FORMER UNION WIRE DIE CORP.

39-40 30th STREET QUEENS, NEW YORK Block 399 Lot 34

REMEDIAL INVESTIGATION WORK PLAN

September 2014

Prepared for: Ganesh Management, LLC 39-40 30th Street, Queens, NY

Prepared By:



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Community Air Monitoring Plan

CERTIFICATION

I, Charles B. Sosik, certify that I am currently a Qualified Environmental Professional as defined in 6 NYCRR Part 375 and that this Remedial Investigation Work Plan was prepared in accordance with all applicable statutes and regulations and in substantial conformance with the DER Technical Guidance for Site Investigation and Remediation (DER-10).

Charle Sosik

1.0 INTRODUCTION

This Remedial Investigation Work Plan (RIWP) was prepared on behalf of Ganesh Management LLC for the property located at 39-40 30th Street, Long Island City, Queens, New York. An application for acceptance into the New York State Brownfield Cleanup Program (BCP) is being submitted with this RIWP.

The Site was undeveloped from 1887 to sometime between 1915 and 1936. By 1936, a gas station with two gasoline tanks occupied the site. The property was redeveloped by 1947 into a 2-story warehouse utilized by Optical Products Corporation for manufacturing, shipping, and as an office. The building has remained since, with several other occupants including Union Wire Die Corp. (1960s-1983), National Tea Packaging Co. Inc. (1962-1991), and a warehouse (1991-2006). A Remedial Investigation performed at the Site, identified chlorinated volatile organic compounds in soil, groundwater and soil vapor. As a result of these findings, which were preformed under the New York City E-Designation Program, the New York City Office of Environmental Remediation (OER) referred the project to the New York State Department of Environmental Conservation.

The purpose of this Remedial Investigation Work Plan is to collect data of sufficient quality and quantity to characterize the nature and extent of chlorinated solvent contamination in on-site soil, groundwater and soil gas, to complete a qualitative exposure assessment for future occupants of the proposed building and the surrounding community and to evaluate alternatives to remediate the contamination.

The overall objectives of the project are to prepare the site for residential use and to remediate known and unknown environmental conditions at the site to the satisfaction of the DEC and the New York State Department of Health (NYSDOH).

1.1 Site Location and Description

The street address for the subject site is 39-40 30th Street, Long Island City, NY 11101 (**Figure 1**). The subject property is located in the City of New York and Borough of Queens (Queens County). The subject site is listed as Block 399, Lot 34 by the City of New York. The lot is located on the northwest corner of the intersection between 40th Avenue and 30th Street. The lot consists of 133 feet of street frontage on 30th Street 100 feet of Street frontage on 40th Avenue for a total of 14,000 square feet (0.32 acres). The lot is currently developed with a two-story commercial warehouse which occupies approximately 70% of the lot (**Figure 2**).

According to the deed and the Automated City Registration Information System (ACRIS) website, the current owner of the lot is Ganesh Management LLC.

The area surrounding the property is highly urbanized and predominantly consists of commercial, industrial and residential buildings with mixed-use buildings (residential w/ first floor retail) along main corridors / thoroughfares.

The elevation of the Site is approximately 28 feet above the National Geodetic Vertical Datum (NGVD). The area topography gradually slopes to the southeast. The depth to groundwater

beneath the Site, as determined from field measurements, is approximately 20 feet below grade. Based on regional groundwater elevation maps, groundwater flows to the west toward the East River.

1.2 Redevelopment Plans

The development project includes the demolition of the existing building and the construction of a new 7-story residential apartment building with a full cellar. The first floor will consist of parking and a lobby with residential units on the upper floors. The building will cover approximately 80% of the lot, leaving a rear courtyard behind the building. The basement level and foundation will require excavation of majority of the site to a total depth of approximately 12 feet below grade.

1.3 Site History

The environmental history of the Site was previously reported by Galli Engineering in a 2013 Phase I Site Assessment.

A review of Sanborn maps showed that the property was occupied by a gas station in 1936. The property was redeveloped by 1947 into a 2-story warehouse utilized by Optical Products Corporation for manufacturing, shipping, and as an office. The building has remained since, with several other occupants including Union Wire Die Corp. (1960s), National Tea Packaging Co. Inc. (1962-1991), and a warehouse (1991-2006).

1.4 Summary of Previous Investigations

Environmental investigations performed at the Site include the following:

- Phase I Environmental Site Assessment Report Galli Engineering (September 2013)
- NYCOER Remedial Investigation Preliminary Data Summary EBC (January 2014)

A summary of the investigations performed is provided in the following sections.

1.4.1 September 2013 – Phase I Environmental Site Assessment Report (Galli)

A Phase I was competed by Galli Engineering, P.C. (Galli) in September, 2013. A history dating back to 1887 was established. According to a review of Sanborn maps, as well as personal interviews, the Site was undeveloped from 1887 to sometime between 1915 and 1936. By 1936, a gas station with two gasoline tanks occupied the site. The property was redeveloped by 1947 into a 2-story warehouse utilized by Optical Products Corporation for manufacturing, shipping, and as an office. The building has remained since, with several other occupants including Union Wire Die Corp (1960s), National Tea Packaging Co. Inc. (1962-1991), and a warehouse (1991-2006).

Based upon reconnaissance of the subject site and surrounding properties, interviews and review of historical records and regulatory agency databases, Galli identified the following environmental concerns:

- The subject property currently contains one warehouse used as an office space and distribution center for electronic goods.
- The property is an E Designation site for hazardous materials, requiring Phase I and Phase II Testing Protocol.
- The subject property was previously used as a gas station and automotive repair shop; an optical product manufacturing facility; a tea packaging facility; and a jewelry manufacturing facility.
- Expected chemicals previously used on the subject property include: petroleum, motor oil, nitrogen and sulfur oxides, TCA, arsenic, ammonia, ammonium chloride, sulfates, cyanide stripping solutions, silica and metallic fine posers, acid and alkaline cleaning solutions, CFCs, HCFCs, cadmium, and chlorinated solvents.
- The property has an active violation in place from the Department of Buildings pertaining to an elevator (Violation Number 9027/416426).
- The property had a boiler removed in 2003. The property is currently serviced with natural gas.
- Asbestos containing materials (ACMs) in roofing and floor tiles, and lead based paint may exist on the subject property.
- No mold or water damage was observed during the site inspection.
- No storage of hazardous materials, distressed vegetation or other adverse environmental conditions were observed on the subject property at the time of inspection.

1.4.2 January 2014 - Preliminary RI Data Summary (EBC)

The remedial investigation was performed from December 9 - 20, 2013 in accordance with the Remedial Action Work Plan approved by the NYCOER as part of the E-designation review process. The investigation included the installation of 8 soil borings to collect 16 soil samples, the installation of five temporary monitoring wells to collect groundwater samples and the installation of five soil gas implants to collect soil gas samples.

The results of the RI identified elevated levels of trichloroethene (TCE) in shallow soil (above the water table), in three of the boring locations at concentrations above unrestricted soil cleanup objectives (SCOs).

Groundwater was encountered at a depth of approximately 20 feet below grade. Chlorinated VOCs including terachloroethene (PCE) and TCE were detected within all five groundwater samples above NYSDEC groundwater standards. Cis-1,2-dichloroethene (DCE) was reported above groundwater standards in one sample. In addition petroleum (gasoline) related VOCs including 1,2,4-trimethylbenzene, 1,3,5-trimethylbenzene, ethylbenzene, isopropylbenzene and mixed xylenes were also reported above standards in one sample.

Both PCE and TCE were reported in soil gas above mitigation levels established within the State DOH soil vapor guidance matrix. TCE concentrations in soil gas ranged from 232 μ g/m3 to a high of 9,400 μ g/m3. PCE concentrations ranged from 3,520 μ g/m3 to 9,760 μ g/m3. TCE was reported at elevated concentrations in all 5 samples obtained. Elevated PCE concentrations were reported in 3 of the 5.

SVOCs including benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, chrysene, dibenzo(a,h)anthracene and ideno(1,2,3-cd)pyrene, were reported above restricted soil cleanup objectives (SCOs) in shallow soil (0-2ft, or 4-6ft) at three of the boring locations.

Metals including arsenic, lead and mercury were reported above restricted residential SCOs in shallow soil in one boring location. Lead, chromium, copper, mercury and Zinc were reported above unrestricted SCOs in many of the boring locations. Elevated SVOCs and metals are commonly associated with historic fill and are not believed to be related to historic use of the property.

The historical use of the site as a wire die manufacturer combined with shallow the site-wide TCE detections in soil, groundwater and soil gas is evidence that it is related to an on-site release and historic use. TCE is known to have been commonly and extensively used to clean and degrease metals and would be expected to be used in a die manufacturing operation.

Since PCE was not reported in soil above unrestricted SCOS, it is likely that the PCE reported in groundwater and soil gas is related to the adjacent property to the north, the Former Bridge Cleaners Site which is a NYS Brownfield Cleanup Program Site. This property has PCE contamination with a known off-site PCE plume.

1.5 Site Geology / Hydrogeology

Subsurface soils at the Site consists of a mixture of a silty non-native fill, to a depth which ranges from 0 to 6 feet below grade. The fill is followed by a silty-sand. A coarse sand was noted at two of the boring locations in place of the silty-sand. Groundwater is present under water table conditions at a depth of approximately 20 feet below the surface and is expected to flow west-southwest (**Figure 3**).

According to the USGS topographic map for the area (Brooklyn Quadrangle), the elevation of the Site 28 feet above the National Geodetic Vertical Datum (NGVD). The area topography gradually slopes to the southeast.

1.6 Site Conceptual Model

CVOC contamination at the Site consists mainly of trichloroethene (TCE) in shallow soil to a depth of at least 6 feet but less then 10 feet below grade along the south and east portions of the building. Groundwater is present at a depth of approximately 20 feet below grade. Chlorinated VOCs including tetrachloroethene (PCE) and TCE were detected throughout the Site above NYSDEC groundwater standards.

Both PCE and TCE were reported in soil gas above mitigation levels established within the State DOH soil vapor guidance matrix.

The historical use of the subject Site as a wire die manufacturer combined with the shallow TCE detections in soil is evidence that this contamination is related to an on-site release and historic use. TCE is known to have been commonly and extensively used to clean and degrease metals and would be expected to be used in a die manufacturing operation.

PCE in groundwater was reported in all of the locations at 3 to 18 times greater than the TCE concentrations suggesting that the TCE is related to dechlorinization of PCE and not a TCE release. Additionally the highest concentrations of PCE and TCE were reported in an upgradient location which is adjacent to and downgradient of the property to the north. This property is known as the Former Bridge Cleaners Site which is a NYS Brownfield Cleanup Program Site. This property has PCE contamination with a known off-site PCE plume.

The release scenario for TCE at the Site is unknown but could have occurred at any time during the occupancy of the wire die company from the 1960's through 1983. Releases would be expected to be from incidental spillage related to milling operations and / or storage of TCE solvent. Under this scenario TCE may have entered the subsurface through cracks or other voids in the concrete floor. Based on the absence of TCE in soil in the 10-12 foot interval and the relatively low concentrations reported in groundwater, 20 feet below grade, it is unlikely that TCE migrated to the groundwater as a solvent. The TCE contamination in groundwater beneath the Site is likely related to the degradation of Bridge Cleaners PCE plume which is migrating beneath the Site.

The elevated TCE levels reported in soil gas are associated with off-gassing from the TCE impacted soil. It would not be expected to be related to off-gassing from the TCE impacted groundwater since the TCE concentrations in groundwater are relatively low. The elevated PCE levels in soil gas are either related to off-gassing from the PCE plume beneath the site or from the migration of vapors from PCE impacted soil on the adjacent Bridge Cleaners property.

This conceptual model will be refined and updated based upon finding of the Remedial Investigation.

RIDGE, NY 11961

2.0 SAMPLING AND ANALYIS PLAN

The purpose of this work plan will be to determine the nature and extent of the on-site contamination and identify all sources of contamination (horizontal/vertical) that may be present at the Site. The investigation must produce data of sufficient quality and quantity to allow NYSDEC and NYSDOH to complete a Significant Threat Determination as per Part 375-2.7 and enable the performance of a qualitative human health exposure assessment as per DER-10, 3.3(c) 4.

The investigation will consist of the following elements:

- Installation of 6 soil borings across the site to delineate the extent of soil impact and to obtain additional information on soil quality with respect to Soil Cleanup Objective (SCOs);
- Installation of 3 additional monitoring wells and collection of groundwater samples from the 3 new and 5 existing monitoring wells to assess CVOC groundwater impacts; and
- Installation of 4 soil gas implants with 2 implants located on the ground level and 2 implants in the cellar to assess vapor phase VOCs.

2.1 **Soil Sampling**

Soil Borings 2.1.1

Six soil borings (B9 through B14) will be advanced across the Site. At all boring locations soil samples will be collected continuously in 5-foot intervals using a GeoprobeTM dual-tube sampling system. The GeoprobeTM uses a direct push hydraulic percussion system to drive and retrieve core samplers. A track-mounted Geoprobe™ model 6620DT will be utilized.

At each boring location sampling will continue to a minimum depth of 15 feet below grade. Collected soil samples will be characterized by an experienced environmental professional (EP) and field screened for the presence of volatile organic compounds (VOCs) using a photoionization detector (PID). The EP will record all observations in a bound project dedicated field book which will be used to prepare a boring log for each soil boring location. Recorded observations will include sample depth, sample recovery, soil type evidence of water (if encountered), PID reading and physical evidence of contamination (odor, staining, sheen, etc.).

A minimum of two samples will be retained from each boring (B9-B14). Retained samples from delineation borings will be collected from the 0 to 2 foot and 13 to 15 foot intervals. The 0-2 foot interval will be analyzed for VOCs and the 13-15 foot interval will be VOCs, SOVCs, metals, pesticides and PCBs. If contaminated soils are encountered additional samples will be retained from the interval and section of the core with the highest PID reading if it is not the same interval as one of the two planned samples. If contaminated soils are encountered in the cellar borings then an additional sample will be retained from the interval and section of the core with the highest PID reading if it is not the same interval as one of the two planned intervals.

A sample matrix showing the number, type and analysis of samples collected during the Remedial Investigation is provided as **Table 1**. The proposed location of the soil borings is shown on **Figure 4**.

2.2 **Groundwater Sampling**

Two monitoring wells, MW6-MW8, be installed to a depth of approximately 6 feet below the water table using a track-mounted GeoprobeTM model 6620DT.

The monitoring well will be constructed of 2-inch diameter pvc casing and 0.010 inch slotted pvc well screen. The well will have 10 feet of screen from 20 to 26 feet below grade. A No. 00 Morie or equivalent filer sand will be placed in the borehole to within 2 feet above the top of the screen. A 1-foot hydrated bentonite seal will be placed on top of the filter sand and the remainder of the borehole will be backfilled to grade.

Groundwater samples will be collected from existing monitoring wells MW1 through MW5 as well as newly installed wells MW6 and MW7. Prior to sample collection well development (purging water from well to remove fine particles and or sediment) will be preformed for MW6 and MW7. Sampling will be performed several days after well development. Sample procurement will be achieved through the use of dedicated polyethylene tubing, a stainless steel check valve and a peristalic pump. Proposed and existing monitoring well locations are shown on **Figure 5**.

All groundwater sampling activities will be recorded in the project dedicated field book. This will include a description of:

- Date and time of sample collection
- Sample location
- Purging time, duration and volume;
- Sample appearance
- Analytical methodology:

Groundwater samples will be collected using a check valve, peristaltic pump and dedicated polyethylene tubing in accordance with standard low-flow sampling procedures as follows:

- Record pump make & model on sampling form.
- Wear appropriate health and safety equipment as outlined in the Health and Safety Plan
- Inspect each well for any damage or evidence of tampering and note condition in field logbook.
- Remove the well cap.
- Lay out plastic sheeting and place the monitoring, purging and sampling equipment on the
- To avoid cross-contamination, do not let any downhole equipment touch the ground.
- Measure well headspace with a PID or FID and record the reading in the field logbook.
- A synoptic water level measurement round should be performed (in the shortest possible time) before any purging and sampling activities begin. Measure and record the depth to

water using a water level meter or interface probe to the nearest 0.01 ft. Record the measurement in the field logbook. Do not measure the depth to the bottom of the well at this time (to avoid disturbing any sediment that may have accumulated). Obtain depth to bottom information from installation information in the field logbook or soil boring logs.

- Collect samples in order from wells with lowest contaminant concentration to highest concentration.
- Connect the polyethylene tubing to the peristaltic pump and lower the tubing into the well to approximately the middle of the screen. Tubing should be a minimum of 2 feet above the bottom of the well as this may cause mobilization of any sediment present in the bottom of the well.
- Start the pump at its lowest speed setting and slowly increase the speed until discharge occurs. Check water level. Adjust pump speed until there is little or no water level drawdown (less than 0.3 feet). If the minimal drawdown that can be achieved exceeds 0.3 feet but remains stable, continue purging until indicator field parameters stabilize.
- There should be at least 1 foot of water over the end of the tubing so there is no risk of entrapment of air in the sample. Pumping rates should be reduced to the minimum capabilities of the pump, if needed, to avoid purging the well dry. However, if the recharge rate of the well is very low and the well is purged dry, then wait until the well has recharged to a sufficient level and collect the appropriate volume of sample.
- During well purging, monitor indicator field parameters (temperature, specific conductance and pH) every three to five minutes (or less frequently, if appropriate). Note: during the early phase of purging emphasis should be put on minimizing and stabilizing pumping stress, and recording those adjustments. Purging is considered complete and sampling may begin when all the above indicator field parameters have stabilized. Stabilization is considered to be achieved when three consecutive readings, taken at three (3) to five (5) minute intervals, are within the following limits:
 - specific conductance (3%),
 - temperature (3%), 0
 - pH (\pm 0.1 unit)
- VOC samples should be collected directly into pre-preserved sample containers. Fill all sample containers by allowing the pump discharge to flow gently down the inside of the container with minimal turbulence. Fill each container with sample to just overflowing so that no air bubbles are entrapped inside. Cap each bottle as it is filled.
- Label the samples, and record them on the chain of custody form. Place immediately into a cooler for shipment and maintain at 4°C.
- Remove the tubing from the well. The polyethylene tubing must either be dedicated to each well or discarded. If dedicated the tubing should be placed in a large plastic garbage bag, sealed, and labeled with the appropriate well identification number.
- Close and lock the well.
- Decontaminate pump either by changing the surgical pump tubing between wells or as follows:
 - 1. Flush the equipment/pump with potable water.
 - 2. Flush with non-phosphate detergent solution. If the solution is recycled, the solution must be changed periodically.
 - 3. Flush with potable or distilled/deionized water to remove all of the detergent solution. If the water is recycled, the water must be changed periodically.

- 4. Flush with isopropyl alcohol (pesticide grade). If equipment blank data from the previous sampling event show that the level of contaminants is insignificant, then this step may be skipped.
- 5. Flush with distilled/deionized water. The final water rinse must not be recycled.

Samples will be collected in pre-cleaned laboratory supplied glassware, stored in a cooler with ice and submitted to a New York State ELAP certified environmental laboratory. Groundwater samples from monitoring well locations will be submitted for analysis of VOCsby EPA Method 8260.

All monitoring wells will be surveyed to determine relative casing elevation to the nearest 0.01 ft and horizontal position to the nearest 0.1ft. Survey data will be used to determine the direction and gradient of groundwater flow at the Site.

2.3 **Soil Vapor Sampling**

Soil vapor samples will be collected in accordance with the Final Guidance for Evaluating Soil Vapor Intrusion in the State of New York (NYSDOH 10/2006) to determine if the medium is contaminated with VOCs. If VOCs are present, the results will be used to evaluate current offsite human exposures and future human exposures within the planned building. The evaluation of current off-site exposure will be useful in determining if further off-site investigation of the exposure pathway is warranted. The evaluation of future on-site exposure will determine whether or not the use of control measures will be necessary to prevent exposure by residents of the new building.

To supplement previously collected soil gas data collected from the interior portions of the Site, 2 soil vapor implants (SG6 and SG7) and 2 sub-slab (SG8 and SG) soil vapor samples will be installed at the perimeter of the Site as shown on Figure 6.

Soil vapor implants will be set at a depth representative of the proposed basement level of the new building which will be approximately twelve feet below surface. Sub slab samples will be installed below the slab of the building.

2.3.1 Soil Vapor Sampling Procedure

The vapor implants will be installed with GeoprobeTM equipment and constructed in the same manner at all locations to minimize possible discrepancies. The implants will be made from stainless steel and fitted with Teflon-lined polyethylene tubing. Coarse sand or glass beads will be added to create a sampling zone of one to two feet in length and sealed above with hydrated bentonite powder for a minimum distance of 3 feet.

The sub-slab soil gas sampling points will be installed by drilling a 1/2 inch hole through the building slab with a handheld drill, and inserting 1/4 inch polyethylene tubing to no more than 2 inches below the base of the slab. The tubing will then be sealed at the surface with hydrated granular bentonite.

After installation of the probes, one to three volumes (i.e., the volume of the sample probe and tube) will be purged prior to collecting the samples to ensure samples collected are representative. Flow rates for both purging and collecting will not exceed 0.2 liters per minute to minimize outdoor air infiltration during sampling. Samples will be collected in Summa® canisters which have been certified clean by the laboratory and analyzed by using USEPA Method TO-15. All samples will be collected over a 2 hour period of time and submitted to a NYSDOH certified laboratory.

A sample log sheet will be maintained summarizing sample identification, date and time of sample collection, sampling depth, identity of samplers, sampling methods and devices, soil vapor purge volumes, volume of soil vapor extracted, vacuum of canisters before and after samples are collected, apparent moisture content of the sampling zone, and chain of custody protocols.

As part of the vapor intrusion evaluation, a tracer gas will be used in accordance with NYSDOH protocols to serves as a quality assurance/quality control (QA/QC) device to verify the integrity of the soil vapor probe seal. Helium will be used as the tracer gas and a box will serve to keep it in contact with the probe during the testing. A portable monitoring device will be used to analyze a sample of soil vapor for the tracer prior to sampling. If tracer sample results show a significant presence of the tracer, the probe seals will be adjusted to prevent infiltration.

After the collection of the analytical sample, a field reading will be recorded at each sampling points utilizing a photoionization detector capable of detecting organic compounds in the parts per billion range.

2.4 Laboratory Analysis

Samples will be submitted to the laboratory for a standard turnaround time, which is estimated to be one to two weeks. The proposed sampling program is summarized in **Table 1**.

2.4.1 Analysis of Soil and Groundwater Samples

Collected soil and groundwater samples will be placed in pre-cleaned laboratory supplied glassware, and placed in a cooler packed with ice for transport to the laboratory. Sample analysis will be provided by a New York State certified environmental laboratory; either York Analytical Laboratories of Stratford Connecticut (NYSDOH Lab I.D. No. 10854) or Phoenix Environmental Laboratories of Manchester Connecticut (NYSDOH Lab I.D. No. 11301). Soil and groundwater samples will be analyzed for the following parameters:

- Volatile organic Compounds (VOCs) by EPA Method 8260
- Semi-volatile organic compounds by EPA Method 8270;
- Pesticides/PCBs by EPA Method 8081/8082; and
- Target Analyte List metals by EPA Method 6010 and 7471;

2.4.2 Analysis of Soil Vapor Samples

Analytical procedures and corresponding reporting limits will be identified when reporting the sampling results. Samples will be analyzed for volatile organic compounds (VOCs) by USEPA

Method TO-15. All samples will be analyzed by a New York State certified environmental laboratory: either York or ALS.

2.5 Management of Investigation Derived Wastes

Investigation derived wastes include contaminated soil, groundwater and disposable sampling equipment generated during the remedial investigation.

Soil from borings will be returned to their original location. Excess soil will be placed in U.S. Department of Transportation (DOT) – approved drums. This material will either be disposed of at an appropriate off-site disposal facility or will be disposed of along with other soil during subsequent remedial activities to be implemented under the RAWP. Purge water generated during groundwater sampling will be containerized in drums and analyzed for VOCs. Final classification and disposal of purge water will be based on the results of this analysis and upon approval of the NYSDEC Project Manager.

Disposable sampling equipment (gloves, tubing, acetate liners, etc.) will be placed in heavy-duty plastic bags and disposed of properly.

3.0 QUALITY ASSURANCE PROJECT PLAN (QAPP)

The fundamental QA objective with respect to accuracy, precision, and sensitivity of analysis for laboratory analytical data is to achieve the QC acceptance of the analytical protocol. The accuracy, precision and completeness requirements will be addressed by the laboratory for all data generated.

