The corporate office reviews this information and will utilize it in performing the regularly scheduled system audits at each lab. If results indicate a significant problem may exist, the network QA director will investigate accordingly.

In addition to participating in the above performance evaluation programs, the corporate office conducts additional performance evaluation studies.

Periodically, performance evaluation samples are submitted to each laboratory for parameters which are not addressed in other performance evaluation programs (ie. TCLP testing). In this type of testing the laboratory is aware the samples are performance check samples but the "true" concentrations are unknown. The results are submitted to corporate QA for evaluation and a report is issued on the findings. Corrective actions are taken if required, as a result of these test findings.

10.4 Independent Audits

IEA network laboratories are routinely audited by state and federal agencies for compliance with government regulations. In addition, several industrial clients conduct systems and performance audits of the facilities prior to project plan approval.

10.5 Subcontracted Services

IEA network laboratories occasionally choose to send selected analyses to a subcontract laboratory outside of the IEA organization. The most common reason for utilization of a subcontract facility is that the procedure is not routinely performed by an IEA network laboratory and the subcontractor has greater experience in day-to-day execution of the method. In such cases, although an IEA lab could in all likelihood conduct the analysis, it is more cost effective for both IEA and the client to utilize a subcontract lab as necessary. All subcontract laboratories utilized by IEA on a continuing basis require approval of the QA department prior to use. The QA manager is responsible for defining the analytical requirements to be met by the subcontract lab. For instance, the QA manager and the subcontracting lab must agree on the specific quality control to be performed with the samples submitted for analysis. Acceptance limits for items such as method blanks and matrix spiking must be determined.

IEA's clients are notified whenever another IEA laboratory or a subcontract laboratory is to be utilized for any portion of the analytical requirements. Although all analytical data appears in the IEA report, all data produced by another IEA laboratory or a subcontract laboratory is identified. In specific cases, states (ie. New Jersey) may have specialized requirements concerning the reporting of subcontracted analyses. In such cases, the laboratory will comply with the stated requirements. Subcontractors are not utilized when specifically restricted in a client's quality assurance project plan.

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APPENDIX, Section 1

PROFESSIONAL PROFILES OF KEY PERSONNEL

The following professional profiles are presented alphabetically and represent the key quality assurance and laboratory management personnel for the network organization. Additional professional profiles are available for review during a site visit to any of our laboratory facilities.





PROFESSIONAL PROFILE Michael V. Bonomo

TITLE: Vice-President and Director of Operations-IEA Connecticut

ACADEMIC ACCOMPLISHMENTS:

Fordham University - Bronx, New York B.S. Biology

Pace University - White Plains, New York M.B.A Marketing

MAJOR AREA OF EXPERTISE:

Environmental Regulations (RCRA, CERCLA, CWA, SDWA, ECRA)

Sampling and Analysis Plan Design

Data Management

SUMMARY OF EXPERIENCE:

Mr. Bonomo has over 15 years experience in environmental monitoring programs. He has functioned in numerous roles including director, co-director, sales manager, project manager, field and laboratory scientist, consultant, and seminar instructor. He has assisted many Fortune 500 companies and consultant/engineers in the design and implementation of sampling and analysis project plans. He has been involved in a wide spectrum of environmental programs for groundwater, soil, and sludge testing as well as monitoring various aquatic biota. Mr. Bonomo is also experienced in data management requirements for large analytical projects. He was instrumental in developing and implementing a data collection through data reporting system that was successfully utilized on many projects. He has also served as a seminar instructor for groundwater monitoring sampling and data tracking for a major waste management company.

PROFESSIONAL EXPERIENCE:

1992 to Present IEA, Inc.

Monroe, CT

Position Vice-President, Director of Operations

Responsibility

Responsible for overall operations and profitability.

<u>1991 to 1992</u> IEA, Inc.

Monroe, CT

<u>Position</u> Co-Director

Responsibilities

Co-responsibility for the profitability and management. Duties included business development, marketing, financial and budget management, sales management, strategic planning and monitoring operations.



1990 to 1991

IEA, Inc.
Monroe, CT

Position

Sales and Marketing Manager

Responsibilities

Responsible for the sales staff, corporate strategic planning, sales management and marketing in the New Jersey, Connecticut, Massachusetts and Vermont laboratories.

1989 to 1990

York Wastewater Consultants (YWC)

Monroe, Connecticut

Position

Executive Director

Responsibilities

Assisted in the growth of an unknown Connecticut based company to one that covered all of the Eastern United States. Participated in a 6-month strategic planning process that provided insight into the tools needed to run a successful company. In spite of severe banking problems, a 2-year Federal EPA investigation, and a downturn in the market, saw YWC through successful acquisition by Aquarion.

1987 to 1989

York Wastewater Consultants (YWC)

Monroe, Connecticut

Position

Sales and Marketing Manager

Responsibilities

Managed three York Laboratories division of YWC, Inc. Responsible for the Northeast, Mid-Atlantic and Midwest United States. Built sales and marketing program where non had existed before. Helped to assimilate five disjointed businesses into a single working division with resource sharing, cross training, budget management, and team building.

1982 - 1987

ETC

Position

National Account Executive

Responsibilities

Developed and managed new business for analytical and data management services. Responsible for marketing to many Fortune 500 chemical, petrochemical, waste, and electronics firms. Efforts included major projects throughout the United States. Worked with clients in regulatory compliance, project design, and data use and interpretation. Developed a client base that was involved in RCRA groundwater monitoring, CERCLA site Remedial Investigation Feasibility Studies, New Jersey ECRA investigations, and Clean Water Act compliance. Involved in one of the first petroleum refinery land treatment demonstrations in the United States. Served as the Chairman of the ETC Technical Product Development Committee.

ists:

1980 - 1982 Lawler, Matusky and Skelly Engineers

Position Assistant Project Manager

Responsibilities

Project Manager for environmental monitoring programs related to the electric utility industry. Responsible for proposal writing, program design, management of staff scientists and technicians for projects, budget control, report writing and technical presentations. These projects included water chemistry, fish population studies, lower trophic level monitoring, and mitigation of the impact of power plants on the river environment. Designed and implemented groundwater sampling programs with customized equipment for a major site in New York State.

1978 - 1980 Lawler, Matusky and Skelly Engineers

<u>Position</u> Project Scientist

Responsibilities

Crew Chief for field survey including sampling of biota, water, soil and sludge. Responsible for maintaining field control of samples, including documentation, custody, and proper sampling techniques. Performed laboratory analysis including wet chemistry procedures, fish taxonomy and other biological studies. Responsible for writing Standard Operating Procedures for laboratory operations.

Rev. 10/95 · Doc#-HRR02101.NET



PROFESSIONAL PROFILE Jeffrey C. Curran

TITLE:

Laboratory Manager

ACADEMIC ACCOMPLISHMENTS:

Southern Connecticut State University - New Haven, Connecticut B.A. Chemistry, 1975
M.S. Chemistry, 1978

MAJOR AREA OF EXPERTISE:

Quality Control/Quality Assurance Hazardous Waste Analyses Classical and Wet Chemistry Analyses PCB Analysis Capillary GC/MS Analysis Industrial Hygiene

Certified Laboratory Director for the States of Connecticut, New York, New Jersey and Massachusetts.

SUMMARY OF EXPERIENCE:

Mr. Curran has extensive experience in analytical chemistry specializing in environmental analysis. He has worked in all areas of the laboratory and has hands-on expertise in general wet chemistry techniques, atomic spectroscopy, gas chromatography, infrared spectroscopy and gas chromatography/mass spectrometry.

PROFESSIONAL EXPERIENCE:

Present

IEA, Inc. - Connecticut

Position Laboratory Manager

Responsibilities

For the past 15 years Mr. Curran has directed and participated in a variety of projects. Some highlights are listed below:

Hazardous Waste Site, East Windsor, CT

At a major Connecticut Hazardous Waste site Mr. Curran participated in the sampling analysis of buried drums of hazardous waste during a state-supervised cleanup project.



Ethylene Oxide Emissions Testing, Sherburn, New York

At a major EtO user in Upstate New York, Mr. Curran directed an on-site testing program for measuring EtO emissions using gas chromatography. Mr. Curran also worked on a testing program in conjunction with the NYSDEC for testing pollutant control equipment for EtO sterilizers.

Canadian Tariff Board Hearings

Mr. Curran provided expert witness testimony at a Canadian Tariff Board Hearing concerning chemical composition of foam packaging material.

Worker Exposure Study, Lynchburg, Virginia

Mr. Curran directed an on-site industrial hygiene study to monitor employee exposure to various solvents and chemicals. Mr. Curran was also part of the team which analyzed the various samples collected using gas chromatography, atomic spectroscopy, and UV-VIS spectroscopy in accordance with NIOSH protocols.

Food Processing Plant, Rochester, New York

Mr. Curran conducted an investigation to determine the cause of stainless steel tubing failures for a national food process company. The results of this study were used in determining alternatives to the current materials used in the process.

Hazardous Breakdown Product Study

Mr. Curran designed a system to identify and measure potentially hazardous breakdown products resulting from the pyrolysis of plastic materials for an international aircraft manufacturer. Results of this study were used to identify what materials were responsible for and how to alleviate the problem.

PROFESSIONAL AFFILIATIONS:

Member of the American Chemical Society

100



PROFESSIONAL PROFILE Marsha Culik

TITLE:

QA Manager

ACADEMIC ACCOMPLISHMENTS:

S.U.N.Y. at Alfred - Alfred, New York A.A.S. Medical, 1976 Laboratory Technology

MAJOR AREA OF EXPERTISE:

Extensive development and "hands on" experience with Gas Chromatography, Atomic Absorption Spectrophotometry, Auto Analyzer, and some computer data stations.

SUMMARY OF EXPERIENCE:

Ms. Culik has over 12 years experience in the environmental laboratory field. Experience ranges from analysis of drinking water with a Grade 3 Water Treatment Plant Operator to gas chromatography chemist with environmental samples. Ms. Culik has experience as supervisor of the Gas Chromatography department.

PROFESSIONAL EXPERIENCE:

1/91 to Present

IEA, Inc. - Connecticut

Position

QA Manager

Responsibilities

Quality Assurance Manager, responsible for monitoring the continuing compliance with the Corporate QA Program and to be a liaison between Corporate QA and laboratory staff.

Additional responsibilities include maintaining certification programs, coordination of external and internal audits, coordinate all inquiries relative to quality issues and follow-up on corrective actions as necessary, maintain files of all QA related documentation include review and approval of all SOP's.

1986 to 1991

Position GC Group Leader



Responsibilities

Supervisor of GC Group, responsible for analysis of environmental samples for pesticides/PCB's according to EPA/NYSDEC CLP Protocols, SW846 Methods and EPA "600" Series Methods. Additional responsibilities include analysis of samples via purge & trap/GC according to various protocols.

Other duties include analysis of air samples, charcoal absorbent tubes and other miscellaneous samples for any parameters requiring gas chromatography analysis. She is also responsible for supervision of the group including sample tracking, data review, etc.

1984 to 1986

Position Chemist

Responsibilities

Experience in sample prep and GC analyses of Pesticides/PCB's in water, oil and soil samples.

1981 to 1984

Position Laboratory Analyst - American Waterworks Service Company

Responsibilities

Experience performing complete laboratory analysis or raw, potable, and waste water including all miscellaneous include Volatile Organics, Trihalomethanes and Aromatics using Purge and Trap techniques; Pesticides and Herbicides by GLC; Transition and Heavy Metals by Flame and Graphite Furnace Atomic Absorption; and Nutrients by Automated and other various wet chemistry procedures. Assisted Lab Director in the development of many methods used in these analyses. Responsible for collection and interpretation of all quality control data.

1978 to 1981

Position Lab Technician - Suffolk County Water Authority

Responsibilities

Laboratory experience in the analysis of potable water for a large water utility. Cooperative studies done in conjunction with state and local health agencies concerning water and wastewater quality. Also monitoring the chemical quality of water and seawater programs for the U.S.G.S. Primary responsibilities were for the analysis of Halogenated and Aromatic organic compounds by Purge and Trap Gas Chromatography. Other areas of experience include the analyses of nutrients by Technicon Auto Analyzer, metals by Flame and Graphite Furnace Atomic Absorption, and microbiological testing using Millipore System.



1976-1978

Position Lab Technician - Hooker Chemicals & Plastics

Responsibilities

Responsible for the analysis of vinyl chloride monomer in PVC Compounds, Resins and Food Packageability studies utilizing Gas Chromatography. Responsible for monitoring the air quality of the plant environment.

SPECIALIZED TRAINING:

1984 Certified Grade 3 Water Treatment Plant Operator

1977 ASCP Registered MLT

Environmental Laboratory Management
Two day seminar on Environmental Laboratory Management
John H. Taylor, Analytical Technology.

Performance Management Workshop
One day seminar
Cynthia Barnet, Human Resources Consultant

Interview Skills Workshop
One day seminar
Cynthia Barnet, Human Resources Consultant

Leadership Development Workshop Four day workshop William Frackler, Ingoldsby, Inc.

Mass Spectral Data Interpretation
One day seminar
Dr. Frank Rutecek, Cornell University

Introduction to Analytical Separations
Four day seminar
Dr. Dhea Habboush, Sacred Heart University

ASQC Course Auditing of Quality Systems

> ASQC Course Introduction to SPC

> > Page 3



PROFESSIONAL PROFILE Lawrence H. Decker

TITLE:

GC/MS Manager

ACADEMIC ACCOMPLISHMENTS:

Franklin Pierce College - Rindge, New Hampshire B.A. Biology 1982

MAJOR AREA OF EXPERTISE:

Final Data Review
Coordination of sample analysis for the GC/MS group
Organics analysis by GC/MS

SUMMARY OF EXPERIENCE:

Lawrence Decker has eight years of GC/MS experience. He has been responsible for operations of the GC/MS group for five years. Presently functioning as the Volatile Group Leader.

PROFESSIONAL EXPERIENCE:

5/92 to Present

IEA, Inc. - Connecticut

Position

GC/MS Manager

Responsibilities

Responsible for the volatile group operations. Duties include: Scheduling workforce, ordering supplies, final data package review, employee reviews, overseeing sample analysis and sample prioritizing, adhering to forecasted budget, dealing with client requests, training employees, updating sample/job status with client service and laboratory directors. Tracking workflow through group.

10/91 to 5/92

Position

GC/MS Section Leader

Responsibilities

Responsibilities included: Sample analysis for both semi-volatile and volatile samples, tracking and scheduling sample analysis, troubleshooting instrumentation, final data package preparation and review. Unknown compound determination (TIC's). Assisting Group Leader with selected tasks. Responsible for tracking and prioritizing sample analysis, reviewing both initial sample batches and final reports, troubleshooting instruments and monitoring of GC/MS operations.

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4/86 to 9/90

Position GC/MS Operator

Responsibilities

Running samples, calibrating instruments, tracking samples, screening, total solids standard preparation, paperwork. Familiarity with EPA/NYSDEC CLP, SW846 and EPA "6--" Series VOA and BNA methods and routine analysis of aqueous and soil samples for VOA and BOA target and non-target (TIC) compounds. Experience in the data review process which involves monitoring surrogate recoveries, internal standard areas, target compounds concentration ranges and matrix spike/matrix spike duplicate performance parameters.

SPECIALIZED TRAINING:

Mass Spectroscopy Data Interpretation
One day Seminar
Dr. Frank Turecek (Cornell University)

Course description included close examination of mass spectra pertaining to identification of molecular ion, stability structure relationship, characteristic ion group effects, fragmentation and identifiable isotope clusters. Further concepts discussed include the nitrogen rule, the picket fence (alkane) series, and common fragment ions.

RTE-VI Procedures File Workshop
Four day seminar
GC/MS HP Aquarius Software Training
Mark Harwick (HP Instructor)

Course description included detailed examination of GC/MS Hardware, theory and function of mass spectroscopy, data acquisition and interpretation. Course emphasized software manipulation to enhance the overall quality and quantity of accurate and legible data.

Hewlett-Packard User I Course Five day seminar Hewlett-Packard, Paramus, New Jersey

Course description included a general overview of the HP computer system, mass spectrometer theory, instrument tuning and utility programs.

Introduction to Analytical Separations

Introduction to Chemical Analysis

Terms associated with chemical analysis; a review of the important considerations in analytical chemistry; sensitivity and detection limit; evaluation of results.

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Analytical Separation

Solvent extraction; emulsions, completeness of extraction; extraction of organic compounds; pH effect; extraction with metal chelator.

Chromatography (General Principles)

Chromatographic behavior of solutes; column efficiency and resolution.

Gas Chromatography

Gas chromatograph; gas chromatographic columns; liquid phases and column selection; detectors for gas chromatography; optimization of experimental conditions; interfacing gas chromatography with mass spectrometry.

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PROFESSIONAL PROFILE John Bennett, Jr.

TITLE:

GC/MS Semi-Volatiles Supervisor

ACADEMIC ACCOMPLISHMENTS:

Southern Connecticut State University - New Haven, CT B.S. Biology 1978 (Chemistry Minor)

MAJOR AREA OF EXPERTISE:

Classical Chemistry Atomic Spectroscopy Organic Extractions Gas Chromatography Microbiology

SUMMARY OF EXPERIENCE:

An extensive background in all phases of laboratory operations. Was responsible for designing, specifying, and hiring staff for a state of the art environmental laboratory. Had day to day responsibility for all phases of operation of the lab. Responsible for writing and conducting performance reviews for staff. Implemented stringent QA/QC program in the lab following USEPA CLP protocols. Had direct responsibility for inorganics section of the laboratory. Functioned as a resource person and problem solver for staff.

Wide ranging experience in the analysis of environmental and hazardous waste samples using EPA, APHA, and ASTM methodologies. Experienced in the analysis of contaminants from stationary sources. Has performed industrial hygiene surveys fore a variety of contaminants, and is familiar with the NIOSH procedures for their analysis. Instrumental expertise is ICP spectroscopy, as well as flame and furnace atomic absorption spectroscopy. In addition, has extensive experience with all basic laboratory apparatus and gas chromatography.

A broad background in microbiology including the identification and enumeration of microorganisms from a wide variety of sources. Familiar with USP and APHA procedures of analysis. Performed studies on the effects of point source contamination of water supplies and has performed characterization of problem microorganisms in sewage treatment plants. Developed a novel procedure for determining the microbial kill effectiveness of ethylene oxide sterilization cycles..

PROFESSIONAL EXPERIENCE:

1988 to Present

IEA, Inc. - Connecticut

Position GC/MS Semi-volatiles Supervisor

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Responsibilities

Responsible for daily operations of organics extractions group. Interacted with other departments in the laboratory concerning the status of client samples. Responsible for the supervision of six staff members. Responsible for the quality of work produced by group as well as meeting turnaround goals.

1987 to 1989

Position Laboratory Director - Chemrox, Inc.

Responsibilities

State of Connecticut Certified Laboratory Director for Chemrox Laboratory Services. Had overall responsibility for the operation of the laboratory, as well as the development of the business. Supervised 10 staff members. Interacted with other departments in the company, as well as outside clients on technical aspects of laboratory analyses. Participated in seminars to educate various groups about environmental issues.

1985 to 1987

Position Senior Chemist

Responsibilities

Responsible for ethylene oxide associated analyses. Performed pilot scale testing on a variety of medical devices to determine optimal de-gassing conditions. Aided in the design and construction of a pilot ethylene oxide. Was a member of the AAMI committee that developed reference test methods for ethylene oxide residues in medical services.

1980 to 1985

Position Senior Microbiologist/Associate Chemist - YWC, Inc.

Responsibilities

Responsible for performing non-routine microbiological analyses as well as providing technical guidance to technicians performing routine work. Instituted strict quality control procedures on all reagents, media and organisms. Was responsible for routine and non-routine chemical analyses on environmental samples. Was heavily involved in atomic spectroscopy analysis. Also performed evaluations on consumer products ranging from air cleaners to home water purification units.

1978 to 1980

<u>Position</u> Senior Chemist - Nutmeg Chemical Company



Responsibilities

Promoted to Assistant Director of Laboratory. Supervised staff in absence of Director. Served as liaison between director and staff. Performed non-routine water and oil analysis, quality control companies products as well as routine water, oil and deposit analysis. Also performed microbiological analysis of water samples.

1978 to 1979

Position Laboratory Technician

Responsibilities

Responsibilities included routine water and oil analyses and quality control of products.

SPECIALIZED TRAINING:

Basic Atomic Spectroscopy
Perkin Elmer
Norwalk, Connecticut 1979

ICP Spectroscopy
Spectra Inc.
Pompton Lakes, New Jersey 1988

Graphite Furnace Atomic Absorption Spectroscopy
Spectra Inc.
Pompton Lanes, New Jersey 1988

Interpretation of Low Resolution Mass Spectra YWC Whippany, New Jersey 1989



PROFESSIONAL PROFILE Daniel W. Helfrich

TITLE:

Inorganics Manager

ACADEMIC ACCOMPLISHMENTS:

Quinnipiac College Sacred Heart University M.S. Chemistry M.B.A. B.A. Biology B.S. Biology, 1985

MAJOR AREA OF EXPERTISE

Four years running ICP on environmental samples.

Two years running Furnace analysis.

Four years sample prep in environmental area.

Three years CLP Data Review.

OSHA trained and certified.

Familiar with EPA & NYSDEC protocols and SW846 Methods relating to inorganic metals analysis.

SUMMARY OF EXPERIENCE:

Mr. Helfrich has over 4 years experience in environmental analysis. He has functioned in numerous analytical roles including: Sample prep, Furnace analysis, ICP analysis and hazardous waste coordinator. Experienced in data review, and familiar with EPA and NYSDEC protocols. OSHA trained and experienced.

PROFESSIONAL EXPERIENCE:

1992 to Present

IEA, Inc. - Connecticut

Position

Group Leader

Responsibilities

Manage daily flow of work, set priorities.

Monitor productivity of group.

CLP data review ensuring QA/QC protocols are followed.

Manage the collection and removal of all hazardous waste generated by IEA-CT.

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<u>1989 to 1992</u>

Position Senior Chemist - IEA, Inc. CT

Responsibilities

ICP & Furnace Operator, manage flow of work, CLP data review ensuring QA/QC protocols are followed.

1987 to 1989

Position Lab Manager - PGP Industries

Responsibilities

ICP Operator and Health & Safety Manager

1984 to 1987

Position Senior Chemist - Handy & Harmon

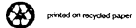
Responsibilities

ICP Operator

SPECIALIZED TRAINING:

OSHA Seminar - 40 hour training + 28 hour update

Clean Harbours - Hazardous Waste Seminar





PROFESSIONAL PROFILE Kimberly A. Maturo

TITLE:

GC/Semi-VOA Group Leader

ACADEMIC ACCOMPLISHMENTS:

Southern Connecticut State University - New Haven, Connecticut B.S. Biology, 1985

SUMMARY OF EXPERIENCE:

Ms. Maturo has over 7 years experience in the environmental field. She started in the organic extractions department as a lab technician and worked her way up to supervisor. From there, she transferred to the Gas Chromatography Department in order to expand her knowledge by learning more about the analysis of environmental samples. She is now Group Leader of the GC Department and is experienced in Pesticide and PCB residue analysis.

PROFESSIONAL EXPERIENCE:

3/91 to Present

IEA, Inc. - Connecticut

Position

GC Group Leader

Responsibilities

Supervisor of GC Group, responsible for analysis of environmental samples for pesticides/PCB's according to EPA/NYSDEC CLP Protocols, SW846 Methods and EPA "600" Series Methods. Additional responsibilities include analysis of samples via purge & trap/GC according to various protocols.

Other duties include analysis of air samples, charcoal absorbent tubes and other miscellaneous samples for any parameters requiring gas chromatography analysis. She is also responsible for supervision of the group including sample tracking, data review, etc.

10/88 to 3/91

Position

GC- Senior Lab Technician

Responsibilities

Ms. Maturo's primary duties are the operation of the gas chromatographs for a variety of analyses. She has experience in pesticide/PCB determinations as well as other miscellaneous analytes such as alcohols, herbicides and solvents in general.



Ms. Maturo's other duties include computer data entry, sample tracking and monitoring QC samples for the group.

10/85 to 10/87

Position Extractions Group

Responsibilities

Over this time period Ms. Maturo was a member of the extractions group and supervised the operations and staff for the last year. Her duties were primarily extraction of environmental samples for semi-volatile organics, pesticides/PCB's and herbicides. She also was responsible for screening of organic extracts via gas chromatography.

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PROFESSIONAL PROFILE Bruno D'Ostilio

TITLE:

Systems Manager - IEA-CT

ACADEMIC ACCOMPLISHMENTS:

Western Connecticut State University - Danbury, CT Computer Science and Business Program September 1978, 1989 - Present

MAJOR AREA OF EXPERTISE:

Various types of computer hardware. Unix and DOS operating systems. Networks. Programming.

SUMMARY OF EXPERIENCE:

Mr. D'Ostilio has over 10 years experience in the computer industry. He has a broad knowledge of computers ranging from mainframes to P.C.'s. He has extensive knowledge of UNIX, XWINDOWS, TCP/IP, DOS and various types of hardware.

PROFESSIONAL EXPERIENCE:

10/92 to Present

IEA, Inc. - Connecticut

Position

Systems Manager-IEA-CT

Responsibilities

Systems Manager-IEA-CT responsible for all systems development and maintenance at CT laboratory. Including LIMS, ADAM and instrument systems support. He will also be responsible for the development and implementation of a new LIMS System scheduled for 1994.

Position

Systems Manager

Responsibilities

Mr. D'Ostilio was responsible for the reintegration, rescaling, checking standards. Mr. D'Ostilio has over 10 years computer related experience on systems ranging from mainframes to personal computers. He is responsible for all systems hardware and software for our CLP 390 system. This system includes a HP9000 workstation with a UNIX Operating System, Envision application software, Ingres RDMS, XWindows and a network of X Terminals and P.C.'s using Advanlink and TCP/IP. His is also responsible for ensuring diskette deliverables meet the requirements specified by the EPA.

1



10/86 to 10/92

Position Senior Customer Engineer - Concurrent Computer Corp., Wallingford, CT

Responsibilities

Install, service and maintain Concurrent mini and microcomputers, as well as many other types and manufacturers of disk drives, tape drives, printers, modems, multiplexers, networks and personal computers. Also responsible for the installation, troubleshooting and upgrading of systems software which involves shell and C programming. System software includes UNIX SYSTEM V, XWINDOWS, MOTIF, TCP/IP, LABWORKBENCH, DOS and many other packages.

1984 to 1986

<u>Position</u> Customer Engineer - Memorex Corporation, Darien, CT

Responsibilities

Install, service and maintain magnetic tapes drives, disk drives, display stations and printers within local Connecticut & New York territories.

SPECIALIZED TRAINING:

IBM compatible mainframe peripherals, include:

High performance disk drives, tape drives, Impact & Laser printers.

Concurrent computer corp. mini computer hardware.

UNIX Systems Manager.

UNIX based Workstation Hardware.

Novell networks.

RDMS;





PROFESSIONAL PROFILE Stephanie N. Plunkett

TITLE:

Client Services Manager

ACADEMIC ACCOMPLISHMENTS:

BA - Biology, 1986 Hartwich College, Oneonta, NY Four year recipient - Hartwick College Merit Scholarship

PROFESSIONAL EXPERIENCE:

1/93 to Present

IEA, Inc. - Connecticut

<u>Position</u>

Client Services Manager

Responsibilities

Responsible for client service representatives/project managers functions. Aid in solving client problems and questions. Discuss technical issues and manage clients through sampling programs. Assist Account Executives on sales calls and project kick-off meetings.

