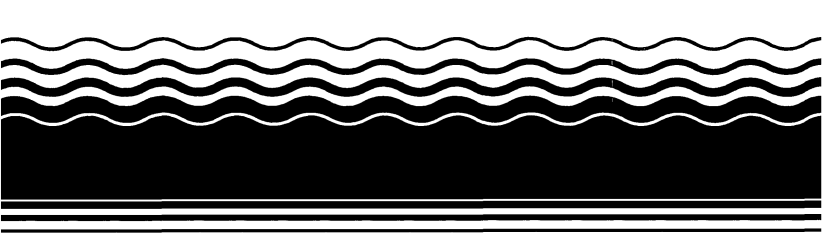
# **SEPA** Superfund Record of Decision:

Pasley Solvents & Chemical, NY



# NOTICE The appendices listed in the index that are not found in this document have been removed at the request of the issuing agency. They contain material which supplement, but adds no further applicable information to the content of the document. All supplemental material is, however, contained in the administrative record for this site.

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15. Supplementary Notes

PB93-963805

### 16. Abstract (Limit: 200 words)

The approximately 0.5-acre Pasley Solvents and Chemical site is a former oil, solvent, and chemical storage tank farm located in the Town of Hempstead, Nassau County, New York. Land use in the area is predominantly industrial, with some commercial and residential areas within one-fourth mile of the site. The only drinking water source in the area consists of aquifers; four public water supply wellfields are located within 2 miles of the site. Prior to 1969, Commander Oil Corporation distributed fuel oil at the site. From 1969 to 1982, Pasley Solvents and Chemicals Company distributed chemicals and used the site for storage of waste and sludge scavenger (Pasley) operations. In 1980, after Pasley requested a chemical storage and removal permit, a state inspection revealed soil contamination with VOCs beneath the above-ground storage tanks and recommended a remedial investigation and clean-up plan. In 1981, ground water monitoring wells were installed and revealed that the ground water was also contaminated with VOCs. All operations onsite ceased in 1982. In 1988, all 12 above-ground storage tanks were removed by the site owners under EPA supervision. A separate site, upgradient from the Pasley facility, was shown to be contributing to the background contamination of the ground water. This ROD addresses the contaminated

(See Attached Page)

### 17. Document Analysis a. Descriptors

Record of Decision - Pasley Solvents & Chemical, NY

First Remedial Action - Final Contaminated Media: soil, gw

Key Contaminants: VOCs (benzene, PCE, TCE, toluene, xylenes) and other organics

b. Identifiers/Open-Ended Terms (naphthalene, PAHs)

c. COSATI Field/Group

a continuation			
18. Availability Statement	19. Security Class (This Report)	21. No. of Pages	
•	None	100	
	20. Security Class (This Page)	22. Price	
	None		

EPA/ROD/RO2-92/171
Pasley Solvents & Chemical, NY
First Remedial Action - Final

### Abstract (Continued)

surface soil and ground water as a final remedy. The primary contaminants of concern affecting the soil and ground water are VOCs, including benzene, PCE, TCE, toluene, and xylenes; and other organics, including PAHs and naphthalene.

The selected remedial action at this site includes soil vacuum extraction for removal of aliphatic hydrocarbons and, as necessary, soil flushing to remove VOCs and semi-VOCs of 13,000 cubic yards of soil with offsite residual disposal, and ground water extraction for onsite treatment with metals precipitation and air stripping followed by granular activated carbon (GAC) polishing of treated water with onsite discharge to the aquifer or to infiltration trenches. Off-gas emissions from the air stripper will be treated by passing the airstream through vapor phase adsorption columns. The estimated present net worth cost for this remedy is \$13,744,000, including an annual O&M cost of \$1,236,000 for a 6-year soil remediation period and a 10- to 40-year ground water treatment operation.

PERFORMANCE STANDARDS OR GOALS: Chemical-specific soil clean-up goals are based on SDWA MCLs to protect ground water by the reduction of mobility and volume of contaminants and include PCE 1.4 mg/kg, TCE 0.7 mg/kg, toluene 1.5 mg/kg, and xylenes 1.2 mg/kg. Chemical-specific ground water clean-up goals may be waived because of the presence of an upgradient contamination source from another site. Clean-up goals are based on the upgradient concentrations of certain contaminants. When the upgradient source has been remediated, drinking water standards, such as SDWA MCLs and state ground water quality regulations, will be assigned.

### **DECLARATION FOR THE RECORD OF DECISION**

### SITE NAME AND LOCATION

Pasley Solvents and Chemicals Site Town of Hempstead Nassau County, New York

### STATEMENT OF BASIS AND PURPOSE

This decision document presents the selected remedial action for the Pasley Solvents and Chemical Site (Site), which was chosen in accordance with the requirements of the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA), as amended by the Superfund Amendments and Reauthorization Act of 1986 (SARA), and the National Oil and Hazardous Substances Pollution Contingency Plan (NCP). This decision document summarizes the factual and legal basis for selecting the remedy for this Site.

The New York State Department of Environmental Conservation (NYSDEC) concurs with the selected remedy. A letter of concurrence from NYSDEC is appended to this document.

The information supporting this remedial action decision is contained in the administrative record for this Site, an index of which is attached as Appendix 5.

### ASSESSMENT OF THE SITE

Actual or threatened releases of hazardous substances from this Site, if not addressed by implementing the response action selected in this Record of Decision, may present an imminent and substantial endangerment to public health, welfare, or the environment.

### **DESCRIPTION OF SELECTED REMEDY**

The remedy presented in this document addresses the treatment of soils and the ground water at the Pasley Solvents and Chemicals Site.

The major components of the selected remedy include:

- Treatment of approximately thirteen thousand (13,000) cubic yards of contaminated soil by soil vacuuming and/or by soil flushing;
- Disposal of treatment residuals at a RCRA Subtitle C facility;
- Remediation of the ground water by extraction/metals precipitation/air stripping with vapor phase granular activated carbon/GAC polishing/recharge;

### ROD FACT SHEET

### SITE

Name:

Pasley Solvents

Location/State:

Hempstead, Nassau Co., New York

EPA Region:

HRS Score (date):

39.65 (6/86)

NPL Rank (date):

510

ROD

Date Signed:

April 24, 1992

Selected Remedy

Groundwater:

Pump and treat (air stripping with vapor phase GAC of off-gas followed by GAC polishing of treated water, if necessary) groundwater with goal of achieving ARARs. Treated groundwater to be recharged if

necessary.

Capital Cost:

\$ 4,280,000 \$ 829,000

O & M: Present Worth:

\$ 12,095,000

Soil:

Soil vacuuming; soil flushing with treated

groundwater if necessary.

Capital Cost:

\$ 921,000

0 & M:

\$ 407,000

Present Worth:

\$ 1,649,000

LEAD

Enforcement, PRP Lead

Primary Contact (phone): Sherrel Henry (212-264-8675)
[4/1/92 thru 10/1/92]>> Mark Granger (212-264-9588)

Secondary Contact (phone): Melvin Hauptman (212-264-7681)

Main PRP:

Commander Oil Co.

PRP Contact (phone):

Gregory Pikul (908-685-4762)

WASTE

Type:

Groundwater -- VOCs and Semi-VOCs.

Soil

-- VOCs and Semi-VOCs.

Medium:

Groundwater, soil.

Origin:

Up until 1969 the Site was operated as a fuel oil distribution facility. The Site was a chemical distribution and storage

facility for oils and solvents from

1969-1982.

- Pumping of contaminated ground water from three extraction wells at combined flow rate of approximately 450 gpm. The actual pumping rate will be determined during the Remedial Design;
- Implementation of a long-term monitoring program to track the migration and concentrations of the contaminants of concern; and
- Implementation of a system monitoring program that includes the collection and analysis of the influent and effluent from the treatment systems and periodic collection of well-head samples.

### **DECLARATION OF STATUTORY DETERMINATIONS**

This selected remedy is protective of human health and the environment, complies with Federal and State requirements that are legally applicable or relevant and appropriate to the remedial action, and is cost effective. This remedy utilizes permanent solutions and alternative treatment technologies to the maximum extent practicable for this Site. Because treatment is being used to address the principal threats at the Site, this remedy satisfies the statutory preference for treatment as a principal element of the remedy.

Due to the existence of an upgradient source of contamination, the selected ground water remedy, by itself, will not meet chemical-specific ARARs nor be capable of restoring the area ground water to applicable ground water quality standards until these upgradient source areas are removed.

As the remedy will result in hazardous substances remaining on site above health-based levels, a review will be conducted within five (5) years after commencement of the remedial action, and every five years thereafter, to ensure that the remedy continues to provide adequate protection of human health and the environment

Constantine Sidamon-Eristoff

Regional Administrator

Date

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### **DECISION SUMMARY**

# PASLEY SOLVENTS AND CHEMICALS SITE TOWN OF HEMPSTEAD, NEW YORK

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY
REGION II
NEW YORK

### I. SITE LOCATION AND DESCRIPTION

The Pasley Solvents and Chemicals Site (Site) is located at 556 Commercial Avenue, Town of Hempstead, Nassau County, New York. The Site lies between the borders of the political subdivisions of the Village of Garden City and Uniondale, in the Town of Hempstead (see Figure 1). The immediate area has light industrial and commercial properties; residential communities are located within 1/4 mile of the Site. The Site measures 75' by 275', and is fenced on the north, east and south. A building and loading platform border the Site to the west (see Figure 2).

