REMEDIAL WORK PLAN

FORMER VOGT MANUFACTURING FACILITY BROWNFIELD CLEANUP PROGRAM (BCP) NYSDEC SITE #C828119 100 FERNWOOD AVENUE ROCHESTER, NEW YORK

> Prepared For: Conifer Development, Inc. 183 East Main Street, 6th Floor

183 East Main Street, 6th Floor Rochester, New York 14604

Prepared By: Day Environmental, Inc.

40 Commercial Street

Rochester, New York 14614

Project No.: 4014R-07

Date: January 2008

New York State Department of Environmental Conservation

Division of Environmental Remediation, Region 8 6274 East Avon-Lima Road. Avon. New York 14414-9519

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June 27, 2008

Ms. Eileen Broderick Conifer Realty Corporation 183 East Main Street, Suite 104 Rochester, New York 14604

Re: Brownfield Cleanup Program - Remedial Work Plan

Former Vogt Manufacturing (C828119)

100 Fernwood Avenue (et. seq.) Rochester (C), Monroe (Co)

Dear Ms. Broderick:

The New York State Department of Environmental Conservation (NYSDEC), in consultation with the New York State Department of Health (NYSDOH) and Monroe County Health Department (MCHD), has completed its review of the January 2008 Remedial Work Plan (RWP) for the subject site. No public comments were received during the 45-day public comment period. Based on our review, the Department is providing the following modifications to the work plan:

- 1. Section 2.2 LNAPL Monitoring and Recovery After a six month period of LNAPL Monitoring and Recovery, an evaluation is to be performed to determine the effectiveness of the four proposed recovery wells in capturing available free product. The need to install additional recovery wells and/or a recovery trench will be evaluated.
- 2. Section 2.4 Monitored Natural Attenuation MNA is to be conducted for a minimum period of five years at the site. After such time, an evaluation is to be performed to determine whether or not MNA will need to continue.
- 3. Section 2.5.3 Periodic Certification The periodic certification is to be provided annually at the same time that the annual MNA Report is submitted. Once MNA reporting is no longer required, an extended periodic certification period can be considered.
- 4. Section 3.0 Final Engineering Report The FER is to be submitted following completion of the active remediation tasks at the site (i.e., implementation of the LNAPL recovery and MNA programs; completion of confirmatory soil and groundwater sampling in the IRM area and supplemental bioremediation treatment, if necessary).

Ms. Eileen Broderick June 27, 2008 Page 2

Pursuant to the Brownfield Cleanup Agreement, Conifer Development may a) choose to accept the above modifications as requested by the Department; b) implement any other Department-approved work plans (not applicable); c) invoke dispute resolution; or d) terminate the BCA. Please notify the Department in writing within 20 days of receipt of this letter as to which of these options is chosen. If accepting the modifications, this modification letter is to be attached to, and become part of, the final approved Remedial Work Plan. A copy of the final approved RWP (with this modification letter) needs to be made available at the project document repository prior to implementation of the fieldwork.

A fact sheet announcing the start of remediation is to be sent out by Conifer to the Site Contact List at least 10 days in advance of the start of fieldwork. Please submit a proposed fact sheet with your response to these modifications.

If you should have any questions regarding this letter or I can be of further assistance, please contact me at (585) 226-5356 or via email at gbmaclea@gw.dec.state.ny.us.

Sincerely,

Gregory B. MacLean, P.E. Environmental Engineer 2

Division of Environmental Remediation

ec: Bart Putzig, P.E., NYSDEC

Bob Knizek, P.E., NYSDEC
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REMEDIAL WORK PLAN

FORMER VOGT MANUFACTURING FACILITY BROWNFIELD CLEANUP PROGRAM (BCP) NYSDEC SITE #C828119 100 FERNWOOD AVENUE ROCHESTER, NEW YORK



Jeffrey A. Danzinger Project Manager

David D. Day, P.E. President

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EXECUTIVE SUMMARY

Day Environmental, Inc. (DAY) prepared this Remedial Work Plan (RWP) for an approximate 8.14-acre property consisting of eleven contiguous parcels addressed as: 100 and 142 Fernwood Avenue; 31, 35 and 41 Rosemary Drive; and 25, 29, 33, 39, 43, 49, and 55 Ilex Place, City of Rochester, County of Monroe, New York (Site). A project locus map (Figure 1) and a Site Plan (Figure 2) are provided at the end of the text of this report. For the purposes of this report, the Site address will generally be referenced as 100 Fernwood Avenue, Rochester, New York. This RWP summarizes the environmental conditions that exist at the Site, and the technical and administrative corrective actions that will be taken to address the environmental conditions.

There are two buildings on the Site. The main building was constructed between 1926 and 1930 and is an approximately 120,000-square foot, one-story concrete block building that has a partial basement. The smaller building is an approximately 3,000-square foot, one-story brick building with a basement that was constructed between 1910 and 1922.

Elmer W. Davis, Inc. currently uses the main building at the Site for the storage of insulation panels; however, it has no full time employees stationed on-site. The main building was originally constructed as Vogt Manufacturing Corporation, which manufactured auto trimmings (e.g., textile trimmings spinning and weaving). Vogt Manufacturing Corporation later became known as Voplex Corporation. The main building was later converted for multi-tenant light industrial/commercial use. Former uses of the main building by tenants include: plastic products manufacturer, tool and die makers, machine shops, painters, printers, graphics companies, and sheet metal contractors. The building was vacant between approximately 2002 and 2004.

The smaller building was originally constructed as, and until recently was used as, a church. However, the smaller building has also been occupied in the past by light industrial/commercial tenants such as Empire Engraving Company (metal cutting allied services) and Phoenix Equipment Co.

The Site is zoned industrial, and is located in a mixed-use urban area. The Site is bounded to the north and west by commercial, industrial and residential properties, and bounded to the south and east by residential properties. The area of the Site is serviced by a public water system.

The Site and surrounding area are generally level. There are no surface water bodies at, or within a 0.5-mile radius of the Site. Surface water appears to flow off the Site via sheet flow toward adjoining streets to the north and to the south (i.e. Rosemary Drive and Fernwood Avenue), into the City of Rochester combined sewer system.

A November 2000 Phase I Environmental Site Assessment (Phase I ESA) report identified the following recognized environmental conditions (RECs) at the Site:

- 1. Abandoned Underground Storage Tanks (USTs)
- 2. Confirmed Local Waste Site/Active New York State Department of Environmental Conservation (NYSDEC) Spill Site on Nearby Property
- 3. Active NYSDEC Spill on Adjoining Property
- 4. Suspect Asbestos-Containing Material [Note: Suspect asbestos-containing material is not addressed as part of this project.]
- 5. Closed NYSDEC Spill on Site
- 6. Transformers/Polychlorinated Biphenyl (PCB) Suspect Equipment
- 7. Historic Use of the Site

In addition to the RECs identified above, the NYSDEC requested that investigative work be included to evaluate whether environmental conditions have been impacted at loading docks equipped with hydraulic lifts. The NYSDEC also requested that a pipe chase in the floor of the main building be further evaluated, and that some limited surface and subsurface evaluation be included on the northern undeveloped portion of the Site.

A Remedial Investigation/Remedial Alternatives Analysis (RI/RAA) Report dated November 2006 as modified by a March 8, 2007 Addendum was prepared by DAY. Tasks performed in 2004 and 2005 as part of the remedial investigation to evaluate or address the RECs identified above included:

- Performing a passive soil gas survey as a screening tool to evaluate the presence of volatile organic compounds (VOCs) at the Site;
- Performing sampling and analysis of various media to evaluate whether PCBs were present at three pad-mounted transformers located east of the main building;
- Performing an evaluation of hydraulic lifts at three loading docks on the main building;
- Performing test pits and magnetic locator work to evaluate the potential presence of abandoned USTs;
- Permanently closing (i.e., removing) four USTs in accordance with applicable regulations;
- Designing and constructing an on-site in-situ bioremediation system within the former tank pit to treat contaminated soils that were displaced/disturbed during the UST closure work;
- Performing post-treatment monitoring to evaluate the effectiveness of the in-situ bioremediation system;
- Evaluating surface soil conditions;
- Evaluating subsurface soil conditions;
- Evaluating groundwater quality conditions and groundwater movement characteristics;
- Performing a vapor intrusion study to evaluate whether VOCs in soil or groundwater were volatilizing and impacting indoor air inside the smaller church building on the Site that is addressed as 142 Fernwood Avenue; and
- Evaluating environmental data for the adjoining former JML Optical, Inc. property located west of the Site.

The locations of test points (e.g., surface soil samples, soil gas points, test pits, test borings, groundwater monitoring wells, etc.) in relation to Site features are shown on Figure 3, Figure 4 and Figure 5.

The findings of the remedial investigation are summarized below:

- The hydraulic loading docks and pad-mounted transformers at the Site do not appear to have adversely impacted environmental conditions at the Site. In addition, evidence of environmental impact was not detected at test boring locations that were completed in proximity to the pipe chase located inside the main building. Therefore, it does not appear that this pipe chase has adversely impacted environmental conditions at the Site.
- Prior to the remedial investigation, a 15,000-gallon UST was removed from the Site. As part of the remedial investigation, one 8,000-gallon UST, two 2,000-gallon USTs, and one 4,000-gallon UST were removed from the Site. These five USTs were located in the same general area north of the northwest corner of the main building (refer to Figure 3 and Figure 4).

- A primary area of soil and groundwater contamination, including the presence of a relatively thin layer (i.e., 0.37 foot or less) of light non-aqueous phase liquid (LNAPL) that is more limited in extent, was detected in proximity to the former UST locations near the northwest corner of the main building (refer to Figure 6). This contamination generally consists of petroleum products and plasticizers that historically leaked from the former USTs. Based on field screening, analytical laboratory test results, and groundwater monitoring, it appears that this impact has migrated radially outward from the former UST area, including beneath the northwest corner of the main building. In addition, the length of the petroleum/plasticizer plume is estimated to be about 60 feet away from the former UST locations. Based on a review of Site data and environmental reports for the adjoining former JML Optical, Inc. property to the west, petroleum and plasticizer contamination attributable to the former UST locations at the Site appears to have also migrated from the Site via groundwater onto an estimated 1,375-square foot area of the adjoining former JML Optical, Inc. property.
- As an interim remedial measure (IRM), petroleum and plasticizer contaminated soils that were displaced during the UST removal work were amended with bioremediation products and placed back into the tank pit excavation as part of an in-situ bioremediation system (refer to Figure 7 and Figure 8). The analysis of post-treatment soil and groundwater samples indicate the in-situ bioremediation system is working, and contaminants have been reduced by approximately 40% (on average). The results of the post-treatment sampling and analytical laboratory testing indicated that these contaminants were still present in soil and groundwater at concentrations exceeding Standards, Criteria and Guidance (SCG) values.
- Two of four surface soil samples collected from the northern undeveloped portion of the Site contained some polyaromatic hydrocarbon (PAH) semi-volatile organic compounds (SVOCs) above December 14, 2006 NYSDEC Part 375 (Environmental Restoration Programs) Track 2 Soil Cleanup Objectives (SCOs) for Restricted Residential Use. However, the concentrations of these SVOCs are comparable to other projects in the City of Rochester where surface soil data has been collected. As such, the NYSDEC concurs that the limited exceedances of the Restricted Residential Use SCOs in surface soil at the Site are attributable to the local geology or urban setting of the Site and are not significant. As such, further actions in relation to surface soil at the Site do not appear warranted.
- Chlorinated VOCs were detected in groundwater samples at some of the monitoring well locations. An on-site source of chlorinated VOCs that could result in contamination of the groundwater was not found during the soil and groundwater studies performed as part of the remedial investigation. It is possible that the chlorinated VOCs are attributable to an off-site source(s) that has resulted in an area-wide groundwater condition. A review of environmental reports indicates a sump and a former degreaser area at the adjoining former JML Optical, Inc. property to the west, and also a nearby NYSDEC Inactive Hazardous Waste Disposal Site located southwest of the Site, could potentially be sources of the chlorinated VOCs that are present at the Site.
- A subsurface soil sample collected from a depth interval of 0-4' at test boring TB-4 contained three PAH SVOCs that exceeded Track 2 SCOs for Restricted Residential Use. This sample was collected beneath the floor of the main building and contained fill material that consisted of reworked soil with some cinders. The PAH SVOCs would presumably be limited in extent to the fill material, and can be a common component of cinders.
- The results of a passive soil gas survey indicate that the SVOC naphthalene and various VOCs identified as aliphatic hydrocarbons and chlorinated VOCs (e.g., tetrachloroethene) were detected

at some soil gas points, many of which are located inside the main building. Subsequent soil and groundwater testing suggest the source of many of the detected constituents in the passive soil gas survey are most likely related to these constituents being present in flooring materials of the main building, and not from impacts to underlying soil or groundwater.

- The results of the vapor intrusion evaluation suggest that chlorinated VOCs detected in nearby groundwater monitoring well MW-1 is not impacting indoor air quality inside the small church building (142 Fernwood Avenue). However, the highest concentration of the chlorinated VOCs (i.e., 51 ug/m³ of trichloroethylene) was detected in an indoor air sample collected from the first floor of this building. Possible sources of the VOCs detected on the first floor during this sampling event may include historic spills to flooring, etc. on the first floor of the building, or VOC residues on former church patron clothing, shoes, etc. In lieu of re-testing to confirm the absence or presence of this condition, Conifer Development, Inc. decided to remove the tenant and leave the building vacant with the understanding that further testing or mitigation would be required in the future before this building can be re-occupied.
- Based on groundwater table elevation data for Site wells, groundwater generally flows radially outward from an unpaved location north of the main building where the five USTs were removed and the in-situ bioremediation system was installed as an IRM. A copy of a potentiometric groundwater contour map for November 14, 2005 that depicts this groundwater flow pattern is included as Figure 9.

Remedial Alternatives Analysis

The November 2006 RI/RAA report identified remedial action objectives (RAOs), contaminants of interest, remediation criteria, and general response actions. In regard to these criteria, four remedial alternatives were developed and evaluated. These alternatives are summarized below:

Alternative #1 No Further Action

Alternative #2 LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls

Alternative #3 Limited In-Situ Remediation, LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls

Alternative #4 Full Excavation, LNAPL Recovery, In-Situ Remediation and Groundwater Monitoring for 5 Years

A detailed evaluation of the four remedial alternatives was performed, and implementation of Alternative #2 (LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls) was recommended in the November 2006 RI/RAA report.

As part of Alternative #2, LNAPL recovery in proximity to existing monitoring well MW-6, and natural attenuation, would be used to remediate and control residual groundwater contamination and reduce risk to exposure. Groundwater monitoring would be implemented to ensure that natural attenuation is adequately controlling and remediating the contamination in the groundwater. Institutional controls and engineering controls would be implemented to protect against exposure to contamination in soil and groundwater. The institutional controls would include a site management plan (SMP), an environmental easement, and periodic certifications. The institutional controls will also require consideration of vapor intrusion (i.e., implementation of engineering controls) for existing and new buildings, if warranted based on construction specifications, etc.

The estimated present worth cost to implement the remedy (Alternative #2) is \$141,703.00. The cost to construct the remedy is estimated to be \$75,240.00, and the estimated average annual costs for five years is about \$13,293.00. The Alternative #2 Opinion of Probable Remedial Costs are further detailed on a table included in Appendix A.

Further details regarding the elements of the proposed remedy (Alternative #2) are as follows:

- 1. <u>LNAPL Monitoring and Recovery</u>: Four new wells will be installed in proximity to existing wells MW-6 and MW-10 (refer to Figure 6). LNAPL will then be monitored/recovered from select existing wells and the four new wells for a period of up to two years. The LNAPL will be further characterized (if necessary) and disposed/recycled off-site in accordance with applicable regulations.
- 2. Confirmatory Sampling and Analysis from In-Situ Bioremediation System Area: At least one round of confirmatory soil and groundwater samples will be collected from the existing in-situ bioremediation system IRM area that is associated with the former UST locations. Depending upon the results of this sampling, additional injection of bioremediation compounds via existing in-situ bioremediation piping in the IRM area and follow-up confirmatory sampling will be considered to the extent required by the NYSDEC (refer to Site Management Plan section, listed as item 4 below).
- 3. <u>Monitored Natural Attenuation:</u> A groundwater monitoring program will be implemented to monitor the effectiveness of natural attenuation. Up to eight existing or new groundwater monitoring wells would be utilized during the monitoring program. Cumulative groundwater data will be evaluated as it is generated to assess the following conditions and trends:
 - Chemical mass, concentrations and toxicity over time;
 - Specific types of natural attenuation; and
 - Evaluate the data through software programs that are useful in evaluating natural attenuation, such as the BIOSCREEN and possibly also the Mass Flux ToolKit that are free software programs developed by GSI Environmental, Inc.

During each monitoring event, groundwater samples will be collected from up to eight wells, and the samples will be analyzed for: VOCs; SVOCs; and, various natural attenuation parameters such as nitrate, iron (II), manganese, sulfate, methane, and chloride. Water quality measurements for dissolved oxygen, oxidation-reduction potential, pH, temperature, conductivity, and turbidity will also be collected.

Groundwater monitoring will be conducted on a bi-annual basis for the first two years, and on an annual basis thereafter. It is currently anticipated that the natural attenuation monitoring will be conducted for a period of up to five years.

4. <u>Site Management Plan:</u> Development of a SMP, which will include the following institutional controls: (a) management to restrict excavation below existing surface soil, pavement, or buildings. [Excavated soil or fill material would be tested, properly handled to protect the health and safety of workers and the nearby community, and would be properly managed in a manner acceptable to the NYSDEC]; (b) continued evaluation of the potential for vapor intrusion for any existing or new buildings developed on the Site, including provision for mitigation of any impacts identified; (c) identification of any use restrictions on the Site; and, (d) provisions for the continued proper operation and maintenance of the components of the remedy.

Subsequent to review of available LNAPL monitoring and recovery, the analytical test results for confirmatory soil and groundwater samples from the existing bioremediation system, and the analytical test results for the first round of monitored natural attenuation samples, a decision will be made with input from the NYSDEC regarding whether additional remediation products should be applied within and/or around the existing in-situ bioremediation system. If additional application of remediation products is deemed warranted, the SMP will outline the scope of the application, and any subsequent confirmatory monitoring, sampling and analysis. It is anticipated that the same, or similar types of bioremediation products that were previously used, would be used during the additional application.

- 5. Environmental Easement: Imposition of an institutional control in the form of an environmental easement that will require: (a) limiting the use and development of the property to restricted residential use, which would also permit commercial or industrial uses; (b) compliance with the approved site management plan; (c) restricting the use of groundwater as a source of potable or process water, without necessary water quality treatment as determined by New York State Department of Health (NYSDOH); and (d) the property owner to complete and submit to the NYSDEC a periodic certification of institutional controls.
- 6. Periodic Certification: The property owner will provide certification of institutional controls and engineering controls (if warranted), prepared and submitted by a professional engineer or such other expert acceptable to the NYSDEC, until the NYSDEC notifies the property owner in writing that this certification is no longer needed. It is anticipated that the periodic certification will be provided every three years. This submittal will: contain certification that the institutional controls and engineering controls (if warranted) put in place are still in place and are either unchanged from the previous certification or are complaint with NYSDEC-approved modifications; allow the NYSDEC access to the Site; and, state that nothing has occurred that would impair the ability of the control to protect public health or the environment, or constitute a violation of failure to comply with the site management plan unless otherwise approved by the NYSDEC.

Conifer Development, Inc. is proposing Alternative #2 (LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls) as the remedy for this Site. This proposed remedy is based on the results of the November 2006 RI/RAA report as modified by the March 8, 2007 addendum. Alternative #2 is being proposed because, as described below, it satisfies the threshold criteria described in Section 8.0 of the November 2006 RI/RAA report given the fact that the in-situ bioremediation system has already been installed as an IRM to address the on-site source area of contamination (i.e., the former UST area). Alternatives #2, #3 and #4 would achieve varying degrees of remediation goals for the site and include addressing the LNAPL that creates the most significant threat to public health and the environment, greatly reducing the source of contamination to groundwater, and creating the conditions needed to restore groundwater quality to the extent practicable.

The level of risk associated with short-term impacts is lowest for Alternatives #1 and #2, minimal for Alternative #3, and highest for Alternative #4. Excavation of contaminated material is the factor that increases the risks associated with short-term impacts for Alternative #4. The short-term impacts for these alternatives can be controlled by implementing a Health and Safety Plan. The time needed to achieve the remediation goals would be longest for Alternative #2, moderate for Alternative #3, and shortest for Alternative #4.

Achieving long-term effectiveness is best accomplished by excavation and removal of the contaminated overburden soils. Alternative #4 contains excavation and removal components.

Alternatives #1, #2, and #3 do not contain excavation and removal components; however, Alternatives #2 and #3 will require institutional controls, engineering controls, and long-term monitoring. Alternative #4 would result in the constituents of concern remaining on-site for less time than Alternatives #2 and #3. Alternatives #2, #3, and #4 will reduce long-term risk associated with the contamination though aggressive remediation and/or institutional controls and engineering controls. Alternative #1 would not reduce risk.

Alternative #2 is favorable in that it is readily implementable in relation to the future restricted residential, commercial or industrial use of the Site. Alternatives #1 and #3 are also implementable. Alternative #4 is not readily implementable. Spatial requirements for Alternatives #2 and #3 are considerably less than the spatial requirements that would be required for Alternative #4. There are no spatial requirements for Alternative #1.

Physical remediation components of Alternatives #3 and #4 would result in the largest reductions of contaminant toxicity, mobility or volume. Alternative #2 that includes a contingency for additional in-situ remediation applications relies more heavily on natural attenuation and other factors such as advection, dispersion, sorption, diffusion, etc., but also results in reductions of contaminant toxicity, mobility or volume. Alternative #1 would result in the least amount of contaminant toxicity, mobility or volume. However, it should be noted that contaminant toxicity, mobility or volume at the Site has already been greatly reduced as a result of removing the USTs and their contents, and installing the in-situ bioremediation system in this area as an IRM.

The cost of the alternatives varies significantly. There is no cost associated with Alternative #1; however, this alternative is not a permanent remedy. Alternative #2 costs less than Alternatives #3 and #4. Alternative #4 costs are substantially high in comparison to the costs of the other alternatives, and are considered excessive in relation to the benefits gained. Costs associated with Alternatives #2 and #3 are considered reasonable in terms of the benefits gained.

Alternative #1 is not protective of human health or the environment and does not address RAOs for this Site. Alternatives #2, #3, and #4 are protective of human health and the environment, and risks associated with potential human health exposure pathways would be eliminated or adequately controlled. Remedial action objectives are generally addressed by Alternatives #2, #3, and #4 in relation to protection of public health and the environment.

Alternative #1 provides no compliance with SCG values. Alternatives #2 and #3 provide varying levels of compliance with SCG values. Alternative #4 provides complete compliance with SCG values and would remediate the Site to pre-release conditions, but at an exorbitant cost in relation to the benefits gained. Alternatives #2, #3, and #4 provide adequate groundwater monitoring to evaluate compliance trends in relation to chemical-specific SCG values.

Alternatives #2, #3 and #4 would be acceptable in relation to the future restricted residential, commercial or industrial use of the Site. Alternative #1 would likely not be acceptable in relation to the future restricted residential, commercial or industrial use of the Site.

In conclusion, Conifer Development, Inc. is recommending Alternative #2 (LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls) as the remedy for the Site. Alternative #2 satisfies the threshold criteria, and the in-situ bioremediation system has already been installed to address source-area contamination.

1.0 INTRODUCTION

DAY prepared this RWP for an approximate 8.14-acre property (Site) consisting of eleven contiguous parcels addressed as 100 and 142 Fernwood Avenue; 31, 35 and 41 Rosemary Drive; and 25, 29, 33, 39, 43, 49, and 55 Ilex Place, City of Rochester, County of Monroe, New York (Tax account #s 106.27-1-5; 91.83-3-19; 91.83-3-20; 91.83-3-21; 106.27-1-87; 106.27-1-88; 106.27-1-89; 106.27-1-90; 106.27-1-91; 106.27-1-92; and 106.27-1-93). A project locus map (Figure 1) and a Site Plan (Figure 2) are provided at the end of the text of this report. For the purposes of this report, the Site address will generally be referenced as 100 Fernwood Avenue, Rochester, New York.

This RWP was developed in accordance with the requirements of Brownfield Site Cleanup Agreement Index #B8-06660-04-05 between the NYSDEC and Conifer Development, Inc. (identified as BCP Site #C828119) and from guidance provided in Section 4.0 of the NYSDEC document titled "Draft Brownfield Cleanup Program Guide" dated May 2004. The RWP summarizes the environmental conditions that exist at the Site, and the technical and administrative corrective actions that will be taken to address the environmental conditions.

1.1 Background

There are two buildings on the Site. The main building was constructed between 1926 and 1930 and is an approximately 120,000-square foot, one-story concrete block building that has a partial basement. The smaller building is an approximately 3,000-square foot, one-story brick building with a basement that was constructed between 1910 and 1922.

Elmer W. Davis, Inc. currently uses the main building at the Site for the storage of insulation panels; however, it has no full time employees stationed on-site. The main building was originally constructed as Vogt Manufacturing Corporation, which manufactured auto trimmings (e.g., textile trimmings spinning and weaving). Vogt Manufacturing Corporation later became known as Voplex Corporation. The main building was later converted for multi-tenant light industrial/commercial use. Former uses of the main building by tenants include: plastic products manufacturer, tool and die makers, machine shops, painters, printers, graphics companies, and sheet metal contractors. The building was vacant between approximately 2002 and 2004.

The smaller building was originally constructed as, and until recently was used as, a church. However, the smaller building has also been occupied in the past by light industrial/commercial tenants such as Empire Engraving Company (metal cutting allied services) and Phoenix Equipment Co.

The Site is zoned industrial, and is located in a mixed-use urban area. The Site and surrounding area are serviced by a public water system. The Site is bounded to the north and west by commercial, industrial and residential properties, and bounded to the south and east by residential properties.

The Monroe County Department of Public Health (MCDPH) has no records of public or private drinking water wells or process water wells within a 0.25-mile radius of the Site. A review of the document titled "Ground Water Resources of Monroe County, New York", 1935 by R.M. Leggette, L.O. Gould and B.H. Dollen identifies an "industrial" groundwater well (#2131) on the Site. This well is listed for Vogt Manufacturing Company on Fernwood Ave, and is identified as being drilled 116 feet deep and set in the Reynales Limestone (i.e., water-bearing bedrock unit). The depth to groundwater in this well is listed as 20 feet below the ground surface. The depth to rock is listed as 12 feet below the ground surface. [Note: The condition and location of this well on the Site is unknown].

The Site and surrounding area are generally level. There are no surface water bodies at, or within a 0.5-mile radius of the Site. Surface water appears to flow off the Site via sheet flow toward adjoining streets to the north and to the south (i.e. Rosemary Drive and Fernwood Avenue), into the City of Rochester combined sewer system. Based on a preliminary review of a 1980 Generalized Groundwater Contour Map, Rochester East Quadrangle, groundwater in the area of the Site appears to flow to the east-northeast toward Irondequoit Bay, which is located approximately three miles from the Site. This flow direction may be modified locally due to buried utilities, seasonal conditions, or other factors. Based on groundwater table elevation data for Site wells, groundwater on the Site itself generally flows radially outward from an unpaved location north of the main building where the five USTs were removed and an in-situ bioremediation system was installed as an IRM. A copy of a potentiometric groundwater contour map for November 14, 2005 that depicts this groundwater flow pattern is included as Figure 9.

1.2 Previous Environmental Studies

DAY completed a Phase I ESA report dated November 15, 2000 for the Site in general accordance with American Society for Testing and Materials (ASTM) Practice E1527-00. Information obtained from the Phase I ESA indicates that each of the buildings on the Site is currently heated with steam boiler systems fueled with natural gas. The main building on the Site was heated with coal and oil in the past. In addition, the Site buildings were connected to the public sewer system at the time they were constructed. The Site buildings are also connected to the public water system.

The November 2000 Phase I ESA report identified the following RECs at the Site:

- 1. Abandoned USTs
- 2. Confirmed Local Waste Site/Active NYSDEC Spill Site on Nearby Property
- 3. Active NYSDEC Spill on Adjoining Property
- 4. Suspect Asbestos-Containing Material [Note: Suspect asbestos-containing material is not addressed as part of this work plan.]
- 5. Closed NYSDEC Spill on Site
- 6. Transformers/PCB Suspect Equipment
- 7. Historic Use of the Site

In addition to the RECs identified above, the NYSDEC requested that investigative work be included to evaluate whether environmental conditions had been impacted at loading docks equipped with hydraulic lifts. The NYSDEC also requested that a pipe chase in the floor of the main building be further evaluated and that some limited surface and subsurface evaluation be included on the northern undeveloped portion of the Site.

A RI/RAA Report dated November 2006, as modified by an addendum dated March 8, 2007, was prepared by DAY. The primary objective of the remedial investigation was to perform environmental work at the Site in accordance with the requirements of the Brownfield Cleanup Program (BCP) to evaluate the nature and extent of contamination at the Site. Other objectives included: performing an exposure assessment; confirming and/or further delineating contamination in areas identified as RECs in the Phase I ESA; evaluating fate and transport of contaminants; identifying remedial alternatives; performing a detailed analysis of selected remedial alternatives; and selecting a remedial alternative.

Specific tasks performed as part of the remedial investigation to evaluate or address the RECs identified above included:

- Performing a passive soil gas survey as a screening tool to evaluate the presence of VOCs at the Site;
- Performing sampling and analysis of various media to evaluate whether PCBs were present at three pad-mounted transformers located east of the main building;
- Performing an evaluation of hydraulic lifts at three loading docks on the main building;
- Performing test pits and magnetic locator work to evaluate the potential presence of abandoned USTs;
- Permanently closing (i.e., removing) four USTs in accordance with applicable regulations;
- Designing and constructing an on-site in-situ bioremediation system within the former tank pit as an IRM to treat contaminated soils that were displaced/disturbed during the UST closure work;
- Performing post-treatment monitoring to evaluate the effectiveness of the in-situ bioremediation system;
- Evaluating surface soil conditions;
- Evaluating subsurface soil conditions;
- Evaluating groundwater quality conditions and groundwater movement characteristics;
- Performing a vapor intrusion study to evaluate whether VOCs in soil or groundwater were volatilizing and impacting indoor air inside the smaller church building on the Site that is addressed as 142 Fernwood Avenue; and
- Evaluating environmental data for the adjoining former JML Optical, Inc. property located west of the Site.

Figure 3, Figure 4, and Figure 5 include the locations of soil gas survey points, the transformers area that was sampled, surface soil samples, test borings, groundwater monitoring wells and test pits that were completed as part of the above studies.

Physical Characteristics of Site

Based on the findings of the remedial investigation, heterogeneous fill material generally consisting of reworked soil (e.g., silt, sand, gravel, and clay) and/or cinders with lesser amounts of brick, concrete, asphalt, organics, and wood is present over many locations of the Site from the ground surface to depths ranging between approximately 0.5 feet and 6.6 feet. The average thickness of fill material is 2.3 feet, and the thinner layers of fill encountered appear to have been used for sub-base material beneath buildings or paved parking lots. At most test locations, the indigenous soil predominantly consists of varying grades of silts and sands, with lesser amounts of gravel and clay. As measured during the remedial investigation, groundwater generally flows radially outward away from the former UST tank pit/current in-situ bioremediation system, located within the former UST tank pit north of the northwest corner of the main building. The RI/RAA report indicates that bedrock underlying the overburden deposits in proximity to the Site consists of Rochester Shale belonging to the Clinton Group, Upper Silurian Period, Paleozoic Era.

Nature and Extent of Contamination

The nature and extent of contamination at this Site is summarized below:

<u>Underground Storage Tanks:</u> One 15,000-gallon UST, one 8,000-gallon UST, two 2,000-gallon USTs, and one 4,000-gallon UST were permanently closed (i.e., removed). Contaminated soil removed from the tank pit excavation was amended with bioremediation agents and placed back into the tank pit excavation as part of an IRM in-situ bioremediation system. Laboratory analysis of post-treatment soil and groundwater samples collected in 2005 indicates the in-situ bioremediation system is working; however, VOC and/or SVOC concentrations in some of these soil and groundwater samples exceeded SCG values at some locations.

<u>Passive Soil Gas Survey:</u> The results of the passive soil gas survey indicate that the SVOC naphthalene and various VOCs identified as aliphatic hydrocarbons and chlorinated VOCs (e.g., tetrachloroethene) were detected at some soil gas points, many of which are located inside the main building. Subsequent soil and groundwater testing suggest the source of detected constituents in the passive soil gas survey are most likely related to these constituents being present in flooring materials of the main building, and not from impacts to underlying soil or groundwater.

<u>Transformers</u>: Oil samples from transformers, wipe samples from the exterior of transformers, and concrete samples from the pad that supports the transformers did not contain concentrations of PCBs exceeding SCG values.

Hydraulic Loading Docks: Although some minor leakage was observed on the concrete pads beneath the hydraulic cylinders associated with the three hydraulic loading docks located within the main building, this leakage did not appear to have the potential to impact soil or groundwater. A sample of the hydraulic oil from the lift located at the southwest corner of the main building contained 2,800,000 mg/kg or parts per million (ppm) of total petroleum hydrocarbon (TPH) that best matched chromatograms for weathered diesel and motor oil (i.e., concentration detected implies pure product).

Surface Soil: Two of four surface soil samples collected from the northern undeveloped portion of the Site contained some PAH SVOCs above December 14, 2006 NYSDEC Part 375 (Environmental Restoration Programs) Track 2 SCOs for Restricted Residential Use. However, the concentrations of these SVOCs are comparable to other projects in the City of Rochester where surface soil data has been collected. As such, the NYSDEC concurs that the limited exceedances of the Restricted Residential Use SCOs in surface soil at the Site are attributable to the local geology or urban setting of the Site and are not significant.

<u>Subsurface Soil:</u> In general, contaminants attributable to petroleum products and plasticizers were detected in subsurface soil samples in proximity to the location where five USTs were removed (i.e., in proximity to the northwest corner of the main building) and also at other miscellaneous areas of the Site. The analytical laboratory test results of field samples are summarized below:

- Target VOCs were detected in 9 of 20 subsurface soil samples, but at concentrations below available NYSDEC Track 2 SCOs for Restricted Residential Use. Most of the VOCs detected appear to be related to petroleum products, and solvents to a lesser degree.
- TPH was detected in 2 of 11 subsurface soil samples, and the TPH detected did not resemble the TPH fingerprint of the oil sample collected from one of the hydraulic loading dock lift cylinders.
- Target SVOCs were detected in 11 of the 19 subsurface soil samples, and most are related to petroleum products or plasticizers. Only the concentrations of benzo(a)anthracene, benzo(b)fluoranthene and benzo(a)pyrene detected in Sample 033 from TB-4(0-4') that was collected immediately beneath the floor of the main building (and which partly consisted of fill that contained cinders) exceeded their available NYSDEC Track 2 SCOs for Restricted

Residential Use. The concentrations of the SVOCs in Sample 033 are comparable to other projects in the City of Rochester where surface soil data has been collected. As such, the presence of these SVOCs appears attributable to the local geology or urban setting of the Site and is not significant. Concentrations of the SVOC bis(2-ethylhexyl)phthalate detected in 6 of the 19 subsurface soil samples were more than one or two orders of magnitude higher than other SVOCs detected in these samples. The higher concentrations of bis(2-ethylhexyl)phthalate were detected in samples from the proximity to the former UST area (i.e., up to 350 ppm at a test location outside the IRM in-situ bioremediation system, and up to 1,400 ppm on a November 2005 post-treatment soil sample collected from within the in-situ bioremediation system). This is the area of the Site where an 8,000-gallon UST containing high concentrations of bis(2-ethylhexyl)phthalate, as well as other USTs containing petroleum products, had been removed. NYSDEC Track 2 SCOs are not available for the SVOC bis(2-ethylhexyl)phthalate.

- Target analyte list (TAL) metals and cyanide test results for subsurface soil samples did not exceed available NYSDEC Track 2 SCOs for Restricted Residential Use. Naturally occurring concentrations of metals in soil at the Site may be contributing to the detected concentrations of metals in the subsurface soil samples (e.g., calcium, iron, magnesium, zinc).
- PCBs were not detected at concentrations above the reported analytical laboratory detection limit in the six subsurface soil samples that were tested.
- Pesticides were detected in 3 of 10 subsurface soil samples that were tested, but at concentrations below their respective NYSDEC Track 2 SCOs for Restricted Residential Use.
- Formaldehyde was detected in 2 of 4 subsurface soil samples at concentrations of 0.27 ppm and 0.43 ppm. NYSDEC SCOs are not available for formaldehyde.

<u>Groundwater:</u> In general, contaminants attributable to solvents were detected in groundwater samples from wells on the southern portion of the Site. In addition, contaminants attributable to petroleum products and plasticizers were detected in groundwater samples in proximity to the location where five USTs were removed (i.e., in proximity to the northwest corner of the main building). The analytical laboratory test results of field samples are summarized below:

Target VOCs and tentatively identified compounds (TICs) were detected in one or more groundwater sample from wells MW-1, MW-2, MW-3, MW-4, MW-5, MW-8 and MW-10. Target VOCs and TICs were also detected in one or more groundwater sample from monitoring wells MWIRM-1 through MWIRM-3. Types of VOCs detected in most wells are generally attributable to solvents, and the VOCs detected in a groundwater sample from well MW-10 are generally attributable to petroleum products. The measured concentrations of the VOCs 1,1,1trichlroethane, trichloroethene, benzene, xylene, toluene, and tetrahydrofuran exceeded NYSDEC Technical and Operational Guidance Series (TOGS 1.1.1) groundwater standards and guidance values in one or more groundwater sample. Based on the concentrations detected [i.e., between 2 and 15 ug/l or parts per billion (ppb)], and since an on-site source was not documented at the Site for these types of VOCs, it is possible that the chlorinated VOCs detected in the groundwater samples at the Site may represent area-wide contamination in groundwater due to an off-site source(s). The petroleum-related VOCs generally appear attributable to the on-site USTs that were formerly located north of the northwest corner of the main building. tetrahydrofuran was detected in groundwater samples collected from wells within the footprint of the former tank excavation after construction of the in-situ bioremediation system. Tetrahydrofuran was not detected in groundwater samples from any of the other wells on the Site; thus, it does not appear that tetrahydrofuran has migrated away from the apparent source area.

- Target SVOCs and TICs were detected in one or more groundwater sample from wells MW-1, MW-2, MW-3, MW-4, MW-5, MW-7, MW-8, MW-9, MW-10 and MW-11. Target SVOCs detected in one or more of these samples included: naphthalene; phenol; caprolactam; 2-methylnaphthalene; 1,1-biphenyl; acenaphthene; dibenzofuran; fluorene; phenanthrene; anthracene; carbazole; and bis(2-ethylhexyl)phthalate. Only the concentration of bis(2-ethylhexyl)phthalate detected in the groundwater samples from well MW-5, and the concentrations of naphthalene and bis(2-ethylhexyl)phthalate detected in the groundwater sample from well MW-10, exceeded their NYSDEC TOGS 1.1.1 groundwater standards or guidance values. [Note: Wells MW-5 and MW-10 are in proximity to the former UST locations where the in-situ bioremediation system was constructed]
- TAL metals detected in one or more groundwater sample included: aluminum; antimony; arsenic; barium; beryllium; calcium; chromium; cobalt; copper; iron; lead; magnesium; manganese; nickel; potassium; selenium; sodium; and vanadium. NYSDEC TOGS 1.1.1 groundwater standards and guidance values were exceeded in one or more groundwater sample for: antimony; iron; magnesium; manganese; and, sodium. Based on local geology, naturally occurring background conditions may be contributing to the detected concentrations of most of these metals.
- Cyanide concentrations detected in two groundwater samples did not exceed its NYSDEC TOGS
 1.1.1 groundwater standard or guidance value.
- PCBs, pesticides and TPH were not detected in groundwater samples at concentrations above reported analytical laboratory detection limits.
- Formaldehyde was only detected in groundwater samples from wells MW-1, MW-4 and MW-8 at concentrations ranging between 5.2 ug/l (ppb) and 5.6 ug/l (ppb). There is no TOGS 1.1.1 groundwater standard or guidance value for formaldehyde.

LNAPL: During this remedial investigation, measurable LNAPL was detected at groundwater monitoring well locations MW-6 and MW-10. Between approximately 0.21 and 0.37 foot of LNAPL was observed on top of the groundwater in well MW-6, and approximately 0.1 foot of LNAPL was observed on top of the groundwater in well MW-10. The LNAPL detected in these wells appeared dark brown. A June 2005 LNAPL sample collected from monitoring well MW-6 primarily contained: VOCs related to petroleum products, and SVOCs related to petroleum products and plasticizers. The LNAPL sample also contained some TAL metals and pesticides. TPH test results best matched a chromatogram for #2 fuel oil.

Petroleum/Plasticizer Plume: Based on field findings and analytical laboratory testing of soil, groundwater and LNAPL samples, the length of the petroleum/plasticizer plume located in proximity to the former UST locations at the northwest corner of the main building is estimated to be about 60 feet. The plume likely extended radially away from the former UST locations, and a zone of more extensive contamination, including LNAPL, was documented to trend south/southwest of the former UST locations, which includes migration beneath the northeast portion of the main building. Based on a review of Site data and environmental reports for the adjoining former JML Optical, Inc. property to the west, petroleum and plasticizer contamination attributable to the former UST locations at the Site appears to have also migrated via groundwater onto the adjoining former JML Optical, Inc. property. This contamination is estimated to be present under an approximate 55-foot by 25-foot area (i.e., estimated 1,375 square-foot area) of the adjoining former JML Optical, Inc. property (refer to Figure 6).

Vapor Intrusion Evaluation: The results of the vapor intrusion evaluation suggest that chlorinated VOCs detected in nearby groundwater monitoring well MW-1 are not impacting indoor air quality inside the small church building (142 Fernwood Avenue). However, the highest concentration of chlorinated VOCs (i.e., 51 ug/m³ of trichloroethylene) was detected in an indoor air sample collected from the first floor of this building. Possible sources of the VOCs detected on the first floor during this sampling event may include historic spills to flooring, etc. on the first floor of the building, or VOC residues on former church patron clothing, shoes, etc. In lieu of re-testing to confirm the absence or presence of this condition, Conifer Development, Inc. decided to remove the tenant and leave the building vacant with the understanding that further testing or mitigation would be required in the future before this building can be re-occupied.

<u>Soil Gas Evaluation:</u> The results of a passive soil gas survey indicate that the SVOC naphthalene and various VOCs identified as aliphatic hydrocarbons and chlorinated VOCs (e.g., tetrachloroethene) were detected at some soil gas points, many of which are located inside the main building. Subsequent soil and groundwater testing suggest the source of many of the detected constituents in the passive soil gas survey are most likely related to these constituents being present in flooring materials of the main building, and not from impacts to underlying soil or groundwater.

<u>DNAPL Monitoring:</u> Evidence of dense non-aqueous phase liquid (DNAPL) was not detected at test boring, test pit, or monitoring well locations during this study.

Contaminant Fate and Transport

Potential routes of migration identified for this Site include:

- SVOCs in surface soil migrating overland during precipitation events;
- SVOCs absorbed onto surface soil particles and becoming windborne;
- VOCs and SVOCs in soil leaching and impacting groundwater through precipitation or contact with groundwater;
- VOCs, SVOCs and possibly some metals migrating in a dissolved groundwater plume;
- VOCs, SVOCs and possibly pesticides in LNAPL migrating on top of the water table;
- VOCs migrating as a vapor in the unsaturated zone;
- VOC volatilization from groundwater or soil to indoor air inside buildings [Note: the vapor intrusion evaluation does not suggest this is occurring at the 142 Fernwood Avenue building that is located on the Site]; and
- Indirect migration pathways may include: volatilization to air, transportation on construction equipment/workers, windborne processes, etc., if the impacted soil were to be disturbed in the future.