7/91 to 9/91

IEA, Inc. - Monroe, CT

Position

Inside Sales/Project Manager

Responsibilities

While continuing to perform project management services for an established list of laboratory clients, I also assumed marketing and sales responsibilities. Duties include surveying various trade journals, The Federal Register, etc. identifying regulatory trends and predicting future business opportunities. Follow-up includes determining which industries would most likely be impacted by pending legislation and the development of a marketing strategy. Strategies implemented include telemarketing campaigns, mass mailings, and a seminar series. Also responsible for surveying existing clients periodically to assess IEA's strengths and weaknesses. The results of these surveys are compiled, graphically displayed and distributed to all employees.

200



10/88 to 3/90 IEA, Inc., Monroe, CT

Client Services/Project Manager

Operational responsibilities in environmental investigations, including scheduling and workload projection, technical supervision, sample tracking, contract compliance, data review, and report preparation and interpretation. Interface with clients on project design, including sampling and analytical program requirements, responsible for coordinating project specific quality control/methodology compliance requirements.

1987/1988

Massachusetts General Hospital

Boston, MA

Technologist

Cardiac Unit/Department of Molecular Research. Utilized knowledge of genetics and biology in the screening of DNA libraries with variously prepared probes, plasmid construction, and single and double stranded Sangar dideoxy chain termination sequencing. Duties also include maintaining laboratory supplies and the training of new personnel.

Related Course Work

Chemistry, Organic Chemistry, Environmental History, Ecology



- 2 -

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Date: 03/24/95

APPENDIX, Section 2

IEA CHAIN-OF-CUSTODY FORM

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Date: 03/24/95

APPENDIX, Section 3

IEA NETWORK SAMPLE PRESERVATION AND HOLDING TIME REQUIREMENTS

Date: 7/28/94 Page 1 of 18

		Me	tals in Water		
Parameter ³	Technique	Method	Holding Time	Container	Preservation
Aluminum	flame	202.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	202.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7020	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Antimony	flame	204.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	204.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7040	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7041	6 Months	500 ml P,G	HNO3 to pH < 2
•	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Arsenic	furnace	206.2	6 Months	500 ml P,G	HNO3 to pH < 2
	AA, hydride	206.3	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	AA, hydride	7061	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7060	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Barium	flame	208.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	208.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7080	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7081	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Beryllium	flame	210.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	210.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7090	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7091	_6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Boron	colorimetric	212.3	6 Months	500 ml P,G	HNO3 to pH < 2
 _	ICP	6010	6 Months	500 mt P,G	HNO3 to pH < 2

Date: 7/28/94 Page 2 of 18

		Metals in	Water-Continued		
Parameter 3	Technique	Method	Holding Time	Container	Preservation
Cadmium	flame	213.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	213.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7130	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7131	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Calcium	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	215.1	6 Months	500 ml P,G	HNO3 to pH < 2
	titrimetric	215.2	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7140	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Chromium	flame	218.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	218.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7190	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7191	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Chromium	Coprecipitation	7195	24 Hours	500 mi P,G	Cool, 4 C.
Hexavalent	colorimetric	7196	24 Hours	500 ml P,G	Cool, 4 C.
	flame	7197	24 Hours	500 ml P,G	Cool, 4 C.
	DPP	7198	24 Hours	500 ml P,G	Cool, 4 C.
Cobalt	flame	219.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	219.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7200	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7201	6 Months	500 ml P,G	HNO3 to pH < 2
- 	ICP	6010	6 Months	500 ml P.G	HNO3 to pH < 2

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		Metals in	Water-Continued		
Parameter ⁵	Technique	Method	Holding Time	Container	Preservation
Copper	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7210	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7211	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Iron	flame	236.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	236.2	6 Months	500 ml P,G	HNO3 to pH<2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7380	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7381	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Lead	flame	239.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	239.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH <2
	flame	7420	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7421	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Magnesium	flame	242.1	6 Months	500 ml P,G	HNO3 to pH < 2
í• -	furnace	239.2	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7450	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7421	6 Months	500 ml P,G	HNO3 to pH<2
Manganese	flame	243.1	6 Months	500 ml P,G	HNO3 to pH<2
	furnace	243.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	74 60	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7461	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Mercury	cold vapor-manual	245.1	28 Days	500 ml P,G	HNO3 to pH < 2
	cold vapor-automated	245.2	28 Days	500 ml P,G	HNO3 to pH < 2
	cold vapor-manual	7470	28 Days	500 ml P.G	HNO3 to pH < 2

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		Metals in	Water-Continued		
Parameter 3	Technique	Method	Holding Time	Container	Preservation
Molybdenum	flame	246.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	246.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7480	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7481	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Nickel	flame	249.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	249.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7520	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7521	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Potassium	flame	258.1	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7610	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Selenium	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
-	furnace	270.2	6 Months	500 ml P,G	HNO3 to pH < 2
	AA, hydride	270.3	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7740	6 Months	500 ml P,G	HNO3 to pH < 2
	AA, hydride	7741	6 Months	500 ml P,G	HNO3 to pH < 2
Silica	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
Silver	flame	272.1	6 Months	500 mi P,G	HNO3 to pH < 2
	furnace	272.2	6 Months	_500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7760	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7761	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	_6 Months	500 ml P.G	HNO3 to pH < 2

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		Metals in	Water-Continued	г т	
Parameter 3	Technique	Method	Holding Time	Container	Preservation
Sodium	flame	273.1	6 Months	500 ml P,G	HNO3 to pH < 2
	1CP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7770	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Thallium	flame	279.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	279.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7840	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	7841	6 Months	500 ml P,G	HNO3 to pH < 2
	·ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Tin	flame	282.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	282.2	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7870	6 Months	500 ml P,G	HNO3 to pH < 2
Titanium	flame	283.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	283.2	6 Months	500 ml P,G	HNO3 to pH < 2
Vanadium	flame	286.1	6 Months	500 ml P,G	HNO3 to pH < 2
	furnace	286.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP ·	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7910	6 Months	500 ml P,G	HNO3 to pH<2
	furnace	7911	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2
Zinc	flame	289.1	6 Months	500 ml P,G	HNO3 to pH < 2_
	furnace	289.2	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	200.7	6 Months	500 ml P,G	HNO3 to pH < 2
	flame	7950	6 Months	500 ml P,G	HNO3 to pH<2
	furnace	7951	6 Months	500 ml P,G	HNO3 to pH < 2
	ICP	6010	6 Months	500 ml P,G	HNO3 to pH < 2

	<u> </u>	Wet Ch	emistries in Water		
Parameter 3	Technique	Method	Holding Time	Container	Preservation
Acidity	titrimetric	305.1	14 Days	100 ml P,G	Cool 4 C.
Alkalinity	titrimetric	310.1	14 Days	100 ml P,G	Cool 4 C.
Biochemical Oxygen Demand (BOD)	5 days, 20 C.	405.1	48 Hours	1000 ml P,G	Cool 4 C.
Bromide	titrimetric	320.1	28 Days	100 ml P,G	none required
Chemical Oxygen	titrimetric, mid-level	410.1	28 Days	50 ml P,G	Cool 4 C, H2SO4 to pH <
Demand (COD)	titrimetric, low-level	410.2	28 Days	50 ml P,G	Cool 4 C, H2SO4 to pH <
	titrimetric, high-level	410.3	28 Days	50 ml P,G	Cool 4 C, H2SO4 to pH <
	automated-colorimetric	410.4	28 Days	50 ml P,G	Cool 4 C, H2SO4 to pH <
Chloride	colorimetric	325.2	28 Days	50 ml P,G	none required
	colorimetric	9250	28 Days	50 ml P,G	none required
	titrimetric	9252	28 Days	50 ml P,G	none required
	colorimetric	9257	28 Days	50 ml P,G	none required
Cyanide	amenable to chlorine	335.1	14 Days ²	500 ml P,G	Cool 4 C, NaOH to pH>
	spectrophotometric	335.2	14 Days ²	500 ml P,G	Cool 4 C, NaOH to pH>
	Total, UV	335.3	14 Days ²	500 ml P,G	Cool 4 C, NaOH to pH > Ascorbic Acid 1
	colorimetric	9012	14 Days ²	500 ml P,G	Cool 4 C, NaOH to pH>
Fluoride	distillation	340.1	28 Days	500 ml P,G	none required
	ion selective electrode	340.2	28 Days	500 ml P,G	none required
	colorimetric	340.3	28 Days	500 ml P,G	none required
Hardness, Total	colorimetric	130.1	6 Months	100 ml P,G	HNO3 to pH < 2
	titrimetric	130.2	6 Months	100 ml P,G	HNO3 to pH < 2
Iodide	titrimetric	345.1	24 Days	100 ml P,G	Cool 4 C.
Methylene Blue Active Substances	colorimetric	425.1	48 Hours	500 ml P,G	Cool 4 C.
Nitrogen	colorimetric,phenate	350.1	28 Days	500 ml P,G	Cool 4 C, H2SO4 to pH <
Ammonia	distillation	350.2	28 Days	500 ml P,G	Cool 4 C, H2SO4 to pH <
	ion selective electrode	350.3	28 Days	500 ml P.G	Cool 4 C. H2SO4 to pH <

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		Wet Chemistri	es in Water-Continue	×d	
Parameter 3	Technique	Method	Holding Time	Container	Preservation
Nitrogen-TKN	colorimetric,phenate	351.1	28 Days	500 ml P,G	Cool 4 C, H2SO4 to pH <2
	block digestor	351.2	28 Days	500 ml P,G	Cool 4 C, H2SO4 to pH <2
	colorimetric	351.3	28 Days	500 ml P,G	Cool 4 C, H2SO4 to pH <2
	ion selective electrode	351.4	28 Days	500 ml P,G	Cool 4 C, H2SO4 to pH <2
Nitrate	colorimetric, brucine	352.1	48 Hours	100 ml P,G	Cool 4 C.
	colorimetric, brucine	9200	48 Hours	100 ml P,G	Cool 4 C.
Nitrate-Nitrite	colorimetric, hydrazine	353.1	28 Days	100 mi P,G	Cool 4 C, H2SO4 to pH <2
	cadmium reduction, auto	353.2	28 Days	100 ml P,G	Cool 4 C, H2SO4 to pH <2
	cadmium reduction, manual	353.3	28 Days	100 ml P,G	Cool 4 C, H2SO4 to pH <2
Nitrite	spectrophotometric	354.1	48 Hours	100 ml P,G	Cool 4 C.
Oil & Grease, Total	gravimetric	413.1	28 Days	1000 ml G only	Cool 4 C, HCL or H2SO4 to pH <2
	IR ·	413.2	28 Days	1000 ml G only	Cool 4 C, HCL or H2SO4 to pH <2
	gravimetric	9070	28 Days	1000 ml G only	Cool 4 C, HCL or H2SO4 to pH <2
	gravimetric-sludge	9071	28 Days	1000 ml G only	Cool 4 C, HCL or H2SO4 to pH <2
Petroleum Hydrocarbons	IR	418.1	14 Days	1000 ml G only	Cool 4 C, HCL to pH <2
рН	electrode	150.1	in-field	50 ml P,G	not applicable
	electrode	9040	in-field	50 ml P,G	not applicable
	test paper	9041	in-field	50 ml P,G	not applicable
Phenolics,	spectrophotometric	420.1	28 Days	500 ml G only	Cool 4 C, H2SO4 to pH <2
T-Recoverable	colorimetric	420.2	28 Days	500 ml G only	Cool 4 C, H2SO4 to pH <2
	4AAP, Manual, Distillation	9065	28 Days	500 ml G only	Cool 4 C, H2SO4 to pH <2
	4AAP, Auto, Distillation	9066	28 Days	500 ml G only	Cool 4 C, H2SO4 to pH <2
	MBTH, Distillation	9067	28 Days	500 ml G only	Cool 4 C, H2SO4 to pH <
Phosphorus, Ortho	colorimetric, auto	365.1	48 Hours	50 ml P,G	Filter immediately, Cool 4 (
	colorimetric, single	365.2	48 Hours	50 ml P,G	Filter immediately, Cool 4 0
	colorimetric-dual	365.3	48 Hours	50 ml P,G	Filter immediately, Cool 4
	total, auto, block	365.4	48 Hours	50 ml P,G	Filter immediately, Cool 4

Parameter 3	Technique	Method	Holding Time	Container	Preservation
Phosphorus, Total	colorimetric, auto	365.1	28 Days	50 ml P,G	Cool 4 C, H2SO4 to pH <
	colorimetric, single	365.2	48 Hours	50 ml P,G	Cool 4 C, H2SO4 to pH <
	colorimetric-dual	365.3	48 Hours	50 ml P,G	Cool 4 C, H2SO4 to pH <
	total, auto, block digester	365.4	48 Hours	50 ml P,G	Cool 4 C, H2SO4 to pH <
Residue (Solids)	filterable (TDS)	160.1	7 Days	100 ml P,G	Cool 4 C.
	non-filterable (TSS)	160.2	7 Days	100 ml P,G	Cool 4 C.
	total (TS)	160.3	7 Days	100 ml P,G	Cool 4 C.
	volatile	160.4	7 Days	100 ml P,G	Cool 4 C.
	settleable	160.5	48 Hours	100 ml P,G	Cool 4 C.
Specific	meter	120.1	28 Days	100 ml P,G	Cool 4 C.
Conductance	meter	9050	28 Days	100 ml P,G	Cool 4 C.
Sulfate	ion chromatography	300.0	28 Days	50 ml P,G	Cool 4 C.
	colorimetric	375.1	28 Days	50 ml P,G	Cool 4 C.
	gravimetric	375.3	28 Days	50 ml P,G	Cool 4 C.
	turbidimetric	375.4	28 Days	50 ml P,G	Cool 4 C.
	colorimetric	9035	28 Days	50 ml P,G	Cool 4 C.
	colorimetric	9036	28 Days	50 ml P,G	Cool 4 C.
	turbidimetric	9038	28 Days	50 ml P,G	Cool 4 C.
Sulfide	titrimetric	376.1	7 Days	500 ml P,G	Cool 4 C, ZnAc/NaOH t
	colorimetric	376.2	7 Days	500 ml P,G	Cool 4 C, ZnAc/NaOH t
	colorimetric	9030	7 Days	500 ml P,G	Cool 4 C, ZnAc/NaOH t
Total Organic Carbon (TOC)	combustion or oxidation	415.1	28 Days	50 ml P,G	Cool 4 C, HCL or H2SO to pH <2
	combustion or oxidation	9060	28 Days	50 ml P,G	Cool 4 C, HCL or H2SC to pH <2
Total Organic Halides (TOX)	titrimetric	9020	28 Days	1000 ml G only ⁴ No Headspace	Cool 4 C, H2SO4 to pH <2 ⁵ Sodium Sulfite
Turbidity	nephelometric	180 1	48 Hours	100 ml P.G	Cool A.C

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Parameter 3	Technique	Method	Holding Time	Container	Preservation
Ialogenated	gas chromatography	601	14 Days	3x40 ml vials	Cool 4 C.,Thiosulfate 6
Volatile Organics	gas chromatography	8010	14 Days	3x40 ml vials	Cool 4 C.,Thiosulfate 6
Non-Halogenated Volatile Organics	gas chromatography	8015	14 Days	3x40 ml vials	Cool 4 C., HCL to pH <
Purgeable Aromatics	gas chromatography	602	7/14 Days 7	3x40 ml vials	Cool 4 C., HCL to pH <
	gas chromatography	8020	7/14 Days 7	3x40 ml vials	Cool 4 C., HCL to pH <
Acrolein & Acrylonitrile	gas chromatography	603	14 Days	3x40 ml vials	Cool 4 C., HCL to pH 5
	gas chromatography	8030	14 Days	3x40 ml vials	Cool 4 C., HCL to pH 5
Phenois	gas chromatography	604	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
	gas chromatography	8040	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
Phthalate Esters	gas chromatography	606	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
	gas chromatography	8060	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
Nitrosamines	gas chromatography	607	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
Organochlorine Pesticides	gas chromatography	608	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
and PCB's	gas chromatography	8080	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate ⁶
Polynuclear Aromatic	gas chromatography/LC	610	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
Hydrocarbons (PNA's)	gas chromatography	8100	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6
	HPLC	8310	ext7 Days	l L, Amber G	Cool 4 C.,Thiosulfate

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	Parameters by Gas Chromatography in Water-Continued								
Parameter 3	Technique	Method	Holding Time	Container	Preservation				
Haloethers	gas chromatography	611	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 4				
Chlorinated Hydrocarbons	gas chromatography	612	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6				
	gas chromatography	8120	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C., Thiosulfate 6				
Organophosphoru s Pesticides	gas chromatography	8140	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6				
Chlorinated Herbicides	gas chromatography	8150	ext7 Days	1 L, Amber G	Cool 4 C., Thiosulfate 6				

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		Parameters	by GC/MS in Water	•	
Parameter 3	Technique	Method	Holding Time	Container	Preservation
Purgeables	GC/MS-624 list	624	7/14 Days 1	3x40 ml vials	Cool 4 C., HCL to pH < 2, Thiosulfate ⁶
	Priority Pollutant list	624	7/14 Days ⁷	3x40 ml vials	Cool 4 C., HCL to pH < 2, Thiosulfate ⁶
	Hazardous Substance	624	7/14 Days 7	3x40 ml vials	Cool 4 C., HCL to pH < 2, Thiosulfate 6
	Target Compound list (TCL)	624	7/14 Days 7	3x40 ml vials	Cool 4 C., HCL to pH < 2,
	Appendix IX list	624	7/14 Days ⁷	3x40 ml vials	Cool 4 C., HCL to pH < 2,
	Priority Pollutant list	8240	7/14 Days 1	3x40 ml vials	Cool 4 C., HCL to pH < 2,
	Hazardous Substance	8240	7/14 Days ⁷	3x40 ml vials	Cool 4 C., HCL to pH < 2, Thiosulfate ⁶
	Target Compound list (TCL)	8240	7/14 Days 7	3x40 ml vials	Cool 4 C., HCL to pH < 2, Thiosulfate 6
	Appendix IX list	8240	7/14 Days ⁷	3x40 ml vials	Cool 4 C., HCL to pH < 2,

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Parameters by GC/MS in Water							
Parameter 3	Technique	Method	Holding Time	Container	Preservation		
Base-Neutral & Acid	625 list	625	ext7 Days anal40 Days	I L, Amber G	Cool 4 C.,Thiosulfate 6		
Extractables	Priority Pollutant list	625	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate		
	Hazardous Substance	625	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C., Thiosulfate		
	Target Compound list (TCL)	625	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate 6		
	Appendix IX list	625	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate		
	Priority Pollutant list	8250	ext7 Days anal40 Days	I L, Amber G	Cool 4 C., Thiosulfate		
	Hazardous Substance	8250	ext7 Days anal40 Days	i L, Amber G	Cool 4 C.,Thiosulfate		
	Target Compound list (TCL)	8250	ext7 Days anal40 Days	I L, Amber G	Cool 4 C.,Thiosulfate		
	Appendix IX list	8250	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate		
	Priority Pollutant list	8270	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate		
	Hazardous Substance	8270	ext7 Days anal40 Days	1 L, Amber G	Cool 4 C.,Thiosulfate		
	Target Compound list (TCL)	8270	ext7 Days anal40 Days	I L, Amber G	Cool 4 C.,Thiosulfate		
	Appendix IX list	8270	ext7 Days	1 L, Amber G	Cool 4 C.,Thiosulfate		

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Metals in Soil						
Parameter 3	Technique	Method	Holding Time	Container	Preservation	
Aluminum	flame	7020	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Antimony	flame	7040	6 Months	100 g P,G	Cool 4 C.	
	furnace	7041	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Arsenic	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
	furnace	7060	6 Months	100 g P,G	Cool 4 C.	
	AA, hydride	7061	6 Months	100 g P,G	Cool 4 C.	
Barium	flame	7080	6 Months	100 g P,G	Cool 4 C.	
	furnace	7081	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Beryllium	flame	7090	6 Months	100 g P,G	Cool 4 C.	
	furnace	7091	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Вогоп	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Cadmium	flame	7130	6 Months	100 g P,G	Cool 4 C.	
	furnace	7131	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Calcium	flame	7140	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Chromium	flame	7190	6 Months	100 g P,G	Cool 4 C.	
	furnace	7191	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P.G	Cool 4 C	

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Metals in Soil-Continued						
Parameter ³	Technique	Method	Holding Time	Container	Preservation	
Cobalt	flame	7200	6 Months	100 g P,G	Cool 4 C.	
	furnace	7201	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Copper	flame	7210	6 Months	100 g P,G	Coo <u>l</u> 4 C.	
	fu rna ce	<i>7</i> 211	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Iron	flame	7380	6 Months	100 g P,G	Cool 4 C.	
	furnace	7381	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Lead	flame	7420	6 Months	100 g P,G	Cool 4 C.	
	furnace	7421	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Magnesium	flame	7450	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Manganese	flame	7460	6 Months	100 g P,G	Cool 4 C.	
-	furnace	7461	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Mercury	cold vapor-manual	7470	28 Days	100 g P,G	Cool 4 C.	
	cold vapor-manual	7471	28 Days	100 g P,G	Cool 4 C.	
Molybdenum	flame	7480	6 Months	100 g P,G	Cool 4 C.	
	furnace	7481	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Nickel	flame	7520	6 Months	100 g P,G	Cool 4 C.	
	furnace	7521	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Potassium	flame	7610	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Selenium	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
	furnace	7740	6 Months	100 g P,G	Cool 4 C.	
	A.A. hydride	7741	6 Months	100 g P,G	Cool 4 C	

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Metals in Soil-Continued						
Parameter 3	Technique	Method	Holding Time	Container	Preservation	
Silver	flame	7760	6 Months	100 g P,G	Cool 4 C.	
	furnace	7761	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Sodium	flame	7770	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Thallium	flame	7840	6 Months	100 g P,G	Cool 4 C.	
	furnace	7841	6 Months	100 g P,G	Cool 4 C.	
_	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Γin	flame	7870	6 Months	100 g P,G	Cool 4 C.	
Vanadium	flame	7910	6 Months	100 g P,G	Cool 4 C.	
	furnace	7911	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P,G	Cool 4 C.	
Zinc	flame	7950	6 Months	100 g P,G	Cool 4 C.	
	furnace	7951	6 Months	100 g P,G	Cool 4 C.	
	ICP	6010	6 Months	100 g P.G	Cool 4 C	

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Wet Chemistries in Soil						
Parameter 3	Technique	Method	Holding Time	Container	Preservation	
Cyanide	spectrophotometric	9010	14 Days	100 g P,G	Cool 4 C.	
	colorimetric	9012	14 Days	100 g P,G	Cool 4 C.	
Sulfate	colorimetric	9035	28 Days	100 g P,G	Cool 4 C.	
	colorimetric	9036	28 Days	100 g P,G	Cool 4 C.	
	turbidimetric	9038	28 Days	100 g P,G	Cool 4 C.	
Sulfide	colorimetric	9030	<u> </u>	100 g.P.G	Cool 4 C	

Date: 7/28/94 Page 17 of 18

Parameter ³	Technique	Method	Holding Time	Container	Preservation
Halogenated Volatile Organics	gas chromatography	8010	14 Days	3x40 ml vials ⁸	Cool 4 C.
Non-Halogenated Volatile Organics	gas chromatography	8015	14 Days	3x40 ml vials	Cool 4 C.
Purgeable Aromatics	gas chromatography	8020	14 Days	3x40 ml vials	Cool 4 C.
Acrolein & Acrylonitrile	gas chromatography	8030	14 Days	3x40 ml vials	Cool 4 C.
Phenois	gas chromatography	8040	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Phthalate Esters	gas chromatography	8060	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Nitrosamines	gas chromatography	8070	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Organochlorine Pesticides and PCB's	gas chromatography	8080	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Polynuclear Aromatic Hydrocarbons (PNA's)	gas chromatography	8100	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
	HPLC	8310	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Chlorinated Hydrocarbons	gas chromatography	8120	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Organophosphoru s Pesticides	gas chromatography	8140	ext14 Days anal40 Days	100 g ,G	Cool 4 C.
Chlorinated Herbicides	gas chromatography	8150	ext14 Days	100 g ,G	Cool 4 C.

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Parameters by GC/MS in Soil						
Parameter 3	Technique	Method	Holding Time	Container	Preservation	
Volatile organics	packed column	8240	14 Days	3x40 ml vials ⁶	Cool 4 C.	
	capillary column	8260	14 Days	3x40 ml vials	Cool 4 C.	
Base-Neutral & Acid	semi-vol packed	8250	ext14 Days anal40 Days	100 g ,G	Cool 4 C.	
Extractables	semi-vol capillary	8270	ext14 Days anal40 Days	100 g ,G	Cool 4 C.	

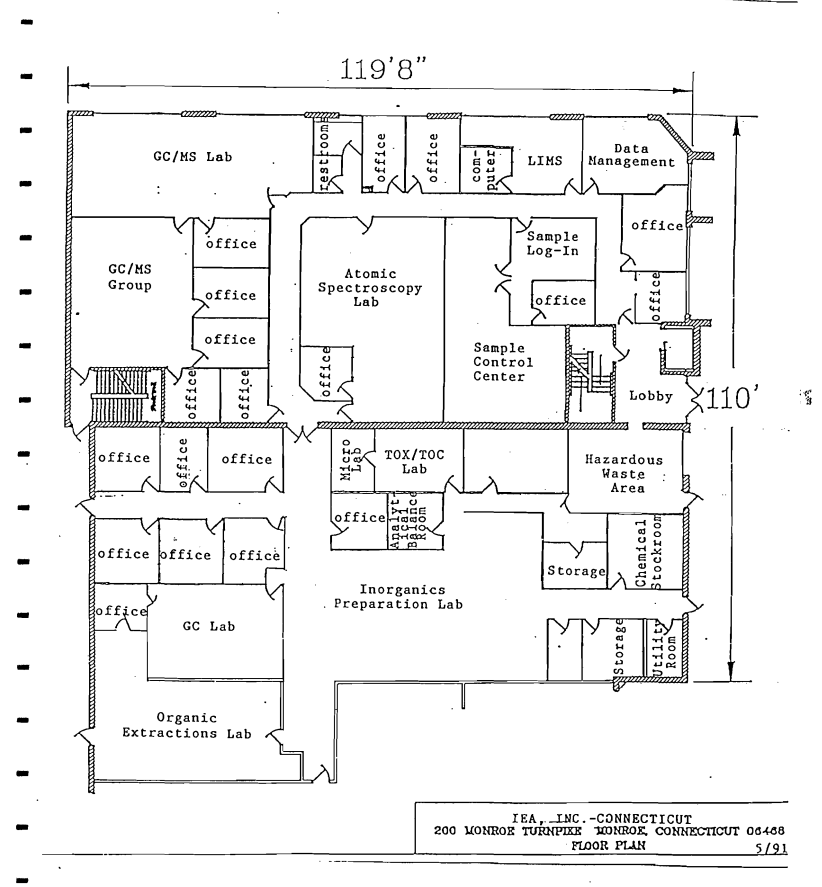
Footnotes

- 1 If residual chlorine is present in the sample, 0.6 g of ascorbic acid is utilized. Ascorbic acid is only used if residual chlorine is present.
- ² Maximum holding time is 24 hours when sulfide is present. Optionally, all samples may be tested with lead acetate paper before pH adjustments in order to determine if sulfide is present. If sulfide is present, it can be removed by the addition of cadmium nitrate powder until a negative spot test is obtained. The sample is filtered and then NaOH is added to pH 12.
- The following information is based upon EPA requirements as outlined in Table II, Part 136, Title 40 of the Code of Federal Regulations, July 1991. This reference should be consulted if further clarification is desired. Various state agencies have differing requirements for both holding times and preservation from those listed above. In such cases, the local requirements supercede the EPA information.
- 4 All samples should be collected in bottles with teflon septa and be protected from light. If this is not possible, use 250 ml bottles fitted with teflon lined caps. Samples should contain no headspace.
- If samples contain residual chlorine, it must be removed in the field by adding sulfite to the sample bottle (5 mg sodium sulfite crystals per liter of sample).
- If samples contain residual chlorine, 0.008% sodium thiosulfate must be added at the time of sampling and should only be used if residual chlorine is present.
- If samples do not receive pH adjustment, the holding time is 7 days. With pH adjustment, the holding time is 14 days.
- Alternatively, wide mouth glass jars designed for volatile samples may be utilized with teflon lined caps.