Collected samples will be appropriately packaged, placed in coolers and shipped via overnight courier or delivered directly to the analytical laboratory by field personnel. Samples will be containerized in appropriate laboratory provided glassware and shipped in plastic coolers. Samples will be preserved through the use of ice or cold-pak(s) to maintain a temperature of 4°C.

Dedicated disposable sampling materials will be used for both soil and groundwater samples (if collected), eliminating the need to prepare field equipment (rinsate) blanks. However, if non-disposable equipment is used, (stainless steel scoop, etc.) field rinsate blanks will be prepared at the rate of 1 for every eight samples collected.

Decontamination of non-dedicated sampling equipment will consist of the following:

- Gently tap or scrape to remove adhered soil;
- Rinse with tap water;
- Wash with alconox® detergent solution and scrub;
- Rinse with tap water;
- Rinse with distilled or deionized water.

Prepare field blanks by pouring distilled or deionized water over decontaminated equipment and collecting the water in laboratory provided containers. Trip blanks will accompany samples each time they are transported to the laboratory. Matrix spike and matrix spike duplicates (MS/MSD) will be collected at the rate of one per 20 samples submitted to the laboratory. Laboratory reports will be upgradeable to ASP category B deliverables for use in the preparation of a data usability report (DUSR). The DUSR will be applicable to all confirmation samples and final round samples. Performance monitoring samples will be in a results-only format. The QAPP prepared for the Site is provided in **Attachment A**.

3.1 Soil Samples

Dedicated disposable acetate liners will be utilized for sampling, therefore, field equipment (rinsate) blanks will not be part of the QA/QC program. Trip blanks will accompany samples each time they are transported to the laboratory.

3.2 Soil Vapor Samples

Extreme care will be taken during all aspects of sample collection to ensure that sampling error is minimized and high quality data are obtained. The sampling team members will avoid actions (e.g., using permanent marker pens and wearing freshly dry-cleaned clothes or personal fragrances) which can cause sample interference in the field. A tracer gas, helium, will be used in accordance with NYSDOH sampling protocols to serve as a QA/QC device to verify the integrity

of the soil vapor probe seals. QA/QC protocols will be followed for sample collection and laboratory analysis, such as use of certified clean sample devices, meeting sample holding times and temperatures, sample accession, and chain of custody.

Samples will be delivered to the analytical laboratory as soon as possible after collection. The laboratory analyzes QC samples with each analytical batch, including a Method Blank (MB), Laboratory Control Sample (LCS), and a Laboratory Control Sample Duplicate (LCSD). Internal standards are added to all calibration standards, samples, and blanks to verify that the analytical system is in control.

3.3 Reporting of Results

Sample analysis will be provided by a New York State certified environmental laboratory. Laboratory reports will include ASP category B deliverables for use in the preparation of a data usability summary report (DUSR). All results will be provided in accordance with the NYSDEC Environmental Information Management System (EIMS) electronic data deliverable (EDD) format.

3.4 DUSR

The DUSR provides a thorough evaluation of analytical data without third party data validation. The primary objective of a DUSR is to determine whether or not the data, as presented, meets the site/project specific criteria for data quality and data use. Verification and/or performance monitoring samples collected under this RIWP will be reviewed and evaluated in accordance with the Guidance for the Development of Data Usability Summary Reports as presented in Appendix 2B of DER-10. The completed DUSR for verification/performance samples collected during implementation of this RIWP will be included in the Remedial Investigation report. DUSRs will be evaluated by Alpha Geosciences.

4.0 HEALTH AND SAFETY PLAN

The Health and Safety Plan (HASP) takes into account the specific hazards inherent in conducting the RI, and presents the minimum requirements which are to be met by Environmental Business Consultants (EBC), its subcontractors, and other personnel in order to avoid and, if necessary, protect against health and/or safety hazards. A HASP has been prepared and is provided in **Attachment B** of this work plan.

Sub-contractors will have the option of adopting this HASP or developing their own site-specific document. If a subcontractor chooses to prepare their own HASP, it must meet the minimum requirements as detailed in the RI HASP prepared by EBC and must be made available to EBC and the NYSDEC.

Activities performed under the HASP will comply with applicable parts of OSHA Regulations, primarily 29 CFR Parts 1910 and 1926. Modifications to the HASP may be made with the approval of the EBC Site Safety Manager (SSM) and/or Project Manager (PM).

5.0 COMMUNITY AIR MONITORING PLAN

The Community Air Monitoring Plan (CAMP) provides measures for protection for on-site workers and the downwind community (i.e., off-site receptors including residences, businesses, and on-site commercial workers) from potential airborne contaminant releases resulting from investigation activities.

The action levels specified require increased monitoring, corrective actions to abate emissions, and/or work shutdown. Additionally, the CAMP helps to confirm that the investigation work did not spread contamination off-site through the air.

The primary concerns during the investigation are odors from VOCs. The CAMP for this investigation is provided as **Attachment C**.

6.0 REMEDIAL INVESTIGATION REPORT

Following completion of the investigation and receipt of the analytical data, EBC will prepare a Remedial Investigation Report (RIR) in accordance with DER10. The RIR will which will include the following:

- 1. A description of the work which was performed under the RI.
- 2. Any modification from this work scope and the reason for the modifications
- 3. The nature and extent of the off-site groundwater plume
- 4. Soil, and groundwater conditions that were observed
- 5. Analytical data in tabular form comparing results to part 375-6 SCOs
- 6. Cross sections and data figures
- 7. Laboratory analytical data, sampling logs and well completion logs for all samples and areas covered by the investigation
- 8. Scaled drawings showing the locations of temporary sampling points, monitoring wells and surface water sampling locations
- 9. A Qualitative Human Health Exposure Assessment



7.0 SCHEDULE

Implementation of the RI will be performed prior to building demolition. Since the building is currently vacant, access to most of the proposed sampling locations will not pose any challenges. However, some preparation may be required such as creating overhead clearance to access some the locations.

Mobilization for the field work is anticipated to begin approximately 2 weeks following NYSDEC approval of the RI Work Plan and 44 days following the public notification of the BCP application. The estimated duration of the full RI activity is one weeks total field time.

The anticipated schedule of events is as follows:

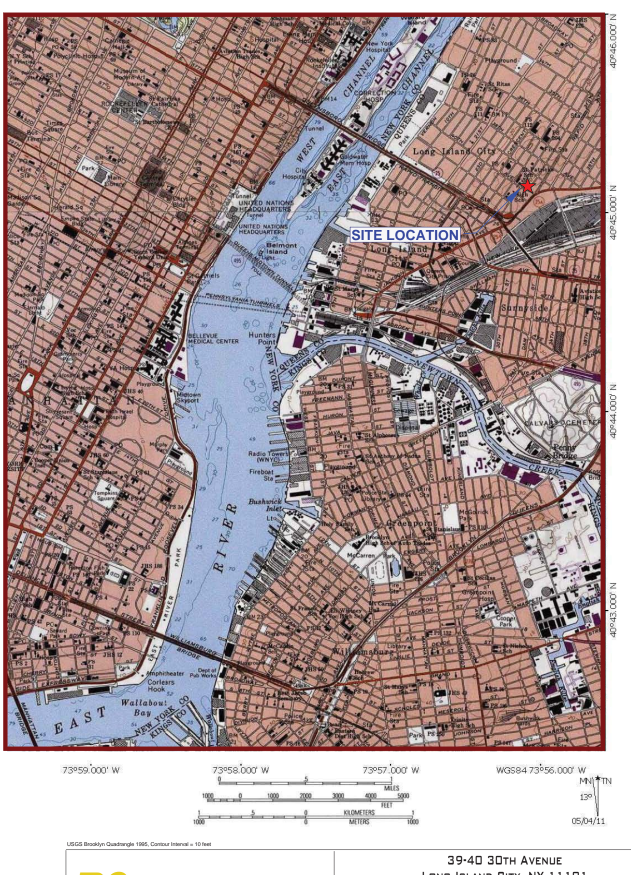
Schedule Task	Estimated Date	
End of 30 day Public Comment Period	September1, 2014	
NYSDEC Approval of RIWP	Week of September 22, 2014	
Mobilize equipment to the Site (begin)	Week of September 29, 2014	
Complete Field Work	Week of September 29, 2014	
Receive all Laboratory Reports	Week of October 6, 2014	
Submit Remedial Investigation Report	Week of October 20, 2014	
Distribute Fact Sheet on RI Results and Comment period on RAWP (if submitted with RIR)	Within 6 weeks of completion of field work	

TABLES

TABLE 1 SUMMARY OF SAMPLING PROGRAM RATIONALE AND ANALYSIS

Matrix	Location	Approximate Number of Samples	Rationale for Sampling	Laboratory Analysis
Subsurface soil (0 to 15 feet bgs)	Six borings located across the Site. Two samples from each boring including one sample from 0-2 feet and one from 13-15 ft.	12	To supplement previous sampling and delineate CVOC affected soil at the Site.	VOCs EPA Method 8260B, SVOCs EPA Method 8270, metals EPA Method 6010 and 7471, Pesticides and PCBs EPA Method 8081/ 8082
Subsurface soil (0 to 15 feet bgs)	If stained or contmainated soil is encounted on site, additional contingency samples will be collected	1 or 2	If stained or contmainted samples are encournted during the investigation, these contingency samples will be collected.	VOCs EPA Method 8260B
Total (Soils)		12		
Groundwater (water table)	From five existing monitoring wells on-site and two new monitoring wells to be installed on-site.	7	Define nature and extent of impacted groundwater	VOCs EPA Method 8260B
Total (Groundwater)		7		
Sub Slab and Soil Gas implants (12 ft below existing grade)	Sub Slab and Soil Gas implants and two sub slab implants (12 ft below existing locations to be installed on the Site.	4	Evaluate soil gas at perimeter of Site.	VOCs EPA Method TO15
Total (Soil Gas)		4		
MS/MSD	Matrix spike and Matrix spike duplicates at the rate 5%	2	To meet requirements of QA / QC program	2 MS/MSD (1 soil 1 GW) for VOCs EPA Method 8260B
Trip Blanks	One laboratory prepared trip blank to accompany samples each time they are delivered to the laboratory.	2	To meet requirements of QA / QC program	VOCs EPA Method 8260B
Total (QA / QC Samples)				

FIGURES

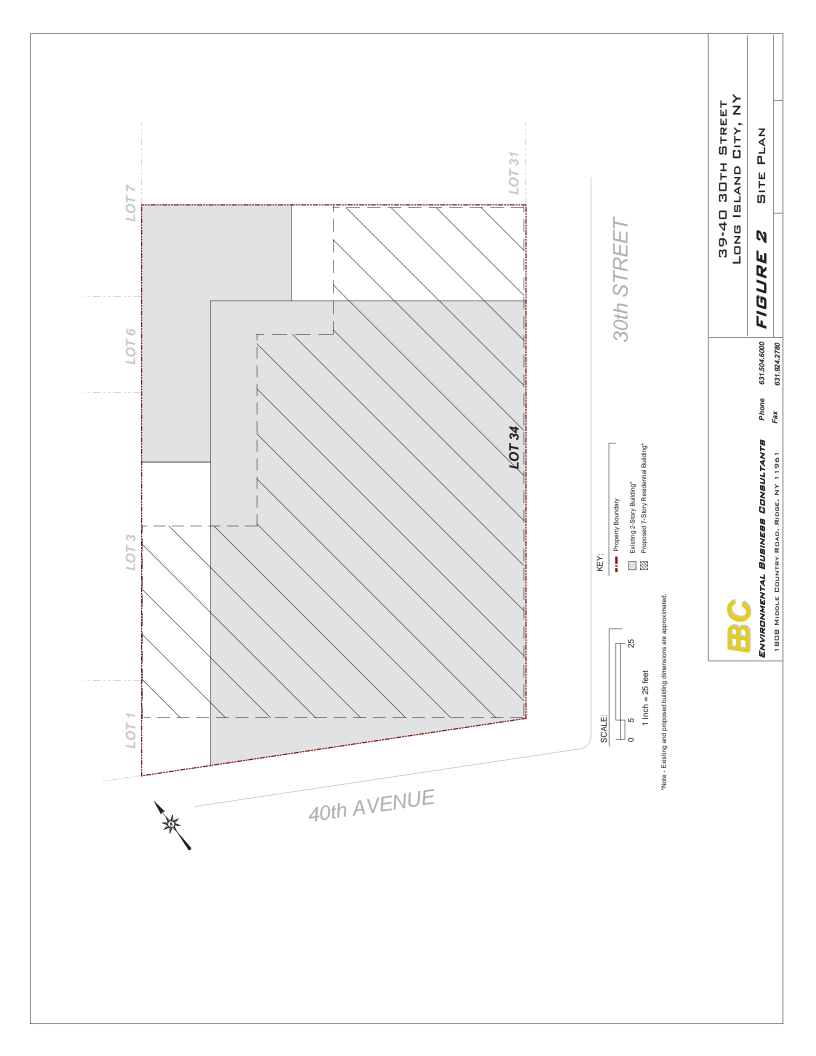


631.504.6000 631. 924 .2870 ENVIRONMENTAL BUSINESS CONSULTANTS

LONG ISLAND CITY, NY 11101

FIGURE 1

SITE LOCATION MAP



FORMER UNION WIRE DIE SITE 39-40 30TH STREET, LONG ISLAND CITY, NY

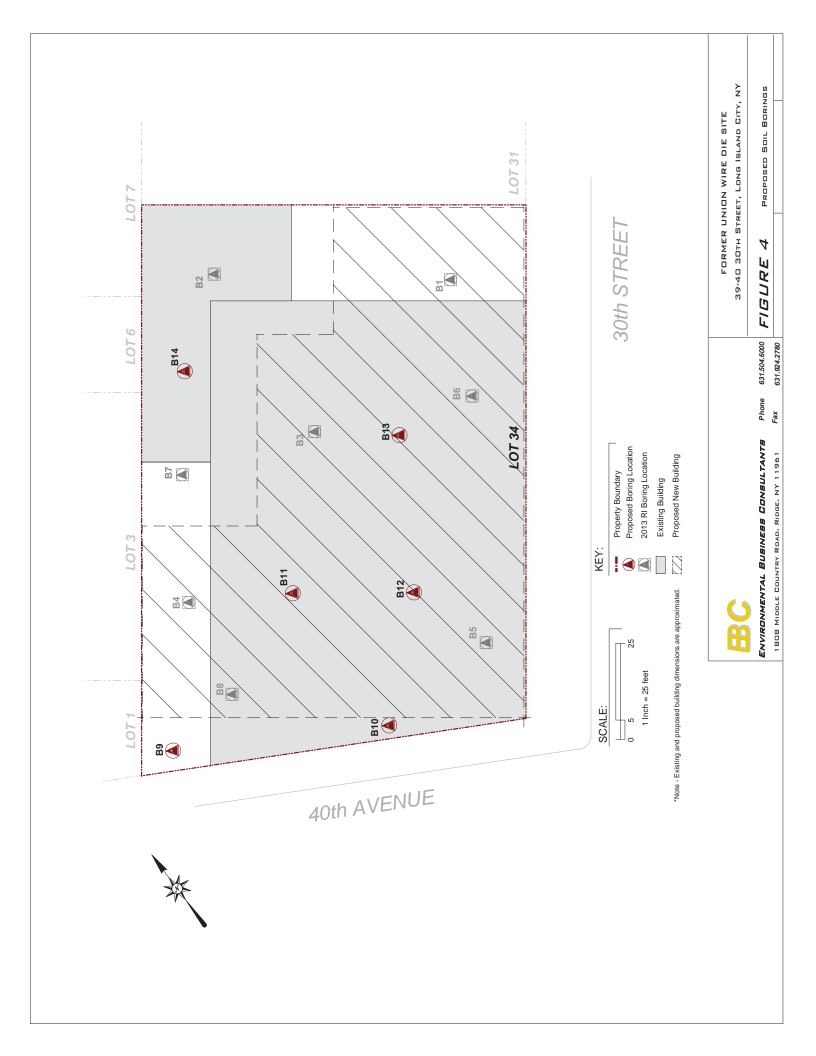
FIGURE

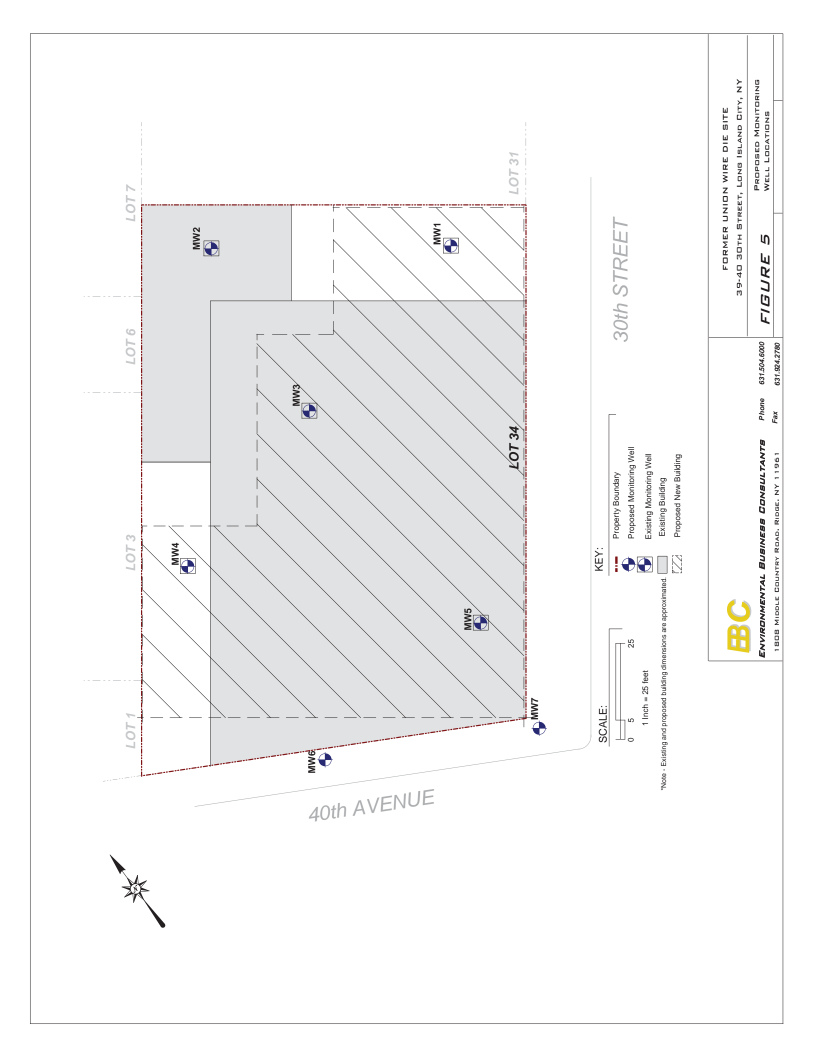
η

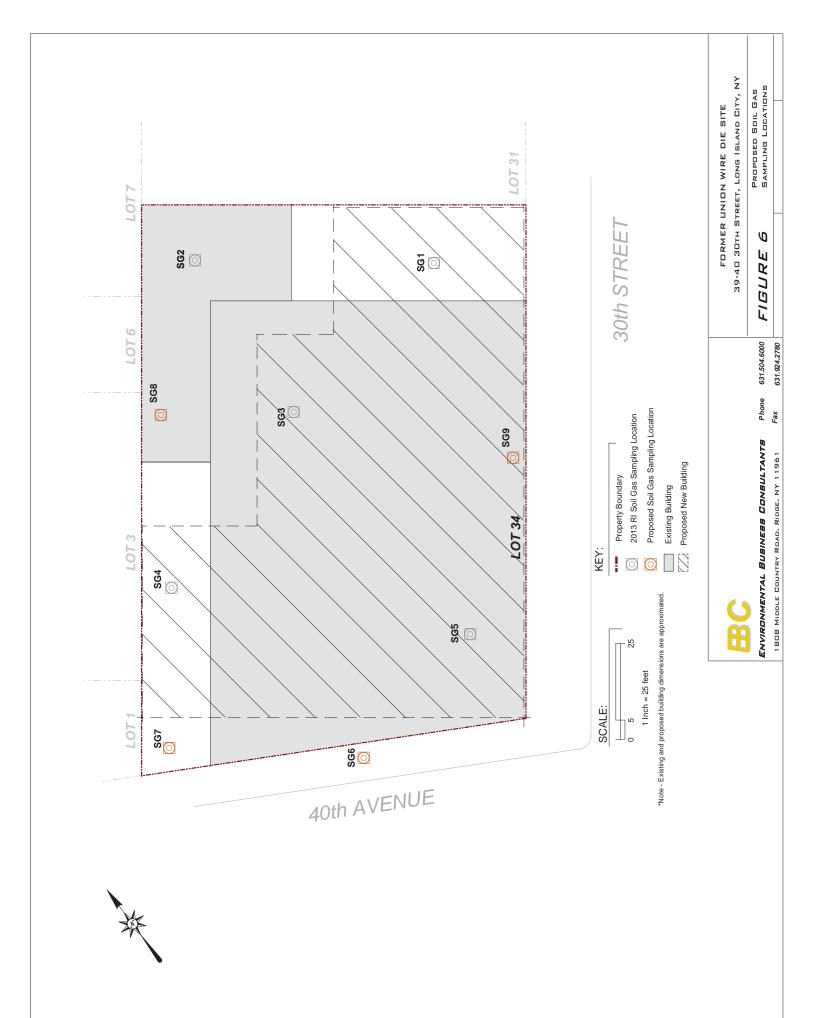
GROUNDWATER ELEVATION

ENVIRONMENTAL BUSINESS CONSULTANTS
1808 MIDDLE COUNTRY ROAD. RIDGE. NY 11961

Phone 631.504.6000 Fax 631.924.2780







ATTACHMENT A Quality Assurance Project Plan

QUALITY ASSURANCE PROJECT PLAN Former Union Die Corp. 39-40 30th Street, Long Island City, NY

Prepared on behalf of:

Ganesh Management LLC 39-40 30th Street Long Island City, NY

Prepared by:

ENVIRONMENTAL BUSINESS CONSULTANTS
RIDGE, NY 11961

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39-40 30th Street, Long Island City, NY

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1.0 INTRODUCTION

This Quality Assurance Project Plan (QAPP) has been prepared in accordance with DER-10 to detail procedures to be followed during the course of the sampling and analytical portion of the project, as required by the approved work plan.

To ensure the successful completion of the project each individual responsible for a given component of the project must be aware of the quality assurance objectives of his / her particular work and of the overall project. The EBC Project Director, Charles Sosik will be directly responsible to the client for the overall project conduct and quality assurance/quality control (QA/QC) for the project. The Project Director will be responsible for overseeing all technical and administrative aspects of the project and for directing QA/QC activities. As Project Director Mr. Sosik will also serve as the Quality Assurance Officer (QAO) and in this role may conduct:

- conduct periodic field and sampling audits;
- interface with the analytical laboratory to resolve problems; and
- interface with the data validator and/or the preparer of the DUSR to resolve problems.

Chawinie Miller will serve as the Project Manager and will be responsible for implementation of the Remedial Investigation and coordination with field sampling crews and subcontractors. Reporting directly to the Project Manager will be the Field Operations Officer, Kevin Waters; who will serve as the on-Site qualified environmental professional who will record observations, direct the drilling crew and be responsible for the collection and handling of all samples.

1.1 Organization

Project QA will be maintained under the direction of the Project Manager, in accordance with this QAPP. QC for specific tasks will be the responsibility of the individuals and organizations listed below, under the direction and coordination of the Project Manager

GENERAL RESPONSIBILITY	SCOPE OF WORK	RESPONSIBILITY OF QUALITY CONTROL
Field Operations	Supervision of Field Crew, sample collection and handling	K. Waters, EBC
Project Manager	Implementation of the RI according to the RIWP.	Chawinie Miller, EBC
Laboratory Analysis	Analysis of soil samples by NYSDEC ASP methods Laboratory	NYSDOH-Certified Laboratory
Data review	Review for completeness and compliance	3 rd party validation

2.0 QUALITY ASSURANCE PROJECT PLAN OBJECTIVES

2.1 Overview

Overall project goals are defined through the development of Data Quality Objectives (DQOs), which are qualitative and quantitative Statements that specify the quality of the data required to support decisions; DQOs, as described in this section, are based on the end uses of the data as described in the work plan.

In this plan, Quality Assurance and Quality Control are defined as follows:

- Quality Assurance The overall integrated program for assuring reliability of monitoring and measurement data.
- Quality Control The routine application of procedures for obtaining prescribed standards of performance in the monitoring and measurement process.