According to the Town of Hempstead's Public Information Division, the population of the Town of Hempstead is approximately 735,000. The predominant form of land use in the vicinity is industrial with the nearest off-site building adjacent to the Site. It is estimated that 75 homes are located within a 1/4 mile radius of the Site and 1,800 homes within one mile of the Site. The only source of drinking water for residences in the Town of Hempstead is ground water. All public water supply wells in the Site area draw water from the deeper aquifer, the Magothy Aquifer. Four public water supply well fields are located within approximately 2 miles of the Site.

There are no surface water bodies or wetlands within the vicinity of the Site. There is no designated New York State Significant Habitat, agricultural land, historic or landmark site directly or potentially affected. There are no endangered species or critical habitats within close proximity to the Site.

### II. SITE HISTORY AND ENFORCEMENT ACTIVITIES

### A. <u>Site History</u>

The Site is a former tank farm used for the storage of oils, solvents and chemicals. From 1969 to 1982 the Site was occupied by Pasley Solvents and Chemicals Company (Pasley) and was used as a chemical distribution facility. The principal activity at the Site included the delivery of various chemicals to the Site, storage of chemicals in the tanks located there and eventual transfer of the chemicals to 55-gallon drums for delivery to customers. These chemicals reportedly included a wide range of aromatic and halogenated aliphatic hydrocarbons, various solvents, ketones and alcohols. Pasley also operated as a "scavenger" that transported waste and sludge, containing hazardous substances that may have been transported to the Site. The Site is owned by Commander Oil Corporation (Commander). Prior to 1969, the Site was occupied by Commander, which distributed fuel oils.

In response to Pasley's request for a New York State Department of Environmental Conservation (NYSDEC) permit to store and remove chemicals, the Nassau County Department of Health (NCDH) conducted a preliminary site inspection in 1980 and collected soil samples from the area beneath the above-ground storage tanks at depths ranging from six to 36 inches. The soil collected was contaminated with halogenated and non-halogenated hydrocarbons, including trichloroethene (TCE), tetrachloroethene (PCE), 1,1,1-trichloroethane, xylene and

toluene. These chemicals were being stored on-site at the time. NCDH then referred the Site to NYSDEC. NCDH and NYSDEC recommended that Pasley submit a plan for a Phase I and Phase II remedial investigation and a cleanup plan.

Lakeland Engineering of Port Washington (Lakeland), New York was hired by Pasley to perform a limited well drilling and ground water sampling program. In August 1981, Lakeland, through its subcontractor, Slack Well Drilling Company installed five (5) on-site monitoring wells. One additional monitoring well was installed off-site. Ground water samples were collected and samples from wells 2, 5, and 6 were analyzed by the NCDH as well as by Lakeland. Contaminants including methylene chloride, PCE, benzene, toluene and xylene were detected at levels exceeding State Drinking Water Standards.

A comparison of the two sets of data from NCDH and Lakeland showed widely divergent results. In February, 1982 Commander was notified by NCDH that the site investigation would continue. In May 1982, Pasley operations ceased when the company filed for bankruptcy.

NYSDEC and NCDH were unsuccessful in their efforts to persuade Commander and Pasley to do additional work at the Site. In 1983, NYSDEC issued a Notice of Hearing and Complaint alleging violations of the New York State Environmental Conservation Law, Articles 17, 27 and 71.

On June 10, 1986, the Site was placed on the National Priorities List (NPL). NYSDEC was the lead agency until January 1987. Then, with NYSDEC's concurrence, EPA assumed responsibility for the cleanup of the Site.

### B. <u>History of Surrounding Sites</u>

Two major ground water contamination sites are adjacent to the Site. One is Roosevelt Field, a former airfield that is now a large shopping mall. The Roosevelt Field site was extensively studied by the United States Geological Survey (USGS) from 1982 to 1984. As a result of this study, the USGS identified three volatile organic ground water contamination plumes. Two of the contamination plumes exist in the Upper Glacial aquifer, and the third is present in both the Upper Glacial aquifer and the Magothy Formation. The plumes were reported in 1986 to extend at least 1,000 feet to the south-southwest of Roosevelt Field, and within 400 feet of the Pasley Site. The report states that the ground water in the Upper Glacial aquifer flows at approximately 1 ft./day. At that rate, it is likely that the plume is responsible for the contamination detected in the upgradient Pasley well cluster. The Roosevelt Field Site was listed as a Class II site on the New York State Registry in July 1991.

The Purex/Mitchell Field Transit Facility site (Purex) is the second major ground water contamination site in the area and is approximately 800 feet east of the Site. An investigation conducted by Camp, Dresser and McKee in 1984 showed that contaminants in the upper

Magothy aquifer associated with the Purex Site include: PCE; TCE; 1,1-dichloroethene; and methylene chloride. The ground water contamination from this site is currently being remediated by the Purex company pursuant to a New York State Consent Order.

### C. Enforcement

EPA identified two potentially responsible parties (PRP's) as owners and/or operators. Special notice letters informing the PRPs of their potential liabilities were mailed on February 12, 1988 to Commander and Pasley for conducting a Remedial Investigation and Feasibility Study (RI/FS) for the Site. Several negotiations were held to discuss technical and legal issues relating to the Administrative Order on Consent (AO) for the conduct of the RI/FS.

On August 19, 1988, EPA entered into an AO, Index NO. II- CERCLA-80212, with Commander. The AO required Commander to perform an RI/FS to determine the nature and extent of contamination at the Site and to remove the 12 above-ground tanks that were located on-site. Pasley declined to participate in the settlement.

The tank farm removal was completed in November of 1988 by ABC Demolition and was supervised by EA Engineering, a former consultant of Commander. Metcalf & Eddy, Inc. performed the RI/FS for Commander. The RI Report was approved by EPA in November, 1991. The revised FS Report was submitted to EPA February, 1992.

In February, 1992 EPA sent information request letters regarding generation of wastes found at the Site to 20 parties.

### III. HIGHLIGHTS OF COMMUNITY PARTICIPATION

The RI/FS Reports and the Proposed Plan for the Site were released to the public for comment on February 14, 1992. These two documents were made available at two information repositories maintained at the EPA Region II Office in New York City and the Nassau Library System. The notice of availability for these documents was published in Newsday on February 14, 1992. A public comment period on the documents was held from February 14, 1992 through March 15, 1992. In addition, a public meeting was held on March 5, 1992. At this meeting, representatives from EPA answered questions about problems at the Site and the remedial alternatives under consideration. Responses to the comments and questions are included in the Responsiveness Summary, which is attached as Appendix 4.

### IV. SCOPE AND ROLE OF RESPONSE ACTION

The objective of this remedy is to address the source of contamination at the Site, the contamination in the surface soils, and ground water contamination attributable to the Site. The selected remedy will treat ground water until the influent contaminant concentrations at the extraction wells equal the upgradient concentrations. For the soil remediation alternative, the

contaminated soil will be treated until the recommended soil cleanup objectives as outlined in Table 13 are met or until no more VOCs can be effectively removed from the unsaturated zone.

Contamination upgradient of the Site is suspected to be contributing to the ground water contamination at the Site. The Roosevelt Field site, which is one of the major suspected sources of contamination detected in the Pasley upgradient Glacial aquifer ground water well, was listed as a Class II site on the New York State Registry in July 1991. The EPA and NYSDEC will ensure that any sources contributing to contamination at the Site are addressed. In addition, during the remedial design process, EPA and NYSDEC will also ensure that the effectiveness of the Pasley remediation is not influenced by the ground water recovery system at the adjacent Purex Site.

### V. SUMMARY OF SITE CHARACTERISTICS

### A. <u>Site Geology and Hydrology</u>

Based on soil borings performed during the field investigation, borings for the 30 foot monitoring wells and borings for the 60 foot monitoring wells, revealed only unconsolidated sands and gravels with some silty material at depth. The unconsolidated sediments encountered to a depth of 60 feet belong to the upper Pleistocene undifferentiated glacial outwash deposits or Upper Glacial aquifer. All of the 90 foot wells were screened in the upper portion of the Magothy aquifer (Upper Cretaceous). The Magothy formation consists of fine sand often containing thin, discontinuous layers of silt and clay. The thickness of the Magothy aquifer is estimated at 400 to 500 feet in the Pasley study area. The Upper Glacial aquifer overlies the Magothy aquifer and the two may act as distinct aquifers, or as one, depending upon the degree of hydraulic connection between the two. It is also reported that there is a downward ground water flow direction from the Glacial aquifer to the Magothy aquifer. This downward flow was not always evident throughout the Site. However, in the Site area, it is believed that the two are hydraulically connected. Ground water flows in the Upper Glacial aquifer in a southwesterly direction. The ground water in the Upper Magothy aquifer has a more southerly flow direction than in the Glacial aquifer.

### B. Nature and Extent of Contamination

### Ground Water

Eighteen monitoring wells were installed to evaluate ground water conditions. The monitoring wells were clustered in six locations (three wells each, screened at depths of 30, 60, and 90 feet). The ground water quality of the aquifer underlying the Site, downgradient and upgradient of the Site was assessed by two rounds of water quality sampling in 1990 and a third round of partial sampling in 1991. The on-site shallow ground water monitoring well (MW-2S) indicated highest contamination as compared to the other seventeen monitoring wells. Tables 1 through 3 present the results of the three rounds of ground water sampling. As

Tables 1 through 3 present the results of the three rounds of ground water sampling. As shown in these Tables, the most prevalent Volatile Organic Compound (VOC) was trans - 1,2-dichloroethene at a maximum concentration of 37,000 parts per billion (ppb).