The contamination at the Site is identified as generally consisting of organic constituents (VOCs and SVOCs), and also various metals. The persistence of these constituents is further discussed below.

Organic Constituents

The VOCs and SVOCs detected at the Site are generally associated with weathered petroleum products and/or plasticizers. Much of the non-target VOCs and SVOCs detected in soil and groundwater samples may reflect biodegradation products of the petroleum and/or plasticizer contamination or other non-target compounds typically associated with these types of products. Petroleum-type VOCs detected in soil and groundwater may be attributable to products such as diesel fuel or heating oil. The SVOCs detected in the soil and groundwater are generally

considered PAHs and phthalates. The VOCs and SVOCs encountered at the Site biodegrade aerobically and anaerobically. The constituents commonly detected in soil or groundwater will generally biodegrade faster under aerobic conditions when compared to biodegradation rates under anaerobic conditions. In addition, chlorinated VOCs were detected in groundwater samples, which suggest an area-wide condition that is likely attributable to an off-site source(s). These chlorinated VOCs are more persistent in the environmental than petroleum-type VOCs, and degrade aerobically and/or anaerobically.

In addition to biodegradation, VOC and SVOC concentrations in the groundwater would presumably decrease as the distance from the source area is increased due to processes such as advection, dispersion, sorption, diffusion, etc.

<u>Inorganics</u>

Various metals were detected in samples of surface soil, subsurface soil and groundwater. Some of the metals detected may be associated with contamination from past uses of the Site, and other metals may be associated with naturally occurring concentrations of metals in soil or groundwater for the area of the Site. Metals can change form (e.g., Fe⁺², Fe⁺³), but are persistent in the environment and do not degrade.

No metals were detected in soil samples at concentrations exceeding Track 2 SCOs for Restricted Residential Use. Only the metals antimony, iron, magnesium, manganese and sodium were detected in one or more groundwater sample at concentrations exceeding TOGS 1.1.1 groundwater standards or guidance values.

Processes such as advection, dispersion, sorption, diffusion, etc, can result in decreases in metals concentrations dissolved in groundwater as the distance away from their source is increased.

The petroleum and plasticizers within the soil and groundwater at the Site are detected at highest concentrations in proximity to the former UST locations near the northwest corner of the main building on the Site. Based on field screening, analytical laboratory test results, and groundwater monitoring, it appears that this contamination has migrated radially outward from the former UST area, including beneath the northwest corner of the main building. In addition, the length of the petroleum/plasticizer plume is estimated to be about 60 feet away from the former UST locations. Based on a review of Site data and environmental reports for the adjoining former JML Optical, Inc. property, petroleum and plasticizer contamination attributable to the former UST locations at the Site appears to have also migrated via groundwater onto the adjoining former JML Optical, Inc. property located west of the Site.

A source of the chlorinated VOCs detected in groundwater samples from various monitoring wells is unknown, but seems attributable to an area-wide groundwater condition.

Factors affecting contaminant migration include: groundwater flow; advection; mechanical dispersion; molecular diffusion; partitioning between air, soil and groundwater; and adsorption of constituents onto soil particles or particles suspended in groundwater.

The type of contamination present at the Site generally consists of SVOCs with lesser amounts of VOCs, and is basically related to petroleum products and/or plasticizers. In general, the detected VOCs are more soluble in water than the detected SVOCs; thus, the VOCs tend to be more mobile in the environment (e.g., migrating through the groundwater and vaporizing into the unsaturated zone).

The estimated average linear groundwater flow velocity for the Site is calculated to range between 0.0006 ft/day and 0.6 ft/day (i.e., 0.22 ft/year to 219 ft/year). The factors described above impact the contaminant flow rates, and the physical properties of the contaminants can impact migration rates.

Exposure Assessment

Under current site conditions, no complete human health exposure pathway has been identified, and it was determined that a Fish and Wildlife Resources Impact Analysis was not needed. However, the following potential future activities have been identified as potential human health exposure pathways:

- Future construction workers and occupants of existing or future buildings on portions of the Site and the adjoining former JML Optical, Inc. property that are constructed over areas of soil, groundwater or LNAPL could be exposed primarily to VOCs, SVOCs and possibly metals that are present at concentrations exceeding SCG values. Examples of exposure include: during disturbance of contaminated material; potential volatilization of VOCs into existing or future site structures; etc. Routes of exposure could include inhalation, ingestion, dermal contact, eye contact, and puncture/injection.
- Future potential on-site or off-site use of groundwater that originates from the Site could pose a potential exposure pathway to VOCs, SVOCs and possibly metals that are present in groundwater at concentrations exceeding SCG values. The primary potential route of exposure would be ingestion; however, other potential routes of exposure include inhalation, dermal contact, eye contact, and puncture/injection.

Conclusions

The hydraulic loading docks and pad-mounted transformers at the Site do not appear to have adversely impacted environmental conditions at the Site. In addition, evidence of environmental impact was not detected at test boring locations TB-17 through TB-19 that were completed in proximity to a pipe chase located inside the main building. Therefore, it does not appear that this pipe chase has adversely impacted environmental conditions at the Site.

Prior to the remedial investigation, a 15,000-gallon UST was removed from the Site. As part of the remedial investigation, one 8,000-gallon UST, two 2,000-gallon USTs and one 4,000-gallon UST were removed from the Site. These five USTs were located in the same general area north of the northwest corner of the main building.

A primary area of soil and groundwater contamination, including the presence of a relatively thin layer (i.e., 0.37 foot or less) of LNAPL that is more limited in extent, was detected in proximity to the former UST locations near the northwest corner of the main building. This contamination generally consists of petroleum products and plasticizers that historically leaked from the former USTs. Based on field screening, analytical laboratory test results, and groundwater monitoring, it appears that this impact has migrated radially outward from the former UST area, including beneath the northwest corner of the main building. In addition, the length of the petroleum/plasticizer plume is estimated to be about 60 feet away from the former UST locations. Based on a review of Site data and environmental reports for the adjoining former JML Optical, Inc. property, petroleum and plasticizer contamination attributable to the former UST locations at the Site appears to have also migrated via groundwater onto an estimated 1,375-square foot area of the adjoining former JML Optical, Inc. property located west of the Site.

As an IRM, petroleum and plasticizer contaminated soils that were displaced during the UST removal work were amended with bioremediation products and placed back into the tank pit excavation as part

of an in-situ bioremediation system. The analysis of post-treatment soil and groundwater samples indicate the in-situ bioremediation system is working, and contaminants have been reduced by approximately 40% (on average). The results of the post-treatment sampling and analytical laboratory testing indicated that contaminants were still present in soil and groundwater at concentrations exceeding SCG values. Based on the work conducted to date, further actions may be warranted to address the residual petroleum and plasticizer contamination located in proximity to the northwest corner of the main building.

Two of four surface soil samples collected from the northern undeveloped portion of the Site, and a subsurface soil sample collected from a depth interval of 0-4' at test boring TB-4 contained some PAH SVOCs above December 14, 2006 NYSDEC Part 375 (Environmental Restoration Programs) Track 2 SCOs for Restricted Residential Use. However, the concentrations of these SVOCs are comparable to other projects in the City of Rochester where surface soil data has been collected. As such, the limited exceedances of the Restricted Residential Use SCOs in soil at the Site are not significant and are likely attributable to the local geology or urban setting of the Site.

Chlorinated VOCs were detected in groundwater samples at some of the monitoring well locations. An on-site source of chlorinated VOCs that could result in contamination of the groundwater was not found during the soil and groundwater studies performed as part of the remedial investigation. It is possible that the chlorinated VOCs are attributable to an off-site source(s) that has resulted in an areawide groundwater condition. A review of environmental reports indicates a sump and a former degreaser area at the adjoining former JML Optical, Inc. property to the west, and also an NYSDEC Inactive Hazardous Waste Disposal Site located southwest of the Site, could potentially be the sources of the chlorinated VOCs that are present at the Site. Depending upon future use of the Site, further actions may be warranted in relation to chlorinated VOCs in groundwater at the Site.

Remedial Alternatives Analysis

The RI/RAA report identified Remedial Action Objectives (RAOs), contaminants of interest, remediation criteria, and general response actions. In regard to these criteria, four remedial alternatives were developed and evaluated. These alternatives are summarized below:

Alternative #1 No Action

Alternative #2 LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls

Alternative #3 Limited In-Situ Remediation, LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls

Alternative #4 Full Excavation, LNAPL Recovery, In-Situ Remediation and Groundwater Monitoring for 5 Years

A detailed evaluation of the four remedial alternatives was performed, and implementation of Alternative #2 (LNAPL Recovery, Monitored Natural Attenuation, Institutional Controls and Engineering Controls) was recommended in the RI/RAA report for the Site. Alternative #2 is summarized in the Executive Summary, and further details are provided in subsequent sections of this Remedial Work Plan.

1.3 Proposed Future Use of Site

The Site will be used for restricted residential, commercial or industrial use. Actual redevelopment plans were not available at the time this RWP was submitted.

1.4 Objectives

The objectives of this remediation project are to implement a remedial alternative that is protective of human health and the environment, addresses current environmental conditions to the satisfaction of the NYSDEC and the general public, and allows for restricted residential, commercial or industrial use of the Site.

1.5 Applicable Project SCGs

Applicable SCGs that may be used for this project are outlined below:

- Guidelines referenced in the NYSDEC document titled "Draft Brownfield Cleanup Program Guide", May 2004.
- Guidelines referenced in the NYSDEC document titled "Draft DER-10 Technical Guidance for Site Investigation and Remediation", December 2002.
- Appropriate SCOs as set forth in the NYSDEC document titled "6 NYCRR Part 375 Environmental Remediation Programs" dated December 14, 2006.
- Groundwater standards and guidance values as referenced in the NYSDEC Division of Water Technical and Operational Guidance Series 1.1.1 document titled "Ambient Water Quality Standards and Guidance Values and Groundwater Effluent Limitations" (TOGS 1.1.1), June 1998 (as amended by an April 2000 addendum).

1.6 Citizen Participation

In accordance with NYSDEC BCP requirements, a Citizen Participation Plan (CPP) dated November 2004 was developed for this project. This CPP is available for review at the document repositories (i.e., NYSDEC Region 8 offices located at 6274 East Avon-Lima Road, Avon, New York 14414; and the Central Library of Rochester and Monroe County located at 115 South Avenue, Rochester, New York 14604). As part of the CPP, Fact Sheets are provided to entities on the CPP mailing list to keep the public informed of project activities and documents that are available for review and/or comment. The CPP allows the general public and other interested parties to provide comments on plans for the Site, including this RWP.

2.0 REMEDIAL ACTIONS

The source of petroleum and plasticizer contamination in proximity to the northwest corner of the main building has been addressed by removing five USTs and installing an in-situ bioremediation system at this former UST area. The remedial alternative selected for the Site consists of various technical and administrative actions that are intended to: supplement the existing in-situ bioremediation system by monitoring and recovery of LNAPL; reduce exposure to Site contaminants; provide monitoring of natural attenuation processes to ensure that the contamination is not migrating; and provide a contingency for additional in-situ applications of bioremediation products in and/or around the in-situ bioremediation system. This remedial alternative is considered a Track 4 cleanup that allows for restricted residential, commercial and industrial use. This section of the RWP provides details on the actions that will be conducted as part of this remedial alternative. In general, the remedial actions will include:

- LNAPL monitoring and recovery;
- Development and recording of an environmental easement;
- Development of a SMP to address the residual contamination and any use restrictions;
- Confirmatory soil and groundwater sampling at the IRM area, and supplemental bioremediation treatment and additional confirmatory sampling if deemed necessary by the NYSDEC;
- Monitored natural attenuation of contaminants in groundwater; and
- Periodic certification of the institutional controls (ICs).

A site-specific health and safety plan (HASP) for this project is included in Appendix B. This HASP outlines the policies and procedures necessary to protect workers and the public from potential environmental hazards posed during project activities.

The technical and administrative actions associated with the selected remedy are further presented in this RWP. It is currently planned that Mitkem Laboratories, a Division of Spectrum Analytical, Inc. (Mitkem), which is a NYSDOH Environmental Laboratory Approval Program (ELAP)-certified analytical laboratory (NYSDOH ELAP ID #11522), will analyze samples of soil and groundwater that are generated as part of this project. A quality assurance project plan (QAPP) for this project is included as Appendix C.

2.1 Site Preparation and Signage

Site preparation and signage tasks will be completed prior to performing additional on-site remedial tasks. Components will include:

- A sign will be conspicuously posted at the Site prior to the start of remedial work. An example of the proposed signage and associated NYSDEC instructions are included in Appendix D. The sign will reflect the actual holders of the positions indicated on the sign at the time the sign is installed.
- A utility stakeout will be completed and field checked prior to installing LNAPL recovery wells described in Section 2.2. With concurrence from the NYSDEC site representative, and to the extent deemed warranted, adjustments may be made in the field concerning the actual locations of the recovery wells to account for the location of buried utilities, as well as other factors.

2.2 LNAPL Monitoring and Recovery

A drilling subcontractor will be retained to install four 2-inch diameter groundwater wells for use in monitoring and recovery of LNAPL. As shown on Figure 6, two of the wells will be installed inside the existing main building, and two of the wells will be installed at exterior locations west of the northwest corner of the main building. These wells are in the area where LNAPL is suspected. Based on field conditions encountered, the actual locations of the wells may vary from those shown on Figure 6 with input from the NYSDEC Site representative. If field evidence of contamination is not encountered at a specific well location, a decision may be made in the field to backfill the specific borehole, and possibly install a new borehole at a different location for installation of the specific well. Well installation and construction information is provided in Section 3.0 of the QAPP included as Appendix C.

The DAY representative will conduct health and safety air monitoring for VOCs and particulates during the well installation work in accordance with provisions of the HASP and Community Air Monitoring Plan (CAMP) (refer to Appendix B). The contractor will either utilize this HASP or the components of its own HASP (accepted by regulatory agencies) for the protection of its on-site workers.

At least one week following installation, the groundwater wells will be developed using the protocol outlined in Section 3.0 of the QAPP included as Appendix C. A licensed surveyor will also measure the horizontal and vertical coordinates of the new wells using the same coordinate system that was used during the survey of the existing wells.

Initially, the fourteen existing monitoring wells (i.e., MW-1 through MW-11 and MWIRM-1 through MWIRM-3) and the four new wells will be monitored for the presence of LNAPL using a Heron Model H01.1 oil water interface probe (refer to Figure 5 and Figure 6 for locations of existing and new wells). After establishing which wells contain LNAPL, the LNAPL will be removed by bailing, pumping, passive absorbent socks, etc. The actual recovery methods used will be based on the amount of LNAPL and associated recharge rate at each well location. Over time, the number of wells to be monitored may be decreased based on the previous LNAPL monitoring and recovery data that is generated (i.e., it is currently anticipated that LNAPL may be present at existing wells MW-6 and MW-10, and at the four new wells to be installed). It is anticipated that LNAPL monitoring and recovery will be completed on a monthly basis; however, this schedule will be adjusted accordingly with input from the NYSDEC site representative. It is anticipated that LNAPL monitoring and recovery will be completed for a period of two years.

Recovered LNAPL will be placed in 55-gallon drums that will be stored in a secure location at the Site (e.g., inside the existing main building, etc.). Full drums will be shipped off-site for recycling or disposal in accordance with applicable regulations.

2.3 Confirmatory Sampling and Analysis from In-Situ Bioremediation System Area

Approximately two months after starting LNAPL monitoring and recovery work, confirmatory soil and groundwater sampling and analysis will completed to assist in evaluating the effectiveness of the existing in-situ bioremediation system. The results of the confirmatory sampling and analysis will be used to assist in determining (with input from the NYSDEC site representative) whether injection of additional remediation products at the existing in-situ bioremediation system is necessary (refer to Section 2.5.1).

Soil

A total of four test borings (designated as TBC-1 through TBC-4) will be advanced within the in-situ bioremediation system. Tentative locations of these test borings are shown on Figure 7; however, final locations may vary from those shown based on field conditions encountered and input from the NYSDEC site representative. Tape measurements from control points or existing site structures will be used to record the locations of test borings for subsequent transfer to a geographic information system (GIS) or computer-aided design (CAD). Based on the results of the previous remedial investigation, it is anticipated that the test borings will be advanced to depths up to approximately 20 feet below the ground surface. However, these depths may be modified based on field observations, input from the NYSDEC site representative during their advancement, and equipment refusal at depths less than 20 feet. Soil samples will be collected throughout the entire depth of each test boring.

The recovered soil samples will be collected, observed, monitored and documented in accordance with the protocols outlined in Section 3.0 of the QAPP included as Appendix C, which includes recording pertinent information on test boring logs. Each test boring will be backfilled with grout upon completion, and soil cuttings will be placed in New York State Department of Transportation (NYSDOT)-approved drums that will be characterized and disposed off-site in accordance with applicable regulations.

With input from the NYSDEC Site representative, one soil sample from each test boring (i.e., total of four samples) with the greatest field evidence of petroleum and/or plasticizer impact (i.e., elevated photoionization detector [PID] readings, staining, odors, etc.) will be selected for analytical laboratory testing. The analytical laboratory testing program for these samples is identified on Table 1 of the QAPP included as Appendix C. Portions of the samples will be provided to the respective analytical laboratories under chain-of-custody control. As shown, Mitkem will analyze a portion of each soil sample for the following parameters:

- Target compound list (TCL) VOCs including TICs using NYSDEC Analytical Services Protocol (ASP) Method OLM04.3;
- TCL SVOCs including TICs using NYSDEC ASP Method OLM04.3; and
- TPH using NYSDOH Method 310.13.

In addition, Osprey Biotechnics (Osprey) in Sarasota, Florida will analyze a portion of each soil sample for total plate count and pseudomonas plate count using United States Environmental Protection Agency (USEPA) Method 9215C. Osprey is the analytical laboratory that provides plate count testing on behalf of CL Solutions, which may provide additional Petrox 1 and Petrox 3 bioremediation amendment products for injection in the in-situ bioremediation system, if warranted. Each sample will be serially diluted into previously prepared buffer solutions. The dilutions will be aseptically transferred and plated according to the method onto Trypticase Soy Agar (TSA) media. The plates will be incubated at 25°C with counts taken at 48 hours. The counts will be reported as Colony Forming Units per milliliter of sample (CFU/ml).

The test results will be summarized on data tables that also include a comparison to previous post-treatment data and available/applicable SCGs.

Groundwater

Three monitoring wells (designated as wells MWIRM-1 through MWIRM-3) were installed within the former UST excavation during the construction of the in-situ bioremediation system. The approximate locations of these wells are shown on Figure 7 and Figure 9. These vertical wells were installed to intercept the amended soil beneath the water table for monitoring purposes and possible future delivery of bioremediation products.

Confirmatory groundwater samples will be collected from monitoring wells MWIRM-1 through MWIRM-3. Groundwater sampling will be conducted using the low-flow purge and sample protocol outlined in Section 3.0 of the QAPP included as Appendix C. Monitoring Well Sampling Logs will be developed that document the procedures and equipment used during the purging and groundwater sampling, and the field measurement data that is obtained.

The analytical laboratory testing program for these samples is identified on Table 1 of the QAPP included as Appendix C. An aliquot of each sample will be provided to the respective analytical laboratories under chain-of-custody control. As shown, Mitkem will analyze the aliquot of each groundwater sample for the following parameters:

- TCL VOCs including TICs using NYSDEC ASP Method OLM04.3; and
- TCL SVOCs including TICs using NYSDEC ASP Method OLM04.3.

In addition, Osprey will analyze an aliquot of each groundwater sample for total plate count and pseudomonas plate count using USEPA Method 9215C.

The test results will be summarized on data tables that also include a comparison to previous post-treatment data and available/applicable SCGs.

2.4 Monitored Natural Attenuation

A monitored natural attenuation (MNA) program will be implemented for the Site. The objective of MNA is to collect and document site-specific data that can be used to evaluate the effectiveness of natural attenuation in controlling contaminants that are present at the Site (i.e., specifically the petroleum/plasticizer contaminated area located at the northwest corner of the main building). The MNA for this project will include the following components:

- Prior to collecting a round of MNA groundwater samples, static water level measurements will be collected from wells MW-1 through MW-11, MWIRM-1 through MWIRM-3, and the four new LNAPL monitoring/recovery wells. Using static water level measurements from these wells, and the surveyed well elevations, DAY will calculate groundwater elevations for each groundwater monitoring event. With assistance of the Surfer 8 software program, the well locations and corresponding groundwater elevations will be used to develop a groundwater potentiometric map for each monitoring event.
- Up to eight groundwater wells that are located at upgradient and downgradient positions from the most contaminated portion of the Site (i.e., upgradient and downgradient of the area in proximity to the northwest corner of the main building) will be sampled during each natural attenuation monitoring event. It is anticipated that wells that could be used during the MNA program could include: MW-2 through MW-6; MW-10; MWIRM-1 through MWIRM-3, and the four new LNAPL monitoring/recovery wells (refer to Figure 5, Figure 6, Figure 7 and Figure 9). This well field should result in a transect of monitoring points across the plume. The NYSDEC will approve the actual wells to be used during the MNA program.

- It is currently anticipated that the natural attenuation monitoring will be conducted for a period of up to five years. It is assumed that the eight wells will be sampled on a bi-annual basis during the 1st and 2nd years, and on an annual basis for the 3rd through 5th years. Samples will be collected using the low-flow purge and sample protocol outlined in Section 3.0 of the QAPP included as Appendix C. As shown on Table 1 of the QAPP included as Appendix C, each round of MNA groundwater samples will be tested for the following parameters:
 - TCL VOCs including TICs using NYSDEC ASP Method OLM04.3;
 - TCL SVOCs including TICs using NYSDEC ASP Method OLM04.3; and
 - Natural attenuation parameters such as nitrate, iron (II), manganese, sulfate, methane, and chloride (Methods SM3500D, E300IC, ILM04.1, and RSK175).
- Water quality measurements such as dissolved oxygen, oxidation-reduction potential, pH, temperature, conductivity, and turbidity using a Horiba U-22 water quality meter (or equivalent) will also be obtained for each round of MNA groundwater samples.
- The cumulative data will be evaluated as it is generated in order to determine if natural attenuation processes are occurring in a manner that controls migration of contamination away from the Site and at a rate that is acceptable to the NYSDEC. For instance, hydrologic data, geochemical data, chemical data and/or biological data would be used to assist in evaluating the following conditions and trends:
 - Chemical mass, concentrations, and toxicity at appropriate monitoring wells over time.
 - Specific types of natural attenuation processes that are, or may be, occurring such as
 advection, adsorption, mechanical dispersion, dissolution, aerobic decay (e.g., aerobic
 respiration involving oxygen) and anaerobic decay (e.g., denitrification, ferric reduction
 sulfate reduction, and methanogenesis).
- The detected concentrations of VOCs and SVOCs for each monitoring event will be compared on a summary table to TOGS 1.1.1 groundwater standards and guidance values. The test results will also be evaluated on a cumulative basis.
- Under this alternative, free computer software programs, such as GSI Environmental, Inc.'s BIOSCREEN model and possibly also Mass Flux ToolKit, will be used to assist in evaluating the effectiveness of natural attenuation, make projections on the estimated time of remediation (i.e., rate of natural attenuation), and calculating contaminant mass flux based on site-specific data. BIOSCREEN is a screening model that simulates remediation through natural attenuation of dissolved hydrocarbons at petroleum release sites. The software utilizes the Domenico analytical solute transport model, has the ability to simulate advection, dispersion, adsorption, aerobic decay, and anaerobic reactions/decay. BIOSCREEN model types that can be run include: solute transport without decay; solute transport with biodegradation modeled as a first-order decay process (simple, lumped-parameter approach); and, solute transport with biodegradation modeled as an instantaneous biodegradation reaction with multiple soluble electron acceptors including dissolved oxygen, nitrate, and sulfate. The BIOSCREEN model is designed to simulate biodegradation by both aerobic and anaerobic reactions and can perform mass flux calculations. Using the Mass Flux ToolKit, the calculated contaminant mass flux data can be used to demonstrate the progress of natural attenuation.

With approval from regulatory agencies, the duration and frequency of the natural attenuation groundwater monitoring, as well as the list of parameters to be tested, may be adjusted as the monitoring program progresses.

2.5 Institutional Controls and Engineering Controls

ICs will be used to address residual contamination that may remain in soil, fill or groundwater, including protecting against exposure to residual contamination. Development of ICs will start once the RWP is approved by the NYSDEC. The ICs are provided in the subsections that follow.

2.5.1 Site Management Plan

A SMP will be developed and implemented to address the characterization, handling, and disposal/reuse of residual contaminated media (e.g., soil, fill, groundwater) that is disturbed during any future site activities. The SMP will also require evaluation of the potential for vapor intrusion into any future buildings to be constructed on the Site (and also the two existing buildings if they are to be occupied), including requirements to mitigate such potential vapor intrusions through use of environmental engineering controls (e.g., sub-slab depressurization system, etc.) or other means, as warranted. In addition, the SMP will identify use restrictions for the Site (e.g., property development and groundwater use restrictions, etc.). The SMP will also include a generic HASP for the Site, and require that this HASP (or a project-specific HASP) be implemented when known or suspected impacted media at the Site have the potential to be disturbed.

2.5.1.1 Additional In-Situ Remediation Application (Contingency)

Subsequent to review of LNAPL monitoring and recovery outlined in Section 2.2, the analytical test results for confirmatory soil and groundwater samples outlined in Section 2.3, and the analytical test results for at least one round of monitored natural attenuation samples outlined in Section 2.4, a decision will be made with input from the NYSDEC regarding whether additional remediation products should be applied within and/or around the existing in-situ bioremediation system. If additional application of remediation products is deemed warranted, the SMP will outline the scope of the application, and any subsequent confirmatory monitoring, sampling and analysis. It is anticipated that the same, or similar types of remediation products will be used during the additional application as those used during construction of the in-situ bioremediation system.

2.5.2 Environmental Easement

An environmental easement will be developed for the Site. The environmental easement will: limit use of the Site to restricted residential, commercial and industrial applications; require compliance with the SMP; restrict use of groundwater as a source of potable water or process water without necessary water quality treatment as determined by the NYSDOH; and, require the property owner to complete and submit to the NYSDEC the periodic certification of institutional controls and also engineering controls (if installed).

2.5.3 Periodic Certification

The property owner will provide certification of institutional controls and engineering controls (if warranted), prepared and submitted by a professional engineer or such other expert acceptable to the NYSDEC, until the NYSDEC notifies the property owner in writing that this certification is no longer needed. It is anticipated that the periodic certification will be provided every three years. This submittal will: contain certification that the institutional controls and engineering controls (if warranted) put in place are still in place and are either unchanged from the previous certification or are complaint with NYSDEC-approved modifications; allow the NYSDEC access to the Site; and, state that nothing has occurred that would impair the ability of the controls to protect public health or the environment, or constitute a violation of failure to comply with the site management plan unless otherwise approved by the NYSDEC.

2.5.4 City Code Restricting Groundwater Use

Chapter 59 (Health and Sanitation), Article III (Nuisances and Sanitation) § 59-27 (Water Supply) of the current Charter and Code of the City of Rochester, New York states:

- A. No person shall use for drinking purposes, or in the preparation of food intended for human consumption, any water except the potable water supply authorized for public use by the City of Rochester; and
- B. Other water supplies, wells or springs used for cooling and washing purposes only, where food is prepared or sold for human consumption, shall be tested and approved by the Monroe County Health Director. All auxiliary water supplies used for commercial or industrial use shall have all hydrants and faucets conspicuously posted indicating that such water is not for drinking use, and such water supplies shall not be cross-connected or interconnected with the public water supply."

This City Code has been interpreted to represent an IC that prohibits groundwater within the city limits to be used as a source of potable water.

2.6 Engineering Controls

Once the actual redevelopment plan for the Site has been identified, an Engineering Control Design Plan (EC Design Plan) will be submitted to regulatory agencies if mitigation of potential vapor intrusions into buildings is warranted. Depending upon timing, the EC Design Plan may be included in the SMP or may be a stand-alone document. The EC Design Plan must be accepted by the NYSDEC and the NYSDOH prior to conducting the associated redevelopment. Depending upon specific redevelopment plans, the engineering controls may consist of a sub-slab depressurization system on new or existing buildings, a sub-slab membrane system, other means, or a combination of technologies.

2.7 Remediation-Derived Wastes

It is anticipated that soil cuttings, well development water, well sampling water, decontamination water, LNAPL, and solid waste will be generated during various stages of this project. These wastes will be handled, characterized, and disposed off-site in accordance with applicable regulations. It is currently anticipated that soil will be transported and disposed off-site at a NYSDEC-approved regulated landfill facility, and that containerized well development water, well purge water, and decontamination water will be disposed off-site through publicly owned treatment works (POTW) system or other NYSDEC-approved disposal facility. LNAPL will be transported off-site for recycling or disposal in accordance with applicable regulations.

3.0 FINAL ENGINEERING REPORT

A final engineering report (FER) will be developed for this project. This FER will include: a summary of the work completed; field documentation; scaled figures depicting monitoring well and LNAPL monitoring/recovery well locations, confirmatory soil sample locations, a groundwater potentiometric map(s); analytical laboratory sampling documentation and test results; data tables; and documentation concerning the transport and disposal of remediation-derived wastes (if documentation is available at the time the FER is submitted).

Information and data for the first round of MNA groundwater samples will be included in the FER. Information and data for subsequent rounds of MNA groundwater samples will be provided in annual MNA reports.

3.1 Certificate of Completion

It is anticipated that the NYSDEC will issue a Certificate of Completion once the FER and SMP are completed and accepted by regulatory agencies, and the environmental easement is executed and recorded.

4.0 PROJECT SCHEDULE

A schedule for the first year of this project is included in Appendix E. Not shown on this schedule are the following components:

- Natural Attenuation Monitoring and associated annual reports for years two through five.
- Periodic certification of ICs that are anticipated to be developed and provided every three years subsequent to receipt of the certificate of completion for the project.

5.0 REFERENCES

Previous Reports and Plans

Phase I Environmental Site Assessment (100 and 142 Fernwood Avenue; 31, 35 and 41 Rosemary Drive; 25,29,33,39,43,48 and 55 Ilex Place; Rochester, New York); November 15, 2000; Day Environmental, Inc.

Remedial Investigation/Remedial Alternatives Analysis Report; Former Vogt Manufacturing Facility; Brownfield Cleanup Program (BCP); NYSDEC Site ID C828119; 100 Fernwood Avenue, Rochester, New York; November 2006; Day Environmental, Inc.

Addendum to Remedial Investigation/Remedial Alternatives Analysis Report; Former Vogt Manufacturing Facility; Brownfield Cleanup Program (BCP); NYSDEC Site ID C828119; 100 Fernwood Avenue, Rochester, New York; March 8, 2007; Day Environmental, Inc.

Citizen Participation Plan; Brownfield Cleanup Program; Former Vogt Manufacturing Facility; 100 and 142 Fernwood Avenue; 31, 35 and 41 Rosemary Drive; 25, 29, 33, 39, 43, 49 and 55 Ilex Place; Rochester, New York; NYSDEC BCP Project #C828119; November 2004; Day Environmental, Inc.

Regulatory Documents

NYSDEC Division of Water Technical and Operational Guidance Series 1.1.1 document titled "Ambient Water Quality Standards and Guidance Values and Groundwater Effluent Limitations" (TOGS 1.1.1) dated June 1998, including April 2000 Addendum Table 1.

NYSDEC DER Draft Brownfield Cleanup Program Guide; May 2004.

NYSDEC 6 NYCRR Part 375 Environmental Remediation Programs; effective December 14, 2006.

City of Rochester, New York Charter and Code; Chapter 59 (Health and Sanitation), Article III (Nuisances and Sanitation) § 59-27 (Water Supply).

NYSDEC Draft DER-10 Technical Guidance for Site Investigation and Remediation, December 2002.

Reference Materials

Ground Water Resources of Monroe County; 1935; R.M. Leggette, L.O. Gould and B.H. Dollen USGS topographic map for the Rochester East, New York quadrangle, 1995

Generalized Groundwater Contour Map (Rochester East Quadrangle); 1980; Dr. R. A. Young

Internet References

GSI Environmental BIOSCREEN Natural Attenuation Computer Model (Free Software) developed for the U.S. Air Force; (http://www.epa.gov/ada/csmos/models/bioscrn.html)

GSI Environmental Mass Flux ToolKit Computer Model (Free Software) developed for the Department of Defense ESTCP program (http://www.gsi-net.com/Software/massfluxtoolkit.asp)

6.0 ACRONYMS

ASP Analytical Services Protocol

ASTM American Society for Testing and Materials

BCP Brownfield Cleanup Program
CAD Computer-Aided Design

CAMP Community Air Monitoring Plan
CFU/ML Colony Forming Units Per Milliliter

CPP Citizen Participation Plan DAY Day Environmental, Inc.

DNAPL Dense Non-Aqueous Phase Liquid EC Design Plan Engineering Control Design Plan

ELAP Environmental Laboratory Approval Program

FER Final Engineering Report
GIS Geographic Information System

HASP Health And Safety Plan
IC Institutional Control
IRM Interim Remedial Measure
LNAPL Light Non-Aqueous Phase Liquid

MCDPH Monroe County Department of Public Health

Mitkem Laboratories, a Division of Spectrum Analytical, Inc.

MNA Monitored Natural Attenuation

NYCRR New York Codes, Rules and Regulations

NYSDEC New York State Department of Environmental Conservation

NYSDOH New York State Department of Health

NYSDOT New York State Department of Transportation

Osprey Osprey Biotechnics

PAH Polyaromatic Hydrocarbon PCB Polychlorinated Biphenyl

Phase I ESA Phase I Environmental Site Assessment

PID Photoionization Detector

POTW Publicly Owned Treatment Works

PPB Parts Per Billion PPM Parts Per Million

QAPP Quality Assurance Project Plan RAO Remedial Action Objective

REC Recognized Environmental Condition

RI/RAA Remedial Investigation/Remedial Alternatives Analysis

RWP Remedial Work Plan

SCG Standards, Criteria and Guidance

SCO Soil Cleanup Objective SMP Site Management Plan

SVOC Semi-Volatile Organic Compound

TAL Target Analyte List
TCL Target Compound List

TIC Tentatively Identified Compound

TOGS Technical and Operational Guidance Series

TPH Total Petroleum Hydrocarbon

TSA Trypticase Soy Agar

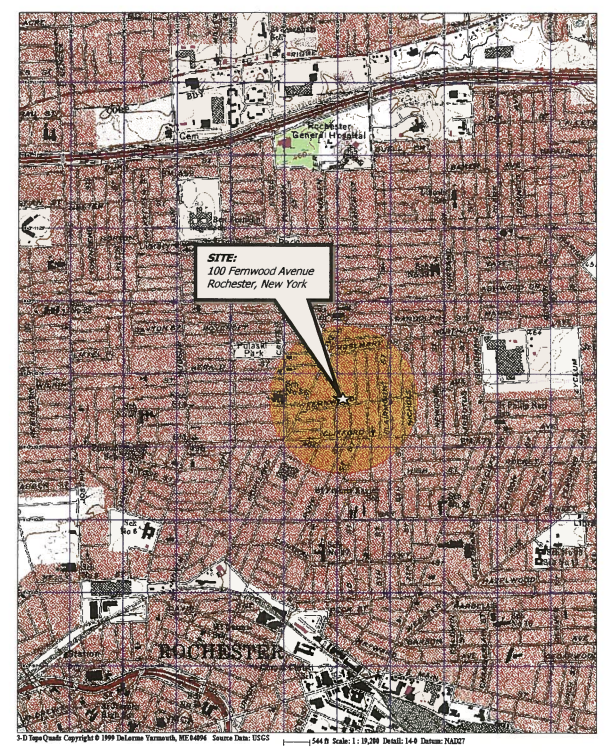
USEPA United States Environmental Protection Agency

USGS United States Geological Survey
UST Underground Storage Tank
VOC Volatile Organic Compound

FIGURES

Day Environmental, Inc. JD5970 / 4014R-07





Drawing Produced From: 3-D TopoQuads, DeLorme Map Co., referencing USGS quad map Rochester East (NY) 1995. Site Lat/Long: N43d-10.66' - W77d-35.22'

DATE 12-04-2007

DRAWN BY

SCALE 1" = 2000' day

DAY ENVIRONMENTAL, INC. ENVIRONMENTAL CONSULTANTS ROCHESTER, NEW YORK 14623-2700

PROJECT TITLE

100 FERNWOOD AVENUE ROCHESTER, NEW YORK

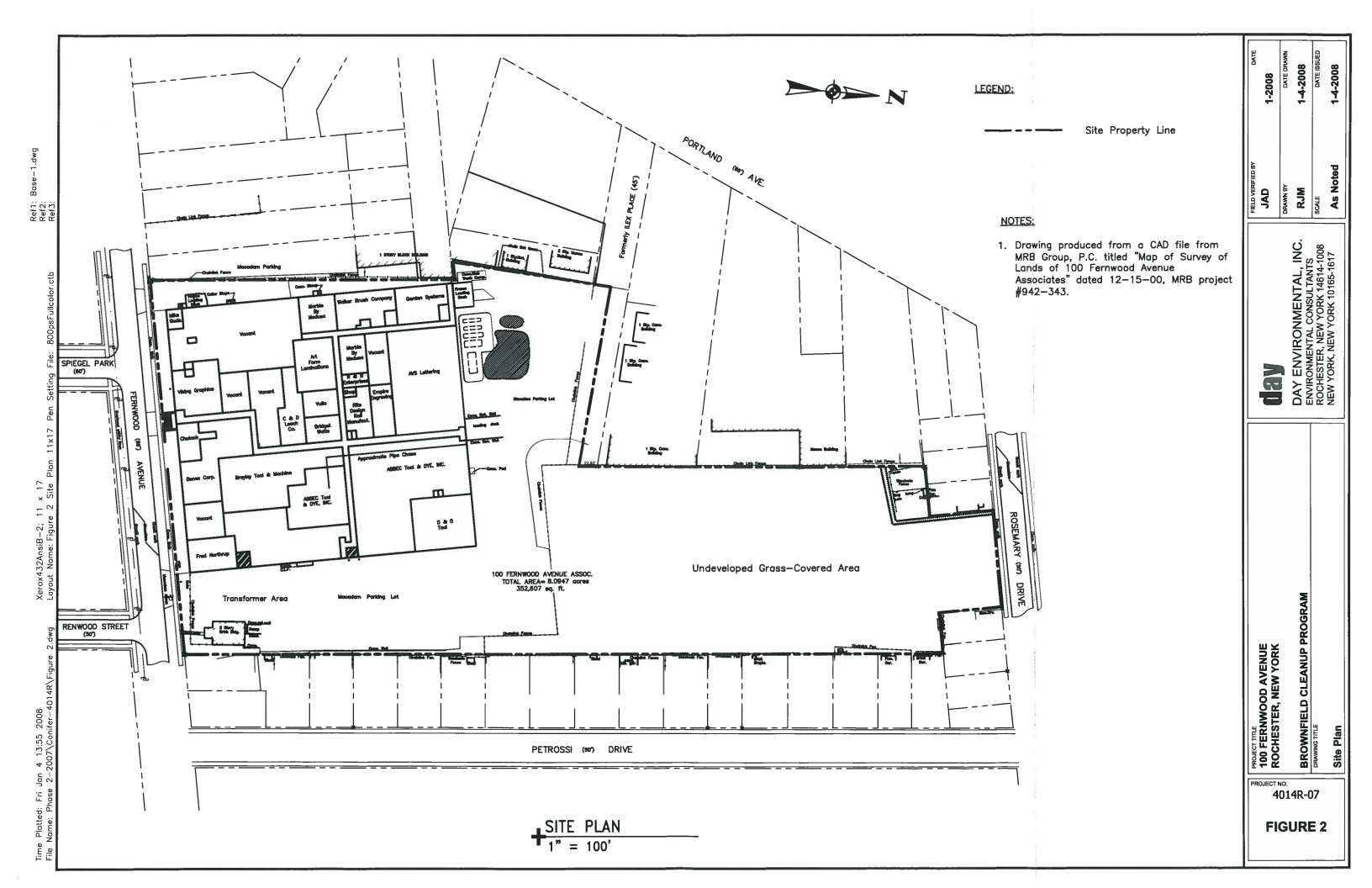
BROWNFIELD CLEANUP PROGRAM

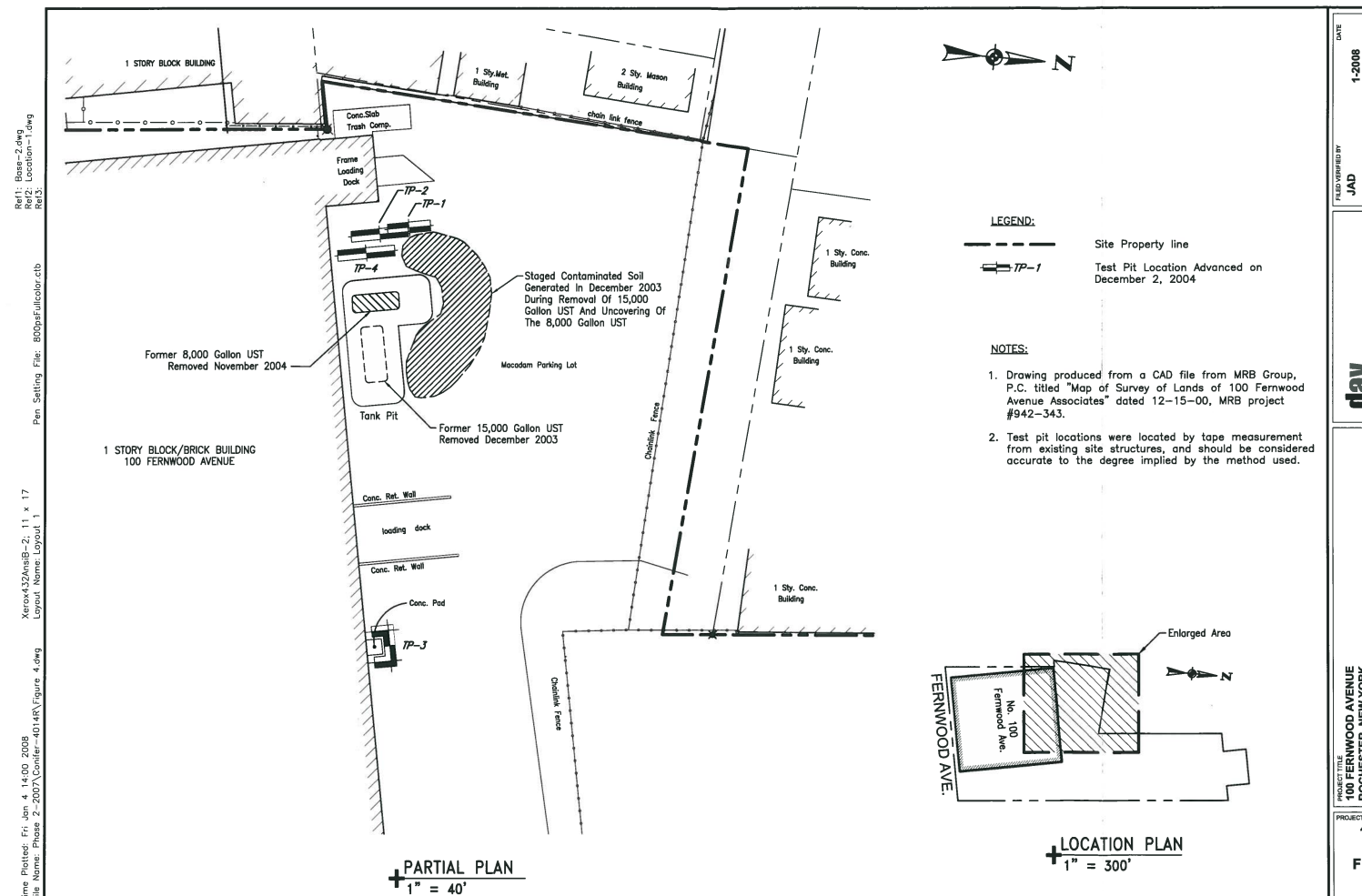
DRAWING TITLE
PROJECT LOCUS MAP

PROJECT NO.