2.2 OA / OC Requirements for Analytical Laboratory

Samples will be analyzed by a New York State Department of Health (NYSDOH) certified laboratory. Data generated from the laboratory will be used to evaluate contaminants such as chlorinated and other volatile organic compounds (VOCs) in soil, soil gas and groundwater, SVOCs in soil, metals in soil and PCBs and pesticides in soil. The QA requirements for all subcontracted analytical laboratory work performed on this project are described below. QA elements to be evaluated include accuracy, precision, sensitivity, representativeness, and completeness. The data generated by the analytical laboratory for this project are required to be sensitive enough to achieve detection levels low enough to meet required quantification limits as specified in NYSDEC Analytical Services Protocol (NYSDEC ASP, 07/2005. The analytical results meeting the required quantification limits will provide data sensitive enough to meet the data quality objectives of this remedial program as described in the work plan. Reporting of the data must be clear, concise, and comprehensive. The QC elements that are important to this project are completeness of field data, sample custody, sample holding times, sample preservation, sample storage, instrument calibration and blank contamination.

2.2.1 Instrument Calibration

Calibration curves will be developed for each of the compounds to be analyzed. Standard concentrations and a blank will be used to produce the initial curves. The development of calibration curves and initial calibration response factors must be consistent with method requirements presented in the most recent version of NYSDEC ASP 07/2005).

2.2.2 Continuing Instrument Calibration

The initial calibration curve will be verified every 12 hrs by analyzing one calibration standard. The standard concentration will be the midpoint concentration of the initial calibration curve. The calibration check compound must come within 25% relative percent difference (RPD) of the average response factor obtained during initial calibration. If the RPD is greater than 25%, then corrective action must be taken as provided in the specific methodology.

2.2.3 Method Blanks

Method blank or preparation blank is prepared from an analyte-free matrix which includes the same reagents, internal standards and surrogate standards as me related samples. II is carried through the

entire sample preparation and analytical procedure. A method blank analysis will be performed once for each 12 hr period during the analysis of samples for volatiles. An acceptable method blank will contain less than two (2) times the CRQL of methylene chloride, acetone and 2-butanone. For all other target compounds, the method blank must contain less than or equal to the CRQL of any single target compound. For non-target peaks in the method blank, the peak area must be less than 10 percent of the nearest internal standard. The method blank will be used to demonstrate the level of laboratory background and reagent contamination that might result from the analytical process itself.

2.2.4 Trip Blanks.

Trip blanks consist of a single set of sample containers filled at the laboratory with deionized. laboratory-grade water. The water used will be from the same source as that used for the laboratory method blank. The containers will be carried into the field and handled and transported in the same way as the samples collected that day. Analysis of the trip blank for VOCs is used to identify contamination from the air, shipping containers, or from other items coming in contact with the sample bottles. (The bottles holding the trip blanks will be not opened during this procedure.) A complete set of trip blanks will be provided with each shipment of samples to the certified laboratory.

2.2.5 Surrogate Spike Analysis

For organic analyses, all samples and blanks will be spiked with surrogate compounds before purging or extraction in order to monitor preparation and analyses of samples. Surrogate spike recoveries shall fall within the advisory limits in accordance with the NY5DEC ASP protocols for samples falling within the quantification limits without dilution.

2.2.6 Matrix Spike / Matrix Spike Duplicate / Matrix Spike Blank (MS/MSDIMSB) Analysis MS, MSD and MSB analyses will be performed to evaluate the matrix effect of the sample upon the analytical methodology along with the precision of the instrument by measuring recoveries. The MS / MSD / MSB samples will be analyzed for each group of samples of a similar matrix at a rate of 5% (one for every 20 field samples). The RPD will be calculated from the difference between the MS and MSD. Matrix spike blank analysis will be performed to indicate the appropriateness of the spiking solution(s) used for the MS/MSD.

2.3 Accuracy

Accuracy is defined as the nearness of a real or the mean (x) of a set of results to the true value. Accuracy is assessed by means of reference samples and percent recoveries. Accuracy includes both precision and recovery and is expressed as percent recovery (% REC). The MS sample is used to determine the percent recovery. The matrix spike percent recovery (% REC) is calculated by the following equation:

$$\%REC = \frac{SSR - SR}{SA} \times 100$$

Where:

SSR = spike sample results

SR = sample results

SA = spike added from spiking mix



2.4 Precision

Precision is defined as the measurement of agreement of a set of replicate results among themselves without a Precision is defined as the measurement of agreement of a set of replicate results among themselves without assumption of any prior information as to the true result. Precision is assessed by means of duplicate/replicate sample analyses.

Analytical precision is expressed in terms of RPD. The RPD is calculated using the following formula:

$$RPD = \frac{D^{1} - D^{2}}{(D^{1} - D^{2})/2} \times 100$$

Where:

RPD = relative percent difference

 D^1 = first sample value

 D^2 = second sample value (duplicate)

2.5 Sensitivity

The sensitivity objectives for this plan require that data generated by the analytical laboratory achieve quantification levels low enough to meet the required detection limits specified by NYSDEC ASP and to meet all site-specific standards, criteria and guidance values (SGCs) established for this project.

2.6 Representativeness

Representativeness is a measure of the relationship of an individual sample taken from a particular site to the remainder of that site and the relationship of a small aliquot of the sample (i.e., the one used in the actual analysis) to the sample remaining on site. The representativeness of samples is assured by adherence to sampling procedures described in the Remedial Investigation Work Plan.

2.7 Completeness

Completeness is a measure of the quantity of data obtained from a measurement system as compared to the amount of data expected from the measurement system. Completeness is defined as the percentage of all results that are not affected by failing QC qualifiers, and should be between 70 and 100% of all analyses performed. The objective of completeness in laboratory reporting is to provide a thorough data support package. The laboratory data package provides documentation of sample analysis and results in the form of summaries, QC data, and raw analytical data. The laboratory will be required to submit data packages that follow NYSDEC ASP reporting format which, at a minimum, will include the following components:

- 1. All sample chain-of-custody forms.
- 2. The case narrative(s) presenting a discussion of any problems and/or procedural changes required during analyses. Also presented in the case narrative are sample summary forms.
- 3. Documentation demonstrating the laboratory's ability to attain the contract specified detection limits for all target analytes in all required matrices.
- 4. Tabulated target compound results and tentatively identified compounds.
- 5. Surrogate spike analysis results (organics).
- 6. Matrix spike/matrix spike duplicate/matrix spike blank results.
- 7. QC check sample and standard recovery results
- 8. Blank results (field, trip, and method).



9. Internal standard area and RT summary.

2.8 Laboratory Custody Procedures

The following elements are important for maintaining the field custody of samples:

- Sample identification
- Sample labels
- Custody records
- Shipping records
- Packaging procedures

Sample labels will be attached to all sampling bottles before field activities begin; each label will contain an identifying number. Each number will have a suffix that identifies the site and where the sample was taken. Approximate sampling locations will be marked on a map with a description of the sample location. The number, type of sample, and sample identification will be entered into the field logbook. A chain-of-custody form, initiated at the analytical laboratory will accompany the sample bottles from the laboratory into the field. Upon receipt of the bottles and cooler, the sampler will sign and date the first received blank space. After each sample is collected and appropriately identified, entries will be made on the chain-of-custody form that will include:

- Site name and address
- Samplers' names and signatures

3.0 ANALYTICAL PROCEDURES

3.1 Laboratory Analysis

Samples will be analyzed by the NYSDOH ELAP laboratory for one or more of the following parameters: VOCs in soil / groundwater by USEPA Method 8260, SVOCs in soil by USEPA Method 8270BN, Target Analyte List (TAL) Metals in soil, pesticides / PCBs by USEPA Method 8081/8082 and VOCs in air by USEPA Method TO15. If any modifications or additions to the standard procedures are anticipated. and if any nonstandard sample preparation or analytical protocol is to be used, the modifications and the nonstandard protocol will be explicitly defined and documented. Prior approval by EBC's PM will be necessary for any nonstandard analytical or sample preparation protocol used by the laboratory, i.e., dilution of samples or extracts by greater than a factor of five (5).



4.0 DATA REDUCTION, REVIEW, AND REPORTING

4.1 Overview

The process of data reduction, review, and reporting ensures the assessments or a conclusion based on the final data accurately reflects actual site conditions. This plan presents the specific procedures, methods, and format that will be employed for data reduction, review and reporting of each measurement parameter determined in the laboratory and field. Also described in this section is the process by which all data, reports, and work plans are proofed and checked for technical and numerical errors prior to final submission.

4.2 Data Reduction

Standard methods and references will be used as guidelines for data handling, reduction, validation, and reporting. All data for the project will be compiled and summarized with an independent verification at each step in the process to prevent transcription/typographical errors. Any computerized entry of data will also undergo verification review.

Sample analysis will be provided by a New York State certified environmental laboratory. Laboratory reports will include ASP category B deliverables for use in the preparation of a data usability summary report (DUSR). All results will be provided in accordance with the NYSDEC Environmental Information Management System (EIMS) electronic data deliverable (EDD) format. Analytical results shall be presented on standard NYSDEC ASP-B forms or equivalents, and include the dates the samples were received and analyzed, and the actual methodology used. Note that if waste characterization samples are analyzed they will be in results only format and will not be evaluated in the DUSR.

Laboratory QA/QC information required by the method protocols will be compiled, including the application of data QA/QC qualifiers as appropriate. In addition, laboratory worksheets, laboratory notebooks, chains-of-custody, instrument logs, standards records, calibration records, and maintenance records, as applicable, will be provided in the laboratory data packages to determine the validity of data. Specifics on internal laboratory data reduction protocols are identified in the laboratory's SOPs.

Following receipt of the laboratory analytical results by EBC, the data results will be compiled and presented in an appropriate tabular form. Where appropriate, the impacts of QA/QC qualifiers resulting from laboratory or external validation reviews will be assessed in terms of data usability.

4.3 Laboratory Data Reporting

All sample data packages submitted by the analytical laboratory will be required to be reported in conformance to the NYSDEC ASP (7/2005), Category B data deliverable requirements as applicable to the method utilized. All results will be provided in accordance with the NYSDEC Environmental Information Management System (EIMS) electronic data deliverable (EDD) format. Note that waste characterization samples if analyzed will be in results only format and will not be evaluated in the DUSR.

5.0 CORRECTIVE ACTION

Review and implementation of systems and procedures may result in recommendations for corrective action. Any deviations from the specified procedures within approved project plans due to unexpected site-specific conditions shall warrant corrective action. All errors, deficiencies, or other problems shall be brought to the immediate attention of the EBC PM, who in turn shall contact the Quality Assurance/Data Quality Manager or his designee (if applicable).

Procedures have been established to ensure that conditions adverse to data quality are promptly investigated, evaluated and corrected. These procedures for review and implementation of a change are as follows:

- Define the problem.
- Investigate the cause of the problem.
- Develop a corrective action to eliminate the problem, in consultation with the personnel who defined the problem and who will implement the change.
- Complete the required form describing the change and its rationale (see below for form requirements).
- Obtain all required written approvals.
- Implement the corrective action.
- Verify that the change has eliminated the problem.

During the field investigation, all changes to the sampling program will be documented in field logs/sheets and the EBC PM advised.

If any problems occur with the laboratory or analyses, the laboratory must immediately notify the PM, who will consult with other project staff. All approved corrective actions shall be controlled and documented.

All corrective action documentation shall include an explanation of the problem and a proposed solution which will be maintained in the project file or associated logs. Each report must be approved by the necessary personnel (e.g., the PM) before implementation of the change occurs. The PM shall be responsible for controlling, tracking, implementing and distributing identified changes.

TABLE 1 SUMMARY OF SAMPLING PROGRAM RATIONALE AND ANALYSIS

Matrix	Location	Approximate Number of Samples	Rationale for Sampling	Laboratory Analysis
Subsurface soil (0 to 15 feet bgs)	Six borings located across the Site. Two samples from each boring including one sample from 0-2 feet and one from 13-15 ft.	12	To supplement previous sampling and delineate CVOC affected soil at the Site.	VOCs EPA Method 8260B, SVOCs EPA Method 8270, metals EPA Method 6010 and 7471, Pesticides and PCBs EPA Method 8081/ 8082
Subsurface soil (0 to 15 feet bgs)	If stained or contmainated soil is encounted on site, additional contingency samples will be collected	1 or 2	If stained or contmainted samples are encournted during the investigation, these contingency samples will be collected.	VOCs EPA Method 8260B
Total (Soils)		12		
Groundwater (water table)	From five existing monitoring wells on-site and two new monitoring wells to be installed on-site.	7	Define nature and extent of impacted groundwater	VOCs EPA Method 8260B
Total (Groundwater)		7		
Sub Slab and Soil Gas implants (12 ft below existing grade)	Sub Slab and Soil Gas implants and two sub slab implants (12 ft below existing locations to be installed on the Site.	4	Evaluate soil gas at perimeter of Site.	VOCs EPA Method TO15
Total (Soil Gas)		4		
MS/MSD	Matrix spike and Matrix spike duplicates at the rate 5%	2	To meet requirements of QA / QC program	2 MS/MSD (1 soil 1 GW) for VOCs EPA Method 8260B
Trip Blanks	One laboratory prepared trip blank to accompany samples each time they are delivered to the laboratory.	2	To meet requirements of QA / QC program	VOCs EPA Method 8260B
Total (QA / QC Samples)				

SAMPLE COLLECTION AND ANALYSIS PROTOCOLS **TABLE 2**

		. I	-		G1	A14	/ 1040	TT 1.11.
Samble	Matrix	Samping	rarameter	Sample	Sample	Analyucal	CKQL/	Holding
Type		Device		Container	Preservation	Method#	MDLH	Time
Soil	Soil	doooS	NOCs	(1) 2 oz Jar	Cool to 4° C	EPA Method 8260	Compound specific	14 days
		Direct into Jar			HCL		(1-5 ug/kg)	
Soil	Soil	Scoop	SVOCs	(1) 8 oz jar	Cool to 4° C	EPA Method 8270 BN	Compound specific	14 day ext/40 days
		Direct into Jar					(1-5 ug/kg)	
Soil	Soil	Scoop	Pest/PCBs	from 8oz jar	Cool to 4° C	EPA Method	Compound specific	14 day ext/40 days
		Direct into Jar		above		8081/8082	(1-5 ug/kg)	
Soil	Soil	Scoop	Metals	from 8oz jar	Cool to 4° C	TAL Metals	Compound specific	6 months
		Direct into Jar		above			(01-1 mg/kg)	
Groundwater	Water	Pump tubing	VOCs	(3) 40 ml	Cool to 4° C	EPA Method 8260	Compound specific	14 days
				vials			(1-5 ug/L)	
Groundwater	Water	Pump tubing	SVOCs	(1) 1 Liter	Cool to 4° C	EPA Method 8270 BN	Compound specific	14 days
				Amber Bottle			(1-5 ug/L)	
Groundwater	Water	Pump tubing	Pesticides	(2) 1 Liter	Cool to 4° C	EPA Method 8081/	Compound specific	14 days
			and PCBs	Amber Bottle		8082	(1-5 ug/L)	
Groundwater	water	Pump tubing	Total Metals	(1) 100 ml	HNO3	TAL Metals	Compound specific	6 months
							(1-5 mg/L)	
Groundwater	water	Pump tubing	Dissolved	(1) 100 ml	None	TAL Metals	Compound specific	6 months
			Metals				(1-5 mg/L)	

Notes:

All holding times listed are from Verified Time of Sample Receipt (VTSR) unless noted otherwise. * Holding time listed is from time of sample collection. The number in parentheses in the "Sample Container" column denotes the number of containers needed.

Triple volume required when collected MS/MSD samples

The number of trip blanks are estimated. CRQL / MDL = Contract Required Quantitation Limit / Method Detection Limit. MCAWW = Methods for Chemical Analysis of Water and Wastes. NA = Not available or not applicable.

ATTACHMENT B Health and Safety Plan

FORMER UNION WIRE DIE CORP. 39-40 30th STREET

39-40 30th STREET LONG ISLAND CITY, NEW YORK Block 399 Lot 4

HEALTH AND SAFETY PLAN

SEPTEMBER 2014

Prepared for:
Ganesh Management LLC
39-40 30th Street
Long Island City, NY 11101

Prepared By:



ENVIRONMENTAL BUSINESS CONSULTANTS

1808 Middle Country Road Ridge, NY 11961

HEALTH AND SAFETY PLAN

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APPENDIX B SITE SAFETY PLAN AMENDMENTS

APPENDIX C CHEMICAL HAZARDS

APPENDIX D HOSPITAL INFORMATION, MAP AND FIELD ACCIDENT REPORT

PHONE: 631.504.6000 | 631.924.2870

STATEMENT OF COMMITMENT

This Health and Safety Plan (HASP) has been prepared to ensure that workers are not exposed to risks from hazardous materials during the planned Subsurface Investigation at the Site.

This HASP, which applies to persons present at the site actually or potentially exposed to hazardous materials, describes emergency response procedures for actual and potential chemical hazards. This HASP is also intended to inform and guide personnel entering the work area or exclusion zone. Persons are to acknowledge that they understand the potential hazards and the contents of this Health and Safety policy by signing off on receipt of their individual copy of the document. Contractors and suppliers are retained as independent contractors and are responsible for ensuring the health and safety of their own employees.

1.0 INTRODUCTION AND SITE ENTRY REQUIREMENTS

This document describes the health and safety guidelines developed by Environmental Business Consultants (EBC) for the subsurface investigation to be performed to protect on-site personnel, visitors, and the public from physical harm and exposure to hazardous materials or wastes during subsurface investigation activities. In accordance with the Occupational Safety and Health Administration (OSHA) 29 CFR Part 1910.120 Hazardous Waste Operations and Emergency Response Final rule, this HASP, including the attachments, addresses safety and health hazards related to subsurface sample collection activities and is based on the best information available. The HASP may be revised by EBC at the request of the client and/or a regulatory agency upon receipt of new information regarding site conditions. Changes will be documented by written amendments signed by EBC's project manager, site safety officer and/or the EBC health and safety consultant.

1.1 Training Requirements

Personnel entering the exclusion zone or decontamination zone are required to be certified in health and safety practices for hazardous waste site operations as specified in the Federal OSHA Regulations CFR 1910.120e (revised 3/6/90).

Paragraph (e - 3) of the above referenced regulations requires that all on-site management personnel directly responsible for or who supervise employees engaged in hazardous waste operations, must initially receive 8 hours of supervisor training related to managing hazardous waste work.

Paragraph (e - 8) of the above referenced regulations requires that workers and supervisors receive 8 hours of refresher training annually on the items specified in Paragraph (e-1) and/or (e-3).

Additionally all on-site personnel must receive adequate site-specific training in the form of an on-site Health and Safety briefing prior to participating in field work with emphasis on the following:

- Protection of the adjacent community from hazardous vapors and / or dust which may be released during intrusive activities.
- Identification of chemicals known or suspected to be present on-site and the health effects and hazards of those substances.
- The need for vigilance in personnel protection, and the importance of attention to proper use, fit and care of personnel protective equipment.
- Decontamination procedures.

IRONMENTAL BUSINESS CONSULTANTS

- Site control including work zones, access and security.
- Hazards and protection against heat or cold.
- The proper observance of daily health and safety practices, such as entry and exit of work zones and site. Proper hygiene during lunch, break, etc.
- Emergency procedures to be followed in case of fire, explosion and sudden release of hazardous gases.



Health and Safety meetings will be conducted on a daily basis and will cover protective clothing and other equipment to be used that day, potential and chemical and physical hazards, emergency procedures, and conditions and activities from the previous day.

1.2 Site Safety Plan Acceptance, Acknowledgment and Amendments

The project superintendent and the site safety officer are responsible for informing personnel (EBC employees and/or owner or owners representatives) entering the work area of the contents of this plan and ensuring that each person signs the safety plan acknowledging the on-site hazards and procedures required to minimize exposure to adverse effects of these hazards. A copy of the Acknowledgement Form is included in **Appendix A**.

Site conditions may warrant an amendment to the HASP. Amendments to the HASP are acknowledged by completing forms included in **Appendix B**.

1.3 Key Personnel - Roles and Responsibilities

Personnel responsible for implementing this Health and Safety Plan are:

Name	Title	Address	Contact
			Numbers
Ms. Chawinie Miller	EBC	1808 Middle Country	(631) 504-6000
	Project Manager	Road	(631) 827-5007
		Ridge, NY 11961	
Mr. Kevin Waters	Site Safety Officer	1808 Middle Country	(631) 504-6000
		Road	(516) 287-9023
		Ridge, NY 11961	
Mr. Paul Dibone	C-Squared	99 Jericho Turnpike,	(516) 417-8588
	Environmental	Suite 300L	(631) 948-6553
	Services	Jericho, NY	
	Equipment Operator		

The project manager is responsible for overall project administration and, with guidance from the site safety officer, for supervising the implementation of this HASP. The site safety officer will conduct daily (tail gate or tool box) safety meetings at the project site and oversee daily safety issues. Each subcontractor and supplier (defined as an OSHA employer) is also responsible for the health and safety of its employees. If there is any dispute about health and safety or project activities, on-site personnel will attempt to resolve the issue. If the issue cannot be resolved at the site, then the project manager will be consulted.

The site safety officer is also responsible for coordinating health and safety activities related to hazardous material exposure on-site. The site safety officer is responsible for the following:

- 1. Educating personnel about information in this HASP and other safety requirements to be observed during site operations, including, but not limited to, decontamination procedures, designation of work zones and levels of protection, air monitoring, fit testing, and emergency procedures dealing with fire and first aid.
- 2. Coordinating site safety decisions with the project manager.



- 3. Designating exclusion, decontamination and support zones on a daily basis.
- 4. Monitoring the condition and status of known on-site hazards and maintaining and implementing the air quality monitoring program specified in this HASP.
- 5. Maintaining the work zone entry/exit log and site entry/exit log.
- 6. Maintaining records of safety problems, corrective measures and documentation of chemical exposures or physical injuries (the site safety officer will document these conditions in a bound notebook and maintain a copy of the notebook on-site).

The person who observes safety concerns and potential hazards that have not been addressed in the daily safety meetings should immediately report their observations/concerns to the site safety officer or appropriate key personnel.

2.0 SITE BACKGROUND AND SCOPE OF WORK

A Phase II Subsurface Investigation is being conducted at the site to identify and characterize potential contaminants within the surface/subsurface soils, groundwater and soil gas at the site.

The results from this investigation will help determine what actions may be required, if any, to prevent exposure to contaminants from the change in use of the site. The work will be conducted in accordance with the procedures as required under the New York State Department of Environmental Conservation Brownfield Cleanup Program (NYSDEC BCP).

The Site was undeveloped from 1887 to sometime between 1915 and 1936. By 1936, a gas station with two gasoline tanks occupied the site. The property was redeveloped by 1947 into a 2-story warehouse utilized by Optical Products Corporation for manufacturing, shipping, and as an office. The building has remained since, with several other occupants including Union Wire Die Corp (1960s), National Tea Packaging Co. Inc. (1962-1991), and a warehouse (1991-2006). A Remedial Investigation performed at the Site, identified chlorinated volatile organic compounds in soil, groundwater and soil vapor.

The results of the RI identified elevated levels of trichloroethene (TCE) in shallow soil (above the water table), in three of the boring locations at concentrations above unrestricted soil cleanup objectives (SCOs). Chlorinated VOCs were detected above NYSDEC groundwater standards. In addition petroleum related VOCs were also reported above standards in one sample.

Both PCE and TCE were reported in soil gas above mitigation levels established within the State DOH soil vapor guidance matrix.

SVOCs including benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, chrysene, dibenzo(a,h)anthracene and ideno(1,2,3-cd)pyrene, were reported above restricted soil cleanup objectives (SCOs) in shallow soil (0-2ft, or 4-6ft).

Metals including lead, chromium, copper, mercury and Zinc were reported above unrestricted SCOs in many of the boring locations. Elevated SVOCs and metals are commonly associated with historic fill and are not believed to be related to historic use of the property.

2.1 **Scope of the Investigation**

The investigation will consist of the following elements:

- Installation of 6 soil borings across the site to delineate the extent of soil impact and to obtain additional information on soil quality with respect to Soil Cleanup Objective (SCOs);
- Installation of 2 additional temporary monitoring wells and collection of groundwater samples from the 3 new and 5 existing monitoring wells to assess CVOC groundwater impacts; and
- Installation of 4 soil gas implants with 2 implants located on the ground level and 2 implants in the cellar to assess vapor phase VOCs.