A contaminant plume could not be defined by plotting the Total Volatile Organic Compounds (TVOC) associated with the Site study area. This was due in part to the fact that contamination was detected entering the Site at the upgradient well cluster, MW-1 (Figure 3). Therefore, a group of VOCs which were found at the Site but which were not detected in upgradient well cluster well MW-1 were chosen to define the plume associated with the Site.

The total volatile organic index compounds (TVOIC) chosen to define the plume for the Site are the following: chloroform, 1,1 dichloroethene, 1,1 dichloroethane, trans - 1,2-dichloroethene, 1,1,1 trichloroethane, ethylbenzene, toluene, chlorobenzene, and xylene. The TVOIC compounds were found to contribute a major part (88%) of the contamination found in the monitoring well cluster located on-site (MW-2). However, the use of TVOIC does not imply that non-index compounds (TCE, PCE) are absent from the Site.

Through the use of the index compounds, a well defined contaminant plume could be identified for the Site. Figures 4 through 6 display the plume detected based on the data collected.

Figure 4 is a map of the TVOIC plume for the 20 to 30 foot depth in the Upper Glacial aquifer. It appears that the contaminant plume extends approximately 400 feet to the southwest, parallel to the ground water flow direction and the contaminant plume is approximately 390 feet wide. The maximum level of TVOC contamination detected was 37,000 ppb for trans - 1,2, dichloroethene, 370 times the Federal MCL. TCE, although not part of the TVOIC plume, was also detected at a maximum concentration of 320 ppb, 64 times the federal MCL.

Figure 5 is a map of the TVOIC plume for the 50 to 60 foot depth in the Lower Glacial aquifer. The areal extent of the plume at this depth was found to be much smaller, and centered on MW-4I, directly downgradient of the Site. The maximum level of TVOIC contamination in this portion of the plume was 15 ppb for trans-1,2, dichloroethene. TCE was also detected at 15 ppb.

Figure 6 is a map of the TVOIC contamination plume for the 80 to 90 foot depth in the Upper Magothy aquifer, directly downgradient of the Site. No TVOIC contamination was found directly downgradient or on-site. However, 13 ppb of a TVOIC (trans-1,2, dichloroethene) was found at the eastern edge of the study area at monitoring wells MW-3D and MW-5D. This contamination did not appear to result from the Site and did not follow the south southwesterly direction of ground water flow from the Site.

Samples collected from upgradient off-site monitoring wells showed a maximum level of 27 ppb of PCE (monitoring well location MW-1S) and 15 ppb for TCE (monitoring well location MW-1D). Benzene was also detected at a maximum level of 38 ppb (monitoring well location MW-1I).

Semi-volatile compounds were detected at low levels in the ground water. The only metal detected above the MCL was chromium at 255 ppb.

### 2. Soils

Fifty (50) surface soil grab samples were collected and analyzed for volatile organic compounds. These samples were collected from an approximate 30 foot grid pattern at a depth of 6 to 12 inches below grade. Samples were then collected and composited for metals and semi-volatile organic analyses. Each composite sample consisted of soil from five adjacent discrete sample locations. Figure 7 illustrates surface soil sampling locations. There were eight VOCs that appeared at high concentrations in the surface soil that were also detected in the ground water. These were trans-1,2-dichloroethene, 1,1,1-trichloroethane, TCE, PCE, toluene, xylenes, ethylbenzene and chloroform.

Data from the surface soil samples revealed elevated levels of VOCs originating from three primary locations. The concentrations of TVOCs, primarily PCE and trans-1,2-dichloroethene, were detected in concentrations of 1,000 ppb up to concentrations of 603,000 ppb. Additionally, total semi-volatile organic compounds were detected in composite samples collected from ten locations. The highest concentrations of total semi-volatiles were detected in composite samples 8 and 9 (204,000 ppb and 126,500 ppb, respectively) collected on the eastern edge of the Site. The results of the analyses for the soil samples collected are presented in Table 4.

Subsurface samples were also collected from eight locations on-site and five locations off-site. On-site, two samples were collected from each of eight borings at depths of 12 to 14 feet and 23 to 25 feet (or the first two feet below the water table). A total of sixteen samples were collected. These boring locations are identified on Figure 8. Boring BH-8 was subsequently converted into a 90 foot deep monitoring well (MW-2D).

Table 5 contains the results of the on-site subsurface soil samples. Elevated levels of total VOCs (greater than 1,000 ppb) were detected in six of the sixteen samples. Table 6 identifies the boring number, depth, primary contaminant detected and total VOC concentrations.

Analytical results for semi-volatile compounds indicated that two of the eight samples collected at the 12 to 14 foot depth exhibited elevated total semi-volatile concentrations (12,500 ppb at BH-2A, and 18,000 ppb at BH-3A). There was only one location (BH-7B) that exhibited a total semi-volatile concentration greater than 10,000 ppb (12,710 ppb) at the 23 to 25 feet depth. This data suggest limited downward migration of semi-volatile compounds. The ground water data supports this. MW-2S (the 30 foot shallow well) exhibited 380,000 ppb of total semi-volatile compounds but MW-2I (the 60 foot intermediate well) and MW-2D (the 90 foot well) did not exhibit any semi-volatile contamination.

The levels of metals in the subsurface on-site samples were within the common range for soil and were not significantly different from the off-site results.

### VI. SUMMARY OF SITE RISKS

EPA conducted a Risk Assessment of the "no-action" alternative to evaluate the potential risk to human health and the environment associated with the current conditions. The Risk Assessment began by selecting chemicals of concern that would be representative of site risks. These chemicals were identified based on factors such as potential for exposure to receptors, toxicity, concentration and frequency of occurrence. Table 7 summarizes the chemical of potential concern selected for each sampled media at the Site. The frequency of detection and concentration range for the contaminants of concern are referenced in Table 8.

EPA's Risk Assessment identified several potential exposure pathways by which the public may be exposed to contaminants released from the Pasley site under current and future land-use scenarios. The actual and potential pathways and population potentially affected are shown in Table 9.

Since access is restricted to the public and the Site is covered by gravel, it is not considered likely that direct contact with the contaminated soil would occur. Therefore, the only complete exposure pathway under current land use conditions is inhalation exposure to chemicals that volatilize from the soil. The reasonable maximum exposure was evaluated. The following pathways were selected for evaluation under the future land use conditions:

- direct contact and incidental ingestion exposure with chemicals present in surface soils,
- ingestion exposures to chemicals present in ground water,
- ingestion and inhalation exposures during home use to chemicals present in ground water, and
- inhalation exposures to chemicals that have volatilized from surface soils.

The potentially exposed populations in all cases were the residents (adult and children) of the neighborhood surrounding the Site and future workers on-site.

Under current EPA guidelines, the likelihood of carcinogenic (cancer causing) and non-carcinogenic effects due to exposure to site chemicals are considered separately. It was assumed that the toxic effects of the site-related chemicals would be additive. Thus, carcinogenic and non-carcinogenic risks associated with exposures to individual compounds of concern were added to indicate the potential risks associated with mixtures of potential carcinogens and non-carcinogens, respectively.

Non-carcinogenic risks were assessed using a hazard index (HI) approach, based on a comparison of expected contaminant intakes and safe levels of intake, or Reference Doses (RfDs). RfDs have been developed by EPA for indicating the potential for adverse health effects. RfDs, which are expressed in units of mg/kg-day, are estimates of daily exposure levels for humans which are thought to be safe over a lifetime (including sensitive individuals). Estimated intakes of chemicals from environmental media (e.g., the amount of a chemical ingested from contaminated drinking water) are compared with the RfD to derive the hazard quotient for the contaminant in the particular medium. The HI is obtained by adding the hazard quotients for all compounds across all media that impact a common receptor.

An HI greater than 1 indicates that the potential exists for non-carcinogenic health effects to occur as a result of site-related exposures. The HI provides a useful reference point for gauging the potential significance of multiple contaminant exposures within a single medium or across media. The RfDs for the chemicals of potential concern at the Pasley site are presented in Table 10.

A summary of the non-carcinogenic risks associated with the chemicals of potential concern across various exposure pathways is found in Table 11. It can be seen from Table 11 that the greatest non-carcinogenic risk from the Site is associated with ingestion of on-site Upper Glacial aquifer water by on-site workers. The noncarcinogenic effects, exceed 1.0 due primarily to chromium and TCE. The hazard index for soil was calculated to be less than 1.0.

Potential carcinogenic risks were evaluated using the cancer slope factors (Sfs) developed by EPA for the chemicals of potential concern. Sfs have been developed by EPA's Carcinogenic Risk Assessment Verification Endeavor (CRAVE) for estimating excess lifetime cancer risks associated with exposure to potentially carcinogenic chemicals. Sfs, which are expressed in units of (mg/kg-day)<sup>-1</sup>, are multiplied by the estimated intake of a potential carcinogen, in mg/kg-day, to generate an upper-bound estimate of the excess lifetime cancer risk associated with exposure to the compound at that intake level. The term "upper bound" reflects the conservative estimate of the risks calculated from the SF. Use of this approach makes the underestimation of the risk highly unlikely. The SF for each indicator chemical is presented in Table 8.

For known or suspected carcinogens, EPA considers excess upper bound individual lifetime cancer risks of between 10<sup>-4</sup> to 10<sup>-6</sup> to be acceptable. This level indicates that an individual has not greater than a one in ten thousand to one in a million chance of developing cancer as a result of site-related exposure to a carcinogen over a 70-year period under specific exposure conditions at the Site. The total cancer risks at the Pasley Site are outlined in Table 9. The total cancer risk for on-site occupants is 4 x10<sup>-4</sup>, based on ingesting untreated ground water from the Upper Glacial aquifer in the vicinity of the Site. The total cancer risk for children is 9 x10<sup>-4</sup> in the vicinity of the Site, based on ingesting untreated ground water from the Upper Glacial aquifer.