4014R-07

FIGURE 1





1-4-2008 DATE ISSU 1-4-2008 RJM

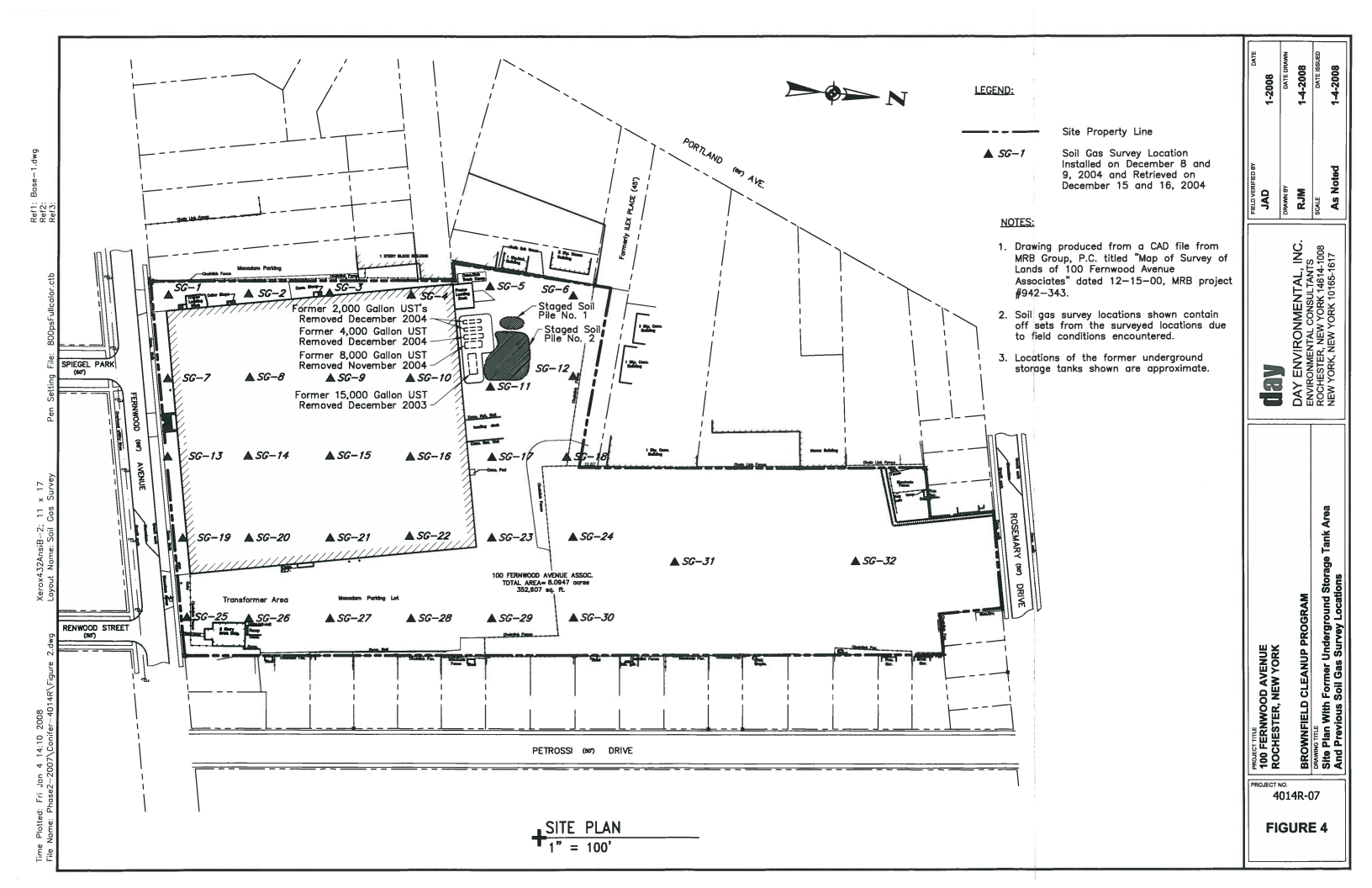
1-2008

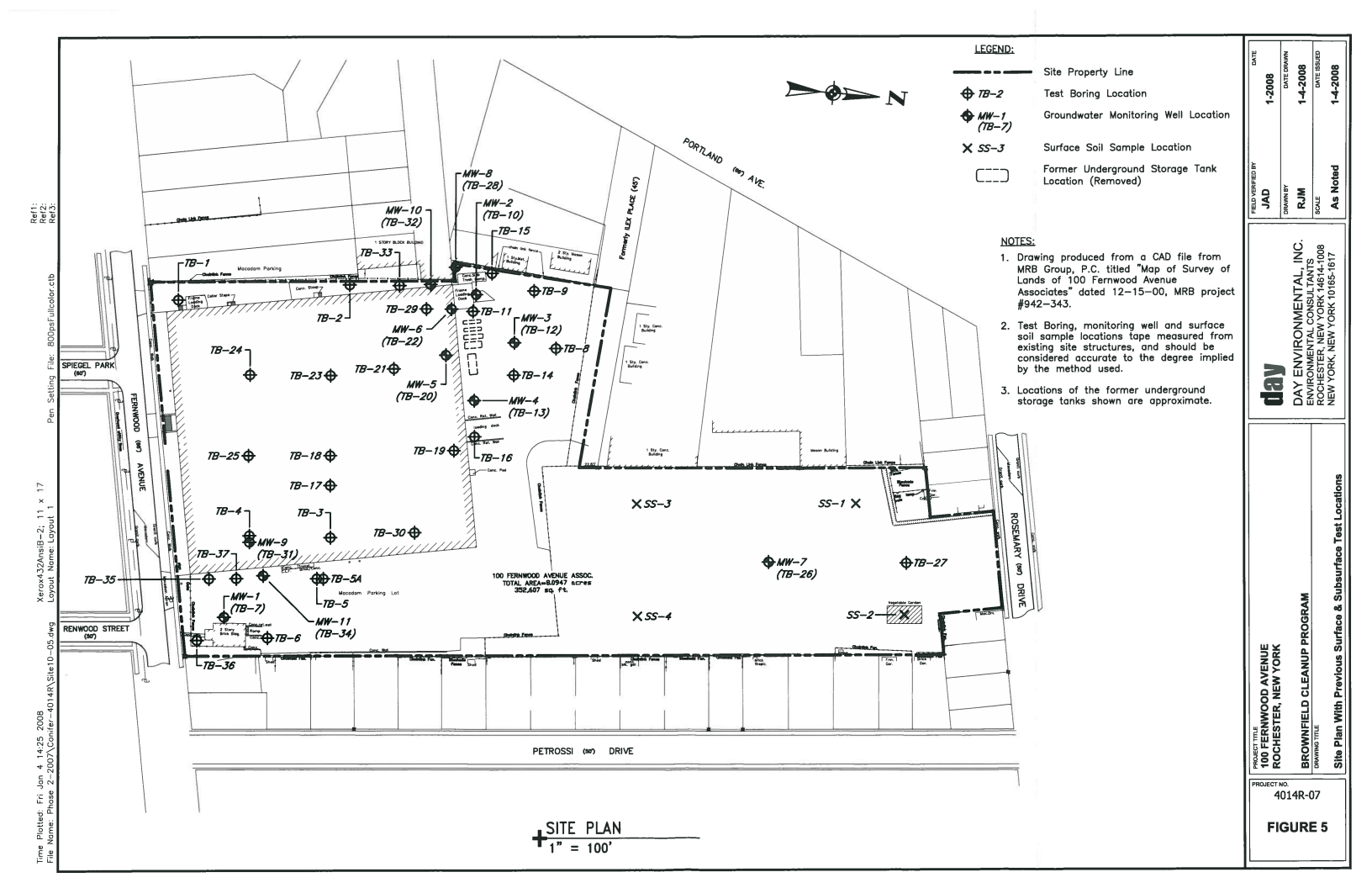
Underground Storage Tank Area evious Test Pit Locations

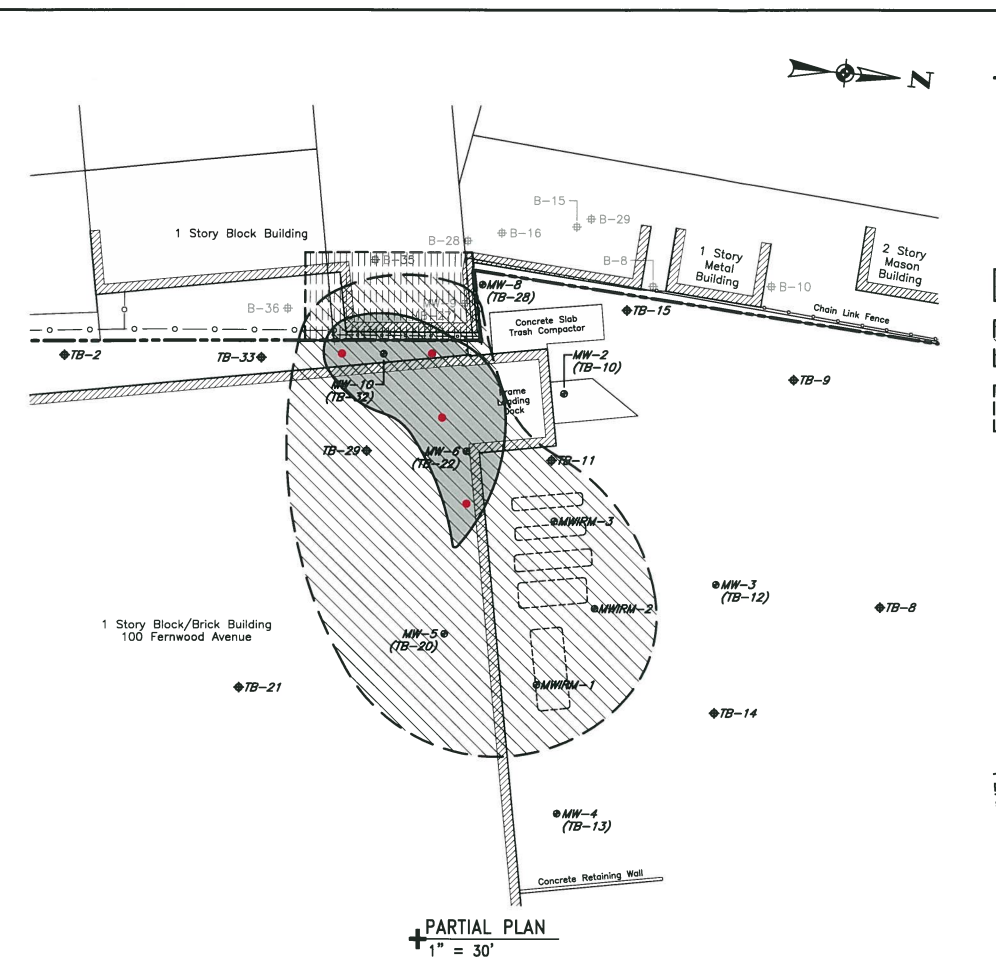
PROJECT TITLE
100 FERNWOOD AVENUE
ROCHESTER, NEW YORK

4014R-07

FIGURE 3







Ref1: Ref2: Ref3:

: Plotted: Fri Jan 4 14:45 2008 Name: Phase 2-2007\Conifer-4014R\UST

LEGEND:

⊕ B-35

Site Property Line

Proposed LNAPL Monitoring And Recovery

Well Location

⊕ 7B−2 **Existing Test Boring Location**

⊕ MW-10 Existing Groundwater Monitoring Well Location (TB-32)

Existing Offsite Test Location By Others

Former Underground Storage Tank Location (Removed)

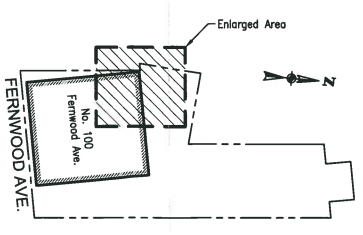
Approximate Extent Of LNAPL That May Be Associated With The Former Underground Storage Tanks

Approximate Extent Of Dissolved Constituents Exceeding TOGS 1.1.1 Groundwater Standards Or Guidance Values That May Be Associated With The Former Underground Storage Tanks

Approximate 55' x 25' Area (1,375 Square Feet) Of Offsite Impact That Appears Attributable To The Former Underground Storage Tank Area

NOTES:

- 1. Drawing produced from a CAD file from MRB Group, P.C. titled "Map of Survey of Lands of 100 Fernwood Avenue Associates" dated 12-15-00, MRB project #942-343.
- 2. Test Boring and monitoring well locations tape measured from existing site structures, and should be considered accurate to the degree implied by the method used.
- 3. Locations of the former underground storage tanks shown are approximate.



LOCATION PLAN

1" = 300'

1-4-2008 DATE ISSU JAD RJM

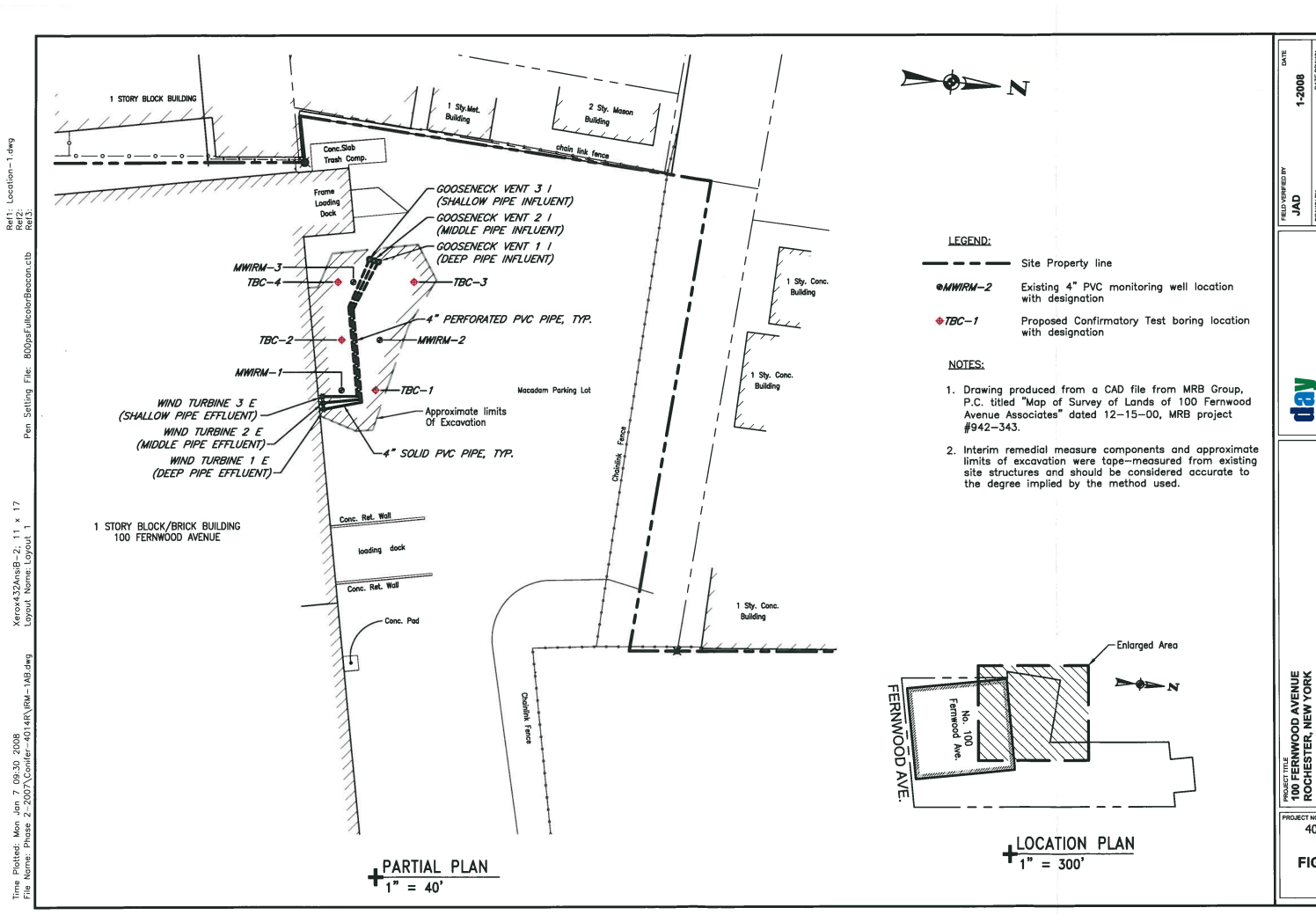
DAY ENVIRONMENTAL, INC. ENVIRONMENTAL CONSULTANTS ROCHESTER, NEW YORK 14614-1008 NEW YORK, NEW YORK, 10165-1617

Monitoring And Recovendenge Underground Storage BROWNFIELD CLEANUP PROGRAM DAWNING TITLE Proposed LNAPL Monitoring And ReNear The Former Underground Stora

PROJECT TITLE 100 FERNWOOD AVENUE ROCHESTER, NEW YORK

4014R-07

FIGURE 6



1-7-2008 As Noted JAD

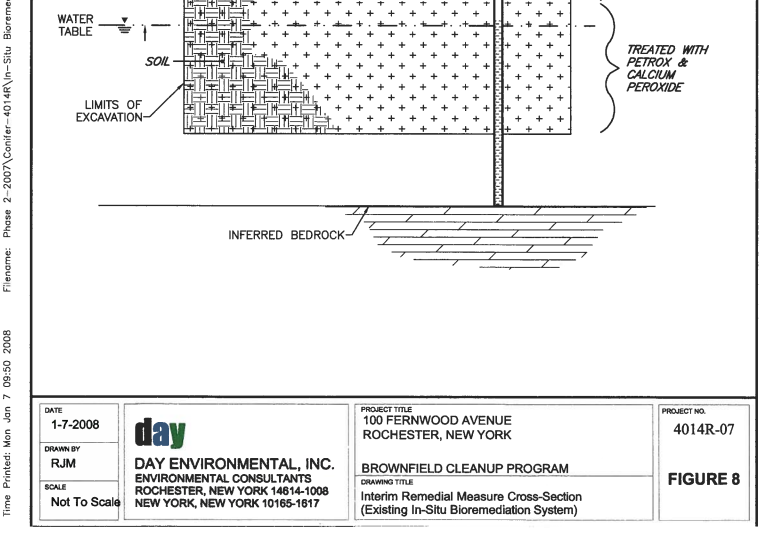
DAY ENVIRONMENTAL, INC. ENVIRONMENTAL CONSULTANTS ROCHESTER, NEW YORK 14614-1008 NEW YORK, NEW YORK 10165-1617

prowing TITLE InterIm Remedial Measure Plan - Record Drawing (Existing In-Situ Bioremediation System)

PROJECT NO. 4014R-07

FIGURE 7

Ref1:



EQUIP WITH WIND TURBINE

PVC WELL

TREATED AS BIOPILE AMEND WITH

FERTILIZER AND BULKING AGENT

EQUIP WITH GOOSENECK VENT

| 4" PERFORATED | PVC PIPE, TYP.

BARRIER MATERIAL (MINIMUM 10 MIL.

1' MIN.

POLYETHLENE SHEETING)

CLEAN SOIL

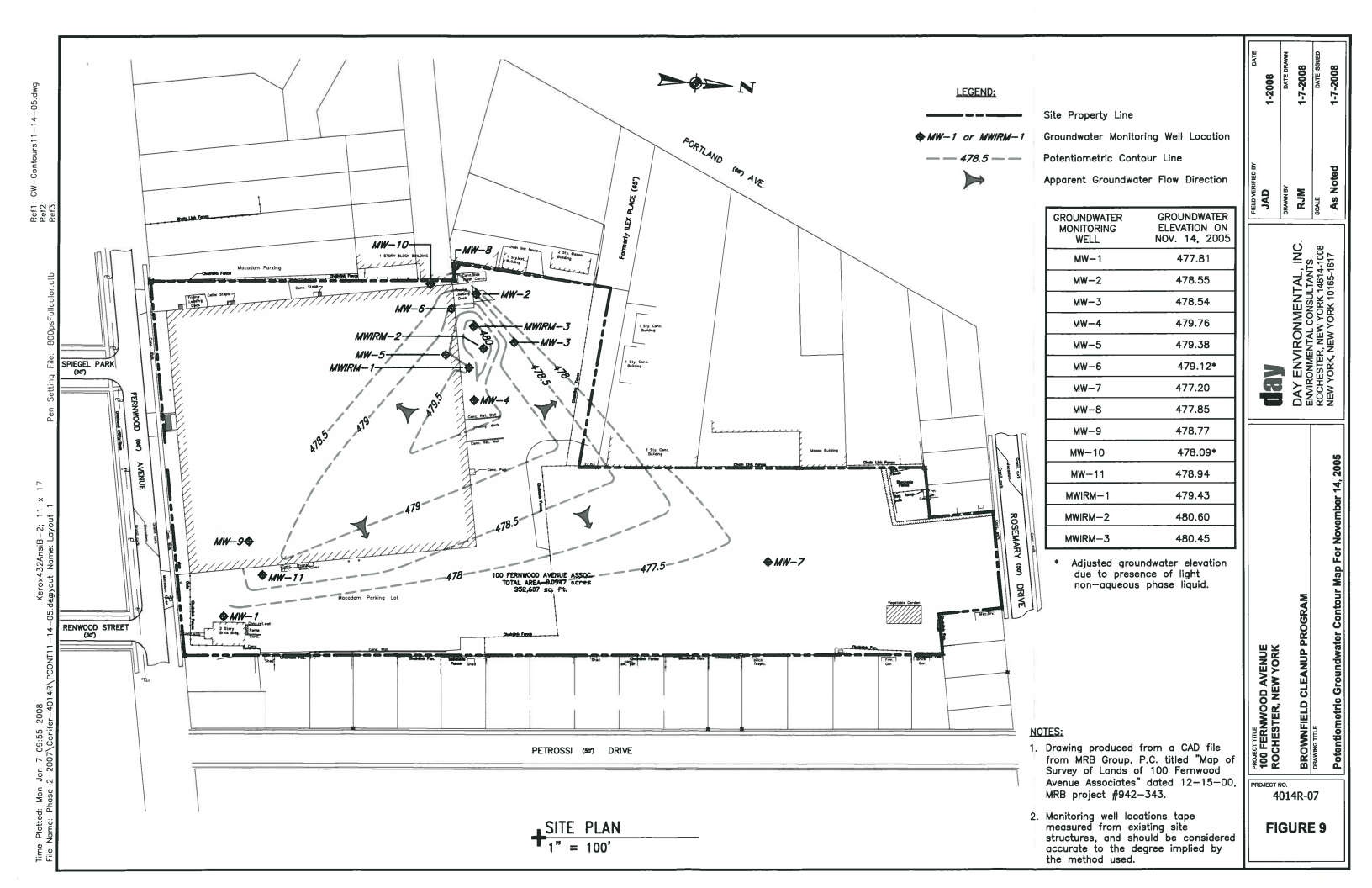
FILTER FABRIC, TYP.

6" #2 STONE, TYP.

2' MIN.

3'-4' OF SOIL

3'-4' OF SOIL



APPENDIX A

Alternative #2 Opinion of Probable Remedial Costs

LNAPL Recovery, MNA, ICs and ECs (Present Worth of Project Costs)

100 Fernwood Avenue Rochester, New York

Opinion of Probable Remedial Costs

Capital/Initial Costs	
Design, Work Plan, HASP, CPP, Fact Sheet	\$ 15,000
Install + Develop Four New LNAPL Recovery Wells	\$ 8,100
Institutional Controls (EE, SMP and 2 Periodic Certifications)	\$ 10,000
Confirmatory Sampling and Analysis from In-Situ Bioremediation System	\$ 9,600
FER, 12 MPRs, and up to five annual MNA reports	\$ 20,000
20% Contingency	\$ 12,540
Total	\$ 75,240
Operation/Maintenance/Annual Costs	
Years 1-2 LNAPL Recovery (\$7,700 X 2 yrs)	\$ 15,400
Years 1-2 Groundwater Monitoring (\$15,900 X 2 yrs)	\$ 31,800
Years 3-5 Groundwater Monitoring (\$8,000 X 3 yrs)	\$ 24,000
QA/QC and DUSR Associated with MNA (assumed on 5th yr data)	\$ 3,600
Total Operation/Maintenance/Annual Costs	\$ 74,800
Present Worth Cost	
Capital/Initial Costs	\$ 75,240
Years 1-2 LNAPL Recovery Present Worth (F=1.8594)	\$ 14,317
Years 1-2 Groundwater Monitoring Present Worth (F=1.8594)	\$ 29,564
Years 3-5 Groundwater Monitoring Present Worth (F=4.3295-1.8594)	\$ 19,761
QA/QC and DUSR Associated with MNA (F=0.7835)	\$ 2,821
Total Present Worth Cost	\$ 141,703

Assumptions

- 5 years at 5% discount factor
- Develop detailed remedial work plan for Site
- Develop and implement institutional controls
- F = Discount Factor of 5% at the nthyear of the project
- Conduct MNA groundwater sampling/analysis for 5 years (biannually at 8 wells for yrs 1-2, annually at 8 wells for yrs 3-5)
- Develop and submit necessary reports to document work completed
- Quotes provided by contractors, etc. to develop the cost estimate will be accurate at the time the work is conducted
- The costs provided are for comparative purposes only, and actual costs will vary
- LNAPL monitoring/recovery for two years, 12 events per year
- QA/QC samples and DUSR associated with MNA will only be conducted during the last monitoring event (i.e., in fifth year)
- Does not include cost of contingency application(s) to existing in-situ bioremediation system, if warranted
- Due to unknown redevelopment plans at this time, the Opinion of Probable Remedial Costs does not include cost to design, construct or operate engineering controls on existing or new buildings, if warranted
- Disposal of LNAPL and remediation-derived wastes will not exceed \$2,400
- Only includes cost to develop and submit two periodic certifications
- No contingency needs to be added to the groundwater monitoring costs and LNAPL recovery costs because it is assumed that the scope of the groundwater monitoring and LNAPL recovery will decrease with time (i.e., the number of monitoring wells that need to be sampled will decrease with time and/or the number of parameters that will be analyzed will decrease with time). If this is not the case, additional monitoring costs could be incurred due to inflation (e.g., increase in analytical costs and hourly rates).

APPENDIX B

Health and Safety Plan

HEALTH AND SAFETY PLAN

BROWNFIELD CLEANUP PROGRAM 100 FERNWOOD AVENUE **ROCHESTER, NEW YORK NYSDEC SITE ID C828119**

Prepared for:

Conifer Development, Inc. 183 East Main Street, 6th Floor Rochester, New York 14604

Prepared by:

Day Environmental, Inc. 40 Commercial Street

Rochester, New York 14614

Project No.:

4014R-07

Date:

January 2008

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ATTACHMENTS

Attachment 1 Figure 1- Route for Emergency Service

1.0 INTRODUCTION

This Health and Safety Plan (HASP) outlines the policies and procedures necessary to protect workers and the public from potential environmental hazards posed during remediation activities under the New York State Department of Environmental Protection (NYSDEC) Brownfield Cleanup Program (BCP). The subject property (Site) consists of eleven contiguous parcels totaling approximately 8.14 acres. The parcels are addressed as: 100 and 142 Fernwood Avenue; 31, 35 and 41 Rosemary Drive; and 25, 29, 33, 39, 43, 49, and 55 Ilex Place, City of Rochester, County of Monroe, New York (Tax account #s 106.27-1-5; 91.83-3-19; 91.83-3-20; 91.83-3-21; 106.27-1-87; 106.27-1-88; 106.27-1-89; 106.27-1-90; 106.27-1-91; 106.27-1-92; and 106.27-1-93). Figure 1 included in Attachment 1 depicts the general location of the Site. As outlined in this HASP, the remedial activities shall be conducted in a manner to minimize the probability of injury, accident, or incident occurrence.

Although the HASP focuses on the specific work activities planned for this Site, it must remain flexible due to the nature of this work. Conditions may change and unforeseen situations can arise that require deviations from the original HASP.

1.1 Site History/Overview

There are two buildings on the Site. The main building was constructed between 1926 and 1930 and is an approximately 120,000-square foot, one-story concrete block building that has a partial basement. The smaller building is an approximately 3,000-square foot, one-story brick building with a basement that was constructed between 1910 and 1922.

Elmer W. Davis, Inc currently uses the main building at the Site for the storage of insulation panels; however, it has no full time employees stationed on-site. The main building was originally constructed as Vogt Manufacturing Corporation, which manufactured auto trimmings (e.g., textile trimmings spinning and weaving). The main building was later converted for multitenant light industrial/commercial use. Former uses of the main building by tenants include: plastic products manufacturer, tool and die makers, machine shops, painters, printers, graphics companies, and sheet metal contractors. The building was vacant between approximately 2002 and 2004.

The smaller building was originally constructed as, and until recently was used as, a church. However, the smaller building has also been occupied in the past by light industrial/commercial tenants such as Empire Engraving Company (metal cutting allied services) and Phoenix Equipment Co.

The Site is located in an urban area that is serviced by a public water system. The Site and surrounding area are generally level. There are no surface water bodies at, or within a 0.5-mile radius of the Site. Surface water appears to flow off the Site via sheet flow toward adjoining streets to the north and to the south (i.e. Rosemary Drive and Fernwood Avenue), into the City of Rochester combined sewer system. Groundwater at the Site generally flows radially outward from an unpaved location north of the main building where five underground storage tanks (USTs) were removed and an in-situ bioremediation system was installed (see below for further information). This flow direction may be modified locally due to buried utilities, seasonal conditions, or other factors.

The Site is zoned industrial, and is located in a mixed-use urban area. The Site is bounded to the north and west by commercial, industrial and residential properties, and bounded to the south and east by residential properties.

A November 2000 Phase I Environmental Site Assessment (Phase I ESA) report identified the following recognized environmental conditions (RECs) at the Site:

- 1. Abandoned Underground Storage Tanks (USTs)
- 2. Confirmed Local Waste Site/Active New York State Department of Environmental Conservation (NYSDEC) Spill Site on Nearby Property
- 3. Active NYSDEC Spill on Adjoining Property
- 4. Suspect Asbestos-Containing Material (ACM) [Note: ACM is not addressed as part of this project.]
- 5. Closed NYSDEC Spill on Site
- 6. Transformers/Polychlorinated Biphenyl (PCB) Suspect Equipment
- 7. Historic Use of the Site

In addition to the RECs identified above, the NYSDEC requested that investigative work be included to evaluate whether environmental conditions have been impacted at loading docks equipped with hydraulic lifts. The NYSDEC also requested that a pipe chase in the floor of the main building be further evaluated, and that some limited surface and subsurface evaluation be included on the northern undeveloped portion of the Site.

A Remedial Investigation/Remedial Alternatives Analysis (RI/RAA) Report dated November 2006 as modified by a March 8, 2007 Addendum was prepared by Day Environmental, Inc. (DAY). Tasks performed as part of the remedial investigation to evaluate or address the RECs identified above included:

- Performing a passive soil gas survey as a screening tool to evaluate the presence of volatile organic compounds (VOCs) at the Site;
- Performing sampling and analysis of various media to evaluate whether PCBs were present at three pad-mounted transformers located east of the main building;
- Performing an evaluation of hydraulic lifts at three loading docks on the main building;
- Performing test pits and magnetic locator work to evaluate the potential presence of abandoned USTs;
- Permanently closing (i.e., removing) four USTs in accordance with applicable regulations;
- Designing and constructing an on-site in-situ bioremediation system within the former tank
 pit to treat contaminated soils that were displaced/disturbed during the UST closure work;
- Performing post-treatment monitoring to evaluate the effectiveness of the in-situ bioremediation system;

- Evaluating surface soil conditions;
- Evaluating subsurface soil conditions;
- Evaluating groundwater quality conditions and groundwater movement characteristics;
- Performing a vapor intrusion study to evaluate whether VOCs in soil or groundwater were volatilizing and impacting indoor air inside the smaller church building on the Site that is addressed as 142 Fernwood Avenue; and
- Evaluating environmental data for the adjoining former JML Optical, Inc. property located west of the Site.

The findings of the remedial investigation are summarized below:

- The hydraulic loading docks and pad-mounted transformers at the Site do not appear to have adversely impacted environmental conditions at the Site. In addition, evidence of environmental impact was not detected at test boring locations that were completed in proximity to a pipe chase located inside the main building. Therefore, it does not appear that this pipe chase has adversely impacted environmental conditions at the Site.
- Prior to the remedial investigation, a 15,000-gallon UST was removed from the Site. As part of the remedial investigation, one 8,000-gallon UST, two 2,000-gallon USTs and one 4,000-gallon UST were removed from the Site. These five USTs were located in the same general area north of the northwest corner of the main building.
- A primary area of soil and groundwater contamination, including the presence of a relatively thin layer (i.e., 0.37 foot or less) of LNAPL that is more limited in extent, was detected in proximity to the former UST locations near the northwest corner of the main building. This contamination generally consists of petroleum products and plasticizers that historically leaked from the former USTs. Based on field screening, analytical laboratory test results, and groundwater monitoring, it appears that this impact has migrated radially outward from the former UST area, including beneath the northwest corner of the main building. In addition, the length of the petroleum/plasticizer plume is estimated to be about 60 feet away from the former UST locations. Based on a review of Site data and environmental reports for the adjoining former JML Optical, Inc. property to the west, petroleum and plasticizer contamination attributable to the former UST locations at the Site appears to have also migrated from the Site via groundwater onto an estimated 1,375-square foot area of the adjoining former JML Optical, Inc. property.
- As an interim remedial measure (IRM), petroleum and plasticizer contaminated soils that were displaced during the UST removal work were amended with bioremediation products and placed back into the tank pit excavation as part of an in-situ bioremediation system. The analysis of post-treatment soil and groundwater samples indicate the in-situ bioremediation system is working, and contaminants have been reduced by approximately 40% (on average). The results of the post-treatment sampling and analytical laboratory testing indicated that contaminants were still present in soil and groundwater at concentrations exceeding SCG values.

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- Two of four surface soil samples collected from the northern undeveloped portion of the Site contained some polyaromatic hydrocarbon (PAH) semi-volatile organic compounds (SVOCs) above December 14, 2006 NYSDEC Part 375 (Environmental Restoration Programs) Track 2 Soil Cleanup Objectives (SCOs) for Restricted Residential Use. However, the concentrations of these SVOCs are comparable to other projects in the City of Rochester where surface soil data has been collected. As such, the NYSDEC concurs that the limited exceedances of the Restricted Residential Use SCOs in surface soil at the Site are attributable to the local geology or urban setting of the Site and are not significant.
- Chlorinated VOCs were detected in groundwater samples at some of the monitoring well locations. An on-site source of chlorinated VOCs that could result in contamination of the groundwater was not found during the soil and groundwater studies performed as part of this investigation. It is possible that the chlorinated VOCs are attributable to an off-site source(s) that has resulted in an area-wide groundwater condition. A review of environmental reports indicates a sump and a former degreaser area at the adjoining former JML Optical, Inc. property to the west, and also a nearby NYSDEC Inactive Hazardous Waste Disposal Site located southwest of the Site, could potentially be sources of the chlorinated VOCs that are present at the Site.
- A subsurface soil sample collected from a depth interval of 0-4' at test boring TB-4 contained three PAH SVOCs that exceeded Track 2 BCP SCOs for restricted residential use. This sample was collected beneath the floor of the main building and contained fill material that consisted of reworked soil with some cinders. The PAH SVOCs would presumably be limited in extent to the fill material, and can be a common component of cinders.

1.2 Planned Activities Covered by HASP

This HASP is intended to be used during this NYSDEC BCP project for remedial activities. Currently, identified activities include:

- Site preparation activities (e.g., put up NYSDEC Remediation Project sign);
- Installation of four wells for monitoring and recovery of light non-aqueous phase liquid (LNAPL);
- Groundwater sampling associated with monitored natural attenuation; and
- Miscellaneous on-site tasks that may arise during this project.

This HASP can be modified to cover other site activities as deemed appropriate. The owner of the property, its contractors, and other site workers will be responsible for the development and/or implementation of health and safety provisions associated with normal construction activities or site activities.

2.0 KEY PERSONNEL AND MANAGEMENT

The Project Manager (PM) and Site Safety Officer (SSO) are responsible for formulating and enforcing health and safety requirements, and implementing the HASP.

2.1 Project Manager

The PM has the overall responsibility for the project and will coordinate with the SSO to ensure that the goals of the remedial program are attained in a manner consistent with the HASP requirements.

2.2 Site Safety Officer

The SSO has responsibility for administering the HASP relative to site activities conducted by DAY personnel, and will be in the field full-time while site activities are in progress. The SSO's operational responsibilities will be monitoring, including personal and environmental monitoring, ensuring personal protective equipment maintenance, and assignment of protection levels. The SSO will be the main contact in any on-site emergency situation. The SSO will direct the safety aspects of field activities conducted by DAY personnel and will be responsible for stopping work when unacceptable health or safety risks exist. The SSO is responsible for ensuring that on-site personnel understand the safety requirements in this HASP.

2.3 Employee Safety Responsibility

Each employee is responsible for personal safety as well as the safety of others in the area. The employee will use the equipment provided in a safe and responsible manner as directed by the SSO.

2.4 Key Safety Personnel

The following individuals are anticipated to share responsibility for health and safety at the site.

Project Manager Jeffrey A. Danzinger

Site Safety Officer Mathew K. Dickinson, Glenn R. Miller, Kelly

A. Crandall, or Samuel C. Price

3.0 SAFETY RESPONSIBILITY

Contractors, consultants, state or local agencies, or other parties, and their employees, involved with this project will be responsible for their own safety while on-site. Their employees will be required to understand the information contained in this HASP, and must follow the recommendations that are made in this document. As an alternative, contractors, consultants, state or local agencies, or other parties, and their employees, involved with this project can utilize their own health and safety plan for this project as long as it is found acceptable to the New York State Department of Health (NYSDOH) and/or the MCDPH.

4.0 JOB HAZARD ANALYSIS

There are many hazards associated with remedial work on a site, and this HASP discusses some of the anticipated hazards for this Site. The hazards listed below deal specifically with those hazards associated with the management of potentially contaminated media (e.g., soil, groundwater, fill, etc.).

4.1 Chemical Hazards

Chemical substances can enter the unprotected body by inhalation, skin absorption, ingestion, or injection (i.e., a puncture wound, etc.). A contaminant can cause damage to the point of contact or can act systemically, causing a toxic effect at a part of the body distant from the point of initial contact.

A list of selected VOCs, SVOCs, and metals that have been detected at the Site and which exceed soil or groundwater standards, criteria and guidance (SCG) values or were detected in one or more media at high concentrations in relation to other constituents are presented below. This list also presents the Occupational Safety and Health Administration (OSHA) 8-hour Time-Weighted Average (TWA) Permissible Exposure Limits (PELs), the National Institute for Occupational Safety and Health (NIOSH) 8-hour TWA Recommended Exposure Limits (RELs), and the NIOSH Immediately Dangerous to Life or Health (IDLH) levels.

CONSTITUENT	OSHA PEL	NIOSH REL	IDLH
Benzene (Ca)	1 ppm	0.1 ppm	500 ppm
Toluene	200 ppm	100 ppm	500 ppm
Ethylbenzene	100 ppm	100 ppm	800 ppm
Xylenes	100 ppm	100 ppm	900 ppm
Trichloroethene (Ca)	100 ppm	25 ppm	1000 ppm
1,1,1-Trichloroethane	350 ppm	350 ppm	700 ppm
Naphthalene	10 ppm	10 ppm	250 ppm
1,1-Biphenyl (Ca)	0.2 ppm	0.2 ppm	15.8 ppm
Phenanthrene (Ca)	0.2 mg/m ³	0.1 mg/m ³	80 mg/m ³
Anthracene (Ca)	0.2 mg/m ³	0.1 mg/m ³	80 mg/m ³
Pyrene (Ca)	0.2 mg/m^3	0.1 mg/m ³	80 mg/m ³
Bis(2-ethylhexyl)phthalate	5 mg/m ³	5 mg/m ³	4000 mg/m ³
Chrysene (Ca)	0.2 mg/m ³	0.1 mg/m ³	80 mg/m ³
Benzo(b)fluoranthene (Ca)	0.2 mg/m ³	0.1 mg/m ³	80 mg/m ³
Benzo(a)pyrene (Ca)	0.2 mg/m^3	0.1 mg/m ³	80 mg/m ³
Gamma-Chlordane (Ca)	0.5 mg/m ³	0.5 mg/m ³	100 mg/m ³
Antimony	0.5 mg/m ³	0.5 mg/m ³	50 mg/m ³
Iron (dust-fume)	10 mg/m ³	5 mg/m ³	2,500 mg/m ³
Magnesium (fume)	15 mg/m ³	10 mg/m ³	750 mg/m ³
Manganese (fume)	5 mg/m ³ (ceiling)	1 mg/m ³ (15-min)	500 mg/m ³

Ca = Potential carcinogen

The potential routes of exposure for these analytes and chemicals include inhalation, ingestion, skin absorption and/or skin/eye contact. The potential for exposure through any one of these routes will depend on the activity conducted. The most likely routes of exposure for the activities that are performed during remedial activities at the Site include inhalation and skin/eye contact.

4.2 Physical Hazards

There are physical hazards associated with this project, which might compound the chemical hazards. Hazard identification, training, adherence to the planned remedial measures, and careful housekeeping can prevent many problems or accidents arising from physical hazards. Potential physical hazards associated with this project and suggested preventative measures include:

- <u>Slip/Trip/Fall Hazards</u> Some areas may have wet surfaces that will greatly increase the
 possibility of inadvertent slips. Caution must be exercised when using steps and stairs due to
 slippery surfaces in conjunction with the fall hazard. Good housekeeping practices are essential
 to minimize the trip hazards.
- <u>Small Quantity Flammable Liquids</u> Small quantities of flammable liquids will be stored in "safety" cans and labeled according to contents.
- <u>Electrical Hazards</u> Electrical devices and equipment shall be de-energized prior to working near them. All extension cords will be kept out of water, protected from crushing, and inspected regularly to ensure structural integrity. Temporary electrical circuits will be protected with ground fault circuit interrupters. Only qualified electricians are authorized to work on electrical circuits. Heavy equipment (e.g., excavator, backhoe, drill rig) shall not be operated within 10 feet of high voltage lines, unless proper protection from the high voltage lines is provided by the appropriate utility company.
- <u>Noise</u> Work around large equipment often creates excessive noise. The effects of noise can include:
 - Workers being startled, annoyed, or distracted.
 - Physical damage to the ear resulting in pain, or temporary and/or permanent hearing loss.
 - Communication interference that may increase potential hazards due to the inability to warn of danger and proper safety precautions to be taken.

Proper hearing protection will be worn as deemed necessary. In general, feasible administrative or engineering controls shall be utilized when on-site personnel are subjected to noise exceeding an 8-hour TWA sound level of 90 dBA (decibels on the A-weighted scale). In addition, whenever employee noise exposures equal or exceed an 8-hour TWA sound level of 85 dBA, employers shall administer a continuing, effective hearing conservation program as described in the OSHA Regulation 29 CFR Part 1910.95.

• <u>Heavy Equipment</u> - Each morning before start-up, heavy equipment will be inspected to ensure safety equipment and devices are operational and ready for immediate use.

• <u>Subsurface and Overhead Hazards</u> - Before any excavation activity, efforts will be made to determine whether underground utilities and potential overhead hazards will be encountered. Underground utility clearance must be obtained prior to subsurface work.

4.3 Environmental Hazards

Environmental factors such as weather, wild animals, insects, and irritant plants can pose a hazard when performing outdoor tasks. The SSO shall make every reasonable effort to alleviate these hazards should they arise.