3.0 SITE HAZARD EVALUATION

This section identifies the hazards associated with the proposed scope of work, general physical hazards that can be expected at most sites; and presents a summary of documented or potential chemical hazards at the site. Every effort must be made to reduce or eliminate these hazards. Those that cannot be eliminated must be guarded against using engineering controls and/or personal protective equipment.

This HASP has been developed for work performed at the site in association with a Phase II subsurface investigation. The primary hazards to the field crew will be physical hazards related to sample collection procedures and equipment, and chemical exposures to the sampling crew from exposure to potential contaminants which may be present at the site.

3.1 Physical Hazards

3.1.1 Tripping Hazards

An area of risk associated with on-site activities are presented by uneven ground, concrete, curbstones or equipment which may be present at the site thereby creating a potential tripping hazard. During intrusive work, care should be taken to mark or remove any obstacles within the exclusion zone

3.1.2 Cuts and Lacerations

Field activities that involve drilling and boring equipment may result in cuts or lacerations from machinery and tools used in collecting samples, cutting disposable tubing and opening acetate sleeves and liners. A first aid kit approved by the American Red Cross will be available during all subsurface investigative activities.

3.1.3 Lifting Hazards

Improper lifting by workers is one of the leading causes of industrial injuries. Field workers and drillers may be required to lift heavy objects such as drilling tools, buckets of decontamination water, cement, etc. Therefore, all members of the field crew should be trained in the proper methods of lifting heavy objects. All workers should be cautioned against lifting objects too heavy for one person.

3.1.4 Utility Hazards

Before conducting any subsurface boring or sampling, the drilling contractor will be responsible for locating and verifying all existing utilities at each excavation.

3.1.5 Traffic Hazards

All traffic, vehicular and pedestrian, shall be maintained and protected at all times consistent with local, state and federal agency regulations regarding such traffic and in accordance with NYCDOT guidelines. The drilling contractor shall carry on his operations without undue interference or delays to traffic. The drilling contractor shall furnish all labor, materials, guards, barricades, signs, lights, and anything else necessary to maintain traffic and to protect his work and the public, during operations.



3.2 Work in Extreme Temperatures

Work under extremely hot or cold weather conditions requires special protocols to minimize the chance that employees will be affected by heat or cold stress.

3.2.1 Heat Stress

The combination of high ambient temperature, high humidity, physical exertion, and personal protective apparel, which limits the dissipation of body heat and moisture, can cause heat stress.

The following prevention, recognition and treatment strategies will be implemented to protect personnel from heat stress. Personnel will be trained to recognize the symptoms of heat stress and to apply the appropriate treatment.

1. Prevention

- a. Provide plenty of fluids. Available in the support zone will be a 50% solution of fruit punch and water or plain water.
- b. Work in Pairs. Individuals should avoid undertaking any activity alone.
- c. Provide cooling devices. A spray hose and a source of water will be provided to reduce body temperature, cool protective clothing and/or act as a quick-drench shower in case of an exposure incident.
- d. Adjustment of the work schedule. As is practical, the most labor-intensive tasks should be carried out during the coolest part of the day.

2. Recognition and Treatment

a. Heat Rash (or prickly heat):

Cause: Continuous exposure to hot and humid air, aggravated by chafing

clothing.

Symptoms: Eruption of red pimples around sweat ducts accompanied by

intense itching and tingling.

Treatment: Remove source or irritation and cool skin with water or wet cloths.

b. Heat Cramps (or heat prostration)

Cause: Profuse perspiration accompanied by inadequate replenishment of

body water and electrolytes.

Symptoms: Muscular weakness, staggering gait, nausea, dizziness, shallow

breathing, pale and clammy skin, approximately normal body

temperature.

Treatment: Perform the following while making arrangement for transport to a

medical facility. Remove the worker to a contamination reduction zone. Remove protective clothing. Lie worker down on back in a cool place and raise feet 6 to 12 inches. Keep warm, but loosen all clothing. If conscious, provide sips of salt-water solution, using one teaspoon of salt in 12 ounces of water. Transport to a medical

facility.

c. Heat Stroke

Cause: Same as heat exhaustion. This is also an extremely serious

condition

Symptoms: Dry and hot skin, dry mouth, dizziness, nausea, headache and rapid

pulse.



Treatment: Cool worker immediately by immersing or spraying with cool

water or sponge bare skin after removing protective clothing.

Transport to hospital.

3.2.2 Cold Exposure

Exposure to cold weather, wet conditions and extreme wind-chill factors may result in excessive loss of body heat (hypothermia) and /or frostbite. To guard against cold exposure and to prevent cold injuries, appropriate warm clothing should be worn, warm shelter must be readily available, rest periods should be adjusted as needed, and the physical conditions of on-site field personnel should be closely monitored. Personnel and supervisors working on-site will be made aware of the signs and symptoms of frost bite and hypothermia such as shivering, reduced blood pressure, reduced coordination, drowsiness, impaired judgment, fatigue, pupils dilated but reactive to light and numbing of the toes and fingers.

3.3 **Chemical Hazards**

Urban fill, present throughout the New York City area, typically contain elevated levels of semivolatile organic compounds and metals. These "contaminants" are not related to a chemical release occurring on the site, but are inherent in the reworked fill material in the area which contains ash bits or tar and asphalt.

Based on the findings of the Phase I and use of adjacent properties, and the inherent properties of urban fill, the following compounds are considered for the site as potential contaminants: volatile organic compounds (VOCs) related to spills of petroleum fuels or industrial solvents, semivolatile organic compounds (SVOCs) related to petroleum fuel spills or inherent in historic fill, pesticides related to historic use of the site and heavy metals such as arsenic, chromium, lead and mercury related to historic fill materials.

The primary routes of exposure to these contaminants are inhalation, ingestion and absorption. **Appendix** C includes information sheets for suspected chemicals that may be encountered at the site.

3.3.1 Respirable Dust and Direct Contact with Soil and Groundwater

Dust may be generated from drilling activities. If visible observation detects elevated levels of dust, a program of wetting will be employed by the site safety officer. If elevated dust levels persist, the site safety office will employ dust monitoring using a particulate monitor (Miniram or equivalent). If monitoring detects concentrations greater than the OSHA action level of 100 μg/m³ over daily background, the site safety officer will take corrective actions as defined herein, including the use of water for dust suppression and if this is not effective, requiring workers to wear APRs with efficiency particulate air (HEPA) cartridges.

Absorption pathways for dust and direct contact with soil and groundwater will be mitigated with the implementation of latex gloves, hand washing and decontamination exercises when necessary.

3.3.2 Organic Vapors

Considering the past and present use of the properties, VOCs may be encountered at the site in soil and/or groundwater. Therefore, soil boring activities may cause the release of organic vapors to the atmosphere. The site safety officer will periodically monitor organic vapors with a Photoionization Detector (PID) during drilling activities to determine whether organic vapor concentrations exceed action levels shown below.

PID Response	Action
Sustained readings of 5 ppm or greater	Shut down equipment and allow area to vent. Resume when readings return to background
Sustained readings of 5 ppm or greater that do not subside after venting	Implement Vapor Release Plan (Section 6.8). Re-evaluate respiratory protection as upgrade may be required.

4.0 PERSONAL PROTECTIVE EQUIPMENT

Personal protective equipment (PPE) shall be selected in accordance with the site air monitoring program, OSHA 29 CFR 1910.120(c), (g), and 1910.132. Protective equipment shall be NIOSH approved and respiratory protection shall conform to OSHA 29 CFR Part 1910.133 and 1910.134 specifications; head protection shall conform to 1910.135; eye and face protection shall conform to 1910.133; and foot protection shall conform to 1910.136. The only true difference among the levels of protection from D thru B is the addition of the type of respiratory protection. It is anticipated that work will be performed in Level D PPE.

4.1 Level D

Level D PPE shall be donned when the atmosphere contains no known hazards and work functions preclude splashes, immersion, or the potential for inhalation of, or contact with, hazardous concentrations of harmful chemicals. Level D PPE consists of:

- standard work uniform, coveralls, or tyvek, as needed;
- steel toe and steel shank work boots:
- high visibility safety vest;
- hard hat;
- gloves, as needed;
- safety glasses;
- hearing protection;
- equipment replacements are available as needed.

4.2 Level C

Level C PPE shall be donned when the concentrations of measured total organic vapors in the breathing zone exceed background concentrations (using a portable OVA, or equivalent), but are less than 5 ppm. The specifications on the APR filters used must be appropriate for contaminants identified or expected to be encountered. Level C PPE shall be donned when the identified contaminants have adequate warning properties and criteria for using APR have been met. Level C PPE consists of:

- chemical resistant or coated tyvek coveralls;
- steel-toe and steel-shank workboots;
- high visibility safety vest;
- chemical resistant overboots or disposable boot covers;
- disposable inner gloves (surgical gloves);
- disposable outer gloves;
- full face APR fitted with organic vapor/dust and mist filters or filters appropriate for the identified or expected contaminants;
- hard hat;
- splash shield, as needed; and,
- ankles/wrists taped with duct tape.

The site safety officer will verify if Level C is appropriate by checking organic vapor concentrations using compound and/or class-specific detector tubes.

The exact PPE ensemble is decided on a site-by-site basis by the Site Safety Officer with the intent to provide the most protective and efficient worker PPE.

4.3 **Activity-Specific Levels of Personal Protection**

The required level of PPE is activity-specific and is based on air monitoring results (Section 4.0) and properties of identified or expected contaminants. It is expected that site work will be **performed in Level D.** If air monitoring results indicate the necessity to upgrade (i.e dust above 5,000 µg/m³ or sustained VOCs above 5 ppm in the breathing zone) the level of protection engineering controls (i.e. Facing equipment away from the wind and placing site personnel upwind of excavations, active venting, etc.) will be implemented before requiring the use of respiratory protection.



5.0 SITE CONTROL

5.1 Work Zones

The primary purpose of site controls is to establish the perimeter of a hazardous area, to reduce the migration of contaminants into clean areas, and to prevent access or exposure to hazardous materials by unauthorized persons. When operations are to take place involving hazardous materials, the site safety officer will establish an exclusion zone, a decontamination zone, and a support zone. These zones "float" (move around the site) depending on the tasks being performed on any given day. The site safety officer will outline these locations before work begins and when zones change. The site safety officer records this information in the site log book. It is expected that for soil boring and sampling activities, identification of an exclusion zone, decontamination zone, and support zone will not be necessary.

Tasks requiring OSHA 40-hour Hazardous Waste Operations and Emergency Response Operations training are carried out in the exclusion zone. The exclusion zone is defined by the site safety officer but will typically be a 50-foot area around work activities. Gross decontamination (as determined by the site Health and Safety Officer) is conducted in the exclusion zone; all other decontamination is performed in the decontamination zone or trailer.

Protective equipment is removed in the decontamination zone. Disposable protective equipment is stored in receptacles staged in the decontamination zone, and non-disposable equipment is decontaminated. All personnel and equipment exit the exclusion zone through the decontamination zone. If a decontamination trailer is provided the first aid equipment, an eye wash unit, and drinking water are kept in the decontamination trailer.

The support zone is used for vehicle parking, daily safety meetings, and supply storage. Eating, drinking, and smoking are permitted only in the support zone. When a decontamination trailer is not provided, the eye wash unit, first aid equipment, and drinking water are kept at a central location designated by the site safety officer.

6.0 CONTINGENCY PLAN/EMERGENCY RESPONSE PLAN

Site personnel must be prepared in the event of an emergency. Emergencies can take many forms: illnesses, injuries, chemical exposure, fires, explosions, spills, leaks, releases of harmful contaminants, or sudden changes in the weather.

Emergency telephone numbers and a map to the hospital will be posted in the command post. Site personnel should be familiar with the emergency procedures, and the locations of site safety, first aid, and communication equipment.

6.1 **Emergency Equipment On-site**

Private telephones: Site personnel.

Site personnel where necessary. Two-way radios:

Emergency Alarms: On-site vehicle horns*. First aid kits: On-site, in vehicles or office.

Fire extinguisher: On-site, in office or on equipment.

6.2 **Emergency Telephone Numbers**

General Emergencies	911
New York City Police	911
Harlem Hospital Center	1-212-939-1000
NYSDEC Spills Division	1-800-457-7362
NYSDEC Division of Env. Remediation	1-518-402-9768
NYCDEP	1-718-699-9811
NYC Department of Health	1-212-788-4711
NYC Fire Department	911
National Response Center	1-800-424-8802
Poison Control	1-212-340-4494
Site Safety Officer	1-631-504-6000
Alternate Site Safety Officer	1-631-504-6000

6.3 Personnel Responsibilities During an Emergency

The project manager is primarily responsible for responding to and correcting any emergency situations. However, in the absence of the project manager, the site safety officer shall act as the project manager's on-site designee and perform the following tasks:

- Take appropriate measures to protect personnel including: withdrawal from the exclusion zone, evacuate and secure the site, or upgrade/downgrade the level of protective clothing and respiratory protection;
- Ensure that appropriate federal, state, and local agencies are informed and emergency response plans are coordinated. In the event of fire or explosion, the local fire department

^{*} Horns: Air horns will be supplied to personnel at the discretion of the project superintendent or site safety officer.

should be summoned immediately. If toxic materials are released to the air, the local authorities should be informed in order to assess the need for evacuation;

- Ensure appropriate decontamination, treatment, or testing for exposed or injured personnel;
- Determine the cause of incidents and make recommendations to prevent recurrence; and,
- Ensure that all required reports have been prepared.

The following key personnel are planned for this project:

Project Manager
Site Safety Officer
Alternate
Ms. Chawinie Miller (631) 504-6000
Mr. Kevin Waters (631) 504-6000
Mr. Dominic Mosca (631) 504-6000

6.4 Medical Emergencies

A person who becomes ill or injured in the exclusion zone will be decontaminated to the maximum extent possible. If the injury or illness is minor, full decontamination will be completed and first aid administered prior to transport. First aid will be administered while waiting for an ambulance or paramedics. A Field Accident Report (**Appendix D**) must be filled out for any injury.

A person transporting an injured/exposed person to a clinic or hospital for treatment will take the directions to the hospital (**Appendix D**) and information on the chemical(s) to which they may have been exposed (**Appendix C**).

6.5 Fire or Explosion

In the event of a fire or explosion, the local fire department will be summoned immediately. The site safety officer or his designated alternate will advise the fire commander of the location, nature and identification of the hazardous materials on-site. If it is safe to do so, site personnel may:

- use fire fighting equipment available on site; or,
- remove or isolate flammable or other hazardous materials that may contribute to the fire.

6.6 Evacuation Routes

Evacuation routes established by work area locations for each site will be reviewed prior to commencing site operations. As the work areas change, the evacuation routes will be altered accordingly, and the new route will be reviewed.

Under extreme emergency conditions, evacuation is to be immediate without regard for equipment. The evacuation signal will be a continuous blast of a vehicle horn, if possible, and/or by verbal/radio communication. When evacuating the site, personnel will follow these instructions:

- Keep upwind of smoke, vapors, or spill location.
- Exit through the decontamination corridor if possible.
- If evacuation through the decontamination corridor is not possible, personnel should remove contaminated clothing once they are in a safe location and leave it near the exclusion zone or in a safe place.
- The site safety officer will conduct a head count to ensure that all personnel have been evacuated safely. The head count will be correlated to the site and/or exclusion zone entry/exit log.
- If emergency site evacuation is necessary, all personnel are to escape the emergency situation and decontaminate to the maximum extent practical.

6.7 Spill Control Procedures

Spills associated with site activities may be attributed to project equipment and include gasoline, diesel and hydraulic oil. In the event of a leak or a release, site personnel will inform their supervisor immediately, locate the source of spillage and stop the flow if it can be done safely. A spill containment kit including absorbent pads, booms and/or granulated speedy dry absorbent material will be available to site personnel to facilitate the immediate recovery of the spilled material. Daily inspections of site equipment components including hydraulic lines, fuel tanks, etc. will be performed by their respective operators as a preventative measure for equipment leaks and to ensure equipment soundness. In the event of a spill, site personnel will immediately notify the NYSDEC (1-800-457-7362), and a spill number will be generated.

6.8 Vapor Release Plan

If work zone organic vapor (excluding methane) exceeds 5 ppm, then a downwind reading will be made either 200 feet from the work zone or at the property line, whichever is closer. If readings at this location exceed 5 ppm over background, the work will be stopped.

If 5 ppm of VOCs are recorded over background on a PID at the property line, then an off-site reading will be taken within 20 feet of the nearest residential or commercial property, whichever is closer. If efforts to mitigate the emission source are unsuccessful for 30 minutes, then the designated site safety officer will:

- contact the local police;
- continue to monitor air every 30 minutes, 20 feet from the closest off-site property. If two successive readings are below 5 ppm (non-methane), off-site air monitoring will be halted.
- All property line and off site air monitoring locations and results associated with vapor releases will be recorded in the site safety log book.



APPENDIX A SITE SAFETY ACKNOWLEDGEMENT FORM

DAILY BREIFING SIGN-IN SHEET

ite: Person Conducting Briefing:	
Project Name and Location:	
AWARENESS (topics discussed, special safety	concerns, recent incidents, etc):
2. OTHER ISSUES (HASP changes, attendee comm	nents, etc):
3. ATTENDEES (Print Name):	
1.	11.
2.	12.
3.	13.
4.	14.
5.	15.
6.	16.
7.	17.
8.	18.
9.	19.
10.	20.

APPENDIX B SITE SAFETY PLAN AMENDMENTS

SITE SAFETY PLAN AMENDMENT FORM

Site Safety Plan Amendment #:		
Site Name:		
Reason for Amendment:		
Alternative Procedures:		
Required Changes in PPE:		
Project Superintendent (signature)	Date	
Health and Safety Consultant (signature)	 Date	
Health and Safety Consultant (Signature)	Date	
Site Safety Officer (signature)	Date	

APPENDIX C CHEMICAL HAZARDS

CHEMICAL HAZARDS

The attached International Chemical Safety Cards are provided for contaminants of concern that have been identified in soils and/or groundwater at the site.

Material Safety Data Sheet

cis-1,2-Dichloroethylene, 97%

ACC# 97773

Section 1 - Chemical Product and Company Identification

MSDS Name: cis-1,2-Dichloroethylene, 97%

Catalog Numbers: AC113380000, AC113380025, AC113380100

Synonyms: cis-Acetylene dichloride.

Company Identification: Acros Organics N.V. One Reagent Lane Fair Lawn, NJ 07410

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

Section 2 - Composition, Information on Ingredients

CAS#	Chemical Name	Percent	EINECS/ELINCS
156-59-2	cis-1,2-Dichloroethylene	97	205-859-7

Section 3 - Hazards Identification

EMERGENCY OVERVIEW

Appearance: Clear liquid. Flash Point: 6 deg C.

Warning! Flammable liquid and vapor. Harmful if inhaled. Unstabilized substance may polymerize. Causes eye and skin irritation. May be harmful if swallowed. May cause respiratory tract irritation.

Target Organs: Central nervous system, respiratory system, eyes, skin.

Potential Health Effects

Eye: Causes moderate eye irritation.

Skin: Causes moderate skin irritation. May cause dermatitis.

Ingestion: May cause gastrointestinal irritation with nausea, vomiting and diarrhea. May be harmful if

swallowed. May cause central nervous system depression.

Inhalation: May cause respiratory tract irritation. May cause narcotic effects in high concentration. Eye irritation, vertigo, and nausea were reported in humans exposed at 2200 ppm.

Chronic: Not available. Some German investigators reported fatty degeneration of the liver upon repeated

narcotic doses in rats and

Section 4 - First Aid Measures

Eyes: In case of contact, immediately flush eyes with plenty of water for a t least 15 minutes. Get medical aid. Skin: In case of contact, flush skin with plenty of water. Remove contaminated clothing and shoes. Get medical aid if irritation develops and persists. Wash clothing before reuse.

Ingestion: If swallowed, do not induce vomiting unless directed to do so by medical personnel. Never give anything by mouth to an unconscious person. Get medical aid.

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

Notes to Physician: Treat symptomatically and supportively.

Section 5 - Fire Fighting Measures

General Information: As in any fire, wear a self-contained breathing apparatus in pressure-demand, MSHA/NIOSH (approved or equivalent), and full protective gear. Vapors may form an explosive mixture with air. Use water spray to keep fire-exposed containers cool. Flammable liquid and vapor. Fire or excessive heat may result in violent rupture of the container due to bulk polymerization. Vapors are heavier than air and may travel to a source of ignition and flash back. Vapors can spread along the ground and collect in low or confined areas. Hazardous polymerization may occur under fire conditions.

Extinguishing Media: Use water fog, dry chemical, carbon dioxide, or regular foam.

Flash Point: 6 deg C (42.80 deg F)

Autoignition Temperature: 440 deg C (824.00 deg F)

Explosion Limits, Lower: 9.70 vol %

Upper: 12.80 vol %

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

Section 6 - Accidental Release Measures

General Information: Use proper personal protective equipment as indicated in Section 8.

Spills/Leaks: Absorb spill with inert material (e.g. vermiculite, sand or earth), then place in suitable container. Remove all sources of ignition. Use a spark-proof tool. Provide ventilation.

Section 7 - Handling and Storage

Handling: Wash thoroughly after handling. Remove contaminated clothing and wash before reuse. Ground and bond containers when transferring material. Use spark-proof tools and explosion proof equipment. Avoid contact with eyes, skin, and clothing. Empty containers retain product residue, (liquid and/or vapor), and can be dangerous. Avoid ingestion and inhalation. Do not pressurize, cut, weld, braze, solder, drill, grind, or expose empty containers to heat, sparks or open flames. Use only with adequate ventilation. Pure vapor will be uninhibited and may polymerize in vents or other confined spaces.

Storage: Keep away from sources of ignition. Store in a tightly closed container. Flammables-area. Store protected from light and air.

Section 8 - Exposure Controls, Personal Protection

Engineering Controls: Use process enclosure, local exhaust ventilation, or other engineering controls to control airborne levels below recommended exposure limits. Facilities storing or utilizing this material should be equipped with an eyewash facility and a safety shower.

Exposure Limits

Chemical Name	ACGIH	NIOSH	OSHA - Final PELs
cis-1,2-Dichloroethylene	200 ppm TWA	none listed	none listed

OSHA Vacated PELs: cis-1,2-Dichloroethylene: No OSHA Vacated PELs are listed for this chemical.

Personal Protective Equipment

Eyes: Wear chemical splash goggles.

Skin: Wear appropriate protective gloves to prevent skin exposure.

Clothing: Wear appropriate protective clothing to prevent skin exposure.

Respirators: Follow the OSHA respirator regulations found in 29 CFR 1910.134 or European Standard EN 149. Use a NIOSH/MSHA or European Standard EN 149 approved respirator if exposure limits are exceeded or if irritation or other symptoms are experienced.

Section 9 - Physical and Chemical Properties

Physical State: Liquid Appearance: Clear Odor: Pleasant odor pH: Not available.

Vapor Pressure: 201 mm Hg @ 25 deg C

Vapor Density: 3.34 (air=1) Evaporation Rate:Not available.

Viscosity: Not available.

Boiling Point: 60 deg C @ 760 mm Hg **Freezing/Melting Point**:-80 deg C

Decomposition Temperature: Not available.

Solubility: Insoluble.

Specific Gravity/Density:1.2800 Molecular Formula:C2H2Cl2 Molecular Weight:96.94

Section 10 - Stability and Reactivity

Chemical Stability: Stable under normal temperatures and pressures. This material is a monomer and may polymerize under certain conditions if the stabilizer is lost.

Conditions to Avoid: Light, ignition sources, exposure to air, excess heat.

Incompatibilities with Other Materials: Strong oxidizing agents, strong bases, copper.

Hazardous Decomposition Products: Hydrogen chloride, phosgene, carbon monoxide, carbon dioxide.

Hazardous Polymerization: May occur.

Section 11 - Toxicological Information

RTECS#:

CAS# 156-59-2: KV9420000

LD50/LC50: CAS# 156-59-2:

Inhalation, rat: LC50 = 13700 ppm;

Carcinogenicity:

CAS# 156-59-2: Not listed by ACGIH, IARC, NTP, or CA Prop 65.

Epidemiology: No data available. **Teratogenicity:** No data available.

Reproductive Effects: No data available.

Mutagenicity: No data available. **Neurotoxicity:** No data available.

Other Studies:

Section 12 - Ecological Information

No information available.

Section 13 - Disposal Considerations

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

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

Section 14 - Transport Information

	US DOT	Canada TDG
Shipping Name:	DOT regulated - small quantity provisions apply (see 49CFR173.4)	1,2-DICHLOROETHYLENE
Hazard Class:		3
UN Number:		UN1150
Packing Group:		II

Section 15 - Regulatory Information

US FEDERAL

TSCA

CAS# 156-59-2 is listed on the TSCA inventory.