The cumulative upperbound risks at the Site for on-site occupants under a future potential land use scenario associated with ground water is 9 x10<sup>-1</sup> which exceed EPA's risk criteria. In addition, MCLs are currently exceeded for several hazardous substance in ground water. Although the risk posed by the soils are within EPA's acceptable risk criteria, contaminants in the soils, if not addressed, will likely continue to contribute to further contamination of the ground water at the Site.

### **UNCERTAINTIES**

The procedures and inputs used to assess risks in this evaluation, as in all such assessments, are subject to a wide variety of uncertainties. In general, the main sources of uncertainty include:

- environmental chemistry sampling and analysis
- environmental parameter measurement
- fate and transport modeling
- exposure parameter estimation
- toxicological data

Uncertainty in environmental sampling arises in part from the potentially uneven distribution of chemicals in the media sampled. Consequently, there is significant uncertainty as to the actual levels present. Environmental chemistry analysis error can stem from several sources including the errors inherent in the analytical methods and characteristics of the matrix being sampled.

Uncertainties in the exposure assessment are related to estimates of how often an individual would actually come in contact with the chemicals of potential concern, the period of time over which such exposure would occur, and in the models used, to estimate the concentrations of the chemicals of potential concern at the point of exposure.

Uncertainties in toxicological data occur in extrapolating both from animals to humans and from high to low doses of exposure, as well as from the difficulties in assessing the toxicity of a mixture of chemicals. These uncertainties are addressed by making conservative assumptions concerning risk and exposure parameters throughout the assessment. As a result, the Risk Assessment provides upper bound estimates of the risk to populations near the site.

A specific uncertainty inherent in the Site risk assessment is that the methodology used to calculate the site risks are site-wide averages, which give a clear overall understanding of site risks. However, as previously stated, EPA has taken into account the sensitivity of the on-site and neighboring populations and has determined that the target risk for the site should be on the order of 10<sup>-6</sup>.

Therefore, actual or threatened releases of hazardous substances from this site, if not addressed by the selected alternative or one of the other remedial measures considered, may

present an imminent and substantial endangerment to the public health, welfare, and the environment. More specific information concerning public health risks, including a quantitative evaluation of the degree of risk associated with various exposure pathways, is presented in the Risk Assessment which can be found in the Administrative Record.

### VII. DESCRIPTION OF ALTERNATIVES

Two media-specific remedial actions are required to protect human health and the environment because of the nature of the contamination at the Site. They are numbered to correspond with their presentation in the FS report. On-site soil has been determined to be a source of contamination. Contaminants were found to move from the unsaturated soil to the ground water. Once in the ground water, the contaminants, under the influence of the ground water gradient, migrate from the Site to potential receptors.

Specific remedial action objectives for this Site include:

Ground water - Restoration of ground water quality to its intended use (Class IIb and GA-potential of drinking water) by reducing contaminant levels below State and Federal drinking water standards where possible (see Table 12). In the case where upgradient concentrations prohibit such restoration for a particular compound, the contaminant level will be reduced to the upgradient level.

Soil - In order for the soil not to be a contributor to ground water contamination, the degree to which the contaminants have to be reduced is different for each component (see Table 13). For VOCs (components of interest, trans-1,2-dichloroethene, 1,1,1-trichloroethane, TCE, PCE, toluene and, xylenes), the contaminated soil will be treated until the recommended soil cleanup objectives are met or until no more VOCs can be effectively removed from the unsaturated zone. For the semi- volatile compounds of interest, the contaminants di-n-butyl phthalate, naphthalene, bis-(2-ethylhexyl) phthalate and floranthene have to be reduced below 50 ppm.

The time to implement refers only to the actual construction and remedial action (time to achieve clean up) time and excludes the time needed to design the remedy, procure contracts, and negotiate with the PRPs, all of which can take 15-30 months.

The alternatives identified for both soil and ground water are presented below:

### **Soil Remediation Alternatives:**

### Alternative 1: No Action

CERCLA requires EPA to consider the "No Action" alternative at every Superfund site to provide a baseline of comparison among alternatives. Under this alternative, the contaminated soil would be left in place without treatment. A long-term monitoring program would be implemented to track the migration of contaminants from the soil into the ground water. In

accordance with Section 121 of CERCLA, remedial actions that leave hazardous substances above health-based levels at a site are to be reviewed at least once every five years to assure that the action is protective of human health and the environment. Accordingly, the no action alternative would have to be reviewed by EPA at least once every five years.

Capital cost:

\$0

Annual Operation

& Maintenance:

\$7,000

30-year Present

Worth:

\$66,000

Time to Implement:

Construction: 2 Months Remedial Action: 30 years

### Alternative 2- Excavation with Off-site Disposal

This alternative involves the excavation and off-site disposal of the contaminated soil from the eastern and western portions of the Site.

The soil excavation would extend to a depth of 2 feet on the eastern section of the Site, and to a depth of 20 feet on the western portion of the Site, where the soils are highly contaminated. Approximately 10,083 cubic yards of soil contaminated with volatile organic and semi-volatile organic compounds would be excavated and the excavated soil would then be disposed of off-site at a RCRA-permitted landfill.

However, the soil will be tested using the Toxicity Characteristic Leaching Procedure (TCLP), to determine if treatment is necessary prior to disposal to insure that RCRA land disposal restrictions are met. The Land Disposal Restrictions set treatment standards which are based on the best demonstrated available technology (BDAT) for treatment of a given waste. In the case of VOCs in soil, the BDAT treatment method is generally incineration. If incineration is necessary to meet the Land Disposal Restriction's, a dry ash material would be produced which may require further RCRA-permitted disposal to protect the environment. This alternative would then be essentially equivalent to Alternative 3. The actual quantity of soil requiring treatment would be refined during the remedial design.

Capital cost:

\$8,675,000

Annual Operation

& Maintenance:

Present Worth:

\$8,675,000

Time to Implement: 1-2 Months

### Alternative 3- Excavation with Off-site Incineration

This alternative involves the same excavation of contaminated soil as described in Alternative 2. However, the excavated soil would be transported to an off-site facility for incineration. This alternative produces a dry ash material high in metals that would require further RCRA-permitted disposal to protect the environment.

Capital cost:

\$43,970,000

Annual Operation

& Maintenance:

Present Worth:

\$43,970,000

Time to Implement: 1-2 Months

### Alternative 4- Excavation with Solidification/Stabilization

This alternative involves the same excavation of contaminated soil described in Alternatives 2 However, instead of transporting the soil off-site for treatment/disposal, the and 3. solidification/ stabilization process would involve construction of a treatment facility on-site.

The process would involve mixing of the excavated contaminated soils with a solidifying matrix to bind chemically the contaminants to form a "soil concrete." A solidifying matrix might include the use of lime, fly ash or cement to bind the contaminants in a solid block of treated soil. After the soils have been mixed with the solidification matrix, the resulting concrete-like substance would be placed back on the Site for hardening and final compaction.

Before the treatment technology is applied to the area, a treatability study would be performed on the soil to determine the effectiveness of different binders and to obtain additional information required for the development of preliminary design considerations.

Capital cost:

\$2,108,000

Annual Operation

& Maintenance:

Present Worth:

\$2,108,000

Time to Implement: 6 - 8 Months

### Alternative 5- Soil Flushing

This alternative would work in conjunction with the selected ground water remedial alternative. This alternative entails installation of an infiltration system to effect soil flushing for removing the VOCs and semi-volatile organics from the soil. This process would involve injection of water or an aqueous solution into the area of soil contamination utilizing infiltration trenches. The injected water would flush the soil contamination into the ground water. The contaminated ground water would be pumped to the surface, treated and recharged to continue the process.

The infiltration trench system would consist of 3 excavated trenches approximately 2 feet in depth backfilled with a coarse stone aggregate. The treated water from the ground water treatment system would be distributed through the gravel trenches by a 4 inch PVC perforated pipe. The 3 trenches would transverse the length of the site and have 20 foot spacing between each trench. The aggregate fill material for the infiltration trenches would be completely surrounded with filter fabric to prevent soil movement into the aggregate. An observation well would be installed in each infiltration trench.

The organic contaminants in the soil at the Site have high solubilities in water and are therefore expected to be flushed from the soil using treated ground water as the washing agent.

Capital cost:

\$137,000

Annual Operation

& Maintenance:

\$15,000

Present Worth: \$185,000

Time to Implement:

Construction: 6 Months Remedial Action: 4 Years

### Alternative 6- Soil Vacuuming

Soil vacuuming would involve the installation of vents in the contaminated unsaturated soil zone. A vacuum would be applied through these vents to volatilize and extract organic compounds from the soil. The organic vapors would be drawn into a collection system where they would be removed through an activated carbon off-gas treatment system.

Circulation of air through the soil also would enhance the biodegradation of semi-volatiles in the unsaturated zone.

A small amount of liquid condensate would be generated during the vapor extraction process. With an on-site ground water treatment alternative operating in conjunction with ground water remediation, the condensate may be treated on-site at minimal cost. Off-site disposal of condensate would be necessary if this alternative was implemented before a ground water treatment system was constructed.