4.3.1 Heat Stress

The combination of warm ambient temperature and protective clothing increases the potential for heat stress. In particular:

- Heat rash
- Heat cramps
- Heat exhaustion
- Heat stroke

Site workers will be encouraged to increase consumption of water or electrolyte-containing beverages such as Gatorade[®] when the potential for heat stress exists. In addition, workers are encouraged to take rests whenever they feel any adverse effects that may be heat-related. The frequency of breaks may need to be increased upon worker recommendation to the SSO.

4.3.2 Exposure to Cold

With outdoor work in the winter months, the potential exists for hypothermia and frostbite. Protective clothing greatly reduces the possibility of hypothermia in workers. However, personnel will be instructed to wear warm clothing and to stop work to obtain more clothing if they become too cold. Employees will also be advised to change into dry clothes if their clothing becomes wet from perspiration or from exposure to precipitation.

5.0 SITE CONTROLS

To prevent migration of contamination caused through tracking by personnel or equipment, work areas and personal protective equipment staging/decontamination areas will be specified prior to beginning operations.

5.1 Site Zones

In the area where contaminated materials present the potential for worker exposure (work zone), personnel entering the area must wear the mandated level of protection for the area. A "transition zone" shall be established where personnel can begin and complete personal and equipment decontamination procedures. This can reduce potential off-site migration of contaminated media. Contaminated equipment or clothing will not be allowed outside the transition zone (e.g., on clean portions of the Site) unless properly containerized for disposal. Operational support facilities will be located outside the transition zone (i.e., in a "support zone"), and normal work clothing and support equipment are appropriate in this area. If possible, the support zone should be located upwind of the work zone and transition zone.

5.2 General

The following items will be requirements to protect the health and safety of workers during implementation of activities that disturb contaminated material.

- Eating, drinking, chewing gum or tobacco, smoking, or any practice that increases the probability of hand to mouth transfer and ingestion of contamination shall not occur in the work zone and/or transition zone during disturbance of contaminated material.
- Personnel admitted in the work zone shall be properly trained in health and safety techniques and equipment usage.
- No personnel shall be admitted in the work zone without the proper safety equipment.
- Proper decontamination procedures shall be followed before leaving the Site.

6.0 PROTECTIVE EQUIPMENT

This section addresses the various levels of personal protective equipment (PPE), which are or may be required at this job site. Personnel entering the work zone and transition zone shall be trained in the use of the anticipated PPE to be utilized.

6.1 Anticipated Protection Levels

TASK	PROTECTION LEVEL	COMMENTS/MODIFICATIONS
Site mobilization	D	
Site preparation	D	
Extrusive work (e.g., surveying, etc.)	D	
Intrusive work (e.g., well installation, collecting samples, etc.)	C/Modified D/D	Based on air monitoring, and SSO discretion
Support zone	D	
Site breakdown and demobilization	D	

It is anticipated that work conducted as part of this project will be performed in Level D or modified Level D PPE. If conditions are encountered that require Level A or Level B PPE, the work will immediately be stopped. The appropriate government agencies (e.g., NYSDEC, NYSDOH, MCDPH, etc.) will be notified and the proper health and safety measures will be implemented (e.g., develop and implement engineering controls, upgrade in PPE, etc.).

6.2 Protection Level Descriptions

This section lists the minimum requirements for each protection level. Modifications to these requirements can be made upon approval of the SSO. If Level A, Level B, and/or Level C PPE is required, Site personnel that enter the work zone and/or transition zone must be properly trained and certified in the use of those levels of PPE.

6.2.1 Level D

Level D consists of the following:

- Safety glasses
- Hard hat when working with heavy equipment
- Steel-toed or composite-toed work boots
- Protective gloves during sampling or handling of potentially contaminated media
- Work clothing as prescribed by weather

6.2.2 Modified Level D

Modified Level D consists of the following:

- Safety glasses with side shields
- Hard hat when working with heavy equipment
- Steel-toed or composite-toed work boots
- Work gloves
- Outer protective wear, such as Tyvek coverall [Tyveks (Sarans) and polyvinyl chloride (PVC) acid gear will be required when workers have a potential to be exposed to contaminated liquids and/or particulates].

6.2.3 Level C

Level C consists of the following:

- Air-purifying respirator with appropriate cartridges
- Outer protective wear, such as Tyvek coverall [Tyveks (Sarans) and PVC acid gear will be required when workers have a potential to be exposed to contaminated liquids and/or particulates].
- Hard hat when working with heavy equipment
- Steel-toed or composite-toed work boots
- Nitrile, neoprene, or PVC overboots, if appropriate
- Nitrile, neoprene, or PVC gloves, if appropriate
- Face shield (when projectiles or splashes pose a hazard)

6.2.4 Level B

Level B protection consists of the items required for Level C protection with the exception that an air-supplied respirator is used in place of the air-purifying respirator. Level B PPE is not anticipated to be required during this project. If the need for level B PPE becomes evident, site remediation activities will be stopped until site conditions are further evaluated, and any necessary modifications to the HASP have been approved by the PM and SSO. Subsequently, the appropriate safety measures (including Level B PPE) must be implemented prior to commencing site activities.

6.2.5 Level A

Level A protection consists of the items required for Level B protection with the addition of a fully-encapsulating, vapor-proof suit capable of maintaining positive pressure. Level A PPE is not anticipated to be required during this project. If the need for level A PPE becomes evident, site remediation activities will be stopped until site conditions are further evaluated, and any necessary modifications to the HASP have been approved by the PM and SSO. Subsequently, the appropriate safety measures (including Level A PPE) must be implemented prior to commencing site activities.

6.3 Respiratory Protection

Any respirator used will meet the requirements of the OSHA 29 CFR 1910.134. Both the respirator and cartridges specified shall be fit-tested prior to use in accordance with OSHA regulations (29 CFR 1910). Air purifying respirators shall not be worn if contaminant levels exceed designated use concentrations. The workers will wear respirators with approval for: organic vapors <1,000 ppm; and dusts, fumes and mists with a TWA < 0.05 mg/m³.

No personnel who have facial hair, which interferes with respirator sealing surface, will be permitted to wear a respirator and will not be permitted to work in areas requiring respirator use.

Only workers who have been certified by a physician as being physically capable of respirator usage shall be issued a respirator. Personnel unable to pass a respiratory fit test or without medical clearance for respirator use will not be permitted to enter or work in areas that require respirator protection.

7.0 DECONTAMINATION PROCEDURES

This section describes the procedures necessary to ensure that both personnel and equipment are free from contamination when they leave the work site.

7.1 Personnel Decontamination

Personnel involved with activities that involve disturbing contaminated media will follow the decontamination procedures described herein to ensure that material which workers may have contacted in the work zone and/or transition zone does not result in personal exposure and is not spread to clean areas of the Site. This sequence describes the general decontamination procedure. The specific stages can vary depending on the Site, the task, and the protection level, etc.

- 1. Leave work zone and go to transition zone
- 2. Remove soil/debris from boots and gloves
- 3. Remove boots
- 4. Remove gloves
- 5. Remove Tyvek suit and discard, if applicable
- 6. Remove and wash respirator, if applicable
- 7. Go to support zone

7.2 Equipment Decontamination

Contaminated equipment shall be decontaminated in the transition zone before leaving the Site. Decontamination procedures can vary depending upon the contaminant involved, but may include sweeping, wiping, scraping, hosing, or steam cleaning the exterior of the equipment. Personnel performing this task will wear the proper PPE.

7.3 Disposal

Disposable clothing will be disposed in accordance with applicable regulations. Liquids (e.g., decontamination water, etc.) or solids (e.g., soil) generated by remedial activities will be disposed in accordance with applicable regulations.

8.0 AIR MONITORING

Air monitoring will be conducted in order to determine airborne particulate and contamination levels. This ensures that respiratory protection is adequate to protect personnel against the chemicals that are encountered and that chemical contaminants are not migrating off-site. Additional air monitoring may be conducted at the discretion of the SSO. Readings will be recorded and be available for review.

The following chart describes the direct reading instrumentation that will be utilized and appropriate action levels.

Monitoring Device	Action level	Response/Level of PPE	
	< 1 ppm in breathing zone, sustained 5 minutes	Level D	
PID Volatile Organic Compound Meter	1-25 ppm in breathing zone, sustained 5 minutes	Level C	
	26-250 ppm in breathing zone, sustained 5 minutes	Level B, Stop work, evaluate the use of engineering controls	
	>250 ppm in breathing zone	Level A, Stop work, evaluate the use of engineering controls	
RTAM Particulate Meter	≤ 150 µg/m³ over an integrated period not to exceed 15 minutes.	Continue working	
	> 150 μg/m ³	Cease work, implement dust suppression, change in way work performed, etc. If levels can not be brought below 150 μg/m³, then upgrade PPE to Level C.	

8.1 Particulate Monitoring

During activities where contaminated materials (e.g., fill) may be disturbed, air monitoring will include real-time monitoring for particulates using a real-time aerosol monitor (RTAM) particulate meter at the perimeter of the work zone in accordance with the 1989 NYSDEC Technical and Administrative Guidance Memorandum (TAGM) #4031 entitled, "Fugitive Dust Suppression and Particulate Monitoring Program at Inactive Hazardous Waste Sites." The TAGM uses an action level of 150 μ g/m³ (0.15 mg/m³) over an integrated period not to exceed 15 minutes. If the action level is exceeded, or if visible dust is encountered, then work shall be discontinued until corrective actions are implemented. Corrective actions may include dust suppression, change in the way work is performed, and/or upgrade of personal protective equipment.

8.2 Volatile Organic Compound Monitoring

During activities where contaminated materials may be disturbed, a photoionization detector (PID) will be used to monitor total VOCs in the ambient air. The PID will prove useful as a direct reading instrument to aid in determining if current respiratory protection is adequate or needs to be upgraded. The SSO will take measurements before operations begin in an area to determine the amount of VOCs naturally occurring in the air. This is referred to as a background level. Levels of VOCs will periodically be measured in the air at active work sites, and at the transition zone when levels are detected above background in the work zone.

8.3 Community Air Monitoring Plan

This Community Air Monitoring Plan (CAMP) includes real-time monitoring for VOCs and particulates (i.e., dust) at the downwind perimeter of each designated work area when activities with the potential to release VOCs or dust are in progress at the Site. This CAMP is based on the NYSDOH Generic CAMP included as Appendix 1A of the NYSDEC document titled "Draft DER-10, Technical Guidance for Site Investigation and Remediation" dated December 2002. The CAMP is not intended for use in establishing action levels for worker respiratory protection. Rather, its intent is to provide a measure of protection for the downwind community (i.e., off-site receptors including residences and businesses and on-site workers not directly involved with the subject work activities) from potential airborne contaminant releases as a direct result of project activities. The action levels specified herein require increased monitoring, corrective actions to abate emissions, and/or work shutdown. Additionally, the CAMP helps to confirm that work activities did not spread contamination off-site through the air. Reliance on the CAMP should not preclude simple, common sense measures to keep VOCs, dust, and odors at a minimum around the work areas.

<u>Continuous monitoring</u> will be conducted during ground intrusive activities. Ground intrusive activities include, but are not limited to, soil/waste excavation and handling, test pitting or trenching, advancement/installation of test borings or monitoring wells, etc.

Periodic monitoring for VOCs will be conducted during non-intrusive activities such as the collection of soil and sediment samples or the collection of groundwater samples from existing monitoring wells. Periodic monitoring during sample collection might reasonably consist of taking a reading upon arrival at a sample location, monitoring while opening a well cap or overturning soil, monitoring during well baling/purging, and taking a reading prior to leaving a sample location. In some instances, depending upon the proximity of potentially exposed individuals, continuous monitoring may be required during sampling activities. Examples of such situations include groundwater sampling at wells on the curb of a busy urban street, in the midst of a public park, or adjacent to a school or residence.

8.3.1 VOC Monitoring, Response Levels, and Actions

VOCs must be monitored at the downwind perimeter of the immediate work area (i.e., the work zone) on a continuous basis or as otherwise specified. Upwind concentrations should be measured at the start of each workday and periodically thereafter to establish background conditions. The monitoring work should be performed using equipment appropriate to measure

the types of contaminants known or suspected to be present. The equipment should be calibrated at least daily for the contaminant(s) of concern or for an appropriate surrogate. The equipment should be capable of calculating 15-minute running average concentrations, which will be compared to the levels specified below.

- If the ambient air concentration of total organic vapors at the downwind perimeter of the work area or exclusion zone exceeds 5 ppm above background for the 15-minute average, work activities must be temporarily halted and monitoring must be continued. If the total organic vapor level readily decreases (per instantaneous readings) below 5 ppm over background, work activities can resume with continued monitoring.
- If total organic vapor levels at the downwind perimeter of the work area or exclusion zone persist at levels in excess of 5 ppm over background but less than 25 ppm, work activities must be halted, the source or vapors identified, corrective actions taken to abate emissions, and monitoring continued. After these steps, work activities can resume provided that the total organic vapor level 200 feet downwind of the exclusion zone or half the distance to the nearest potential receptor or residential/commercial structure, whichever is less (but in no case less than 20 feet), is below 5 ppm over background for the 15-minute average.
- If the organic vapor level is above 25 ppm at the perimeter of the work area, activities must be shutdown.

The 15-minute readings must be recorded and made available for NYSDEC and NYSDOH personnel to review. Instantaneous readings, if any, used for decision purposes should also be recorded.

8.3.2 Particulate Monitoring, Response Levels, and Actions

Particulate concentrations should be monitored continuously at the upwind and downwind perimeters of the work zone at temporary particulate monitoring stations. The particulate monitoring should be performed using real-time monitoring equipment capable of measuring particulate matter less than 10 micrometers in size (PM-10) and capable of integrating over a period of 15 minutes (or less) for comparison to the airborne particulate action level. The equipment must be equipped with an audible alarm to indicate exceedance of the action level. In addition, fugitive dust migration should be visually assessed during work activities.

- If the downwind PM-10 particulate level is 100 micrograms per cubic meter (μg/m³) greater than background (upwind perimeter) for the 15-minute period or if airborne dust is observed leaving the work area, then dust suppression techniques must be employed. Work may continue with dust suppression techniques provided that downwind PM-10 particulate levels do not exceed 150 μg/m³ above the upwind level and provided that no visible dust is migrating from the work area.
- If, after implementation of dust suppression techniques, downwind PM-10 particulate levels are greater than 150 μg/m³ above the upwind level, work must be stopped and a re-evaluation of activities initiated. Work can resume provided that dust suppression measures and other controls are successful in reducing the downwind PM-10 particulate concentration to within 150 μg/m³ of the upwind level and in preventing visible dust migration.

Readings will be recorded and made available for NYSDEC and NYSDOH personnel to review.

9.0 EMERGENCY RESPONSE

To provide first-line assistance to field personnel in the case of illness or injury, the following items will be made immediately available on the Site:

- First-aid kit;
- Portable emergency eye wash; and
- Supply of clean water.

9.1 Emergency Telephone Numbers

The following telephone numbers are listed in case there is an emergency at the Site:

Fire/Police Department:	011
Fire/Police Department:	911

Poison Control Center: (800) 222-1222

NYSDEC

Greg MacLean (585) 226-5356 Spills (585) 226-2466

NYSDOH

Debra McNaughton (585) 423-8069

MCDPH

Joe Albert (585) 753-5904

CONIFER DEVELOPMENT, INC.

Eileen Broderick (585) 324-0503

DAY ENVIRONMENTAL, INC.

Jeff Danzinger (585) 454-0210 x114 Ray Kampff (585) 454-0210 x108

Nearest Hospital Rochester General Hospital

1425 Portland Avenue Rochester, NY 14621 (585) 922-4000 (Main)

(585) 922-2000 (Emergency Department)

Directions to the Hospital (refer Figure 1): Turn right (west) onto Fernwood Avenue and

travel approximately 0.18 miles. Turn right (north) onto Portland Avenue and travel approximately 1.17 miles. Turn left (west) into Rochester General Hospital and follow

signs to the Emergency Department.

9.2 Evacuation

A log of each individual entering and leaving the Site will be kept for emergency accounting practices. Although unlikely, it is possible that a site emergency could require evacuating personnel from the site. If required, the SSO will give the appropriate signal for site evacuation (i.e., hand signals, alarms, etc.).

All personnel shall exit the site and shall congregate in an area designated by the SSO. The SSO shall ensure that all personnel are accounted for. If someone is missing, the SSO will alert emergency personnel. The appropriate government agencies will be notified as soon as possible regarding the evacuation, and any necessary measures that may be required to mitigate the reason for the evacuation.

9.3 Medical Emergency

In the event of a medical emergency involving illness or injury to one of the on-site personnel, the Site should be shut down and immediately secured. The appropriate government agencies should be notified immediately. The area in which the injury or illness occurred shall not be entered until the cause of the illness or injury is known. The nature of injury or illness shall be assessed. If the victim appears to be critically injured, administer first aid and/or cardio-pulmonary resuscitation (CPR) as needed. Instantaneous real-time air monitoring shall be done in accordance with air monitoring outlined in Section 8.0 of this HASP.

9.4 Contamination Emergency

It is unlikely that a contamination emergency will occur; however, if such an emergency does occur, the Site shall be shut down and immediately secured. If an emergency rescue is needed, notify Police, Fire Department and Emergency Medical Service (EMS) Units immediately. Advise them of the situation and request an expedient response. The appropriate government agencies shall be notified immediately. The area in which the contamination occurred shall not be entered until the arrival of trained personnel who are properly equipped with the appropriate PPE and monitoring instrumentation as outlined in Section 8.0 of this HASP.

9.5 Fire Emergency

In the event of a fire on-site, the Site shall be shut down and immediately secured. The area in which the fire occurred shall not be entered until the cause can be determined. All non-essential site personnel shall be evacuated from the site to a safe, secure area. Notify the Fire Department immediately. Advise the Fire Department of the situation and the identification of any hazardous materials involved. The appropriate government agencies shall be notified as soon as possible.

The four classes of fire along with their constituents are as follows:

Class A: Wood, cloth, paper, rubber, many plastics, and ordinary combustible

materials.

Class B: Flammable liquids, gases and greases.

Class C: Energized electrical equipment.

Class D: Combustible metals such as magnesium, titanium, sodium, potassium.

Small fires on-site may be actively extinguished; however, extreme care shall be taken while in this operation. Approaches to the fire shall be done from the upwind side if possible. Distance from on-site personnel to the fire shall be close enough to ensure proper application of the extinguishing material, but far enough away to ensure that the personnel are safe. The proper extinguisher shall be utilized for the Class(s) of fire present on the site. If possible, the fuel source shall be cut off or separated from the fire. Care must be taken when performing operations involving the shut-off values and manifolds, if present.

Examples of proper extinguishing agent as follows:

Class A: Water

Water with 1% AFFF Foam (Wet Water)
Water with 6% AFFF or Fluorprotein Foam

ABC Dry Chemical

Class B: ABC Dry Chemical

Purple K

Carbon Dioxide

Water with 6% AFFF Foam

Class C: ABC Dry Chemical

Carbon Dioxide

Class D: Metal-X Dry Powder

No attempt shall be made against large fires. These shall be handled by the Fire Department.

9.6 Spill or Air Release

In the event of spills or air releases of hazardous materials on-site, the Site shall be shut down and immediately secured. The area in which the spills or releases occurred shall not be entered until the cause can be determined and site safety can be evaluated. Non-essential site personnel shall be evacuated from the Site to a safe and secure area. The appropriate government agencies shall be notified as soon as possible. The spilled or released materials shall be immediately identified and appropriate containment measures shall be implemented, if possible. Real-time air monitoring shall be implemented as outlined in Section 8.0 of this HASP. If the materials are unknown, Level B protection is mandatory. Samples of the materials shall be acquired to facilitate identification.

9.7 Locating Containerized Waste and/or Underground Storage Tanks

In the event that unanticipated containerized waste (e.g., drums) and/or USTs are encountered/discovered during remedial activities, the Site shall be shut down and immediately secured. The area where unanticipated containerized wastes and/or tanks are discovered shall not be

entered until site safety can be evaluated. Non-essential Site personnel shall be evacuated from the Site to a safe and secure area. The appropriate government agencies shall be notified as soon as possible. The SSO shall monitor the area as outlined in Section 8.0 of this HASP.

Prior to any handling, containers that are encountered will be visually assessed by the SSO to gain as much information as possible about their contents. As a precautionary measure, personnel shall assume that unlabelled containers and/or tanks contain hazardous materials until their contents are characterized. To the extent possible based upon the nature of the containers encountered, actions may be taken to stabilize the area and prevent migration (e.g., placement of berms, etc.). Subsequent to initial visual assessment and any required stabilization, properly trained personnel will sample, test, remove, and dispose of any containers and/or tanks, and their contents. After visual assessment and air monitoring, if the material remains unknown, Level B protection is mandatory.

10.0 ABBREVIATIONS

BCP Brownfield Cleanup Program

CAMP Community Air Monitoring Program
CPR Cardio-Pulmonary Resuscitation

DAY Day Environmental, Inc.

dBA Decibels on the A-Weighted Scale DNAPL Dense Non-Aqueous Phase Liquid

EMS Emergency Medical Service HASP Health and Safety Plan

IDLH Immediately Dangerous to Life or Heath

LNAPL Light Non Aqueous Phase Liquid

MCDPH Monroe County Department of Public Health

mg/m³ Milligram Per Meter Cubed

NIOSH National Institute for Occupational Safety and Health

NYSDEC New York State Department of Environmental Conservation

NYSDOH New York State Department of Health

OSHA Occupational Safety and Health Administration

PAH Polyaromatic Hydrocarbon PEL Permissible Exposure Limit PID Photoionization Detector

PM Project Manager

PM-10 Particulate Matter Less Than 10 Micrometers In Diameter

PPE Personal Protection Equipment

ppm Parts Per Million PVC Polyvinyl Chloride

REC Recognized Environmental Condition

REL Recommended Exposure Limit

RI/RAA Remedial Investigation/Remedial Alternatives Analysis

RTAM Real-Time Aerosol Monitor SCG Standards, Criteria and Guidance

SCO Soil Cleanup Objective SSO Site Safety Officer

SVOC Semi-Volatile Organic Compound

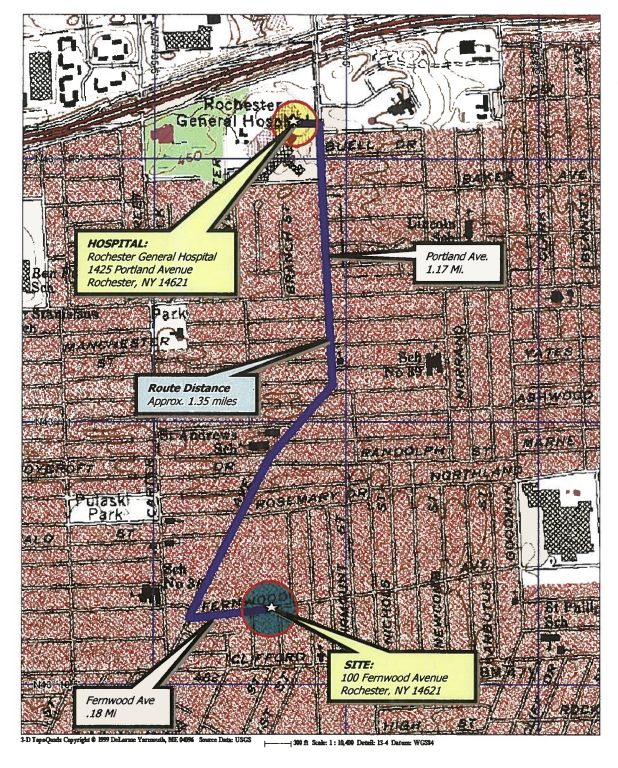
TAGM Technical and Administrative Guidance Memorandum

TOGS Technical and Operational Guidance Series

TWA Time-Weighted Average
 μg/m³ Micrograms Per Meter Cubed
 UST Underground Storage Tank
 VOC Volatile Organic Compound

ATTACHMENT 1

Figure 1- Route for Emergency Services



Drawing Produced From: 3-D TopoQuads, DeLorme Map Co., referencing USGS quad maps Rochester East (NY) 1995.

12-04-2007

DRAWN BY **RJM**

SCALE As Shown

DAY ENVIRONMENTAL, INC. **ENVIRONMENTAL CONSULTANTS** ROCHESTER, NEW YORK 14614-1008

PROJECT TITLE

100 FERNWOOD AVENUE ROCHESTER, NEW YORK

HEALTH AND SAFETY PLAN

DRAWING TITLE
ROUTE FOR EMERGENCY SERVICE

PROJECT NO.

4014R-07

FIGURE 1

APPENDIX C

Quality Assurance Project Plan

QUALITY ASSURANCE PROJECT PLAN

This project-specific Quality Assurance Project Plan (QAPP) was prepared in accordance with Section 2.2 of the New York State Department of Environmental Conservation (NYSDEC) draft DER-10 document for NYSDEC Site ID C828119 (Site). The QAPP provides quality assurance/quality control (QA/QC) protocols and guidance that are to be followed when implementing the remedy for the Site to ensure that data of a known and acceptable precision and accuracy are generated. The QAPP also provides a summary of the remedial project, identifies personnel responsibilities, and provides procedures to be used during sampling of environmental media, other field activities, and the analytical laboratory testing of samples. The components of the QAPP are provided herein.

1.0 Project Scope and Project Goals

The QAPP applies to the aspects of the project associated with implementing a physical remedy and the collection of field data, the collection and analytical laboratory testing of field samples and QA/QC samples, and the evaluation of the quality of the data that is generated. Specifically, the physical remediation will include: light non-aqueous phase liquid (LNAPL) monitoring, recovery and off-site disposal or recycling; confirmatory soil and groundwater sampling and analysis at the existing in-situ bioremediation system; and monitored natural attenuation that involves analytical laboratory testing of groundwater samples and the collection of groundwater quality measurements. After review of the data, the site management plan (SMP) that is developed for the Site may include a contingency that involves additional application of remediation products at the existing in-situ bioremediation system including subsequent additional confirmatory soil or groundwater sampling and analysis.

2.0 Project/Task Organization

Project organization and tentative personnel to implement the work are outlined in this section of the QAPP.

Principal in Charge

The Principal in Charge is responsible for review of project documents and ensuring the project is completed in accordance with applicable work plans. Mr. David D. Day, P.E., a Day Environmental, Inc. (DAY) representative, will serve as the Principle-in-Charge on this project.

Project Manager

The Project Manager has the overall responsibility for implementing the project and ensuring that the project meets the objectives and quality standards as presented in this QAPP. Mr. Jeffrey A. Danzinger, a DAY representative, will serve as the Project Manager on this project, and will serve as the primary point of contact and control for the project.

Quality Assurance Officer

The Quality Assurance Officer is responsible for QA/QC on this project. The Quality Assurance Officer's responsibilities on this project are not as a project manager or task manager involved with project productivity or profitability as job performance criteria. Ms. Hope Kilmer, a DAY representative, will serve as the Quality Assurance Officer on this project. The Quality Assurance Officer may conduct audits of the operations at the site to ensure that work is being performed in accordance with the QAPP.

Technical Staff, Subconsultants and Subcontractors

DAY's technical staff for this project consists of experienced professionals (e.g., professional engineers, engineers-in-training, scientists, technicians, etc.) that possess the qualifications necessary to effectively and efficiently complete the project tasks. The technical staff will be used to gather and analyze data, prepare various project documentation, etc. Subconsultants and subcontractors used on this project will consist of firms and companies with experience in the services to be provided.

Analytical Laboratory

It is anticipated that Mitkem Laboratories, a Division of Spectrum Analytical, Inc., with facilities at 175 Metro Center Boulevard, Warwick, Rhode Island will be retained to complete the required analytical laboratory testing of samples as part of this project. Mitkem is a New York State Department of Health (NYSDOH) Environmental Laboratory Approval Program (ELAP)-certified analytical laboratory (ELAP ID11522).

Dr. Kin S. Chiu is the Laboratory Director for Mitkem. The laboratory director is responsible for analytical work, and works in conjunction with the Laboratory Manager and QA unit regarding QA and chain-of-custody requirements.

Ms. Agnes Ng of Mitkem will act as the Laboratory Manager on this remediation project. The Laboratory Manager will report to the laboratory director and work in conjunction with the laboratory QA unit regarding QA elements of specific sample analyses tasks.

3.0 Sampling Procedures

This section of the QAPP provides the protocols for collection of confirmatory soil samples from test borings, installation of groundwater wells for monitoring and recovery of LNAPL, well development, and collection of groundwater samples from monitoring wells as part of the remediation project.

Collection of Confirmatory Soil Samples from Test Borings

A subcontractor will be retained to provide vehicle-mounted direct-push soil sampling equipment to advance the test borings. However, if it is determined in the field that such equipment cannot adequately be advanced through the existing overburden soils, then the NYSDEC will be consulted to approve any modifications to the drilling program (i.e., use of rotary drill-rig, etc.).

Based on the results of the previous remedial investigation, it is anticipated that the test borings will be advanced to depths up to approximately 20 feet below the ground surface. Sampling equipment will be used to collect soil samples in two-foot or four-foot intervals throughout the entire depth of the test borings. The soil samples will be collected in new disposable plastic liners.

The recovered soil samples will be visually examined by a DAY representative for evidence of suspect contamination (e.g., staining, unusual odors) and screened with a photoionization detector (PID). Portions of the samples will be placed in containers for possible analytical laboratory testing. Different portions of the soil samples will be placed in sealable Ziploc®-type plastic baggies, and will be field screened the same day the samples are collected. These samples will be agitated and homogenized for at least 30 seconds and allowed to equilibrate for at least three minutes. The ambient headspace air inside the baggies above each soil sample will be screened for total volatile organic compound (VOC) vapors with a RAE Systems MiniRAE 2000 PID equipped with a 10.6 eV lamp (or equivalent). The sampling port for the PID will be placed in the ambient air headspace

inside each baggie by opening a corner of the "locked" portion of the baggie. The PID will monitor air inside each baggie for a period of at least 15 seconds, and the peak readings measured will be recorded on a log sheet or log book.

Pertinent information for each boring will be recorded on a test boring log. The recorded information will include:

- Date, boring identification, and project identification.
- Name of individual developing the log.
- Name of drilling company.
- Drill make and model.
- Identification of any alternative drilling methods used.
- Depths recorded in feet and fractions thereof (tenths of inches) referenced to ground surface.
- The length of the sample interval and the percentage of the sample recovered.
- The depth of the first encountered water table, along with the method of determination, referenced to ground surface.
- Drilling and borehole characteristics.
- Sequential stratigraphic boundaries.
- Initial PID screening results of soil samples, and/or PID screening results of ambient headspace air above selected samples.

Each test boring will be backfilled with grout upon completion. Soil cuttings, disposable materials and decontamination water will be placed in New York State Department of Transportation (NYSDOT)-approved drums that will be characterized and disposed off-site in accordance with applicable regulations.

Installation of Groundwater Wells for Monitoring and Recovery of LNAPL

A subcontractor will be retained to provide vehicle-mounted Geoprobe Systems Model 6000 series or equivalent direct-push soil sampling equipment to advance test borings for the subsequent installation of groundwater wells. However, if it is determined in the field that such equipment cannot adequately be advanced through the existing overburden soils, then the NYSDEC will be consulted to approve any modifications to the drilling program and installation of associated wells.

Based on the results of the previous remedial investigation, it is anticipated that the test borings will be advanced to depths up to approximately 20 feet below the ground surface. Sampling equipment will be used to collect soil samples in two-foot or four-foot intervals throughout the entire depth of the test borings. The soil samples will be collected in new disposable plastic liners. The soil samples will be collected ahead of 4.25-inch inner diameter hollow stem augers. The soil sampling equipment and hollow stem auger equipment will be advanced to equipment refusal (i.e., inferred top of bedrock).

The recovered soil samples will be visually examined and screened with a PID in accordance with the protocol specified above for "Collection of Confirmatory Soil Samples from Test Borings". This information will be recorded on test boring logs.

Following the completion of drilling, a Schedule 40 polyvinyl chloride (PVC) monitoring well will be constructed within each completed test boring. Each monitoring well will consist of a pre-cleaned two-inch inner diameter, threaded, flush-jointed, five-foot to ten-foot long No. 10 slot screen that is attached to solid riser casing that will extend from the top of the screened section to the ground surface. Each well

screen will be installed to intercept the top of the uppermost water-bearing unit. A washed and graded sand pack surrounding the screen and extending up to one foot below it and about one to two feet above it will be placed in the annulus. A minimum two-foot bentonite seal will be placed above the sand pack and the remaining annulus will be filled with cement/bentonite grout. A steel protective casing with locking cap, or flush-mounted curb box with bolted cover will be placed over each well and cemented in place, and a concrete seal will be installed at the ground surface.

Pertinent information will be recorded on test boring logs and well construction diagrams, which will include:

- Date, boring/well identification, and project identification;
- Name of individual developing the log;
- Name of drilling contractor;
- Drill make and model, auger size, and sampling method;
- Identification of alternative drilling methods used;
- Depths recorded in feet and fractions thereof (tenths of inches) referenced to ground surface.
- The length of the sample interval and the percentage of the sample recovered.
- The depth of the first encountered water table, along with the method of determination, referenced to ground surface.
- Drilling and borehole characteristics;
- Sequential stratigraphic boundaries;
- Well specifications (materials; screened interval; amount of Portland cement, bentonite and water used to mix grout; etc.); and
- Initial PID screening results of soil samples, and/or PID screening results of ambient headspace air above selected samples.

Soil cuttings, disposable materials, and decontamination water will be placed in NYSDOT-approved drums that will be characterized and disposed off-site in accordance with applicable regulations.

Well Development

At least one week following installation, new groundwater wells will be developed by utilizing either a new dedicated disposable bailer with dedicated cord and/or a pump and new dedicated disposable tubing. No fluids will be added to the wells during development, and non-dedicated well development equipment will be decontaminated prior to development of each well. The development procedure will be as follows:

- Obtain pre-development static water level readings with a static water level indicator or oil/water interface meter;
- Calculate water/sediment volume in the well;
- Obtain initial field water quality measurements (e.g., pH, conductance, turbidity, temperature) using a Horiba U-22 water quality meter (or similar);
- Select development method and set up equipment depending on method used;
- Alternate water agitation methods (e.g., moving a bailer or pump tubing up and down inside the screened interval) and water removal methods (e.g., pumping or bailing) in order to suspend and remove solids from the well;

- Obtain field water quality measurements using a Horiba U-22 water quality meter (or similar) for every one to five gallons of water removed. Record water quantities and rates removed;
- Stop development when water quality criteria listed below have been met;
- Obtain post-development water level readings using a Horiba U-22 water quality meter (or similar); and
- Document development procedures, measurements, quantities, etc.

To the extent feasible, development will continue until the following criteria are achieved:

- Water is clear and free of sediment and turbidity is less than 50 nephelometric turbidity units (NTUs);
- Monitoring parameters have stabilized (i.e., parameters are ±10%); and/or
- A minimum of five well volumes has been removed.

The field measurement data will be presented on Monitoring Well Development Logs.

Collection of Groundwater Samples from Monitoring Wells

Static water level measurements will be obtained from each well using an oil/water interface meter. The presence of LNAPL will be monitored by using visual observations and the oil/water interface meter at each well location. The results of this work will be documented.

Subsequent to obtaining static water level measurements and monitoring the wells for LNAPL, the following low-flow purge and sample techniques will be used to collect a groundwater sample from each well:

- A portable bladder pump connected to new disposable polyethylene tubing will be lowered and positioned at or slightly above the mid-point of the water column within the well screen when the screened interval is set in relatively homogeneous material. When the screened interval is set in heterogeneous materials, the pump will be positioned adjacent to the zone of highest hydraulic conductivity (as defined by geologic samples). Care will be taken to install and lower the bladder pump slowly in order to minimize disturbance of the water column.
- The pump will be connected to a control box that is operated on compressed gas (nitrogen, air, etc.) and is capable of varying pumping rates. An in-line flow-through cell attached to a Horiba U-22 water quality meter (or similar equipment) will be connected to the bladder pump effluent tubing to measure water quality data.
- The pump will be started at a pumping rate of 100 ml/min or less (for pumps that can not achieve a flow rate this low, the pump will be started at the lowest pump rate possible). The water level in the well will be measured and the pump rate will be adjusted (i.e., increased or decreased) until the drawdown is stabilized. In order to establish the optimum flow-rate for purging and sampling, the water level in the well will be measured on a periodic basis (i.e., every one or two minutes) using an electronic water level meter or an oil/water interface meter. When the water level in the well has stabilized (i.e., use goal of <0.33 ft of constant drawdown), the water level measurements will be collected less frequently.

- While purging the well at the stabilized water level, water quality indicator parameters will be monitored on a three to five minute basis with the Horiba U-22 water quality meter (or similar equipment). Water quality indicator parameters will be considered stabilized when the parameter readings listed below are generally achieved after three consecutive readings:
 - $pH(\pm 0.1)$;
 - specific conductance (± 3%);
 - dissolved oxygen (± 10 %);
 - oxidation-reduction potential (± 10 mV);
 - temperature (± 10%); and
 - turbidity (+ 10%, when turbidity is greater than 10 NTUs)
- Following stabilization of the water quality parameters, the flow-through cell will be disconnected and a groundwater sample will be collected from the bladder pump effluent tubing. The pumping rate during sampling will remain at the established purging rate or it may be adjusted downward to minimize aeration, bubble formation, or turbulent filling of sample containers. A pumping rate below 100 ml/min. will be used when collecting VOC samples.
- The procedures and equipment used during the purging and groundwater sampling, and the field measurement data obtained, will be documented in the field and recorded on Monitoring Well Sampling Logs.

During sampling, the following parameters will be measured using a water quality meter(s) and will later be presented on Monitoring Well Sampling Logs:

- Dissolved Oxygen
- Conductivity
- Oxidation/Reduction Potential (redox)
- pH
- Temperature
- Turbidity

4.0 Decontamination Procedures

In order to reduce the potential for cross-contamination of samples collected during this project, the following procedures will be implemented to ensure that the data collected (primarily the laboratory data and groundwater quality measurement data) is acceptable.

It is anticipated that most of the materials used to assist in obtaining samples will be disposable oneuse materials (e.g., sampling containers, bailers, rope, pump tubing, latex gloves, etc.). When equipment must be re-used (e.g., static water level indicator, oil/water interface meter, drilling equipment, etc.), it will be decontaminated by at least one of the following methods:

- Steam clean the equipment; or
- Rough wash in tap water; wash in mixture of tap water and alconox-type soap; double rinse with deionized or distilled water; and air dry and/or dry with clean paper towel.

Split-spoon samplers used during rotary drilling, Macrocore cutting shoes used during direct-push drilling, and other re-usable equipment, will be decontaminated between each use.

When deemed necessary, a temporary decontamination pad will be constructed for decontamination of equipment. Any decontamination pad will be removed following completion of associated activities. Decontamination pad materials, liquids, disposable equipment, and personal protective equipment will be containerized in NYSDOT-approved 55-gallon drums and left on-site until the disposal method is determined.

5.0 Operation and Calibration of On-Site Monitoring Equipment

The field personnel will be familiar with the equipment being used. Volatile vapor monitoring will be conducted using a PID. It is anticipated that a RAE Systems MiniRAE 2000 PID equipped with a 10.6 eV lamp, or equivalent, will be used during this project. The PID will be calibrated in accordance with the manufacturer's specifications using an isobutylene gas standard prior to use and as necessary during fieldwork. Measurements will be collected in accordance with the protocols outlined in the Health and Safety Plan (HASP).

Other miscellaneous field instruments that may be used during this project include:

- An electronic static water level indicator;
- An oil/water interface meter;
- LNAPL recovery equipment;
- A low-flow bladder pump system;
- A global positioning system (GPS);
- Survey equipment; and
- A Horiba U-22 water quality meter, or similar.

These meters will be calibrated, operated, and maintained in accordance with the manufacturer's recommendations.

Mitkem's preventative maintenance procedures and calibration procedures for laboratory equipment are provided in its Quality Assurance Plan (QAP) included in Attachment 1.

6.0 Sample Handling and Custody Requirements

During sampling activities, personnel will wear disposable latex or nitrile gloves. Between collection of samples, personnel performing the sampling will discard used latex gloves and put on new gloves to preclude cross-contamination between samples. As few personnel as possible will handle samples or be in charge of their custody prior to shipment to the analytical laboratory.

New laboratory-grade sample containers will be used to collect soil and groundwater samples. Sufficient volume (i.e., as specified by the analytical laboratory and on Table 7.2 of Mitkem's QAP included in Attachment 1) will be collected to ensure that the laboratory has adequate sample to perform the specified analyses.

Samples will be preserved as specified by the analytical laboratory for the type of parameters and matrices being tested. Table 7.2 of Mitkem's QAP included in Attachment 1 provides sample preservation requirements. Sample holding times and preservation protocols will be adhered to during this project in accordance with the requirements that are also presented on Mitkem's Table 7.2.

Chain-Of-Custody

Samples that are collected for subsequent testing as part of this project will be handled using chain-of-custody control. Chain-of-custody documentation will accompany samples from their inception to their analysis, and copies of chain-of-custody documentation will be included with the laboratory's report. The chain-of-custody will include the date and time the sample was collected, the sample identity and sampling location, the requested analysis, and any request for accelerated turnaround time.

Sample Labels

Sample labels for field samples and QC samples with adhesive backing will be placed on sample containers in order to identify the sample. Sample information will be clearly written on the sample labels using waterproof ink. Sufficient sample information will be provided on the label to allow for cross-reference with the field sampling records or sample logbook.

The following information will be provided on each sample label:

Name of company;

Initials of sampler;

Date and time of collection;

Sample identification;

Intended analyses; and

Preservation required.

Custody Seals

Custody seals are preprinted adhesive-backed seals that are designed to break if disturbed. Seals will be signed and dated before being placed on the shipping cooler. Seals will be placed on one or more location on each shipping cooler as necessary to ensure security. Shipping tape will be placed over the seals on the coolers to ensure that the seals are not accidentally broken during shipment. Sample receipt personnel at the laboratory will check and document whether the seals on the shipping coolers are intact when received.

Sample Identification

The following format will be used on the labels affixed to sample containers to identify samples:

Each sample will be numbered starting at 001, and continue in succession (i.e., 001, 002, 003, etc.). The sample test location will also be provided after the sample number using the following test location designations:

TBC-(x') Confirmatory soil sample from test boring location with depth or depth interval in

parentheses.

MW- Existing or new monitoring well location

MWIRM- Existing monitoring well location
TBxx/xx/xx- Trip Blank with day/month/year

FBxx/xx/xx- Field Blank (equipment rinsate) with day/month/year

As an example, assuming the first project sample is a confirmatory soil sample collected from test boring TBC-1 at a depth of 10 feet, the sample will be designated as 001/TBC-1(10').

Transportation of Samples

Samples will be handled, packaged and shipped in accordance with applicable regulations, and in a manner that does not diminish their quality or integrity. Samples will be delivered to the laboratory no later than 48 hours from the day of collection.

7.0 Analytical Quality Assurance/Quality Control

Analytical laboratory testing will be completed by Mitkem (NYSDOH ELAP ID #11522). The analytical laboratory test results for confirmatory soil and groundwater samples and monitoring natural attenuation groundwater samples will be reported in NYSDEC ASP Category B deliverable reports. Analytical laboratory test results for soil samples will be reported on a dry-weight basis. Mitkem will analyze the samples using the lowest practical quantitation limits (PQLs) possible.

Mitkem will provide internal QA/QC checks that are required by NYSDEC ASP and/or United States Environmental Protection Agency (USEPA) CLP protocol, such as analyses performed, spike blanks, internal standards, surrogate samples, calibration standards, and reference standards. Laboratory reports will be reviewed by Mitkem as outlined in its 2007 QAP that is included in Attachment 1, and also by the Quality Assurance Officer.

Mitkem's laboratory results will be compared to data quality indicators in accordance with Mitkem's QAP included in Attachment 1 and NYSDEC ASP. Data quality indicators include: precision, accuracy, representation, completeness, and comparability.