Health & Safety Reporting List

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

Chemical Test Rules

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

Section 12b

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

TSCA Significant New Use Rule

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

CERCLA Hazardous Substances and corresponding RQs

None of the chemicals in this material have an RQ.

SARA Section 302 Extremely Hazardous Substances

None of the chemicals in this product have a TPQ.

Section 313 No chemicals are reportable under Section 313.

Clean Air Act:

This material does not contain any hazardous air pollutants.

This material does not contain any Class 1 Ozone depletors.

This material does not contain any Class 2 Ozone depletors.

Clean Water Act:

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

None of the chemicals in this product are listed as Priority Pollutants under the CWA.

None of the chemicals in this product are listed as Toxic Pollutants under the CWA.

OSHA:

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

STATE

CAS# 156-59-2 can be found on the following state right to know lists: Pennsylvania, Massachusetts.

California Prop 65

California No Significant Risk Level: None of the chemicals in this product are listed.

European/International Regulations

European Labeling in Accordance with EC Directives

Hazard Symbols:

XN F

Risk Phrases:

R 11 Highly flammable.

R 20 Harmful by inhalation.

R 52/53 Harmful to aquatic organisms, may cause long-term adverse effects in the aquatic environment.

Safety Phrases:

S 16 Keep away from sources of ignition - No smoking.

S 29 Do not empty into drains.

S 7 Keep container tightly closed.

S 61 Avoid release to the environment. Refer to special instructions /safety data sheets.

WGK (Water Danger/Protection)

CAS# 156-59-2: No information available.

Canada - DSL/NDSL

CAS# 156-59-2 is listed on Canada's NDSL List.

Canada - WHMIS

WHMIS: Not available.

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

Canadian Ingredient Disclosure List

Section 16 - Additional Information

MSDS Creation Date: 2/09/1998 Revision #5 Date: 3/16/2007

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

TRICHLOROETHYLENE











1,1,2-Trichloroethylene Trichloroethene Ethylene trichloride Acetylene trichloride C₂HCl₃ / ClCH=CCl₂ Molecular mass: 131,4

ICSC # 0081 CAS # 79-01-6 RTECS # <u>KX4550000</u>

UN # 1710

EC # 602-027-00-9 April 10, 2000 Validated







ICSC: 0081

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS/ SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Combustible under specific conditions. See Notes.		In case of fire in the surroundings: all extinguishing agents allowed.
EXPLOSION		Prevent build-up of electrostatic charges (e.g., by grounding).	In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE		PREVENT GENERATION OF MISTS! STRICT HYGIENE!	
•INHALATION	Dizziness. Drowsiness. Headache. Weakness. Nausea. Unconsciousness.	Ventilation, local exhaust, or breathing protection.	Fresh air, rest. Artificial respiration may be needed. Refer for medical attention.
•SKIN	Dry skin. Redness.	Protective gloves.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain.	Safety spectacles, or eye protection in combination with breathing protection.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Abdominal pain. (Further see Inhalation).	Do not eat, drink, or smoke during work.	Rinse mouth. Do NOT induce vomiting. Give one or two glasses of water to drink. Rest.

SPILLAGE DISPOSAL **STORAGE** PACKAGING & LABELLING Do not transport with food and feedstuffs. Ventilation. Personal protection: filter Separated from metals (see Chemical respirator for organic gases and vapours Dangers), strong bases, food and feedstuffs . Marine pollutant. adapted to the airborne concentration of the Dry. Keep in the dark. Ventilation along the T symbol R: 45-36/38-52/53-67 substance. Collect leaking and spilled liquid floor. Store in an area without drain or sewer in sealable containers as far as possible. access. S: 53-45-61 Absorb remaining liquid in sand or inert UN Hazard Class: 6.1 absorbent and remove to safe place. Do NOT UN Packing Group: III let this chemical enter the environment.

SEE IMPORTANT INFORMATION ON BACK

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the

ICSC: 0081

OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

TRICHLOROETHYLENE

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID , WITH CHARACTERISTIC ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation and by ingestion.				
M P	PHYSICAL DANGERS: The vapour is heavier than air. As a result of flow, agitation, etc., electrostatic charges can be generated.	INHALATION RISK: A harmful contamination of the air can be reached rather quickly on evaporation of this substance at 20°C.				
0	CHEMICAL DANGERS: On contact with hot surfaces or flames this substance	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes and the skin.				
R	decomposes forming toxic and corrosive fumes (phosgene, hydrogen chloride). The substance	Swallowing the liquid may cause aspiration into the lungs with the risk of chemical pneumonitis. The				
Т	decomposes on contact with strong alkali producing dichloroacetylene, which increases fire hazard. Reacts	substance may cause effects on the central nervous system, resulting in respiratory failure. Exposure could				
A	violently with metal powders such as magnesium, aluminium, titanium, and barium. Slowly decomposed	cause lowering of consciousness.				
N	by light in presence of moisture, with formation of corrosive hydrochloric acid.	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE:				
Т	OCCUPATIONAL EXPOSURE LIMITS: TLV: 50 ppm as TWA; 100 ppm as STEL; A5; BEI	Repeated or prolonged contact with skin may cause dermatitis. The substance may have effects on the central nervous system, resulting in loss of memory.				
D	issued; (ACGIH 2004). MAK:	The substance may have effects on the liver and kidneys (see Notes). This substance is probably carcinogenic to				
A	Carcinogen category: 1; Germ cell mutagen group: 3B; (DFG 2007).	humans.				
Т	OSHA PEL±: TWA 100 ppm C 200 ppm 300 ppm (5-minute maximum peak in any 2 hours)					
A	NIOSH REL: Ca See Appendix A See Appendix C NIOSH IDLH: Ca 1000 ppm See: 79016					
PHYSICAL PROPERTIES	Boiling point: 87°C Melting point: -73°C Relative density (water = 1): 1.5 Solubility in water, g/100 ml at 20°C: 0.1 Vapour pressure, kPa at 20°C: 7.8 Relative vapour density (air = 1): 4.5	Relative density of the vapour/air-mixture at 20°C (air = 1): 1.3 Auto-ignition temperature: 410°C Explosive limits, vol% in air: 8-10.5 Octanol/water partition coefficient as log Pow: 2.42 Electrical conductivity: 800pS/m				
ENVIRONMENTAL	The substance is harmful to aquatic organisms. The substance environment	ance may cause long-term effects in the				

DATA

aquatic environment.



ICSC: 0081

NOTES

Combustible vapour/air mixtures difficult to ignite, may be developed under certain conditions. Use of alcoholic beverages enhances the harmful effect. Depending on the degree of exposure, periodic medical examination is suggested. The odour warning when the exposure limit value is exceeded is insufficient. Do NOT use in the vicinity of a fire or a hot surface, or during welding. An added stabilizer or inhibitor can influence the toxicological properties of this substance, consult an expert.

Transport Emergency Card: TEC (R)-61S1710

NFPA Code: H2; F1; R0;

Card has been partially updated in October 2004: see Occupational Exposure Limits, EU Classification, Emergency Response. Card has been partially updated in April 2010: see Occupational Exposure Limits, Ingestion First Aid, Storage.

ICSC: 0081 TRICHLOROETHYLENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

TETRACHLOROETHYLENE











1,1,2,2-Tetrachloroethylene Perchloroethylene Tetrachloroethene C_2Cl_4 / $Cl_2C=CCl_2$ Molecular mass: 165.8

ICSC # 0076 CAS # 127-18-4 RTECS # <u>KX3850000</u>

UN # 1897

EC # 602-028-00-4 April 13, 2000 Validated







ICSC: 0076

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS/ SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Not combustible. Gives off irritating or toxic fumes (or gases) in a fire.		In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION			
EXPOSURE		STRICT HYGIENE! PREVENT GENERATION OF MISTS!	
•INHALATION	Dizziness. Drowsiness. Headache. Nausea. Weakness. Unconsciousness.	Ventilation, local exhaust, or breathing protection.	Fresh air, rest. Artificial respiration may be needed. Refer for medical attention.
•SKIN	Dry skin. Redness.	Protective gloves. Protective clothing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain.	Safety goggles, face shield.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Abdominal pain. (Further see Inhalation).	Do not eat, drink, or smoke during work.	Rinse mouth. Do NOT induce vomiting. Give plenty of water to drink. Rest.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
	Dangers), food and feedstuffs . Keep in the dark. Ventilation along the floor.	Do not transport with food and feedstuffs. Marine pollutant. Xn symbol N symbol R: 40-51/53 S: (2-)23-36/37-61 UN Hazard Class: 6.1 UN Packing Group: III

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0076

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

TETRACHLOROETHYLENE

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID , WITH CHARACTERISTIC ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation and by ingestion.				
M	PHYSICAL DANGERS:	INHALATION RISK:				
P	The vapour is heavier than air.	A harmful contamination of the air will be reached rather slowly on evaporation of this substance at 20°C.				
О	CHEMICAL DANGERS: On contact with hot surfaces or flames this substance	EFFECTS OF SHORT-TERM EXPOSURE:				
R	decomposes forming toxic and corrosive fumes (hydrogen chloride, phosgene, chlorine). The substance	The substance is irritating to the eyes, the skin and the respiratory tract. If this liquid is swallowed, aspiration				
Т	decomposes slowly on contact with moisture producing trichloroacetic acid and hydrochloric acid. Reacts with	into the lungs may result in chemical pneumonitis. The substance may cause effects on the central nervous				
A	metals such as aluminium, lithium, barium, beryllium.	system. Exposure at high levels may result in unconsciousness.				
N	OCCUPATIONAL EXPOSURE LIMITS: TLV: 25 ppm as TWA, 100 ppm as STEL; A3	EFFECTS OF LONG-TERM OR REPEATED				
Т	(confirmed animal carcinogen with unknown relevance to humans); BEI issued; (ACGIH 2004).	EXPOSURE: Repeated or prolonged contact with skin may cause				
D	MAK: skin absorption (H); Carcinogen category: 3B;	dermatitis. The substance may have effects on the liver and kidneys. This substance is probably carcinogenic to				
A	(DFG 2004). OSHA PEL <u>†</u> : TWA 100 ppm C 200 ppm 300 ppm (5-	humans.				
T	minute maximum peak in any 3-hours) NIOSH REL: Ca Minimize workplace exposure					
A	concentrations. See Appendix A NIOSH IDLH: Ca 150 ppm See: 127184					
	Boiling point: 121°C	Vapour pressure, kPa at 20°C: 1.9				
PHYSICAL	Melting point: -22°C	Relative vapour density (air = 1): 5.8				
PROPERTIES	Relative density (water = 1): 1.6 Solubility in water, g/100 ml at 20°C: 0.015	Relative density of the vapour/air-mixture at 20°C (air = 1): 1.09				
		Octanol/water partition coefficient as log Pow: 2.9				
ENVIRONMENTAL DATA	The substance is toxic to aquatic organisms. The substance environment.	te may cause long-term effects in the aquatic				
NOTES						
Depending on the degree of exposure, periodic medical examination is suggested. The odour warning when the exposure limit value is exceeded is insufficient. Do NOT use in the vicinity of a fire or a hot surface, or during welding. An added stabilizer or inhibitor can influence the toxicological properties of this substance, consult an expert. Card has been partly updated in April 2005. See section Occupational Exposure Limits.						
	Transport Emergency Card: TEC (R)-61S1897					
		NFPA Code: H2; F0; R0;				

ADDITIONAL INFORMATION

ICSC: 0076 TETRACHLOROETHYLENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE: Neither NIOSH, the CEC or the IPCS nor any person acting on behalf of NIOSH, the CEC or the IPCS is responsible for the use which might be made of this information. This card contains the collective views of the IPCS Peer Review Committee and may not reflect in all cases all the detailed requirements included in national legislation on the subject. The user should verify compliance of the cards with the relevant legislation in the country of use. The only

ICSC: 0076

ICSC:NENG0076 International Chemical Safety Cards (WHO/IPCS/ILO) | CDC/NIOSH

modifications made to produce the U.S. version is inclusion of the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

TOLUENE ICSC: 0078











 $\begin{array}{c} \text{Methylbenzene} \\ \text{Toluol} \\ \text{Phenylmethane} \\ \text{C}_6\text{H}_5\text{CH}_3 \, / \, \text{C}_7\text{H}_8 \end{array}$

Molecular mass: 92.1

ICSC # 0078 CAS # 108-88-3 RTECS # <u>XS5250000</u>

UN # 1294

EC # 601-021-00-3

October 10, 2002 Peer reviewed

Personal protection: self-contained breathing

apparatus



	34334 15, 2332 1011 1011 1011					
TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ SYMPTO		PREVENTION		FIRST AID/ FIRE FIGHTING	
FIRE	Highly flammable.		NO open flames, NO sparks, ar smoking.	nd NO	Powder, AFFF, foam, carbon dioxide.	
EXPLOSION	Vapour/air mixtures are explosive.		Closed system, ventilation, explosion-proof electrical equipment and lighting. Prevent build-up of electrostatic charges (e.g., by grounding). Do NOT use compressed air for filling, discharging, or handling. Use non-sparking handtools.		In case of fire: keep drums, etc., cool by spraying with water.	
EXPOSURE			STRICT HYGIENE! AVOID EXPOSURE OF (PREGNANT WOMEN!	")		
•INHALATION	Cough. Sore throat. Dizziness. Drowsiness. Headache. Nausea. Unconsciousness.		Ventilation, local exhaust, or breathing protection.		Fresh air, rest. Refer for medical attention.	
•SKIN	Dry skin. Redness.		Protective gloves.		Remove contaminated clothes. Rinse and then wash skin with water and soap. Refer for medical attention.	
•EYES	Redness. Pain.		Safety goggles.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.	
•INGESTION	Burning sensation. Abd (Further see Inhalation)			Rinse mouth. Do NOT induce vomiting. Refer for medical attention.		
SPILLAGE DISPOSAL		STORAGE PA		CKAGING & LABELLING		
Evacuate danger area in large spill! Consult an expert in large spill! Remove all ignition sources. Ventilation. Collect leaking liquid in sealable containers. Absorb remaining liquid in sand or inert absorbent and remove to safe place. Do NOT wash away into sewer. Do NOT let this chemical enter the environment.		Fireproof. Sep	parated from strong oxidants.	S: 2-36 UN Ha		

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0078

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

TOLUENE ICSC: 0078

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID, WITH CHARACTERISTIC	ROUTES OF EXPOSURE: The substance can be absorbed into the body by
M	ODOUR.	inhalation, through the skin and by ingestion.
P	PHYSICAL DANGERS: The vapour mixes well with air, explosive mixtures are	INHALATION RISK: A harmful contamination of the air can be reached rather
О	formed easily. As a result of flow, agitation, etc., electrostatic charges can be generated.	quickly on evaporation of this substance at 20°C.
R	CHEMICAL DANGERS:	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes and the respiratory
Т	Reacts violently with strong oxidants causing fire and explosion hazard.	tract The substance may cause effects on the central nervous system If this liquid is swallowed, aspiration
A	OCCUPATIONAL EXPOSURE LIMITS:	into the lungs may result in chemical pneumonitis. Exposure at high levels may result in cardiac
N	TLV: 50 ppm as TWA (skin) A4 BEI issued (ACGIH 2004).	dysrhythmiaandunconsciousness.
Т	MAK: 50 ppm 190 mg/m³ H Peak limitation category: II(4) Pregnancy risk group: C (DFG 2004).	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE: The liquid defats the skin. The substance may have
D	OSHA PEL±: TWA 200 ppm C 300 ppm 500 ppm (10-minute maximum peak)	effects on the central nervous system Exposure to the substance may enhance hearing damage caused by
A	NIOSH REL: TWA 100 ppm (375 mg/m ³) ST 150 ppm (560 mg/m ³)	exposure to noise. Animal tests show that this substance possibly causes toxicity to human reproduction or
T	NIOSH IDLH: 500 ppm See: <u>108883</u>	development.
A		
PHYSICAL PROPERTIES	Boiling point: 111°C Melting point: -95°C Relative density (water = 1): 0.87 Solubility in water: none	Relative density of the vapour/air-mixture at 20°C (air = 1): 1.01 Flash point: 4°C c.c. Auto-ignition temperature: 480°C
INOLEKTIED	Vapour pressure, kPa at 25°C: 3.8 Relative vapour density (air = 1): 3.1	Explosive limits, vol% in air: 1.1-7.1 Octanol/water partition coefficient as log Pow: 2.69
ENVIRONMENTAL DATA	The substance is toxic to aquatic organisms.	

NOTES

Depending on the degree of exposure, periodic medical examination is suggested. Use of alcoholic beverages enhances the harmful effect.

Transport Emergency Card: TEC (R)-30S1294

NFPA Code: H 2; F 3; R 0;

ADDITIONAL INFORMATION

ICSC: 0078 TOLUENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

m-XYLENE ICSC: 0085











meta-Xylene 1,3-Dimethylbenzene m-Xylol $C_6H_4(CH_3)_2/C_8H_{10}$ Molecular mass: 106.2

ICSC # 0085 CAS # 108-38-3 RTECS # <u>ZE2275000</u> UN # 1307

EC # 601-022-00-9 August 03, 2002 Validated



TYPES OF HAZARD/ EXPOSURE	ACUTE HAZA SYMPTOM		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Flammable.		NO open flames, NO sparks, and smoking.		Powder, water spray, foam, carbon dioxide.
EXPLOSION	Above 27°C explosive va mixtures may be formed.		Above 27°C use a closed system, ventilation, and explosion-proof electrical equipment. Prevent bui of electrostatic charges (e.g., by grounding).		In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE			STRICT HYGIENE!		
•INHALATION	Dizziness. Drowsiness. H Nausea.	leadache.	Ventilation, local exhaust, or breathing protection.		Fresh air, rest. Refer for medical attention.
•SKIN	Dry skin. Redness.		Protective gloves.		Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain.		Safety spectacles.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Burning sensation. Abdor (Further see Inhalation).	minal pain.	Do not eat, drink, or smoke durin work.	0	Rinse mouth. Do NOT induce vomiting. Refer for medical attention.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
	Fireproof. Separated from strong oxidants	N. C
	6	Note: C
containers as far as possible. Absorb		Xn symbol
remaining liquid in sand or inert absorbent		R: 10-20/21-38
and remove to safe place. Do NOT let this		S: 2-25
chemical enter the environment. (Extra		UN Hazard Class: 3
personal protection: filter respirator for		UN Packing Group: III
organic gases and vapours.)		

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0085

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

m-XYLENE ICSC: 0085

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID, WITH CHARACTERISTI ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation, through the skin and by ingestion.			
M		, 8			
P	PHYSICAL DANGERS: As a result of flow, agitation, etc., electrostatic charge can be generated.	INHALATION RISK: A harmful contamination of the air will be reached rather slowly on evaporation of this substance at 20°C.			
0		, 1			
R	CHEMICAL DANGERS: Reacts with strong acids strong oxidants	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes and the skin The substance may cause effects on the central nervous			
T	OCCUPATIONAL EXPOSURE LIMITS: TLV: 100 ppm as TWA 150 ppm as STEL A4 (ACGI	system If this liquid is swallowed, aspiration into the			
A	2001). BEI (ACGIH 2001). MAK: 100 ppm 440 mg/m ³	EFFECTS OF LONG-TERM OR REPEATED			
N	Peak limitation category: II(2)	EXPOSURE:			
Т	skin absorption (H); Pregnancy risk group: D (DFG 2005).	The liquid defats the skin. The substance may have effects on the central nervous system Animal tests show that this substance possibly causes toxicity to human			
D	EU OEL: 50 ppm as TWA 100 ppm as STEL (skin) (1 2000).	EU reproduction or development.			
A	OSHA PEL±: TWA 100 ppm (435 mg/m³) NIOSH REL: TWA 100 ppm (435 mg/m³) ST 150 pp	m			
Т	(655 mg/m ³) NIOSH IDLH: 900 ppm See: <u>95476</u>				
A					
PHYSICAL PROPERTIES	Boiling point: 139°C Melting point: -48°C Relative density (water = 1): 0.86 Solubility in water: none Vapour pressure, kPa at 20°C: 0.8	Relative vapour density (air = 1): 3.7 Relative density of the vapour/air-mixture at 20°C (air = 1): 1.02 Flash point: 27°C c.c. Auto-ignition temperature: 527°C Explosive limits, vol% in air: 1.1-7.0 Octanol/water partition coefficient as log Pow: 3.20			
ENVIRONMENTAL DATA	The substance is toxic to aquatic organisms.				
NOTES					
	Depending on the degree of exposure, periodic medical examination is indicated. The recommendations on this Card also apply to technical xylene. See ICSC 0084 o-Xylene and 0086 p-Xylene.				
	NFPA Code: H 2; F 3; R 0; Transport Emergency Card: TEC (R)-30S1307-III				
ADDITIONAL INFORMATION					

ICSC: 0085 m-XYLENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

O-XYLENE ICSC: 0084











ortho-Xylene 1,2-Dimethylbenzene o-Xylol $C_6H_4(CH_3)_2/C_8H_{10}$ Molecular mass: 106.2

ICSC # 0084 CAS # 95-47-6 RTECS # <u>ZE2450000</u> UN # 1307

EC # 601-022-00-9 August 03, 2002 Validated

remaining liquid in sand or inert absorbent

and remove to safe place. Do NOT let this

chemical enter the environment. (Extra



TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ SYMPTO		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Flammable.		NO open flames, NO sparks, ar smoking.	nd NO	Powder, water spray, foam, carbon dioxide.
EXPLOSION	Above 32°C explosive vapour/air mixtures may be formed. Above 32°C use a closed system, ventilation, and explosion-proof		f uild-up	In case of fire: keep drums, etc., cool by spraying with water.	
EXPOSURE			STRICT HYGIENE! AVOID EXPOSURE OF (PREGNANT WOMEN!)	
•INHALATION	Dizziness. Drowsiness. Headache. Nausea.		Ventilation, local exhaust, or breathing protection.		Fresh air, rest. Refer for medical attention.
•SKIN	Dry skin. Redness.		Protective gloves.		Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain.				First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Burning sensation. Abd (Further see Inhalation)			Rinse mouth. Do NOT induce vomiting. Refer for medical attention.	
SPILLAG	SPILLAGE DISPOSAL		STORAGE PACKAGING & LABEL		CKAGING & LABELLING
Ventilation. Remove all ignition sources. Collect leaking and spilled liquid in sealable containers as far as possible. Absorb		parated from strong oxidants	n strong oxidants Note: C Xn symbol		

personal protection: filter respirator for organic gases and vapours.) UN Packing Group: III

SEE IMPORTANT INFORMATION ON BACK

R: 10-20/21-38

UN Hazard Class: 3

S: 2-25

ICSC: 0084

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

o-XYLENE ICSC: 0084

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID, WITH CHARACTERISTIC ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation, through the skin and by ingestion.
M	PHYSICAL DANGERS: As a result of flow, agitation, etc., electrostatic charges	INHALATION RISK: A harmful contamination of the air will be reached
P	can be generated.	rather slowly on evaporation of this substance at 20°C.
О	CHEMICAL DANGERS:	EFFECTS OF SHORT-TERM EXPOSURE: The substance is imitating to the guess and the claim The
R	Reacts with strong acids strong oxidants	The substance is irritating to the eyes and the skin The substance may cause effects on the central nervous
Т	OCCUPATIONAL EXPOSURE LIMITS: TLV: 100 ppm as TWA 150 ppm as STEL A4 (ACGIH	system If this liquid is swallowed, aspiration into the lungs may result in chemical pneumonitis.
A	2001). BEI (ACGIH 2001). MAK: 100 ppm 440 mg/m ³	EFFECTS OF LONG-TERM OR REPEATED
N	Peak limitation category: II(2) skin absorption (H);	EXPOSURE: The liquid defats the skin. The substance may have
Т	Pregnancy risk group: D (DFG 2005). EU OEL: 50 ppm as TWA 100 ppm as STEL	effects on the central nervous system. Exposure to the substance may enhance hearing damage caused by exposure to noise. Animal tests show that this substance
D	(skin) (EU 2000).	possibly causes toxicity to human reproduction or development.
A	OSHA PEL±: TWA 100 ppm (435 mg/m ³) NIOSH REL: TWA 100 ppm (435 mg/m ³) ST 150 ppm	
Т	(655 mg/m^3)	
A	NIOSH IDLH: 900 ppm See: <u>95476</u>	
PHYSICAL PROPERTIES	Boiling point: 144°C Melting point: -25°C Relative density (water = 1): 0.88 Solubility in water: none Vapour pressure, kPa at 20°C: 0.7	Relative vapour density (air = 1): 3.7 Relative density of the vapour/air-mixture at 20°C (air = 1): 1.02 Flash point: 32°C c.c. Auto-ignition temperature: 463°C Explosive limits, vol% in air: 0.9-6.7 Octanol/water partition coefficient as log Pow: 3.12
ENVIRONMENTAL DATA	The substance is toxic to aquatic organisms.	
	NOTES	
	ee of exposure, periodic medical examination is indicated. p-Xylene and 0085 m-Xylene.	The recommendations on this Card also apply to technical
	1 J	Transport Emergency Card: TEC (R)-30S1307-III NFPA Code: H 2; F 3; R 0;

ADDITIONAL INFORMATION

ICSC: 0084 o-XYLENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

p-XYLENE ICSC: 0086











para-Xylene 1,4-Dimethylbenzene p-Xylol $C_6H_4(CH_3)_2/C_8H_{10}$ Molecular mass: 106.2

ICSC # 0086 CAS # 106-42-3 RTECS # <u>ZE2625000</u> UN # 1307

EC # 601-022-00-9 August 03, 2002 Validated



TYPES OF HAZARD/ EXPOSURE	ACUTE HAZA SYMPTOM		PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Flammable.		NO open flames, NO sparks, and NO smoking.	Powder, water spray, foam, carbon dioxide.
EXPLOSION	Above 27°C explosive var mixtures may be formed.	pour/air	Above 27°C use a closed system, ventilation, and explosion-proof electrical equipment. Prevent build-up of electrostatic charges (e.g., by grounding).	In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE			STRICT HYGIENE! AVOID EXPOSURE OF (PREGNANT) WOMEN!	
•INHALATION	Dizziness. Drowsiness. He Nausea.	eadache.	Ventilation, local exhaust, or breathing protection.	Fresh air, rest. Refer for medical attention.
•SKIN	Dry skin. Redness.		Protective gloves.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain.		Safety spectacles.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Burning sensation. Abdon (Further see Inhalation).	ninal pain.	Do not eat, drink, or smoke during work.	Rinse mouth. Do NOT induce vomiting. Refer for medical attention.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
Ventilation. Remove all ignition sources.	Fireproof. Separated from strong oxidants,	
Collect leaking and spilled liquid in sealable	strong acids	Note: C
containers as far as possible. Absorb		Xn symbol
remaining liquid in sand or inert absorbent		R: 10-20/21-38
and remove to safe place. Do NOT let this		S: 2-25
chemical enter the environment. (Extra		UN Hazard Class: 3
personal protection: filter respirator for		UN Packing Group: III
organic gases and vapours.)		