Date: 03/24/95

APPENDIX, Section 4

LABORATORY FLOOR PLAN



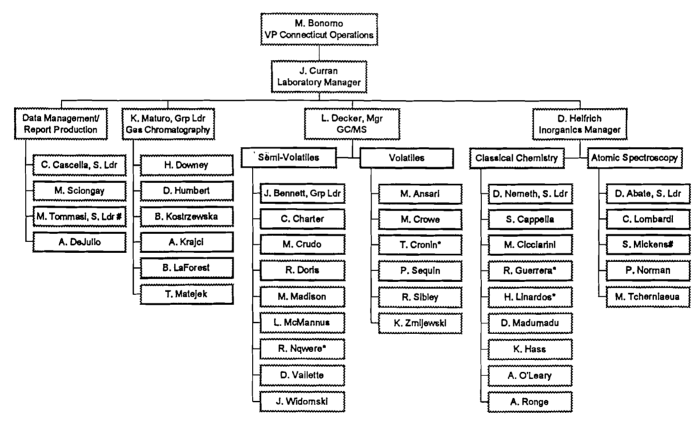
Date: 03/24/95

APPENDIX, Section 5

ORGANIZATIONAL CHART

IEA, Inc. - CT An Aquarion Company

Doc #QAC00104.CT Date: 01/23/96



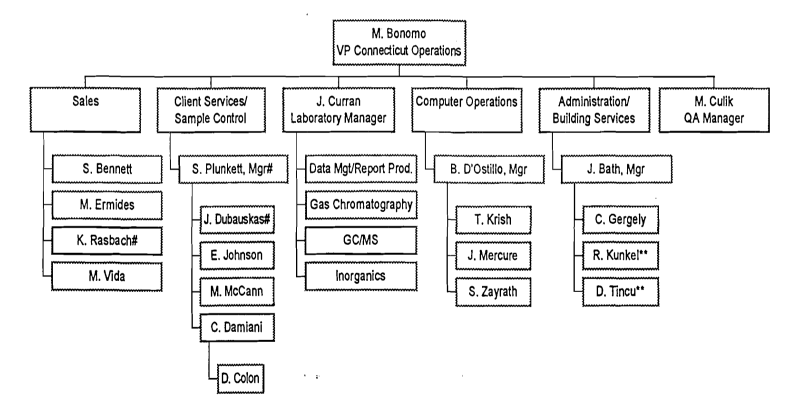
*Part Time/Temp

**25% charged to AMS

#30 hour work week



Doc #QAC00104.CT Date: 01/23/96



*Part Time/Temp **25% charged to AMS #30 hour work week

APPENDIX, Section 6

CORRECTIVE ACTION FORM



CORRECTIVE ACTION FORM

	ation			Client Inquiry_
Client:		Job/0	Case:	
			ole Number(s):	
	ntact:		Time Response Due:_	
Detailed Description of	f Potential Problem:			
•				_
		•	•	
B. Quality Assurance	Information		Corrective Acti	on ID#
Recommended Correct	ive Action:			
·	· · · · · · · · · · · · · · · · · · ·			
Groups Involved:	Sample Control	Wet Chemistry		
		Wet Chemistry	Metals	
	Sample Control	Wet Chemistry	Metals	
Groups Involved:	Sample ControlGas Chromatography	Wet Chemistry Mass Spectrometry	Metals	
	Sample ControlGas Chromatography	Wet Chemistry Mass Spectrometry	Metals	
Groups Involved: C. Final Resolution	Sample ControlGas Chromatography	Wet Chemistry Mass Spectrometry Sample Preparation	Metals Report Generation	
Groups Involved: C. Final Resolution	Sample ControlGas ChromatographyClient Service	Wet Chemistry Mass Spectrometry Sample Preparation	Metals Report Generation	
Groups Involved: C. Final Resolution	Sample ControlGas ChromatographyClient Service	Wet Chemistry Mass Spectrometry Sample Preparation	Metals Report Generation	
Groups Involved: C. Final Resolution Describe What Happen	Sample ControlGas ChromatographyClient Service	Wet Chemistry Mass Spectrometry Sample Preparation	Metals Report Generation	
Groups Involved: C. Final Resolution Describe What Happer	Sample ControlGas ChromatographyClient Service ned and Long Term Correct	Wet Chemistry Mass Spectrometry Sample Preparation	Metals Report Generation	
Groups Involved: C. Final Resolution Describe What Happer	Sample ControlGas ChromatographyClient Service ned and Long Term Correct	Wet Chemistry Mass Spectrometry Sample Preparation	Metals Report Generation	
Groups Involved: C. Final Resolution Describe What Happer Supervisor Signature:	Sample ControlGas ChromatographyClient Service ned and Long Term Correct	Wet Chemistry Mass Spectrometry Sample Preparation tive Action Taken:	Metals Report Generation	
Groups Involved: C. Final Resolution Describe What Happen Supervisor Signature: D. Quality Assurance	Sample ControlGas ChromatographyClient Service ned and Long Term Correct	Wet Chemistry Mass Spectrometry Sample Preparation tive Action Taken: Date	Metals Report Generation	
Groups Involved: C. Final Resolution Describe What Happer Supervisor Signature: D. Quality Assurance	Sample ControlGas ChromatographyClient Service ned and Long Term Correct Final Approval (QA Man	Wet Chemistry Mass Spectrometry Sample Preparation tive Action Taken: Date	Metals Report Generation	

APPENDIX, Section 7

EXAMPLE LISTING OF LABORATORY STANDARD OPERATING PROCEDURES (SOPs)

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SAMPLE CONTROL

Standard Operating Procedure	Code	Date Generated
SOP for Bottle Order Preparation	SMS00100.CT	02/15/95
SOP for Sample Processing and Sample Arrival	SMS00402.CT	05/15/92
SOP for Log-in of CLP Samples	SMS00502,CT	05/15/92
SOP for Storing Water and Soil Samples	SMS00602.CT	05/12/92
SOP for Generating Labels/Labeling Containers	SMS00700.CT	05/15/92
SOP for Documenting Sample Removal from Laboratory	SMS00802.CT	05/15/92
SOP for Securing the Laboratory and Samples	SMS00903.CT	05/15/92
SOP for Temperature Control Requirements	SMS01001.CT	05/15/92
SOP for Compositing Samples	SMS01100.CT	06/16/94
SOP for Sample Receipt (NJDEPE)	SMS01200.CT	01/24/95
SOP for Operating and Maintaining Fume Hoods	SFS00202.CT	05/15/92
SOP for Hazardous Waste Disposal	SFS00100.CT	05/06/92
SOP for Emergency Procedures	SFS00300.CT	06/21/94
SOP for Hazardous Waste Minimization Plan	SFS00500.CT	07/25/94
SOP for Tracking and Collection of Mixed Waste	RAS00100.CT	02/06/94
SOP for Radioactivity Swpie Tests	RAS00200.CT	08/17/94
SOP for Radiation Screening	RAS00300.CT	08/15/94
SOP for Management/Disposal of Mixed Waste	RAS00400.CT	08/24/94

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DATA MANAGEMENT/HANDLING

Standard Operating Procedure	Code	Date Generated
SOP for Preparation/ Review of Laboratory Reports	RPS00300.CT	02/16/94
SOP for Documentation Policy/Procedures	DM:090191:2	09/01/91
SOP for Data Reduction, Mgt, and Handling - CLP	RPS00200.CT	05/05/92
SOP for Sample Tracking	QAS00200,CT	01/13/92
SOP for Data validation/Self Inspection - CLP	QAS00100.CT	05/06/92
SOP for Data Validation/Self Inspection - OLM02.1	QAS00600.CT	01/17/94
SOP for Data Validation	QAS00700.CT	dft
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EXTRACTIONS

Standard Operating Procedure	Code	Date Generated
SOP for CLP Aqueous BNA Preparation	SPS00303.CT	08/20/91
SOP for CLP Aqueous Pesticide/PCB Preparation	SPS00403.CT	08/19/91
SOP for CLP Soil BNA Preparation	SPS00102.CT	08/23/91
SOP for CLP Soil Pesticide/PCB Preparation	SPS00202.CT	08/26/91
SOP for CLP Extractions Standard Prep	SPS00702.CT	05/07/92
SOP for CLP BNA extract Screening	SPS00803.CT	05/12/92
SOP for CLP GPC BNA Extracts	SPS00502.CT	08/29/91
SOP for CLP GPC Pesticide/PCB Extracts	SPS00602.CT	04/02/92
SOP for Cleaning Glassware	SPS00901.CT	05/13/92
SOP for Hydrocarbon Sample Prep	SPS01000.CT	dft
SOP for Aqueous Herbicides Method 509B	SPS01100.CT	dft
SOP for Prepararion of Chlorinated Herbicides - 8150	SPS02800.CT	dft
SOP for Aqueous BNA Methods 3510/3520	SPS01300.CT	09/10/93
SOP for Aqueous Pest/PCB Methods 3510/3520	SPS01200.CT	09/15/93
SOP for Soil BNA Method 3550	SPS01400.CT	12/10/93
SOP for Soil Pest/PCB Method 3550	SPS01600.CT	01/21/94
SOP for Aqueous OP Pesticides Methods 3510/3520	SPS01700.CT	06/15/94
SOP for SW846 GPC of BNA extracts	SPS01800.CT	12/17/94
SOP for GPC of Pesticide/PCB extracts method 3640	SPS01900.CT	03/04/94
SOP for Soil OP Pesticides Method 3550	SPS02700.CT	03/07/94
SOP for Waste dilution - BNA	SPS03000.CT	03/08/94
SOP for Waste dilution - Pesticides/PCB	SPS03100.CT	03/04/94
SOP for Pesticide/PCB extraction method 608	SPS03200.CT	08/24/94
SOPs for extractions CLP OLM02.1	SPS02000.CT- SPS02600.CT	dft
SOP for Extraction Standard Prep	SPS01500.CT	dft

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EXTRACTIONS - cont.

Standard Operating Procedure	Code	Date Generated
SOP for Prep Soil/Sediment samples for CLP P/P OLM03.1	SPS03300.CT	11/11/94
SOP for GPC of Pesticide extracts OLM03.1	SPS03400.CT	11/11/94
SOP for Prep Soil/Sed samples for CLP BNA's OLM03.1	SPS03500.CT	11/11/94
SOP for GPC of Semivolatile extracts OLM03.1	SPS03600.CT	11/11/94
SOP for Prep of Aqueous samples for CLP BNA's OLM03.1	SPS03700.CT	11/11/94
SOP for Prep of Aqueous samples for CLP P/P OLM03.1	SPS03800.CT	11/11/94
SOP for Prep of Low Level PCB Analysis 3510A	SPS03900.CT	11/29/94
SOP for Alumina Column C/U Method 3611A	SPS02900.CT	Dft
SOP for Prep Aqueous SV OLC10/92	SPS04000.CT	Dft
SOP for Prep of Semivolatiles in Tissue samples	SPS04200.CT	10/21/95
SOP for Prep of Pesticides/PCBs in Tissue samples	SPS04300.CT	10/21/95
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GC/MS

Standard Operating Procedures	Code	Date Generated
SOP for CLP Volatiles (GC/MS)	MSS00601.CT	09/04/91
SOP for Semi-volatile CLP OLM01.8	MSS01001.CT	09/10/91
SOP for Volatile Std Prep CLP	MSS00100.CT	05/05/92
SOP for Semi-volatile Std Prep CLP	MSS00200.CT	05/05/92
SOP for Cleaning AS vials	MSS01200.CT	02/15/93
SOP for Analysis of BNA Method 8270A	MSS00700.CT	05/23/94
SOP for Analysis of Volatiles Method 8240A	MSS00400.CT	04/30/93
SOP for Volatile Standard Prep	MSV:120588:1	12/05/88
SOP for BNA standard Prep	MSSV:112686:2	11/26/86
SOP for GC/MS Semi-volatiles CLP OLM02.1	MSS00800.CT	01/14/94
SOP for GC/MS Volatiles CLP OLM02.1	MSS00900.CT	01/14/94
SOP for Volatile Std Prep CLP OLM02.1	MSS01300.CT	01/14/94
SOP for Semi-volatile Std Prep CLP OLM02.1	MSS01400.CT	01/14/94
SOP for GC/MS Volatiles in Air	MSS00300.CT	dft .
SOP for GC/MS Volatile in Air - Summa Canister	MSS01100.CT	dft
SOP for GC/MS Volatile 524.2 Rev. 3	MSS01500.CT	dft
SOP for GC/MS Semivolatiles OLM03.1	MSS01600.CT	11/12/94
SOP for GC/MS Semivolatile Standard Prep OLM03.1	MSS01700.CT	11/12/94
SOP for GC/MS Volatiles OLM03.1	MSS01800.CT	11/12/94
SOP for GC/MS Volatile Standard Prep OLM03.1	MSS01900.CT	11/12/94
SOP for GC/MS Analysis Method 625	MSS02000.CT	07/13/94
SOP for GC/MS Analysis Method 624	MSS02100.CT	02/27/95
SOP for GC/MS Semivolatile OLC10/92	MSS02200.CT	Dft

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GAS CHROMATOGRAPHY

Standard Operating Procedures	Code	Date Generated
SOP for GC CLP OLM01.8	GCS00200.CT	09/11/91
SOP for Standard Prep CLP- Pesticides	GCS00100.CT	05/05/92
SOP for Sulfur Removal	GCS00300.CT	04/30/93
SOP for Pest/PCB Method 8080A	GCS00600.CT	02/15/94
SOP for Analysis of OP Pesticides Method 8141	GCS00500.CT	02/28/94
SOP for HP3350A LAS System	GCS00400.CT	06/08/93
SOP for Misc. Volatiles Method 8015 (DAI)	GCS00700.CT	02/14/94
SOP for Herbicide analysis Method 8150	GCS00800.CT	02/14/94
SOP for Analysis of Hydrocarbon Fingerprinting	GCS01300.CT	08/02/94
SOP for GC/ECD Pesticides/PCB CLP OLM02.1	GCS00900.CT	01/14/94
SOP for Pesticide/PCB Standard Prep OLM02.1	GCS01000.CT	01/14/94
SOP for Pesticides/PCB Method 608	GCS01100.CT	02/15/94
SOP for Sulfur Removal - CLP OLM01.8	GCS01200.CT	06/10/94
SOP for GC/ECD Pesticides/PCB analysis OLM03.1	GCS01400.CT	11/11/94
SOP for Pesticide/PCB Standard Prep OLM03.1	GCS01500.CT	11/11/94
SOP for Low Level Pesticide/PCB analysis - 8080	GCS01600.CT	11/29/94

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METALS

Standard Operating Procedures	Code	Date Generated
SOP for SW846 Method 3005	MES00800.CT	04/21/93
SOP for SW846 Method 3010	MES00900.CT	04/21/93
SOP for SW846 Method 3020A	MES00701.CT	04/21/93
SOP for SW846 Method 3050	MES01001.CT	04/21/93
SOP for CLP SOW Digestion (S)	MES01100.CT	04/21/93
SOP for CLP SOW Digestion (W)	MES01200.CT	04/21/93
SOP for Method 200.7 with TJA 61 Operation	MES00600.CT	04/16/93
SOP for GFAAS 200 series methods	MES00501.CT	04/16/93
SOP for Tracking Metals and Cyanide Samples	IN:050189:1	05/01/89
SOP for Standards Preparations	AS:092988:1	09/29/88
SOP for Determination of Mercury in Water ILM03.0	MES01300.CT	06/10/94
SOP for Determination of Mercury in Soils ILM03.0	MES01400.CT	06/10/94
SOP for Determination of Mercury in Water - 7470	MES01500.CT	09/12/94
SOP for Determination of Mercury in Soils - 7471	MES01600.CT	09/12/94
SOP for Method 6010A with TJA 61	MES00400.CT	09/12/94
SOP for GFAAS SW846 series methods	MES00300.CT	09/12/94
SOP for Microwave Digestion Method 3015 (W)	MES01700.CT	04/20/95
SOP for Microwave Digestion Method 3051 (S)	MES01800.CT	04/20/95
SOP for Digestion of AS/SE (GFAA)	MES01900.CT	10/02/95

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COMPUTER SYSTEMS

Standard Operating Procedures	Code	Date Generated
SOP for PCB EPA CLP Forms and Disk File	SYS00100.CT	05/25/89
SOP for LIMS Data Entry	SYS00201.CT	02/23/89
SOP for LIMS Data Entry Errors	SYS00301.CT	05/12/92
SOP for LIMS Data Base Security and Backup	SYS00400.CT	08/24/91
SOP for Testing, Modifying and Implementing Changes to Existing Computer Systems	SYS00502.CT	08/25/91
SOP for System Maintenance Operations and Response Time	SYS00600.CT	08/26/91
SOP for Lotus Diskette Deliverable	SYS00700.CT	02/25/92
SOP for Volatile Data Filter Program	SSY00800.CT	03/25/92
SOP for Metals Data Filter Program	SYS00900.CT	03/26/92
SOP for Classical Chemistry Results Program	SYS01000.CT	03/24/92
SOP for LIMS to PC File Transfer	SYS01100.CT	03/27/92
SOP for Classical Chemistry Completion Date Entry Program	SYS01200.CT	03/31/92
SOP for Hamilton Standard Diskette Deliverable	SYS01300.CT	04/01/92
SOP for Envision Software - Organic Deliverables	SYS01400.CT	03/27/92
SOP for Acres Diskette Deliverable	SYS01501.CT	12/01/92
SOP for Control Charts	SYS01600.CT	dft
SOP for CH2MHILL Diskette Deliverable	SYS01701.CT	02/23/93
SOP for AAS File Filter Program	SYS01800.CT	dft
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QUALITY ASSURANCE

Standard Operating Procedures	Code	Date Generated
SOP for Document Control	QAS00301.CT	05/08/92
SOP for Corrective Action Reports	QAS00501.CT	07/08/91
SOP for Internal Quality Assurance	QAS00400.CT	07/18/92
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SALES/MARKETING

Standard Operating Procedures	Code	Date Generated
SOP for Taking Client Orders	MKS00100.CT	02/21/94
SOP for LIMS Log-in	MKS00200.CT	02/21/94
SOP for Preparation for Price Quotations	MKS00300.CT	02/14/94
SOP for Telephone Logs	MKS00400.CT	06/22/94
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Date: 04/19/94 Page 11 of 13

CLASSICAL CHEMISTRY

Standard Operating Procedures	Code	Date Generated
Analysis of Tannins and Ligins in Environmental Samples	WC:042091:0	04/20/91
Analysis of Acidity (Method 305.2)	WC:033191:0	03/31/91
Analysis of Acidity (Method 305.1)	CVS00800.CT	03/24/94
Bromide (Method 405)	WC040791:0	04/07/91
Analysis of Hydrocarbons (418.1)	WC:041891:0	04/18/91
Analysis of Oil & Grease (Gravimetric)- 413.1	CVS01000.CT	03/29/94
Analysis of Salinity in Water	WC:070891:0	07/08/91
Analysis of Temperature in Water	WC:070591:0	07/05/91
Analysis of Grain Size	WC:071591:0	07/15/91
Measurement of Conductivity	WC:082190:0	08/21/90
Analysis of Dissolved Oxygen in Water	WC:071691:0	07/16/91
Analysis of Phosphorus in Water	WC:053191:0	05/31/91
Analysis of Alkalinity in Water - 310.1	CVS00700.CT	02/22/94
Analysis of Ammonia (method 350.1) in Water	WC:070791:0	07/07/92
Analysis of MBAS in Water	CVS00600.CT	03/31/94
Measurement of pH	CVS00900.CT	03/31/94
Analysis of Sulfide (9030)	CVS01700.CT	Dft
Analysis of Biochemical Oxygen Demand	CVS00500.CT	02/22/94
Analysis of COD (Method 410.4)	CVS01201.CT	08/17/94
Analysis of Hexavalent Chromium in cromite ore samples	WC:911205:0	12/05/91
Analysis of Samples for Total Cyanide CLP Protocol	CVS01100.CT	07/01/87
Analysis of Fluroide in Water (Method 340.2)	WC:051590.0	05/15/90
Total Organic Halides Analysis in Water Samples	WC:051490:0	05/14/90
Analysis of Total Organic Carbon in Water	WC:021390:1	02/13/90

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CLASSICAL CHEMISTRY (cont.)

Standard Operating Procedures	Code	Date Generated
Analysis of Hexavalent Chromium Colorimetric	WC:090192:0	09/01/92
Analysis of Hexavalent Chromium Alkaline digestion of Soil Samples	· WC:083192:0	08/31/92
Analysis of TOC Soil Samples	WC:102692:0	10/26/92
Analysis of TKN in Environmental Samples	WC:081090:0	08/10/90
Analysis of Hardness in Water	WC:081390:1	08/13/90
Analysis of Chloride (325.2) in Water	WC:081190:2	08/11/90
Analysis of Chloride (325.3) in Water	WC:040991:0	04/09/91
Analysis of Ammonia-Nitrogen in Environmental Samples	WC:021690:0	02/16/90
Standard Operating Procedure for Reactivity	CVS01900.CT	09/29/94
Standard Operating Procedure for Corrosivity	WC:011069:0	01/10/69
Standard Operating Procedure for Ignitability	WC:011889:0	01/18/89
Manual Spectrophotometric Method for Hexavalent Chromium	WC:110889:4	11/08/89
Analysis of Total Suspended Solids in Water	CVS00200.CT	08/21/93
Analysis of Sulfate in Water (Method 375.3)	CVS01300.CT	03/04/89
Analysis of Sulfate in Water (Method 375.4)	CVS01400.CT	Dft
EPTOX Leachate Procedure in Environmental Samples	WC:081090:0	08/10/90
Analysis of Total Dissolved Solids in Water	CVS00100.CT	08/16/93
Analysis of Nitrate and Nitrite for Water Samples (Method 353.2)	CVS02500.CT	05/03/90
Gravimetric Determination of Lube Oils in Solids	WC:062889:0	06/28/89
Total Recoverable Phenols Automated 4-AAP Method	WC:111389:0	11/13/89
Analysis of Environmental Samples for T- Phenols	WC:080186:1	08/01/86
Analysis of Samples for Chloride (SM407A)	WC:031189:0	03/11/89
Analysis of Environmental Samples for Formaldehyde	WC:072489:0	07/24/89
SOP for Total Cyanide - Method 335.4	CV\$02000.CT	10/04/94
SOP for Amenable Cyanide - Method 335.1	CVS02100.CT	10/04/94
SOP for Toxicity Characteristic Leaching Procedure - 1311	CVS01500.CT	09/28/94

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CLASSICAL CHEMISTRY (cont.)