Under this alternative approximately thirteen thousand (13,000) cubic yards of contaminated soil would be treated until no more VOCs can be effectively removed from the unsaturated vadose zone.

Subsurface soil sampling would be required to monitor the progress of the soil vapor extraction process.

Capital cost:

\$882,000

Annual Operation

& Maintenance:

\$664,000

Present Worth:

\$1,562,000

Time to Implement:

Construction:

6 Months

Remedial Action: 2 Years

### Alternative 7- Soil Vacuuming and Soil Flushing

This alternative combines Alternatives 5 and 6. The soil flushing technology would remove most volatile and semi-volatile compounds but may not be as effective in removing a group of volatile compounds known as monocyclic aromatic hydrocarbons. Soil vacuuming, however, would perform well in removing monocylic and aliphatic hydrocarbons but may not be as effective for semi-volatile compounds. However, it should be noted that the circulation of air through the soil as part of the vacuuming procedure would enhance the biodegradation of the semi-volatiles in the soil.

Under this alternative, soil vacuuming would be performed initially to remove the volatile and semi-volatile compounds. A soil sampling and analysis program would then be implemented to evaluate the success of the soil vacuuming. Soil flushing, used to flush any remaining water-soluble contaminants from the soil, would be performed after soil vacuuming to achieve soil cleanup goals. However, if it is found after the soil vacuuming that concentrations of semivolatile compounds are decreasing in the soil and are not impacting ground water, the soil flushing technique may be abandoned. Periodic subsurface soil sampling and analysis would be required to monitor the progress of both processes.

Capital cost:

\$921,000

Annual Operation

& Maintenance:

\$407,000

Present Worth:

\$1,649,000

Time to Implement:

Construction:

1 Year

Remedial Action:

6 Years

### **Ground Water Treatment Alternatives:**

All of the remedial ground water alternatives, except the No Action alternative, involve extraction, treatment and recharge of the treated water to the ground water. The contaminated ground water is recovered using extraction wells at the downgradient end of the contaminant plume. The extracted ground water is treated and returned to the aquifer via a series of recharge wells located upgradient of the contaminant plume and/or infiltration trenches located in the area of soil contamination.

Recent studies have indicated that pumping and treatment technologies may contain uncertainties in achieving the ppb concentrations required under ARARs over a reasonable period of time. However, these studies also indicate significant decreases in contaminant concentrations early in the system implementation, followed by a leveling out. For these reasons, the selected ground water treatment alternative stipulates contingency measures, whereby the groundwater extraction and treatment system's performance will be monitored on a regular basis and adjusted as warranted by the performance data collected during operation. Modifications may include any or all of the following:

- a) at individual wells where cleanup goals have been attained, pumping may be discontinued;
- b) alternating pumping at wells to eliminate stagnation points;
- c) pulse pumping to allow aquifer equilibration and to allow adsorbed contaminants to partition into groundwater; and
- d) installation of additional extraction wells to facilitate or accelerate cleanup of the contaminant plume.

If it is determined, on the basis of the preceding criteria and the system performance data, that certain portions of the aquifer cannot be restored to their beneficial use in a reasonable time frame, all or some of the following measures involving long-term management may occur, for an indefinite period of time, as a modification of the existing system:

- a) engineering controls such as physical barriers, source control measures, or longterm gradient control provided by low level pumping, as containment measures;
- b) chemical-specific ARARs may be waived for the cleanup of those portions of the aquifer based on the technical impracticability of achieving further contaminant reduction;

- c) institutional controls, in the form of local zoning ordinances, may be recommended to be implemented and maintained to restrict access to those portions of the aquifer which remain above remediation goals;
- d) continued monitoring of specified wells; and
- e) periodic reevaluation of remedial technologies for groundwater restoration.

The decision to invoke any or all of these measures may be made during a periodic review of the remedial action, which will occur at intervals of no less often than every five years.

### Alternative 1- No Action

CERCLA, as amended, requires that the "no-action" alternative be considered at every site. Under this alternative, no remediation measures would be implemented at this time. This alternative allows for natural attenuation of the contaminants and includes institutional controls and monitoring. This alternative also would include restrictions on future ground water use and a pubic awareness program.

Periodic ground water sampling and analysis would be required to monitor the progress of natural attenuation. In effect, this no action alternative is essentially equivalent to the no action alternative under the soil remediation alternative section of this ROD.

Capital cost:

\$0

**Annual Operation** 

& Maintenance:

\$7.000

10-year

\$43,000

30-year Present

Worth:

\$66,000

Time to Implement:

Construction:

2 Months

Remedial Action: 30 Years

## Alternative 2- Metals Precipitation/ Powdered Activated Carbon Treatment (PACT)/GAC Polishing

This alternative utilizes three collection wells for the extraction of contaminated ground water followed by on-site treatment. To contain and remove ground water from the contamination plume, it is estimated that it would be necessary to pump 450 gallons per minute (GPM) from three extraction wells placed at depths of 60 feet. Ground water would be pumped from the extraction well system to a holding/ equalization tank. The pumped ground water would then enter the treatment plant where it would go through an initial two-stage precipitation

and clarification/filtration unit for the removal of all heavy metals. The heavy metals treatment would be followed by powdered activated carbon treatment (PACT) to remove volatile organic and semi-volatile organic compounds.

The granular activated carbon (GAC) adsorption system that follows the PACT would be used, if necessary, as a final polishing step to remove any remaining organic compounds in order to achieve ARARs. Carbon adsorption would remove organic compounds from waste water onto the activated carbon. The exact amount of treated water that would be recharged to the ground water either by the recharge wells or by the infiltration trenches would be determined in the remedial design.

The by-products resulting from the treatment system include metals sludge, filtered solids, and spent granular activated carbon. The sludge would be transported off-site for treatment and disposal at a RCRA-permitted facility.

Periodic sampling and analysis of the influent and effluent would be required to monitor the progress of this treatment alternative.

Capital cost:

\$6,465,000

Annual Operation

& Maintenance:

\$1,623,000

10-year Present Worth: \$16,438,00 30-year Present Worth: \$ 21,765,000

Time to implement:

Construction:

6 Months

Remedial Action:

10-40 Years

Alternative 3- Metals Precipitation/Air Stripping with Fume Incineration/Granular Activated Carbon(GAC) Polishing

Under this alternative, the same extraction system is used to withdraw the contaminated ground water as that of Alternative 2. This alternative differs in that after metals removal, the effluent from the metals system would be pumped into an air stripper that would be effective in removing the VOCs from the water. Air stripping is a mass transfer process in which volatile contaminants in water are transferred to the gaseous phase.

Fume incineration would be used to treat any gaseous discharge from the air stripper. Fume incineration units are chambers heated by supplemental fuel which provide high enough temperatures and retention time to combust the contaminants in the off-gas. Temperatures in the combustion chamber range from 1200°F to 1800°F.

The liquid phase from the air stripper would be pumped into the granular activated carbon (GAC) adsorption system that would be used as a final polishing step to remove any remaining organic compounds. Treatment residuals include spent carbon from the fume incinerator and spent carbon from the liquid phase carbon polishing.

Periodic sampling and analysis of the influent and effluent would be required to monitor the progress of the treatment alternative. During the periodic sampling and analyses of the influent, if it is determined that metals concentrations are below standards and low enough not to cause malfunction of the air stripper, the metals precipitation portion of the treatment train may be eliminated.

Capital cost:

\$3,199,000

Annual Operation

& Maintenance: \$1,069,000 10-year Present Worth: \$9,768,00 30-year Present Worth: \$13,276,000

Time to Implement:

Construction: 2 Years Remedial Action: 10-40 Years

Alternative 4-Metals Precipitation/Air Stripping with Vapor Phase Granular Activated Carbon/GAC Polishing

This treatment alternative is the same as Alternative 3 except that the off-gas emissions from the air-stripper would be treated by passing the air stream through vapor phase carbon adsorption columns, instead of the fume incinerator. In this alternative, contaminated air flows through the columns or carbon bed, and organics adsorb onto the carbon. The treated air then leaves the carbon bed with reduced concentrations of contaminants until the carbon adsorbent cannot take on additional organics. Removal efficiencies utilizing vapor phase activated carbon have been reported at greater than 98 percent.

Additional sludges would be generated from the carbon adsorption columns.

Capital cost:

\$4,280,000

Annual Operation

& Maintenance: \$829,000 10-year Present Worth: \$9,374,000 30-year Present Worth: \$ 12,095,00

Time to implement:

Construction: 2 Years Remedial Action: 10-40 Years

### Alternative 5- Metals Precipitation/UV Peroxidation

Under this alternative, the same extraction system is used to withdraw the contaminated ground water as that of Alternative 2. UV Peroxidation is an innovative technology for cleanup and destruction of organic compounds in ground water. In this process, ultraviolet light reacts with hydrogen peroxide to form hydroxyl radicals. These powerful chemical oxidants then react with the organic contaminants in water. The end products of the oxidation process are carbon dioxide (CO<sub>2</sub>), water, and hydrochloric acid. Chemical oxidation would reduce the toxicity and volume of contaminated ground water at the Site.

Periodic sampling and analysis of the influent and effluent would be required to monitor the progress of this treatment alternative.

Capital cost:

\$4,421,000

Annual Operation

& Maintenance:

\$1,459,000

10-year Present Worth: \$13,386,000 30-year Present Worth: \$18,175,000

Time to Implement:

Construction:

1 Year

Remedial Action: 10-40 Years

### VIII. SUMMARY OF COMPARATIVE ANALYSIS OF ALTERNATIVES

In accordance with the NCP, a detailed analysis of each alternative is required. The purpose of the detailed analysis is to assess objectively the alternatives with respect to nine evaluation criteria that encompass statutory requirements and include other gauges of the overall feasibility and acceptability of remedial alternatives. This analysis is comprised of an individual assessment of the alternatives against each criterion and a comparative analysis designed to determine the relative performance of the alternatives and identify major trade-offs, that is, relative advantages and disadvantages, among them.