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0086

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

p-XYLENE ICSC: 0086

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID, WITH CHARACTERISTIC ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by	
M	ODOUR.	inhalation, through the skin and by ingestion.	
P	PHYSICAL DANGERS: As a result of flow, agitation, etc., electrostatic charges	INHALATION RISK: A harmful contamination of the air will be reached	
0	can be generated.	rather slowly on evaporation of this substance at 20°C.	
R	CHEMICAL DANGERS: Reacts with strong acids strong oxidants	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes and the skin The substance may cause effects on the central nervous	
Т	OCCUPATIONAL EXPOSURE LIMITS: TLV: 100 ppm as TWA 150 ppm as STEL A4 (ACGIH	system If this liquid is swallowed, aspiration into the	
A	2001). BEI (ACGIH 2001). MAK: 100 ppm 440 mg/m ³	EFFECTS OF LONG-TERM OR REPEATED	
N	Peak limitation category: II(2)	EXPOSURE:	
Т	skin absorption (H); Pregnancy risk group: D (DFG 2005).	The liquid defats the skin. The substance may have effects on the central nervous system. Animal tests show that this substance possibly causes toxicity to human	
D	EU OEL: 50 ppm as TWA 100 ppm as STEL (skin) (EU 2000).	reproduction or development.	
A	OSHA PEL±: TWA 100 ppm (435 mg/m³) NIOSH REL: TWA 100 ppm (435 mg/m³) ST 150 ppm		
Т	(655 mg/m ³) NIOSH IDLH: 900 ppm See: <u>95476</u>		
A			
PHYSICAL PROPERTIES	Boiling point: 138°C Melting point: 13°C Relative density (water = 1): 0.86 Solubility in water: none Vapour pressure, kPa at 20°C: 0.9	Relative vapour density (air = 1): 3.7 Relative density of the vapour/air-mixture at 20°C (air = 1): 1.02 Flash point: 27°C c.c. Auto-ignition temperature: 528°C Explosive limits, vol% in air: 1.1-7.0 Octanol/water partition coefficient as log Pow: 3.15	
ENVIRONMENTAL DATA	The substance is toxic to aquatic organisms.		
NOTES			
Depending on the degree of exposure, periodic medical examination is indicated. The recommendations on this Card also apply to technical xylene. See ICSC 0084 o-Xylene and 0085 m-Xylene. Transport Emergency Card: TEC (R)-30S1307-III NFPA Code: H 2; F 3; R 0;			
ADDITIONAL INFORMATION			

ICSC: 0086 p-XYLENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

1,2,4-TRIMETHYLBENZENE











 $\begin{array}{c} \text{Pseudocumene} \\ \text{C}_9 \text{H}_{12} \end{array}$

Molecular mass: 120,2

ICSC # 1433 CAS # 95-63-6 RTECS # <u>DC3325000</u>

UN # 1993

EC# 601-043-00-3

March 06, 2002 Peer reviewed



ICSC: 1433

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS/ SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Flammable.	NO open flames, NO sparks, and NO smoking.	Alcohol-resistant foam, dry powder, carbon dioxide.
EXPLOSION	Above 44°C explosive vapour/air mixtures may be formed.	Above 44°C use a closed system, ventilation, and explosion-proof electrical equipment. Prevent build-up of electrostatic charges (e.g., by grounding).	In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE		PREVENT GENERATION OF MISTS!	
•INHALATION	Confusion. Cough. Dizziness. Drowsiness. Headache. Sore throat. Vomiting.	Ventilation, local exhaust, or breathing protection.	Fresh air, rest. Refer for medical attention.
•SKIN	Redness. Dry skin.	Protective gloves.	Rinse skin with plenty of water or shower.
•EYES	Redness. Pain.	Safety spectacles.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	(See Inhalation).	Do not eat, drink, or smoke during work.	Rinse mouth. Do NOT induce vomiting. Refer for medical attention.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
		Xn symbol N symbol R: 10-20-36/37/38-51/53 S: 2-26-61 UN Hazard Class: 3 UN Packing Group: III

SEE IMPORTANT INFORMATION ON BACK

ICSC: 1433

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

1,2,4-TRIMETHYLBENZENE

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID, WITH CHARACTERISTIC	ROUTES OF EXPOSURE: The substance can be absorbed into the body by	
M	ODOUR.	inhalation.	
P	PHYSICAL DANGERS:	INHALATION RISK:	
О		A harmful contamination of the air will be reached rather slowly on evaporation of this substance at 20°C;	
R	CHEMICAL DANGERS: The substance decomposes on burning producing toxic	on spraying or dispersing, however, much faster.	
Т	and irritating fumes Reacts violently with strong oxidants causing fire and explosion hazard.	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes the skin and the respiratory tract If this liquid is swallowed, aspiration	
A	OCCUPATIONAL EXPOSURE LIMITS: TLV: (as mixed isomers) 25 ppm as TWA (ACGIH	into the lungs may result in chemical pneumonitis. The substance may cause effects on the central nervous	
N	2004).	system	
Т	MAK: (as mixed isomers) 20 ppm 100 mg/m³ Peak limitation category: II(2) Pregnancy risk group: C (DFG 2004).	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE:	
D	OSHA PEL±: none NIOSH REL: TWA 25 ppm (125 mg/m ³)	The liquid defats the skin. Lungs may be affected by repeated or prolonged exposure, resulting in chronic	
A	NIOSH IDLH: N.D. See: IDLH INDEX	bronchitis The substance may have effects on the central nervous system blood See Notes.	
Т			
A			
PHYSICAL PROPERTIES	Boiling point: 169°C Melting point: -44°C Relative density (water = 1): 0.88 Solubility in water: very poor Relative vapour density (air = 1): 4.1	Relative density of the vapour/air-mixture at 20°C (air = 1): 1.01 Flash point: 44°C c.c. Auto-ignition temperature: 500°C Explosive limits, vol% in air: 0.9-6.4 Octanol/water partition coefficient as log Pow: 3.8	
ENVIRONMENTAL	The substance is toxic to aquatic organisms. Bioaccumulation of this chemical may occur in fish.		

ENVIRONMENTAI DATA



ICSC: 1433

NOTES

Use of alcoholic beverages enhances the harmful effect. Depending on the degree of exposure, periodic medical examination is suggested. See also ICSC 1155 1,3,5-Trimethylbenzene (Mesitylene), ICSC 1362 1,2,3-Trimethylbenzene (Hemimellitene), ICSC 1389 Trimethylbenzene (mixed isomers). 1,3,5-Trimethylbenzene (Mesitylene) is classified as a marine pollutant.

Transport Emergency Card: TEC (R)-30GF1-III NFPA Code: H0; F2; R0;

ADDITIONAL INFORMATION

ICSC: 1433 1,2,4-TRIMETHYLBENZENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

ETHYLBENZENE











Ethylbenzol Phenylethane EB C_8H_{10} / $C_6H_5C_2H_5$ Molecular mass: 106.2

ICSC # 0268 CAS # 100-41-4 RTECS # <u>DA0700000</u>

UN # 1175

EC # 601-023-00-4 March 13, 1995 Validated



ICSC: 0268

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS/ SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Highly flammable.	NO open flames, NO sparks, and NO smoking.	Powder, AFFF, foam, carbon dioxide.
EXPLOSION	Vapour/air mixtures are explosive.		In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE		PREVENT GENERATION OF MISTS!	
•INHALATION	Cough. Dizziness. Drowsiness. Headache.	Ventilation, local exhaust, or breathing protection.	Fresh air, rest. Refer for medical attention.
•SKIN	Dry skin. Redness.	Protective gloves.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain. Blurred vision.	Face shield or eye protection in combination with breathing protection.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	(Further see Inhalation).	Do not eat, drink, or smoke during work.	Rinse mouth. Give a slurry of activated charcoal in water to drink. Refer for medical attention.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
Ventilation. Collect leaking liquid in covered containers. Absorb remaining liquid in sand or inert absorbent and remove to safe place. Do NOT wash away into sewer. Personal protection: A filter respirator for organic gases and vapours.		F symbol Xn symbol R: 11-20 S: 2-16-24/25-29 UN Hazard Class: 3 UN Packing Group: II

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0268

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

ETHYLBENZENE ICSC: 0268

I M	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID , WITH AROMATIC ODOUR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation of its vapour, through the skin and by ingestion.
P	PHYSICAL DANGERS:	ingestion.
О	The vapour mixes well with air, explosive mixtures are easily formed.	INHALATION RISK: A harmful contamination of the air will be reached rather slowly on evaporation of this substance at 20°C.
R	CHEMICAL DANGERS:	, 1
Т	Reacts with strong oxidants. Attacks plastic and rubber.	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes the skin and the
A	OCCUPATIONAL EXPOSURE LIMITS: TLV: 100 ppm as TWA 125 ppm as STEL A3 (confirmed animal carcinogen with unknown relevance	respiratory tract Swallowing the liquid may cause aspiration into the lungs with the risk of chemical pneumonitis. The substance may cause effects on the
N	to humans); BEI issued (ACGIH 2005).	central nervous system Exposure far above the OEL
Т	MAK: skin absorption (H); Carcinogen category: 3A; (DFG 2004).	could cause lowering of consciousness. EFFECTS OF LONG-TERM OR REPEATED
D	OSHA PEL±: TWA 100 ppm (435 mg/m³) NIOSH REL: TWA 100 ppm (435 mg/m³) ST 125 ppm	EXPOSURE: Repeated or prolonged contact with skin may cause dermatitis.
A	(545 mg/m ³) NIOSH IDLH: 800 ppm 10%LEL See: <u>100414</u>	dermatitis.
T		
A		
PHYSICAL PROPERTIES	Boiling point: 136°C Melting point: -95°C Relative density (water = 1): 0.9 Solubility in water, g/100 ml at 20°C: 0.015 Vapour pressure, kPa at 20°C: 0.9 Relative vapour density (air = 1): 3.7	Relative density of the vapour/air-mixture at 20°C (air = 1): 1.02 Flash point: 18°C c.c. Auto-ignition temperature: 432°C Explosive limits, vol% in air: 1.0-6.7 Octanol/water partition coefficient as log Pow: 3.2
ENVIRONMENTAL DATA	The substance is harmful to aquatic organisms.	
	NOTES	
The odour warning who	en the exposure limit value is exceeded is insufficient.	
I	-	

Transport Emergency Card: TEC (R)-30S1175 or 30GF1-I+II

NFPA Code: H2; F3; R0

ADDITIONAL INFORMATION

ICSC: 0268 ETHYLBENZENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

1,3,5-TRIMETHYLBENZENE











Molecular mass: 120.2

ICSC # 1155 CAS # 108-67-8 RTECS # <u>OX6825000</u> UN # 2325

EC # 601-025-00-5

March 06, 2002 Peer reviewed



ICSC: 1155

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Flammable.	NO open flames, NO sparks, and NO smoking.	Alcohol-resistant foam, dry powder, carbon dioxide.
EXPLOSION	Above 50°C explosive vapour/mixtures may be formed.	Above 50°C use a closed system, ventilation, and explosion-proof electrical equipment. Prevent build-up of electrostatic charges (e.g., by grounding).	In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE		PREVENT GENERATION OF MISTS!	
•INHALATION	Confusion. Cough. Dizziness. Drowsiness. Headache. Sore th Vomiting.	Ventilation, local exhaust, or breathing protection.	Fresh air, rest. Refer for medical attention.
•SKIN	Redness. Dry skin.	Protective gloves.	Remove contaminated clothes. Rinse skin with plenty of water or shower.
•EYES	Redness. Pain.	Safety spectacles.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	(See Inhalation).	Do not eat, drink, or smoke during work.	Rinse mouth. Do NOT induce vomiting. Refer for medical attention.
CDILI A CI	FDISPOSAL	STOPACE PA	CKACING & LARFILING

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
		Marine pollutant. Xi symbol N symbol R: 10-37-51/53 S: 2-61 UN Hazard Class: 3 UN Packing Group: III

SEE IMPORTANT INFORMATION ON BACK

ICSC: 1155

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

1,3,5-TRIMETHYLBENZENE

I	PHYSICAL STATE; APPEARANCE: COLOURLESS LIQUID, WITH CHARACTERISTIC	ROUTES OF EXPOSURE: The substance can be absorbed into the body by		
M	ODOUR.	inhalation.		
P	PHYSICAL DANGERS:	INHALATION RISK: A harmful contamination of the air will be reached		
0	CHEMICAL DANGERS	rather slowly on evaporation of this substance at 20°C;		
R	CHEMICAL DANGERS: The substance decomposes on burning producing toxic	on spraying or dispersing, however, much faster.		
Т	and irritating fumes. Reacts violently with strong oxidants causing fire and explosion hazard.	EFFECTS OF SHORT-TERM EXPOSURE: The substance is irritating to the eyes the skin and the		
A	OCCUPATIONAL EXPOSURE LIMITS:	respiratory tract If this liquid is swallowed, aspiration into the lungs may result in chemical pneumonitis. The		
N	TLV (as mixed isomers): 25 ppm; (ACGIH 2001). MAK (all isomers): 20 ppm; 100 mg/m ³ ; class II 1 ©	substance may cause effects on the central nervous system.		
Т	(2001) OSHA PEL <u>†</u> : none	EFFECTS OF LONG-TERM OR REPEATED		
D	NIOSH REL: TWA 25 ppm (125 mg/m ³) NIOSH IDLH: N.D. See: IDLH INDEX	EXPOSURE: The liquid defats the skin. Lungs may be affected by		
		repeated or prolonged exposure, resulting in chronic bronchitis. The substance may have effects on the		
A		central nervous system blood See Notes.		
T				
A				
PHYSICAL	Boiling point: 165°C Melting point: -45°C Relative density (water = 1): 0.86	Relative vapour density (air = 1): 4.1 Relative density of the vapour/air-mixture at 20°C (air = 1): 1.01		
PROPERTIES	Solubility in water:	Flash point: 50°C (c.c.)		
	very poor Vapour pressure, kPa at 20°C: 0.25	Auto-ignition temperature: 550°C Octanol/water partition coefficient as log Pow: 3.42		
ENVIRONMENTAL	The substance is harmful to aquatic organisms. Bioaccumulation of this chemical may occur in fish.			

DATA



ICSC: 1155

NOTES

Use of alcoholic beverages enhances the harmful effect. Depending on the degree of exposure, periodic medical examination is indicated. See ICSC 1433 1,2,4-Trimethylbenzene (Pseudocumene), ICSC 1362 1,2,3-Trimethylbenzene (Hemimellitene), ICSC 1389 Trimethyl benzene (mixed isomers).

Transport Emergency Card: TEC (R)-30S2325

NFPA Code: H0; F2; R0

ADDITIONAL INFORMATION

ICSC: 1155 1,3,5-TRIMETHYLBENZENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

INDENO(1,2,3-cd)PYRENE











ICSC: 0730

ICSC: 0730

o-Phenylenepyrene 2,3-Phenylenepyrene $C_{22}H_{12}$

Molecular mass: 276.3

ICSC # 0730 CAS # 193-39-5 RTECS # <u>NK9300000</u>

March 25, 1999 Peer reviewed

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE					In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION					
EXPOSURE			AVOID ALL CONTACT!		
•INHALATION			Local exhaust or breathing protect	ction.	Fresh air, rest.
•SKIN			Protective gloves. Protective clot	hing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES			Safety spectacles or eye protection in combination with breathing protection.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION			Do not eat, drink, or smoke during work.		Rinse mouth. Refer for medical attention.
SPILLAGE	E DISPOSAL		STORAGE	PA	CKAGING & LABELLING
Sweep spilled substance into covered containers; if appropriate, moisten first to prevent dusting. Carefully collect remainder, then remove to safe place. Do NOT let this chemical enter the environment.		ontain effluent from fire . Well closed. R: S:			
SEE IMPORTANT INFORMATION ON BACK					
Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs,					

International Chemical Safety Cards

NIOSH RELs and NIOSH IDLH values.

INDENO(1.2.3-cd)PYRENE

TIDE TO(1,2,5-cu)1 1 KENE	
I	PHYSICAL STATE; APPEARANCE:	ROUTES OF EXPOSURE:
	YELLOW CRYSTALS	The substance can be absorbed into the body by inhalation
\mathbf{M}		of its aerosol and through the skin.
	PHYSICAL DANGERS:	•
P		INHALATION RISK:

O R T A N T D A T	CHEMICAL DANGERS: Upon heating, toxic fumes are formed. OCCUPATIONAL EXPOSURE LIMITS: TLV not established. MAK: Carcinogen category: 2; (DFG 2004).	Evaporation at 20°C is negligible; a harmful concentration of airborne particles can, however, be reached quickly. EFFECTS OF SHORT-TERM EXPOSURE: EFFECTS OF LONG-TERM OR REPEATED EXPOSURE: This substance is possibly carcinogenic to humans.		
PHYSICAL PROPERTIES	Boiling point: 536°C Melting point: 164°C Solubility in water: none	Octanol/water partition coefficient as log Pow: 6.58		
ENVIRONMENTAL DATA	Ilwater quality Ringerumulation of this chemical may occur in tich			
NOTES				

Indeno(1,2,3-cd)pyrene is present as a component of polycyclic aromatic hydrocarbons (PAH) content in the environment usually resulting from the incomplete combustion or pyrolysis of organic matters, especially fossil fuels and tobacco. ACGIH recommends environment containing Indeno(1,2,3-c,d)pyrene should be evaluated in terms of the TLV-TWA for coal tar pitch volatile, as benzene soluble 0.2 mg/m³. Insufficient data are available on the effect of this substance on human health, therefore utmost care must be taken.

ADDITIONAL INFORMATION ICSC: 0730 INDENO(1,2,3-cd)PYRENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

CHRYSENE ICSC: 1672











 $\begin{array}{c} Benzoaphenanthrene\\ 1,2\text{-Benzophenanthrene}\\ 1,2,5,6\text{-Dibenzonaphthalene}\\ C_{18}H_{12} \end{array}$

Molecular mass: 228.3







ICSC # 1672 CAS # 218-01-9 RTECS # <u>GC0700000</u> UN # 3077

EC # 601-048-00-0 October 12, 2006 Validated

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ	PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Combustible.	NO open flames.		Water spray. Dry powder. Foam. Carbon dioxide.
EXPLOSION	Finely dispersed particle explosive mixtures in air	Prevent deposition of dust; closed system, dust explosion-proof election equipment and lighting.		
EXPOSURE	See EFFECTS OF LONG REPEATED EXPOSUR	AVOID ALL CONTACT!		
•INHALATION		Local exhaust or breathing protection.		Fresh air, rest.
•SKIN		Protective gloves. Protective clot	hing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES				First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION		Do not eat, drink, or smoke during work.		Rinse mouth.
SPILLAGE DISPOSAL STORAGE PACKAGING & LABER		CKAGING & LABELLING		

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
Personal protection: P3 filter respirator for	Separated from strong oxidants, Provision to	
toxic particles. Do NOT let this chemical enter		T symbol
the environment. Sweep spilled substance into	in an area without drain or sewer access.	N symbol
sealable containers; if appropriate, moisten first		R: 45-68-50/53
to prevent dusting. Carefully collect remainder,		S: 53-45-60-61
then remove to safe place.		UN Hazard Class: 9
		UN Packing Group: III
		Signal: Warning
		Aqua-Cancer
		Suspected of causing cancer
		Very toxic to aquatic life with long lasting
		effects
		Very toxic to aquatic life

SEE IMPORTANT INFORMATION ON BACK

ICSC: 1672

International Chemical Safety Cards

CHRYSENE ICSC: 1672

I	PHYSICAL STATE; APPEARANCE:	ROUTES OF EXPOSURE:			
M	COLOURLESS TO BEIGE CRYSTALS OR POWDER	The substance can be absorbed into the body by inhalation of its aerosol, through the skin and by ingestion.			
P	PHYSICAL DANGERS: Dust explosion possible if in powder or granular form,	INHALATION RISK:			
О	mixed with air.	A harmful concentration of airborne particles can be reached quickly when dispersed			
R	CHEMICAL DANGERS: The substance decomposes on burning producing toxic	EFFECTS OF SHORT-TERM EXPOSURE:			
Т	fumes Reacts violently with strong oxidants				
A	OCCUPATIONAL EXPOSURE LIMITS: TLV: A3 (confirmed animal carcinogen with unknown	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE:			
N	relevance to humans); (ACGIH 2006). MAK not established.	This substance is possibly carcinogenic to humans.			
Т					
D					
A					
Т					
A					
PHYSICAL PROPERTIES	Boiling point: 448°C Melting point: 254 - 256°C Density: 1.3 g/cm ³	Solubility in water: very poor Octanol/water partition coefficient as log Pow: 5.9			
ENVIRONMENTAL DATA	lite etrangly advised that this substance does not enter the environment				
	NOTES				

Depending on the degree of exposure, periodic medical examination is suggested. Do NOT take working clothes home. This substance does not usually occur as a pure substance but as a component of polyaromatic hydrocarbon (PAH) mixtures. Human population studies have associated PAH's exposure with cancer and cardiovascular diseases.

Transport Emergency Card: TEC (R)-90GM7-III

	ADDITIONAL INFORMATION	
ICSC: 1672		CHRYSENE
	(C) IPCS, CEC, 1994	

IMPORTANT LEGAL NOTICE:

BENZO(k)FLUORANTHENE











Dibenzo(b,jk)fluorene 8,9-Benzofluoranthene 11.12-Benzofluoranthene $C_{20}H_{12}$

Molecular mass: 252.3





ICSC: 0721

ICSC# 0721 CAS# 207-08-9 RTECS # DF6350000 EC# 601-036-00-5 March 25, 1999 Peer reviewed

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS/ SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE			In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION			
EXPOSURE		AVOID ALL CONTACT!	
•INHALATION		Local exhaust or breathing protection.	Fresh air, rest.
•SKIN		Protective gloves. Protective clothing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES		Safety spectacles or eye protection in combination with breathing protection if powder.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
·INGESTION		Do not eat, drink, or smoke during work.	Rinse mouth. Refer for medical attention.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
Sweep spilled substance into covered	Provision to contain effluent from fire	
containers; if appropriate, moisten first to	extinguishing. Well closed.	T symbol
prevent dusting. Carefully collect remainder,		N symbol
then remove to safe place. Do NOT let this		R: 45-50/53
chemical enter the environment.		S: 53-45-60-61

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0721

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

BENZO(k)FLUORANTHENE

ICSC: 0721

PHYSICAL STATE; APPEARANCE:

YELLOW CRYSTALS

ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation of its aerosol and through the skin.