Standard Operating Procedures	Code	Date Generated
Measurement of Turbidity in Water Samples	WC:082190:0	08/21/90
Shake Extraction of Solids for Wet Chemistry Analysis	WC:041391:0	04/13/91
COD (410.1)	WC:082290:1	08/22/90
SOP for WC Data Reporting/Validation	CVS00400.CT	08/29/93
SOP for Total Solids	CVS00300.CT	08/21/93
SOP for Flashpoint - Method 1010	CVS01600.CT	09/28/94
SOP for Waste Extraction Test (WET) Procedure	CVS01800.CT	09/28/94
SOP for Cation/Anion Balance	CVS02800.CT	3/20/95
SOP for CEC Method 9081	CVS02900.CT	3/20/95
SOP for Soil Homogenization	CVS03000.CT	3/20/95
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APPENDIX, Section 8

LISTING OF ANALYTICAL METHODS AND ASSOCIATED DETECTION LIMITS

COMPONENT	LCS/QC CHECK % RECOVERY	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD)	MDL ug/L	PQL ug/L
	Method 8080 Organoc	hlorine Pesticides i	n Water		
alpha-BHC	37-134	-	-	.005	0.05
beta-BHC	17-147	-	-	.017	0.05
delta-BHC	19-140	-		.004	0.05
gamma-BHC (Lindane)	32-127	56-123	20	.004	0.05
Heptachlor	34-111	40-131	20	.012	0.05
Aldrin	42-122	40-120	20	.008	0.05
Heptachlor epoxide	37-142			.005	0.05
Endosulfan I	45-153	-		.005	0.05
Dieldrin	36-146	52-126	20	.006	0.1
4,4'-DDE	30-145		-	.008	0.1
Endrin	30-147	56-121	20	.005	0.1
Endosulfan II	D-202		-	.015	0.1
4,4' DDD	31-141	-	-	.010	0.1
Endosulfan sulfate	26-144		-	.022	0.1
4,4'-DDT	25-160	38-127	20	.011	0.1
Methoxychlor	50 - 168	-	-	.019	0.5
Toxaphene		-	-	2.0	0.5
Aroclor 1016	-	-	-	.008	1.0
Aroclor 1221		-	-	.014	2.0
Aroclor 1232			•	.031	1.0
Aroclor 1242	33-128	-		.029	1.0
Aroclor 1248	-	·	-	.022	1.0
Aroclor 1254	-	-	-	.060	1.0
Aroclor 1260	41-116	15-1 <u>75</u>	20	.025	1.0
Chlordane (technical)	-	<u> </u>	-	.0.158	0.2
Endrin aldehyde	44-154	<u> </u>	-	.010	0.1
Endrin ketone	30-150	<u>-</u>		.006	0.1
<u> </u>	Method 8150 Chlorin	l ated Herbicide	l s in Water		
2,4-D	_ 50-176	10-200	20	0.173	0.50
Silvex (2,4,5-TP)	10-134	10-197	20	0.015	0.50
2,4,5-T	10-146	_	_	0.018	0.50

COMPONENT	LCS/QC CHECK % RECOVERY	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD)	MDL ug/Kg	PQL ug/Kg			
Method 8080 Organochlorine Pesticides in Soil								
alpha-BHC	37-134	-	-	0.195	1.7			
beta-BHC	17-147	-	-	0.396	1.7			
deita-BHC	19-140	-	<u>-</u>	0.094	1.7			
gamma-BHC (Lindane)	32-127	46-127	20	0.196	1.7			
Heptachlor	34-111	35-130	20	0.466	1.7			
Aldrin	42-122	40-120	20	0.146	1.7			
Heptachlor epoxide	37-142	-	-	0.142	1.7			
Endosulfan I	45-153	-		0.270	1.7			
Dieldrin	36-146	31-134	20	0.176	3.3			
4,4'-DDE	30-145	-		0.188	3.3			
Endrin	30-147	42-139	20	0.205	3.3			
Endosulfan II	D-202	-	<u> </u>	2.03	3.3			
4,4' DDD	31-141	<u>-</u>		2.30	3.3			
Endosulfan sulfate	26-144	· -	-	0.224	3.3			
4,4'-DDT	25-160	23-134	20	1.580	3.3			
Methoxychlor	50 - 168	-	-	0.511	17			
Toxaphene	-			3.86	17			
Aroclor 1016	•		-	1.28	33			
Aroclor 1221	-	_		3.69	67			
Aroclor 1232	-		-	2.54	33			
Aroclor 1242	33-128	<u>-</u>	-	1.19	_ 33			
Aroclor 1248			-	4.79	33			
Aroclor 1254		<u>-</u>	-	2.62	33			
Aroclor 1260	41-116	10-175	20	1.05	33			
Chlordane (technical)		<u>-</u>	-	ns	6.7			
Endrin aldehyde	44-154			0.81	3.3			
Endrin ketone	30-150	-	-	0.247	3.3			
	Method 8150 Chlori	nated Herbicid	es in Soil					
2,4-D	50-176	10-200	20	9.25	20			
Silvex (2,4,5-TP)	10-134	10-197	20	13.0	20			
2,4,5-T	10-146		<u>-</u>	1.11	5.0			

		,_ 	Metals	,. 	_ 	
COMPONENT	SAMPLE MATRIX	ANALYTICAL METHOD	PRECISION %RSD	ACCURACY % RECOVERY	UNITS	PQL
Aluminum	Water	200.7	0-20	90-110	ug/l	200
	Water	6010	0-20	90-110	ug/L	200
	Soil	6010	0-20	90-110	mg/Kg	40
Antimony	Water	200.7	0-20	90-110	ug/L	60
	Water	6010	0-20	90-110	ug/L	60
	Soil	6010	0-20	90-110.	mg/Kg	12
	Water	204.2	0-20	80-120	ug/L	
	Soil	7412	0-20	80-120	mg/Kg	
Arsenic	Water	200.7	0-20	90-110	ug/L	10
	Water	6010	0-20	90-110	ug/L	10
	Water	206.2	0-20	80-120	ug/L	10
	Water	7060	0-20	80-120	ug/L	10
	Soil	7060	0-20	80-120	mg/ Kg	2.0
	Soil	6010	0-20	90-110	mg/Kg	2.0
Barium	Water	200.7	0-20	90-110	ug/l	200
	Water	6010	0-20	90-110	ug/L	200
	Soil	6010	0-20	90-110	mg/Kg	40
Beryllium	Water	200.7	0-20	90-110	ug/l	5.0
	Water	6010	0-20	90-110	ug/L	5.0
	Soil	6010	0-20	90-110	mg/Kg	1.0
Cadmium	Water	200.7	0-20	90-110	ug/l	5.0
	Water	6010	0-20	90-110	ug/L	5.0
	Soil	6010	0-20	90-110	mg/Kg	1.0
Calcium	Water	200.7	0-20	90-110	ug/l	5000
	Water	6010	0-20	90-110	ug/L	5000
	Soil	6010	0-20	90-110	mg/Kg	1000
Cobalt	Water	200.7	0-20	90-110	ug/l	50
<u> </u>	Water	6010	0-20			
	Soil	6010	0-20	90-110	ug/L	50
Chenerius				90-110	mg/Kg	10
Chromium	Water	200.7	0-20	90-110	ug/l	10
	Water	6010	0-20 0-20_	90-110	ug/L	10

			Metals	T		
COMPONENT	SAMPLE MATRIX	ANALYTICAL METHOD	PRECISION %RSD	ACCURACY % RECOVERY	UNITS	PQL
Соррег	Water	200.7	0-20	90-110	ug/l	25
	Water	6010	0-20	90-110	ug/L	25
	Soil	6010	0-20	90-110	mg/Kg	<u>5</u> .0
Iron	Water	200,7	0-20	90-110	ug/l	100
	Water	6010	0-20	90-110	ug/L	100
	Soil	6010	0-20	90-110	mg/Kg	20
Lead	Water	200.7	0-20	90-110	ug/L	3.0
	Water	239.2	0-20	80-120	ug/L	3.0
	Water	7421	0-20	80-120	ug/l	3.0
	Water	6010	0-20	90-110	ug/L	3.0
	Soil	6010	0-20	90-110	mg/Kg	0.6
	Soil	7421	0-20	90-110	mg/Kg	0.6
Magnesium	Water	200.7	0-20	90-110	ug/l	5000
	Water	6010	0-20	90-110	ug/L	5000
	Soil	6010	0-20	90-110	mg/Kg	1000
Manganese	Water	200.7	0-20	90-110	ug/L	15
	Water	6010	0-20	90-110	ug/L	15
	Soil	6010	0-20	90-110	mg/Kg	3.0
Molybdenum	Water	200.7	0-20	90-110	ug/l	20
	Water	6010	0-20	90-110	ug/L	20
	Soil	6010	0-20	90-110	mg/Kg	4.0
Mercury	Water	245.1	0-20	80-120	ug/L	0.2
	Water	7470	0-20	80-120	ug/L	0.2
	Soil	7471	0-20	80-102	mg/Kg	0.1
Nickel	Water	200.7	0-20	90-110	ug/L	40
	Water	6010	0-20	90-110	ug/L	40
	Soil	6010	0-20	90-110	mg/Kg	8.0
Potassium	Water	200.7	0-20	90-110	ug/L	5000
	Water	6010	0-20	90-110	ug/L	5000
	Soil	6010	0-20	90-110	mg/Kg	1000
Selenium	Water	200.7	0-20	90-110	ug/L	5.0
	Water	270.2	0-20	80-120	ug/L	5.0
	Water	6010	0-20	90-110	ug/L	5.0

COMPONENT	SAMPLE MATRIX	ANALYTICAL METHOD	PRECISION %RSD	ACCURACY % RECOVERY	UNITS	PQL			
	Water	7740	0-20	80-120	ug/L	5.0			
	Soil	7740	0-20	80-120	mg/Kg	1.0			
	Soil	6010	0-20	90-110	mg/Kg	1.0			
<u>Sil</u> ver	Water	200.7	0-20	90-110	ug/L	10			
	Water	6010	0-20	90-110	ug/L	10			
	Soil	6010	0-20	90-110	mg/Kg	2.0			
Sodium	Water	200.7	0-20	90-110	ug/L	5000			
	Water	6010	0-20	90-110	ug/L	5000			
	Soil	6010	0-20	90-110	mg/Kg	1000			
Thallium	Water	200.7	0-20	90-110	ug/L	10			
	Water	6010	0-20	90-110	ug/L	10			
	Water	279.2	0-20	80-120	ug/L	10			
	Water	7841	0-20	80-120	ug/L	10			
	Soil	7841	0-20	80-120	mg/Kg	2.0			
	Soil	6010	0-20	90-110	mg/Kg	2.0			
Tin	Water	200.7	0-20	90-110	ug/L	50			
	Water	6010	0-20	90-110	ug/L	50			
	Soil	6010	0-20	90-110	mg/Kg	10			
Titanium	Water	200.7	0-20	90-110	ug/L	20			
	Water	6010	0-20	90-110	ug/L	20			
	Soil	6010	0-20	90-110	mg/Kg	4.0			
Zinc	Water	200.7	0-20	90-110	ug/L	20			
	Water	6010	0-20	90-110	ug/L	20			
	Soil	6010	0-20	90-110	mg/Kg	4.0			
Vanadium	Water	200.7	0-20	90-110	ug/L	50			
	Water	6010	0-20	90-110	ug/L	50			
	Soil	6010	0-20	90-110	mg/Kg	10			

⁽¹⁾ Acceptance limits are those indicated in the published method data.

	_	W	et Chemistry			
COMPONENT	SAMPLE MATRIX	ANALYTICAL METHOD	PRECISION %RSD	ACCURACY % RECOVERY	UNITS	PQL
Acidity	Water	305.1	0-20-	NA NA	mg/L	1.0
Alkalinity	Water	310.1	0-20	NA NA	mg/L	2.0
Ammonia-N	Water	350.3	0-20	75-125	mg/L	0.04
Bicarbonate	Water	406C	0-20	NA	mg/L	1.0
Biochemical Oxygen Demand (BOD)	Water	405.1	0-20	75-125	mg/L	2.0
Bromide	Water	320.1	0-20	75-125	mg/L	2.0
Bromide	Water	405	0-20	75-125	mg/L	0.50
Chloride	Water	325.2	0-20	75-125	mg/L	3.0
Chlorine Demand	Water	3-364	0-20	_NA	mg/L	1.0
Chlorine Residual	Water	330.4	0-20	NA	mg/L	0.1
Chemical Oxygen Demand (COD)	Water	410.4	0-20	75-125	mg/L	10.0
Color	Water	110.2	0-20	NA	Pt-Co	5.0
Conducitivity	Water	120.1	0-20	NA	umho/cm	NA NA
Chromium (VI)	Water	7196	0-20	75-125	mg/L	0.01
Cyanide-Total	Water	335.4	0-20	75-125	ug/L	10.0
Cyanide-Total	Water	9012	0-20	75-125	ug/L	10.0
Cyanide-Amenable	Water	335.1	0-20	75-125	ug/L	10.0
Cyanide-CLP	Water	ILM04	0-20	75-125	ug/L	10.0
Dissolved Oxygen	Water	360.1	0-20	NA	mg/L	0.1
Flashpoint	Water	1010	0-20	75-125		-
Fluoride	Water	340.2	0-20	75-125	mg/L	0.10
Grain Size	Water	D442-63	0-20	NA NA		
Hardness	Water	130.2	0-20	75-125	mg/L	1.0
Hydrocarbons (Grav.)	Water	503E	0-20	75-125	mg/L	1.0
Hrdrocarbons (IR)	Water	418.1	0-20	75-125	mg/L	1.0
MBAS	Water	425,1	0-20	<u>75-1</u> 25	mg/L	0.04
Nitrate-Nitrite-N	Water	353.2	0-20	75-125	mg/L	0.10
Nitrate-N	Water	353.2	0-20	<u>75</u> -125	mg/L	0.005
Odor	Water	140.1	0-20	<u>N</u> A	NA_	-
Oil & Grease (Grav.)	Water	413.1	0-20	75-125	mg/L	1.0
Oil &Grease (IR)	Water	413.2	0-20	75-125	mg/L	1.0

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-		w	et Chemistry			
COMPONENT	SAMPLE MATRIX	ANALYTICAL METHOD	PRECISION %RSD	ACCURACY % RECOVERY	UNITS	PQL
Paint Filter Test	Water	9095	0-20	75-125	NA	-
рН	Water	150.1	0-20	NA	NA	-
рН	Water	9040	0-20	NA_	NA	-
Phenols	Water	420.2	0-20	75-125	mg/L	0.005
Phenois	Water	9066	0-20	75-125	mg/L	0.005
Phosphorus	Water	365.2	0-20	75-125	mg/L	0.10
Phosphate (Ortho)	Water	365.2	0-20	75-125	mg/L	0.10
Settable solids	Water	160.5	0-20	NA	mL/L	1.0
Silica	Water	370.1	0-20	75-125	mg/L	1.0
Specific Gravity	Water	3-61	0-20	75-125	NA	_
Sulfate	Water	375.3	0-20	75-125	mg/L	10.0
Sulfate	Water	375.4	0-20	75-125	mg/L	10.0
Sulfide	Water	376.1	0-20	75-125	mg/L	1.0
Sulfide	Water	9030	0-20	75-125	mg/L	1.0_
Sulfite	Water	377.1	0-20	-	mg/L	1.0
Sludge Volume Index	Water	213C	0-20	-	ml/mg	1.0
Total Kjeldahl Nirogen	Water	351.2	0-20	75-125	mg/L	1.0
Total Kjeldahl Nitrogen	Water	351.1	0-20	75-12 5	mg/L	1.0
Total Soilds	Water	160.3	0-20	N/A	mg/L	1.0
Total Dissolved Solids	Water	160.1	0-20	N/A	mg/L	5.0
Total Suspended Solids	Water	160.2	0-20	N/A	mg/L	5.0
Total Volatile Solids	Water	160.4	0-20	N/A	mg/L	1.0
Total Organic Carbon	Water	415.2	0-20	75-125	mg/L	0.5
Total Organic Halides	Water	9020	0-20	75-125	ug/L	10.0
Turbidity	Water	180.1	0-20	-	NTU	0.10
Cyanide	Soil	ILM04	0-20	75-125	mg/Kg	0.5
Total Organic Carbon	Soil	9060M	0-20	75-1 <u>25</u>	mg/Kg	100
Corrosivity Char.	Soil	9045	-		•	-
Ignitability Char.	Soil	BRT			•	
TCLP	W/S	1311	-	-	-	•
SPLP	W/S	1312	•	_		

^{*} Acceptance limits are those indicated in the published method data.

COMPONENT	ACCURACY % RECOVERY	MATRIX SPIKE % RECOVERY	MDL ug/L	PQL ug/L				
Method 608 Organochlorine Pesticides in Water								
alpha-BHC	37-134	26-126	.005	.005				
beta-BHC	17-147	54-140	.017	.017				
delta-BHC	19-140	3-113	.004	.004				
gamma-BHC (Lindane)	32-127	47-123	.004	.004				
Heptachlor	<u>34-</u> 111	26-119	.012	.012				
Aldrin	42-122	53-104	.008	.008				
Heptachlor epoxide	37-142	59-125	.005	.005				
Endosulfan I	45-153	69-138	.005	.005				
Dieldrin	36-146	50-136	.006	.006				
4,4'-DDE	30-145	73-104	.008	.008				
Endrin	30-147	52-154	.005	.005				
Endosulfan II	D-202	18-124	.015	.015				
4,4' DDD	31-141	10-163	.010	.010				
Endosulfan sulfate	26-144	59-152	.022	.022				
4,4'-DDT	25-160	51-140	.011	.011				
Methoxychlor	62-181	62-181	.019	.019				
Toxaphene	41-126		2.0	2.0				
Aroclor 1016	50-114		008	.008				
Aroclor 1221	15-178		.014	.014				
Aroclor 1232	10-215	-	.031	.031				
Aroclor 1242	39-150	-	.029	.029				
Aroclor 1248	38-158	-	.022	.022				
Aroclor 1254	29-131		.060	.060				
Aroclor 1260	8-127	·	.025	.025				
Chlordane (technical)	45-119		0.158	0.158				
Endrin aldehyde	30-164	30-164	.010	.010				
Endrin ketone	30-150	30-150	.006	.006				
<u> </u>				L				

GC/MS Volatile Organics								
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/Kg)	PQL (ug/Kg)			
	Method	8240 Purgeables	in Soil					
Acetone	14-187	<u> </u>	-	4.3	10			
Benzene	70-145	66-142	21	0.5	5			
Bromodichloromethane	70-125		-	0.4	5			
Bromoform	45-169		-	1.0	5			
Bromomethane	13-145	-	•	2.9	10			
2-Butanone	D-251	-	-	3.5	10			
Carbon disulfide	D-475	-	-	1.0	5			
Carbon tetrachloride	70-149	-	-	0.6	5			
Chlorobenzene	90-135	60-133	21	0.8	5			
Dibromochloromethane	70-130	-	-	0.4	5			
Chloroethane	14-230	-	-	1.6	10			
2-Chloroethylvinyl ether	D-305	<u> </u>	-	0.9	10			
Chloroform	80-135	-	-	1.0	5			
Chloromethane	D-273		-	2.1	10			
1,1-Dichloroethane	75-135	59-172	22	0.8	5			
1,2-Dichloroethane	<u>6</u> 5-135	-	-	1.0	5			
1,1-Dichloroethene	70-125	-	-	1.4	5			
1,2-Dichloroethene (total)	68-132	-	-	1.4	5			
1,2-Dichloropropane	75-145	-	-	0.7	5			
cis-1,3-Dichloropropene	70-113	<u> </u>	-	0.5	5			
trans-1,3-Dichloropropene	70-113	<u>-</u>		0.8	5			
Ethylbenzene	75-130	<u> </u>	•	0.6	5			
2-Hexanone	28-170	<u>-</u>	-	1.4	10			
Methylene chloride	50-160	•	-	4.1	5			
4-Methyl-2-pentanone	60-170	-	-	1.0	10			
Styrene	80-120	-	-	0.7	5			
1,1,2,2-Tetrachloroethane	65-130	•	-	0.8	5			
Tetrachloroethene	64-148		_	1.1	5			
Toluene	31-130	50-139	21	0.9	5			
1,1,1-Trichloroethane	80-120	<u> </u>	_	0.6	5			

GC/MS Volatile Organics								
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/Kg)	PQL (ug/Kg)			
1,1,2-Trichloroethane	85-130	-	-	1.0	5			
Trichloroethene	70-135	62-137	24	1.0	5			
Vinyl acetate	8-118	-	-	1.3	10			
Vinyl chloride	1-240	•		1.1	10			
Xylenes (total)	55-172	-	_	2.6	5			

GC/MS Extractable Organics							
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/l)	PQL (ug/l)		
	Method 82	70 Extractables in	n Water				
Acenaphthene	47-145	46-118	31	2.5	10		
Acenaphthylene	33-145	-		2.3	10		
Anthracene	27-133	-	-	1.6	10		
Benzoic acid	D-473	-	-	ns	50		
Benzo(a)anthracene	33-143	-	-	1.7	10		
Benzo(b)fluoranthene	24-139	-	-	1.6	10		
Benzo(k)fluoranthene	11-162	<u> </u>	-	3.3	10		
Benzo(g,h,i)perylene	D-219	<u>-</u>	-	6.4	10		
Benzo(a)pyrene	17-163	-	-	1.8	10		
Benzyl alcohol	D-130	-	-	1.5	10		
bis(2-Chloroethoxy)methane	33-184	•	-	2.9	10		
bis(2-Chloroethyl)ether	12-138	-	-	1.7	10		
bis(2-Chloroisopropyl)ether	36-166	-	-	5.4	10		
bis(2-Ethylhexyl)phthalate	8-138	-	_	2.9	10		
4-Bromophenyl phenyl ether	53-127	-	-	1.1	10		
Benzyl butyl phthalate	D-132	-	_	2.1	10		
4-Chloroaniline	1-78	-	-	3.2	10		
2-Chloronaphthalene	60-118	-	-	2.4	10		
4-Chloro-3-methylphenol	44-294	23-97	42	1.6	10		
2-Chlorophenol	46-268	27-123	40	0.8	10		
4-Chlorophenyl phenyl ether	25-138	-	-	2.5	10		
Chrysene	17-168	-	-	2.5	10		
Dibenzo(a,h)anthracene	D-227	_	_	4.1	10		
Dibenzofuran	D-170	_	-	2.4	10		
Di-n-butylphthalate	1-118	-	-	1.9	10		
1,3-Dichlorobenzene	D-172	-	-	1.9	10		
1,4-Dichlorobenzene	20-124	36-97	28	2.4	10		
1,2-Dichlorobenzene	32-129	-	-	2.3	10		
3,3'-Dichlorobenzidine	D-52	•		2.8	10		
2,4-Dichlorophenol	78-270			2.1	10		
Diethyl phthalate	D-114	-		2.1	10		

	GC/M	IS Extractable Organic	cs		\ -
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/l)	PQL (ug/l)
2,4-Dimethylphenol	64-238	-	<u> </u>	1.3	10
Dimethyl phthalate	D-112			1.8	10
4,6-Dinitro-2-methylphenol	D-362	-	<u> </u>	2.3	25
2,4-Dinitrophenol	D-382	-	<u> </u>	1.7	25
2,4-Dinitrotoluene	39-139	24-96	38	2.2	10
2,6-Dinitrotoluene	50-138		<u> </u>	2.8	10?
Di-n-octylphthalate	4-146		-	1.0	10
Fluoranthene	26-137	-	<u>-</u>	1.4	_10
Fluorene	59-121	<u>.</u>		3.0	10
Hexachlorobenzene	D-132			0.8	10
Hexachlorobutadiene	24-116			3.0	10
Hexachlorocyclopentadiene	D-59		<u> </u>	1.1	10
Hexachloroethane	40-113 .		-	1.5	10
Indeno(1,2,3-cd)pyrene	D-171		-	5.4	10
Isophorone	21-196	<u> </u>	-	2.6	10
2-Methylnaphthalene	D-127	-	<u>-</u>	2.7	10
2-Methylphenol (o-cresol)	40-189	<u>-</u>	-	1.4	10
4-Methylphenol (p-cresol)	28-198		<u> </u>	7.3	10
Naphthalene	21-133		<u> </u>	2.6	_10
2-Nitroaniline	D-127	<u>-</u>	<u> </u>	2.0	25
3-Nitroaniline	D-91	<u>-</u>	<u> </u>	20.6	25
4-Nitroaniline	D-108		<u>-</u>	2.5	20
Nitrobenzene	35-180	<u> </u>		2.5	10
2-Nitrophenol	58-364		<u> </u>	1.5	10
4-Nitrophenol	D-264	10-80	50	0.6	25
N-Nitroso-di-n-propylamine	D-230	41-116	38	3,3	10
N-Nitrosodiphenylamine	D-114		<u>-</u>	1.2	10
Pentachlorophenol	28-352	9-103	50	22.6	25
Phenanthrene	54-120	<u> </u>		1.4	10
Phenol	10-224	12-110	42	0.9	10
Pyrene	52-113	26-127	31	2.4	10
1,2,4-Trichlorobenzene	44-142_	39-98	28	2.7	10

GC/MS Extractable Organics					
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/l)	PQL (ug/l)
2,4,5-Trichlorophenol	82-354			1.4	25
2,4,6-Trichlorophenol	74-288		<u>-</u>	1.4	25_

·····	GC/M	IS Extractable Organi	cs		-
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/Kg)	PQL (ug/Kg)
	Method 8	270 Extractables	in Soil		
Acenaphthene	47-145	31-137	19	12.6	330
Acenaphthylene	33-145	<u> </u>	<u> </u>	12.8	330
Anthracene	27-133	-	<u> </u>	13.0	330
Benzoic acid	D-473		<u> </u>	100	1600
Benzo(a)anthracene	33-143	.	<u> </u>	14.6	330
Benzo(b)fluoranthene	24-139	-	<u> </u>	23.0	330
Benzo(k)fluoranthene	11-162	_	<u> </u>	16.5	330
Benzo(g,h,i)perylene	D-219			17.8	330
Benzo(a)pyrene	17-163		<u> </u>	16.2	330_
Benzyl alcohol	D-130	<u> </u>	_	13.4	330
bis(2-Chloroethoxy)methane	33-184			11.7	330
bis(2-Chloroethyl)ether	12-138	<u>-</u>	-	17.0	330
bis(2-Chloroisopropyl)ether_	36-166	<u> </u>	-	15.2	330
bis(2-Ethylhexyl)phthalate	8-138	<u> </u>		20.8	330
4-Bromophenyl phenyl ether	53-127	<u> </u>	<u> </u>	12.9	330
Benzyl butyl phthalate	D-132	<u>-</u>	<u> </u>	11.4	330
4-Chloroaniline	1-78		<u> </u>	47.8	330
2-Chloronaphthalene	60-118	-	<u> </u>	14.3	330
4-Chloro-3-methylphenol	44-294	26-103	33	14.5	330
2-Chlorophenol	46-268	25-102	50	12.0	330
4-Chlorophenyl phenyl ether	25-138	-		14.6	330
Chrysene	17-168		<u> </u>	15.1	330
Dibenzo(a,h)anthracene	D-227	-	<u> </u>	18.5	330
Dibenzofuran	D-170	-	-	16.7	330
Di-n-butylphthalate	1-118	-		16.8	330
1,3-Dichlorobenzene	D-172	<u> </u>		14.0	330
1,4-Dichlorobenzene	20-124	28-104	27	16.7	330
1,2-Dichlorobenzene	32-129		-	13.4	330
3,3'-Dichlorobenzidine	D-52			10.4	660_
2,4-Dichlorophenol	78-270		-	10.2	330
Diethyl phthalate	D-114			17.1	330

	GC	/MS Volatile Organi	cs		
	LFB % RECOVERY LIMIT	LAB MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/l)	PQL (ug/l)
	Method 524.2 I	ow Level Purge	ables in Water		
Benzene	80-120	80-120	13	0.31	1.0
Bromobenzene	80-120	_80-120	13	0.34	1.0_
Bromochloromethane	80-120	80-120	13	0.43	1.0_
Bromodichloromethane	80-120	80-120	13	0.40	1.0
Bromoform	80-120	80-120	13	0.43	1.0
Bromomethane	80-120	80-120	13	0.27	1.0
n-Butylbenzene	80-120	80-120	13	0.33	1.0
sec-Butylbenzene	80-120	80-120	13	0.30	1.0
tert-Butylbenzene	80-120	80-120	13	0.28	1.0
Carbon tetrachloride	80-120	80-120	13	0.28	1.0
Chlorobenzene	80-120	80-120	13	0.32	1.0
Chloroethane	80-120	80-120	13	0.28	1.0
Chloroform	80-120	80-120	13	0.48	1.0
Chloromethane	80-120	80-120	13	0.37	1.0
2-Chlorotoluene	80-120	80-120	13	0.33	1.0
4-Chlorotoluene	80-120	80-120	13	0.34	1.0
Dibromochloromethane	80-120	80-120	13	0.45	1.0
1,2-Dibromo-3- chloropropane	80-120	80-120	13	0.46	1.0
1,2-Dibromoethane	80-120	80-120	13	0.46	1.0
Dibromomethane	80-120	80-120	13	0.43	1.0
1,2-Dichlorobenzene	80-120	80-120	13	0.36	1.0
1,3-Dichlorobenzene	80-120	80-120	13	0.35	1.0
1,4-Dichlorobenzene	80-120	80-120	13	0.37	1.0
Dichlorodifluoromethane	80-120	80-120	13	0.29	1.0
1,1-Dichloroethane	80-120	80-120	13	0.33	1.0
1,2-Dichloroethane	80-120	80-120	13	0.42	1.0_
1,1-Dichloroethene	80-120	80-120	13	0.31	1.0
cis-1,2-Dichloroethene	80-120	80-120	13	0.33	1.0
trans-1,2-Dichloroethene	80-120	80-120	13	0.31	1.0

	7 F7D		DO.		
	LFB % RECOVERY LIMIT	LAB MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/l)	PQL (ug/l)
1,2-Dichloropropane	80-120	80-120	13	0.38	1.0
1,3-Dichloropropane	80-120	80-120	13	0.48	1.0
2,2-Dichloropropane	80-120	80-120	13	0.27	1.0
1,1-Dichloropropene	80-120	80-120	13	0.32	1.0
Ethylbenzene	80-120	80-120	13	0.30	1.0
Hexachlorobutadiene	80-120	80-120	13	0.47	1.0
Isopropylbenzene	80-120	80-120	13	0.28	1.0
p-Isopropyltoluene	80-120	80-120	13	0.29	1.0
Methylene chloride	80-120	80-120	13	0.38	1.0
Naphthalene	80-120	80-120	13	0.50	1.0
n-Propylbenzene	80-120	80-120	13	0.28	1.0
Styrene	80-120	80-120	. 13	0.31	1.0
1,1,1,2-Tetrachloroethane	80-120	80-120	13	0.36	1.0
1,1,2,2-Tetrachloroethane	80-120	80-120	13	0.40	1.0
Tetrachloroethene	80-120	80-120	13	0.29	1.0
Toluene	80-120	80-120	13	0.30	1.0
1,2,3-Trichlorobenzene	80-120	80-120	13	0.38	1.0
1,2,4-Trichlorobenzene	80-120	80-120	13	0.50	1.0
1,1,1-Trichloroethane	80-120	80-120	13	0.34	1.0
1,1,2-Trichloroethane	80-120	80-120	13	0.45	1.0
Trichloroethene	80-120	80-120	13	0.38	1.0
Trichlorofluoromethane	80-120	80-120	13	0.31	1.0
1,2,3-Trichloropropane	80-120	80-120	13	0.38	1.0
1,2,4-Trimethylbenzene	80-120	80-120	13	0.31	1.0
1,3,5-Trimethylbenzene	80-120	80-120	13	0.30	1.0
Vinyl chloride	80-120	80-120	13	0.35	1.0
o-Xylene	80-120	80-120	13	0.31	1.0
m/p-xylene	80-120	80-120	13	0.56	1.0