The analytical methods to be used by Mitkem for each type of sample and sample matrix are identified on Table 1 included in Attachment 2. These exclude analytical methods required by regulated landfill facilities or Monroe County Pure Waters (MCPW) for the purposes of waste disposal. As shown, sample methods include the following:

- Target compound list (TCL) VOCs including tentatively identified compounds (TICs) using NYSDEC ASP Method OLM04.3;
- TCL semi-volatile organic compounds (SVOCs) including TICs using NYSDEC ASP Method OLM04.3;
- Total petroleum hydrocarbons (TPH) using NYSDOH Method 310.13; and
- Natural attenuation parameters such as nitrate, iron (II), manganese, sulfate, methane, and chloride (Methods SM3500D, E300IC, ILM04.1, and RSK175).

In order to provide control over the collection, analysis, review, and interpretation of analytical laboratory data, the following QA/QC samples will be included as part of this project (refer to Table 1 in Attachment 2):

- During the confirmatory groundwater sampling and one round of monitored natural attenuation groundwater sampling, one trip blank will be included per 20 liquid samples, or per shipment if less than 20 samples, when the shipment contains liquid field samples (i.e., groundwater samples) that are to be analyzed by Mitkem for VOCs. These trip blanks will be analyzed for VOCs.
- One matrix spike/matrix spike duplicate (MS/MSD) will be analyzed during confirmatory soil sampling, during confirmatory groundwater sampling, and also during one round of monitored natural attenuation groundwater sampling event for each 20 samples of each matrix that are shipped within a seven-day period. Specific parameters that MS/MSD samples will be tested for by Mitkem will be dependent upon the test parameters of the samples that are being analyzed.

One field blank (i.e., rinsate sample) will be collected from reusable groundwater sampling equipment and reusable soil sampling equipment for each sampling event of 20 samples, or per shipment if less than 20 samples. It is anticipated that a field blank will be collected during confirmatory soil sampling, confirmatory groundwater sampling, and one round of monitored natural attenuation groundwater sampling. It is anticipated that equipment rinsate samples will be tested for the test parameters of the samples that are being analyzed by Mitkem.

As an exception, Osprey will perform plate count analyses on select confirmatory soil and groundwater samples, and will provide the test results on its standard data report format. As shown on Table 1 included in Attachment 2, Osprey will analyze the samples for total plate count and pseudomonas plate count using USEPA Method 9215C.

Data Usability Summary Report

Ms. Hope Kilmer of DAY will complete a data usability summary report (DUSR) on some of the analytical laboratory data that is generated as part of the scope of work in the remedial work plan, to the extent required by the NYSDEC (i.e., analytical laboratory results for confirmatory soil and groundwater samples, and up to two rounds of monitored natural attenuation groundwater sampling). The DUSR will be conducted in accordance with the provisions set forth in Appendix 2B of the Draft DER-10 Technical Guidance for Site Investigation and Remediation dated December 25, 2002. The findings of the DUSR conducted on available data at the time will be incorporated in the Final Engineering Report (FER) and/or subsequent annual MNA reports. A copy of Ms. Kilmer's resume is included in Attachment 3.

Reporting

Analytical and QC data will be included in the FER. The FER will summarize the remedial work and provide evaluation of the data that is generated, including the validity of the results in the context of OA/OC procedures.

8.0 Record Keeping and Data Management

DAY will document project activities in a bound field book on a daily basis. Information that will be recorded in the field book will include:

- Dates and time work is performed;
- Details on work being performed;
- Details on field equipment being used;
- Visual and olfactory observations during field activities;
- Field meter measurements collected during monitoring activities;
- Sampling locations and depths;
- Measurements of sample locations, and test locations, excavations, etc.;
- Personnel and equipment on-site;
- Weather conditions; and
- Other pertinent information as warranted.

Alternatively, DAY may record such information from test locations on designated logs (e.g., sample logs, boring logs, well construction diagrams, etc.). Well development data and well sampling data will also be presented on designated logs.

The analytical data will be reported as electronic data deliverables (EDDs) and as hard copies. Differential GPS, swing ties from existing surveyed site structures, and/or a licensed surveyor will be used to collect spatial data. The spatial data will be plotted using integrated geographic information system (GIS) and/or computer-aided design (CAD) mapping. Electronic and hard copy files will be maintained by DAY.

ATTACHMENT 1

Mitkem Quality Assurance Plan (QAP)

Day Environmental, Inc. JD5970 / 4014R-07

Mitkem Corporation

QUALITY ASSURANCE PLAN 2007

Approved By:	
Shaup & Lewle	1/30/07
Sharyn B. Lawler	
QA/QC Director	Date
Vihai Dig	1/30/07
Yihai Ding · · ·	
Laboratory Manager	Date

MITKEM Corporation 175 Metro Center Boulevard, Warwick, RI 02886 Phone Number: (401) 732-3400, Fax Number: (401) 732-3499

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3.0 INTRODUCTION

MITKEM Corporation is a minority-owned small business environmental services company, incorporated in the State of Rhode Island.

Offices and laboratories are located in Warwick, Rhode Island. The laboratory occupies approximately 11,000 square feet of laboratory space.

This Quality Assurance Plan (QAP) describes the policies, organization, objectives, quality control activities. It also specifies quality assurance functions employed at MITKEM and demonstrates MITKEM's dedication to the production of accurate, consistent data of known quality. This QAP is developed by following the guidelines discussed in the EPA Requirements for Quality Assurance Project Plans for Environmental Data Operations, EPA QA/R-5, Interim Final, Jan., 1994 and the National Environmental Laboratory Accreditation Conference (NELAC) standards, July 12, 2002.

4.0 QUALITY ASSURANCE POLICY STATEMENT

MITKEM is firmly committed to the production of valid data of known quality through the use of analytical measurements that are accurate, reproducible and complete. To ensure the production of such data, MITKEM has developed a comprehensive Quality Assurance/Quality Control Program that operates throughout the entire organization.

MITKEM Management considers Quality Assurance/Quality Control to be of the highest importance in the success of its Analytical Testing Laboratory and therefore fully supports the staff in the implementation and maintenance of a sound and thorough Quality Assurance Program.

MITKEM's corporate success is based on its participation in the most rigorous and quality-focused environmental testing programs, such as the EPA Contract Laboratory Program, US Department of Defense programs, NELAC, and other nationwide and state-specific certification and approval programs. These programs require consistent application of the QA/QC procedures described in this document. MITKEM's ability to demonstrate and document that analyses were performed in this manner is one of the foundations of its business. The other foundation of its business is to provide superior levels of customer service, above and beyond the norm for laboratories performing at this level of quality.

MITKEM's approach to customer service is to aggressively meet or exceed customer expectations, particularly in terms of turnaround time for results. While the production of rapid turnaround time data may require MITKEM employees to "go the extra mile" for the customer, without quality, the data are useless. MITKEM constantly strives to manage its business to rapidly provide data to meet all the requirements of its quality program.

- MITKEM management works to insure: that employees understand the primary importance of quality in its day to day operations,
- that employees will not be subjected to pressure to sacrifice quality for turnaround, financial or other considerations,
- that employees understand the importance of their ethical responsibilities in terms of data manipulation, falsification or other illegal or improper actions,
- that the company avoids involvement in activities that diminish its competence, impartiality, judgment or operational integrity.
- that employees maintain all client information in a confidential manner, and
- that employees understand that any short-term gain realized by disregarding the QA/QC program will be more than wasted by the serious penalties for these actions.
- That the laboratory has the technical personnel to identify occurrences of departure from the quality system and to initiate actions to prevent or minimize such departures.

All employees receive training in these issues as part of the initial orientation process, and are required to acknowledge that they understand their responsibilities in these areas.

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These issues are also discussed among all laboratory staff at company meetings and retraining sessions. The QA Officer, Technical Director and other senior company management are readily available to all staff through their daily presence, "open door" policy and approachable manner. This allows any employee to readily discuss any questions, concerns or issues that may occur.

Quality Control is defined as an organized system of activities whose purpose is to demonstrate that quality data are being produced through documentation. Quality Assurance is more broadly defined as a system of activities designed to ensure that the quality control program is actually effective in producing data of the desired quality.

Quality Control is included as part of Quality Assurance. In supporting government regulatory and enforcement proceedings, a high degree of attention to quality is essential. Thorough application of quality control principles and routine quality assurance audits is required.

The basic components of the MITKEM QA/QC Program are control, evaluation and correction.

<u>Control</u> ensures the proper functioning of analytical systems through the implementation of an orderly and well-planned series of positive measures taken prior to and during the course of analysis including quality control practices, routine maintenance and calibration of instruments, and frequent validation of standards.

<u>Evaluation</u> involves the assessment of data generated during the control process. For example, precision and accuracy are determined from the results of duplicates and spikes, and other check samples. Long-term evaluation measures include performance and systems audit conducted by regulatory agencies, as well as the MITKEM quality assurance group.

<u>Correction</u> includes the investigation, diagnosis and resolution of any problems detected in an analytical system. Proper functioning of the system may be restored through method re-evaluation, analysis of additional check samples, trouble-shooting and repair of instrumentation or examination and comparison with historical data. Corrective actions are documented and reviewed to make sure they are implemented.

Certain situations may occur when there are occasional departures or exceptions from documented policies and procedures or standard specifications due to client or project specific protocols, unusual sample matrix, or special non-target analyte or non-routine analyses. MITKEM's policy is to fully document all such procedures and their associated QC, and notify the client or regulatory agency. If the situation is to continue, a Standard Operating Procedure will be written and implemented.

5.0 QUALITY ASSURANCE MANAGEMENT, ORGANIZATION AND RESPONSIBILITY

Quality Assurance at MITKEM is a company-wide function that depend on:

- (1) cooperative working relationships at all levels within the laboratory and
- (2) multi-level review through all working levels of responsibility.

Responsibilities for QA/QC functions begin with the bench scientist and extend to the chief executive officer.

The primary level of quality assurance resides with the bench scientist. After completion of the documented training program, his/her responsibilities include:

- complying with all aspects of formally approved analytical methods and SOPs,
- carefully documenting each step of the analytical process,
- conscientiously obtaining peer review as required,
- promptly alerting laboratory supervisors and/or QA staff members to problems or anomalies that may adversely impact data quality, and
- participation in corrective actions as directed by the laboratory supervisor or QA Director.

The supervisor of each laboratory is responsible for ensuring thorough oversight of the quality of the data generated by the bench scientists. The laboratory supervisor implements and monitors the specific QC protocols and QA programs with the laboratory to ensure a continuous flow of data meeting all control protocols and Mitkem QA requirements. The laboratory supervisor's responsibilities include providing the bench chemist with adequate resources to achieve the desired quality of performance.

The MITKEM organizational structure is shown in the Organization Chart. Resumes of the CEO/Technical Director, Quality Assurance Director, Operations Manager, Laboratory Manager, MIS Director, Project Manager, Laboratory Supervisors, and other key personnel are included.

Mitkem's lines of communication flow upward on the Organizational Chart. Mitkem's open door policy allows all employees access to anyone on the organization chart. If an employee has an issue with his/her immediate supervisor, he or she may, at any time, speak with someone in management higher up in the Organizational Chart.

Implementation of the entire Quality Assurance Program is the responsibility of the QA Director. While interacting on a daily basis with laboratory staff members, the QA Director remains independent of the laboratories and reports directly to the Chief Executive Officer/Technical Director. The QA Director evaluates laboratory compliance with respect to the QA program through informal and formal systems and performance

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audits as described in Section 13.0. Remedial action, to alleviate any detected problems, is suggested and/or discussed with the appropriate parties and implemented when necessary.

With input from the appropriate staff members, the QA Director writes, edits and archives QA Plans, QC protocols, safety procedures, and Standard Operating Procedures (SOPs) in accordance with US EPA approved methodologies, and GLP procedures. If sites-specific or project-specific QA Plans and/or QC protocols are required, these will be generated as needed.

An essential element of the QA program is record keeping and archiving all information pertaining to quality assurance including QA/QC data, pre-award check sample results, performance test sample results, scores, and follow-up; state certifications of the laboratory; external and internal audits with resolution of EPA and other audit team comments, recommendations and reports. The QA Director also plays an important role in the corrective action mechanism described in Section 16.

In addition, the QA Director works with scientists and management to continuously upgrade procedures and systems to improve the laboratory's efficiency and data quality.

Ultimately, the success of the QA program depends on the cooperation and support of the entire organization. MITKEM's most valuable resource is its staff of dedicated professionals who take personal pride in the quality of their performance.

Laboratory management works to ensure the competence of all who operate equipment, perform tests and calibrations, evaluate data and sign reports. When employees are in training, appropriate supervision will be provided until the employee has demonstrated the appropriate level of understanding, training, and skill.

Mitkem Corporation's personnel job descriptions:

Responsibilities of each staff area in the laboratory include:

Bench Scientist / Preparation Laboratory Areas:

- Analysis of samples through compliance with all aspects of formally approved analytical methods and laboratory SOPs.
- Carefully documenting each step of the analytical process.
- Noting in the appropriate logbook area any unusual occurrences or sample matrix problems.
- Conscientiously obtaining peer review as required.
- Promptly alerting laboratory supervisors and/or QA staff members to problems or anomalies, that may adversely impact data quality.
- Routine housekeeping duties for their laboratory area.

Bench Scientist / Instrument Laboratory Areas:

- Analysis of samples through compliance with all aspects of formally approved analytical methods and laboratory SOPs.
- Routine maintenance of instrumentation.
- Preparation of analytical standards and spiking solutions which are documented and traceable to their original source.
- Carefully documenting each step of the analytical process.
- Noting in the appropriate logbook area any unusual occurrences or sample matrix problems.
- Conscientiously obtaining peer and supervisor review as required.
- Promptly alerting laboratory supervisors and/or QA staff members to problems or anomalies that may adversely impact data quality.
- Documenting the initial review of analysis data to determine compliance with established company QA/QC protocols and any project-specific QA criteria, and noting any unusual occurrences or discrepancies on the data review checklist.
- Routine housekeeping duties for their laboratory area.

Data Reporting Staff:

- Assemble CLP-format data reports by organizing data report forms and raw data in proper order to allow for technical data review.
- Enter data into LIMS or other data reporting computer programs.
- Provide non-technical typographical review of data entered into computer systems by other individuals.
- Deliver data reports to customers by FAX or electronic mail.
- Paginate, photocopy, scan, archive Mitkem's copies of customer reports or other documentation to be retained by the laboratory.
- Ship, or organize for courier delivery, final data reports to customers.
- Assist the QA Director in management of the document control system.

Supervisor:

- Oversight of bench scientists in their laboratory areas.
- Monitors the status of all work in their laboratory area to insure compliance with holding time and turnaround time requirements.
- Training new scientists in the appropriate procedures and methods in the laboratory.
- Works with laboratory managers and the QA staff to review, revise and implement SOPs.
- Insures adequate resources to perform the needed tasks by working with administrative personnel to order needed supplies.

- Insures all supplies and reagents meet the QC requirements of their intended task prior to their use in the laboratory.
- Insures all staff are using proper safety protocols.
- Works with laboratory managers on the annual review of personnel performance.
- Interviews prospective new employees to insure they have the minimal level of qualifications, experience, education and skills necessary to perform their tasks, as well as the appropriate work ethic and social skills necessary for proper teamwork and productivity.
- Review of analytical data to insure compliance with method/SOP requirements prior to release to the client.
- Documents any non-compliance or other unusual occurrences noted during sample analysis and data review such that these can be included in the report narrative and explained to the client.

Senior Scientists:

- Review of analytical data to insure compliance with method/SOP requirements prior to release to the client.
- Documents any non-compliance or other unusual occurrences noted during sample analysis and data review such that these can be included in the report narrative and explained to the client.
- Assist laboratory Managers and Supervisors in other tasks as required.

Laboratory Manager:

- Works with laboratory Supervisors to coordinate laboratory areas in the completion of analytical projects.
- Review of analytical data to insure compliance with method/SOP requirements prior to release to the client.
- Works with QA Director to implement new SOPs and to annually review and revise existing SOPs.
- Works with the QA Director and laboratory Supervisors to develop and implement corrective action when needed.
- Works with management and supervisory staff to continuously improve the quality and efficiency of all company procedures.
- Assists laboratory Supervisors in the annual review of personnel performance.
- Supervises laboratory Supervisors to insure compliance with company QA policies and other company procedures.

Operations Manager:

Prioritizes work in the laboratory areas to insure projects are completed on a timely basis.

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- Works with laboratory Managers and Supervisors to coordinate laboratory areas in the completion of analytical projects.
- Review of analytical data to insure compliance with method/SOP requirements prior to release to the client.
- Writes project report narratives to document any unusual occurrences noted during sample analysis.
- Works with management and supervisory staff to continuously improve the quality and efficiency of all company procedures.
- Works with clients to insure all questions and concerns are addressed and answered.
- Assists laboratory Managers and Supervisors in the annual review of personnel performance.
- Supervises laboratory Managers and Supervisors to insure compliance with company QA policies and other company procedures.

Project Manager:

- Works with the client to completely understand the requirements of all incoming work.
- To evaluate the client's requirements as compared to the abilities of the laboratory as stated in Mitkem's Standard Operating Procedure (SOP); Project Management, SOP 110.0023.
- To communicate the customer's requirements to all laboratory staff working on the project.
- Works with the customer to determine the number and type of sample containers required for the project.
- Works with the Sample Custodian to resolve and communicate to the client any problem or discrepancies with incoming samples.
- Maintains open, responsive and continuous communication with the customer.
- Follows up with the client to assess level of satisfaction, and insure all project goals have been accomplished.

QA Director:

- Implements the entire QA program.
- Interact on a daily basis with laboratory staff.
- Evaluates compliance with the QA program through formal and informal reviews of data and processes.
- Implements the corrective action system.
- Works with laboratory Managers and Supervisors to implement new SOPs and to annually review and revise existing SOPs.
- Interfaces with certification authorities and agencies to maintain existing certifications and obtain new certifications.
- Maintains records of employee training and certification.

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- Instructs laboratory personnel on ethics in the workplace.
- Oversees analytical trends that need to be evaluated and corrected.
- Oversees the implementation of MDLs and control limit studies.
- Directs both the internal and external audit programs.

CEO/Technical Director:

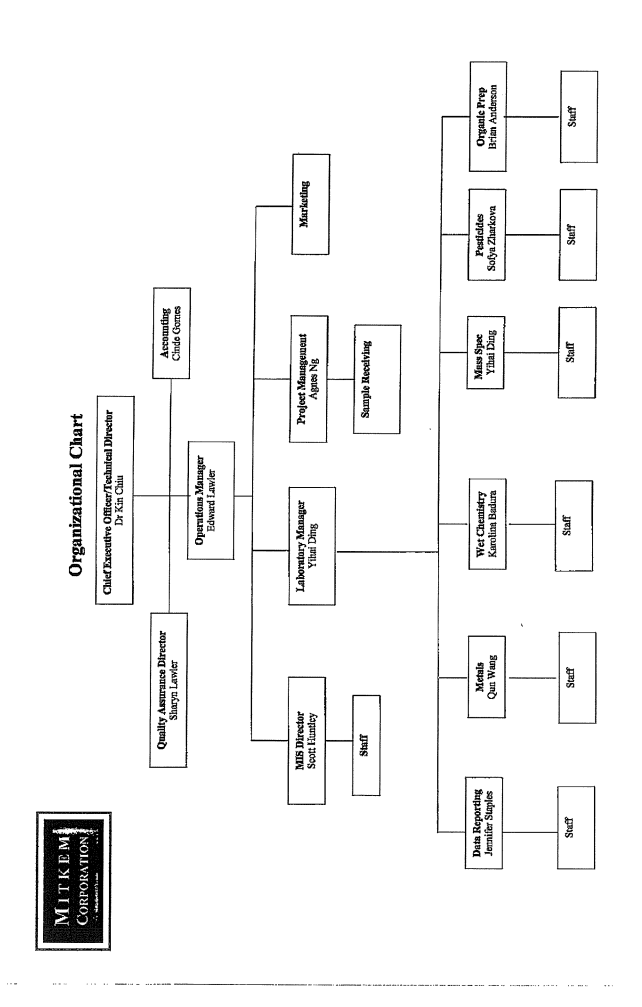
- Review of analytical data to insure compliance with method/SOP requirements prior to release to the client.
- Supervises all Management, QA and Supervisory staff to insure compliance with company QA policies and other company procedures.
- Provides technical assistance to all areas of the laboratory staff.
- Works with clients to insure their understanding of complex technical issues.
- Performs final review of select analytical data to ensure compliance with method/SOP requirements prior to release to the client.
- Acts as technical consultant for chemistry related issues that arise in the lab.
- Provides assistance with instrument optimization or performance issues as needed.
- Offers input on the purchase and operation of new instrumentation.
- Trains other analysts in procedures and methodologies.

In Mitkem's organizational structure, the CEO/Technical Director is one of the principal owners of the company. He is the ultimate authority for all chemistry-related aspects of the company. The QA Director reports directly to the CEO/Technical Director. She has the authority within the management system to bring any issue to the highest levels of the company management and ownership, as well as to halt the release of data she believes to be questionable or suspend the performance of an analysis she believes to be unreliable. The Operations Manager is a Vice President of the company, and works with the project management and marketing staff and with the laboratory Supervisors to prioritize and coordinate work within the laboratories.

The personnel training records are located in the QA department. All individual training is documented including new employee training, individual training, annual retraining procedures, and Health and Safety training.

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Figure 5-1
MITKEM Corporation's Organizational Chart



6.0 QUALITY ASSURANCE OBJECTIVES FOR MEASUREMENT DATA IN TERMS OF PRECISION, ACCURACY, REPRESENTATION, COMPLETENESS AND COMPARABILITY AND QA REPORTING

As part of the evaluation component of the overall QA Program, laboratory results are compared with the data quality indicators defined as follows:

- Precision: the agreement of reproducibility among individual measurements of the same property usually made under identical conditions.
- Accuracy: the degree of agreement of a measurement with the true or accepted value.
- Representation: the degree to which data accurately and precisely represent a characteristic of a population, parameter variations of a sample of a finite process condition, or of a finite environmental condition.
- Completeness: a measure of the amount of valid data obtained from a measurement system compared with the amount that was expected to be obtained under normal conditions.
- Comparability: an expression of the confidence with which one laboratory data set can be compared with another laboratory data set in regard to the same property and laboratory sample population.

Quality Assurance objectives may vary by project and requested parameters. The accuracy, precision, and representation of data will be functions of the origins of the sample material, the procedures used to analyze sample and generate data, and the specific sample matrices involved in each project. Quality control practices utilized in the evaluation of these data quality indicators include blanks, replicates, spikes, standards, check samples, calibrations and surrogates. The process for quantifying or assessing the above indicators for data quality is addressed in Section 15.

6.1 Precision and Accuracy:

For each parameter analyzed, the QA objectives for precision and accuracy will be determined from:

- Published historical data;
- Method validation studies;
- MITKEM experience with similar samples and/or;
- Project-specific requirements, such as those stipulated by the USEPA in the CLP protocols and control documents.

6.2 Representation:

Analytical data should represent the sample analyzed regardless of the heterogeneity of the original sample matrix. In most cases, representation is achieved by mixing the laboratory sample well before removing a portion for analysis. On occasion, multi-phase laboratory samples may require that each phase be analyzed individually and reported in relation to its proportion in the whole sample.

6.3 Completeness:

The completeness goal is 100% in all cases and includes:

- Analysis of all samples;
- Generation and analysis of all required QC samples;
- Sufficient documentation of associated calibration, tuning and standardization;
- Records of data reduction processes, including manual calculations.

While the laboratory staff is responsible for achieving the completeness objective stated above, assigning each project a specific project manager whose functions include sample management and tracking ensures completeness.

6.4 Comparability:

To assure comparability, MITKEM employs established and approved analytical methods (e.g. USEPA protocols), consistent analytical bases (dry weight, volume, etc.) and consistent reporting units (mg/Kg, μ g/L, etc.). Where data from different samples must be comparable, the same sample preparation and analysis protocols are used for all of the samples of interest.

6.5 QA Reporting

General QA procedures require that an MS/MSD or DUPLICATE/MS be reported with each sample batch up to 20 samples. In addition, each batch requires a method blank (MB) and laboratory control sample (LCS).

An acceptance criterion for the MB depends upon the method criteria. In-house control limits dictate the acceptability of the LCS. A high bias LCS is considered acceptable if the analyte is not present in the samples above the reporting limit. A low bias LCS will require re-extraction (if sample volume allows) and re-analysis.

DUP, MS, and MSD recoveries and calculated RSD's are specified in the methods of analyses. Recoveries outside the limits require some form of corrective action, whether that includes a post-digestion/distillation/extraction

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spike, re-extraction, re-analysis and/or notification to the client in the project narrative.

Omega LIMS will flag any QA samples outside method criteria on the reporting forms. Formal written corrective action reports are required for any incident that does not meet method criteria and cannot be remedied at the laboratory. The QA Officer signs off on any corrective actions and can also track QA trends in this manner.

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7.0 SAMPLING PROCEDURES

For most projects, outside sampling teams deliver or send samples to the MITKEM laboratory. When sampling by MITKEM personnel is required, the sampling team follows the sampling procedures outlined in the EPA *Test Methods for Evaluating Solid Wastes*, SW-846, 3rd Edition, or procedures found in the EPA "Handbook for Sampling and Sample Preservation of Water and Wastewater".

Appropriately prepared sample containers are supplied by MITKEM at clients' request. When required, preservatives are added to the sample containers. Tables 7-1 through 7-3 provide the MITKEM Recommended Container, Preservation Techniques and Holding Times. Additional sample volumes may be required if additional QC functions are to be performed.

Holding times for SW846, CLP Methods, Standard Methods and certain USEPA methods are different and are presented in Tables 7-1 to 7-3. Holding times for most methods are calculated from the date of sample collection. Holding times for CLP methods are calculated from the Validated Time of Sample Receipt (VTSR). It should be noted that the CLP analysis program combines chemical analyses and contract compliance procedures in one document. For laboratory analysis and contract compliance purposes, holding times are calculated from VTSR, while post-analysis data usability and validation (generally performed by the client or a third party) compares holding times to the SW-846 method holding times calculated from date of sample collection.

Representative portions of samples are taken for analysis by following Mitkem SOP 110.0039, Standard Operating Procedure for Sub-Sampling.

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Table 7-1
Recommended Container, Preservation Techniques and Holding Times for SW-846 Analyses

Analyte Volatile	es Organics	Method	<u>Containers</u>	Required* Volume	Preservation	Holding <u>Times</u>
Volume	Solid	8260C, 5030B	Amber glass jar with Teflon lining	Minimal head- space in jar	4°C	14 days
	Solid ^a	8260C, 5035	40mL vial or Encore with Teflon lining	5.0gram ± 0.5	4°C, unpreserved	48 hours
			with 10tion mining		DI Water -10 to -20°C	14 days
					Sodium bisulfate -10 to -20°C, 4°C	
					Methanol 4°C	14 days
	Aqueous	8260C, 5030B	40mL VOA Vials with Teflon septum	40mL	4°C HCl, pH<2	14 days
Semivo	latile Organics					
	Solid	3540C, 3550B 8270D	Amber glass jar with Teflon lining	30gram	4°C	Extraction within 14 days Analysis within 40 days
	Aqueous	3510C, 3520C 8270D	Amber glass bottles with Teflon lining	1L	4°C	Extraction within 7 days Analysis within 40 days
Polychie	orinated Biphenyl	s				
2 00,	Solid	3540C, 3550B 8082	Amber glass jar with Teflon lining	30gram	4°C	Extraction within 14 days Analysis within 40 days
	Aqueous	3510C, 3520C 8082	Amber glass bottle with Teflon lining	1 L	4°C	Extraction within 7 days Analysis within 40 days
Oroano	chlorine Pesticides	4				
o.g.mv.	Solid	3540C, 3550B 8081A	Amber glass jar with Teflon lining	30gram	4°C	Extraction within 14 days Analysis within 40 days
	Aqueous	3510C, 3520C 8081A	Amber glass bottle with Teflon lining	1L	4°C	Extraction within 7 days Analysis within 40 days
Chlorin	ated Herbicides					
	Solid	8151A	Amber glass jar with Teflon lining	30gram	4°C	Extraction within 14 days Analysis within 40 days
	Aqueous	8151A	Amber glass bottle with Teflon lining	1L	4°C	Extraction within 7 days Analysis within 40 days

Table 7-1 (cont'd)

Recommended Containers, Preservation Techniques and Holding Times for SW846 Analyses

Analytes Total Pe	troleum Hydroca	<u>Method</u> arbons	Containers	Required* Volume	<u>Preservation</u>	Holding <u>Times</u>				
Gasoline	e Range Organics Solid	s, including Maine 8015M, 5030B ME 4.1.17	→GRO** Amber glass jar With Teflon lining	Minimal head- space in jar	4°C	14 days				
	Solid ^a	8015M, 5035	40mL vial or Encore with Teflon lining	5.0gram ± 0.5	4°C, unpreserve	d 48 hours				
			wat tolder assuing		4°C, Methanol	14days				
	Aqueous	8015M, 5030B ME 4.1.17	40mL VOA vials With Teffon septum	40mL	4°C HCl, pH<2	14 days				
Diesel R	ange Organics, i	ncluding Maine-D	RO							
	Solid	3540C, 3550B 8015M ME 4.1.25	Amber glass jar with Teflon lining	30gram	4°C	Extraction within 14 days Analysis within 40 days				
	Aqueous	3510C, 3520C 8015M ME 4.1.25	Amber glass bottle with Teflon lining	1L	4°C H₂SO₄, pH<2	Extraction within 7 days Analysis within 40 days				
Total Me	Total Metals except Mercury and Chromium (VI)									
	Solid	3050B	Amber glass jar	10g	4°C	180 days				
est.	A.	6010C	with Teflon lining							
	Aqueous	3005A, 3010A	Polyethylene bottle	100mL	HNO ₃ , pH<2	180 days				
Chromiu	m (VI)									
	Solid	7196A	Amber glass jar with Teflon lining	10g	4°C	Digestion within 30 days Analysis within 96 hours				
	Aqueous	7196A	Polyethylene bottle	25mL	4°C	24 hours				
Mercury										
	Solid	7471A	Amber glass jar	10g	4°C	28 days				
	Aqueous	7470A	Polyethylene bottle	100mL	4°C HNO₃, pH<2	28 days				
Cyanide	Solid	9012	Amber glass jar with Teflon lining	10g	4°C	14 days				
	Aqueous	9012	Polyethylene bottle	50mL	4°C NaOH, pH≥12	14 days				
Flashpoir	nt									
-	Aqueous	1010	Amber glass bottle	30mL	4°C	28 days				

Table 7-2

Recommended Container, Preservation Techniques and Holding Times
For
CLP/ASP Analyses

Analyte		Method	Containers	Required* Volume	Preservation	Holding <u>Times</u>	
Volatile	e Organics Solid	CLP/ASP	Amber glass jar with Teflon lining	Minimal head- space in jar	4°C	10 days from VTSR	
	Aqueous	CLP/ASP	40mL VOA vials with Teflon septum	40mL	4°C HCl, pH<2	10 days from VTSR	
		CLP Low	40mL VOA vials with Teflon septum	40mL	4°C HC l, pH<2	10 days from VTSR	
Comisso	latile Organics						
Seimvo	Solid	CLP/ASP	Amber glass jar with Teflon lining	30gram	4°C	10 days from VTSR Analysis within 40 days	
	Aqueous	CLP/ASP	Amber glass bottle with Teflon lining	1L	4°C	5 days from VTSR Analysis within 40 days	
		CLP Low	Amber glass bottle with Teflon lining	1L	4°C	5 days from VTSR Analysis within 40 days	
Omerana	chlorine Pesticide	and.					
Organo	Solid	CLP/ASP	Amber glass jar with Teflon lining	30gram	4°C	10 days from VTSR Analysis with 40 days	
	Aqueous	CLP/ASP	Amber glass bottle with Teflon lining	1L	4°C	5 days from VTSR Analysis within 40 days	
		CLP Low	Amber glass bottle with Teflon lining	1L	4°C	5 days from VTSR Analysis within 40 days	
Cyanid	٠.						
Сушна	Solid	CLP/ASP	Amber glass jar	10gram	4°C	12 days from VTSR	
	Aqueous	CLP/ASP	Polyethylene bottle	50mL	4°C NaOH, pH>12	12 days from VTSR	
Total N	fetals except Mere	curv					
1 0001 1	Solid	CLP/ASP	Amber glass jar	10gram	4°C	180 days from VTSR	
	Aqueous	CLP/ASP	Polyethylene bottle	100mL	HNO ₃ , pH<2	180 days from VTSR	

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Table 7-2 (con't)

Recommended Container, Preservation Techniques and Holding Times For CLP/ASP Analyses

Analytes Mercury	Method	Containers	Required* <u>Volume</u>	Preservation	Holding <u>Times</u>
Solid	CLP/ASP	Amber glass jar	10gram	4°C	26 days from VTSR
Aqueous	CLP/ASP	Polyethylene bottle	100mL	4°C HNO₃, pH<2	26 days from VTSR

Table 7-3

Recommended Containers, Preservation Techniques and Holding Times for Other Analyses

Analytes Volatile Organics Aqueous	olatile Organics		Required* Volume 40mL	Preservation 4°C HCl, pH<2	Holding <u>Times</u> 14 days			
Semivolatile Organics Aqueous	3510C, 3520C 625	Amber glass bottle with Teflon lining	1L	4°C	Extraction within 7 days Analysis within 40 days			
Organochlorine Pesticid				***	Produce at the model in 7 days			
Aqueous	3510C, 3520C 608	Amber glass bottle with Teflon lining	1L	4°C	Extraction within 7 days Analysis within 40 days			
EDB/DBCP								
Aqueous	504.1	40mL VOA vials with Teflon septum	35mL	4°C HCl, pH<2	28 days			
MA Extractable Petroleum Hydrocarbons (EPH)								
Solid	3540C, 3550B MADEP	Amber glass jar with Teflon lining	10gram	4°C	Extraction within 7 days Analysis within 40 days			
Aqueous	3510C, 3520C MADEP	Amber glass bottle with Teflon lining	1L	4°C HCl, pH<2	Extraction within 14 days Analysis withinn 40 days			
MA Volatile Petroleum	Hydrocarbons (VP		,					
Solid	MADEP	Amber glass jar with Teflon lining	10gram	4°C 10mL Methanol	14 days			
Aqueous	MADEP	40mL VOA vial with Teflon lining	40mL	4°C HCl, pH<2	14 days			
Oil & Grease			11	A9.C1	38 dayre			
Aqueous	1664	Amber glass bottle with Teflon lining	IL	4°C HCl, pH<2	28 days			
Alkalinity								
Aqueous	SM2320	Polyethylene bottle	100mL	4°C	14 days			
Ammonia		_		405	an James			
Aqueous	SM4500NH3B	Polyethylene bottle	100mL	4°C H₂SO₄, pH<2	28 days			
Chloride Aqueous	EPA 325.2	Polyethylene bottle Table 7-3 (100mL (cont'd)	4°C	28 days			

Recommended Containers, Preservation Techniques and Holding Times for Other Analyses

	A walusta	_	16-AL-J	Cantainan	Required	Buseaution	Holding
	<u>Analyte</u>	<u>s</u>	Method	<u>Containers</u>	Volume	Preservation	Times
	Chloride	•	E300.0	Polyethylene bottle	50mL	4°C	28 days
	COD			•			
		Aqueous	SM5220D	Amber VOA vial	40mL	4°C H2SO4, pH<2	28 days
	Color					11250-1, pri 42	
		Aqueous	E110.2Modified	Polyethylene bottle	50mL	4°C	Immediate
	Nitrate/I	Vitrite					
		Aqueous	E353.2	Polyethylene bottle	50mL	4°C H₂SO₄, pH<2	28 days
	Nitrate/1						40.1
		Aqueous	E300.0	Polyethylene bottle	50mL	4°C	48 hours
	Nitrite						
		Aqueous	SM4500NO2B E300.0	Polyethylene bottle	50mL	4C	48 hours
	Orthoph	osphate	13500.0				
		Âqueous	SM4500-P, E E300.0	Polyethylene bottle	50mL	4°C	48 hours
	Total ph	-					
ris.		Aqueous	SM4500-P B,E	Polyethylene bottle	50mL 50mL	4°C H₂SO₄, pH<2	28 days
	Phenols	.	en secoon	Datamaka tana 1 anti-	000	4073	t. 90
		Aqueous	SM5530B	Polyethylene bottle	250mL	4°C H ₂ SO ₄ , pH<2	28 days
	Sulfates						
		Aqueous	SM4500SO4 E E300.0	Polyethylene bottle	50mL	4°C	28 days
	Sulfide Total						
		Aqueous	SM4500-S D	Polyethylene bottle	50mL	4°C NaOH, pH>12 ZnAc	28 days
	Reactivi	•	Of and a d	A \$ \$	10	400	20 3
		Solid	Chapter 7 SW846	Amber glass jar	10gram	4°C	28 days
		Aqueous	Chapter 7	Polyethylene bottle	250mL	4°C	28 days
	Total Or	ganic Carbon (TO					
		Solid	Lloyd Kahn	Amber glass jar	10g	4°C	14 days

Table 7-3 (cont'd)

Recommended Containers, Preservation Techniques and Holding Times For Other Analyses

Analytes	Method	Containers	Required* Volume	Preservation	Holding <u>Times</u>
Total Organic Carbon Aqueous	E415.1	40mL VOA vials	40mL	4°C HCl, pH<2	28 days
TKN Aqueous	SM4500Norg C	Polyethylene bottle or Amber glass bottle	50mL	4°C H₂SO₄, pH<2	28 days
Total Solids (TS) Aqueous	SM2540B	Polyethylene bottle	200mL	4°C	7 days
Total Dissolved Solids ('I Aqueous	DS) SM2540C	Polyethylene bottle	200mL	4°C	7 days
Total Suspended Solids (Aqueous	TSS) SM2540D	Polyethylene bottle	200mL	4°C	7 days
Settleable Solids Aqueous	SM2540F	Polyethylene bottle	200mL	4°C	48 hours

EPA SW-846 Method 5035 provides several options for preservation of soil samples for volatile organics. Certain state jurisdictions (NY for example) have not adopted these options to-date, and continue to recommend the collection of unpreserved soil sample aliquots for volatiles analysis. Mitkem's preference for low-level analysis is to collect approximately 5 grams of soil into 5mL of organic-free DI water and to preserve by freezing within 48hours of collection. A separate container with approximately 5 grams of soil into 5mL of methanol is also collected for potential medium-level analysis. A separate container of unpreserved soil also must be collected to perform percent moisture analysis.

^{*} These represent minimum required volume. Additional sample volumes should be collected to minimize headspace loss for volatile analysis. Additional sample aliquot are also required to perform QA/QC functions (e.g. spikes, duplicates), % moisture for solid samples and sample re-analysis (if needed).

^a For Massachusetts analyses, the Volatile Organics soil samples are preserved in Methanol in the field.

^{**} Maine GRO soil analysis requires a medium level methanol extraction. A 10 gram sample and 10mL methanol volume is used.

8.0 SAMPLE CUSTODY

8.1 Chain of Custody:

Samples are physical evidence collected from a facility or the environment. In hazardous waste investigations, sample data may be used as evidence in (EPA) enforcement proceedings. In support of potential litigation, laboratory chain-of-custody procedures have been established to ensure sample traceability from time of receipt through the disposal of the sample.

A sample is considered to be in the custody under the following conditions:

- It is in an authorized person's actual possession, or
- It is in an authorized person's view, after being in that person's physical possession, or
- It was in an authorized person's possession and then was locked or sealed to prevent tampering, or
- It is in a secure area.

Chain-of-custody originates as samples are collected. Chain-of-custody documentation accompanies the samples as they are moved from the field to the laboratory with shipping information and appropriate signatures indicating custody changes along the way.

Laboratory chain-of-custody is initiated as samples are received and signed for by the Sample Custodian or his/her designated representative at MITKEM. Documentation of sample location continues as samples are signed in and out of the central storage facility for analysis in the several MITKEM departments, using the Sample Tracking Forms (Fig 8.4-1). After analysis, any remaining sample is held in the central storage area to await disposal. Mitkem's policy is to hold spent samples for a period of at least thirty days from submittal of final report, unless other arrangements are agreed upon with the client.

8.2 Laboratory Security:

Samples and all data generated from the analyses of samples at MITKEM are kept within secure areas during all stages of residence, including the periods of time spent in preparation for analysis, while undergoing analysis, and while in storage.

The entire laboratory is designated as a secure area. The doors to the laboratory are under continuous surveillance, are kept locked after regular business hours and may only be accessed by key or keypad entry. Only authorized personnel are allowed to enter the secure areas. The central laboratory facility and IT office are

only accessed through keypad entry. A MITKEM staff member must accompany visitors to the laboratory.

8.3 Duties and Responsibilities of Sample Custodian:

Duties and responsibilities of the Sample Custodian include:

- 8.3.1 Receiving samples.
- 8.3.2 Inspecting and documenting sample shipping containers for presence/absence and condition of:
 - 8.3.2.1 Custody seals, locks, "evidence tape", etc.;
 - 8.3.2.2 Container breakage and/or container integrity, including air space in aqueous samples, or proper preservation for soil samples for Volatiles analysis.
- 8.3.3 Recording condition of both shipping containers and sample containers (cooler temperature, bottles, jars, cans, etc.).
- 8.3.4 Signing documents shipped with samples (i.e. air bills, chain-of-custody record(s), Sample Management Office (SMO) Traffic Reports, etc.)
- 8.3.5 Verifying and recording agreement or non-agreement of information on sample documents (i.e. sample tags, chain-of-custody records, traffic reports, air bills, etc.). If there is non-agreement, recording the problems, contacting the project manager for direction, and notifying appropriate laboratory personnel. (Client's corrective action directions shall be documented in the case file.)
- 8.3.6 Initiating the paper work for sample analyses on laboratory documents (including establishing sample workorder files) as required for analysis or according to laboratory standard operating procedures.
- 8.3.7 Label samples with laboratory sample identification numbers and cross-referencing laboratory numbers to client numbers and sample tag numbers.
- 8.3.8 Placing samples and spent samples into appropriate storage and/or secure areas.
- 8.3.9 Where applicable, making sure that sample tags are removed from the sample containers and included in the workorder file.

- 8.3.10 Where applicable, accounting for missing tags in a memo to the file or documenting that the sample tags are actually labels attached to sample containers or were disposed of, due to suspected contamination.
- 8.3.11 Monitoring storage conditions for proper sample preservation such as refrigeration temperature and prevention of cross-contamination.
- 8.3.12 Sending shipping containers with prepared sample bottles and sample instructions to clients who request them.
- 8.3.13 Recording temperatures of freezers and refrigerators in the laboratories.
- 8.3.14 Calibrating the non-contact infrared temperature gun quarterly.
- 8.3.15 Disposal of samples after a specified time period determined by contract or client request.

8.4 Sample Receipt:

The Sample Custodian or his/her designated representative receives sample shipments at MITKEM. Unless the shipment is a continuation of a previous workorder, a new workorder file is started for the sample. The information is logged into the Sample Receipt Logbook (Figure 8.4-1).

The cooler is inspected for the following (if applicable) and findings are documented on the Sample Login Form (Figure 8.4-2) for USEPA CLP samples, and on the Sample Condition Form (Figure 8.4-3) for all other samples:

- Custody seal (conditions and custody number)
- Air bill (courier and air bill #)

The cooler is then opened and the following items are checked (in order). Make sure the hood is turned on when the cooler is opened.

- Chain of custody (COC) records (or traffic report). These are usually taped to the inside of the cooler cover.
- Radioactivity using the Geiger counter, which continuously monitors the receiving area for radiation
- Cooler temperature using the non-contact infrared temperature gun. Record the temperature of a temperature blank if available, using a calibrated thermometer. Record each temperature on the COC.