M

I

P O R T A N T D A	PHYSICAL DANGERS: CHEMICAL DANGERS: Upon heating, toxic fumes are formed. OCCUPATIONAL EXPOSURE LIMITS: TLV not established. MAK: Carcinogen category: 2; (DFG 2004).	INHALATION RISK: Evaporation at 20°C is negligible; a harmful concentration of airborne particles can, however, be reached quickly. EFFECTS OF SHORT-TERM EXPOSURE: EFFECTS OF LONG-TERM OR REPEATED EXPOSURE: This substance is possibly carcinogenic to humans.		
A				
PHYSICAL PROPERTIES	Boiling point: 480°C Melting point: 217°C Solubility in water: none	Octanol/water partition coefficient as log Pow: 6.84		
ENVIRONMENTAL DATA	This substance may be hazardous to the environment; special attention should be given to air quality and water quality. Bioaccumulation of this chemical may occur in crustacea and in fish.			
NOTES				
Ranzo(k) fluoranthana is present as a component of polycyclic gromatic hydrocarbons (PAH) content in the anyironment usually resulting from				

Benzo(k)fluoranthene is present as a component of polycyclic aromatic hydrocarbons (PAH) content in the environment usually resulting from the incomplete combustion or pyrolysis of organic matters, especially fossil fuels and tobacco. ACGIH recommends environment containing benzo(k)fluoranthene should be evaluated in terms of the TLV-TWA for coal tar pitch volatile, as benzene soluble 0.2 mg/m³. Insufficient data are available on the effect of this substance on human health, therefore utmost care must be taken.

ADDITIONAL INFORMATION ICSC: 0721 BENZO(k)FLUORANTHENE (C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

BENZO(b)FLUORANTHENE











Benz(e)acephenanthrylene 2,3-Benzofluoroanthene Benzo(e)fluoranthene 3,4-Benzofluoranthene $C_{20}H_{12}$

Molecular mass: 252.3





ICSC: 0720

ICSC # 0720 CAS # 205-99-2 RTECS # <u>CU1400000</u> EC # 601-034-00-4 March 25, 1999 Peer reviewed

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ	PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE				In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION				
EXPOSURE		AVOID ALL CONTACT!		
•INHALATION		Local exhaust or breathing protect	ction.	Fresh air, rest.
•SKIN		Protective gloves. Protective clot	hing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES		Safety spectacles or eye protection combination with breathing protections.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION		Do not eat, drink, or smoke durinwork.	ng	Rinse mouth. Refer for medical attention.
SPILLAGE DISPOSAL STORAGE PACKAGING & LAB		CKAGING & LABELLING		

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
containers; if appropriate, moisten first to prevent dusting. Carefully collect remainder, then remove to safe place. Do NOT let this		T symbol N symbol R: 45-50/53
chemical enter the environment.		S: 53-45-60-61

SEE IMPORTANT INFORMATION ON BACK

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

BENZO(b)FLUORANTHENE

ICSC: 0720

M P O R T A N T D A T A	PHYSICAL DANGERS: CHEMICAL DANGERS: Upon heating, toxic fumes are formed. OCCUPATIONAL EXPOSURE LIMITS: TLV: A2 (suspected human carcinogen); (ACGIH 2004). MAK: Carcinogen category: 2; (DFG 2004).	of its aerosol and through the skin. INHALATION RISK: Evaporation at 20°C is negligible; a harmful concentration of airborne particles can, however, be reached quickly. EFFECTS OF SHORT-TERM EXPOSURE: EFFECTS OF LONG-TERM OR REPEATED EXPOSURE: This substance is possibly carcinogenic to humans. May cause genetic damage in humans.
PHYSICAL PROPERTIES	Boiling point: 481°C Melting point: 168°C Solubility in water: none	Octanol/water partition coefficient as log Pow: 6.12
ENVIRONMENTAL DATA	This substance may be hazardous to the environment; speci water quality. NOTES	al attention should be given to air quality and

Benzo(b)fluoranthene is present as a component of polycyclic aromatic hydrocarbons (PAH) content in the environment usually resulting from the incomplete combustion or pyrolysis of organic matters, especially fossil fuels and tobacco. ACGIH recommends environment containing benzo(b)fluoranthene should be evaluated in terms of the TLV-TWA for coal tar pitch volatile, as benzene soluble 0.2 mg/m³. Insufficient data are available on the effect of this substance on human health, therefore utmost care must be taken.

ADDITIONAL INFORMATION ICSC: 0720 BENZO(b)FLUORANTHENE (C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

BENZO(a)PYRENE







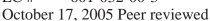




Benz(a)pyrene 3,4-Benzopyrene Benzo(d,e,f)chrysene $C_{20}H_{12}$

Molecular mass: 252.3

ICSC # 0104 CAS # 50-32-8 RTECS # <u>DJ3675000</u> EC # 601-032-00-3







ICSC: 0104

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZARDS/ SYMPTOMS	PREVENTION	FIRST AID/ FIRE FIGHTING
FIRE	Combustible.	NO open flames.	Water spray, foam, powder, carbon dioxide.
EXPLOSION			
EXPOSURE	See EFFECTS OF LONG-TERM OR REPEATED EXPOSURE.	AVOID ALL CONTACT! AVOID EXPOSURE OF (PREGNANT) WOMEN!	
•INHALATION		Local exhaust or breathing protection.	Fresh air, rest.
•SKIN	MAY BE ABSORBED!	Protective gloves. Protective clothing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES		Safety goggles or eye protection in combination with breathing protection.	First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION		Do not eat, drink, or smoke during work.	Induce vomiting (ONLY IN CONSCIOUS PERSONS!). Refer for medical attention.

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
Evacuate danger area! Personal protection: complete protective clothing including self-contained breathing apparatus. Do NOT let this chemical enter the environment. Sweep spilled substance into sealable containers; if appropriate, moisten first to prevent dusting. Carefully collect remainder, then remove to safe place.		T symbol N symbol R: 45-46-60-61-43-50/53 S: 53-45-60-61

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0104

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

BENZO(a)PYRENE

I	PHYSICAL STATE; APPEARANCE:	ROUTES OF EXPOSURE:
	PALE-YELLOW CRYSTALS	The substance can be absorbed into the body by inhalation
\mathbf{M}		of its aerosol, through the skin and by ingestion.
	PHYSICAL DANGERS:	
P		INHALATION RISK:
		Evaporation at 20°C is negligible; a harmful concentration
0	CHEMICAL DANGERS:	of airborne particles can, however, be reached quickly
	Reacts with strong oxidants causing fire and explosion	when dispersed.
R	hazard.	when dispersed.
K	nazaru.	EFFECTS OF SHOPE TERM EXPOSIDE.
T		EFFECTS OF SHORT-TERM EXPOSURE:
1	OCCUPATIONAL EXPOSURE LIMITS:	
A	TLV: Exposure by all routes should be carefully controlled	THE CEG OF LOVE THE LOD DEDE LEED
A	to levels as low as possible A2 (suspected human	EFFECTS OF LONG-TERM OR REPEATED
N	carcinogen); (ACGIH 2005).	EXPOSURE:
N	MAK:	This substance is carcinogenic to humans. May cause
T.	Carcinogen category: 2; Germ cell mutagen group: 2;	heritable genetic damage to human germ cells. Animal tests
T	(DFG 2005).	show that this substance possibly causes toxicity to human
		reproduction or development.
D		
\mathbf{A}		
T		
\mathbf{A}		
	D 11. 1. 40.000	0.11777
DIII GAA	Boiling point: 496°C	Solubility in water: none (<0.1 g/100 ml)
PHYSICAL	Melting point: 178.1°C	Vapour pressure :
PROPERTIES	Density: 1.4	negligible
	g/cm³	Octanol/water partition coefficient as log Pow: 6.04
	The substance is very toxic to aquatic organisms. Bioaccumu	ulation of this chemical may occur in fish in
ENVIRONMENTAL	plants and in molluscs. The substance may cause long-term	
DATA		
	Momed	
	NOTES	

Do NOT take working clothes home. Benzo(a)pyrene is present as a component of polycyclic aromatic hydrocarbons (PAHs) in the environment, usually resulting from the incomplete combustion or pyrolysis of organic matters, especially fossil fuels and tobacco.

ADDITIONAL INFORMATION ICSC: 0104 BENZO(a)PYRENE

(C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

BENZ(a)ANTHRACENE











1,2-Benzoanthracene Benzo(a)anthracene 2,3-Benzphenanthrene Naphthanthracene $C_{18}H_{12}$

Molecular mass: 228.3





ICSC: 0385

ICSC # 0385 CAS # 56-55-3 RTECS # <u>CV9275000</u> EC # 601-033-00-9 October 23, 1995 Validated

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ SYMPTO		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Combustible.				Water spray, powder. In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION	explosive mixtures in air.		Prevent deposition of dust; closed system, dust explosion-proof electrical equipment and lighting.		
EXPOSURE			AVOID ALL CONTACT!		
•INHALATION			Local exhaust or breathing protection	ction.	Fresh air, rest.
•SKIN					Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES			Safety goggles face shield or eye protection in combination with breathing protection.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION		Do not eat, drink, or smoke during work. Wash hands before eating.		Rinse mouth.	
SPILLAGE DISPOSAL		STORAGE	PA	CKAGING & LABELLING	
Sweep spilled substance into sealable containers; if appropriate, moisten first to prevent dusting. Carefully collect remainder, then remove to safe place. Personal protection: complete protective clothing including self-contained breathing apparatus. Well closed.			T symb N symb R: 45-5 S: 53-4	bol	

SEE IMPORTANT INFORMATION ON BACK

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

ICSC: 0385

BENZ(a)ANTHRACENE

PHYSICAL STATE; APPEARANCE:

Ι

M	COLOURLESS TO YELLOW BROWN FLUORESCENT FLAKES OR POWDER.	The substance can be absorbed into the body by inhalation, through the skin and by ingestion.		
P	PHYSICAL DANGERS: Dust explosion possible if in powder or granular form,	INHALATION RISK: Evaporation at 20°C is negligible; a harmful concentration		
О	mixed with air.	of airborne particles can, however, be reached quickly.		
R	CHEMICAL DANGERS:	EFFECTS OF SHORT-TERM EXPOSURE:		
T				
A	OCCUPATIONAL EXPOSURE LIMITS: TLV: A2 (suspected human carcinogen); (ACGIH 2004). MAK:	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE: This substance is probably carcinogenic to humans.		
N	Carcinogen category: 2 (as pyrolysis product of organic	This substance is probably careinogenic to numans.		
Т	materials) (DFG 2005).			
D				
A				
Т				
A				
PHYSICAL PROPERTIES	Sublimation point: 435°C Melting point: 162°C Relative density (water = 1): 1.274 Solubility in water: none	Vapour pressure, Pa at 20°C: 292 Octanol/water partition coefficient as log Pow: 5.61		
ENVIRONMENTAL DATA	Bioaccumulation of this chemical may occur in seafood.			
NOTES				
This substance is one of many polycyclic aromatic hydrocarbons - standards are usually established for them as mixtures, e.g., coal tar pitch volatiles. However, it may be encountered as a laboratory chemical in its pure form. Insufficient data are available on the effect of this substance on human health, therefore utmost care must be taken. Do NOT take working clothes home. Tetraphene is a common name. Card has been partly updated in October 2005 and August 2006: see sections Occupational Exposure Limits, EU classification.				

ROUTES OF EXPOSURE:

ADDITIONAL INFORMATION

ICSC: 0385

(C) IPCS, CEC, 1994

BENZ(a)ANTHRACENE

IMPORTANT LEGAL NOTICE:

NICKEL ICSC: 0062











Ni Atomic mass: 58.7 (powder)

ICSC # 0062 CAS # 7440-02-0 RTECS # <u>QR5950000</u> EC # 028-002-00-7

October 17, 2001 Peer reviewed

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZA SYMPTON		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Flammable as dust. Toxio be released in a fire.	c fumes may			Dry sand. NO carbon dioxide. NO water.
EXPLOSION	Finely dispersed particles explosive mixtures in air.		Prevent deposition of dust; closed system, dust explosion-proof election equipment and lighting.		
EXPOSURE			PREVENT DISPERSION OF DUAVOID ALL CONTACT!	UST!	
•INHALATION	Cough. Shortness of brea	th.	Local exhaust or breathing protection.		Fresh air, rest.
•SKIN			Protective gloves. Protective clot	hing.	Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES			combination with breathing protection.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION			Do not eat, drink, or smoke during work.		Rinse mouth.
SDILLAGE DISPOSAL STODAGE DACKAG		CKACING & LARELLING			

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
Vacuum spilled material. Carefully collect	Separated from strong acids.	
remainder, then remove to safe place. Personal		Xn symbol
protection: P2 filter respirator for harmful		R: 40-43
particles.		S: 2-22-36

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0062

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

International Chemical Safety Cards

NICKEL ICSC: 0062

PHYSICAL STATE; APPEARANCE:

SILVERY METALLIC SOLID IN VARIOUS FORMS.

ROUTES OF EXPOSURE:

The substance can be absorbed into the body by inhalation of the dust.

Ι

PHYSICAL DANGERS:

M	Dust explosion possible if in powder or granular form, mixed with air.	INHALATION RISK: Even portion at 20°C is madicible a hormful concentration					
P		Evaporation at 20°C is negligible; a harmful concentration of airborne particles can, however, be reached quickly					
О	CHEMICAL DANGERS: Reacts violently, in powder form, with titanium powder	when dispersed.					
R	and potassium perchlorate, and oxidants such as ammonium nitrate, causing fire and explosion hazard. Reacts slowly with non-oxidizing acids and more rapidly	EFFECTS OF SHORT-TERM EXPOSURE: May cause mechanical irritation. Inhalation of fumes may cause pneumonitis.					
Т	with oxidizing acids. Toxic gases and vapours (such as nickel carbonyl) may be released in a fire involving nickel.	-					
A		EXPOSURE:					
N	OCCUPATIONAL EXPOSURE LIMITS: TLV: (Tabelella fraction)	Repeated or prolonged contact may cause skin sensitization. Repeated or prolonged inhalation exposure					
Т	(Inhalable fraction) 1.5 mg/m³ as TWA A5 (not suspected as a human carcinogen); (ACGIH 2004).	may cause asthma. Lungs may be affected by repeated or prolonged exposure. This substance is possibly carcinogenic to humans.					
D	MAK: (Inhalable fraction) sensitization of respiratory tract and skin (Sah);						
A	Carcinogen category: 1; (DFG 2004).						
Т	OSHA PEL*±: TWA 1 mg/m ³ *Note: The PEL does not apply to Nickel carbonyl.						
A	NIOSH REL*: Ca TWA 0.015 mg/m³ See Appendix A *Note: The REL does not apply to Nickel carbonyl. NIOSH IDLH: Ca 10 mg/m³ (as Ni) See: 7440020						
PHYSICAL PROPERTIES	Boiling point: 2730°C Melting point: 1455°C Density: 8.9 g/cm3	Solubility in water:					
ENVIRONMENTAL DATA							
	NOTES						
symptoms of asthma oft	ickel oxide fumes will be formed. Depending on the degree of ten do not become manifest until a few hours have passed and re essential. Anyone who has shown symptoms of asthma due	d they are aggravated by physical effort. Rest and medical					

ADDITIONAL INFORMATION ICSC: 0062 **NICKEL** (C) IPCS, CEC, 1994

IMPORTANT LEGAL NOTICE:

ZINC POWDER ICSC: 1205











Blue powder
Merrillite
Zn
Atomic mass: 65.4
(powder)

ICSC # 1205

CAS # 7440-66-6 RTECS # ZG8600000

UN # 1436 (zinc powder or dust)

EC# 030-001-00-1

October 24, 1994 Peer reviewed









·					
TYPES OF HAZARD/ EXPOSURE	ACUTE HAZA SYMPTON		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Highly flammable. Many reactions may cause fire or explosion. Gives off irritating or toxic fumes (or gases) in a fire. NO open flames, NO sparks, and NO smoking. NO contact with acid(s), base (s) and incompatible substances (see Chemical Dangers).		, base	Special powder, dry sand, NO other agents. NO water.	
EXPLOSION	Risk of fire and explosion on contact with acid(s), base(s), water and proof electrical equipment and lighting.		In case of fire: cool drums, etc., by spraying with water but avoid contact of the substance with water.		
EXPOSURE			PREVENT DISPERSION OF DU STRICT HYGIENE!	JST!	
•INHALATION	Metallic taste and metal fume fever. Symptoms may be delayed (see Notes).		Local exhaust.		Fresh air, rest. Refer for medical attention.
•SKIN	Dry skin.		Protective gloves.		Rinse and then wash skin with water and soap.
•EYES			Safety spectacles.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Abdominal pain. Nausea.			Rinse mouth. Refer for medical attention.	
SPILLAGE	SPILLAGE DISPOSAL STORAGE PACKAGING & LABELLING				

SPILLAGE DISPOSAL STORAGE PACKAGING & LABELLING Extinguish or remove all ignition sources. Do Fireproof. Separated from acids, bases oxidants Airtight. NOT wash away into sewer. Sweep spilled Dry. F symbol substance into containers. then remove to safe N symbol place. Personal protection: self-contained R: 15-17-50/53 S: 2-7/8-43-46-60-61 breathing apparatus. UN Hazard Class: 4.3 UN Subsidiary Risks: 4.2

SEE IMPORTANT INFORMATION ON BACK

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

ICSC: 1205 ZINC POWDER

I	PHYSICAL STATE; APPEARANCE: ODOURLESS GREY TO BLUE POWDER.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation					
M		and by ingestion.					
P	PHYSICAL DANGERS: Dust explosion possible if in powder or granular form,	INHALATION RISK:					
0	mixed with air. If dry, it can be charged electrostatically by swirling, pneumatic transport, pouring, etc.	Evaporation at 20°C is negligible; a harmful concentration of airborne particles can, however, be reached quickly when dispersed.					
R	CHEMICAL DANGERS:	when dispersed.					
Т	Upon heating, toxic fumes are formed. The substance is a strong reducing agent and reacts violently with oxidants. Reacts with water and reacts violently with acids and bases	EFFECTS OF SHORT-TERM EXPOSURE: Inhalation of fumes may cause metal fume fever. The effects may be delayed.					
A	forming flammable/explosive gas (hydrogen - see	enests may see detayed.					
N	ICSC0001) Reacts violently with sulfur, halogenated hydrocarbons and many other substances causing fire and	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE:					
Т	explosion hazard.	Repeated or prolonged contact with skin may cause dermatitis.					
	OCCUPATIONAL EXPOSURE LIMITS: TLV not established.						
D	TEV not established.						
A							
T							
A							
PHYSICAL	Boiling point: 907°C	Solubility in water: reaction					
PROPERTIES	Melting point: 419°C	Vapour pressure, kPa at 487°C: 0.1					
	Relative density (water = 1): 7.14	Auto-ignition temperature: 460°C					
ENVIRONMENTAL DATA							
NOTES							
Zinc may contain trace amounts of arsenic, when forming hydrogen, may also form toxic gas arsine (see ICSC 0001 and ICSC 0222). Reacts violently with fire extinguishing agents such as water, halons, foam and carbon dioxide. The symptoms of metal fume fever do not become manifest until several hours later. Rinse contaminated clothes (fire hazard) with plenty of water.							
	Transport Emergency Card: TEC (R)-43GWS-II+III NFPA Code: H0; F1; R1;						
ADDITIONAL INFORMATION							

IMPORTANT LEGAL NOTICE:

ICSC: 1205

Neither NIOSH, the CEC or the IPCS nor any person acting on behalf of NIOSH, the CEC or the IPCS is responsible for the use which might be made of this information. This card contains the collective views of the IPCS Peer Review Committee and may not reflect in all cases all the detailed requirements included in national legislation on the subject. The user should verify compliance of the cards with the relevant legislation in the country of use. The only modifications made to produce the U.S. version is inclusion of the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

(C) IPCS, CEC, 1994

ZINC POWDER

COPPER ICSC: 0240



ICSC: 0240









Cu (powder)

ICSC # 0240 CAS # 7440-50-8 RTECS # <u>GL5325000</u>

September 24, 1993 Validated

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ SYMPTO		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Combustible.		NO open flames.		Special powder, dry sand, NO other agents.
EXPLOSION					
EXPOSURE			PREVENT DISPERSION OF D	UST!	
•INHALATION	Cough. Headache. Short Sore throat.	ness of breath.	Local exhaust or breathing prote	ection.	Fresh air, rest. Refer for medical attention.
•SKIN	Redness.		Protective gloves.		Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES	Redness. Pain.		Safety goggles.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION	Abdominal pain. Nausea	. Vomiting.	Do not eat, drink, or smoke duri work.	ng	Rinse mouth. Refer for medical attention.
SPILLAGE DISPOSAL			STORAGE PA		ACKAGING & LABELLING
Sweep spilled substance into containers. Carefully collect remainder. Then remove to safe place. (Extra personal protection: P2 filter respirator for harmful particles).		Separated from	parated from - See Chemical Dangers. R: S:		
SEE IMPORTANT INFORMATION ON BACK					

International Chemical Safety Cards

NIOSH RELs and NIOSH IDLH values.

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Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs,

COPPER ICSC: 0240

Ţ	PHYSICAL STATE; APPEARANCE: RED POWDER, TURNS GREEN ON EXPOSURE TO MOIST AIR.	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation and by ingestion.
M	PHYSICAL DANGERS:	INHALATION RISK: Evaporation at 20°C is negligible; a harmful concentration
P	CHEMICAL DANGERS:	of airborne particles can, however, be reached quickly when dispersed.

1					
0	Shock-sensitive compounds are formed with acetylenic				
	compounds, ethylene oxides and azides. Reacts with strong	EFFECTS OF SHORT-TERM EXPOSURE:			
R	oxidants like chlorates, bromates and iodates, causing	Inhalation of fumes may cause metal fume fever. See			
	explosion hazard.	Notes.			
T					
	OCCUPATIONAL EXPOSURE LIMITS:	EFFECTS OF LONG-TERM OR REPEATED			
A	TLV: 0.2 mg/m ³ fume (ACGIH 1992-1993).	EXPOSURE:			
N	TLV (as Cu, dusts & mists): 1 mg/m ³ (ACGIH 1992-1993).				
IN .	Intended change 0.1 mg/m ³	sensitization.			
T	Inhal.,				
1	A4 (not classifiable as a human carcinogen); MAK: 0.1 mg/m³ (Inhalable fraction)				
	Peak limitation category: II(2) Pregnancy risk group: D				
D	(DFG 2005).				
	OSHA PEL*: TWA 1 mg/m ³ *Note: The PEL also applies				
A	to other copper compounds (as Cu) except copper fume.				
	NIOSH REL*: TWA 1 mg/m ³ *Note: The REL also				
T	applies to other copper compounds (as Cu) except Copper				
	fume.				
A	NIOSH IDLH: 100 mg/m ³ (as Cu) See: 7440508				
	1410511 1DE11. 100 mg/m (as cu) sec. 7440308				
	Boiling point: 2595°C	Solubility in water:			
PHYSICAL	Melting point: 1083°C	none			
PROPERTIES	Relative density (water = 1): 8.9				
ENVIRONMENTAL					
DATA					
	NOTES				
The symptoms of metal fume fever do not become manifest until several hours.					
The symptoms of meta	Tunic level do not become maintest until several nours.				
	ADDITIONAL INFORMA	ΓΙΟΝ			
ICSC: 0240		COPPER			

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CHROMIUM ICSC: 0029











Chrome Cr Atomic mass: 52.0 (powder)

ICSC # 0029 CAS # 7440-47-3 RTECS # <u>GB4200000</u>

October 27, 2004 Peer reviewed

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Combustible under specific conditions.				In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION			Prevent deposition of dust; closed system, dust explosion-proof electrical equipment and lighting.		
EXPOSURE			PREVENT DISPERSION OF D	UST!	
•INHALATION	Cough.		Local exhaust or breathing protection	ction.	Fresh air, rest.
•SKIN			Protective gloves.		Remove contaminated clothes. Rinse skin with plenty of water or shower.
•EYES	Redness.		Safety goggles.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION			Do not eat, drink, or smoke durir work.	ng	Rinse mouth.
SPILLAGE DISPOSAL		STORAGE	PA	ACKAGING & LABELLING	
Sweep spilled substant appropriate, moisten fi Personal protection: Pa harmful particles.	rst to prevent dusting. 2 filter respirator for		NT INFORMATION ON BAC	R: S:	

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0029

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International Chemical Safety Cards

CHROMIUM ICSC: 0029

_	PHYSICAL STATE; APPEARANCE:
1	GREY POWDER

M PHYSICAL DANGERS:

P Dust explosion possible if in powder or granular form, mixed with air.