GC/MS Volatile Organics						
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	MDL (ug/l)	PQL (ug/l)		
	Method 624 Pur	geables in Water				
Benzene	37-151	78-122	1.2	5		
Bromodichloromethane	35-155	82-117	1.1	5		
Bromoform	45-169	61-136	1.0	5		
Bromomethane	d-242	67-122	1.5	10		
Carbon tetrachloride	70-140	76-127	1.1	5		
Chlorobenzene	37-160	78-117	1.0	5		
Chloroethane	14-230	79-118	2.2	10		
2-Chloroethylvinyl ether	D-305	10-305	2.0	5		
Chloroform	51-138	83-114	1.5	5		
Chloromethane	D-273	35-152	1.2	10		
Dibromochloromethane	53-149	78-122	1.2	5		
1,2-Dichlorobenzene	18-190	18-190	1.0	5		
1,3-Dichlorobenzene	59-156	59-156	0.4	5		
1,4-Dichlorobenzene	18-190	60-145	0.8	5		
1,1-Dichloroethane	59-155	81-181	1.2	_5		
1,2-Dichloroethane	49-155	80-123	1.2	5		
1,1-Dichloroethene	D-234	79-121	1.2	5		
1,2-Dichloroethene (total)	54-156	<u>85</u> -113	1.6	5		
1,2-Dichloropropane	D-210	77-124	1.4	5		
cis-1,3-Dichloropropene	D-227	75-110	1.4	5		
trans-1,3-Dichloropropene	17-183	73-132	1.1	5		
Ethylbenzene	37-162	83-112	1.4	5		
Methylene chloride	D-221	83-115	4.1	5		
1,1,2,2-Tetrachloroethane	46-157	69-138	1.3	5		
Tetrachloroethene	64-148	76-121	0.8	5_		
Toluene	47-150	77-117	1.0	5		
1,1,1-Trichloroethane	52-162	72-130	1.2	5		
1,1,2-Trichloroethane	52-150	71-126	1.1			

·	GC/MS Volatile Organics							
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	MDL (ug/l)	PQL (ug/l)				
Trichloroethene	71-157	82-116	0.8	5				
Trichlorofluoromethane	17-181	17-181	1.3	5				
Vinyl Chloride	D-251	65-127	1.3	10				

	GC	MS Volatile Organic	<u>s</u>	_	
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/L)	PQL (ug/l)
	Method 8	240 Purgeables in	n Water		
Acetone	28-374	·	_ ·	9.9	10
Benzene	14.2-25.8	76-127	11	0.4	5
Bromodichloromethane	13.1-26.9	<u> </u>		1.2	5
Bromoform	14.2-25.8_			1.2	5
Bromomethane	D-37.2	<u>-</u> _		1.0	10
2-Butanone	D-502		-	2.2	10
Carbon disulfide	D-95	<u> </u>		1.2	5
Carbon tetrachloride	14.6-25.4		-	0.9	5
Chlorobenzene	13.2-26.8	75-130	13	0.7	5
Dibromochloromethane	13.5-26.5	<u> </u>	-	0.8	5
Chloroethane	7.6-32.4	<u>-</u>		_1.0	10
2-Chloroethylvinyl ether	D-44.8			0.9	10
Chloroform	13.5-26.5	_	-	1.8	5
Chloromethane	D-40.8	<u>-</u>	-	1.1	10
1,1-Dichloroethane	14.5-25.2	61-145	14	0.9	5
1,2-Dichloroethane	13.6-26.4	_		0.9	5
1,1-Dichloroethene	10.1-29.9		-	1.2	5
1,2-Dichloroethene (total)	13.9-26.1		<u></u>	0.9	5
1,2-Dichloropropane	6.8-33.2	<u> </u>	-	0.8	5_
cis-1,3-Dichloropropene	1.8-13.3		<u> </u>	_0.6	5
trans-1,3-Dichloropropene	9.9-48	<u> </u>	<u>-</u>	0.5	5
Ethylbenzene	11.8-28.2	<u>-</u>		0.4	5
2-Hexanone	11-68	<u>-</u>	-	1.4	10
Methylene chloride	12.1-27.9	<u> </u>		1.5	5
4-Methyl-2-pentanone	24-68			1.0	10
Styrene	16-24	<u>-</u>		0.6	5
1,1,2,2-Tetrachloroethane	12.1-27.9	<u> </u>		1.1	5
Tetrachloroethene	14.7-25.3	<u> </u>		1.7	5
Toluene	14.9-25.1	76-125	13	0.5	5
1,1,1-Trichloroethane	13-25			0.9	_ 5 _

Date: 03/24/95

GC/MS Volatile Organics					
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/L)	PQL (ug/I)
1,1,2-Trichloroethane	12.8-27.2	-	-	0.7	5
Trichloroethene	13.3-26.9	71-120	14	0.8	5
Vinyl acetate	3.2-47	_	-	2.1	10
Vinyl chloride	0.8-39.2		-	1.1	10
Xylenes (total)	33-103	-		1.3	5



IEA - CT LABORATORY QUALIFICATIONS AND EXPERIENCE STATEMENT

200 Monroe Turnpike Monroe, CT 06468 Phone (203) 261-4458 Fax (203) 268-5346

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I INTRODU**CTIO**N

IEA is full service environmental laboratory serving both the public and private sectors with facilities in Whippany, NJ, Monroe, CT, Schaumburg, IL, Sunrise, N. Billerica, MA, and 2 in Research Triangle Park, NC, including a Radiological facility.

IEA performs analyses following EPA protocols and produces deliverable packages to comply with the current regulatory requirements of CERCLA, RCRA, NPDES, Safe Drinking Water Act, Clean Air Act and State ECRA type programs. Our services start with a technical Marketing/Sales force that can tailor a proposal to your project needs, and includes dedicated project managers, documented bottle preparation, sample pick-up, and expert data interpretation, in addition to the actual analyses and data report.

The following pages contain a closer look at IEA's capabilities and credentials.

II. LABORATORY CAPABILITIES

There are a number of environmental regulations that require analytical testing. Each of these has specific objectives and most have established test methodologies. IEA has developed testing capabilities to conform with the requirements of these regulations. In order to implement this type of program, we have also developed a technically oriented Client Service and Account Executive Staff. These individuals understand the differences in the protocols and deliverables and can clearly communicate the requirements to our laboratory staff.

Some of the regulatory programs where IEA is typically involved, are discussed below:

A. Hazardous Waste Assessments

1. CERCLA (Superfund)

Superfund site investigations, whether sponsored by Federal, State, or Principally Responsible Parties (PRP's) utilize the analytical guidelines established by the USEPA Contract Laboratory Program (CLP). Typical analytical requirements performed at Superfund sites include:

- . Full Target Compound List (TCL's formerly known as the Hazardous Substance List-HSL's) air, soil and water.
- . Tentatively Identified Compounds (TIC's) performed by GC/MS library search with analyst review.
- . Site specific compounds, often times requiring some level of method development.
- . Air analysis for volatile constituents potentially migrating from the site.

IEA has extensive experience providing these services to the New York State Department of Environmental Conservation, Connecticut Department of Transportation, New York City Department of Sanitation, as well as numerous confidential industrial clients.

2. RCRA

The Resource Conservation and Recovery Act has a number of requirements for analytical testing. These requirements generally fall into two categories: analysis of waste to determine if the material is hazardous; and analysis of environmental samples (soils, water, air) to determine if hazardous constituents are present.

Analysis of waste material usually involves the Toxicity Characteristic Leaching Procedure Tests including:

- . Volatiles
- . Semivolatiles
- . Pesticides
- . Herbicides
- . Metals

· It is also common practice to test this material for:

- . Priority Pollutants
- . Appendix IX constituents
- . Other characteristics required by waste disposal companies

The 1986 RCRA Amendments also list a plan, phasing out certain hazardous materials from land disposal. The RCRA land ban program includes a number of analytical requirements. Some of these are:

- . TCLP Solvent list (F001-F005 wastes)
- . California List
- . Appendix III

Analysis of environmental samples at RCRA facilities generally involves:

- . Groundwater monitoring for RCRA indicators, Appendix IX, and specific constituent monitoring
- . Soil analysis for Appendix IX or priority pollutants
- . Air analysis for off-site migration of volatile organics

The guidelines for analytical testing for RCRA programs is provided in the SW846 Manual (Third Edition). IEA has reviewed the requirements of SW846 and has developed the analytical capabilities to comply with this document. We have carefully reviewed and are able to deliver all of the QA/QC described in SW846. Our understanding of the application and limitations of these methods helps us provide guidance to our clients. We are also able to clearly define, review, and manage our laboratory with the knowledge that we are providing quality data that meets the regulatory requirements.

B. Effluent Monitoring

1. NPDES

The Clean Water Act has certain requirements for effluent monitoring. This includes both the NPDES permit process for direct discharges and the Pretreatment program for effluents to POTW's. The NPDES permit generally involves priority pollutant analysis upon permit renewal (5 year) or on an annual basis, plus monthly reporting of "conventional pollutants". The Pretreatment program is based on industry specific categorical standards and the requirements of the POTW.

The analytical guidelines for these programs are described in 40 CFR Part 136, specifically, Methods 624/625 and 608 for organics and the 200 series for inorganics. The QA/QC requirements are described in these methods.

IEA has been providing effluent monitoring services since 1977. We provide sampling and analytical program design, sample collection and pick-up, and analysis using the appropriate methodologies and QA/QC.

C. Potable Water Analysis

1. Safe Drinking Water Act (SDWA)

The SDWA requires water purveyors test their supply at least once annually. Those supplying larger populations must test more frequently. There are a number of parameter groups that are tested under SDWA including the primary and secondary drinking water standards. These include:

PRIMARY	SECONDARY
INORGANICS	INORGANICS
Arsenic, Barium, Cadmium, Silver, Sodium, Chromium Lead, Mercury, Selenium, Fluoride, Nitrate	Chloride, Color, Copper, Iron, Langelier Index, Manganese, Odor, TDS, Sulfate, Surfactants, Zinc
ORGANICS	ORGANICS
Vinyl Chloride, Endrin, Lindane Methoxychlor, Toxaphene 2,4-D, 2,3,5-TP (Silvex)	Vinyl Chloride

PRIMARY	SECONDARY
BIOLOGICAL	
Total Coliform	
TRIHALOMETHANES and VOLATILE ORGANICS	
RADIOACTIVITY	
Gross alpha, beta, Radium 226, 228	

The EPA promulgated amendments to the SDWA in 1986. The amendments include a "phase in" program for the analysis of volatile synthetic organic chemicals (SOC's), inorganic chemicals (IOC's), microbiological and radionuclide contaminants. The analytical protocols for the organic constituents are defined in the EPA 500 series methods.

IEA has provided drinking water analysis to many clients. We have also been contracted by the USEPA to perform a Special Analytical Services (SAS) project for the analysis of drinking water detection limit volatile organics. This required the use of capillary column purge and trap GC/MS using Method 524.2.

D. Real Estate Transfers

Many state environmental agencies have promulgated regulations requiring a site assessment prior to the sale of an industrial facility. The extent and complexity of the assessment varies due to different state regulatory requirements as well as site specific conditions. These transfers of property are governed by specific state environmental guidelines such as ECRA (NJ), Superlien (CT), and proposed regulations in Illinois.

IEA has provided laboratory services for numerous property transfers, particularly in New Jersey, under ECRA regulations, and CT under the Superlien Laws. The data generated from these projects have to withstand close scrutiny by state regulators. The data must also be of "courtroom quality" should there be any dispute over the findings.

We have the capability to test for specific chemicals and chemical groups as well as broad base chemical lists such as Priority Pollutants, RCRA characteristics, Target Compound List and Appendix IX. We can provide rush turnaround time service for "hot" transactions as quickly as 24 hours for volatile organics. The most important aspect of this type of testing, however, is the dat quality and its usability for the site assessment.

E. Air Testing Capabilities

IEA-CT has been providing air testing services since the 1970's. We specialize in the analysis of volatile organic compounds for ambient air monitoring programs. We are also experienced in the analysis of other organic compounds, as well as inorganics.

Our experience includes several different sampling media such as Tenax tubes, Tedlar bags, SUM. Canisters, impinger solutions, and a variety of adsorptive traps. EPA has not published air testing methodologies of the same detail as water and soil methods. Existing air methods, for example T0¹ and T02 volatile organics, do not have the rigorous procedures and QA/QC detail. This require that the laboratory have sufficient working knowledge and experience to provide quality, defensible data.

IEA has performed air testing for several clients, including the USEPA. We were contracted under the EPA's Special Analytical Services (SAS) program to provide trace analysis of volatile organic from SUMA canisters. We have conducted stack gas monitoring at a midwest steel mil formaldehyde analysis on impinger solutions from a chemical manufacturing facility, soil vent gas testing from landfills in New Jersey, as well as numerous other projects.

As part of our air testing services, we provide consultation on sampling program design, including the selection of the most appropriate sampling media.

We have investigated the adsorptive properties of many different traps and are able to pass this information, as well as our experience, to our customers. We are very proficient in the analysis c Tenax/adsorptive media traps using cryofocus GC/MS techniques. Using this knowledge, and the customers knowledge of site conditions, we can develop the most effective sampling plan.

We believe that our understanding and experience, combined with our familiarity with EPA-CLP QA/QC requirements, makes us uniquely qualified to provide air testing services.

A summary of the IEA-CT analytical capabilities is presented in TABLE II-1

TABLE II - 1

IEA-CT ANALYTICAL CAPABILITIES

I. ORGANICS-GC/MS

Volatile Organics - 524 Volatile Organics - 8240 Volatile Organics - CLP Volatile Organics - TO1/TO2 Volatile Organics - Appendix IX Acid & Base/Neutrals - 8270 Acid & Base Neutrals - CLP

Acid & Base/Neutrals - Appendix IX

III. INORGANIC METALS

ICP Metals Furnace Metals CLP Metals

V. INORGANIC WET CHEMISTRY

Acidity Alkalinity Ammonia Bicarbonate

Biochemical Oxygen Demand (BOD)

Bromide Chloride Chlorine Demand Chlorine Residual Chemical Oxygen Demand

Color Conducitivity Chromium (VI) Cyanide - Amenable Cyanide - Total Cyanide (CLP) Dissolved Oxygen Flashpoint

Fluoride Grain Size

Hydrocarbon analysis

MBAS Nitrate Nitrite Odor

Oil and Grease Paint Filter Test

pН Phenois

II. ORGANICS-GC

Organohalide Pesticides & PCBs - 608 Organohalide Pesticides & PCBs - 8080 Organohalide Pesticides & PCBs - CLP Organophosphate Pesticides - 8140 Organohalide Pesticides & PCBs - Appendix IX Chlorinated Herbicides - 8150

Chlorinated Herbicides - Appendix IX

IV. BIOLOGICAL ANALYSES

Total Coliform Enterococci Fecal Coliform Fecal Streptococcus Standard Plate Count

Phosphate Phosphorus Settleable Solids Silica

Specific Gravity Sulfate Sulfide Sulfite

Sludge Volume Index Tannins and Lignins Total Dissolved Solids Total Kjeldahl Nitrogen Total Organic Carbon Total Organic Halides **Total Solids**

Total Suspended Solids

Turbidity Volatile Solids

Corrosivity Characteristics Ignitability Characteristics

EPTOX TCLP

III. REPRESENTATIVE ENVIRONMENTAL PROJECTS

<u>United States Environmental Protection Agency</u> - U.S. EPA Contract Laboratory Program contract for organic and inorganic analysis of 60 environmental samples per month for 30 months. Analysis by rigid protocols and QA/QC policies for Target Compound List (TCL) organics by GC/MS and GC/ECD procedures.

<u>United States Environmental Protection Agency</u> - many SAS contracts, including PCB analysis for a major enforcement action and low detection level drinking water analyses by GC/MS.

<u>U.S. Navy, Philadelphia, PA</u> - Analysis of bay samples, river water, groundwater, and surface water for priority pollutants, xylenes, and petroleum hydrocarbons.

Northville Industries - Melville, NY - Analysis of air samples associated with cleanup and vapor monitoring of contaminated groundwater. Adsorptive tubes are analyzed for volatile organics utilizing very strict QA/QC requirements. Approximately 1000 samples are tested annually.

<u>URS Consultants - Buffalo, NY - NYC Landfills, Fountain Avenue and Pennsylvania Avenue Analysis of approximately 700 samples, over a 9 month period, consisting of water, soil, sediment, waste, and air. Parameters included Full TCL/TAL and miscellaneous wet chemistry tests, all in accordance with '91NYSDEC ASP/CLP with Superfund deliverables and customized diskette deliverable.</u>

Wehran Engineering - Middletown, NY - NY State manufacturing facility. Analysis of approximately 600 samples consisting of water and soil for Full TCL/TAL analysis by '91NYSDEC ASP CLP Protocol and Category B deliverables with customized diskette deliverables.

McLaren-Hart - Pittsburgh, PA - American Cyanamid, Willow Island, W.VA. Analysis of approximately 75 water and soil samples for Full TCL/TAL by CLP 3/90 protocols and deliverables.

Fanning, Phillips and Molnar\NY Urban Development Corporation/Donald Trump - Analysis of over 100 soils, sediments, and groundwater samples for volatiles, base/neutrals, petroleum hydrocarbons, and metals.

TABLE II - 1

IEA-CT ANALYTICAL CAPABILITIES

I. ORGANICS-GC/MS

Volatile Organics - 524 Volatile Organics - 8240 Volatile Organics - CLP Volatile Organics - TO1/TO2 Volatile Organics - Appendix IX Acid & Base/Neutrals - 8270 Acid & Base Neutrals - CLP Acid & Base/Neutrals - Appendix IX

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III. INORGANIC METALS

ICP Metals Furnace Metals CLP Metals

IV. BIOLOGICAL ANALYSES

Total Coliform Enterococci Fecal Coliform Fecal Streptococcus Standard Plate Count

V. INORGANIC WET CHEMISTRY

Acidity Alkalinity Ammonia Bicarbonate

Biochemical Oxygen Demand (BOD)

Bromide Chloride Chlorine Demand

Chlorine Residual Chemical Oxygen Demand

Color Conducitivity Chromium (VI) Cyanide - Amenable Cyanide - Total Cyanide (CLP) Dissolved Oxygen Flashpoint Fluoride Grain Size

Hydrocarbon analysis

MBAS Nitrate Nitrite Odor Oil and Grease

Paint Filter Test

pН Phenols

Phosphate Phosphorus Settleable Solids Silica

Specific Gravity Sulfate Sulfide Sulfite

Sludge Volume Index Tannins and Lignins Total Dissolved Solids Total Kjeldahl Nitrogen Total Organic Carbon Total Organic Halides

Total Solids Total Suspended Solids

Turbidity

Volatile Solids

Corrosivity Characteristics Ignitability Characteristics

EPTOX TCLP

III. REPRESENTATIVE ENVIRONMENTAL PROJECTS

<u>United States Environmental Protection Agency</u> - U.S. EPA Contract Laboratory Program contract for organic and inorganic analysis of 60 environmental samples per month for 30 months. Analysis by rigid protocols and QA/QC policies for Target Compound List (TCL) organics by GC/MS and GC/ECD procedures.

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Fanning, Phillips and Molnar\NY Urban Development Corporation/Donald Trump - Analysis of over 100 soils, sediments, and groundwater samples for volatiles, base/neutrals, petroleum hydrocarbons, and metals.

<u>CH₂M Hill - Alachua, FL</u> - Analysis of 60 samples for full Appendix IX. Results were provided both electronically (diskette) and in full CLP-like hardcopy.

<u>Fortune 500 Company - New England Locations</u> - Analysis of over 1,000 samples over a two year period for PCB's in soils, sediments, groundwaters, surface waters, concrete cores, and transformer oils.

<u>Various Connecticut</u>, <u>New York</u>, <u>Massachusetts and New Jersey Consulting Engineering Firms</u> - Groundwater and effluent analysis for priority pollutants and conventional indicator parameters.

<u>Numerous Electroplaters in the State of Connecticut</u> - NPDES and SPDES analyses for compliance with Connecticut DEP guidelines. Analyses include heavy metals and volatile priority pollutants.

IV. DATA DELIVERABLES

Just like there are many different analytical protocols for a specific parameter list, there are also different report formats or deliverables. These deliverables may be based on State or Federal requirements or client preference. The USEPA Contract Laboratory Program (CLP) also has a defined set of requirements for reporting.

IEA offers several Report Levels. These include Data Summaries, CLP, NJ Regulatory and Reduced Format, and client specified report packages.

Custom report packages are also available. We recognize that clients have different needs depending on data quality objectives, data validation criteria, and individual preferences. Our data processing system affords us the capabilities to produce this variety of formats.

IEA-CT is especially adept in the production of Full CLP reports. We have over 8 years experience in the USEPA CLP program and consider this level of reporting part of our routine services.

V. CUSTOMER SERVICE/PROJECT MANAGEMENT

One of IEA's strengths is our dedication to customer service. Our Client Service Staff is experienced in laboratory testing and understand technical and logistical issues. These three individuals serve as the point contact for sample coordination with the project team. One Client Service Representative is assigned to each customer, so there is consistency in project management.

The Client Service Representatives coordinate bottle preparation and delivery, order entry, monitor project status, and notify the client of any analytical difficulties. Upon receipt of bottles, they will notify the client as to the condition of bottles, chain-of-custody questions, or any other sample receipt issues.

Technical questions about data packages are also directed to our Client Service staff. They can address most data package issues immediately. If the question requires further assistance, a Corrective Action Report (CAR) is generated documenting the question and the date/time a response is due. This expedites clients inquiries with one phone call.

VI. DATA MANAGEMENT

IEA-CT currently uses a Perkin Elmer Laboratory Information Management System (LIMS). This LIMS is primarily used for order processing and sample tracking. Samples are pre-logged at the time a client calls with an order. All information detailing the project requirements are entered at this time. Upon receipt of actual samples, the original PreLog-In information is checked against the actual samples that arrived. If necessary, edits are made and the project is upgraded to Logged-In status. This system has greatly enhanced the timeliness of Log-In. Group Leaders use LIMS to track sample status, due dates, and holding times.

In addition to PE-LIMS, IEA-CT utilizes an ORACLE data base system, called Seedpack II/III for analytical data processing, report generation, and diskette deliverables. This system utilizes SP II/III development tools to customize the laboratories information management needs. Data, including results and all associated QC, moves from the instruments to the centralized data base, where it is available for final report. Report programs have been written to satisfy customer needs. These include USEPA-CLP, NYSDEC-ASP, Level I Summary Reports, and other customer specific formats. A menu driven diskette program was developed to accommodate a wide variety of electronic deliverable formats. These diskettes, typically used as data summaries, can be delivered by overnight mail. They are transmitted in ASCII or LOTUS format, generally useful for a wide range of applications.

VII. QA/QC SUMMARY

The staff at IEA has an understanding of the requirements of each analytical protocol. Our QA/QC program is based on the EPA methodologies and their specific requirements. These methods are numerous and are constantly changing. We have defined the QC programs and continue to monitor them. This is an assurance to our clients of compliance to regulatory requirements.

The responsibility of the Inorganics and Organics Group Leaders at IEA is to produce analytical data that meets the appropriate and specific regulatory requirement in terms of completeness, precision, accuracy, representativeness, documentation, and comparability of data.

The QA/QC program at IEA is administered by the QA/QC officer. It is the responsibility of the QA/QC officer to monitor and guarantee, to the extent possible, the quality of all measurements to the Director of each IEA Laboratory. The QA/QC officer accomplishes the above objectives by implementing a formal QA/QC program and periodically monitoring laboratory produced analytical data, Corrective Action Reports, and the results of analysis of QC and QA samples.

The QA/QC officer implements the correction of any problems by notifying laboratory management using a QA/QC Deficiency Report. This report specifies deficiencies in analytical procedures, quality control procedures, bench data, and/or standard operating procedures. An implementation due date is also specified.

Our QC program is an integrated approach consisting of:

Good Laboratory Practice -

High standards of excellence by the IEA Chemists and Technicians, in a well maintained analytical laboratory performing duplicate analyses, blanks, standard additions, internal standards, and using the highest quality reagents. Legally defensible chain-of-custody procedures for analysis as well as sampling are used. Standard Operating Procedures are used and are available at the bench for quick reference by the analyst.

An Internal Quality Control Program -

Consisting of sample chain-of-command, laboratory notebook review, quality control sample analysis, technique review, and continued education programs.

External Quality Control Audits -

Done on a quarterly basis by the USEPA-CLP States of Connecticut, New York and New Jersey, which encompass organics, metals, nutrients, and demand series. Also, the QA/QC officer regularly introduces "blind" samples from outside consultants and other laboratory "round-robin" series.

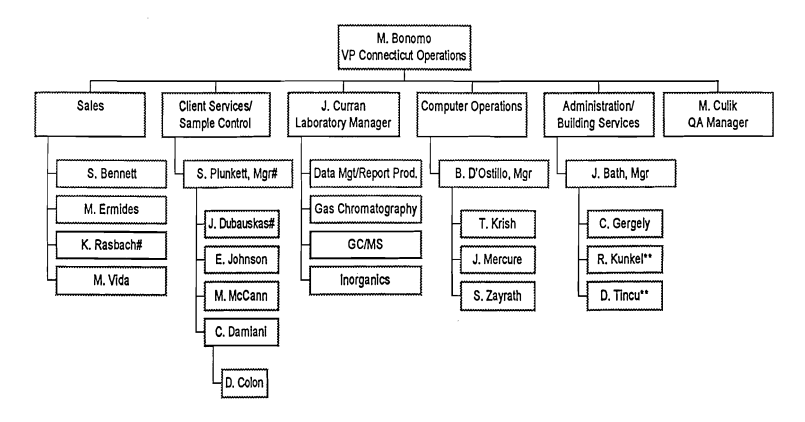
VIII. IEA LABORATORIES ORGANIZATIONAL STRUCTURE

IEA maintains full-time staff of over 320 professionals with approximately 75% experienced in the areas of organic and inorganic sample preparation and analysis. We also have professional profiles for all laboratory managers and group leaders. This information will be made available upon request.

The IEA-CT Organization structure is presented in Table VIII-1.