The Sample Custodian will perform the following:

- Remove the sample containers and arrange them in the same order as documented in the chain of custody report.
- Inspect condition of the sample containers.
- Assign laboratory sample ID and cross-reference the laboratory ID to the client ID.
- Remove tags and place in the workorder file.
- Check preservative and document in the Sample Condition Form (Figure 8.4-3) if needed. If additional preservative is needed, it is added at this time.
- Check for air bubbles in aqueous samples and for proper preservation and immersion of soil samples designated for volatile organic analysis.
- Ensure peer review occurs for proper cross-referencing and labeling of sample containers.

Any discrepancies or problems are noted in the Sample Condition Notification Form (Figure 8.4-4).

The sample custodian conveys the information to the project manager who will in turn inform the client, or may directly inform the client of the discrepancies.

Samples can be rejected at Mitkem for any of the following reasons:

- 1. Complete and proper documentation was not sent with the samples.
- 2. Sample labels cannot be identified because indelible ink was not used during the sampling procedure.
- 3. Hold times had already been exceeded when samples arrived at the laboratory.
- 4. Inadequate sample volume.
- 5. Potential cross-contamination has occurred among samples.
- 6. Samples are inadequately preserved.
- 7. The samples or shipping container is badly destroyed during shipping.
- 8. The samples are potentially radioactive.
- 9. The samples represent untreated fecal waste for which Mitkem employees are currently not inoculated against.

In all instances, the client is contacted initially before any action is taken at Mitkem.

The Sample Custodian signs the Sample Receipt Form and originates a file folder for the set of samples. The following forms are included in the file: the Sample Receipt Form, chain of custody records, shipping information, and an orange Sample Condition Notification Form if any problems or discrepancies need to be addressed.

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When the Sample Custodian is not available to receive samples, another MITKEM staff member signs for the sample container. The time, date and name of the person receiving the container are recorded on the custody records. In addition, the cooler temperature is measured and recorded on the Sample Condition Form. The samples are then stored in the centralized walk-in refrigerator in the sample receipt area. The sample receipt area is located in the secure central storage facility of the laboratory. VOA samples are stored in the VOA analysis laboratory. The samples are officially received and documented by the Sample Custodian or designee before the next business day.

At times, samples will be sent to another lab for analysis not performed at MITKEM. These subcontracted analyses are performed by laboratories certified to perform the analyses. The use of a subcontractor laboratory is discussed with the client prior to sending samples, per Mitkem's Project Management Standard Operating Procedure.

These samples are packed to prevent breakage and stored in a cooler in the walk-in or stored in the small refrigerator in the central storage facility. The samples are either hand delivered to a local sub-contract lab, or shipped with sufficient coolant to maintain a 4 degree temperature by air courier under MITKEM's chain-of-custody (Figure 8.4-5).

8.5 Sample Log-in Identification:

8.5.1 Sample Identification:

To maintain sample identity, each sample received at MITKEM is assigned a unique sample identification (Sample ID) number. Samples are logged into MITKEM via the Omega Laboratory Information Management System (LIMS).

After inspecting the samples, the Sample Custodian logs each sample into the Omega LIMS, which assigns a MITKEM Sample ID Number. These Numbers are assigned sequentially in chronological order. MITKEM Sample Identification Numbers appear in the following format: YXXXX-NNF

In which: Y – represents the current year with A for 2002, B for 2003, C for 2004, etc.

XXXX – represents a four-digit work order number that is assigned sequentially to each submittal of samples

NN – represents the sample number within the group or workorder. F – represents the fraction. All sample portions that are received in identical bottles with identical preservatives are grouped into one fraction.

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For example, the first fraction of the fifth sample of the 20th workorder of 2003 would have the number: B0020-05A

The MITKEM Sample ID Numbers are recorded on the Sample Login Form (Figure 8.4-2) for USEPA CLP samples, and on the Sample Condition Form (Figure 8.4-3) for all other samples. Information on these forms cross-reference the Sample ID Numbers with SDG numbers, sample tag numbers and/or other client identifiers. Each sample is clearly labeled with its MITKEM Sample ID Number by the Sample Custodian. The same sample ID Number appears on the LIMS status report, on each sample preparation container and extract vial associated with the sample.

8.5.1.1 Sample Extract Identification:

As described in Section 8.5.1, a sample extract is identified with the same unique sample identification number as the sample from which it derives

8.5.2 Sample Login:

Sample login system at MITKEM consists of computerized entry using Omega LIMS (Figure 8.5-1). The information recorded onto the Workorder Report includes:

- Workorder number
- Client name
- Project name and location
- Final data report format
- Date of receipt
- Date sample collected
- Due date, fax and/or hardcopy
- EDD requirements
- Comments or notes on the workorder
- MITKEM Sample Identification numbers
- Client Sample Identification numbers
- Sample matrix
- Analyses required
- Case number, where used by the client
- SDG number, where used by the client

8.5.3 Sample Information:

After sample information is properly recorded (Sample Receipt Logbook, Sample Receipt Forms) and the samples have been properly logged into

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the LIMS, bottle labels are generated and applied to the sample containers. The Sample Custodian notifies the Project Manager or peer or supervisor to review the sample bottle labeling. This person reviews all the information associated with the samples. He/she verifies (by initialing) the correctness of the information on the Sample Condition Form or Sample Log-In Form. Sample login information is available through the Omega LIMS to all appropriate laboratory staff.

The Sample Custodian initiates a red workorder file. This file contains the original Sample Log-In Form or Sample Condition Form, air bills, SMO traffic reports, sample tags, workorder reports and all correspondence with the Client or SMO or others. The red workorder file is forwarded to the Project Manager for review of the login paperwork, and for updating status of the workorder in the LIMS. Once the login information is thoroughly reviewed for correctness, the red workorder file is stored in the data reporting area. Analytical data are placed in this as analyses are completed and data are reviewed.

8.6 Sample Storage and Disposal:

Samples at MITKEM are stored in a central storage facility. After sample receipt and login procedures are completed, the Sample Custodian places the samples in the centralized walk-in refrigerator. Volatile Organic sample aliquots are released to the volatile organic lab with documentation (Figure 8.6-1).

The central storage facility is for samples only; no standards or reagents are to be stored there. Access to the centralized sample storage facility is limited by keypad entry at all times.

All sample/extract refrigerators are maintained at $4^{\circ}\text{C} \pm 2^{\circ}\text{C}$. Standards are kept in freezers maintained at -10 to -20°C . They are monitored twice every working day and once daily on the weekends. Temperatures are recorded in the Temperature Log (Figure 8.6-2).

When analysis is complete, any remaining sample is retained in the central storage facility until it may be removed for disposal (see SOP 30.0024 for Sample Disposal). Broken and damaged samples are promptly disposed in a safe manner. Unless there is a specific request by the client, excess, unused sample aliquots are stored for at least 30 days after the submission of compliant data. The samples are then disposed after such period. USEPA and NYS ASP extracts are stored under refrigeration for at least one year. Other extracts are stored under refrigeration for up to three months, unless there is a specific agreement with the client. After such time, the extracts are disposed. All disposals are performed in a manner compliant with federal and state regulations.

8.6.1 Extract Transfer:

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The extracts generated during the preparation for the organic analyses are transferred from the Organic Prep Lab to the Analysis Labs. The extracts, for Semivolatiles, TPH, Pesticides and PCBs, are checked in the Analysis Lab by entries in the appropriate Extract Transfer Logbook (Figures 8.6-3 and 8.6-4).

Metals analysis samples that are transferred from the prep area to the analysis room are signed for by the metals analyst. This entry occurs in the Metals Preparation Logbooks at the time of the transfer (Figures 8.6-5).

There is no extract transfer that occurs with either Wet Chemistry or VOA samples.

8.6.2 Extract Storage:

Semivolatile, Pesticide/PCB, and TPH extracts, which are contained in crimp top vials or screw cap vials with Teflon lined septa, are stored at 4°C±2°C. Semivolatile and Pesticide/PCB extracts are stored in refrigerators in the Organic Analysis room. They are catalogued numerically by workorder number that approximates chronological order, according to date of receipt. USEPA CLP extracts are stored separately within the refrigerator from sample extracts of other clients.

Excess Pesticide extracts, not analyzed, are stored in screw cap vials with Teflon lined septa in the Organic Prep Lab. In most instances, they consist of the remaining 8 mL portions of aqueous and soil sample extracts and are stored chronologically by workorder.

8.7 Sample Tracking:

When a sample is removed from storage, the analyst who has custody signs the Sample Receipt Log. The Sample Receipt Log records the initials of the sample custodian or other authorized lab personnel who relinquishes custody of the sample(s) to the analyst, as well as the initials of the analyst who receives the sample. When the sample(s) are returned to the central storage facility, the analyst relinquishes the sample to the sample custodian or other authorized lab personnel. In addition to the individual's initials, the date is recorded. This information indicates the location of the sample at any point in time.

Chain-of-custody of a sample ensures that the sample is traceable from the field, where it was taken, through laboratory receipt, preparation, analysis and finally disposal. The primary chain-of-custody documents are used to locate a sample at any point in time.

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- 1. The chain-of-custody form from the field describes the origin and transportation of a sample;
- 2. The MITKEM Sample Receipt Logbook and supporting login records document acceptance of a sample by the Mitkem laboratory; and
- 3. The MITKEM Sample Receipt Logbook documents which analyst has custody of the sample after removal from storage.
- 4. The sample preparation logs and/or extract transfer logs document when the extracts or digestates were received by the analytical labs and where they are stored..

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Figure 8.4-1 Sample Receipt Tracking Logbook Form

MITKEM CORPORATION

Sample Receiving Logbook

Date Recv'd	•	mple #s		Storage	Locations:		
Date Recv'd		mple #s			Locations:		
Date Recv'd		mple #s		Storage	Locations:_		
Date Recv'd	Sa	mple #s		Storage	Locations:_		
Date Recv'd	Sa	mple #s		Storage	Locations:_		
		OUT				IN	
Relinquishe	d By	I	Received By	Relin	quished By		Received By
tate: I	nit:	Date:	Init:	Date:	Init:	Date:	<u>Init:</u>
amp. #s			-				
ate: I	nit:	Date:	Init:	Date:	Init:	Date:	Init:
amp. #s							
ate: I	n it:	Date:	Init:	Date:	Init:	Date:	Init:
amp. #s							
ate: I	nit:	Date:	Init:	Date:	Init:	Date;	Init:
amp. #s							
ate: In	nit:	Date:	Init:	Date:	Init:	Date:	Init:
amp. #s							
ate: Ii	nit:	Date:	Init:	Date:	Init:	Date:	Init:
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afe: In	nit:	Date:	Init:	Date:	Init:	Date:	Init:
amp. #s			•				
omments:	· · · · · · · · · · · · · · · · · · ·	-	•				
lease record analyst's i	nitials, da	te, and sample #	removed. Add	any comments	if necessary (broken bottl	es, empty ja
clude the abbreviated clude bottle or jar nur		he test to be perf	ormed., ie: SVO	A, PCBnear	the "samp. #	s ¹¹ .	

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Figure 8.4-2 USEPA CLP Sample Login Form

•		SWIND DOWN	w oncor		
Lab Name Miłkem	Corporation		,		Page of
Received By (Print	4				Log-in Date
Received By (Signata	ure)			•	
Case Number		Sample Deli	very Group No.	······································	SAS Number
Remarks: (1) Please Se	e associated		Corr	esponding	·
Sample/extract transfir logbook pages submitted with this data package.		EPA Sample #	Sample Tag	Assigned Lab #	Remarks: Condition of Sample Shipment, etc.
1. Custody Seal(s)	Present/Absent* Intact/Broken	·			
2. Custody Seal Nos.				·	
3. Chain of Custody Records	Present/Absent*				
4. Traffic Reports or Packing Lists	Present/Absent*		·		
5. Airbill	Airbill/Sticker Present/Absent*	·			
6. Airbill No.	·······				
7. Sample Tags	Present/Absent*				
Sample Tag Numbers	Listed/Not Listed on Chain- of-Custody	·	·		_
B. Sample Condition	Intact/Broken*/ Leaking			-	
9. Cooler Temperature			,		<u>.</u>
10.Does information on custody records, traffic reports, and sample tags	Yes/No*				
gree? 1.Date Received at Lab					
2.Time Received					
Sample Tr					
raction BNA & Pest pcb	Fraction VOA (1)				
rea # RI .	Area # vea Lab		·	·	
У	Bý				
n .	On	·			
Contact SMO and atta	ach record of resolu				
eviewed By			Logbook No.		A STATE OF THE STA
ate			Logbook Page No	o. ·	

SAMPLE LOG-IN SHEET FORM DC-1

Lab Name Mitkem (Corporation			•	Page of
Received By (Print Na		-	***		Log-in Date
Received By (Signatur	e)				
Case Number		Sample Delive	ery Group No.		Mod. Ref. No.
Remarks:	and a lawtenet		Corre	esponding	
Remarks: (1) Please see associated sample extract transfer logbook pages submitted with this data package.		EPA Sample #	Sample Tag #	Assigned Lab #	Remarks: Condition of Sample Shipment, etc.
1. Custody Seal(s)	Present/Absent* Intact/Broken			•	
2. Custody Seal Nos.	<u></u>			••	
3. Traffic Reports/ Chain of Custody Records (TR/COCs) or Packing Lists	Present/Absent*				•
4. Airbill	Airbill/Sticker Present/Absent*				
5. Airbill No.	A A A A A A A A A A A A A A A A A A A				
6. Sample Tags	Present/Absent*			·	
Sample Tag Numbers	Listed/Not Listed on Chain-of- Custody				
7. Sample Condition	Intact/Broken*/ Leaking				
8. Cooler Temperature Indicator Bottle	Present/Absent				
9. Cooler Temperature					
10. Does information on TR/COCs and sample tags agree?	Yes/No*			·	
II. Date Received at Laboratory	4				
12. Time Received					
Sample Tra	Sample Transfer				
Fraction VOA (1)	Fraction VOA (1) Fraction SNA Production				
Area #	Area #			•	
Ву	B y				
On	On				
* Contact SMO and attac	h record of resoluti	on.			
Reviewed By			Logbook No.		
Date			Logbook Page No).	

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Figure 8.4-3 Sample Condition Form

MITKEM CORPORATION

Sample Condition Form

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Received By: Reviewed By			•	Date:			MITKEM Workorder #:			
Client Project:					Client:					Soil Headspace
							ation (pH)		VOA	or Air Bubbles
			Lab Sam	ole ID	HNO ₃	H ₂ SO ₄	HCI	NaOH	Matrix	≥ 1/4"
1) Cooler Sealed Yes / N	No OF									
0) O t - t - O (-)	Present /	Aleman		-						
2) Custody Seal(s)				<u> </u>						
	Coolers /				-					
,	Intact / Br	oken		<u> </u>	ļ					
3) Custody Seal Number(s)										
				ļ						
										
4) Chain-of-Custody	Present /	Absent								<u>.</u>
					<u> </u>					
5) Cooler Temperature										
Coolant Condition										
	•									
A. A. I. 1967.).	D1	A 1			1					
8) Airbill(s)	Present /	Absent	<u> </u>	<u> </u>						
Airbill Number(s)				1	 					
				<u></u>	<u> </u>					
								<u> </u>		
7) Campia Dattica	Intest/Bro	ken/Leaking								
7) Sample Bottles	IIIIACEDIU	noin Loaniig								
								!		
8) Date Received	•						<u> </u>	L	<u> </u>	L
				1		1	I			
9) Time Received						1	VOA	Matrix K	ву:	
							US = 1	Unpreser	ved Soil	A = Air
Preservative Name/Lot No:							UA =	Unpreser	ved Aqu.	H = HCl
						1	M= M	еΟН		E = Encore
·					1	ı	aHSO₄		F = Freeze	
•					 	-	L	·•		
						1				
See Sample Cond	ition Notific	ation/Correc	tive Action F	orm y	es / no					
				_			Rad C	K yes/	no	

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Figure 8.4-4 Sample Condition Notification Form

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Sample Condition Notification

Mitkem Project#:	Date of Receipt:
Client:	Received By:
Client project #/name:	
Unusual Occurance Description:	
Client Contacted:	
Contacted via: Phone/Fax/E-mail	
Date:Time:	
Contacted By:	NATIONAL CONTRACTOR OF THE PROPERTY OF THE PRO
Name of person contacted:	
Client Response:	
Responded via: Phone/Fax/E-mail	
Name of person responding:	
Responding to:	
<u> </u>	
Mitkem Action Taken:	4.1

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Figure 8.4-5
MITKEM Chain-of-custody Form

MITKEN Corporation

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175 Metro Center Boulevard Warwick, Rhode Island 02886-1755 (401) 732-3400 • Fax (401) 732-3499 email: mitkern@mitkern.com

CHAIN-OF-CUSTODY RECORD

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		REPORT TO	RT TO								1			É	INVOICE TO	OT.	l	l					
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Figure 8.5-1 Workorder Information Form

Mitkem Corporation	20/Dec/06 11:35	11:35 WorkOrder: E1468	rder:	E1468
Client ID: MITKEM WARWICK	Case:		Level:	Report Level: LEVEL 2
Project: WW 9/25	SDG:		EDD:	
Location:	PO: -	ЭН	HC Due:	10/10/06
Comments: N/A		Fa	fax Due:	

Sample ID	Sample ID Client Sample ID	Collection Date	Date Recy'd Matrix Test Code	Matrix		Lab Test Comments	Hold MS SEL Storage
E1468-01A	E1468-01A WW-9/25-C	09/25/2006 0:00	09/26/2006 Aqueous SM5220	Aqueous			Disposed
E1468-01B	E1468-01B WW-9/25-C	09/25/2006 0:00	09/26/2006 Aqueous E200.7	Aqueous		Cd, Cr, Cu, Pb, Ni, Ag, Zn	Disposed
E1468-02A	B1468-02A WW-9/25-G	09/25/2006 0:00	09/26/2006 Aqueous E624	Aqueous	E624		Disposed
E1468-02B	W.W-9/25-G	09/25/2006 0:00	09/26/2006 Aqueous E335.4	Aqueous	E335.4		☐ ☐ Disposed

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Client Rep:

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Figure 8.6-1 Volatiles Receiving Logbook Form

MITKEM CORPORATION: VOLATILES RECEIVING LOGBOOK

1.641				ACIAIIAO FOODO	
VOA Log-In Date	Workorder	Client	Sample Numbers	Relinquished By	Comments
					, , , , , , , , , , , , , , , , , , ,
				-	
		V			

Logbook ID 90.0191-04/05

Reviewed By:

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Figure 8.6-2 Temperature Logbook Form

Date: ___ Analyst _____ Comments Time1: Time 2: Time 3: Refrigerator Freezer ID R-Temp F-Temp F-Temp R-Temp F-Temp R-Temp D N/A R-1-Front R-1- Back R2 F2 F3 R3 F4 R4 F5 R5 **F7** R7 R8 F8 R9 F9 R10 F10 N/A R11 R12 F12 R13 F13 NA R14 F15 NA Se F16 F17 **R17** F18 WA R19 R20 Temperature Requirements

MITKEM CORPORATION: Refrigerator/Freezer Temperature Logbook

Reviewed by:

Freezers between-10 and -20 degree C Refrigerators between 2 and 6 degree C

Logbook ID: 30.0108-12/06

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Figure 8.6-3
Extracts Transfer Logbook Form – Semivolatile Analysis

MITK	EM CORPORA	TION E	XTRACT TRA	NSFER LO	GBOOK: SEMIV	OLATILE ANALYSIS
Date Transferred from Prep Lab	Lab ID		Transferred By	Received By	Storage Location	Comments
					•	1
				.,		
			1,000			

Logbook l	D 70.0	0141-1	12/06
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Figure 8.6-4
Extracts Transfer Logbook Form – Pesticide/PCB Analysis

MITKEM CORPO	RATION EXTRAC	TS TRANSFER L	OGBOOK: PE	STICIDES/PCB	ANALYSIS
Date Transferred from Prep Lab	Lab ID	Transferred by	Received by	Storage Location	Comments
				•	
	1				
				-	
				:	
				:	
			<u> </u>		
				1	

Logbook ID: 60.0132 - 07/06 Reviewed by_____

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Figure 8.6-5
Preparation Logbook Form – Metals Analysis

	Analyst									D 0			1
	Comments												
	Final Volume (ml)									Digestion Temp:			
ok	Sample Clarity After	·								Diges			REVIEWED BY:
MITKEM CORPORATION: Aqueous Metals Preparation Logbook	Sample Color After											ed to:	REVIEW
repar	E H E										,	inquish	
etals P	Conic. HCI											Digestate Relinquished to:	
ous M	Conc. HNO ₃											Digest	
ON: Aque	Clarity Before									Method:	SOP#:		
RPORATI	Sample Color Before			•									
(CO)	Hd										,		
MITKE	Sample Vol (ml)									1 1			
	Client ID												
											ä		11/06
	Sample ID									# vot#	LCSS/Spike ID:		7
	Date									HCI Lot# HN03 Lot#			

9.0 CALIBRATION PROCEDURES AND FREQUENCIES

9.1 Instruments:

Specific calibration and check procedures are given in the analytical methods referenced in Section 10. The frequencies of calibration and the concentrations of calibration standards are determined by the cited methods and any special project or contract-specific requirements. Standard calibration curves of signal response versus concentration are generated on each analytical instrument used for a project, prior to analysis of samples. A calibration curve of the appropriate linear range is established for each parameter that is included in the analytical procedure employed and is verified on a regular basis with check standards as specified in the appropriate CLP Protocols. For non-CLP work, MITKEM adheres to the calibration criteria specified by SW-846 and/or Standard Methods for both organic and inorganic analyses. Where requested, other method specific calibration criteria are used.

For organic analyses whenever possible, unless otherwise specified in the individual methods, the initial calibration standards (ICAL), continuing calibration verification standards (CCV), laboratory control sample spike (LCS) and matrix spike (MS) will all be from the same source. The initial calibration verification (ICV) standards are prepared from a separate source. The following are examples of calibration procedures for various instrumental systems. Refer to the Standard Operating Procedures for the specific calibration requirements.

GC/ECD and GC/FID — An initial calibration is performed using five different concentration levels for each parameter of interest for SW-846 analyses. The initial calibration is done on each column and each instrument, and is repeated each time a new column is installed or whenever a major change is made to the chromatographic system.

An initial calibration verification (ICV), near mid level concentration for all analytes, is performed immediately after the calibration. If the ICV does not meet method specific criteria, a new calibration curve is generated and an ICV is analyzed. If repeated ICV failures are encountered, the system is checked to find the cause of these failures, and the problem is corrected. For certain GC/FID analyses (i.e. GRO or DRO), the instrument is calibrated using individual compounds while the laboratory control sample or ICV uses a petroleum product (diesel or gasoline).

A continuing calibration verification (CCV), near a mid-level concentration for all analytes, is run at ten (10) sample intervals. If CCV values are determined outside the upper limit of the method specified range and if no analytes were detected in the samples, the run will be accepted as valid and 'No Detects' reported for the sample. If an analyte is detected and the CCV is out at the high

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end, the problem will be identified and corrected and the affected samples will be re-analyzed with a compliant CCV.

If a CCV value is out of the method specified limits at the lower limit, the cause of the problem will be identified and corrected, and all samples affected by the out of control CCV will be rerun with a compliant CCV.

For CLP-type analyses, the continuing calibration takes place at the beginning of the analytical sequence and once every twelve (12) hours throughout the analytical sequence. The percent difference in calibration factors for each standard must not exceed the criteria specified by the method.

If a CCV fails to meet criteria limits, a new calibration curve will be generated and all samples affected will be re-analyzed.

GC/MS – For CLP methods, a minimum of five-level calibration (four-level for selected semivolatile compounds) is carried out for each analyte per system before analysis of samples take place.

Continuing calibrations, near midpoint levels, are analyzed every twelve hours of instrument analysis time for CLP analyses.

Re-calibration takes place whenever a major change occurs in the system, such as a column change in the GC or a source cleaning of the mass spectrometer or when the continuing calibration fails to meet method specific requirements.

Tunes are performed once every twelve (12) hours. The GC/MS system is tuned to USEPA specifications for bromofluorobenzene (BFB) or decafluorotriphenylphosphine (DFTPP) for volatile and semivolatile analyses, respectively. Verification of tuning criteria occurs every twelve hours of instrument run time for all CLP-type and SW846 analyses.

More detailed instrument and method-specific calibration procedures and criteria are described in the individual analysis SOPs.

ICAP – Instrument calibration, for each wavelength used, occurs at the start of each analysis. The calibration curve is constructed per method specification.

An initial calibration verification and initial calibration blank (ICB) are analyzed before analysis of samples. If the ICV and ICB do not meet method specific criteria for an analyte, the analyte is re-analyzed with a new calibration.

During the analysis, a continuing calibration verification (CCV) and continuing calibration blank (CCB) is analyzed at least every ten (10) samples. If either the CCV or CCB fails to meet method specific criteria for an analyte, the source of the problem is investigated. If it can be determined that the failed CCV and/or

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CCB is not representative (such as for instrument carryover from previous sample or from an empty autosampler tube), the CCV and/or CCB are re-analyzed and the reason for the failure documented. If a failure still occurs, further corrective action is performed, and the analyte is re-analyzed with a new calibration.

The CCV is obtained from a source independent from that of the standards. The CCV concentration for the different analytes are at method specified levels.

The Flow Injection Mercury System (FIMS) - Instrument calibration occurs at the start of each analysis. The calibration curve is constructed per method specification.

An initial calibration verification (ICV) and initial calibration blank (ICB) are analyzed before analysis of samples. If the ICV and ICB do not meet method specific criteria for Mercury, re-calibration and reanalysis are required.

During the analysis, a continuing calibration verification (CCV) and continuing calibration blank (CCB) is analyzed at least every ten (10) samples. If either the CCV or CCB fails to meet method specific criteria for Mercury, the source of the problem is investigated. If it can be determined that the failed CCV and/or CCB is not representative (such as for instrument carryover from previous sample or from an empty autosampler tube), the CCV and/or CCB are re-analyzed and the reason for the failure documented. If a failure still occurs, further corrective action is performed, and the analyte is re-analyzed with a new calibration.

The CCV is obtained from a source independent from that of the standards. The CCV concentration for Mercury is at method specified levels.

Other instrumentation:

pH- the meter is calibrated at two pH levels (4.0 and 10.0) before analyses of samples. The pH 7.0 buffer is analyzed as an LCS and recovery is calculated.

Lachat 8000- automated flow-through spectrophotometer is calibrated per method specification before the analyses of samples.

An initial calibration verification and initial calibration blank (if required) are analyzed before analysis of samples. If the ICV and/or ICB do not meet method specific criteria for an analyte, re-calibration must occur.

During the analyses, a continuing calibration verification and continuing calibration blank is analyzed at least every ten (10) samples. If either the CCV or CCB fails to meet specified criteria for an analyte, the source of the problem is investigated. If it can be determined that the failed CCV and/or CCB is not representative (such as for instrument carryover from previous sample or from an empty autosampler tube), the CCV and/or CCB are re-analyzed and the reason for

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the failure documented. If a failure still occurs, further corrective action is performed, and the analyte is re-analyzed with a new calibration.

The CCV is obtained from a source independent from that of the standards. The CCV concentration for the different analytes are at method specified levels.

SpecGenesys- manual spectrophotometer is calibrated per method specification.

A calibration curve calibration verification is analyzed at the beginning, end, and at least every 10 samples. The verification standard is from an independent source. If the calibration verification does not meet method specific criteria for an analyte, it is re-analyzed once. If failure still occurs, a new calibration curve is established and any affected samples are reanalyzed. Calibration curves are established at least quarterly.

Balances: are calibrated by an outside source on an annual basis. The balances are calibrated with Class "S" weights each day of use. A calibration check is performed with NIST Class "1" traceable weights monthly. The Class "1" weights are NIST certified by an outside certified service on a regular basis.

Thermometers are calibrated once a year against a NIST-verified thermometer or as they are replaced. The NIST-verified thermometers are certified by an outside certified service annually.

Gel Permeation Chromatography is used to clean samples according to CLP and client requirements. GPCs are calibrated using a calibration standard provided by Ultra Scientific, Cat. # CLP-340. Once a successful calibration is achieved it is valid for a period of seven days.

9.2 Standards and Reagents:

Standard reference materials used for routine calibration, calibration checks, and accuracy are obtained from commercial manufacturers. These reference materials are traceable to the source and readily compared to EPA references. Most standards are traceable to NIST; however, certain projects, especially those involving pesticide registration, may necessitate the use of reference standards supplied by the client. New standards are also routinely validated against known standards that are traceable to EPA or NBS reference materials.

Standards are purchased from valid vendors with proven expertise in their field. All standards come with a Certificate of Analysis which is kept on record in the appropriate laboratories. Intermediate standards, if necessary, are prepared in the labs and then QA'd by spiking reagent water with the standard. The spike sample is then carried through the normal extraction and analysis procedures. Criteria for the intermediate spike must meet the method or in-house criteria. If acceptable,

the spike is able to be used. If unacceptable, another intermediate standard is prepared and the same steps repeated.

Intermediate and working standards are prepared in the same solvent or solution as the samples that the standard will be spiked.

Primary, intermediate and working standards are all named with specific nomenclature as designated in the QA Department SOP No. 80.0013, Reagent Purchasing and Tracking.

Standards are dated and labeled upon arrival. Any material exceeding its shelf life as described by the methods in QAP Section 10 is discarded and replaced. Standards are periodically analyzed for concentration changes/degradation and inspected for signs of deterioration such as color change and precipitate formation. Standards Receiving and Preparation Logbooks, which contain all pertinent information regarding the source and preparation of each analytical standard, are maintained by each of the MITKEM laboratory departments (Examples, Figures 9.2-1 to 9.2-4).

See Mitkem individual analytical SOPs, sections 7 and 8 for standards preparation procedures.

Solvents are examined for purity prior to use to ensure there is no external source of contamination. For organic solvents, each lot number of solvent is QC'd prior to use. This is accomplished by concentrating or extracting an aliquot of solvent or reagent media in the same manner as the samples and analyzing it for contamination. Any detectable analyte could render the solvent or reagent unsuitable for use. Supervisors make the final decision as to the suitability of the solvent or reagent.

Reagents are stored in the respective laboratories during use. Backup supplies are stored in Mitkem's stockroom. All chemicals and reagents are given a 3-year expiration period unless designated otherwise by the manufacturer. Sometimes the viability of the reagent does not remain throughout the entire 3-year period. In this case, the chemical or reagent is readily discarded.

Chemicals and reagents are logged into the laboratory and each bottle is given a unique ID. The ID is based upon the date of its arrival at Mitkem. The only exceptions include cases/cycletainers of solvents and cases of acids.

Any applicable certificates of analysis (COA) are stored in the individual laboratories or in the QA Department. When a bottle is opened in the laboratory, it is inspected to ensure it meets the requirements of the method. The analyst records his or her initials on the bottle along with the date opened and the ID.

9.3. Lab Pure Water:

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For wet chemistry, most standards are prepared in DI reagent water. For inorganic analyses Mitkem uses a US Filter mixed-bed deionization system followed by particle and carbon filters. This is followed by a polishing system using Barnstead E-Pure cartridges optimized for removal of inorganic constituents. Purity is monitored each day of use, using an on-line electrical resistivity meter while drawing water through the DI system, as well as reading the conductivity of the water with a hand-held conductivity meter.

Mitkem uses several systems to generate analyte-free water for use in the Organics laboratory. These systems generate high quality, analyte free water dedicated to the needs of specific analyses. The extractable organics laboratory uses a Barnstead E-Pure system optimized for removal of organic constituents. The volatile organics laboratory uses an in-house activated carbon filtration system to provide analyte free water. As organic contaminants are not measured by a resistivity meter, this is not relied-upon to monitor the quality of organic analyte-free water. Instead laboratory method blanks are used, typically several per working day, to monitor the acceptability of the water for its intended use. Any analyte detected above (half of) the reporting limit is investigated. If this can be traced to the water purification system as its source, maintenance is performed on the water purification system.

9.4. All purchased equipment, materials, and services must meet either specific method requirements, standard requirements, or project specific requirements. These requirements are documented in the individual analytical or project SOPs. Reagents requirements are specified in the Mitkem SOP, SOP 80.0013 Reagent Purchasing and Tracking. The equipment requirements are specified in the individual methods and SOPs.

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Figure 9.2-1 Metals Primary Standard Receipt Logbook – Instrument Laboratory

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METALS PRIMARY STD RECEIPT LOGBOOK:INSTRUMENT LAB			ANALYTE(S)												
S PRIMA			LOT#												
METAL			CATALOG#												¥
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MITKEM CORPORATION			PRIMARY STD ID												
MITKE		DATE	REC												

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Figure 9.2-2 Semivolatile Primary Standard Logbook - Preparation Laboratory

MITKEM CORPORATION SEMIVOLATILE PREP WORKING STANDARD LOGBOOK

		,	_	-	,		-	 _	·	<u> </u>	 	 	 	 -	 	 -
EXP	DATE		MV demonstratures													
PREP	ВҰ															
	WORKING STD ID															
Solvent Name:	LOT#															
FINAL	CONC ug/L														-	
FINAL	VOL (ml)															
AMT.	ADDED ml															
INITIAL	CONC.ug/L ADDED ml VOL (ml) CONC ug/L LOT #															
	PRIMARY STD ID					•										
	ANALYTE(S)															
PREP	DATE									•	·			-		

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Figure 9.2-3
Pesticide/PCB Primary Receipt Logbook

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	REC BY				,			,				
GBOOK	ANAL/YTE(S)											
CEIPT LO	CONC.											
PRIMARY STD RECEIPT LOGBOOK	LOT#								•			
CB PRIMA	CATALOG#							-				
STICIDE/P	VENDOR							-				-
MITIKEM CORPORATION PESTICIDE/PCB	PRIMARY STD ID		٠									
MITKEN	DATE											

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Figure 9.2-4
Reagent Preparation Logbook – Inorganic Preparation Laboratory

Mitkem Corporation Inorganic Laboratory Reagent Preparation Logbook

Date:

	1	1			 1	T	 	
Analyst								
Exp. Date						·		
Reagent ID								
Diluent Final Conc.								
Diluent								
Final Volume. (ml)								
g/mL								
Chemical Lot#								
Chemical ID								
Reagent/Analysis								

Reviewed By:

Logbook ID 100.0170 - 10/06

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10.0 ANALYTICAL PROCEDURES

MITKEM uses the methods specified in Tables 10-1 through 10-6 unless otherwise specified by the client.

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Table 10-1 Potable Water Analytical Methods

Parameter	Method Description	Method Reference
1,2-Dibromo-3-chloropropane 1,2-Dibromomethane	Micro extraction GC\ECD Analysis	504.1

Table 10-2 Non-potable Water Priority Pollutant Analytical Methods

Parameter	Method Description	Method Reference		
Metals Aluminum, Antimony, Arsenic, Barium, Beryllium, Cadmium, Chromium, Cobalt, Copper, Iron, Lead, Manganese, Molybdenum, Nickel, Selenium, Silver, Silver, Thallium, Potassium Vanadium, Zinc, Sodium	ICP	200.7		
Mercury	Cold Vapor	245.1		
Cyanide Aqueous	Midi-distillation Automated	EPA 335.4		
Alkalinity	Titration	SM2320		
Anions Chloride Sulfate Nitrate Nitrite Phosphate Bromide	Ion Chromatography	EPA 300.0		
Chloride	Colorimetric	EPA 325.2		
pН	Electrode	SM4500 H+ B		
Sulfate	Turbidimetric	SM4500-SO4 E		
Ammonia	Distillation/Nesslerization	SM4500-NH3 B		
Nitrate	Autoanalyzer	EPA 353.2		
Nitrite	Colorimetric	SM4500-NO2 B		
Orthophosphate	Ascorbic, Manual	SM4500-P E		
Total phosphate	Persulfate, Manual	SM4500-P B3 & E		

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Table 10-2 Non-potable Water Priority Pollutant Analytical Methods (cont.)

<u>Parameter</u>	Method description	Method Reference
Chemical Oxygen Demand	Spectrophotometric(Closed Reflux)	SM5220-D
Total Organic Carbon	Combustion	EPA 415.1
Phenols	Distillation, Color, Automated	.SM5530B
Total Dissolved Solids	Gravimetric	SM2540 C
Total Solids	Gravimetric	SM2540 B
Total Suspended Solids	Gravimetric	SM2540 D
Total Settleable Solids	Imhoff cones	SM2540 F
Volatile Organics Halocarbons Aromatics	Purge & Trap, GC/MS Purge & Trap, GC/MS	624 624
Semivolatile Organics	Extraction, GC/MS	625
Organochlorine Pesticides/ PCBs	Extraction, GC/ECD	608
Oil & Grease	Extraction, Gravimetric	1664

Table 10-3 SW-846 Inorganic Analytical Methods

<u>Parameter</u> Metals	Method Description	Method Reference
Aqueous	Acid digestion ICAP analysis	Method 3005A/3010A Method 6010C
Solid	Acid digestion ICAP analysis	Method 3050B Method 6010C
Mercury		
Aqueous	Permanganate digestion Cold Vapor analysis	Method 7470A
Solid	Permanganate digestion Cold Vapor analysis	Method 7471A
Hexavalent Chromium	_	
Aqueous	Diphenyl Carbazide Colorimetric	SM 3500Cr D
Solid	Acid Digestion colorimetric	Method 3060A/7196A
Cyanide		
Aqueous	Midi-distillation Automated	Method 9012B
Solid	Midi-distillation Automated	Method 9012B
рΉ		
Solid	Electrode	Method 9045C
Ignitability (Flashpoint)		
Aqueous	Pensky-Martens closed cup	Method 1010
Solid	Pensky-Martens closed cup	Method 1010 Mod.
Reactive Cyanide		
Solid & Aqueous	Distillation Automated	SW 846 7.3.3.2
Reactive Sulfide Solid & Aqueous	Distillation Colorimetric	SW 846 7.3.4.2

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Table 10-3 SW-846 Inorganic Analytical Methods (cont.)

<u>Parameter</u>	Method Description	Method Reference
Toxicity Characteristic Leaching Procedure (TCLP) Aqueous	Leachate by Filtration	Method 1311
Solid	Leachate Generation	Method 1311
Synthetic Precipitation Leaching Procedure (SPLP) Aqueous	Leachate by Filtration	Method 1312
Solid	Leachate Generation	Method 1312

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Table 10-4 SW-846 Organic Analytical Methods

Parameter Volatile Organic Compounds	Sample Preparation	Sample Analysis
Aqueous	Method 5030	Method 8260C
Solid	Method 5035	Method 8260C
Semivolatile Organic Compounds		
Aqueous	Method 3510C Method 3520C	Method 8270D
Solid	Method 3540C Method 3550B Method 3545 Method 3570	Method 8270D
Organochlorine Pesticides		
Aqueous	Method 3510C Method 3520C	Method 8081A
Solid	Method 3540C Method 3550B Method 3545 Method 3570	Method 8081A
Polychlorinated Biphenyls (Aroclors and Congeners)		
Aqueous	Method 3510C Method 3520C	Method 8082
Solid	Method 3540C Method 3550B Method 3545	Method 8082
Total Petroleum Hydrocarbons	Method 3570	
Aqueous	Method 3510C	%
riquous	Method 3520C	Method 8015M
Solid	Method 3540C Method 3550B Method 3545 Method 3570	Method 8015M

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Table 10-4 SW-846 Organic Analytical Methods (cont.)

<u>Parameter</u>	Sample Preparation	Sample Analysis
Herbicides Aqueous	Method 8151A	Method 8151A
Solid	Method 8151A	Method 8151A
Toxicity Characteristic Leaching Procedure (TCLP) Aqueous Method 1311		
Solid	Method 1311	
Synthetic Precipitation Leaching Procedure (SPLP) Aqueous Method 1312		
Solid	Method 1312	
Gal Permeation Chromatography	GPC)	

Gel Permeation Chromatography (GPC)

Aqueous Method 3640A

Solid Method 3640A

Florisil Cleanup

Aqueous Method 3620B

Solid Method 3620B

Silica Gel Cleanup

Aqueous Method 3630C

Solid Method 3630C

Sulfur Cleanup

Aqueous Method 3660B

Solid Method 3660B

Sulfuric Acid Cleanup

Aqueous Method 3665A

Solid Method 3665A

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Table 10-5 CLP-Type Analytical Methods

<u>Parameter</u> <u>Method Reference</u>

USEPA CLP Organics OLM04.3, SOM01.1

USEPA CLP Inorganics ILM04.1, ILM05.3

USEPA Low Level Organics OLC03.2

NYS-ASP CLP Organics ASP 2000/2003 SOW

NYS-ASP CLP Organics ASP 2000/2003 SOW

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Table 10-6 Other Analytical Methods

<u>Parameter</u>	Method Reference
Volatile Petroleum Hydrocarbons Aqueous	MADEP VPH 1.1
Solid	MADEP VPH 1.1
Extractable Petroleum Hydrocarbons Aqueous	MADEP EPH 1.1
Solid	MADEP EPH 1.1
New York State Total Petroleum Hydrocarbon Solid	310.13 Mod.
Extractable Total Petroleum Hydrocarbons Aqueous	CT ETPH 99-3
Solid	CT ETPH 99-3
Deisel Range Organics Aqueous	ME 4.1.25
Solid	ME 4.1.25
Gasoline Range Organics Aqueous	ME 4.1.17
Solid	ME 4.1.17

10.1 Analytical References

- Analysis of Extractable Total Petroleum Hydrocarbons (ETPH) Using Methylene Chloride Gas Chromatograph/Flame Ionization Detection, Environmental Research Institute, University of Connecticut, March, 1999
- 2. Analytical Services Protocol, Volume 1-8, New York State Department of Environmental Conservation, 2003.
- 3. Annual Book of ASTM Standards. Part 31-Water. American Society for Testing and Materials, Philadelphia, PA, 1981.
- 4. Chemical Characteristics of Marine Samples, API Publications No. 4307, API, Washington, D. C.
- 5. Federal Register. Vol. 55, No. 61, March 29, 1990
- 6. Methods for Chemical Analysis of Water and Wastes, EPA-600/4-79-020, 3/83 Revision.
- The EPA 600 Series. Methods for Organic Chemical Analysis of Municipal and Industrial Wastewater, Appendix A, 40 CFR Part 136, Federal Register, Vol. 49, No. 209, 1984.
- 8. Methods of Soil Analysis. Part 2, Chemical and Microbiological Properties, Second Edition, American Society of Agronomy, Inc., Soil Science Society of America, Inc., Madison, WI, 1982.
- Standard Methods for the Examination of Water and Wastewater, 20th Edition, APHA, Washington, D. C., 1998.
- Test Methods for Evaluating Solid Waste-Physical/Chemical Methods, SW-846, 3rd Edition Update IV. Office of Solid Waste and Emergency Response, USEPA, Washington, D. C., 1998.
- 11. USEPA Contract Laboratory Program. Statement of Work for Organic Analysis, USEPA, OLM04.3, OLC03.2, and SOM01.1.
- 12. USEPA Contract Laboratory Program. Statement of Work for Inorganic Analysis, USEPA, ILM04.1, ILM05.3.

11.0 DATA COLLECTION, REDUCTION, VALIDATION AND REPORTING

11.1 Data Collection:

Most of Mitkem's data is uploaded into the Omega LIMS systems directly from the instruments. The exception is the GC's and GC/MS's in which data is first processed in Target and then uploaded into the LIMS. Mitkem is making progress in that the elimination of the Target reporting will occur in the near future.

Either the instrument analyst or data reporting group will upload the data into the LIMS. The person who performs the upload does a technical review to ensure recoveries of CCVs, MS, MSD, and LCS all seem to be correct. A completeness review is done at this time to ensure all applicable samples have been uploaded for all the necessary analytes.

Next, an employee with a technical background will perform the QA process of the uploaded data. This person is either a supervisor or someone with extensive experience in environmental chemistry. Corrections to the run are made at this step if necessary. When the review is complete, this technical person authorizes the data to be reported by "QA-ing" the run in the LIMS. For a more detailed view of the LIMS uploading/review procedure, see SOP No. 110.0028.