ROUTES OF EXPOSURE:

INHALATION RISK:

A harmful concentration of airborne particles can be reached quickly when dispersed.

l					
0					
R	CHEMICAL DANGERS: Chromium is a catalytic substance and may cause reaction	EFFECTS OF SHORT-TERM EXPOSURE: May cause mechanical irritation to the eyesand the			
Т	in contact with many organic and inorganic substances, causing fire and explosion hazard.	respiratory tract. EFFECTS OF LONG-TERM OR REPEATED			
A	OCCUPATIONAL EXPOSURE LIMITS: TLV: (as Cr metal, Cr(III) compounds) 0.5 mg/m³ as TWA	EXPOSURE:			
N	A4 (ACGIH 2004). MAK not established.				
Т	OSHA PEL*: TWA 1 mg/m ³ See Appendix C *Note: The PEL also applies to insoluble chromium salts.				
D	NIOSH REL: TWA 0.5 mg/m ³ See Appendix C NIOSH IDLH: 250 mg/m ³ (as Cr) See: 7440473				
A	-				
Т					
A					
PHYSICAL PROPERTIES	Boiling point: 2642°C Melting point: 1900°C Density: 7.15 g/cm ³	Solubility in water: none			
ENVIRONMENTAL DATA					
	NOTES				
The surface of the chromium particles is oxidized to chromium(III)oxide in air. See ICSC 1531 Chromium(III) oxide.					
	ADDITIONAL INFORMA	TION			
ICSC: 0029		CHROMIUM			

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ICSC: 0056 **MERCURY**











Quicksilver Liquid silver Hg Atomic mass: 200.6

ICSC# 0056

CAS# 7439-97-6 RTECS # OV4550000

UN# 2809

EC# 080-001-00-0 April 22, 2004 Peer reviewed









TYPES OF HAZARD/ EXPOSURE	ACUTE HAZA SYMPTOM		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Not combustible. Gives of toxic fumes (or gases) in				In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION	Risk of fire and explosion	1.			In case of fire: keep drums, etc., cool by spraying with water.
EXPOSURE			STRICT HYGIENE! AVOID EXPOSURE OF (PREGNANT) WOMEN! AVOID EXPOSURE ADOLESCENTS AND CHILD	OF	IN ALL CASES CONSULT A DOCTOR!
•INHALATION	Abdominal pain. Cough. Shortness of breath. Vom or elevated body temperat	iting. Fever	Local exhaust or breathing prote	ction.	Fresh air, rest. Artificial respiration if indicated. Refer for medical attention.
•SKIN	II II		Protective gloves. Protective clo	thing.	Remove contaminated clothes. Rinse and then wash skin with water and soap. Refer for medical attention.
•EYES			combination with breathing protection.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.
•INGESTION			Do not eat, drink, or smoke during work. Wash hands before eating.		Refer for medical attention.

SPILLAGE DISPOSAL **STORAGE** PACKAGING & LABELLING Provision to contain effluent from fire Evacuate danger area in case of a large spill! Special material. Do not transport with food and feedstuffs. Consult an expert! Ventilation. Collect leaking extinguishing. Separated from food and and spilled liquid in sealable non-metallic feedstuffs Well closed. T symbol containers as far as possible. Do NOT wash N symbol away into sewer. Do NOT let this chemical R: 23-33-50/53 enter the environment. Chemical protection S: 1/2-7-45-60-61 suit including self-contained breathing UN Hazard Class: 8 apparatus. UN Packing Group: III

SEE IMPORTANT INFORMATION ON BACK

Prepared in the context of cooperation between the International Programme on Chemical Safety & the Commission of the ICSC: 0056 European Communities (C) IPCS CEC 1994. No modifications to the International version have been made except to add the OSHA PELs, NIOSH RELs and NIOSH IDLH values.

MERCURY ICSC: 0056

I .	Ú-						
I	PHYSICAL STATE; APPEARANCE: ODOURLESS, HEAVY AND MOBILE SILVER	ROUTES OF EXPOSURE: The substance can be absorbed into the body by inhalation					
M	LIQUID METAL.	of its vapour and through the skin, also as a vapour!					
P	PHYSICAL DANGERS:	INHALATION RISK: A harmful contamination of the air can be reached very					
О	CHEMICAL PANCEDS	quickly on evaporation of this substance at 20°C.					
R	CHEMICAL DANGERS: Upon heating, toxic fumes are formed. Reacts vio with ammonia and halogens causing fire and expl						
T	hazard. Attacks aluminium and many other metals forming amalgams.	vapours may cause pneumonitis. The substance may cause effects on the central nervous systemandkidneys. The					
A	OCCUPATIONAL EXPOSURE LIMITS:	effects may be delayed. Medical observation is indicated.					
N	TLV: 0.025 mg/m ³ as TWA (skin) A4 BEI issued (ACGIH 2004).	EFFECTS OF LONG-TERM OR REPEATED EXPOSURE:					
Т	MAK: 0.1 mg/m³ Sh Peak limitation category: II(8) Carcinogen catego (DFG 2003).	The substance may have effects on the central nervous					
D	OSHA PEL‡: C 0.1 mg/m ³	speech disorders. Danger of cumulative effects. Animal					
A	NIOSH REL: Hg Vapor: TWA 0.05 mg/m ³ skin Other: C 0.1 mg/m ³ skin	tests show that this substance possibly causes toxic effects upon human reproduction.					
T	NIOSH IDLH: 10 mg/m ³ (as Hg) See: <u>7439976</u>						
A							
PHYSICAL PROPERTIES	Boiling point: 357°C Melting point: -39°C Relative density (water = 1): 13.5 Solubility in water: none	Vapour pressure, Pa at 20°C: 0.26 Relative vapour density (air = 1): 6.93 Relative density of the vapour/air-mixture at 20°C (air = 1): 1.009					
ENVIRONMENTAL DATA	litakes place specifically in fish						
NOTES							
Depending on the degree of exposure, periodic medical examination is indicated. No odour warning if toxic concentrations are present. Do NOT take working clothes home. Transport Emergency Card: TEC (R)-80GC9-II+III							
ADDITIONAL INFORMATION							

IMPORTANT LEGAL NOTICE:

ICSC: 0056

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MERCURY

LEAD ICSC: 0052











Lead metal Plumbum Pb Atomic mass: 207.2 (powder)

ICSC # 0052 CAS # 7439-92-1 RTECS # <u>OF7525000</u>

October 08, 2002 Peer reviewed

TYPES OF HAZARD/ EXPOSURE	ACUTE HAZ SYMPTO		PREVENTION		FIRST AID/ FIRE FIGHTING
FIRE	Not combustible. Gives or toxic fumes (or gases				In case of fire in the surroundings: use appropriate extinguishing media.
EXPLOSION	Finely dispersed particle explosive mixtures in air		Prevent deposition of dust; close system, dust explosion-proof electrical equipment and lightin		
EXPOSURE	See EFFECTS OF LON REPEATED EXPOSUI		OR PREVENT DISPERSION OF DUST! AVOID EXPOSURE OF (PREGNANT) WOMEN!		
•INHALATION			Local exhaust or breathing prote	ection.	Fresh air, rest.
•SKIN			Protective gloves.		Remove contaminated clothes. Rinse and then wash skin with water and soap.
•EYES		Safety spectacles.		First rinse with plenty of water for several minutes (remove contact lenses if easily possible), then take to a doctor.	
•INGESTION	Abdominal pain. Nausea. Vomiting. Do not eat, drink, or smoke during work. Wash hands before eating.		Rinse mouth. Give plenty of water to drink. Refer for medical attention.		
SPILLAGE DISPOSAL		STORAGE	PA	CKAGING & LABELLING	
Sween spilled substan	gen spilled substance into containers: if Separated from food and feedstuffs				

SPILLAGE DISPOSAL	STORAGE	PACKAGING & LABELLING
,	Separated from food and feedstuffs incompatible materials See Chemical Dangers.	R: S:

SEE IMPORTANT INFORMATION ON BACK

ICSC: 0052

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International Chemical Safety Cards

LEAD ICSC: 0052

PHYSICAL STATE; APPEARANCE: **ROUTES OF EXPOSURE:** BLUISH-WHITE OR SILVERY-GREY SOLID IN The substance can be absorbed into the body by VARIOUS FORMS. TURNS TARNISHED ON inhalation and by ingestion. EXPOSURE TO AIR. Ι INHALATION RISK: PHYSICAL DANGERS: A harmful concentration of airborne particles can be M Dust explosion possible if in powder or granular form, reached quickly when dispersed, especially if powdered. mixed with air. P EFFECTS OF SHORT-TERM EXPOSURE: CHEMICAL DANGERS: \mathbf{o} On heating, toxic fumes are formed. Reacts with oxidants. Reacts with hot concentrated nitric acid, EFFECTS OF LONG-TERM OR REPEATED R boiling concentrated hydrochloric acid and sulfuric acid. **EXPOSURE:** Attacked by pure water and by weak organic acids in the The substance may have effects on the blood bone \mathbf{T} presence of oxygen. marrow central nervous system peripheral nervous system kidneys, resulting in anaemia, encephalopathy OCCUPATIONAL EXPOSURE LIMITS: (e.g., convulsions), peripheral nerve disease, abdominal TLV: 0.05 mg/m³ A3 (confirmed animal carcinogen cramps and kidney impairment. Causes toxicity to N with unknown relevance to humans); BEI issued human reproduction or development. (ACGIH 2004). T MAK: Carcinogen category: 3B; Germ cell mutagen group: 3A; (DFG 2004). D EU OEL: as TWA 0.15 mg/m³ (EU 2002). OSHA PEL*: 1910.1025 TWA 0.050 mg/m³ See Appendix C *Note: The PEL also applies to other lead compounds (as Pb) -- see Appendix C. NIOSH REL*: TWA 0.050 mg/m³ See Appendix C *Note: The REL also applies to other lead compounds A (as Pb) -- see Appendix C. NIOSH IDLH: 100 mg/m³ (as Pb) See: 7439921 Boiling point: 1740°C Density: 11.34 g/cm3 **PHYSICAL** Solubility in water: none **PROPERTIES** Melting point: 327.5°C Bioaccumulation of this chemical may occur in plants and in mammals. It is strongly advised that this ENVIRONMENTAL substance does not enter the environment. DATA NOTES Depending on the degree of exposure, periodic medical examination is suggested. Do NOT take working clothes home. Transport Emergency Card: TEC (R)-51S1872 ADDITIONAL INFORMATION

IMPORTANT LEGAL NOTICE:

ICSC: 0052

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LEAD

APPENDIX D HOSPITAL INFORMATION AND MAP FIELD ACCIDENT REPORT

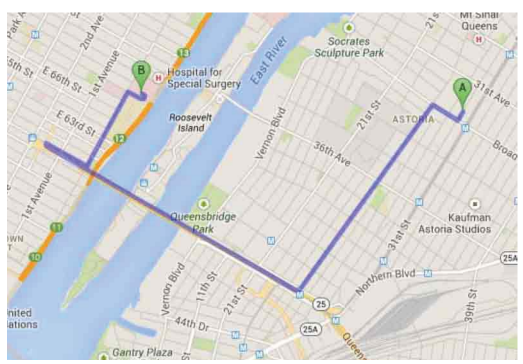
HOSPITAL INFORMATION AND MAP

The nearest emergency room to the site is:

NY Presbyterian/Weill Cornell Medical Center 525 E. 68th Street, New York, NY 10065

525 E. 68th Street, New York, NY 10065 (212) 746-5454

3.4 Miles – About 10 Minutes

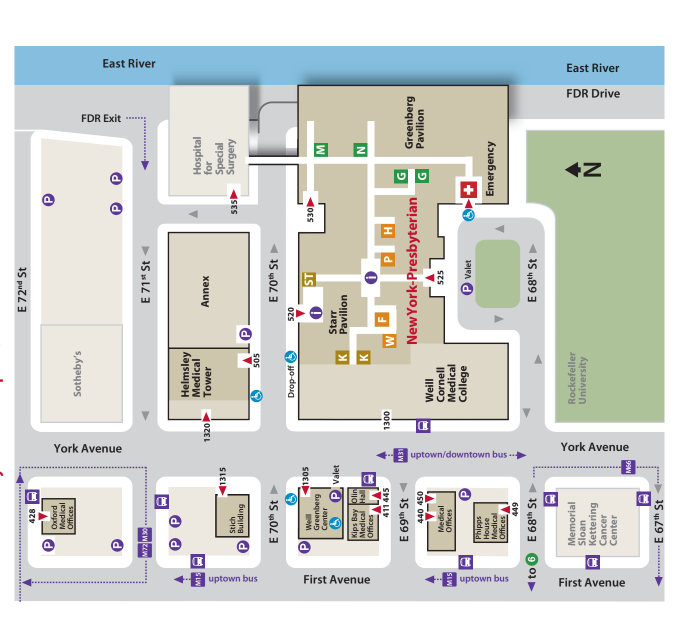




31-70 30th St, New York, NY 11106

	1. Head southwest on 30th St toward Broadway	go 256 ft total 256 ft
Ļ	2. Take the 1st right onto Broadway	go 0.1 mi total 0.2 mi
٦	 Turn left at the 2nd cross street onto Crescent St About 3 mins 	go 1.1 mi total 1.3 mi
25)	 Take the ramp onto New York 25 W/Ed Koch Queensboro Bridge About 3 mins 	go 1.2 mi total 2.4 mi
	5. Take the exit toward E 60th St	go 390 ft total 2.5 mi
Ļ	6. Keep right at the fork, follow signs for 1 Avenue N/FDR Drive	go 0.1 mi total 2.6 mi
Ļ	7. Turn right onto E 60th St	go 0.2 mi total 2.8 mi
٦	Take the 2nd left onto York Ave About 1 min	go 0.4 mi total 3.2 mi
Ļ	9. Turn right onto E 68th St	go 433 ft total 3.3 mi
4	10. Turn left to stay on E 68th St Destination will be on the left	go 180 ft total 3.4 mi
B	525 E 68th St, New York, NY 10065	

NewYork-Presbyterian Hospital/Weill Cornell Medical Center



Visitor's Entrance

Accessible Entrance

Emergency

Information Desk

Subway Entrance Visitor Parking

(I)

Elevators

Greenberg Pavilion

Starr Pavilion, J Corridor & L Corridor S

K Wing & F Wing, floors 2-9 ¥

Baker Pavilion / F Wing, floors 9-24

Baker Pavilion / F Wing, floors 9-23 I

Payson Pavilion a Whitney Pavilion

M Wing 3 Z Z

N Wing

Additional Medical Offices

Weill Cornell Medical Assoc. Eastside

Weill Cornell Medical Assoc. Westside

Iris Cantor Women's Health Center 12 West 72nd Street 425 East 61st Street

at NewYork-Presbyterian Weill Cornell Imaging

520 East 70th Street, lobby level 425 East 61st Street, 9th Floor 1305 York Avenue, 3rd Floor 416 East 55th Street

Hospital information: 212 746 5454

FIELD ACCIDENT REPORT

This report is to be filled out by the designated Site Safety Officer after EVERY accident.

PROJECT NAME	PROJECT. NO							
Date of Accident T	Time Report By							
Type of Accident (Check One):								
() Vehicular () Person	nal () Property							
Name of Injured	DOB or Age							
How Long Employed								
Names of Witnesses								
Description of Accident								
Action Taken								
Did the Injured Lose Any Time? F	How Much (Days/Hrs.)?							
Was Safety Equipment in Use at the Tim Shoes, etc.)?		ty						
(If not, it is the EMPLOYEE'S sole response Welfare Fund.) INDICATE STREET NAMES, DESCRIPTIO	onsibility to process his/her claim through his/her Health at	nd						

ATTACHMENT C Community Air Monitoring Plan

COMMUNITY AIR MONITORING PLAN

FORMER UNION WIRE DIE CORP. 39-40 30th STREET LONG ISLAND CITY

MARCH - 2014

FORMER UNION WIRE DIE CORP.

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APPENDICES

Appendix A Action Limit Report

This Community Air Monitoring Plan (CAMP) has been prepared for the drilling and sampling activities to be performed under a Remedial Investigation Work Plan (RIWP) at the Former Union Wire Die Corp. Site. The CAMP provides measures for protection for the downwind community (i.e., off-site receptors including residences, businesses, and on-site workers not directly involved in the investigation activities) from potential airborne contaminant releases resulting from investigative activities at the site.

Compliance with this CAMP is required during all activities associated with drilling and sampling activities that have the potential to generate airborne particulate matter and volatile organic compounds (VOCs). These activities include drilling and soil and groundwater sampling. This CAMP has been prepared to ensure that investigation activities do not adversely affect passersby, residents, or workers in the area immediately surrounding the Site and to preclude or minimize airborne migration of investigation-related contaminants to off-site areas.

1.1 **Regulatory Requirements**

This CAMP was established in accordance with the following requirements:

- New York State Department of Health's (NYSDOH) Generic Community Air Monitoring Plan as presented in DER-10 Technical Guidance for Site Investigation and Remediation (NYSDEC May 3, 2010). This guidance specifies that a community air-monitoring program shall be implemented to protect the surrounding community and to confirm that the work does not spread contamination off-site through the air;
- New York State Department of Environmental Conservation (NYSDEC) Technical and Guidance Memorandum (TAGM) #4031 - Fugitive Dust Suppression and Particulate Monitoring Program at Inactive Hazardous Waste Sites: This guidance provides a basis for developing and implementing a fugitive dust suppression and particulate monitoring program as an element of a hazardous waste site's health and safety program.

2.0 AIR MONITORING

Chlorinated volatile organic compounds (VOCs) and petroleum VOCs are the constituents of concern at the Site. The appropriate method to monitor air for these constituents during remediation activities is through real-time VOC and air particulate (dust) monitoring.

2.1 **Meteorological Data**

At a minimum, wind direction will be evaluated at the start of each workday, noon of each workday, and the end of each workday. These readings will be utilized to position the monitoring equipment in appropriate upwind and downwind locations.

2.2 **Community Air Monitoring Requirements**

To establish ambient air background concentrations, air will be monitored at several locations around the site perimeter before activities begin. These points will be monitored periodically in series during the site work. When the drilling area is within 20 feet of potentially exposed populations or occupied structures, the perimeter monitoring points will be located to represent the nearest potentially exposed individuals at the downwind location.

Fugitive respirable dust will be monitored using a MiniRam Model PDM-3 aerosol monitor (or equivalent). Air will be monitored for VOCs with a portable Ionscience 3000 photoionization detector (PID), or equivalent. All air monitoring data will be documented in a site log book by the designated site safety officer. The site safety officer or delegate must ensure that air monitoring instruments are calibrated and maintained in accordance with manufacturer's specifications. All instruments will be zeroed daily and checked for accuracy. A daily log will be kept. If additional monitoring is required, the protocols will be developed and appended to this plan.

3.0 **VOC MONITORING, RESPONSE LEVELS, AND ACTIONS**

Volatile organic compounds (VOCs) will be monitored at the downwind perimeter of the immediate work area (i.e., the exclusion 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 parts per million (ppm) above background for the 15minute average, work activities must be temporarily halted and monitoring 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 of 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. All 15-minute readings must be recorded and be available for State (DEC and DOH) personnel to review. Instantaneous readings, if any, used for decision purposes should also be recorded.

All readings will be recorded and made available for NYSDEC and NYSDOH personnel to review. If an exceedance of the Action Limits occurs, an Action Limit Report, as shown in **Appendix A**, will be completed.

Potential Corrective Measures and VOC Suppression Techniques 3.1

If the 15-minute integrated VOC level at the downwind location persists at a concentration that exceeds the upwind level by more than 5 ppm but less than 25 ppm during remediation activities, then vapor suppression techniques will be employed. The following techniques, or others, may be employed to mitigate the generation and migration of fugitive organic vapors:

- Collection of purge water in covered containers;
- storage of excess sample and drill cuttings in drums or covering with plastic



4.0 PARTICULATE MONITORING

Air monitoring for particulates (i.e., dust) will be performed continuously during drilling activities using both air monitoring equipment and visual observation at upwind and downwind locations. Monitoring equipment capable of measuring particulate matter smaller than 10 microns (PM₁₀) and capable of integrating (averaging) over periods of 15 minutes or less will be set up at upwind (i.e., background) and downwind locations, at heights approximately four to five feet above land surface (i.e., the breathing zone). Monitoring equipment will be MIE Data Ram monitors, or equivalent. The audible alarm on the particulate monitoring device will be set at 90 micrograms per cubic meter (μg/m₃). This setting will allow proactive evaluation of worksite conditions prior to reaching the action level of 100 µg/m³ above background. The monitors will be calibrated at least once per day prior to work activities and recalibrated as needed thereafter. In addition, fugitive dust migration will be visually assessed during all intrusive work activities.

The following summarizes particulate action levels and the appropriate responses:

- If the downwind PM-10 particulate level is 100 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 an evaluation of activities initiated. Work can resume provided that dust suppression measures (as described in Section 2.3.1 below) 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.

All readings will be recorded and be available for NYSDEC and NYSDOH personnel to review. If an exceedance of the Action Limits occurs, an Action Limit Report as shown in **Appendix A** will be completed.

4.1 **Potential Particulate Suppression Techniques**

If the integrated particulate level at the downwind location exceeds the upwind level by more than 100 μg/m₃ at any time during drilling activities, then dust suppression techniques will be employed. The following techniques, or others, may be employed to mitigate the generation and migration of fugitive dusts:

- Placement of drill cuttings in drums or covering stockpiles with plastic;
- Misting of the drilling area with a fine water spray from a hand-held spray bottle

Work may continue with dust suppression techniques provided that downwind PM₁₀ levels are not more than 150 µg/m³ greater than the upwind levels.



There may also be situations where the dust is generated by drilling activities and migrates to downwind locations, but is not detected by the monitoring equipment at or above the action level. Therefore, if dust is observed leaving the working area, dust suppression techniques such as those listed above will be employed.

If dust suppression techniques do not lower particulates to below 150 μg/m³, or visible dust persists, work will be suspended until appropriate corrective measures are identified and implemented to remedy the situation.

All air monitoring readings will be recorded in the field logbook and will be available for the NYSDEC and NYSDOH personnel to review.

5.0 DATA QUALITY ASSURANCE

5.1 Calibration

Instrument calibration shall be documented on instrument calibration and maintenance sheets or in the designated field logbook. All instruments shall be calibrated as required by the manufacturer. Calibration checks may be used during the day to confirm instrument accuracy. Duplicate readings may be taken to confirm individual instrument response.

5.2 **Operations**

All instruments shall be operated in accordance with the manufacturer's specifications. Manufacturers' literature, including an operations manual for each piece of monitoring equipment will be maintained on-site by the SSO for reference.

5.3 **Data Review**

The SSO will interpret all monitoring data based the established criteria and his/her professional judgment. The SSO shall review the data with the PM to evaluate the potential for worker exposure, upgrades/downgrades in level of protection, comparison to direct reading instrumentation and changes in the integrated monitoring strategy.

Monitoring and sampling data, along with all sample documentation will be periodically reviewed by the PM.

RECORDS AND REPORTING **6.0**

All air readings must be recorded on daily air monitoring log sheets and made available for review by personnel from NYSDEC and NYSDOH.

APPENDIX A

CAMP ACTION LIMIT REPORT

Project Location:			
Date:	_	Work Start Time:	
Name:	_	Work Glop Time.	
Contaminant:	PM-10:	VOC:	
Wind Speed: AM PM	Wind Dir	rection: AM	PM
Temperature: AM PM		Barometric Pressure	e:
Monitor ID#:			
UPWIND DATA Monitor ID #:			
Monitor ID#:	Location:	Level Reported:	
BACKGROUND CORRECTED LE	EVELS		
Monitor ID #: Location:	Level Reported:		
ACTIONS TAKEN			