Doc #QAC00104.CT Date: 01/23/96



*Part Time/Temp

**25% charged to AMS

#30 hour work week



Doc #QAC00104.CT Date: 01/23/96

M. Bonomo VP Connecticut Operations J. Curran Laboratory Manager Data Management/ K. Maturo, Grp Ldr L. Decker, Mgr D. Helfrich Report Production Gas Chromatography **GC/MS** Inorganics Manager Classical Chemistry Semi-Volatiles Volatiles Atomic Spectroscopy C. Cascella, S. Ldr H. Downey D. Humbert M. Sciongay J. Bennett, Grp Ldr M. Ansari D. Nemeth, S. Ldr D. Abate, S. Ldr M. Tommasi, S. Ldr # B. Kostrzewska C. Charter M. Crowe S. Cappella C. Lombardi A. DeJulio A, Krajci M. Crudo S. Mickens# T. Cronin* M. Cicciarini B. LaForest P. Norman R. Doris R. Guerrera* P. Sequin T. Matejek M. Tcherniaeua M. Madison H. Linardos* R. Sibley L. McMannus K. Zmijewski D. Madumadu R. Nawere* K. Hass D. Vallette A. O'Leary

A. Ronge

J. Widomski

*Part Time/Temp

**25% charged to AMS

#30 hour work week

IX. IEA NETWORK SERVICES

IEA, Monroe, CT is part of the IEA network consisting of 7 labs throughout the Eastern United States:

Monroe, CT
Cary, NC (Headquarters) (2 labs including Radiological)
N. Billerica, MA
Whippany, NJ
Schaumburg, IL
Sunrise, FL

The entire network is used to help facilitate client requests. The Account Executive and Client Service staff will make arrangements with other IEA labs for services. Samples are never sent to another IEA lab, without first consulting with the client. Utilizing the network in this manner gives us the resources of a staff of 320 people and one of the largest analytical capabilities in the country. Any work sent to another IEA lab will be managed, tracked, and reported by IEA-CT, so that the customer has the benefit of dealing with one contact.

The following is a list of the additional services offered by IEA:	
COURIER - SAMPLE PICK UP and DELIVERY	
SAMPLE CONTAINER PREPARATION and TRANSPORT	,
AIR ANALYSIS	
ASBESTOS (PLM, TEM)	
ELECTRON MICROSCOPY	
RADIOLOGICAL ANALYSIS	
MIXED WASTE TESTING	

Data Validation

IEA has performed data validation for several consulting and engineering firms that did not have an internal QA/QC department. We have performed validation services on New Jersey ECRA projects (Tier II reports) and Superfund remediation sites (CLP-Tier I reports).

Courier Service

IEA offers sample pick-up for local clients. This service was designed for customer convenience and has a small fee dependent upon the location of sample pick-up.

Sample Containers and Transport-

IEA provides new, pre-labelled, pre-preserved bottles for sample collection. Chain-of-custody documentation is initiated at the laboratory and included with the bottles. We can list up to 11 samples on a single form. This means less paperwork in the field and fewer mistakes.

There are two types of bottles that can be provided: Standard and I-Chem glassware (as used by the USEPA for CLP projects). Preservatives specified by the analytical methods are added to bottles before delivery to the client. The bottle labels identify both the parameter and preservative.

Bottles are delivered to the site in sample coolers chilled with blue ice, using Federal Express or the IEA courier service. This generally requires 1-2 weeks advance notice. Different size coolers are used depending on the number of containers. Each bottle is wrapped with protective "bubble wrap" prior to shipment. All coolers are sealed with York Labs custody tape. This is the same procedure used by the USEPA for Superfund site (CLP) projects. Coolers are returned to the laboratory by overnight courier, such as Federal Express, or the IEA courier service.

TABLE X - 1

CERTIFICATIONS

In some instances it may be necessary for environmental data to be reported to a regulatory authority with reference to a certified laboratory. For your convenience, the laboratory identification numbers for the IEA-Connecticut laboratory are provided in the following table. Many states certify laboratories for specific parameters or tests within a category (i.e. method 325.2 for wastewater). The information in the following table indicates the lab is certified in a general category of testing such as drinking water or wastewater analysis. The laboratory should be contacted directly, if parameter-specific certification information is required.

IEA-Connecticut Certification Summary (as of June 1993)				
STATE	RESPONSIBLE AGENCY	CERTIFICATION	LAB NUMBER	
Connecticut	Department of Health Services	Drinking Water, Wastewater	PH-0497	
Kansas	Department of Health and Environmental Services	Drinking Water, Wastewater/Solid, Hazardous Waste	E-210/E-1185	
Massachusetts	Department of Environmental Protection	Potable/Non-Potable Water	CT023	
New Hampshire	Department of Environmental Services	Drinking Water, Wastewater	252891	
New Jersey	Department of Environmental Protection	Drinking Water, Wastewater	46410	
New York	Department of Health	CLP, Drinking Water, Wastewater, Solid/Hazardous Waste	10602	
North Carolina	Division of Environmental Management	Wastewater	388	
Rhode Island	Department of Health	ChemistryNon-Potable Water and Wastewater	A43	
California	Department of Health Services	Hazardous Waste	1778	
US Army Corps of Engineers	Missouri River Division	Organics, Inorganics		

XI. INVENTORY OF LABORATORY EQUIPMENT

Major Analytical Instrument Inventory - Monroe, CT

CLASSICAL CHEMISTRY

		Model	Serial
Equipment Name	<u>Manufacturer</u>	<u>Number</u>	<u>Number</u>
S	D 11 D1	1010	10.4.400
Spectrophotometer, IR	Perkin-Elmer	1310	134423
Spectrophotometer, UV-VIS	Perkin-Elmer	35	34630
Turbidimeter	Hach Company	2100A	851017142
TOC Analyzer	Xertex-Dohrmann	DC-80	HF2029
TOX Analyzer	Xertex-Dohrmann	MC3 A,B	MF2106
Fluorometer	Sequoia-Turner	112-003	D01491
pH/ISE Meter	Orion	SA 720	SR45A
pH/ISE Meter	Beckman	12	0232578
Conductivity Meter	Cole-Parmer Instrument	1484-20	1421
Flash Point Apparatus	Precision Scientific	Pensky-Martin	10 Au-12
Oven	Fisher Scientific	55G	291
Oven	VWR	1320	0701090
Incubator	Blue M Electric	100A	IN1-1362
Bio Refrigerator	Frost Queen	R20/L	00029
Centrifuge	DYNAC	0101	16846
Water Bath	Blue M Electric	MW-1220	MX-2520
D.O. Meter	YSI	51A	0241
Autoclave	Market Forge	STM-E	034200
COD Reactor	HACH	45600	920300006892
Muffle Furnace	Thermolyne		
TKN Block Digestor	Scientific Instruments	AD-4020	8915-049
Digital Hot Plate/Stirrer	PMC	730	0298E
Digital Hot Plate/Stirrer	PMC	730	0299E
Semiautomated Analyzer	LACHAT	Quikchem	125-360
BOD Incubator	Precision Scientific	FLC02662	FU199JRW2
BOD Incubator	Precision Scientific	FLC02662	FU178RRWR
Mini Distillation Setup	Andrews Glass Co	110-10-R	A4WO309
Mini Distillation Setup	Andrews Glass Co	110-10-R	A4WO209

BARBARA A. DEBUONO, M.D., M.P.H. Commissioner



Expires 12:01 AM April 1, 1996 ISSUED April 1, 1995 REVISED February 27, 1996

INTERIM CERTIFICATE OF APPROVAL FOR LABORATORY SERVICE

Issued in accordance with and pursuant to section 502 Public Health Law of New York State

Lab ID No.: 10602

Director: MR. JEFFREY CURRAN

Lab Name: IEA INC

Address: 200 MONROE TURNPIKE MONROE CT 06468

is hereby APPROVED as an Environmental Laboratory for the category

ENVIRONMENTAL ANALYSES NON POTABLE WATER

All approved subcategories and/or analytes are listed below:

Wastewater Metals III:

r. Hydrocarbon Pesticides : -DDD 4'-DD8 4'-DDT alpha-BHC 41drin ta-BHC lordane fotal delta-BHC Dieldrin drin aldehyde drin ■udosulfan I Endosulfan II padosulfan sulfate ptachlor ptachlor epoxide Tindane **Methoxychlor** vaphene

Wastewater Miscellaneous : Bro∎ide Boron, Total Cyanide, Total Color Phenols Oil & Grease Total Recoverable Hydrogen Ion (pH) Specific Conductance Sulfide (as S) Surfactant (MBAS) Organic Carbon, Total Wastewater Metals II (ALL) Organophosphate Pesticides (ALL) Phthalate Esters (ALL) Purgeable Halocarbons (ALL)

Cobalt, Fotal
Molybdenum, Total
Fin, Total
Fin, Total
Finanium, Total
Fhallium, Total
Nutrient:
Kjeldahl Nitrogen, Fotal
Nitrate (as N)
Orthophosphate (as P)
Phosphorus, Total
Haloethers (ALL)
Nitroaromatics and Isophorone (ALL)
Polynuclear Aromatics (ALL)
Priority Pollutant Phenols (ALL)
Residue (ALL)

Mineral: Alkalinity Chloride Pluoride, fotal Sulfate (as \$04) Hardness, fotal Acrolein and Acrylonitrile (ALL) Benzidines (ALL) Chlorophenoxy Acid Pesticides (ALL) Chlorinated Hydrocarbons (ALL) Demand (ALL) Wastewater Metals I (ALL) Nitrosoamines (ALL) Polychlorinated Biphenyls (ALL) Purgeable Aromatics (ALL) TCLP Additional Compounds (ALL)

Serial No.: 032214

Wadsworth Center

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DOH-3317 (3/95)

THERE AT CHICOTHY HIDT, MAY IN , MAY THE COMMISSIONER.

BARBARA A. DEBUONO, M. M.P.H.

Commissioner

Expires 12:01 AM April 1, 1996 ISSUED April 1, 1995 REVISED June 29, 1995

INTERIM CERTIFICATE OF APPROVAL FOR LABORATORY SERVICE

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Lab ID No.: 10602

Director: MR. JEFFREY CURRAN

Lab Name: IEA INC

Address : 200 MONROE TURNPIKE MONROE CT 06468

is hereby APPROVED as an Environmental Laboratory for the category

ENVIRONMENTAL ANALYSES/ POTABLE WATER

All approved subcategories and/or analytes are listed below:

king Water Non-Metals : Alkalinity
Calcium Hardness
Chloride
Colot
Corrosivity
Fluoride, Fotal
Mitrate (as N)
Hydrogen Ion (pH)
Solids, Fotal Dissolved
Sulfate (as S04)

Drinking Water Tribalomethane (ALL) Drinking Water Metals I (ALL) Volatile Halocarbons (ALL)

Volatile Aromatics (ALL)

Serial No.: 027221

Wadsworth Center for Laboratories and Research

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BARBARA A. DEBUONO, M.D., M.P.H. Commissioner



Expires 12:01 AM April 1, 1996 ISSUED April 1, 1995 REVISED February 27, 1996

INTERIM CERTIFICATE OF APPROVAL FOR LABORATORY SERVICE

Issued in accordance with and pursuant to section 502 Public Health Law of New York State

Lab ID No.: 10602

Director: MR. JEFFREY CURRAN

Lab Name: IEA INC

Address: 200 MONROE TURNPIKE
MONROE CT 06468

is hereby APPROVED as an Environmental Laboratory for the category

ENVIRONMENTAL ANALYSES NON POTABLE WATER

All approved subcategories and/or analytes are listed below:

r. Hydrocarbon Pesticides : 4'-DDD 4'-DDB 4'-DDT alpha-BHC 4)drin ta-BHC lordane fotal delta-BHC Dieldrin drin aldehyde drin andosulfan I Endosulfan II p-dosulfan sulfate ptachlor ptachlor epoxide Tindane **Methoxychlor** raphene

Wastewater Miscellaneous:
Bromide
Boron, Total
Cyanide, Total
Color
Phenols
Oil & Grease Total Recoverable
Hydrogen Ion (pH)
Specific Conductance
Sulfide (as S)
Surfactant (MBAS)
Organic Carbon, Total
Wastewater Metals II (ALL)
Organophosphate Pesticides (ALL)
Phtbalate Esters (ALL)
Purgeable Halocarbons (ALL)

Wastewater Metals III:
Cobalt, Total
Holybdenum, Total
Tin, Total
Titanium, Total
Thallium, Total
Nutrient:
Kjeldahl Nitrogen, Total
Nitrate (as N)
Orthophosphate (as P)
Phosphorus, Total
Haloethers (ALL)
Nitroaromatics and Isophorone (ALL)
Polynuclear Aromatics (ALL)
Priority Pollutant Phenols (ALL)
Residue (ALL)

Mineral: Alkalinity Chloride Pluoride, Total Sulfate (as SO4) Hardness, fotal Acrolein and Acrylonitrile (ALL) Benzidines (ALL) Chlorophenoxy Acid Pesticides (ALL) Chlorinated Hydrocarbons (ALL) Demand (ALL) Wastewater Metals I (ALL) Nitrosoamines (ALL) Polychlorinated Biphenyls (ALL) Purgeable Aromatics (ALL) TCLP Additional Compounds (ALL)

Serial No.: 032214

Wadsworth Center

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Must be conspicuously posted. Valid certificate has a red serial number.

DOH-3317 (3/95)

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Commissioner

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Lab ID No.: 10602

Director: MR. JEFFREY CURRAN

Lab Name: IEA INC

Address : 200 MONROE TURNPIKE
MONROE CT 06468

is hereby APPROVED as an Environmental Laboratory for the category

ENVIRONMENTAL ANALYSES / POTABLE WATER

All approved subcategories and/or analytes are listed below:

king Water Non-Hetals : Alkalinity
Calcium Hardness
Chloride
Color
Corrosivity
Fluoride, Total
Mitrate (as N)
Hydrogen Ion (pH)
Solids, Total Dissolved
Sulfate (as S04)

Drinking Water Tribalomethane (ALL) Drinking Water Metals I (ALL) Volatile Halocarbons (ALL) Volatile Aromatics (ALL)

Serial No.: 027221

Wadsworth Center for Laboratories and Research

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BARBARA A. DEBUONO, M. M. M.P.H. \ Commissioner

Expires 12:01 AM April ISSUED April 1, 1995 REVISED June 29, 1995

INTERIM CERTIFICATE OF APPROVAL FOR LABORATORY SERVICE

Issued in accordance with and pursuant to section 502 Public Health Law of New York State

Lab ID No.: 10602

Director: MR. JEFFREY CURRAN

Lab Name: IEA INC

Address : 200 MONROE TURNPIKE MONROE CT 06468

is hereby APPROVED as an Environmental Laboratory for the category

ENVIRONMENTAL ANALYSES/AIR AND EMISSIONS

All approved subcategories and/or analytes are listed below:

ble Aromatics (ALL)

Purgeable Halocarbons (ALL)

-Serial No.: 027222

Wadsworth Center for Laboratories and Research

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MADY D. CHACCIN M.D. M.D. M.D.H. COMMICCIONED

BARBARA A. DEBUONO, M. M. P.H.

Commissioner

Expires 12:01 AM April 1, ISSUED April 1, 1995 REVISED June 29, 1995

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Address : 200 MONROE TURNPIKE

MONROE CT 06468

is hereby APPROVED as an Environmental Laboratory for the category

ENVIRONMENTAL ANALYSES/SOLID AND HAZARDOUS WASTE

All approved subcategories and/or analytes are listed below:

acteristic Testing :
Corrosivity
Ignitability
Reactivity
FCLP
E.P. Toxicity
eable Halocarbons (ALL)

Miscellaneous :
 Cyanide, Total
 Hydrogen Ion (pH)
 Sulfide (as S)
Organophosphate Pesticides (ALL)
Phthalate Esters (ALL)

Acrolein and Acrylonitrile (ALL)
Chlor. Hydrocarbon Pesticides (ALL)
Haloethers (ALL)
Metals II (ALL)
Polynuclear Arom. Hydrocarbon (ALL)
Priority Pollutant Phenols (ALL)

Chlorophenoxy Acid Pesticides (ALL)
Chlorinated Hydrocarbons (ALL)
Hetals I (ALL)
Nitroaromatics Isophorone (ALL)
Polychlorinated Biphenyls (ALL)
Purgeable Aromatics (ALL)

Serial No.: 027223

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MIDIC D. CHICCIN M.D. M.D.D. M.D.H. COMMICHONDO

BARBARA A. DEBUONO, M.P.

Expires 12:01 AM April 1, 1996 ISSUED April 1, 1995 REVISED June 29, 1995

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MONROE CT 06468

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CONTRACT LABORATORY PROTOCOL (CLP)

All approved subcategories and/or analytes are listed below:

'I.P Inorganics

CLP PCB/Pesticides

CLP Semi-Volatile Organics

CLP Volatile Organics

Commissioner

Serial No.: 027224

Wadsworth Center for Laboratories and Research

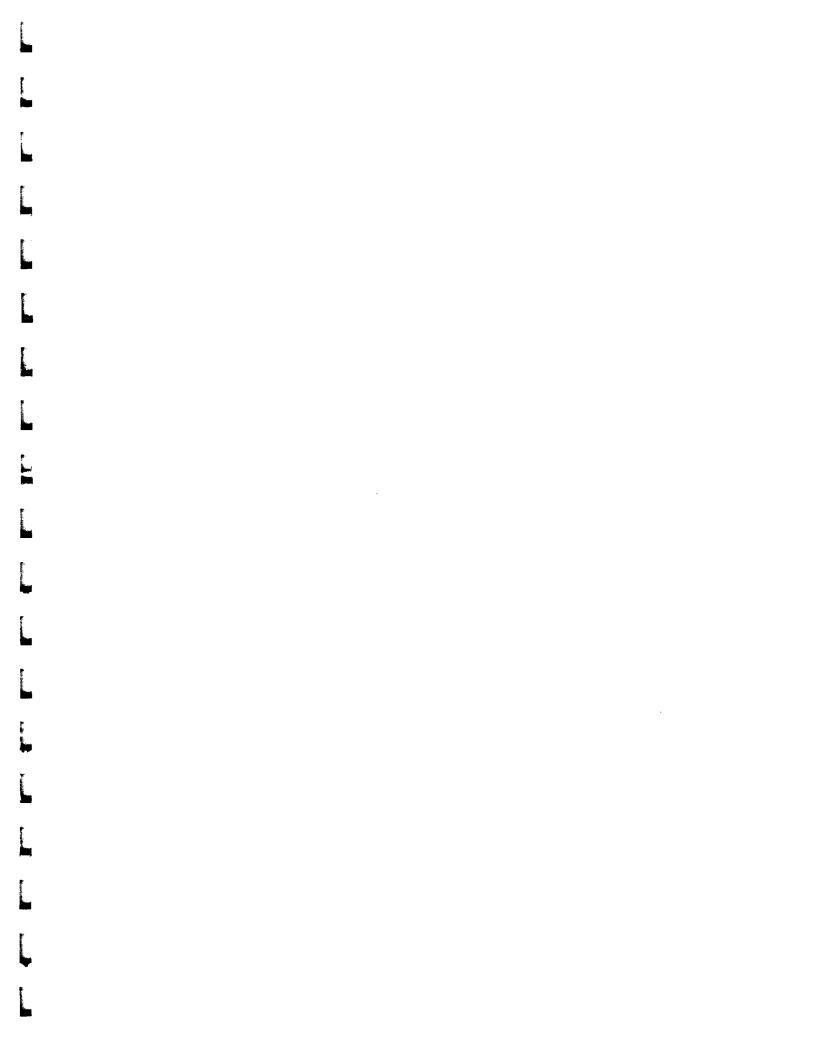
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Date: 03/24/95

GC/MS Extractable Organics					
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/Kg)	PQL (ug/Kg)
2,4-Dimethylphenol	64-238	<u>-</u>	<u>-</u>	39.5	330
Dimethyl phthalate	D-112	-	<u> </u>	9.88	330
4,6-Dinitro-2-methylphenol	D-362		<u> </u>	11.9	1600
2,4-Dinitrophenol	D-382		<u> </u>	105	1600
2,4-Dinitrotoluene	39-139	28-89	47	11.6	330
2,6-Dinitrotoluene	50-138	<u>.</u>		14.0	330
Di-n-octylphthalate	4-146	<u>. </u>		14.8	330
Fluoranthene	26-137		-	11.8	330
Fluorene	59-121	-	<u>-</u>	18.2	330
Hexachlorobenzene	D-132	<u> </u>		13.8	330
Hexachlorobutadiene	24-116			15.4	330
Hexachlorocyclopentadiene	D-59			12.8	330
Hexachloroethane	40-113			21.5	330
Indeno(1,2,3-cd)pyrene	D-171	<u> </u>	-	17.1	330
Isophorone	21-196	-	-	14.9	330
2-Methylnaphthalene	D-127	-	-	14.8	330
2-Methylphenol (o-cresol)	40-189	<u>-</u>	<u>-</u>	11.2	330
4-Methylphenol (p-cresol)	28-198		_	17.8	330
Naphthalene	21-133		-	13.0	330
2-Nitroaniline	D-127	•	_	49.2	1600
3-Nitroaniline	D-91		_	342	1600
4-Nitroaniline	D-108	•	•	138	1600
Nitrobenzene	35-180	-	_	13.5	330_
2-Nitrophenol	58-364	-	_	13.5	330
4-Nitrophenol	D-264	10-80	50	128	1600
N-Nitroso-di-n-propylamine	D-230	41-126	38	13.1	330
N-Nitrosodiphenylamine	D-114	<u> </u>		17,1	330
Pentachlorophenol	28-352	17-109	47	76.7	1600
Phenanthrene	54-120	<u> </u>	-	12.5	330
Phenol	10-224	26-90	35	10.6	330
Pyrene	52-113	35-142	36	18.2	330
1,2,4-Trichlorobenzene	44-142	38-107	23	15.2	330

Date: 03/24/95

GC/MS Extractable Organics						
	QC CHECK/LCS % RECOVERY LIMIT	MATRIX SPIKE % RECOVERY LIMIT	RELATIVE % DIFFERENCE (RPD) LIMIT	MDL (ug/Kg)	PQL (ug/Kg)	
2,4,5-Trichlorophenol	82-354		, <u>-</u>	56.1	1600	
2.4.6-Trichlorophenol	74-288		_	28.9	330	



<u>QAPjP</u> ATTACHMENT B

STANDARD OPERATING PROCEDURES

Standard Operating Procedure Completion of Field Notes

This protocol is designed to ensure that proper techniques are used during the collection and preparation of field notes. Field notes are collected in field notebooks, which are often the only source of "first hand" information regarding activities that were conducted at a site. Field notes may be called into a court of law; therefore, it is imperative that field notes be maintained in a thorough and proper manner.

All field notes should be completed in a water-proof notebook and should not be completed on loose sheets of paper that might get lost or misplaced. All field notes should be completed in permanent ink, rather than pencil and should be neat and orderly. Use of a pencil for collection of field notes is acceptable only in extremely poor weather conditions. All field notes taken during the field activities should be photocopied immediately after completion of the activities and placed in the project file to preserve a permanent record of the activities. In addition, when conducting field activities, the following information should also be collected:

- the date and time of the field activities (both the start and the finish time) including the time that certain "milestones" are achieved;
- weather conditions on the day of the field activities (in some cases it may also be appropriate to include the weather conditions for the previous day, such as when a heavy snow fall has occurred);
- the names and affiliations of all personnel involved in the field activities;
- the purpose of the field activities (e.g., groundwater sampling, site inspection, UST removal);

The field notes should accurately reflect a chronology of the activities that were conducted at the site. The following are examples of information that should be included in the field notes, but might not be applicable in all situations:

- the time that subcontractors, clients, police details, consultants or other persons arrived and left the site;
- a site sketch indicating the approximate location of groundwater observation wells to be sampled, borings to be installed, test pits to be performed, utilities to be located or suspected underground storage tanks, abutters (Note: Site sketches should be included even when a site plan has been provided. If a site sketch is not feasible, the site plan that is being used to locate structures should be referenced);
- reference to any other documents that are completed during the course of the site activities that may include additional information not included in the field book, including: Chain of Custody forms; Test Boring Reports; Test Pit Reports; Manifests and calibration log books;
- a site sketch indicating areas where snow cover, vehicles, debris or other obstructions may have limited a site inspection, prevented well sampling or

- otherwise prohibited the completion of activities;
- where and when field instruments (e.g., PID, OVA) are being used, all calibration and sampling/screening conditions should be logged;
- any unsafe conditions observed by GEC personnel and presented to on-site personnel or subcontractors;
- observations made during site inspections or field activities including, but not limited to: the locations of stained soils or stressed vegetation; noticeable odors; the presence of nonaqueous-phase liquid; and
- types of sampling containers used and preservatives used in sample containers.

It is the responsibility of each GEC employee to maintain his/her own field book. All field books are the property of GEC and in the event that the employee terminates employment with GEC the field books are to remain at GEC.

Standard Operating Procedure Decontamination Procedures for Field Equipment

All field equipment (bailers, well sounder, gloves, etc.) must be decontaminated before each use, between samples and before it is returned to the equipment room. Decontamination procedures vary for the type of analyses to be performed. The following basic procedures should always be used to decontaminate equipment regardless of the type of analysis:

- 1) Scrub equipment with soapy water (Liquinox, Alconox, trisodiumphosphate or equivalent).
- 2) Rinse with tap water, if available.
- 3) Rinse with deionized water from green spray bottle.

For Metals, perform the following additional procedures:

- 4) Rinse with 10% nitric acid (HNO₃).
- 5) Final rinse with deionized water.

For base/neutral/acid extractables, PCB's and pesticides perform the following, additional procedures:

- 4) Rinse with pesticide grade methanol or hexane and let dry.
- 5) Rinse with hexane (if applicable) and let dry.
- 6) Final rinse with deionized water.

For Volatile Organics and all other analyses, perform the following additional procedures:

- 4) Rinse with methanol.
- 5) Final rinse with deionized water

NOTE: When sampling for more than one of the above types of analyses, use the protocol for volatile organics last.

Solvent use should be gauged carefully so that a minimal amount of solvent is left after use. Allow any remaining solvent to evaporate.

Standard Operating Procedure Sample Preservation and Chain of Custody

This protocol is designed to ensure that proper techniques are employed in the preservation and chain-of custody of samples collected for laboratory analyses or for screening. This Protocol is intended to be consistent with Massachusetts Publication #WSC-310-91 (Standard References for Monitoring Wells), and 40 CFR 136 (Guidelines Establishing Test Procedures for the Analysis of Pollutants).

The results of screening and/or laboratory analysis of solid, liquid or gaseous media constitute the basis of evaluation of the majority of the disposal sites under investigation. It is therefore imperative that the preservation of the samples be appropriate to the media being analyzed as well as the analysis which is being performed. In addition, the integrity of the sample is dependent upon the premise that a clear chain of responsibility for the sample integrity has been maintained. Without this "Chain-of-Custody", the integrity of the laboratory results may inevitably come into question.

The preservation and Chain-of-Custody (COC) protocols outlined in the following paragraphs are not intended to be all inclusive, and this protocol is written with the understanding that the sampling of certain media or analyses may require specific sample preservation. This protocol is, however, intended to cover the majority of the media and analyses performed as well as the COC procedures employed at the majority of waste disposal sites.

A COC program must be followed during sampling and handling activities from the field through laboratory operations. This program is designed to assure that each sample is accounted for at all times. Field data sheets, COC records, and sample labels must also be completed by the appropriate sampling and laboratory personnel for each sample. The objective of the sample custody identification and control system is to assure, to the extent practical, that:

- all samples are uniquely identified;
- the correct samples are analyzed for the correct parameters and are traceable through their records;
- important sample characteristics are preserved;
- samples are protected from damage or loss;
- any processing of samples (e.g., filtration, preservation) is documented; and
- client confidentially is maintained.