The nine evaluation criteria against which the alternatives are evaluated are as follows:

Threshold Criteria - The first two criteria must be satisfied in order for an alternative to be eligible for selection.

### 1. Overall Protection of Human Health and the Environment:

This criterion addresses whether or not a remedy provides adequate protection and describes how risks are eliminated, reduced, or controlled through treatment, engineering controls, or institutional controls.

### 2. Compliance with ARARs:

This criterion addresses whether or not a remedy will meet all the ARARs of other federal or State environmental statutes and/or provide grounds for invoking a waiver.

<u>Primary Balancing Criteria</u> - The next five "primary balancing criteria" are to be used to weigh major trade-offs among the different hazardous waste management strategies.

### 3. Long-term Effectiveness and Permanence:

This criterion refers to the ability of the remedy to maintain reliable protection of human health and the environment over time once cleanup goals have been met.

### 4. Reduction of Toxicity, Mobility, or Volume:

This criterion addresses the degree to which a remedy utilizes treatment technologies to reduce the toxicity, mobility, or volume of contaminants.

### 5. Short-term Effectiveness:

This criterion considers the period of time needed to achieve protection and any adverse impacts on human health and the environment that may be posed during the construction and implementation period until cleanup goals are met.

### 6. Implementability:

This criterion examines the technical and administrative feasibility of a remedy, including availability of materials and services needed to implement the chosen solution.

### **7.** Cost:

This criterion includes capital and O&M costs.

<u>Modifying Criteria</u> - The final two criteria are regarded as "modifying criteria," and are to be taken into account after the previous criteria have been evaluated. They are generally to be focused upon after public comment is received.

### 8. State Acceptance:

This criterion indicates whether, based on its review of the FS and Proposed Plan, the State concurs with, opposes, or has no comment on the proposed alternative.

### 9. Community Acceptance:

This criterion indicates whether, based on its review of the FS and Proposed Plan, the public concurs with, opposes, or has no comment on the proposed alternative. Comments received during this public comment period, and the EPA's responses to those comments, are summarized in the Responsiveness Summary which is appended to this ROD.

The following is a summary of the comparison of each alternative's strengths and weaknesses with respect to the nine evaluation criteria.

### 1. Overall Protection of Human Health and the Environment

### Soil Remediation Alternatives

All the soil remediation alternatives are considered protective of human health and the environment except Alternative 1. Alternative 1 is not protective of human health and the environment because it does not eliminate, reduce or control the contaminants at the Site. Since it does not meet this threshold criterion, Alternative 1 will not be discussed further.

Alternatives 2 and 3 would not require any long term maintenance or deed restrictions. However, Alternatives 2 and 3 involve transportation of contaminated soil off-site, and increase the potential risks associated with dust generated during excavation and/or transportation. Alternative 4 would require long-term monitoring to ensure the stability of the solidification/stabilization process. Alternatives 5, 6, and 7 reduce potential human health risks by utilizing treatment to remove contaminants from the soil.

### Ground Water Treatment Alternatives

All the ground water alternatives, except the No Action alternative, are considered protective over the long term and would provide overall protection by effectively removing contaminants so that the ground water could be used for potable purposes, if desired. All the treatment alternatives would result in permanent protection of human health and the environment through the reduction in toxicity, mobility, and volume of the contaminants.

However, Alternative 2, by using the PACT system, has a disadvantage over Alternatives 3, 4, and 5, namely, additional sludges would be produced with the activated carbon system thus posing an added minor risk to workers and the environment, especially during the transportation of the sludges for disposal off-site.

Alternatives 3 and 4 pose additional risks associated with air emissions. However, the vapor phase treatment would eliminate any risk associated with air emissions. Alternative 5, by using UV peroxidation has certain advantages over the other alternatives, since it would provide complete destruction of VOCs, thus reducing waste sludges that would otherwise require further treatment and disposal.

### 2. Compliance With ARARs

### Soil Remediation Alternatives

There are no chemical-specific ARARS for soils. It is anticipated that any action specific ARARS

associated with soil treatment can be met by each alternative. However, Alternative 4 would require that treated soil be tested using the Toxicity Characteristic Leaching Procedure (TCLP), before backfilling, to insure that RCRA land disposal restrictions are met. At this point in time, a determination cannot be made whether these levels can be met. If levels cannot be met, a treatability variance may be required.

### Ground Water Treatment Alternatives

Alternatives 2 through 5 achieve ARARs to a similar degree. None of the alternatives would achieve chemical-specific ARARs for ground water as a potential drinking water supply. Achieving chemical-specific ARARs for ground water is dependent on remediation of upgradient sources. This is due to the fact that regardless of the Site cleanup, upgradient sources will continue to be a source of contamination to the ground water beneath the Pasley Site. EPA believes that the proposed remedial action will result in attainment of chemical specific ground water ARARs providing upgradient sources are remediated so that they no longer impact the Upper Glacial aquifer.

EPA may invoke a technical waiver of the chemical-specific ARARs if the remediation program indicates that reaching MCLs in the glacial aquifer is technically impracticable due to the presence of upgradient sources.

Until upgradient sources are remediated so that they no longer impact the Site, EPA will attain ground water cleanup levels which are equal to upgradient concentrations. The remedial action will attain ground water cleanup levels equal to upgradient concentrations for certain contaminants.

Alternatives 2 through 5 would meet action-specific ARARs as outlined in Table 2-1 of the FS Report. Under these alternatives, treated ground water would meet pertinent federal and state ARARs.

### 3. Long-term Effectiveness

### Soil Remediation Alternatives

Alternatives 5, 6 and 7 afford a greater degree of long-term effectiveness and permanence than Alternatives 2 or 4. Alternative 4 would require institutional controls for land use, which would need to be enforced for complete effectiveness.

Alternative 3 is the only alternative that removes all contaminants from the Site and provides total destruction of the contamination sources.

### **Ground Water Treatment Alternatives**

Long-term effectiveness of the ground water alternatives requires the remediation of upgradient contamination. Alternatives 2 through 5 provide long-term effectiveness because these alternatives are designed to reduce contaminant concentrations in the treated ground water to levels that are protective of human health and the environment before discharge. Alternative 1 may present a long-term risk because it relies on natural attenuation to reduce contaminant concentrations.

### 4. Reduction of Toxicity, Mobility, or Volume

### Soil Remediation Alternatives

Alternative 2 does not utilize treatment to reduce the toxicity, mobility or volume of the contaminants. Alternative 3, excavation and off-site incineration, would provide the greatest degree of destruction of contaminants and therefore, the greatest degree of reduction of toxicity, mobility, and volume. However, Alternative 3 would produce ash that would require disposal. In addition, Alternative 4 would not cause a reduction in toxicity but would result in a reduction in mobility. Alternative 4 would increase the soil volume by the introduction of a solidifying matrix.

Alternatives 5 and 6 may not provide as great a degree of contaminant destruction or reduction in contaminant mobility as Alternatives 3 and 4, respectively. However, they are expected to provide an adequate degree of contaminant destruction by gradual reduction of mobility, toxicity and volume. Alternatives 5 and 7 involves soil flushing and must be done in conjunction with ground water extraction and treatment. These technologies used in combination would provide sufficient reduction of mobility, toxicity and volume.

### **Ground Water Treatment Alternatives**

Alternatives 2 through 5 would control the mobility of contaminants contributed by the Site. These alternatives also would significantly reduce or eliminate the toxicity and volume of contaminated ground water by treatment to remove metals, semi-volatile and volatile organic compounds.

However, Alternative 5 by utilizing the UV peroxidation is more advantageous than Alternatives 2 through 4 because it provides a total chemical breakdown of the VOCs into less toxic compounds without any accumulation of sludges and waste residuals.

### 5. Short-term Effectiveness

### Soil Remediation Alternatives

Alternatives 2, 3, and 4, the excavation alternatives, may potentially increase the risk to the community during their implementation because they remove contaminants and create new potential exposure routes not identified in the Risk Assessment. However, necessary measures, such as implementation of proper safety procedures and on-site monitoring would be taken to minimize any significant risk from exposure to the contaminants.

Alternatives 5, 6 and 7 would have the least short-term effect on the community during implementation, since they would be conducted in-situ. All the alternatives have minor short-term effects on the surrounding community, including increased vehicular traffic, a slight increase in noise level from construction equipment, and fugitive dust emissions.

### **Ground Water Treatment Alternatives**

The extraction and treatment alternatives for ground water involve little disturbance to contaminated subsurface areas; therefore the potential risks to site workers and the surrounding community are minor and can be managed. The potential short-term risks to human health and the environment are also anticipated to be low for each of these alternatives.

### 6. Implementability

### Soil Remediation Alternatives

All the alternatives are technically and administratively feasible. Of the soil remediation alternatives, Alternatives 2 and 3 would require the least time to implement. Alternative 4 would take more time to implement since it would require a treatability study and special equipment to treat the soils.

The potential impacts that Alternatives 5 and 7 may have on ground water flow regimes make these alternatives more complex and difficult to implement than Alternative 6. The soil flushing alternatives, Alternatives 5 and 7, require coordination with the ground water treatment alternative.