11.2 Data Reduction:

Instrument printouts, computer terminal displays, chromatograms, strip chart recordings and physical measurements provide raw data that are reduced to concentrations of analytes through the application of the appropriate calculations.

Equations are generally given within the analytical methods referenced in Section 10. Data reduction may be performed automatically by computerized data systems on the instrument, manually by the analyst, or by PCs using spreadsheet and/or data base software. This software includes Thru-Put's 'TARGET' for the analyses of organic analytes and Omega LIMS for metals, cyanide and mercury analysis. Currently all OLC analyses are processed and reported through Omega at this time. Mitkem expects that all organic data, both CLP and non-CLP, will be processed completely through the LIMS System during 2006.

11.3 Data Verification:

The verification process requires the following checks to be made on data before they are submitted to the client:

- A completeness inspection is required which ensures that all required data are included in the data packages submitted to the client and that the appropriate signatures are present on the data packages.
- A contract compliance screening to ensure that contractual requirements have been satisfied.
- A consistency check to ensure that nominally identical or similar data appearing in different places within a data package are consistent with respect to value and units.
- A correctness check to ensure that reported data have been calculated correctly or transcribed correctly.

11.4 Data Validation:

Data validation is an essential element of the QA evaluation system. Validation is the process of data review and subsequent acceptance or rejection based on established criteria.

The following analytical criteria are employed by MITKEM in the technical evaluation of data:

- Accuracy requirements.
- Precision requirements.
- Detection limit requirements.
- Documentation requirements.

As in the case of EPA/CLP procedures, data acceptance limits may be defined within the method. As one means of tracking data acceptability, quality control charts are plotted for specific parameters determined in similar, homogeneous matrices. Control limits for non-CLP methods are statistically determined annually as analytical results are accumulated.

Upon completion of the evaluation, the evaluator dates and initials the data review checklist as described in Section 11.5 below.

11.5 Data Interpretation and Reporting:

Interpretation of raw data and calculation of results are performed by a scientist experienced in the analytical methodology. Upon completion of data reduction, the scientist signs for the reported results on the data review checklist. For GC/ECD and GC/MS, a technical peer review is performed using the data processing software prior to form generation.

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The laboratory supervisor is responsible for the data generated in that department. The supervisor or other senior technical staff performs an independent review of data and completed report forms. Members of the QA staff also check the results on selected sets of data (usually 10%).

11.5.1 Report Formats:

Mitkem uses a flexible data reporting system where final report format is based on the requirements of the client. The two most common types of data reports generated by Mitkem are Level 2 or "commercial-format" and Level 4 or "CLP-format". Mitkem adapts its data report format, wherever possible, to meet customer requirements. Occasionally reports are generated that are a compromise between "commercial" and CLP-format deliverables or are designed to meet the needs of a particular regulatory format or sampling program.

Commercial data reports are generated using the Omega LIMS or MS EXCEL. For the Omega LIMS system, all instrumental analysis data are uploaded from instruments to the LIMS by electronic data transfer. Non-instrumental analysis data or sample preparation data are manually entered into the LIMS. All manual data entry steps are double-checked to insure they are correct, and instrumental data are spot-checked to insure the proper functioning of the data upload system. For data entered into MS-EXCEL, all the pertinent client information and the analysis results are entered manually. The draft report is subject to a 100% technical and completeness review before it is printed in its final form. All data receive a 100% review before they are released to the client as final.

CLP data reports are generated using specialized software, Thru-Put TARGET for many organics analyses, and the CLP report modules in the Omega LIMS for all inorganic and certain organic analyses. These reports also undergo a 100% review before they are released to the client in their final form.

Records are maintained for all data, even those results that are rejected as invalid.

11.5.2 Data Reporting for Massachusetts Drinking Water Samples:

Drinking water data reports generated for clients in the State of Massachusetts need to be reported on state forms. These reports are sent to the client. The client is responsible for forwarding copies of the report to the regional DEP Offices and local officials.

11.6 Levels of Data Review:

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MITKEM employs five (5) levels of data review. These are based on requirements outlined in several government and other environmental analysis programs including the U. S. Army Corps of Engineers, Air Force Center for Environmental Excellence (AFCEE), Naval Facilities Engineering Service Center (NFESC), HAZWRAP, EPA Contract Laboratory Program (CLP), as well as commercial engineering firm programs.

The data review and evaluation process is structured to insure that all data reported to customers has been thoroughly reviewed and approved using a multistep process designed to identify and correct any error. At any step in the data evaluation and review process, the reviewer has the responsibility and authority to return any data not meeting requirements back to the previous step for re-analysis or correction. No reports are released to the client as final data without successfully passing through each step in the data evaluation and review process. The steps of the data review process are documented, generally using a checklist. Several checklists are used, depending on the type and format of analysis data being reviewed. Any data released prior to the completion of the full review process are released with the statement that the data is preliminary pending final review. The word "Preliminary" is automatically printed on the bottom of all data sheets that are generated prior to completion of data review.

The five levels of data review are detailed in **SOP No. 110.0028**. A Flow chart of the data review process follow in Figure 11.6-1.

11.7 Document Control:

All login sheets, Chains-of-Custody (COC) and Sample Condition Forms (SCF) and other sample transmittal documentation are generated in Sample Receiving. A red Workorder File is initiated to contain all workorder-specific hard copy documents. Samples are signed in/out of the sample receiving area by analysts. In the Prep lab, samples and all pertinent information is recorded into logbooks. Once samples are moved to the instrument lab, the transfer of extracts is documented in the transfer logbook. In the instrument lab, the analysis of extracts is recorded in the instrument run log. All analysis data, including ICAL, CAL and raw data are acquired using computer-controlled instruments, and stored on the hard drive of the computer performing data acquisition. Data are automatically copied to the company file server after acquisition. Organics analysis data are processed using Thru-Put Systems' Target software. This system creates a folder on the file server for each analysis fraction for each work order or SDG. This folder contains raw data, processed analysis results, instrument tune, initial calibration and continuing calibration results as well as a copy of the data processing method used. This allows for long-term archiving and complete reconstruction of the data at any time in the future. Data reporting forms and raw data are printed and arranged with all appropriate samplepreparation logbook page copies for technical review.

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Inorganic data files are uploaded into Omega LIMS and reporting forms are printed. The original instrument data files and the processed SDG are stored on the file server where they can later be archived by the LIMS Administrator. Hard copy printouts for reporting forms, instrument data hardcopy output and all associated preparation logbook page copies are assembled for technical data review.

The company file server consists of two separate computers, each with an array of multiple hard disk drives, that are continuously mirrored, such that the failure of any single component or computer will not impact the operation of the system, or the ability to recover data. All new files or data are copied to magnetic tape on a daily basis. On a monthly basis full system back up to tape is performed. Following technical review, and generation of the report narrative results go into the workorder file in data reporting. The original copy of the report is sent to the client. The report is also scanned into an optical file database for long-term archiving. As documents are scanned into the database they are recorded for permanent storage on CD-ROM disks. Mitkem's system includes a "jukebox" to provide access to numerous CD-ROMS on an as-needed basis. All other information associated with the report, including data review checklists are kept in the red workorder file. The workorder files are kept onsite in a storage area for approximately 6 months. The files are then shipped to an offsite storage area where they will remain for a total of 7 years. After this time, the files will be destroyed.

11.7.1 Logbooks:

All logbooks are issued and controlled by the QA Department. Logbooks are given a unique ID that includes the mm/yy the logbook was printed. Laboratory personnel must sign for the logbook when it has been released by the QA Department. When logbooks are complete, the analyst returns them to the QA Department for archiving. At that point, a new logbook is released. The archived logbooks are stored in an on-site storage area for approximately 4-6 months and then are boxed and stored in a locked off-site storage facility. Mitkem will archive logbooks for a minimum of ten (10) years.

11.7.2 Workorder/Data Files:

MITKEM is a secured, limited access building. The doors are secured with a keypad entry system. All hard copy information pertaining to the analysis of samples is maintained and stored in a workorder file folder. This information includes all login sheets, COC, SCF, bench sheets and analytical data. Electronic data are also stored by laboratory workorder number on the company file server, and in the optical file database of completed reports. File folders containing all hard copy data and other

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workorder information are stored in an off-site storage facility for a total of 7 years. The off-site storage facility is a locked storage area. Access is limited to the CFO or his designee and request to retrieve a file will be made to this person.

In the event Mitkem Corporation changes ownership, the maintenance, control, storage and eventual disposal at the end of the appropriate time period, of all records, including client data and QA/QC files, will transfer to the new owners.

In the event Mitkem Corporation decides to cease operations, clients will be notified prior to the cessation of operations and their files/records will be made available to them. Within a designated time period after notification, the client will be responsible for taking custody and the future maintenance of their records. If the client determines they do not want to maintain the records, these will be disposed of properly.

11.7.3 Standard Operating Procedures (SOPs):

SOPs are prepared by the Lab Supervisor and laboratory personnel in conjunction with the QA/QC Director. The QA Director/Staff downloads a copy of the current SOP to the network. The SOPs can be found in Avogadro/Public/QA Public. In addition a .pdf file of the SOP is located in Avogadro/Public/QA Public/SOP-PDF Versions, for sending to clients or for analyst reference.

The laboratory staff revises the SOPs by making changes to the document that is then reviewed by the department supervisor only if the supervisor is not the party responsible for the revisions. Any additional changes are made at this point.

The QA Department is notified that revisions are completed. The QA Director/Staff moves the revised copy of the SOP to the QA directory, QA Safety/SOPs Needing QA Revision. The QA Director makes changes to the document to include revision number and date and title clarification, if necessary.

The QA Director prints a copy of the SOP that is signed by the Lab Manager or Operations Manager, and the QA Director. Copies of the signed SOP are then made for the relevant departments. Each copy is assigned a control number that is recorded on the SOP cover sheet. Copies are distributed to the relevant departments with a review sheet attached. At this time the old copies of the SOP are collected from the labs and destroyed. Each analyst who performs any duties related to the SOP must review the new version and sign that he or she has read and understands the material there. The signed review sheets are then returned

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to the QA Department. The SOP copy is stored in the department for easy reference. A new .pdf file is made to overwrite the "old" version in QA Public/SOP-PDF Versions.

SOP review/revisions occur on an annual basis. The procedure for preparing, reviewing, approving, revising and distributing SOPs as well as the SOP Revision Schedule are described in SOP No. 80.0012.

Minor changes to the SOP between revision dates can be done by making hand-written changes to the document and its copies. The changes must be initialed by the QA Director and incorporated into the next version SOP. Minor changes are recorded in the Minor Revision Record that is a part of the master copy.

11.7.4 Method Updates:

In most cases it is the laboratory's policy to implement new revisions of frequently used methods within six months of the date the method revision is promulgated or published as a final method. The QA/QC Director and Technical Director make the final decision on when a method revision will be adopted by the laboratory. Additionally, if a client specifically requests or mandates that an "older" method, Mitkem will advise the client that it is not the most recent method. If the client still insists upon the older method, Mitkem will comply and make a note in the narrative.

When the laboratory is in the middle of a client's project, the lab will continue using the same revision for the entire sampling event unless advised otherwise by the client. Consequently, once the laboratory has formally adopted a new method revision, both the old and new revision may be in use at the same time, depending on the project.

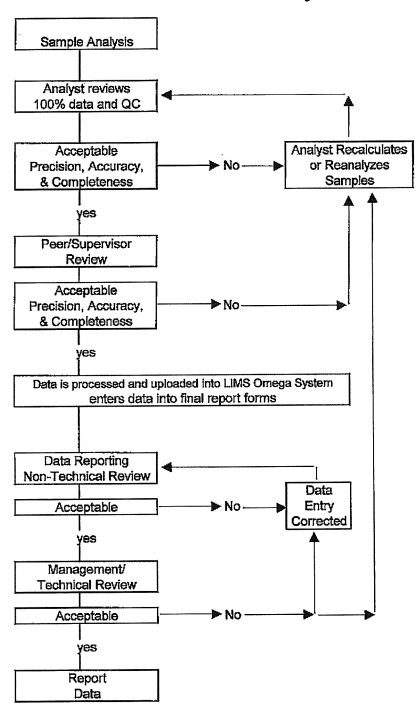
If a client should not specify which methods to be used, the methods employed by the laboratory shall be fully documented and validated. Additionally, the methods shall be published in a reputable technical journal or text or by a reputable technical organization or instrument manufacturer.

Laboratory-developed methods can be used as long as they have been documented and validated by qualified personnel. In all cases the client should be notified.

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Figure 11.6-1 Data Review Flow Diagram

MITKEM CORPORATION Review Process Flow Diagram



12.0 LABORATORY QUALITY CONTROL CHECKS

MITKEM analytical procedures are based on sound quality control methodology, which derives from three primary sources:

- 1. Specific EPA and other approved analytical methods, and
- 2. "Handbook for Analytical Quality Control in Water and Wastewater Laboratories" (EPA 600/4-79-019).
- 3. Standards for Good Laboratory Practice.

In the application of established analytical procedures MITKEM employs, at a minimum, the QC protocols described in the references found in the Analytical Methods section of this document. Specific projects may require additional quality control measures, due to such factors as difficult sample matrices or use of innovative techniques. For those projects MITKEM will recommend and implement, subject to client approval, QC measures to produce data of known quality.

Each of the MITKEM laboratory departments have an individual QC program, which includes, but is not limited to, the practices described below.

12.1 Method Detection Limit Determination/Verification:

Method Detection Limits are developed annually for certain inorganic and many organic analyses. Per NELAC Standards, MDLs are not required where target analytes are not reported below the lowest calibration standard concentration. For these analyses, results are only reported within the calibration range, and MDLs are not appropriate or needed. For certain inorganic analyses and most organic analyses, Mitkem typically reports analytes below the lowest level of the calibration range, but above the MDL, as estimated and are qualified with the "J" flag. For these analyses MDLs are developed. Mitkem reports estimated values below the calibration range for those analyses where results are able to be confirmed as in dual column confirmation, or by two concurrent determinative tests such as retention time and mass spectra as in GC/MS analyses.

To address special project requirements, MDLs can be determined for those tests which are not routinely reported below calibration range. If a client requests results to be reported below the calibration range without an MDL study, this is clearly identified in the workorder narrative.

Following an MDL study, the determined limits are verified by the analysis of an MDL Verification Standard. This standard is analyzed at approximately 2 to 3 times the calculated MDL.

12.2 Personnel Training:

Chemists who begin their employment at MITKEM are to be instructed under the MITKEM Safety Training Program within the first month. The Safety Training Program includes laboratory basics, safety video and testing, and MSDS instruction.

Before performing any analyses, a chemist is required to read the appropriate protocols and SOPs. The chemist is required to complete an SOP review form which lists all the SOPs he or she has read and understands.

The new analyst must become familiar with the laboratory equipment and the analytical methods, and begins a training period during which he or she works under strict supervision. Independent work is only permitted after the chemist successfully completes an accuracy and precision study.

The study is also commonly referred to as a Demonstration of Capability exercise. Upon the successful completion of the Demonstration of Capability exercise, the QA Department issues a Demonstration of Capability Certificate (DOCC) which is signed by both the QA Director and Operations Manager and filed in the employee's personnel folder, which is stored in the QA Department.

Demonstration of Capability studies require the acceptable recovery of 4 LCS samples for each matrix or the acceptable analysis of a blind spike sample such as a Performance evaluation sample. Acceptance limits are established by the method. It is necessary to pass the study whether for extraction and/or analysis.

Initial and on-going personnel training includes data integrity training. The 4 required elements of the data integrity system include: 1) data integrity training, 2) signed data integrity documentation, 3) in-depth, periodic monitoring of data integrity, and 4) data integrity procedure documentation.

Data integrity training topics will include the need for honesty and full disclosure in all analytical reporting, how and when to report integrity issues and what those issues could be. Employees will understand that infractions of data integrity procedures can result in an investigation that could lead to serious consequences which include immediate termination, and civil or criminal prosecution. At the start of employment all new employees read, discuss and sign a Confidentiality, Ethics and Data Integrity Agreement. Annually, an on-going integrity training session is held. An attendance sheet will be generated for every integrity session.

Data integrity procedures are reviewed and updated annually by senior management.

Training for the EPA Statement of Work occurs according to the above requirements. In addition, analysts are required to read the CLP Statement of Work as a part of the documentation training.

12.3 Control Charts:

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For organic and inorganic analyses, the recoveries of analytes in the lab control samples are plotted on control charts. These charts are used to establish control and warning limits.

12.3.1 Control limits are calculated ,compared, and/or updated at least annually from the LCS, MS/MSD, and Surrogate data points for each analyte and matrix using the following equations:

$$Average(\bar{x}) = \frac{\left[\sum_{i=1}^{n} x_i\right]}{n}$$

$$SD = \sqrt{\frac{\sum_{i=1}^{n} (x_i - \overline{x})^2}{n-1}}$$

In which:

SD = Standard Deviation N = number of data points

Warning Limits = Average $\pm 2 * SD$

Control Limits = Average $\pm 3 * SD$

12.3.2 Control limits must be approved by the QA/QC Director and by the Technical Director or Operations Manager prior to adoption by the laboratory. In the event that limits are wider than method recommended

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limits, the method recommended limits may be adopted and the analytical procedure will be re-evaluated and/or re-determined to identify possible causes. Additionally, in the event that control limits are tighter than 15% from the average, the lab may adopt a control limit of ±15% from the average. If in the experience of the laboratory, statistical control limits are unreasonably wide or narrow, alternative limits may be used until appropriate statistical limits are developed. Alternative limits are based on sources such as Department of Defense Quality Systems Manual published guidelines, EPA limits from the specific test method or from similar methods, laboratory experience with the method or other sources.

12.3.3 Control charts are plotted in EXCEL using the Omega LIMS system.

Data from each laboratory is uploaded into the LIMS. The compounds, recoveries, and date analyzed for each test are recorded in the system. In order for LIMS generated control limits to be valid, all data, including data not meeting existing recovery criteria, must be uploaded. As the laboratory uploads data for a wider range of tests, control charts will be available for these tests. Control charts may be generated for each analyte in the inorganic department to include both metals and wet chemistry parameters, and for a representative sampling of analytes in the organic sections. Each control chart is then printed for review by the QA/QC Director and by the Lab Supervisor. Out of control situations noted on the control chart are discussed with the Supervisor or Technical Director by the QA/QC Director.

An example control chart is presented as Figure 12.3-1. LCS data must be reviewed and evaluated daily against the Control Limits to establish that the system is in control.

- 12.3.4 The following situations constitute an out of control situation on a control chart:
 - One data point above or below the Control Limit line.
 - Two consecutive data points above or below the Warning Limit line.
 - Six or more consecutive data points above the Average Line or six or more consecutive data points below the Average Line. This situation suggests a trend and suggests the procedure has been changed in some way (for better or worse). The cause for this trend must be investigated.

12.4.1. Organics Laboratory:

- Trip blanks and holding blanks, when applicable, are analyzed to detect contamination during sample shipping, handling and storage.
- Method blanks, at a minimum of one in every 20 samples, are analyzed to detect contamination during analysis.
- Volatile organic method blanks are analyzed once during each analytical sequence.
- One blank spike (Laboratory Control Sample or LCS) consisting of an analytical sample of laboratory water, anhydrous sodium sulfate, or Ottawa sand with every batch of 20 or fewer samples, is analyzed to determine accuracy.
- Sample spikes and spike duplicates, as requested, are analyzed to
 determine accuracy and the presence of matrix effects. The Relative
 Percent Difference (RPD) is also determined for matrix spike/matrix
 spike duplicates to measure precision. The criteria followed are stated
 in the individual methods. For batches without a sample duplicate (for
 example, if insufficient sample volume is provided), a duplicate blank
 spike (LCSD) is performed to provide for precision measurement.
- Performance evaluation samples from EPA and state agencies are analyzed to verify continuing compliance with EPA QA/QC standards.
- Surrogate standards are added to samples and calculations of surrogate recoveries are performed to determine matrix effect and extraction efficiency.
- Internal standards for GC/MS analysis are added to sample extracts to account for sample-to-sample variation.
- GC analysis of EPA traceable standards to verify working standard accuracy and instrument performance.
- Initial multi-level calibrations are performed to establish calibration curves.
- Instrument calibration is established or verified with every analytical sequence.

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 Tuning of GC/MS systems once every 12 hours for CLP and SW-846 methods or 24 hours for methods 624/625 to method specifications is implemented for consistency in data generation.

When QC limits are not met during an analytical run, the source of the problem must be investigated. Following an evaluation of the data, those samples affected must be re-analyzed after the problem has been solved. If QC limits continue to be out of control, the instrument must be checked and/or a service call made and/or further corrective action implemented.

12.4.2. Inorganic Laboratory:

- Trip blanks are analyzed when applicable, to detect contamination during sample shipping, handling and storage.
- Method blanks are analyzed at a minimum of one every 20 samples, to detect contamination during analysis.
- One matrix spike of an analytical sample or laboratory water or soil is made and spike recoveries are calculated with every batch up to 20 samples to determine accuracy. Duplicate samples are analyzed and the RPD between the sample and duplicate is calculated for every batch up to 20 samples. If insufficient volume of sample is received, a note is made in the appropriate preparation logbook.
- Performance evaluation samples from EPA and state agencies are analyzed to verify continuing compliance with EPA QA/QC standards.
- Metals analysis instruments are calibrated for every analytical run.
- QC/LCS checks samples are analyzed during every analytical batch of up to 20 samples in order to document accuracy.

When QC limits are not met during an analytical run, the source of the problem must be investigated. Following an evaluation of the data, those samples affected must be re-analyzed after the problem has been solved. If QC limits continue to be out of control, the instrument must be checked and/or a service call made and/or further corrective action implemented.

12.5. Lab Pure Water used for method blanks and dilutions:

Mitkem uses several systems to generate analyte-free water for use in the laboratory. These systems generate high quality, analyte free water dedicated to the needs of specific analyses.

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- 12.5.1. For inorganic analyses Mitkem uses a US Filter mixed-bed deionization system followed by particle and carbon filters. This is followed by a polishing system using Barnstead E-Pure cartridges optimized for removal of inorganic constituents. Purity is monitored using an on-line electrical resistivity meter.
- 12.5.2. For organic analyses, the extractable organics laboratory uses a Barnstead E-Pure system optimized for removal of organic constituents. The volatile organics laboratory uses an in-house activated carbon filtration system to provide analyte free water. As organic contaminants are not measured by a resistivity meter, this is not a relied-upon method to monitor the quality of organic analyte-free water. Instead, laboratory method blanks are used, typically several per working day, to monitor the acceptability of the water for its intended use. Any analyte detected above (half of) the reporting limit is investigated. If this can be traced to the water purification system as its source, maintenance is performed on the water purification system.

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Figure 12.3-1
Example Control Chart

Mitkem Corporation

REC QUALITY CONTROL CHART

Date: 20-Dec-06

Test Code: SW8260B_W Analyte: BROMOFLUOROBENZENE

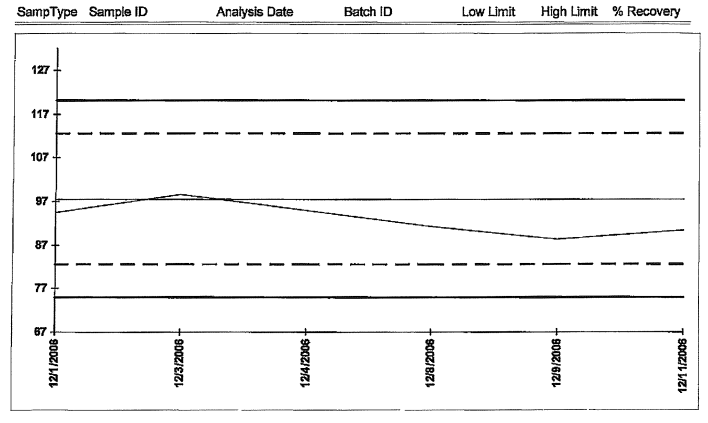
		· · · · · · · · · · · · · · · · · · ·	THO CHODEN IN			
SampType	Sample ID	Analysis Date	Batch ID	Low Limit	High Limit	% Recovery
SAMP	E1838-08A	12/1/2006	27316	75	120	92.6
SAMP	E1838-06A	12/1/2006	27316	75	120	98.9
SAMP	E1838-05A	12/1/2006	27316	75	120	91.7
MBLK	MB-27333	12/3/2008	27333	75	120	92.6
LCS	LCS-27333	12/3/2006	27333	75	120	104.3
SAMP	E1838-01A	12/3/2006	27333	75	120	98.2
SAMP	E1838-03A	12/3/2006	27333	75	120	100.7
SAMP	E1838-02A	12/3/2008	27333	75	120	97.6
SAMP	E1838-04A	12/4/2006	27340	75	120	76.1
SAMP	E1838-07A	12/4/2006	27340	75	120	108.1
LCS	LCS-27340	12/4/2006	27340	75	120	102.2
MBLK	MB-27340	12/4/2006	27340	75	120	95.7
MBLK	MB-27441	12/8/2006	27441	75	120	90.4
SAMP	E1878-04A	12/8/2006	27441	75	120	89.6
SAMP	E1879-05A	12/8/2006	27441	75	120	89.3
SAMP	E1879-04A	12/8/2006	27441	75	120	90.4
SAMP	E1879-03A	12/8/2006	27441	75	120	89.5
SAMP	E1879-02A	12/8/2006	27441	75	120	88.6
SAMP	E1879-01A	12/8/2006	27441	75	120	87.6
SAMP	E1878-07A	12/8/2006	27441	75	120	88.4
SAMP	E1878-06A	12/8/2006	27441	75	120	90.0
SAMP	E1878-05A	12/8/2006	27441	75	120	89.1
SAMP	E1879-06A	12/8/2006	27441	75	120	89.3
LCSD	LCSD-27437	12/8/2006	27437	75	120	97.4
MBLK	MB-27437	12/8/2006	27437	75	120	93.1
LCSD	LCSD-27441	12/8/2006	27441	75	120	91.4
SAMP	E1871-04A	12/8/2006	27437	75	120	98.9
SAMP	E1878-03A	12/8/2006	27441	75	120	89.2
SAMP	E1871-02A	12/8/2006	27437	75	120	92.7
SAMP	E1871-03A	12/8/2006	27437	75	120	92.3
SAMP	E1871-01A	12/8/2006	27437	75	120	91.8
LCS	LCS-27441	12/8/2006	27441	75	120	92.3
SAMP	E1878-02A	12/8/2006	27441	75	120	89.3
LCS	LCS-27437	12/8/2006	27437	75	120	96.9
SAMP	E1879-07A	12/9/2006	27441	75	120	88.2
SAMP	E1843-02A	12/11/2008	27471	75	120	90.7
MBLK	MB-27471	12/11/2006	27471	75	120	89.6
LCS	LCS-27471	12/11/2006	27471	75	120	92.7
LCSD	LCSD-27471	12/11/2006	27471	75	120	90.9
SAMP	E1878-01A	12/11/2006	27471	75	120	87.2

Mitkem Corporation

REC QUALITY CONTROL CHART

Date: 20-Dec-06

Test Code: SW8260B_W Analyte: BROMOFLUOROBENZENE



13.0 QUALITY ASSURANCE SYSTEMS AUDITS, PERFORMANCE AUDITS AND FREQUENCIES

The MITKEM Quality Assurance Director and/or staff performs routine internal audits of the laboratory. The frequency of such audits depends on the workload in-house but is done annually, at a minimum. The audits entail reviewing laboratory logbooks and all appropriate operations to ensure that all laboratory systems including sample control, analytical procedures, data generation and documentation meet contractual requirements and comply with good laboratory practices.

13.1 System Audits:

The QA/QC Director audits each individual laboratory annually in order to detect any sample flow, analytical or documentation problems and to ensure adherence to good laboratory practices as described in MITKEM's Standard Operating Procedures and Quality Assurance Plan. An example checklist used in an internal systems audit at MITKEM is presented in Figure 13.1-1.

Areas covered by the internal audit include logbook documentation and review, standard traceability, standard storage and expiration dates, method criteria adherence, instrument maintenance records, SOP review, and knowledge/training of the analysts. Often, deficiencies that have been noted during "outside" audits and outstanding Corrective Actions will also be reviewed.

Upon the completion of the internal audit, a formal audit report is presented to the laboratory supervisor who is given a specific timeframe to respond in writing regarding the deficiencies. The QA Department will do a follow up audit to check that at least the major deficiencies have been corrected. The follow-up audit occurs within 30-45 days from the date of the lab's audit response.

13.2 Performance Audits:

MITKEM participates in external Performance Test (PT) studies under the National Environmental Accreditation Program (NELAP) through the State of New Jersey (Mitkem Laboratory's Primary Accreditation Authority). The QA department of the laboratory administers the Performance Evaluation Samples for Wastewater/Solid Waste (WW/SHW). The Performance Evaluation Samples generally follow a quartely schedule, with wastewater alternating with soil/solid waste.

Several times a year outside agencies (federal, state, or private) may schedule an audit at Mitkem in order to check the laboratory's processes. Most often these audits begin and end with a meeting between auditors and laboratory management. Each individual laboratory is then examined. The QA Department

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and/or Senior Management Staff are most likely to remain with the auditors at all times during the audit.

Sometime after the audit, Mitkem receives a formal written audit report to which it must respond. The audit report is initially reviewed by the QA Director who may copy and distribute the report to each laboratory supervisor. In several instances, the report is sent electronically and supervisors may receive an electronic version. The supervisors are required to respond to the findings that pertain to his or her department. The QA Director compiles the formal corrective action plan that may undergo several revisions before the auditing authority accepts it.

The QA Director then sends a memo to each supervisor to detail what needs to be done in each department within a specific timeframe. The QA Department then follows up with the labs to ensure procedures have been modified and the corrective actions are in place. In some instances, a LIMS corrective action report is also initiated as a result of an audit finding.

Internally, performance is monitored on a daily basis at MITKEM through the use of surrogate standards, LCS, and MS/MSD samples. Check samples from independent commercial sources are employed routinely in each of the MITKEM laboratory departments and ensure continuing high-level performance. The QA Director may distribute internal blind PE samples to each laboratory department. These blind PE samples can also be used to show on-going analyst proficiency in lieu of 4 LCS studies.

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Figure 13.1-1 QA Systems Audit Checklist

1.7

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~ .	Laboratory Audit		
Date:	Auditor:	Department:	
Category:			
1. Facility			
•			
Adequate work areas, c	ounter space		
Chemical storage areas	Acids, Flammables		
Eyewash, showers, insp	ected?		
Tanks Secured			
Hoods calibrated, adequ	uate		
2. Personnel			
Do analysts follow the SO			Yes/No
	emonstration of proficiency	y study?	Yes/No
Are analysts adequately tr	ained and knowledgeable?		Yes/No
Wearing appropriate PF	Es		
Dressed appropriately_			
Trained in procedure, tr	aining documented?		

Page 2 of 4

Passed proficiency Documented?		
3. SOPs		
Standard_Operating_Procedures		
Are the general SOPs updated annually?		Yes/No
Are SOPs updated annually for each analytical method?		Yes/No
Are SOPs controlled documents?		Yes / No
Are SOPs signed by appropriate individuals?		Yes/No
Notes		
		· · · · · · · · · · · · · · · · · · ·
		And the second s
4. Chemicals		
Labeled correctly?		
Chemicals stored correctly?		
Chamberly broken bolivery .		

Standard ID#	•	
Standards traceable?		
Are standards QC'd against a second source after each ICAL?		Yes / No
Are standards traceable throughout the lab?		Yes / No
Are expired standards present in the lab?		Yes/No
Is there a defined system for assigning expiration dates? Yes		
Is standard freezer temperature monitored? Yes		

Notes

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5. Logbooks				
Does a run logbook exist for each analytical instrument?			Yes/No	
Does an instrument maintenance log exist for each instrument? Are logbooks peer reviewed weekly? Proper correction techniques?			Yes/No	
			Yes/No	
			Yes / No	
Empty spaces "z"d o	ut?			Yes / No
Paginated?				Yes/No
Controlled?				Yes / No
Do logbooks contain	all pertine	nt information to the procedu	ure?	Yes/No
(I.e., method, matrix,	reagent lo	t #, etc.)		
				•
Clear, legible, corre In-dated,	ctions com	plete		
6. Equipment	•			
General_Laboratory_Ec				Yes/No
s an NIST traceable thermometer available?		Yes/No		
Are lab thermometers calibrated annually against the NIST thermometer?				
Are correction factors in use on lab thermometers?			Yes/No	
Are Class "S" weights calibrated NIST every 2 years?		Yes/No		
Are balances serviced annually?		Yes/No		
Are balances calibrated as needed and the calibration recorded?		Yes/No		
s balance calibration a	acceptance	criteria clearly defined and	posted?	Yes / No
Maintenance				

Preventive maintenance	Page 4 of 4	
Calibrated:		
Scales		
Thermometers		The state of the s
Hoods		WO TOWN SHA A class of the section o
Syringes		**
Timers		
Equipment stored correctly, Glassware, syringes, tools e	etc.	
•		
Analytical_Methods		
s ICAL documentation maintained on file in the lab?		Yes / No
When %RSD > 15%, is the average adopted?		Yes/No
s a CCV run at the end of the analytical sequence? (USAC	CE)	Yes/No
Is a Method Blank analyzed after each CCV?		Yes/No
Does analyst review data for false negatives?		Yes/No

14.0 PREVENTIVE MAINTENANCE

Preventive maintenance is a routine practice at MITKEM for all instrumentation. Scheduled preventive maintenance minimizes instrument downtime and subsequent interruption of analysis. All major instrumentation is under service contracts so that downtime (due to catastrophic events) is minimized.

Only those equipment items meeting or exceeding applicable performance requirements are used for data collection. This includes items such as laboratory balances as well as major analytical instruments such as ICPs, GCs and GC/MSs.

MITKEM's laboratory personnel are familiar with the routine and non-routine maintenance requirements of the instruments they operate. This familiarity is based on education, hands-on experience and manufacturer's training courses.

GC Maintenance:

- 1. The injection septum will be replaced once approximately fifty (50) injections or earlier if a leak develops.
- The injection liner will be replaced once approximately fifty (50) injections or when initial and/or continuing calibrations fails repeatedly to meet method requirements.
- 3. The gold seal will be replaced except for septum and liner, and the column will be trimmed whenever an initial calibration is run.
- 4. The column will be replaced if chromatograms show excessive peak tailing and/or initial and continuous calibration verifications fail repeatedly to meet method requirements.

GC/MS Maintenance:

- 1. GC injector and liner are cleaned daily for semivolatiles and monthly for volatiles.
- The column will be replaced if chromatograms show excessive peak tailing and/or initial and continuous calibration verifications fail repeatedly to meet method requirements.
- 3. The ion source will be cleaned when initial and/or continuing calibration repeatedly fail method specified criteria.

4. The pump oil will be replaced once a year.

ICAP Maintenance:

- 1. Peristaltic pump tubing will be replaced every sixteen (16) hours of instrument time or sooner when memory effects are manifested.
- 2. The plasma torch is cleaned with (aqua regia) every 1-2 weeks. If memory effects are manifested the torch will be cleaned immediately.
- 3. The sample introduction (spray chamber and nebulizer) is cleaned every 2-3 weeks.
- 4. Air filters are cleaned each time the torch is cleaned or as needed upon visual inspection.
- 5. Once every six (6) months, under service contract, the instrument undergoes extensive maintenance by a manufacturer's service engineer.

Mercury FIMS 100 Maintenance:

- 1. Pump tubing is replaced every 48 hours of instrument run time.
- 2. Sample loops, gas tubing extensions and sample capillaries are replaced as needed.

Lachat 8000 Maintenance:

W. So

- 1. All pump tubing is replaced every 48 hours of instrument run time.
- 2. Auto sampler arm is lubricated every 48 hours of instrument run time.
- 3. The manifolds, tubing connections, valves, etc. are cleaned or replaced as needed.

TCLP/SPLP Tumbler Maintenance:

- 1. The tumbler is checked at every use for number of rotations per minute (30rpms), the ambient temperature checked and documented in the RPS Logbook.
- 2. If the tumbler is not spinning at 30rpms, motor is cleaned and oiled.
- 3. If tumbler is not spinning at 30rpms after maintenance, the motor will be replaced.

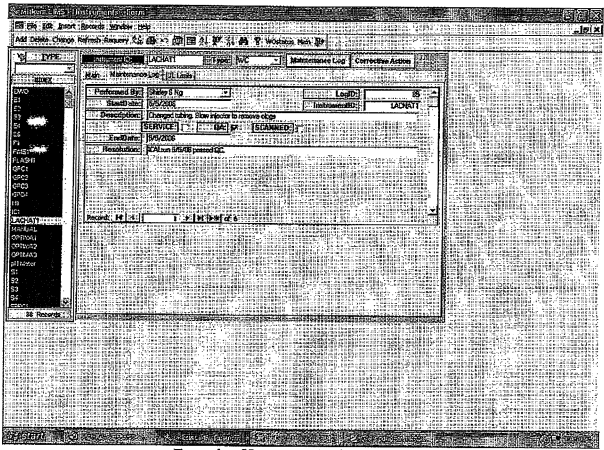
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Instrument maintenance logs are kept for each instrument in the OMEGA LIMS System (figure 14-1). All employees have access to the LIMS system. The person performing the maintenance is required to provide the following information in the online log:

- Equipment identifier
- The inspection, maintenance, calibration or corrective action(s) performed.
- The trigger(s) for the maintenance action(s)
- The identity of the person(s) performing the maintenance
- The date on which the work was performed, and
- The condition of the equipment upon completion of the work.

MITKEM maintains an inventory of replacement parts required for preventive maintenance and spare parts that often need replacement, such as filaments for GC/MS systems and the more mundane electrical fuses and GC column ferrules. To control cost, the appropriate supervisor shall decide the types and numbers of spare parts kept on hand for each equipment item.

Figure 14-1



Example of Instrument Maintenance Log

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Figure 14-2 Instrument Maintenance Schedule Figure 14-2
MITKEM CORPORATION
Preventive MaIntenance Schedule

Instrument	A control of the cont	
	לוואחער	Frequency
Gas Chromatograph (GC)	Injection septum replaced Injection liner replaced The column will be replaced if chromatograms show excessive paak tailing and/or initial and continuing calibration verifications fall repeatedly to meet method requirements.	Every 50 Injections Every 50 Injections As needed
GC/MS	GC injector and liner replaced The column will be replaced if chromatograms show excessive peak tailing and/or initial and continuing calibration verifications fall repeatedly to meet method requirements. The ion source will be cleaned when initial and/or continuing calibration repeatedly fall method specified criteria. The pump oil is replaced.	Dally As needed As needed As needed
Inductively Coupled Plasma (ICP)	Peristaltic pump tubing is replaced The plasma torch is cleaned (aqua regia). The sample introduction (spray chamber and nebulizer) is cleaned Air filters are cleaned. The instrument undergoes extensive maintenance by the manufacturer's service engineer.	Every 16 hours of instrument run time time Weekly Weekly Biweekly Biweekly Semiannually
Mercury FIMS 100	Pump tubing is replaced Sample capillary and tubing are replaced Inside of optical cell is cleaned	Every 48 hours of instrument run time Every 48 hours of instrument run time Every 48 hours of instrument run time time
Lachat 8000	All pump tubing is replaced Autosampler arm is lubricated The instrument undergoes extensive maintenance by the manufacturer's service engineer.	Every 48 hours of Instrument run time Every 48 hours of Instrument run time Semiannually

15.0 SPECIFIC ROUTINE PROCEDURES USED TO ASSESS DATA PRECISION, ACCURACY, COMPLETENESS, METHODS DETECTION LIMIT AND LINEAR DYNAMIC RANGE

These mathematical equations represent the means of calculating analytical figures of merit on a routine basis at MITKEM. However, they may be supplanted with other calculations if requested by the client. Precision, accuracy and completeness are also discussed in Section 6.

15.1 Precision:

Precision is frequently determined by the comparison of replicates, where replicates result from an original sample that has been split for identical analyses. Standard deviations, s, of a sample are commonly used in estimating precision.

Sample standard deviation, s:

$$s = \sqrt{\frac{1}{n-1}\sum_{i=1}^{n}(x_i - \overline{x})^2}$$

where a quantity, x_i (e.g. a concentration), is measured n times with a mean, \bar{x} .

The relative standard deviation, RSD (or sample coefficient of variation, CV), which expresses standard deviation as a percentage of the mean, is generally useful in the comparison of three or more replicates (although it may be applied in the case of n=2).

$$%RSD = 100 (s / \bar{x})$$

or

$$CV = 100 (s / \overline{x})$$

In which: RSD = relative standard deviation, or CV = coefficient of variation s = standard deviation

 $\overline{x} = \text{mean}$

For duplicates (samples that result when an original sample have been split into two for identical analyses), the relative percent difference (*RPD*) between the two samples may be used to estimate precision.

$$RPD = \frac{2(D_1 - D_2)}{(D_1 + D_2)} \times 100\%$$

In which: D_I = first sample value D_2 = second sample value (duplicate)

15.2 Accuracy:

The determination of accuracy of a measurement requires knowledge of the true or accepted value for the signal being measured. Accuracy may be calculated in terms of bias as follows:

$$Bias = X - T$$

$$\%Bias = 100 \frac{(X-T)}{T}$$

In which: X = average observed value of measurement T = "true" value

Accuracy also may be calculated in terms of the recoveries of analytes in spiked samples:

% Re cov ery(%R) =
$$100 \times \frac{(SSR - SR)}{SA}$$

where: SSR = spikes sample result SR = sample result

SA =spike added

15.3 Completeness:

Determine whether a database is complete or incomplete may be quite difficult. To be considered complete, the data set must contain all QC check analyses verifying precision and accuracy for the analytical protocol. Less obvious is whether the data are sufficient to achieve the goals of the project. All data are reviewed in terms of goals in order to determine if the data set is sufficient.

Where possible, the percent completeness for each set of samples is calculated as follows:

15.4 Method Detection Limit:

The method detection limit (MDL) is the minimum concentration of a substance that can be measured and reported with 99% confidence that the analyte concentration is not zero. It is computed as follows from data obtained by repeatedly determining an analyte in a given sample matrix:

- Analyze at least seven samples of a homogeneous matrix spike that
 contains the analyte(s) of interest at concentrations of three to five times
 the expected MDL. The entire sample preparation and analysis protocol
 must be applied in each analysis; simply preparing one sample and
 repeating a measurement three or more times on the sample in not
 acceptable.
- 2. Upload the acceptable data into LIMS Omega.
- 3. The LIMS will compute the standard deviation of the results for each analyte using the following equation:

$$MDL = t_{(n-1, \alpha=0.99)}(s)$$

Where t is the one-sided student's t value appropriate for the number of samples analyzed, n; α is the statistical confidence level; and s is the standard deviation.

The one-sided t-values are presented below:

Number of samples	<u>t-value</u>
7	3.14
8 .	2.996
9	2.90
10	2.82

- 4. The MDL is then checked against 40CFR136 requirements by the QA Department. If the MDL is acceptable then it is uploaded into the LIMS by either the QA Department or LIMS Administrator.
- 5. Immediately following the determination of the MDL, MDL check samples are analyzed at a concentration approximately equal to 2 x the new MDL. The analyte of interest must be detected at this concentration, or the MDL may require raising.
- 6. An elevated MDL can be uploaded if necessary into the LIMS as long as documentation is available to show that the applicable method can produce an MDL at least that low. This can commonly occur for ICP

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analysis in which extremely low MDLs can cause method compliance issues.

15.5 Linear Dynamic Range:

The linear dynamic range is the concentration range over which the instrument response is linear. It is determined by analyzing a series of standard solutions that extends beyond the non-linear calibration region at both the low and high extremes, and selecting that range of standards which demonstrates a linear relationship between instrument response and concentration.

For ICP analysis, the linear dynamic range is determined by analyzing each metal at 3 different concentrations. The concentration which produces results within a 10% error is determined to be the linear dynamic range. This procedure must be performed per individual method requirements.