A sample is considered under a COC if it meets all of the following criteria:

- the sample is in your custody,
- the sample is in your view, after being in your possession,
- the sample is in your possession and then you locked it up to prevent tampering, and
- the sample is in a designated, secured area.

The following paragraphs outline GEC's preservation and COC protocol.

- 1) Prior to initiating any work, the Health and Safety Plan developed for the specific site activities should be reviewed by all field personnel. The indicated measures on the Plan should be enacted prior to initiation of any sampling activities. Any concerns not addressed in the Plan are to be brought immediately to the attention of the Health and Safety Officer. Personnel participating in the excavations will dress with protective equipment appropriate for the anticipated conditions.
- 2) Sample integrity is assured by use of containers appropriate to both the matrix to be sampled and the analytes of interest. Sample containers must be prepared in the laboratory in a manner consistent with USEPA protocols. Unless the proper sample bottle preparation and sample preservation measures are taken in the field, sample composition can be altered by contamination, degradation, biological transformation, chemical interaction, and other factors during the time between sample collection and analysis. Prior to sampling, GEC personnel will ensure that the sample containers obtained from either a laboratory or a commercial supplier have been prepared in accordance with NYSDEC and EPA protocols. Sample containers are to be used once and discarded. Under no circumstance should a soil, water or gaseous media which has been collected for analysis be placed in a previously used sample container unless that container has been recleaned and preserved by a certified laboratory.

As part of the COC protocol, sample containers should have prepared labels for each sample. The label should include sample identification, date and time of collection, sample parameters to be analyzed, any preservatives used, and the name of the sample collector.

Upon collection of the sample(s), documentation of chain-of-custody (i.e. COC form) should be initiated and should include at least the following:

date and time of sampling;

- sampling locations;
- sample bottle identification; and
- specific sample acquisition measures.

The COC and sample description requires:

- a unique identification of each sample;
- the name(s), address(es) and telephone number(s) of the sampler(s) and the person(s) shipping the samples and all subsequent transfers of custody;
- the type and method of analyses requested;
- the date and time of sample collection and transfer of custody; and
- the name(s) of those responsible for receiving the samples at the laboratory.
- 3) In some cases, field filtration of samples may be required. Information regarding the method of filtration should be determined in advance and communicated to the laboratory. Filtering of any sample collected for organic analysis should be avoided. Decanting of a liquid media is a preferred method for the removal of particulate matter. When field filtering is required, an appropriate filter medium must be selected to avoid potential sample contamination during the filtering process.
- 4) Sample holding times are specified for the initiation of chemical analyses, usually beginning at the time of sample collection but occasionally beginning at the time of sample receipt at the laboratory. This determination must be made prior to sampling to allow proper logistical planning for sample shipments. Holding times also vary with the regulatory basis under which analyses are conducted. It is essential that the laboratory be consulted before sampling take place in order to properly schedule work.
- 5) Sample containers are most often packed in plastic, insulated "coolers" for shipment. Bottles are to be packed tightly so that only minimal motion of the sample containers is possible. Materials which are considered to be highly hazardous may require special handling and packing for shipment. Ice, or a similar heat transfer fluid, should be placed over the top of the sample containers and should be placed within a water tight plastic bag to assure that the samples are kept as dry as possible. In addition, all applicable paper work should also be enclosed within a second water-tight bag and included in the cooler. The sample cooler should then be taped shut.
- 6) Upon receipt of the samples at the laboratory, any laboratory identification numbers should also be included on the COC form.

Finally, those responsible for receipt of the samples should be indicated on the COC form as well as the date and time of the sample drop-off.

Standard Operating Procedure Boring/Well Installation

This protocol is designed to insure that proper techniques are used, safety is considered, and quality assurance maintained during soil boring and well installation.

- DIGSAFE, municipalities and the owner are contacted prior to any soil boring
 or well installation to minimize chances of damaging underground utilities
 (DIGSAFE contacts utility companies to mark the location of utilities to the
 site). The Geologist or Inspector surveys the site visually for markings
 delineating the location of underground utilities. If warranted, the inspector
 modifies the drilling program to compensate for field conditions.
- The Geologist or Inspector continuously monitors all drilling activities and is responsible for maintaining independent field notes, well logs and ensuring that proper procedures are followed.
- Drilling equipment is steam cleaned prior to use in any boring and between borings (if necessary), to minimize potential cross contamination. At a minimum the following pieces of equipment are steam cleaned: augers, cutting heads, samplers, drill rods, and forks. The working end of the drill rig is also cleaned and inspected for evidence of hydraulic fluid or diesel fuel leaks.
- Subsurface soil samples are collected at a minimum of five foot intervals in accordance with standard ASTM methods for split spoon sampling. After logging soil characteristics, samples are collected. Two samples are placed in clean jars with an aluminum bladder below the lid for head space screening. Soil sample screening is performed in accordance with the GEC Jar Headspace Screening procedure. Samples with elevated readings (< 10 ppm) soil are quickly transferred into two clean VOA vials with Teflon liners. The vial is half filled and soil particles are removed from the lip of the vial to assure a proper seal with the lid. All samples are labeled in accordance with the GEC standard labeling identification system and handled/stored in compliance with USEPA protocols.
- The split spoon sampler is decontaminated in accordance with GEC's Decontamination Protocol after sample retieval and it is steam cleaned between borings. The Geologist may increase the frequency of steam cleaning as necessary.
- All cuttings from drilling remain on the subject property. If cuttings are designated as uncontaminated fill, via headspace screening, and the boring is not completed as a monitoring well the cuttings are used as backfill.

- Monitoring well screens are set to depths adequate for the required sampling. Monitoring wells are typically constructed with a silica sand filter surrounding and extending a few feet above the screen. The screen extends at least one to two feet above groundwater. The riser extends from the top of the screen to ground level, has a bentonite pellet seal above the screened interval, a cement seal and protective cover at the surface. No glues or solvents are employed in the well construction.
- Soil Logs are to be maintained by the Geologist and should contain the following:
 - Date and Location of boring/well
 - Drilling contractor
 - Job number
 - Depth of sampling
 - Boring number
 - Depth to well point.
 - Soil description includes; soil colors, grain size from greatest percentage to lowest, rock fragments, obvious fill constituents, staining, and odor if obvious.
 - Changes in soil strata and elevation of the water table are also noted.

Standard Operating Procedure HNU HW-101 Photoionization Detector

The HNu Instruments, Inc. HW-101 is a portable field instrument used to detect and approximate the concentrations of gaseous phase volatile compounds. The HW-101 operates by drawing the gas sample into an ionization chamber and exposing the gas to ultraviolet light, effecting photoionization.¹ The ions are collected on positive and negative electrodes, creating a current proportional to the concentration of ions. The HW-101 will detect molecules with ionization potentials of approximately 10.2 eV (electron volts) or less.²

II. Instrument Assembly and Battery Check

After the probe assembly and readout module are removed from the protective case, the end of the probe cable is connected to the 12-pin receptacle on the readout module. The tab on the probe cable and the corresponding notch on the 12-pin receptacle must be aligned for proper assembly. The filter nozzle is then attached to the end of the probe assembly.

The condition of the internal battery is determined prior to each use of the HW-101. Deep discharge damages the internal battery and should be avoided. The control knob on the face of the readout module is turned to the "BATT" position to determine the charge of the internal battery. If the meter needle deflects to within the green arc, battery charge is sufficient. If the meter needle is not within the green arc, the instrument must be recharged prior to use.

III. Calibration

The HW-101 requires calibration on a frequent and regular basis. The instrument should, at a minimum, be calibrated prior to commencing a day of sampling procedures and again at the end of the day to verify instrument response. In addition, the instrument must be calibrated subsequent to cleaning activities, or whenever suspect readings are noted.

Calibration procedures are initiated by rotating the control knob to the "STANDBY" position and observing the meter needle. If the meter does not indicate "0," the zero knob is rotated sufficiently to correct the discrepancy. The tip of the probe assembly is then connected to a cylinder containing a known concentration of isobutylene gas. The cylinder label will indicate a concentration "as benzene," the standard to which the instrument must be calibrated. The valve on the isobutylene cylinder is fully opened and the control knob of the HW-101 is

¹Claff, Roger E. (1991). "An Evaluation of Soil Gas and Geophysical Techniques for Detection of Hydrocarbons", *Health and Environmental Sciences API Publication Number 4509*. American Petroleum Institute, Washington D.C.

²HNu Systems, Inc. (1990) HW-101 Operation Manual.

rotated to the "0-200" parts per million detection scale position. After allowing approximately 1 minute for the reading to stabilize, the span knob is then rotated so as to match the instrument reading with the known concentration of the calibration gas, as printed on the cylinder label.

IV. Operation

Operation of the HW-101 requires two steps. First, the proper detection scale must be determined. If there is uncertainty regarding the correct detection scale, begin with the "0-2000" parts per million scale and increase sensitivity as necessary. Readings significantly higher than the upper limit of the scale may result in damage to the meter needle assembly. If meter needle deflection is less than 10% of the total range, the detection scale should be reduced by one step.

The second step involves placing the tip of the probe assembly within the volume of gas to be sampled and recording the reading. Although specific conditions affect response time, the HW-101 will generally record peak readings 2-3 seconds subsequent to the start of sampling. The peak reading observed for any given volume of gas is recorded as the result.

V. Notes

Detection Limit: Although information provided by HNu, Inc. indicates that the detection limit of the HW-101 is 0.2 parts per million, field experience has shown the practical detection limit to be approximately 1.0 parts per million.

Interferences: The sensitivity of photoionization processes are reduced by the presence of gases not ionized by the lamp. Water vapor or methane gas are two frequently encountered examples of such a gas. The HW-101 operator should be aware of and document the possible presence of such gases when interpreting data generated during photoionization screening.

Standard Operating Procedure Photovac 10S55 Gas Chromatograph

The Photovac 10S55 portable gas chromatograph (GC) allows a field operator to quickly screen gas-phase samples for the presence of volatile organic compounds (VOCs). In addition, the Photovac can tentatively identify and quantify selected VOCs by comparing retention times and response factors for samples to values stored in memory.

Preparation and Set-Up

Several preparatory activities must be completed prior to operating the GC. These activities include:

- Charge the internal battery for 24 hours prior to use.
- Bake sample containers for at least 24 hours, and purge the containers with hydrocarbon-free air prior to use.
- Bake all syringes for at least 24 hours prior to use at a temperature not higher than 37°C.
- Fill the internal carrier gas storage tank with hydrocarbon free air to a pressure of 1,500 pounds per square inch (psi).
- Replace the septum for the GC injection port.
- Gather all accessory materials together at one location and package for easy transportation. A complete listing of all required equipment is provided in Appendix A of this document.
- Immediately prior to departing for the site, prepare a gas-phase benzene standard and a hydrocarbon-free air standard.

After each of these steps has been completed, the GC and all supporting equipment can be transported to the site. Following arrival, a location must be selected from which the GC will be operated. The Photovac is sensitive to changes in ambient temperature and solar rediation; therefore, a location should be selected that is thermally stable and shaded. Indoor operation of the GC will provide the best stability.

Set-up is performed by completing the following steps:

- The GC is placed on a level surface and the case opened.
- Both power supplies are connected to a 120 volt AC outlet. The oven power supply requires the use of an AC/DC power converter.
- The computer module is opened and the oven temperature selector rotated to "30." The computer module is then closed and locked.
- The two (2) flow meter tubes are attached to the outlet ports and the flow meter leveled.

- The lower flow control knob (this knob has a red sticker on it) is rotated in a counterclockwise direction until carrier gas flow is approximately 10 mL/minute, as indicated by the flow meter.
- The computer module is activated by pressing the "on" button in the upper portion of the keypad.
- Site specific parameters are entered into or selected from the GC computer, including time and date, the retention time window (as a percentage), attenuation level, and the appropriate calibration library.

After the GC has been set-up according the procedures outlined above, it must be allowed to warm for a period of approximately one hour prior to further use. This warm up time is necessary to allow oven temperature and carrier gas flow rates to stabilize.

As part of the set-up procedures, the operator should prepare a log to record all activities conducted in the field. The operator enters data regarding set-up conditions, calibration procedures, sample identification and volume injected into the GC so that this data will be incorporated into result evaluation.

Calibration

Before commencing analysis, a stable baseline must be demonstrated and the instrument calibrated to a calibrant prepared from a primary benzene standard. A baseline determination is completed by performing a "run," without an injection. The resulting chromatograph will indicate whether the instrument is free from contamination and if oven temperature has stabilized.

Once the baseline run has been satisfactorily completed, the instrument is calibrated to a gaseous phase standard prepared from a primary benzene standard of approximately 1 part per million (volume/volume). Calibration is started by injecting a 100 μ L aliquot of benzene standard and comparing the observed retention time to the retention time in the computer library. (Procedures for injection are provided for reference in Appendix B). If the observed retention time varies by more than three tenths (0.3) of a second from the objective, the flow rate is adjusted and a benzene standard injected again. This procedure is repeated until the observed retention time is within 0.3 seconds of the objective. Frequently, repeated flow adjustments are required to properly set the instrument flow rate.

After the flow had been adjusted so that the observed retention time is acceptable, three successive 100 μL injections of benzene must be completed that have identical retention times. In addition, variation between the observed detector response for each of the three injections must be less than 10%.

When the above criteria have been met, the calibration library is adjusted to the benzene standard. The computer adjusts the internal retention times and

response factors for each component in the library relative to the observed retention times and response for the benzene standard.

After the above procedures have been followed, the initial calibration of the Photovac GC is complete. However; calibration of the GC also involves an ongoing process of retention time verification and adjustment. A benzene standard is injected before and after each sample is analyzed (known as "bracketing") to ensure that instrument conditions and retention times have not changed. Procedures for bracketing are outlined below, in Section IV.

Operation

After the set-up and initial calibration procedures are complete, analysis of samples can commence. A determination is made regarding the volume of sample to inject into the GC. This information is provided by the sample collectors who screen a sample duplicate with an HNU Instruments, Inc. HW-101 photoionizer. The GC operator utilizes this data to approximate levels of sample contamination and determine an appropriate injection volume.

Once the correct volume of sample has been determined, it is injected into the GC and the instrument is allowed to run for approximately 720 seconds. Immediately following completion of the sample run, a retention time verification, or bracketing, standard is injected. If the observed response time for the bracketing standard is not within 0.3 seconds of the objective, the flow must be adjusted and an additional benzene standard injected. After the observed retention time is within 0.3 seconds of the objective, the retention times stored within the GC computer library are recalibrated to the benzene standard and the sample is injected again. This process is repeated until the bracketing standards are within 0.3 seconds of the objective time. This procedure is repeated for each sample to be analyzed.

Quality Control/Quality Assurance

QA/QC involves determinations of accuracy and precision. For purposes of the screening analysis outlined in this protocol, accuracy is sufficiently determined by the initial and on-going calibration to known standards and by the periodic analysis of hydrocarbon-free air blanks. Precision is determined by the analysis of sample duplicates and by "bracketing" each sample run with injections of a benzene standard.

A minimum of one (1) duplicate and one (1) blank is analyzed for every ten (10) samples. This frequency is in accordance with Massachusetts Department of Environmental Protection (MDEP) Policy #WSC-89-004 (Minimum Standards for Analytical Data for Remedial Response Actions Under M.G.L. c. 21E) and guidelines presented in United States Environmental Protection Agency (USEPA) Policy #540/G-90/004 (Quality Assurance/Quality Control Guidance for Removal

Activities). In addition, retention time verification is performed prior and subsequent to each sample analysis via a secondary benzene standard.

Practical quantitation limits are generated for each compound of interest by injecting progressively smaller volumes of a representative contaminated sample. This injection of progressively smaller volumes simulates the injection of progressively lower concentrations of the compound of interest, allowing the practical quantitation limit to be determined. Smaller volumes are injected until detector response is no longer linear (within 20 percent of expected response). The lowest volume for which response was linear is then repeated three (3) times to verify detector precision at the practical quantitation limit. Results varying more than 20 percent from one another indicate unacceptable precision, and a higher practical quantitation limit must be utilized.

Injection Technique

Injections of gas samples into the GC should be performed in the following manner:

- Always utilize "gas-tight" syringes when performing injections into the GC.
- Select a syringe of the correct size for the injection volume. The sample should occupy between 20% and 80% of total syringe volume.
- Flush the syringe three (3) times with ambient air and three (3) times with hydrocarbon-free air prior to use. Discharge the hydrocarbon-free air into the atmosphere, not back into the storage container.
- Flush the syringe three (3) times with the sample or standard to be injected without removing the syringe from the sample or standard container.
- Injections into the GC must be highly consistent to generate reproducible peaks. Fully insert the syringe prior to injecting, discharge the contents quickly, and be sure the entire contents of the syringe has been discharged before removing the syringe. Withdraw the syringe quickly after the injection is complete.
- If the chromatograph indicates a highly contaminated sample, flush the syringe used to inject the sample repeatedly and inject hydrocarbon-free air to determine whether the syringe has been contaminated.
- Response for replicate injections should be reproducible within 20%. If observed response varies beyond 20% the syringe should be checked for proper mechanical operation. This involves checking the needle for

-	proper seating against the syringe body and checking if the plunger tip has become worn, resulting in a faulty seal.
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Standard Operating Procedure Photovac Tip

I. Theory

The Photovac TIP is a field monitoring instrument capable of detecting a wide range of volatile and semi-volatile organic compounds as well as some inorganic compounds in air and are collectively designated total ionizable compounds (TICs). The instrument uses a 10.6 eV lamp and can detect compounds having ionization potentials less than 10.6 eV. See attached list of compounds and ionization potentials. The detection limit for the Photovac TIP is 1 part per million (ppm) TICs in air. The TIP is calibrated to zero with hydrocarbon free air and 100 ppm isobutylene as a benzene equivalent.

The TIP does not distinguish between different chemical compounds and the signal produced represents a composite reading of all TICs. Gross levels of contamination are those above 10 ppm, as read on the LCD. The correct implication to be drawn when the TIP registers over 10 ppm is that there is a potentially serious situation, depending on suspected contaminants present, and may require further investigation using more specific detection equipment.

Limitations:

- Relative humidity can cause substantial interference and inaccuracies in organic response interpretation.
- The lamp window must be periodically cleaned to ensure ionization of the air containments.
- Instrument response to a gas or vapor may significantly change when the gas or vapor is mixed with air contaminants Individual contaminant levels are not necessarily additive within a mixture of air.

II. Procedures for Use

Field Preparation:

Prior to using the TIP, review the manual.

In advance of using the TIP in the field, both batteries must be fully charged for 16 hours. The TIP has two batteries, one within the instrument itself and a second in the battery pack. Both batteries, fully charged and used together will provide approximately eight hours of continuous use. To maximize the operating time of the TIP, the batteries should be fully discharged prior to recharging. The batteries are fully discharged by running the TIP on both batteries, until the "LOWBAT" signal appears on the read out. DO NOT UNDER CHARGE! as this could reduce the charge held by the batteries and reduce the running time of the TIP.

III. Calibration

Follow the procedure outlined in the TIP User's Manual. Plus the following:

- Always fill the zero air bag first.
- The TIP is to be Zeroed using the reference Zero Air supplied in the canister.
- Prior to filling the bag, purge the regulator and fittings with zero air.
- Then purge the regulator and fittings with isobutylene.
- Fill the isobutylene bag.
- After using the TIP be sure to record use and any maintenance performed on the instrument in the Field Log.

APPENDIX B HEALTH AND SAFETY PLAN

GEC has prepared a site-specific Health and Safety Plan designed to protect personnel involved in the performance of investigatory and remedial activities at the site. This plan addresses health and safety concerns that may be encountered during the sampling and investigation activities proposed in this work plan. The Health and Safety Plan should be updated in the event that additional information regarding site contaminants is encountered or in the event that activities not specified in the plan are conducted.

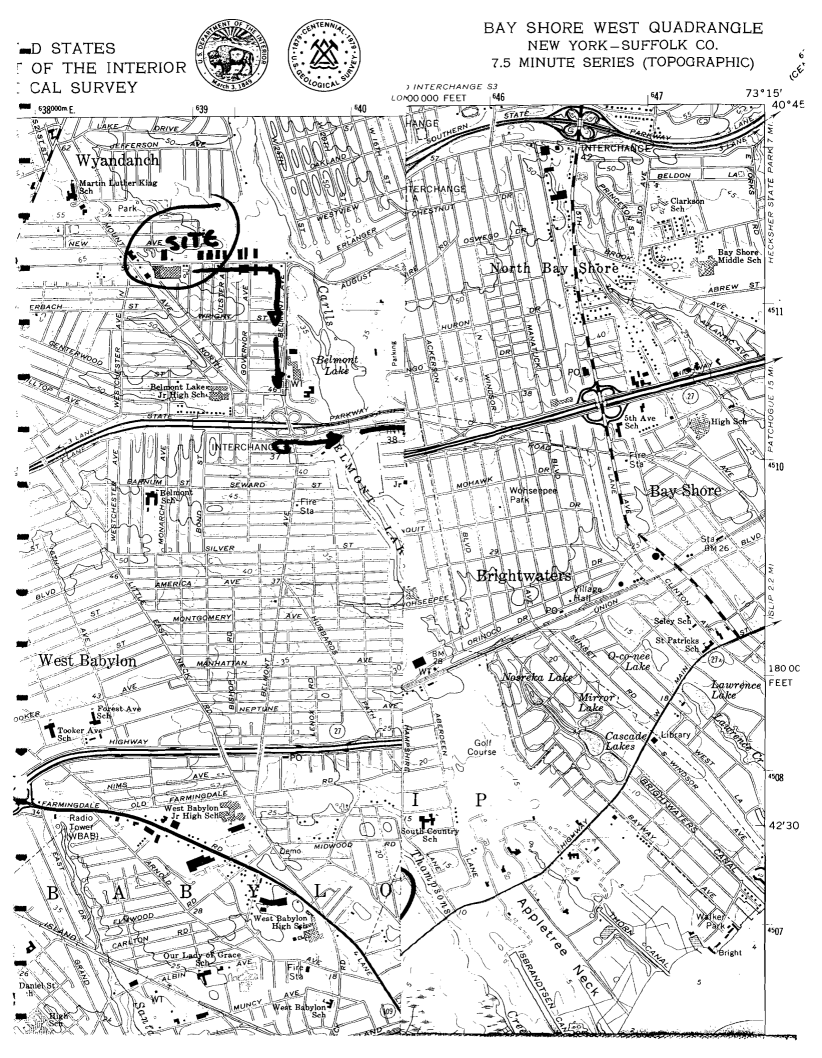
The Health and Safety Plan, included with this Work Plan, does not specifically address health and safety issues associated with personnel not conducting the site investigation but who may be working at the site. Contamination at the site is limited to the presence of metals and volatile organic compounds in subsurface soil and groundwater. As such, the plan is protective of those likely to come in contact with subsurface soil and/or groundwater; i.e. those conducting the investigations. During investigation of remedial activities, GEC will establish an exclusion zone, within which only personnel covered by the health and safety plan will be permitted.

The Health and Safety Plan has been prepared in accordance with 29 CFR 1910 and all other applicable standards by a Health and Safety Professional.

HEALTH AND SAFETY PLAN GOLDMAN ENVIRONMENTAL CONSULTANTS

SITE DESCRIPTION	ON		
Date of Original P Date of Revision a	Plan: <u>9/30/94</u> and Modification: <u>3/23</u>		ber: <u>444-006-94</u>
Site Name: Site Address:			
Site Conditions:	Industrial facility with	unpaved and paved are	<u>as</u>
excavation activities	ties including but not excavations and stoo	perations; soil and grout limited to: test pitting skpiling, and trenching;	<u>ig, UST removals,</u>
hazardous waste		he Massachusetts MCP Yes/No? <u>Yes</u>	or an uncontrolled
	EMERGENCY	INFORMATION	
Nearest Phone &	Location: Inside build	ling - 516-643-3500	
Nearest two-way	radio: <u>None Availabl</u>	<u>e</u>	
	<u>Number</u>	<u>Loca</u>	<u>tion</u>
Fire: Police: Ambulance: Hospital:	854-8100 911	Wyandanch V 1st Precinct, B Wyandanch 0 1000 Montauk Highwa	labylon n, F.D.
Does hospital hav	ve chemical trauma ca	pability? Yes X] No
approximately on Ave. to South Sta Robert Moses Ca	<u>e half mile, take a ric</u> te Parkway. Follow St auseway (south). Foll	rn right onto Wyandanch tht onto Belmont Avenu S Parkway south to exit ow to Exit 27A West ar th three traffic lights an	ue. Follow Belmont 40 (south) and onto nd take right off exit

SEE MAP ATTACHED



Additional important Phone Numbers.	•		
Goldman Environmental DEP Spill Reporting Other State Agency:	(617) 9 ———	961-1200	
Chemtrec National Response Center ATSDR AT & F (Explosive Information) Pesticide Information Service CMA Chemical Referral Center National Poison Control Center U. S. DOT LEPC Contact: Not Applicable Name: Title: Phone Number:	(800) 4 DAY: (4 (800) 4 (800) 8 r (800) 2 r (800) 9 DAY:		
SPECIAL LOCAL EMERGENCY PLAN	INING COMMIT	TEE REQUIRE	MENTS (if any
NA			
DIGSAFE INFORMATION (if warranted LILCO No.: 820113 Ag Date / Time: N/A PERSONAL PROTECTIVE EQUIPMENT The following level of protection will be	encies Contacte ! - NT	d: LILCO-gas NYNEX	& electric
Task to be Level of	0 4004.	Δ	ir Purification
Performed Protection	Coverall Glo	ove In/Out	Cartridge
1) Sample Collection. D	Cotton Lat	tex/Nitrile	<u>None</u>
slug tests, drilling and excavation systems	n activities, and	operations of re	emedial
2) <u>Upgrade</u> C	Tyvek Late	ex/Nitrile	Organic _
for all tasks	· · · · · · · · · · · · · · · · · · ·		
3)			

Oxygen Meter				
Rubber Boots Other Other				
HAZARD DESCRIPTION				
Physical Hazards Heat (A.8) X Cold (A.9) X Noise (A.10) X Underground Utilities x Overhead Utilities x Heavy Equipment x Physical Hazards (continued) Confined Spaces Pressurized Airlines Explosive (A.11) Ladders or Scaffolds Unguarded floor/ground openings Liquids in open containers, ponds, or lagoons Radiation (A.12) Physical Hazards (A.13) x Oxygen Deficiency (A.14) Traffic x Other				
HAZARD EVALUATION				
Suspected Sources of Contamination:				
Ground water has been shown to contain petroleum, volatile organic compounds including chlorinated solvents, and metals typical of foundry use. Concentrations of ground water contaminants have not yet been determined. Free-phase petroleum has been detected in monitoring wells on the site. Respiratory Hazards:				
The chemical contaminants detected on-site can represent an exposure hazard in concentrated form. Inhalation of petroleum vapors emitted from soils or ground water is the primary respiratory hazard. During soil boring and well installation air in the breathing zone will be monitored using a PID calibrated to a benzene equivalent. Readings consistently above 5 ppm TIC threshold limit in the breathing zone will require an upgrade to level C protection. Soil Samples collected during soil boring will also be screened with the PID for TICs. During sample collection, the PID will be used to monitor the breathing zone for TICs. Readings consistently above the 5 ppm TIC threshold limit will require an upgrade to level C. If such a situation exists, personnel who have not been fit tested for work at level C will remain upwind of the area, where TIC threshold cannot be exceeded. Transient exceedences above the 5 ppm TICs in the breathing zone will require Level D work stoppage until levels return to subthreshold levels, after which work in level D may resume. Dermal Hazards:				
Contact to skin during sample collection will be minimized as protective clothing will be worn by workers. Latex and nitrile gloves should provide sufficient protection from the dermal hazards. Workers will adhere to good personal hygiene practices.				