### **Ground Water Treatment Alternatives**

The treatment components of Alternatives 2 through 4 are proven effective for all contaminants of concern and should be easiest to implement because they rely on well understood and readily available commercial components. Alternative 5 relies on an innovative technology for treatment. Treatability studies would be required to determine the level of effectiveness that can be provided by this technology.

### 7. Cost

Individual cost breakdowns are included in the Description of Alternatives section of this ROD. Capital cost is the value for building the remedial action. Annual operation and maintenance (O&M) costs are used to quantify the yearly expense of O&M. The 30 year present worth cost is then calculated and expressed in current value terms.

### Soil Remediation Alternatives

The present worth cost of Alternative 7 for soils is approximately \$1,649,000. The estimated cost range of the alternatives is from a present worth of \$66,000 (no action alternative) to \$43,970,000 (excavation and off-site incineration).

### **Ground Water Alternatives**

The 30-year present worth cost of Alternative 4 for ground water is approximately \$12,095,000. The estimated cost range of the alternatives is from a 30-year present worth of \$66,000 (no action alternative) to \$21,765,000 (PACT).

### 8. State Acceptance

The State of New York supports the selected remedy presented in this ROD. A copy of their concurrence letter is appended to this ROD.

### 9. Community Acceptance

The local community accepts the selected remedy. All comments that were received from the public during the public comment period are addressed in the attached Responsiveness Summary.

### IX. THE SELECTED REMEDY

Based upon consideration of the results of the RI/FS reports and after careful consideration of all reasonable alternatives, EPA recommends the following alternative for cleaning up the contaminated soils and ground water at the Pasley Solvents and Chemicals Superfund Site:

Soil Remediation Alternative 7: Soil Vacuuming and Soil Flushing in conjunction with Ground Water Treatment Alternative 4: Extraction/Metals Precipitation/Air Stripping with Vapor Phase Granular Activated Carbon/GAC Polishing/Recharge.

The soil remediation alternative, soil vacuuming, has been demonstrated to be effective primarily for removal of VOCs from the unsaturated zone. Circulation of air through the soil during the vacuuming process also would enhance the biodegradation of semi-volatiles in the

unsaturated zone. If sampling after the conclusion of soil vacuuming demonstrates that concentrations of semi-volatile compounds are decreasing in the soil and are still not impacting ground water, the soil flushing portion (for the removal of semi-volatiles in soil) of Alternative 7 may be eliminated.

Specifically, the preferred alternatives will involve the following:

- Treatment of approximately thirteen thousand (13,000) cubic yards of contaminated soil by soil vacuuming and/or by soil flushing, as necessary, until the recommended soil cleanup objectives are met or until no more VOCs can be effectively removed from the unsaturated (vadose) zone;
- 2) Disposal of treatment residuals at a RCRA Subtitle C facility;
- 3) Remediation of the ground water by extraction/metals precipitation/air stripping with vapor phase granular activated carbon/GAC polishing/ and recharge to meet Federal and State drinking water MCLs, except in those cases where upgradient concentrations are above such standards:
- 4) Pumping of contaminated ground water from three extraction wells at a combined flow rate of approximately 450 gpm. The actual pumping rate will be determined during the Remedial Design;
- 5) Long-term monitoring to track the migration and concentrations of the contaminants of concern;
- 6) Implementation of a system monitoring program that includes the collection and monthly analysis of the influent and effluent from the treatment systems and periodic collection of well-head samples.
- 7) Evaluation of Site conditions at least once every five years to determine if a modification to the selected alternative is necessary; and
- 8) The option for EPA to invoke a technical waiver of the ground water ARARs if the remediation program indicates that reaching MCLs in the glacial aquifer is technically impracticable.

The selected ground water alternative also stipulates contingency measures, outlined under Ground Water Treatment Alternatives in the Description of Alternatives section of this ROD, whereby the groundwater extraction and treatment system's performance will be monitored on a regular basis and adjusted as warranted by the performance data collected during operation. If it is determined, in spite of any contingency measures that may be taken, that portions of the aquifer cannot be restored to its beneficial use, ARARs may be waived based on technical

impracticability of achieving further contaminant reduction. The decision to invoke a contingency measure may be made during periodic review of the remedy, which will occur at intervals of no less often than every five years.

#### X. STATUTORY DETERMINATIONS

Under its legal authorities, EPA's primary responsibility at Superfund sites is to undertake remedial actions that achieve adequate protection of human health and the environment. In addition, Section 121 of the CERCLA establishes several other statutory requirements and preferences. These specify that, when complete, the selected remedial action for a site must comply with applicable or relevant and appropriate environmental standards established under federal and state environmental laws unless a statutory waiver is justified. The selected remedy also must be cost effective and utilize permanent solutions and alternative treatment technologies to the maximum extent practicable. Finally, CERCLA includes a preference for remedies that employ treatment that permanently and significantly reduces the volume, toxicity, or mobility of hazardous substances as their principal element. The following sections discuss how the selected remedy meets these statutory requirements.

#### 1. Protection of Human Health and the Environment

The selected remedy for ground water is protective of human health and the environment. The selected ground water remedy eliminates all outstanding threats posed by the Site. The selected ground water remedy reduces contamination to health based levels except in those cases where upgradient concentrations exceed those levels. Contamination upgradient of the Site is suspected to be contributing to the ground water contamination at the Site. The Roosevelt Field Site, which is one of the major suspected sources of the contamination detected in the Pasley upgradient ground water monitoring well, was listed as a Class II site on the New York State Registry in July 1991. The EPA and NYSDEC will ensure that any sources contributing to contamination of the Site are addressed.

The selected remedy for soils is also fully protective of human health and the environment. The soil remedy removes a continuing threat to ground water posed by the on-site contaminated soils.

### 2. Compliance with Applicable or Relevant and Appropriate Requirements

At the completion of response actions, the selected remedy will have complied with the following ARARs and considerations:

#### Action-specific ARARs:

Safe Drinking Water Act (SDWA) Maximum Contaminant Levels (40 CFR 141.11-141.16) and 6 NYCRR Ground Water Quality Regulations (Parts 703.5, 703.6, 703.7) and the NYS Sanitary

code (10 NYCRR part 5) provide standards for toxic compounds for public drinking water supply systems. The recharge process for treated ground water will meet underground injection well regulations under 40 C.F.R. 147. The extracted ground water will be treated to meet the above referenced drinking water standards prior to recharge.

Spent carbon from the ground water treatment system for removal of organics will be disposed of off-site, as well as any treatment residuals, consistent with applicable RCRA land disposal restrictions under 40 C.F.R. 268.

#### Chemical-specific ARARs:

Since the ground water at the Site is classified as IIb (GA by NYSDEC), drinking water standards are relevant and appropriate. Again, these include SWDA MCLs and 6NYCRR Ground Water Quality Regulations. However, achieving chemical-specific ARARs for ground water is dependent on remediation of upgradient sources. This is due to the fact that regardless of the Site cleanup, upgradient sources will continue to be a source of contamination to the ground water beneath the Site. EPA believes that the proposed remedial action will result in attainment of chemical specific ground water ARARs providing upgradient sources are remediated so that they no longer impact the Upper Glacial aquifer.

EPA may invoke a technical waiver of the chemical-specific ARARs if the remediation program indicates that reaching MCLs in the Upper Glacial aquifer is technically impracticable.

Until upgradient sources are remediated so that they no longer impact the Site, the remedial action will attain ground water cleanup levels equal to upgradient concentrations for certain contaminants.

#### 3. Cost Effectiveness

The selected remedy is cost effective and provides the greatest overall protectiveness proportionate to costs. Soil vacuuming and soil flushing, at a present worth of \$1,649,000 is more cost effective than excavation with off-site disposal, at a present worth of \$8,675,000, and offers an equivalent degree of protectiveness. The \$12,095,000, 30-year present worth cost associated with the selected ground water treatment, is the most cost effective of all the alternatives. The \$12,095,000 cost associated with ground water treatment is cost effective in that the remedy provides the greatest overall protectiveness compared with the \$66,000 cost associated with no action, which is not considered to be protective.

# 4. Utilization of Permanent Solutions and Alternative Treatment(or Resource Recovery) Technologies to the Maximum Extent Practicable

The selected remedies represent the maximum extent to which permanent solutions and alternative treatment technologies can be utilized in a cost effective manner for the Site. This

is evident by the selection of soil vacuuming, clearly an innovative technology. After treatment is complete, the soil will no longer be contributing contaminants to the underlying aquifer.

The ground water treatment used in the selected remedy will reduce the contaminants of concern to levels protective of human health prior to recharge. In addition, of those alternatives which are protective of human health and the environment and comply with ARARs, EPA has determined that the selected remedy provides the best balance of trade-offs in terms of the five balancing criteria: long-term effectiveness and permanence; reduction of toxicity, mobility, or volume through treatment; short-term effectiveness; implementability; and cost. The modifying considerations of State and community acceptance also played a part in this determination.

The long-term effectiveness and permanence of the selected soil remedy is very high in that the surface soils would be treated and the contaminated areas restored. Ground water treatment also offers long-term effectiveness and permanence in that the remedial goal is to achieve ARARs except in those cases where upgradient concentrations prohibit such restoration.

Reduction of toxicity, mobility, or volume is also evident in the selected remedy. The treatment of on-site soil by soil vacuuming and/or soil flushing will effectively reduce the mobility of contaminants in surface soils. Ground water treatment has the goal of reducing contaminant concentrations in the aquifer to meet ARARs, effectively diminishing both toxicity and volume.