ILM5.3 requires the analysis of the linear dynamic range be determined quarterly, with a 5 % error.

16.0 CORRECTIVE ACTION

An essential element of the QA Program, Corrective Action provides systematic, active measures taken in the resolution of problems and the restoration of analytical systems to their proper functioning.

Corrective actions for laboratory problems are described in MITKEM Corporation laboratory standard operating procedures. Personal experience often is most valuable in alerting the bench scientist to questionable results or the malfunctioning of equipment. Specific QC procedures are designed to help the analyst determine the need for corrective actions (see Section 11, Data Reduction, Validation and Reporting). Corrective actions taken by scientists in the laboratory help avoid the collection of poor quality data. Mitkem's corrective action program divides these issues into routine and non-routine corrective actions as described below.

Routine Corrective Action — A routine corrective action is taken when the out-of-control event encountered is one that is detected at the appropriate level in the QA process. Routine corrective actions are defined in the analytical SOP with specific steps to be taken as corrective action (i.e., low surrogate recovery, continuing calibration verifications, project specific protocols that do not meet acceptance criteria, etc.) Routine corrective actions must be documented as described in the analytical SOP, but do not require further documentation in the corrective action logbook. Examples of routine corrective action situations: surrogate/surrogates out, LCS out, CCV out, ICV out, IS area/areas out, typographical errors, random blank contamination, or false positive hit/spectral ID match corrected during data review.

Non-Routine Corrective Action — A non-routine corrective action is taken when the outof-control event encountered is not typical for the method. For example, QC failures that
pass through the final review to the client, procedural errors — not following the SOP, or a
situation not being detected by normal QA procedures that could adversely impact the
accuracy, precision, etc. of a result. Non-routine corrective actions must be documented
in the Corrective Action Request (CAR) logbook. The analyst, using his/her own
judgement, may deem any corrective action situation non-routine and formally document
it on a CAR. When in doubt about a corrective action, the analysts are instructed to err
on the side of formal CAR documentation. Examples of non-routine corrective action
situations include: bad standard, expired standard mix being used, incorrect equation,
"client-detected" problems, not following SOP protocols, using bad or contaminated lot
of chemical/reagent/solvent, deciding to release data not conforming to SOP
requirements, compound retention time outside of range, or improper library spectrum
that leads to re-occurring mis-identification of compounds.

The essential steps in MITKEM Corporation corrective action system are:

- 1. Identify and define the problem.
- 2. Assign responsibility for investigating the problem. Usually this individual is the department supervisor.
- Investigate and determine the cause of the problem.
- 4. Determine a corrective action to eliminate the problem and prevent recurrence. Any changes that result from the corrective action investigation must be documented.
- Assign and accept responsibility for implementing the corrective action.
- 6. Establish effectiveness of the corrective action and implement it.
- 7. Verify that the corrective action has eliminated the problem.
- 8. Both the laboratory and the QA Department need to monitor the corrective action to ensure it is effective.
- 9. Any corrective actions that cast doubt on the laboratory's compliance with its own policies and procedures may require an internal audit by the QA Department.

This scheme is generally accomplished through the use of Corrective Action Report Forms available to each of MITKEM's laboratories within the OMEGA LIMS system. Use of this report notifies the QA Department of a potential problem as described in SOP No. 80.0007. The QA Director initiates the corrective action by relating the problem to the appropriate laboratory managers and/or project managers who then investigate or assign responsibility for investigating the problem and determine its cause. Once determined, the QA Director will approve appropriate corrective action. Its implementation is later verified through and internal laboratory audit. Once the QA Director feels the system has returned to control, s/he will finalize the CAR using a password protected QA step.

Information contained on corrective action forms is kept confidential within MITKEM and is generally limited to the individuals involved. Severe problems and difficulties may warrant special reports to the President of MITKEM who will ensure that the appropriate corrective actions are taken.

Nonconformance:

1967 TUS.

Any breech of standard protocols is a nonconformance item that is documented on the Corrective Action Request Form and management informed immediately. The following are nonconformance items:

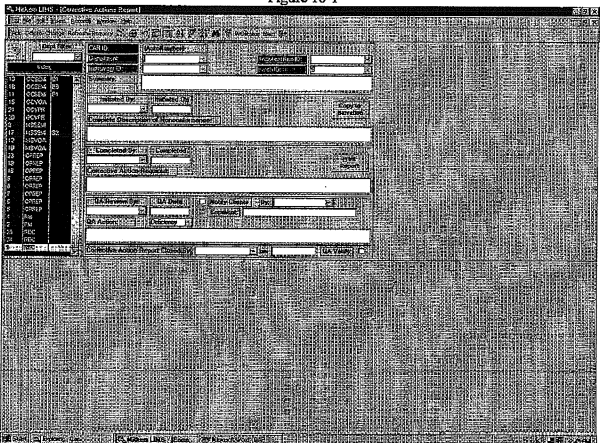
- 1. Sample holding time exceeded.
- 2. Hoods, Class "S" weights, NIST Thermometers, balances, automatic pipettes, being used but not certified.
- 3. Expired standards being used.
- 4. Manual integration being misrepresented.

16.1 Client Complaints:

Mitkem Corporation ensures client complaints are dealt with quickly and completely. The policies are stated in the laboratory Client Complaint Standard Operating procedure (SOP No. 80.0002).

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Figure 16-1



Quality Assurance Corrective Action Request Form

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17.0 QUALITY ASSURANCE REPORTS TO MANAGEMENT

The MITKEM Quality Assurance Director submits a QA report annually to the Operations Manager and the President of the Laboratory. The report should be completed and submitted no later than the 15th of July in any calendar year.

The report contains detailed laboratory information and QA activities during the previous twelve months. Items to include are the status of internal and external audits, client complaints, quality control activities, resources and staffing. See the following pages for the report format.

Management will review the QA report and respond to outstanding issues. Management will add a review of the suitability of policies and procedures, and any other relevent issues. The response report is due within 30 days of the QA Report receipt.

A copy of the report is kept on file in the QA department.

In case of a severe problem or difficulty, a special report is prepared by the QA Director and submitted immediately to management.

MITKEM CORPORATION Annual Quality Assurance Report to Management

1.	Status of Internal Audits.
<u>2.</u>	Status of External Audits
3.	Identification of Quality Control issues in the laboratory.
4.	Discussion of corrective action issues.

5. Proficiency Testing.

6. Changes in volume and type of work undertaken.

7. Client Feedback.

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8. Reports from management and supervisory personnel.

18.0 SAFETY

MITKEM maintains safety program managed by the Health and Safety Officer and the Safety Committee. Responsibilities include many aspects that comply with the Right-to-Know Laws. Training includes:

- Training seminars with information on OSHA safety instruction for new employees.
- Introductory training to include location of fire extinguishers, first aid supplies, etc.
- Chemical Hygiene Plan/Health and Safety manual review when hired initially and then annually thereafter.
- Monthly Safety Committee meetings.
- Centralized MSDS information.
- Maps with safety equipment and all exits noted.
- Posted safety rules.

If a chemical spill occurs, proper actions are described in Mitkem's Contingency Plan. Each department at Mitkem has its own copy of the Contingency Plan. Additionally, the local fire department (Warwick) and hospital (Kent County) also have a copy in case a need arises. All employees are required to review the plan when hired.

Emergency equipment, such as spill control kits, fire extinguishers and fire blankets are located throughout the laboratory areas. The Contingency Plan has instructions for evacuation, notification of emergency authorities and regulatory personnel in the event of a chemical accident.

19.0 WASTE MANAGEMENT

19.1 Pollution Prevention

The waste management option of choice is to prevent pollution by minimizing the amount or types of chemical wastes that are generated. Mitkem's ability to minimize waste generation is limited by the chemical analysis techniques that are required by the EPA or other authors of test methods. As new test methods are utilized in the laboratory, the type and volume of chemical waste generated by the new test is considered. Analysts and Supervisors are encouraged to look for ways to reduce the amount of chemical waste, or the type of chemical waste generated during the testing process; HOWEVER, no method is allowed to be modified without discussion among the Supervisor, Technical Director, QA Director and other management personnel to determine the affect of the change on the resulting data.

19.2. Waste Management

Mitkem has identifies and routinely disposes of chemical wastes in several hazardous waste streams. In general these are acids, caustics, solvent wastes and various laboratory waste solids. No laboratory chemical waste is disposed in the trash or dumped down the drain. All remaining sample volume following testing, and after contract-required disposal date has past, are disposed in one of these waste streams. These wastes are fully described in Mitkem's Waste Management Plan and in Mitkem's Profile Log that has been prepared by Univar, Mitkem's waste hauler. Other hazardous wastes are identified and properly disposed according to these documents.

Continued compliance is monitored monthly by an outside consultant to ensure all RI DEM regulations are met.

20.0 DEFINITIONS, ACRONYMS, ABBREVIATIONS:

ACCURACY: The closeness of agreement between an observed value and An accepted reference value.

BATCH: A group of samples of the same matrix that are processed as a unit.

Unless defined differently by a specific analytical method (such as Oil &

Grease by Method 1664), the maximum batch size is 20 samples.

BIAS: The deviation due to analytical or matrix effects of the measured value

from a known spiked amount.

BLANK: A "clean" matrix analysis. Such as: Equipment Blank, Method Blank,

Trip Blank.

CAS: Chemical Abstracts Service, a registry where chemicals are assigned

identification numbers.

CCB: Continuing Calibration Blank

CCV: Continuing Calibration Verification standard.

CLP: Contract Laboratory Program. A contract used by EPA to purchase

analytical services. Also refers to the test protocols described in that contract. The CLP analyses can be used for EPA or for other clients. CLP-format data reports are arranged as described in the EPA CLP contract, including specified data report pages and all raw data. The CLP analysis scheme includes OLM (Organic Low/Medium-soil and water), OLC (organic low concentration-waters only) and ILM (Inorganic

Low/Medium-soil and water) analyses.

CONTROL A QC sample introduced into a process to monitor the

SAMPLE performance of the system.

DL: Dilution, not used when the initial analysis is performed at dilution, but is

used for a secondary dilution.

DUPLICATE: see Matrix Duplicate, Field Duplicate, and Matrix Spike

Duplicate.

EQUIPMENT A sample of analyte-free water that has been used

BLANK during sample collection to measure any contamination introduced during

sample collection.

ICB: Initial Calibration Blank

ICV:

Initial Calibration Verification standard

IDL:

Instrument Detection Limit. Statistical value similar to MDL, but with analyses performed on standards that have not been through the sample preparation process.

FIELD

Independent samples that are collected as close as DUPLICATES possible to the same point in space and time. They are two separate samples taken from the same source, stored in separate containers, and analyzed independently. These duplicates are useful in documenting the precision of the sampling process.

LAB

A blank spiked with compound(s)

CONTROL

representative of the target analytes. This is used to document laboratory SAMPLE(LCS)performance in a "clean" matrix.

MATRIX:

The component or substrate (e.g., water, soil, air, and oil) which contains the analyte of interest.

MATRIX

A sample split by the laboratory that is used

DUPE (DUP) to document the precision of a method in a given sample matrix.

MATRIX

An aliquot of sample spiked with a known

SPIKE (MS)

concentration of target analyte(s). The spiking occurs prior to sample preparation and analysis. A matrix spike is used to document the bias of a method in a given sample matrix.

MATRIX SPIKE

Laboratory split samples spiked with identical concentrations of target analyte(s). The spiking occurs prior to sample preparation and analysis. DUPE (MSD) They are used to document the precision and bias of a method in a given Sample matrix.

METHOD

An analyte-free matrix to which all reagents are

BLANK (MB) added in the same volumes or proportions as used in sample processing, The method blank should be carried through the complete sample preparation and analytical procedure. The method blank is used to document contamination resulting from the analytical process.

METHOD DETECTION LIMIT (MDL) The minimum concentration of a substance that can be measured and reported with 99% confidence that the analyte concentration is greater than zero and is determined from analysis of a sample in a given matrix type containing the analyte. For operational purposes, when it is necessary to determine the MDL in the matrix, the

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MDL should be determined by multiplying the appropriate one-sided 99% t-statistic by the standard deviation obtained from a minimum of seven analyses of a matrix spike containing the analyte of interest at a concentration estimated to be three to five times the MDL, where the t-statistic is obtained from standard references.

MSA:

Method of Standard Additions

ND:

Not Detected. Used in conjunction with the reporting limit.

ORGANIC-FREE REAGENT WATER: For volatiles, all references to water in the methods refer to water in which an interferent is not observed at the reporting limit of the compounds of interest. Organic-free reagent water can be generated by passing tap water through a carbon filter bed containing about 1 pound of activated carbon. A water purification system may be used to generate organic-free deionized water.

For semivolatiles and nonvolatiles, all references to water in the methods refer to water in which an Interferent is not observed at the reporting limit of the compounds of interest. Organic-free reagent water can be generated by passing tap water through a carbon filter bed containing about 1 pound of activated carbon.

A water purification system may be used to generate organic-free deionized water.

PPB:

Parts Per Billion, ug/L, ug/Kg

PPM:

Parts Per Million, mg/L, mg/Kg

PQL:

Practical Quantitation Limit. Is equivalent to Reporting Limit.

PRECISION: The agreement among a set of replicate analyses.

PS:

Post Spike. Spike added at the analysis level (as opposed to at the beginning of sample preparation) to determine interferences.

REPORTING LIMIT: The lowest concentration that can be reliably achieved within specified limits of precision and accuracy during routine laboratory operating conditions. The RL is generally 5 to 10 times the MDL. However, it may be nominally chosen other than these guidelines to simplify data reporting. For many analytes the RL concentration is selected as the lowest non-zero standard in the calibration curve. Sample RLs are matrix-dependent, and are adjusted by the amount of sample analyzed, dilution, percent moisture.

RE:

Reextraction or Reanalysis

RPD:

Relative Percent Difference, used to determine precision.

RRF:

Relative Response Factor. Used for quantification with the internal

standard procedure.

RT:

Retention Time for a chromatographic peak, as calculated from the time of

injection.

SD:

Serial Dilution

STANDARD ADDITION: The practice of adding a known amount of an analyte

to a sample immediately prior to analysis. It is typically

used to evaluate interferences.

STANDARD CURVE: A plot of concentrations of known analyte standards

versus the instrument response to the analyte. Calibration standards are prepared by successively diluting a standard solution to produce working standards which cover the working range of the instrument. Standards should be

prepared at the frequency specified in the appropriate method. The

calibration standards should be prepared

using the same type of acid or solvent and at the same concentration as will result in the samples following sample preparation. This is applicable to organic and

inorganic chemical analyses.

SURROGATE: An organic compound that is similar to the target

analyte(s) in chemical composition and behavior in the analytical process, but which is not normally found in

environmental samples.

TRIP BLANK: A sample of analyte-free media taken from the laboratory

to the sampling site and returned to the laboratory unopened. A trip blank is used to document contamination attributable to shipping and field handling procedures. This type of blank is useful in documenting contamination

of volatile organics samples.

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MITKEM CORPORATION INSTRUMENTATION and EQUIPMENT LIST APPENDIX A

Weight Set Identification:

- 1. WT1-Organic Prep Weight Set
- 2. WT2-Organic Prep 100g
- 3. WT3-Organic Prep 300g
- 4. WT4-Organic Prep 1kg
- 5. WT5-Inorganics Weight Set
- 6. WT6-VOA Weight Set

Mitkem Corporation Balance List

			Date	Date in	Condition	Equipment	
Equipment	Manufacturer	Serial #	Received	Service	New/Used	. Ω	Location
TOP-LOADING Balance	OHAUS	1121230069	2000	2000	New	TL10	Organic
Analytical Balance	Denver	0077138	1995	1995	New	AB-1	Inorganic
TOP-LOADING Balance	OHAUS Voyager	F2921120391055	2001	2001	New	TL9	Inorganic
TOP-LOADING Balance	Denver	0079896	2000	2000	New	TL1	Metals
TOP-LOADING Balance	OHAUS Precision Std.	C22427176	2002	¥.	New	TL6	Backup
TOP-LOADING Balance	OHAUS Navigator	1121122373	2002	2002	New	TL11	Unit 3
TOP-LOADING Balance	OHAUS	CD8910	2000	2000	New	TL4	VOA
TOP-LOADING Balance	OHAUS Navigator	1122173423	2003	NA	New	TL12	Inorganic
	-						

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Mitkem Corporation Equipment List

Receiving Location Unit 3 Unit 3 Equipment ID New/Used Condition nsed nsed New Service Date In Received Date Department: Receiving TD-12-90-133 Model: TD-00008-A Not Applicable 600011006 Serial # Dura-Stop MP, Dura-Dry MP Manufacturer Thelco Lab Oven Equipment Walk in Cooler % Solid Oven Freeze Dryer

Witkem Corporation Equipment List

Department: Organic Prep

Equipment	Manufacturer	Serial #	Date Received	Date in Service	Condition New/Usec	Equipment ID	Location
Vortex Concentrator	Labconco	000493001C	96-Inf	weN 86-Jnf	New	RVI	O Prep
	1	7405050	0	4			£ (
Vollex Concenitator	Labconco	10080100	ADI-88	Apr-99 New	MeM	= ^	O Prep
Vortex Concentrator	Labconco	011196291E	Jun-01	Jun-01 New	New	RV III	O Prep
Vortex Concentrator	Labconco	246368	Dec-05	Jan-06 Used	Used	RV IV	O Prep
Vortex Concentrator	Labconco	266438	Dec-05	Jan-06 Used	Used	RV V	O Prep
Vortex Concentrator	Labconco	246505	Dec-05	Jan-06 Used	Used	RVVI	O Prep
Vortex Concentrator	Labconco	266818	Dec-05	Jan-06 Used	Used	RVVII	O Prep
Nitrogen Concentrator Bath	Organomations	17033	Jun-97	Jun-97 New	New	NZ1	O Prep
Deionized Water Generator	Barnstead Thermodyne	582941018789	Jun-95	Jun-95 New	New	DI1	O Prep
Pressurized Fluid Extractor	Dionex	98070129	00-unf	Jun-00 New	New	PFE1	O Prep
Ool Dermostion Chromatograph	.12/Aca i Prep	P26D031	90-unf	Jul-05 New	New	GPC3	O Prep
Gel Permeation Chromatograph	J2/AccuPrep	06D-1196-4.1	70-Juľ	Aug-06 New	New	GPC4	O Prep
Misonex Ultrasonic Disruptor	Sonicator/Heat systems	Unable to view			New	OPH1	O Prep
Misonex Ultrasonic Disruptor	Sonic Dismembrator Fisher Model 550	Unable to view			New	OPH2	O Prep

				12/	12/20/2006
	Sonic Dismembrator Fisher				
Misonex Ultrasonic Disruptor	Model 500	Unable to view	New	OPH3	O Dran
	Sonic Dismembrafor Eisher				3
Misonex Ultrasonic Disruptor	Model 500	Unable to view	in Cla	Y THOU	(
			3		

Mitkem Corporation Equipment List

Department: Inorganics: Metals& Wet Chemistry

Wetchem Location Metals Metals Metals Unit 3 Unit 3 Unit 3 Unit 3 Unit 3 Equipment Centrifuge DryKeeper Optima3 ₽ Optima2 Spec 2 Lachat **TOC1** FIMS $\overline{\Omega}$ Condition New/Used Apr-03 Demo Mar-00 Used Apr-02 Used Nov-03 New Nov-98 New Apr-96 New May-03 New Apr-02 New June-06 June-06 New Service Date in Apr-03 Apr-96 Nov-98 Apr-02 Apr-02 May-03 Nov-03 Mar-00 Received Date 95030498E980802 Serial # 077N3102302 069N8060801 3SGD332010 AB3000-1020 US03035002 7M149 none 1131 Beckman Instruments Manufacturer Tekmar/Dohrmann Lachat Instruments Thermospectronic Sanplatec Corp Perkin Elmer Perkin Elmer Perkin Elmer Dionex Equipment Quick Chem 8000 Optima 4300DV GPR Centrifuge Optima 3100XL Apollo 9000 Genesys 20 Dessicator **FIMS 100** O

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Mitkem Corporation Equipment List

Department: Pest/PCB

Pest/PCB Pest/PCB Pest/PCB Pest/PCB Pest/PCB Pest/PCB Location Equipment ID 罚 贸 罚 型 砬 됴 New/Used Condition Oct-94 New Oct-94 New Service Date in Oct-94 Oct-94 Received Date Serial # US00037060 US00001898 3336A55650 3336A59890 3235A45554 US00032017 Manufacturer Hewlett Packard Hewlett Packard Hewlett Packard Hewlett Packard **Hewlett Packard** Agilent Equipment GC/ECD GC/ECD GC/ECD GC/ECD GC/ECD GC/FID

Corporation	nent List
Mitkem Co	Equipmen

Department: VOA

			Date	Date in	Condition	Equipment	
Equipment	Manufacturer	Serial #	Received	Service	New/Used	₽	Location
GC/MS	Hewlett Packard	3336A55963				۷1	VOA
GC/MS	Hewlett Packard	3336A58222				۷2	VOA
GC	Hewlett Packard	3336A56504				V3	VOA
GC	Hewlett Packard	2843A21041				۷4	VOA
GC/MS	Hewlett Packard	US00007055				75	VOA
GC/MS	Hewlett Packard	US00031343				9/	VOA
Ç	Hewlett Dackard	3140437463				7/	VOA

Mitkem Corporation Equipment List

Department: VOA

VOA LAB VOA LAB VOA LAB **VOA LAB VOA LAB** VOA LAB concentrator |VOA LAB VOA LAB VOA LAB Location concentrator | VOA LAB **VOA LAB** VOA LAB VOA LAB concentrator autosampler Autosampler concentrator concentrator Equipment Headspace TYPE Sample TOWER Sample Sample Sample Sample TRAY TRAY DPM MHC ID Number MITKER A/S-32B A/S-32A A/S-22 A/S-23 A/S-24 A/S-25 A/S-26 A/S-28 A/S-29 A/S-30 **AS-27** A/S-31 A/S-33 instrument current \$ 7 * **OUT OF SERVICE** Serial # M943460129 N111460838 US01170015 H340460074 F445464080 1430460188 3651460769 D730416521 Manufacturer Ol Analytical Ol Analytical Ol Analytical OI Analytical Ol Analytical Tekmar Tekmar DPM-16 Discrete Purging MHC-16 Multiple Heat LSC 2000 ALS 2016 Equipment A/S Model 4551-A A/S Model 4552 A/S Model 4560 A/S Model 4552 A/S Model 4560 A/S Model 4560 A/S Model 4552 A/S Model 4560 A/S Model 4560 A/S Model 7000 Multisampler Controller

12/20/2006

Mitkem Corporation Equipment List

Department: SVOA

Agilent 3435A01848 Oct-94 Oct-94 h	Equipment	Manufacturer	Serial #	Date Received	Date in Service	Condition New/Used	Equipment	location
Agilent 3449A02133 Oct-94 Oct-94 New Agilent US72821130 Nov-99 Nov-99 Used Agilent CN10315002 May-03 Nay-03 New		Agilent	3435A01848	Oct-94	Oct-94	New		SVOA
Agilent US72821130 Nov-99 Nov-99 Used Agilent CN10315002 May-03 May-03 New		Agilent	3449A02133	Oct-94	Oct-94	New	S2	SVOA
Agilent CN10315002 May-03 May-03 New		Agilent	US72821130	Nov-99	Nov-99	Used	S3	SVOA
		Agilent	CN10315002	May-03	May-03	New	S4	SVOA
								-

12/20/2006

SVOA LAB

Location

SVOA LAB

SVOA LAB

SVOA LAB

SVOA LAB

Mitkem Corporation **Equipment List**

Equipment TYPE TOWER TOWER TOWER TOWER TOWER TOWER TOWER TRAY TRAY TRAY TRAY TRAY TRAY TRAY ID Number MITKEM A/S-10 **AVS-11** A/S-12 A/S-13 A/S-14 A/S-5 A/S-6 A/S-2 A/S-3 A/S-8 A/S-9 AS-1 AIS-4 A/S-7 Instrument current Department: GC AND GC/MS US514307466 18596C CN143220863 G1513A CN15121474 G1513A CN31623836 G2614A CN31630412 GZ613A US12111699 G2614A US94706562 G2614A US11618592 G2613A US12109082 G2613A US14207448 18596C CN13920644 G1513A US14307475 18596C CN13720586 G1513A 3522A38799 18596M Serial # HEWLETT-PACKARD Manufacturer Equipment AUTOSAMPLER AUTOSAMPLER

SVOA LAB

_							
12/20/2006	SVOA LAB	SVOA LAB	SVOA LAB				
	TRAY	TOWER	TRAY	TOWER	TRAY	TOWER 1	TOWER 2
	A/S-15	A/S-16	A/S-17	A/S-18	A/S-19	A/S-20	A/S-21
	US14207449 18596C	US00001909 G1513A	US92505547 G2614A	US94710320 G2613A	3216A28361 18596B	COULDN'T SEE	COULDN'T SEE
	HEWLETT-PACKARD	HEWLETT-PACKARD	HEWLETT-PACKARD	HEWLETT-PACKARD	HEWLETT-PACKARD	HEWLETT-PACKARD	HEWLETT-PACKARD
	AUTOSAMPLER	AUTOSAMPLER	AUTOSAMPLER	AUTOSAMPLER	AUTOSAMPLER	AUTOSAMPLER	AUTOSAMPLER

Laboratory Information System Equipment

1. Data Collection:

- 1.1. 12 HP chem station software for collecting GC-ECD and GC-MS data
 - 1.1.1. 5 GC-ECD
 - 1.1.2. 4 GC-MS (SVOA)
 - 1.1.3. 4 GC-MS (VOA)
- 1.2. Hardware varies but is x86 compatible
- 1.3. OS is Windows, Various Versions (9x, NT, 2000)

2. Data Storage:

- 2.1. Dell Poweredge servers
 - 2.1.1. Dual P IV Xeon processors
 - 2.1.2. 2 GB RAM
 - 2.1.3. 105 GB Storage expandable to 750 GB internally
 - 2.1.4. OS is Windows, Various Versions (NT and 2003)
- 2.2. LTO tape drive daily backup, long term archiving and data restoration
- 2.3. Tape software is Backup Exec (10.x)

3. Compound Identification:

- 3.1. 12 Target 4.14 chromatographic software
- 3.2. Hardware is Intel based (3GHZ, 512MB RAM) for Target 4.14
- 3.3. OS is Windows Xp

4. Forms Generation:

- 4.1. In house forms generation LIMS modules for SW-846, ILM4 and ILM5 metals
- 4.2. In house forms generation LIMS modules for SW-846, OLC03 and SOM01 organics
- 4.3. Target-based forms generation for OLM04 and SW-846 organics
- 4.4. Hardware varies but is x86 compatible
- 4.5. OS is Windows, Various Versions (2000 and Xp)

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MITKEM CORPORATION CONFIDENTIALITY, ETHICS, and DATA INTEGRITY AGREEMENT APPENDIX B

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CONFIDENTIALITY, ETHICS, AND DATA INTEGRITY

The confidentiality, ethics, and data integrity agreement attached must be signed and dated by all new personnel associated with the data generated by Mitkem Corporation. All said personnel will complete a training course and understand the information stated in the agreement. The course must include the ethical and legal responsibilities including the potential punishments and penalties for improper, unethical, or illegal actions. All personnel must fully understand this information before signing the agreement.

Data Integrity training will be done on an annual basis. If changes to the enclosed integrity agreement are made, then all employees will be required to review and sign. All documents are stored in the employee's personnel file located in the QA Department.

MITKEM CORPORATION

CONFIDENTIALITY, ETHICS AND DATA INTEGRITY AGREEMENT

I.		(Name), state that I understand the standards of grity required of me with regard to the duties I perform and the data I report in action with my employment at Mitkem Corporation.
п.	I agr	ree that in the performance of my duties at Mitkem Corporation:
	A.	I shall not improperly use manual integrations to meet calibration or method QC criteria, such as peak shaving or peak enhancement.
	В.	I shall not intentionally misrepresent the date or time of analysis by resetting computer or instrument date/time.
	C.	I shall not falsify analytical results.
	D.	I shall not report analytical results without proper analysis documentation to support the results; dry-labbing.
	Е.	I shall not selectively exclude data to meet QC criteria, such as calibration points without technical or statistical justification.
	F.	I shall not misrepresent laboratory performance by presenting calibration data or QC limits within data reports that are not linked to the data set reported.
	G.	I shall not represent matrix interference as basis for exceeding acceptance criteria in interference-free matrices, such as method blanks and Laboratory Control Standards (LCS).
	Н.	I shall not manipulate computer software for improper background subtraction or chromatographic baseline manipulations.
	I.	I shall not alter analytical conditions such as EM voltage, GC temperature program, etc. from standards analysis to sample analysis.
	J.	I shall not misrepresent QC samples such as adding surrogates after sample extraction, omitting sample preparation steps, or over-spiking/under-spiking.

I shall not report analytical results from the analysis of one sample for those of

I shall not intentionally represent another individual's work as my own.

K.

L.

another.

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- III. I agree to report immediately any accidental or intentional reporting of non-authentic data either I or another employee may have committed. Such report must be made to any member of Mitkem Corporation's Management (Kin Chiu, Reinier Courant, Edward Lawler, Yihai Ding) or the Quality Assurance Director, either orally or in writing. Every incident will be investigated by senior management. A written corrective action is required of any findings from the investigation.
- IV. Any incidents that violate the standards of data integrity can result in immediate termination of the employee as well as civil or criminal charges.
- V. Questions pertaining to confidentiality, ethics, and integrity may be posed to any of the above individuals.
- VI. I agree not to divulge any pertinent information including but not limited to data and any other information about a project to outside sources without the prior consent from the client.

I understand that failure to comply with the above ethics and data integrity agreement can result in my immediate dismissal from Mitkem Corporation.

(Signature)	(Date)

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MITKEM CORPORATION

SUBCONTRACTORS

CONFIDENTIALITY, ETHICS AND DATA INTEGRITY AGREEMENT

I.	I,	(Name), authorized representative of
	perfe	(Subcontractor) state that I understand the dards of integrity required of me and the Subcontractor with regard to the duties ormed and the data reported in connection with the analysis/analyses contracted by tem Corporation.
П.	Subo	contractor agrees that in the performance of analysis for Mitkem Corporation:
	A.	Subcontractor shall not intentionally report data values or results that are not the actual values measured or observed;
	В.	Subcontractor shall not modify data values unless the modification can be technically justified through a measurable analytical process;
	C.	Subcontractor shall not intentionally report the dates and times of data analyses that are not the true and actual dates and times of analyses; and
	D.	Subcontractor shall not intentionally represent another's work as its own.
III.		contractor agrees to report immediately any accidental or intentional reporting of authentic data to Mitkem.
IV.	data	contractor agrees not to divulge any pertinent information including but not limited to and information about any Mitkem projects to outside sources without the prior ent from Mitkem or its clients.
I unde	erstand mediat	I that failure to comply with the above ethics and data integrity agreement can result e termination of the subcontract relationship with Mitkem Corporation.
(Signa	ture)	(Date)
(Name)	
(Title)		

ATTACHMENT 2

Table 1

(Analytical Laboratory Testing Program)

Table 1

Analytical Laboratory Testing Program

Remedial Work Plan 100 Fernwood Avenue Rochester, New York (NYSDEC Site ID C828119)

Task	Sample Matrix	Parameter	Field Samples	Trip Blanks	MS/MSD	Field Blanks	Analytical Methods	Reporting Levels	Corresponding SCGs
	Soil	TCL VOCs	4	0	1	1	ASP Method OLM04.3	ASP-B	Part 375 Soil Cleanup Objectives
	Soil	TCL SVOCs	4	0	1	1	ASP Method OLM04.3	ASP-B	Part 375 Soil Cleanup Objectives
,	Soil	трн	4	0	1	1	NYSDOH Method 310.13	MDL	Not Applicable
Confirmatory Samples from Existing In-Situ Bioremediation System	Soil	Plate Count	4	0	0	0	USEPA Method 9215C	MDL	Not Applicable
	Water	TCL VOCs	3	1	1	1	ASP Method OLM04.3	ASP-B	TOGS 1.1.1 Groundwater Standards and Guidance Values
	Water	TCL SVOCs	3	0	1	1	ASP Method OLM04.3	ASP-B	TOGS 1.1.1 Groundwater Standards and Guidance Values
	Water	Plate Count	3	0	0	0	USEPA Method 9215C	MDL	Not Applicable
	Water	TCL VOCs	up to 56 (up to 7 rounds, up to 8 samples/round)	П	-	-	ASP Method OLM04.3	ASP-B	TOGS 1.1.1 Groundwater Standards and Guidance Values
	Water	TCL SVOCs	up to 56 (up to 7 rounds, up to 8 samples/round)	0	1	1	ASP Method OLM04.3	ASP-B	TOGS 1.1.1 Groundwater Standards and Guidance Values
	Water	Nitrate	up to 56 (up to 7 rounds, up to 8 samples/round)	0	1	1	USEPA Method E300IC	MDL	Not Applicable
Monitored Natural Attenuation	Water	Iron (II)	up to 56 (up to 7 rounds, up to 8 samples/round)	0	1	-	USEPA Method SM3500D	MDL	Not Applicable
Groundwater Samples	Water	Manganese	up to 56 (up to 7 rounds, up to 8 samples/round)	0	1	-	ASP Method ILM04.1	ASP-B	Not Applicable
	Water	Sulfate	up to 56 (up to 7 rounds, up to 8 samples/round)	0	1	1	USEPA Method E300IC	MDL	Not Applicable
· · · · · · · · · · · · · · · · · · ·	Water	Methane	up to 56 (up to 7 rounds, up to 8 samples/round)	0	-	,	USEPA Method RSK175	MDL	Not Applicable
	Water	Chloride	up to 56 (up to 7 rounds, up to 8 samples/round)	0	1	_	USEPA Method E300IC	MDL	Not Applicable

VOC = Volatile Organic Compound SVOC = Semi-Volatile Organic Compound TCL = Target Compound List TPH = Total Petroleum Hydrocarbon

ASP = Analytical Services Protocol NYSDOH = New York State Department of Health MDL = Method Detection Limit USEPA = United States Environmental Protection Agency

ATTACHMENT 3

Resume of Ms. Hope Kilmer

Day Environmental, Inc. JD5970 / 4014R-07

EXPERIENCE

Day Environmental, Inc.: March 2006 to present

Years with Other Firms: Over 14 years

AREAS OF SPECIALIZATION

- Environmental Compliance
- Quality Assurance Officer and DUSR reporting
- Industrial Hygiene Sampling & Analysis Techniques
- Inorganic and Organic Methods & Analysis
- Radiation Safety & Analysis

EDUCATION

State University of New York at Fredonia; B.S. Chemistry 1989 Additional Chemistry and Industrial Hygiene curricula graduate course work

REGISTRATIONS/AFFILIATIONS

- Certified Hazardous Materials Manager (CHMM), ID# 14070
- 24 hour HAZWOPER Emergency Response Training
- 8 Hour OSHA Hazardous Waste Site Worker Refresher Training

RESPONSIBILITIES AND PROJECT EXPERIENCE

Ms. Kilmer has more than 15 years of experience providing sampling information, calibrated equipment, and report data. Ms Kilmer's experience includes working within environmental laboratories performing multiple analysis techniques on various media including: personnel samples, soil, sludge, air, and water; addressing environmental, health, and safety issues within a manufacturing facility, waste characterization, waste management, annual OSHA, RCRA, and Radiation Safety training.

Regulatory Compliance:

Air Permit Data Management and Compliance Reporting, Industrial Facility, Albion, New York: Maintain Access database containing air permit information including materials used and their VOC and HAP emissions, receive monthly material usage reports from the facility and prepare monthly emissions report as per Title V requirement. Identified opportunities for improved data collection, management of database functions, and evaluation of status of compliance against permit conditions. Submitted semi-annual and annual Title V compliance monitoring reports on timely basis.

Clean Water and Oil Pollution Prevention Regulatory Compliance, Industrial Facilities, New York: Performed storm water permitting assessment. Assisted in the preparation of Storm Water Pollution Prevention Plans (SWPPP) and Spill Control and Countermeasures (SPCC) Plans for facilities.

Investigation of ambient air quality, Manufacturing Facility, Rochester, New York: Performed health and safety monitoring including volatile organic compound sampling and particulate monitoring using various sampler types. Evaluated data, prepared and provided a report.

Investigation of ambient air quality, Manufacturing Facility, Arcade, New York: Conducted noise exposure monitoring and an indoor air quality survey in a manufacturing facility. Five individuals were monitored to determine noise exposure and air samples for three different materials were collected at four locations in the building. Evaluated data, prepared, and provided a report.

Polychlorinated Biphenyl (PCB) Annual Log, Metro North Railroad Yards, New York and Connecticut: Prepared the PCB Annual Log for multiple facilities.

RCRA Hazardous Waste Compliance, Industrial Facility, Rochester, New York: Project activities included waste characterization and disposal, preparation of hazardous waste profiles, manifests, the Special Assessment forms for NYS Tax Department, and the Hazardous Waste Report.

(continued)

RCRA Hazardous Waste Compliance, Multiple Industrial Facilities, Rochester, New York: Preparation of Hazardous Waste Reports.

RCRA Hazardous Waste Compliance, Manufacturing Facility, Rochester, New York: Performed RCRA 40CFR part 265 subpart BB/CC monitoring for a large manufacturing facility.

SARA/EPCRA Regulatory Compliance, Multiple Industrial Facilities, New York: Tasks included preparation of, Toxic Release Inventory and Tier II reports for several facilities.

Site Assessments/Investigations, Rochester, New York: Conducted and prepared associated reports for Phase I site assessment.

Environmental Remediation Activities - Former Manufacturing Facilities, Rochester, New York: Current activities include the evaluation of laboratory data and the preparation of Data Usability Summary Report (DUSR) documentation for submittal to the New York State Department of Environmental Conservation (NYSDEC).

Chemical Technician, Eastman Kodak Company, New York:

Worked within four separate laboratories, Industrial Hygiene Analysis, Inorganic Analysis, Metals Analysis, and Environmental Process Monitoring. Consulted with internal clients to determine needs and provide necessary sampling equipment and media. Assisted in field sampling activities for worker and environment exposure projects. Performed instrument maintenance and calibration.

Lab Analysis: Performed analysis of samples utilizing OSHA, NIOSH, ELAP, and ASTM methods. Samples included Industrial Hygiene personnel dosimeters, silica gel tubes; groundwater, soils, sludge, filters, aqueous solutions, and unknown solid materials. Develop and document methods of analysis for multiple laboratory techniques including Gas, Ion, and HP-Liquid Chromatography techniques; alpha/beta analysis, Segmented flow analysis, Total Organic Carbon, Inorganic Carbon, ICP-Atomic Absorption, FIAS-MHS (Flow Injection Atomic Spectroscopy- Mercury Hydride System) and AA. Developed digestion methods for various materials (waters, solids, sludges, gelatin, bone).

Data Analysis: Designed and wrote reports for various types of sampling, reviewed reports of others for accuracy and data evaluation and validation. Performed analyses using ELAP protocols; stringent quality control programs were followed as determined by state and federal agencies; participated in ELAP proficiency testing.

Project examples:

Cyanide in Air: Determined a method of sampling for cyanide compounds possibly being generated over a development process. The process consisted of several tanks of solutions over which a conveyor system for film operated. The sampling chain was made up of bubblers containing 0.025 M sodium hydroxide solution and calibrated pumps. The air was sampled for 15 and 30 minutes while the process was in operation. The samples were collected into sealed glass vials, analyzed, and results reported to the IH.

Formaldehyde in air: The concern was that formaldehyde was in use in a new manufacturing process. The monitoring was to determine if formaldehyde was being exhausted through a building ventilation system on the roof. Sep-Paks and calibrated pumps were set up at the stacks exits. The exhausts temperatures and velocities were measured and formaldehyde sampled for 5, 15, and 30 minute intervals. Samples were sent to an outside lab for analysis. Upon receipt, the results were checked for data validation and a report generated for the IH.

Methylene Chloride Exposure: An area consisted of several large open vats of methylene chloride and the concern was regarding personnel in the area being exposed to large quantities of the chemical in air. The people were monitored using passive charcoal badges to collect the chemical. The badges were collected after 30 minute and 4 hour intervals and sealed for analysis. The analysis was performed in-house and a report submitted to the IH.

APPENDIX D

Proposed Brownfield Cleanup Program Sign And NYSDEC Instructions



Brownfield Cleanup Program

Former Vogt Manufacturing Site C828119 Conifer Development, Inc.

Eliot Spitzer, Governor Peter Grannis, Commissioner Robert J. Duffy, Mayor Build for the Future Transform the Past....

SITE SIGNS FOR REMEDIAL PROGRAMS

Instructions

Signs are required at sites where remedial actions are being performed under one of the following remedial programs: State Superfund, Voluntary Cleanup Program (VCP), Brownfield Cleanup Program (BCP), and Environmental Restoration Program (ERP). They will not be required during the investigation and design phases. The cost of the sign will be borne by the parties performing the remedial action based on the legal document the activities are being performed under (i.e. volunteers/participants would pay 100% of the cost under the BCP; municipalities would pay 100% and then would be reimbursed for the cost under the ERP).

Sign Requirements

Size:

Horizontal format - 96" wide by 48" high

Construction Materials: Aluminum or wood blank sign boards with vinyl sheeting.

Inserts:

"Site Name", "Site Number", "Name of Party Performing Remedial

Activities" and "Municipal Executive".

Indicate position, size and topography for specific inserts.

Color Scheme:

Copy surrounding DEC logo - "NEW YORK STATE DEPARTMENT

OF ENVIRONMENTAL CONSERVATION" - PMS 355

DEC logo:

PMS 301 Blue

PMS 355 Green

Text:

Program (choose one):

PMS 301

Brownfield Cleanup Program Voluntary Cleanup Program State Superfund Program

1996 Clean Water/Clean Air Bond Act - Environmental Restoration Program

Site Name, Site Number, Party Performing Remedial Activities
Names of Governor, Commissioner, Municipal Executive
PMS 355
Transform the Past......Build for the Future
PMS 355

Type Specifications:

All type is Caslon 540, with the exception of the logotype. Format is: center each line of copy with small caps and

initial caps.

Production Notes:

96" wide x 48" high aluminum blanks will be covered with vinyl sheeting to

achieve background color. Copy and logo will be silk screened on this

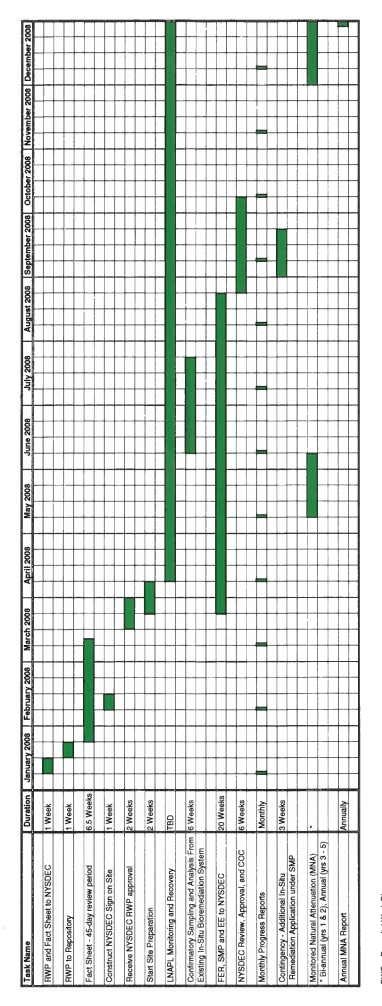
surface.

See attached format

APPENDIX E

Project Schedule

Tentative Remediation Schedule 100 Fernwood Avenue Rochester, New York (NYSDEC Site ID C828119) (First Year)



RWP = Remedial Work Plan
NYSDEC = New York State Department of Environmental Conservation
LNAPL = Light Non-Aqueous Phase Liquid
TBD = To Be Determined
TER = Final Engineering Report
SMP = Site Management Plan
EE = Environmental Easement