Ingestion Hazards:

Ingestion of contaminated water is considered unlikely as hand to mouth contact will be avoided and face shields will be worn during water sampling. Personal hygiene should be sufficient to prevent ingestion of contaminants.

Physical Hazards:

Heavy equipment and obstacles typical of construction sites may be present. Extreme care will be exercised while conducting all on site work with respect to physical hazards. GEC employees will not enter trenches or excavations deeper than 4 feet.

DECONTAMINATION

Step by Step Decontamination Procedures and Solutions:

Personal Protective Equipment (PPE): <u>Tyvek suits will be disposed as solid waste</u>. All other PPE will be rinsed with soapy water, DI water, and methanol and DI water. For gasoline/oil contaminated PPE/sampling equipment, acetone, then hexane should be substituted to remove stubborn petroleum residue.

Sampling Equipment: <u>Scrubbed with soapy water, rinsed with DI water and methanol and DI water.</u>

Other Equipment: See sampling equipment

Disposal of waste clothing, decontamination solution, etc.: <u>Decon solutions</u> will be allowed to evaporate, clothing discarded into the dumpster.

MSDS(s) for Methanol is/are attached.

WORK LIMITATIONS OR PRECAUTIONS

Describe limitations due to time of day, weather, situations, if any: On site work may be suspended due to severe weather conditions, and night time work will be avoided.

Sample Preservatives: Acids and caustics used as preservatives should be handled with gloves and safety glasses.

SIGNATURES

All site personnel have read the above plan and are familiar with its provisions:

Name		Signa	ature
Tenus Dale	V	Som It	kili o
Bun of	reg /	idh at	
SAN BUTTON 2	miller /	The was	
JASON MALPO	1 M		
Written by:	Jamy Daky	Date:	3/27/95
		 -	
Approved by:	athlen W! Mal	Date:	4/12/95

A.8 HEAT STRESS

EFFECTS OF HEAT

Heat produced within the body is brought to the surface largely by the blood stream. When at the surface, the heat escapes the body by conduction, radiation and convection. Interference with heat loss leads to a raised body temperature similar to a fever. This accelerates certain body processes which in turn produce more heat, requiring not only the normal elimination of heat, but an extra requirement for heat loss.

When the temperature of the air becomes equal to or higher than the body temperature, the body must rely on losing it's body heat through the sweating process. As the air becomes more humid, this type of heat loss becomes inefficient and heat loss decreases. It is on such days, hot with high humidity, or a succession of such days, that the conditions are ideal for heat stress. Emergencies due to heat stress are described by three categories: heat cramps, heat exhaustion, and heat stroke.

HEAT STRESS SYMPTOMS AND FIRST AID

1) Heat Cramps

CAUSE

Profuse sweating leads to a loss of body salts and electrolytes. If these are not replaced, painful cramps occur. Cramps could also occur by drinking ice water or other drinks too quickly or in too large a quantity.

SYMPTOMS

- a. Muscle cramps in legs and abdomen.
- b. Pain accompanying cramps.
- c. Faintness.
- d. Profuse perspiration.

FIRST AID

- a. Remove individual to a cool place.
- b. Give individual sips of Gatorade or equivalent to replace salts and electrolytes.
- c. Apply manual pressure to cramped muscle.
- d. Remove patient to a hospital if more serious conditions exist.

2) Heat Exhaustion

CAUSE

Occurs in individuals working in hot environments, and may be associated with heat cramps. It is brought about by the pooling of blood near the surface of the skin. Heat is transported by the blood to the surface of the skin. This process dilates the skin vessels so that a large amount of blood is pooled in the skin and lower extremities. This condition can lead to an inadequate return of blood to

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the heart and then to collapse.

SYMPTOMS

Weak pulse; Rapid and unusually shallow breathing; Generalized weakness; Pale and clammy skin; Profuse sweating; Dizziness; Unconscious; Appearance of having fainted.

FIRST AID

- a. Remove victim to a cool place and remove as much clothing as possible.
- b. Administer cool water, Gatorade, or its equivalent.
- c. If possible, fan the victim to remove heat by convection, but do not allow chilling or overcooling.
- d. Treat for shock if necessary, and remove to a medical facility if serious conditions persist.

3) Heat Stroke

CAUSE

Heat stroke is a profound disturbance of the heat regulating mechanism. It occurs from direct exposure to the sun for prolonged periods. Poor air circulation and poor physical condition add to this threat. This is a serious threat to life and carries a twenty percent mortality rate.

SYMPTOMS

Sudden onset; Dry, hot and flushed skin; Dilated pupils; Early loss of consciousness; Full and fast pulse; Breathing deep at first, later shallow and almost absent; Muscle twitching; Excessively high body temperatures.

FIRST AID

- Transportation to a medical facility should occur as quickly as possible.
- b. Remove patient to a cool environment and remove as much clothing as possible.
- c. Assure an open airway.
- d. Cool body by dousing with water or wrapping in a wet sheet.
- e. Place cold packs or ice under the arms, around the neck, at the ankles, or on forehead.
- f. Protect victim from injury during convulsions.

PREVENTIVE MEASURES

- a. Assure that all employees drink plenty of fluid. A fluid is needed, such as Gatorade, that will replace body salts and electrolytes. Such fluids should be consumed for a period that includes 12 hours before to 12 hours after the anticipated activities which induce heat stress. Tap water is insufficient to replace body salts and electrolytes under heat stress conditions.
- b. Assure frequent breaks to avoid over heating.
- c. Revise work schedules, when necessary, to take advantage of the cooler parts of the day.

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A.9 COLD EXPOSURE

COLD EXPOSURE SYMPTOMS AND FIRST AID

1) Hypothermia

CAUSE

This is a result of the body losing heat faster than it can produce it. Most critical condition for hypothermia is above freezing weather on a wet and windy day. The mind thinks it is too warm for danger and therefore disregards the symptoms. The wetness against the body acts as a conductor which takes excessive amounts of heat from the body. This heat loss affects the internal organs. Hypothermia sets in when the body temperature reduces to 95 degrees Fahrenheit. (Note- Alcohol and drugs accelerate heat loss through vasodilation. This means that the blood vessels dilate.)

STAGES AND SYMPTOMS

- a. Shivering.
- b. Apathy.(Key Symptom)
- c. Unconsciousness.
- d. Freezing.
- e. Death.

FIRST AID

- a. Remove individual from the cold.
- b. Remove wet clothes, get out of the wind.
- c. Rewarm victim:
 - 1. Actively- Apply heat to the body. This can be done with a powered heat source, the sun, or by the heat of another individuals body heat.
 - 2. Passively- Wrap individual in a blanket so that they begin to rewarm themselves.

2) Frostbite

THREE TYPES

- 1) <u>Frostnip- Whitened or grayish area of the skin.</u> A burning sensation is experienced.
- 2) <u>Superficial Frostbite</u>- This is slightly worse than frostnip. The skin is whitened or grayish with a waxy feeling to it. The skin experiences very little pain at this point.
- 3) <u>Deep Frostbite</u>- Completely frozen area, perhaps to the bone. The area is white and hard. The individual loses sensation in that area.

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FIRST AID

1) For Frostnip and Superficial Frostbite

- a. Remove victim from the cold.
- b. Give the individual liquids.
- c. Remove clothing from the area.
- d. Soak affected area in a warm water bath- 102-108°F.

2) For Deep Frostbite

- a. Keep area frozen.
- b. Get the victim to a doctor as quickly as possible.
- c. Do not try to treat this in the field.

PREVENTIVE MEASURES

- a. Dress warmly, but do not overdress. Sweating will only form a conduction to take heat away from the body.
- b. Don't touch cold metal.
- c. Keep moving.
- d. Take shelter whenever possible.
- e. Drink warm soups and liquids periodically.
- f. Wear layered clothing and a hat.
- g. Have a spare dry set of clothing on site.

Page 2 of 2

A.10 NOISE

Exposure Route or Cause

Compressors, machinery, and large equipment.

Suspected Site Specific Sources

Symptoms and Effects

Temporary or permanent hearing loss, aural pain, nausea, reduced muscular control (when exposures are severe), distraction, and interference with communication.

Measurement or Measure Devices

Sound levelmeter and octave band analyzer.

Prevention

- a. Shielding or enclosure of source.
- b. Distance/isolation.
- c. Substitution of equipment/machines generating less noise.

Personal Protection

Ear muffs, ear plugs, or noise-insulating earphone

Additional Comments

a. Use of earphones with communication built-in can improve coordination and warnings.

b. Use of earplugs must include consideration of potential indirect chemical exposures if the earplugs become contaminated.

REFERENCES:

Information on this page extraceted from:

Martin, W. F., Lippit, J. M., Prothero, T. G., Hazardous Waste Handbook for Health and Safety. Butterworth Publishers, pp. 10-11, 1987.





MATERIAL SAFETY DATA SHEET

4011

IDENTIFICATION

Name

Methanol

Grade

Synonyms Methyl alcohol, Wood

alcohol, Carbinol

CAS Name

ethanol

■D. Nos./Codes

NIOSH Registry No. PC1400000

fanufacturer/Distributor

E. I. du Pont de Nemours & Co. (Inc.)

1dress

Wilmington, DE 19898

PHYSICAL DATA

: →iling Point, 760 mm Hg

37.7°C (148.5°F)

Specific Gravity

2 792 at 20°C (68°F)

Paper Density

1.1 (Air = 1)

olatiles by Vol.

.003

-orm

Appearance

. uid Clear

:__nformation

RAZARDOUS COMPONENTS

ic irial(s)

ethanol '

DU PONT 15 A SUPPLIETE OF METHANOL SOLD Chemical Family BY STERLING-CLARK-LURTON CORP.

Alcohol

Formula

CH₃OH

CAS Registry No.

67-56-1

Du Pont Registry No.

Product Information and Emergency Phone ---(302) 774-2421

Transportation Emergency Phone (800) 424-9300

Melting Point

-97.8°C (-144°F)

Vapor Pressure

138 mm Hg at 25°C (77°F)

200 mm Hg at 37.7°C (100°F)

Solubility in H=O

Evaporation Rate (Butyl Acetate = 1) > 1

Color

Odor

Colorless

Faint alcoholic

Octanol/Water Partition Coefficient

Approximate %

100

* TARDOUS REACTIVITY

llity. able:

compatibility Reacts vigorously with strong oxidizers, chromic anhydride, lead : nlorate, perchloric acids.

composition

curs from heat and reaction with materials above.

- v erization
- _mot occur.

5 L7

Date: 7/82

ree Meterial Salety Data Sheet relates only to the specific material designated herein and does not relate to use in combination with any other material or in any proce of th herein is furnished free of charge and is based on technical data that Ou Pont believes to be reliable. It is intended for use by persona having technical still and at ion and risk. Since conditions of use are outside our control, we make no warranties, express or implied, and assume no itability in connection with any and herein is to be taken as a license to operate under or a recommendation to infringe any patenta

AE AND EXPLOSION DATA

Hash Point

Method

Autoignition Temperature

11°C (52°F)

TCC

385°C (725°F)

Flammable Limits in Air, % by Vol.

Lower 6.0% Upper 36 %

Fire and Explosion Hazards Flammable. Flame is invisible in daylight. Methanol-water mixtures will burn unless very dilute; mixtures with 25% or more methanol are DOT Class I flammable liquids.

Extinguishing Media

Dry chemical, CO2, water spray, "alcohol" foam.

Special Fire Fighting Instructions

Use water spray to cool tanks or containers.

HEALTH HAZARD INFORMATION

Exposure Limits

OSHA 8-hour Time Weighted Average (TWA) and ACGIH TLV® TWA = 200 ppm, 260 mg/m¹. ACGIH adds "skin" notation.

Significant Routes and Effects of Exposure

Harmful if inhaled.

May be fatal or cause blindness if swallowed.

Cannot be made nonpoisonous.

May cause irritation.

LD50 (oral, rats) = 12,900 mg/kg; LC50 (rats, 1 hour) = 145,000 ppm.

| Safety Precautions

Avoid contact with eyes, skin or clothing.

Avoid prolonged or repeated breathing of vapor.

· Wash thoroughly after handling.

First Aid

_If swallowed: Induce vomiting immediately by giving two glasses of water and

sticking finger down throat.

If inhaled: Remove to fresh air. If not breathing, give artificial respiration;

preferably mouth-to-mouth. If breathing is difficult, give oxygen.

Call a physician.

mIn case of eye contact: Immediately flush with plenty of water for at least

15 minutes. Call a physician.

_In case of skin contact: Flush with water.

18717

Date: 7/82

JIEU HUN INFURMATION

pod general ventilation should be provided to keep vapor concentrations below dexposure limits.

Personal Protective Equipment

Have available and wear where appropriate: Safety spectacles (side shields preferred), chemical splash goggles, hard hat with brim, face shield (full length), neoprene coated cotton gloves, solvent resistant gloves, rubber safety shoes or rubber overshoes, rubber apron, appropriate respiratory protection (See Reference 2, page 4).

Other

DISPOSAL INFORMATION

: Aquatic Toxicity

TLm 96: > 1000 ppm Spill, Leak or Release

Dike large spills. Flush spill area with plenty of water. Do not flush to sewer. Comply with federal, state, and local regulations on reporting releases.

, Waste Disposal

Comply with federal, state and local regulations. If approved, incineration, bio-oxidation, subsurface injection, or disposal contractor may be used.

SHIPPING INFORMATION

Transportation

DOT Hazard Class.": Flammable Liquid

IMCO Class.: 3.2

DOT Shipping Name: Methyl Alcohol UN No.: 1230

NA No.:

RQ Quantity*:

*49 CFR 172.101 Shipping Containers

Barge, railroad tank cars, tank trucks.

Storage Conditions

Keep away from heat, sparks and flame. Keep container tightly closed. Do not store or mix with strong oxidizers, chromic anhydride, lead perchlorate or perchloric acid. Store in adequately ventilated area.

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DITIONAL INFORMATION AND REFERENCES

Reference 1) Du Pont Methanol Properties, Uses, Storage and Handling Bulletin.

2) DHEW (NIOSH) Publication No. 76-189" A Guide to Industrial Respiratory Protection", available from Dept. HHS/NIOSH, 4676 Columbia Parkway, Cincinnati, OH 45226, phone (513) 684-4287.



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Date: 7/82

APPENDIX C

EQUIPMENT SPECIFICATIONS

Engineering Specifications

Soil Vapor Extraction System

PART I GENERAL

1.1 <u>INTRODUCTION</u>

The following Engineering Specifications for a Soil Vapor Extraction (SVE) system have been prepared by Goldman Environmental Consultants, Inc. (GEC) on behalf of Watts Industries, Inc. (Watts) and in accordance with a NYSDEC-approved Interim Remedial Measure (IRM) Work Plan for the Jameco facility located in Wyandanch, NY. A SVE pilot test has been conducted at the site to determine the feasibility of remediating soils and to identify design parameters for a full-scale soil extraction system.

1.2 <u>SITE DESCRIPTION</u>

Jameco Industries, Inc., a subsidiary of Watts Industries, received a NYSDEC Consent Order which requires remediation of trichloroethylene (TCE) at the site to New York State drinking water standards. GEC conducted a soil gas survey inside the building near plating and degreasing operations and VOC concentrations were detected as high as 1,800 ppm in soil gas.

The SVE system described by these specifications will be installed within the building which has the following available utilities:

Electricity 230/460 VAC, 3 phase, 60 Hz Domestic Water

1.3 <u>MECHANICAL PLANS</u>

The enclosed plans are intended to be diagrammatic only. They are not intended to show every item in its exact location. The contractor shall verify the actual dimensions of the equipment proposed. The installation shall be within the limitations imposed by the architectural, structural and electrical requirements, with adequate space for maintenance.

PART II EXTRACTION WELLS AND MANIFOLD PIPING

2.1 <u>EXTRACTION WELLS</u>

Extraction wells shall be 4-inch in diameter and 8 feet deep with 4 feet Sch 40, 0.010 slot screen from 4 to 8 feet, bentonite seal one foot above screen and 1-1/2 feet cement seal to sump level. See the Extraction Well diagram for more detail.

2.2 <u>PIPING</u>

Manifold piping connects the extraction points to the skid-mounted SVE system. Piping and fittings shall be 4-inch diameter, schedule 80, industrial grade, polyvinyl chloride (PVC) manufactured from Type I, Grade 1 conforming to ASTM D-1784, D-1785, D-2466 and D-2564. All pipe fittings including valves, unions and flanges shall be pressure rated at 150 psi

Each vapor extraction point shall be individually piped, valved and connected to a manifold located upstream of a moisture separator. A flow and vacuum gauge, and air flow control valve shall be installed on each pipe at the extraction well. See Process Piping Layout Drawing No. PP-1 for more details.

Branches shall intersect the main pipe at a reducer and the angle of entry at the intersection shall not exceed 45° to the direction of flow.

ASA 150 psi rated flanges shall be installed on both inlet and outlet piping of the SVE system.

2.3 TRENCHES

Piping shall be placed in shallow utility trenches that lead from the extraction points to the SVE system as shown in Drawing No.

UT-1. Utility trenches shall be 12-inches or 24-inches wide and 12-inches deep and coated with epoxy. Trenches shall be sloped toward the spill collection sump and piping shall be sloped towards the extraction well so that the condensate or entrained groundwater flows back toward the well.

The trenches shall be covered with medium traffic fiberglass or steel grating panels as manufactured by Fibergrate or equivalent. The panel shall be 24-inches wide, 48-inches long and 1-1/2-inches thick.

PART III <u>SVE SYSTEM DESIGN</u>

3.1 SCOPE

Design, furnish and deliver to the Wynadanch, NY job site one complete Soil Vapor Extraction (SVE) System. The SVE system shall be supplied with all the appurtenances and accessories, unless otherwise specifically excluded herein, necessary to make the equipment, when installed per the Vendor's installation recommendations, complete and operable.

3.2 SYSTEM DESCRIPTION

The SVE system shall be a skid-mounted VACPAC, Bisco Environmental, Inc. Model URAI59, or equivalent, which shall be designed to remediate the VOC contaminants from the soil beneath a portion of the building. The system shall be capable of extracting VOC vapors at the rate of 500 acfm with 85 inches w.c. and at a blower discharge pressure of 0.5 psig. The SVE system shall consist of a moisture separator with level control sensors to control the operation of the transfer pump and the positive displacement blower, inlet particulate filter, and discharge silencer. The SVE system shall include provisions to continuously monitor the extraction well vacuum, air flow, and air temperature; and pressure and temperature on the discharge side of the system shall also be monitored. See Drawing No. P&ID-1 for more details for system operation and controls. The following are major components and features of the SVE system.

3.2.1 <u>Positive Displacement Blower</u> (PDB-1)

The blower shall be a positive displacement blower Roots Model URAI 59, or equivalent. The blower shall be capable of providing 350 to 500 CFM at a vacuum of 60 to 80 inches of water (which includes an additional 10 inches of pressure drop for friction losses).

The blower shall consist of two "Figure 8" lobe impellers mounted on parallel shafts, rotating in opposite directions. As each impeller passes the blower inlet, it traps a definite volume of air and carries it around the case to the blower outlet, where the air is discharged. With constant speed operation, the displaced air volume is essentially the same regardless of pressure, temperature or barometric pressure. Timing gears control the relative position of the impellers to each other and

maintain small but definite clearances, which allows operation without lubrication being required inside the air casing.

The blower shall consist of a cast iron casing; carburized and ground alloy steel, spur timing gears that are secured to steel shafts with a taper mounting and locknut; and cast iron involute impellers.

The blower motor shall be a foot-mounted, NEMA A Design B, TEFC, standard industrial continuous duty, ball bearing, variable torque type suitable for operation at 15 HP, 230/460 VAC, 3 phase, and 60 Hertz.

3.2.2 <u>Moisture Separator</u> (MS-1)

The moisture separator shall have a liquid capacity of 120 gal and shall be designed for use in a soil vapor extraction system. The separator shall be capable of continuous operation with a pressure drop of less than six inches of water at the rated flow of 500 SCFM. The separator unit shall be capable of operation under various inlet conditions ranging from a fine mist to slugs of water.

The separator unit shall be equipped with a transfer pump and liquid level controls. The level control switches shall be single polarity, reed-type, float switches. High, low, and mid-level switches shall control on-off functions for the transfer pumps and blower. Liquid levels controlled/monitored shall be independently set and field adjustable with levels identified as:

"High Level" - Blower Shut-off

"Medium Level" - Transfer Pump On

"Low Level" - Transfer Pump Off and Blower On

3.2.3 <u>Inlet Particulate Filter</u> (IPF-1)

The inlet particulate filter shall be installed after the moisture separator and before the blower. The filter unit shall protect the blower from harmful dust and other particles that may be drawn into the blower through the air distribution system. The filter unit shall be capable of removing 8 to 10 micron size particles with 97 - 98% efficiency.

3.2.4 <u>Differential Pressure Transmitter</u> (DPT-1)

DPT-1 shall be a diaphragm (Bourdon tube) operated with a

sensing element motion that is restrained by a calibrated spring affixed with a silicon strain gauge transducer. DPT-1 shall be capable of measuring pressure up to 30 psia and it shall sense a single pressure relative to atmosphere and convert that into a 4-20 mA output signal. When DPT-1 reads the pressure change from the set point, DPT-1 shall transmit the signal to a pressure indicating controller (PIC) which, in turn, controls a motorized three-way valve position to direct flow to a parallel carbon canister.

3.2.5 <u>Vacuum Gauge</u> (VG)

The vacuum gauge shall be a diaphragm-actuated dial for measuring vacuum in the extraction well piping. It shall read from 0 - 100 inches of water in 2-inch divisions.

3.2.6 <u>Temperature Gauge</u> (TI)

Temperature gauges shall be installed in the extraction well piping for measuring the vapor temperature in the piping. The gauge shall be capable of reading 0-200°C (392°F) with ±1% accuracy.

3.2.7 <u>Flow Gauge</u> (Fl)

Flow gauges shall be installed in the extraction well piping for measuring the air flow in CFM. The gauge shall be capable of directly reading 0-740 SCFM.

3.2.8 Vacuum Relief Valve

A vacuum relief valve shall be installed at the blower inlet to prevent excessive system vacuum that could result from line restrictions. The relief valve shall be 2" cast iron and manually adjustable from 1.0 to 4.5 psig.

3.2.9 <u>Transfer Pump</u> (TP-1)

Transfer pump TP-1 shall be a magnetic drive, end suction centrifugal-type pump which shall have pumping capacity of up to 15 gallons per minute (GPM) at 20 ft. total dynamic head (TDH). The pump body shall be polypropylene with no mechanical seals and close-coupled to a 1/3 HP, 3450 RPM, 115/230V, single phase and 60Hz TEFC motor complete with dry cut protector to monitor electrical voltage and amperes to protect the pump from destructive dead-head or run dry

conditions.

3.2.10 <u>Control Panel</u>

A control panel cabinet shall be provided complete with a Programmable Logic Controller (PLC), remote monitoring and fax/pager notification capabilities, motor starter with thermal overload protection, three position "Hand/Off/Auto" (HOA) selector switches for blower and pump, and blower and pump status indicator run lights. Additionally, a high level alarm indicator light for the moisture separator, high temperature indicator and alarm light, and a common manual reset button shall be mounted. The cabinet shall be a NEMA 4, polyvinyl chloride or carbon steel epoxy coated, weathertight and mounted on the skid.

3.3 VOC CONTROL SYSTEM

During the pilot test, VOC concentrations were measured as high as 72 ppm and will probably drop below 10 ppm within two (2) days of the system operation. GEC requires carbon adsorption VOC control technology because of low VOC loading and relatively high air flow needs.

3.3.1 <u>Vapor Phase Carbon Adsorption Unit</u>

The vapor phase carbon adsorption unit shall be a Bisco Model VF-500, or equivalent. The unit shall be packed with 500 lbs per canister of activated carbon which shall provide approximately 46 days of treatment for the first cycle and 60 days thereafter. The unit shall be an epoxy-lined steel vessel with false floor manifold distributor to provide low pressure drop.

3.3.2 <u>Liquid Phase Carbon Unit</u>

One water purification carbon canister shall be a Bisco Model AF-55 or equivalent. The unit shall be constructed of mild steel, with an epoxy-phenolic internal coat and standard dimensions of 24"(D) X 36"(H). The unit shall have a minimum of two canisters in parallel. The unit shall be capable of processing contaminated groundwater at a rate of 10 GPM, and it shall contain 175 pounds of activated carbon.

PART IV EXECUTION

4.1 <u>SVE DOCUMENTATION REQUIREMENTS</u>

The vendor shall furnish three copies of the following documents with their quotation:

- a. Installation drawings showing all details of construction, dimensions including system footprint, process piping, all mechanical and electrical connections, and demands for proper operation and installation per applicable codes and regulations.
- b. Bill of materials
- c. Bound copies of individualized operations and maintenance procedures, listing of recommended spare parts including current price list.

4.2 WORKMANSHIP

All work shall be performed by tradesmen skilled in their field to produce quality work. All work shall be neatly installed, accessible for maintenance, and complete with all accessories required.

4.3 <u>ACCESSIBILITY</u>

All equipment and accessories shall be installed in such a way that all components requiring access are so located that they can be serviced, reset or recalibrated by service people with normal service tools and equipment.

4.4 PERMITS. CODES AND LAWS

All work shall be done in compliance with NEC, NFPA, OSHA, State Building Codes and other state and local codes and regulations that may apply.

PART V FIELD ASSISTANCE

The vendor shall arrange for a qualified service representative to review the installation of the equipment when complete, and to inspect, operate, test and adjust the equipment to meet performance requirements stated herein. In addition, the vendor shall instruct the Owner on the proper operation and maintenance of the equipment. Vendor shall provide up to two (2) days of installation/field assistance to fine-tune/adjust the system.

PART VI <u>ACCEPTANCE TEST</u>

After the installation is complete and the vendor has performed field assistance services noted above, the Engineer may conduct acceptance tests during average and/or maximum operation of the unit to compare to the performance rate specified above. Should the equipment not attain the specified performance, the Owner at his discretion, may request the Vendor to retrofit the equipment to meet the performance specifications, adjust the purchase price of the equipment, or remove the equipment for a full refund of the purchase price.

PART VII GUARANTEE

7.1 The vendor shall guarantee the equipment for a period of one (1) year from the date of the successful acceptance test as noted above. If any defects in materials, workmanship, corrosion resistance, or operation/performance occur within this period, they shall be corrected by the vendor at no extra cost to the Owner. This excludes defects which are shown to the Engineer's satisfaction to be caused by the Owner's lack of adherence to the recommended operation or maintenance of the equipment per the vendor's manual.

PART VIII DELIVERY AND PROJECT CONTACT

8.1 <u>Delivery Requirements</u>:

The system shall be delivered within four weeks from the date of issuance of a purchase order.

8.2 <u>Project Contact</u>:

Technical questions regarding these specifications shall be addressed to the Engineer as follows:

Mr. Harry S. Wahra
Environmental Engineer
Goldman Environmental Consultants, Inc.
60 Brooks Drive
Braintree, MA 02184
Telephone Number: (617) 356-9140
Fax Number: (617) 356-9147

PART IX PRICING

9.1 F.O.B., Wyandanch, NY

For Appendix D and related Oversized Plates, see Project Manager.