The short-term effectiveness and implementability of the selected soil remedy is high in that it would be conducted in-situ. The short-term effectiveness and implementability of the ground water treatment alternative is high in that there is no exposure to contaminated ground water during implementation and the remedy employs standard equipment and well developed technologies. As stated above, the cost associated with the selected remedy is the least costly of each alternative that is protective of human health and the environment and provides for treatment of the most hazardous substances.

#### 5. Preference for Treatment as a Principal Element

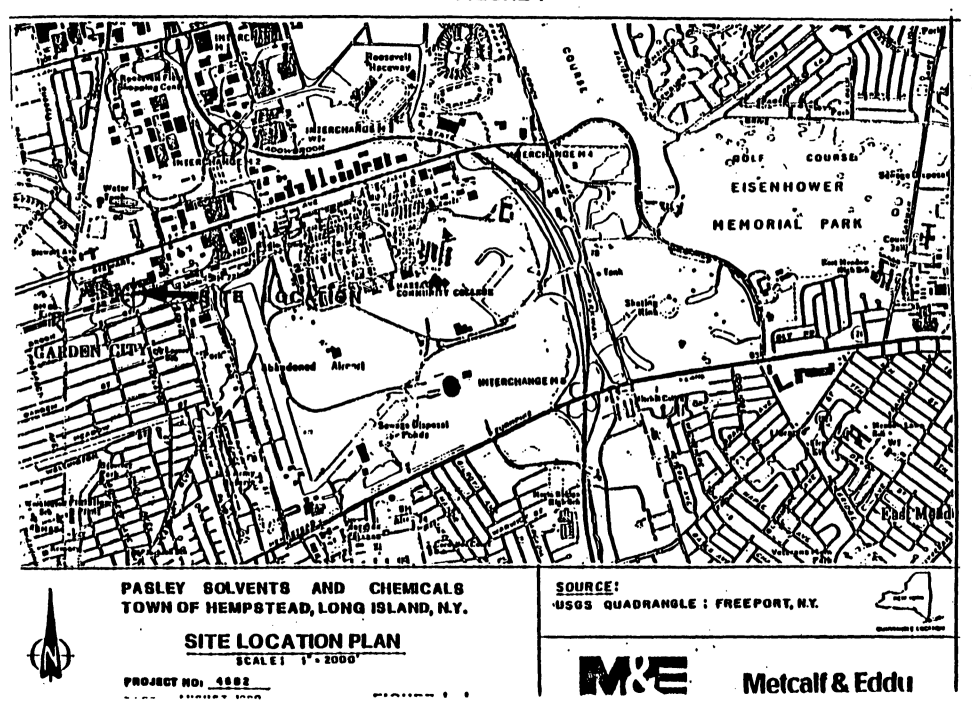
By treating the VOC contaminated soils and ground water by means of in- situ soil vacuuming and/or soil flushing, and air stripping respectively, the selected remedy addresses the principal threat posed by the Site through the use of treatment technologies. Therefore, the statutory preference for remedies that employ treatment as a principal element is satisfied.

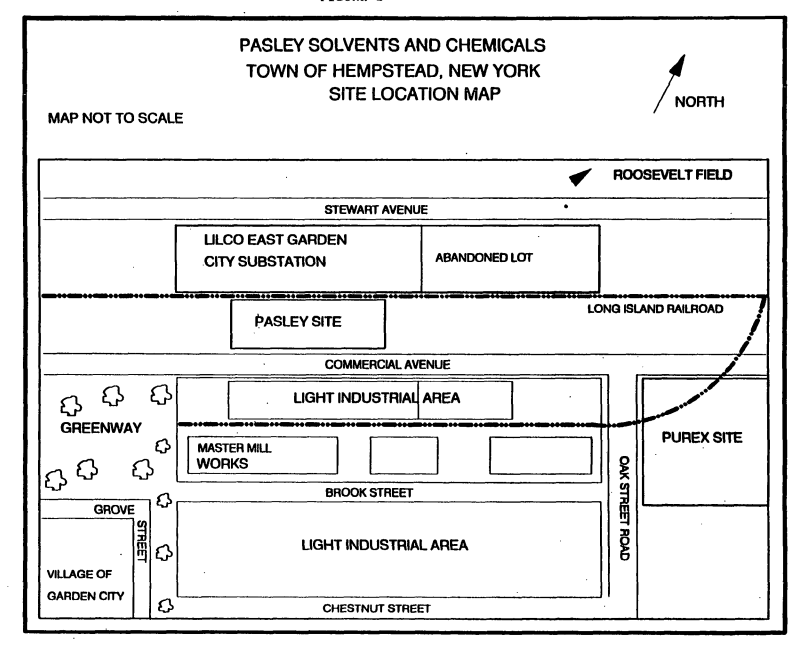
### XI. DOCUMENTATION OF SIGNIFICANT CHANGES

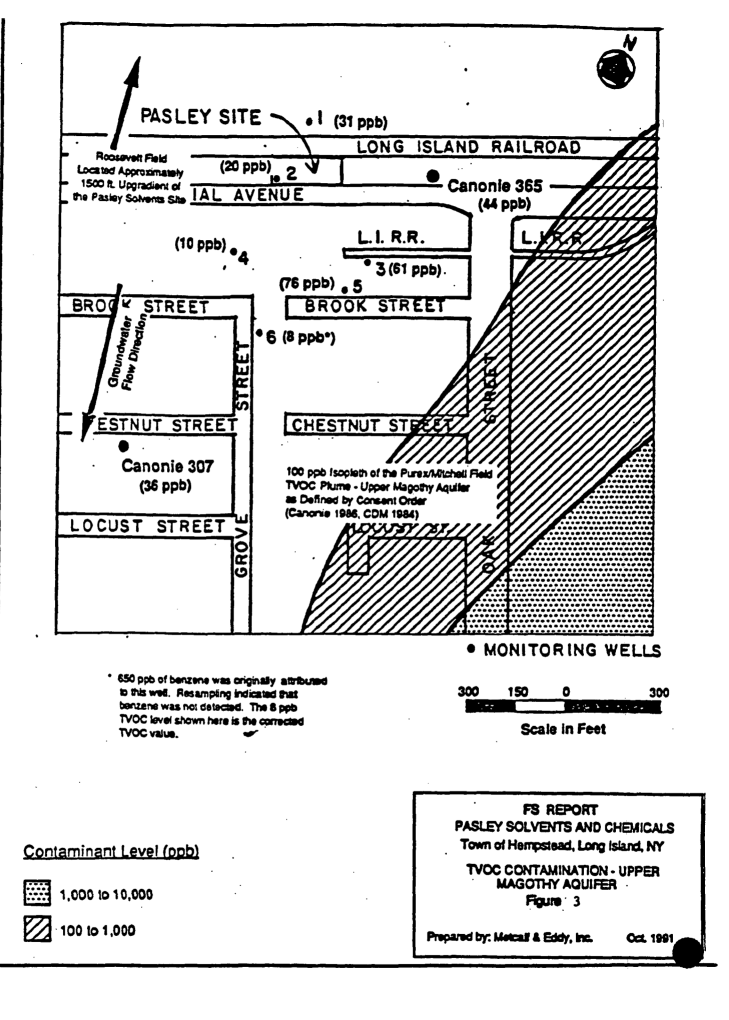
The Proposed Plan for the Pasley Solvents and Chemicals Site was released to the public on February 14, 1992. The Proposed Plan identified soil remediation Alternative 7 and ground water remediation Alternative 4 as the preferred alternatives. EPA reviewed all comments submitted. Upon review of the comments, it was determined that no significant changes to the preferred remedy, as it was originally identified in the Proposed Plan, were necessary.

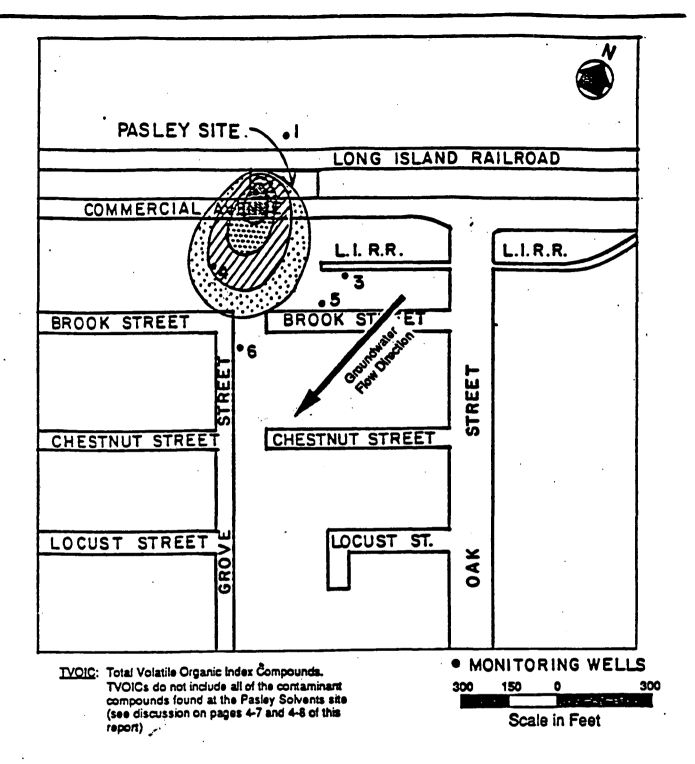
**APPENDIX 1** 

## FIGURE 1









## TVOIC Level (ppb)



> 10,000



1,000 to 10,000



100 to 1,000

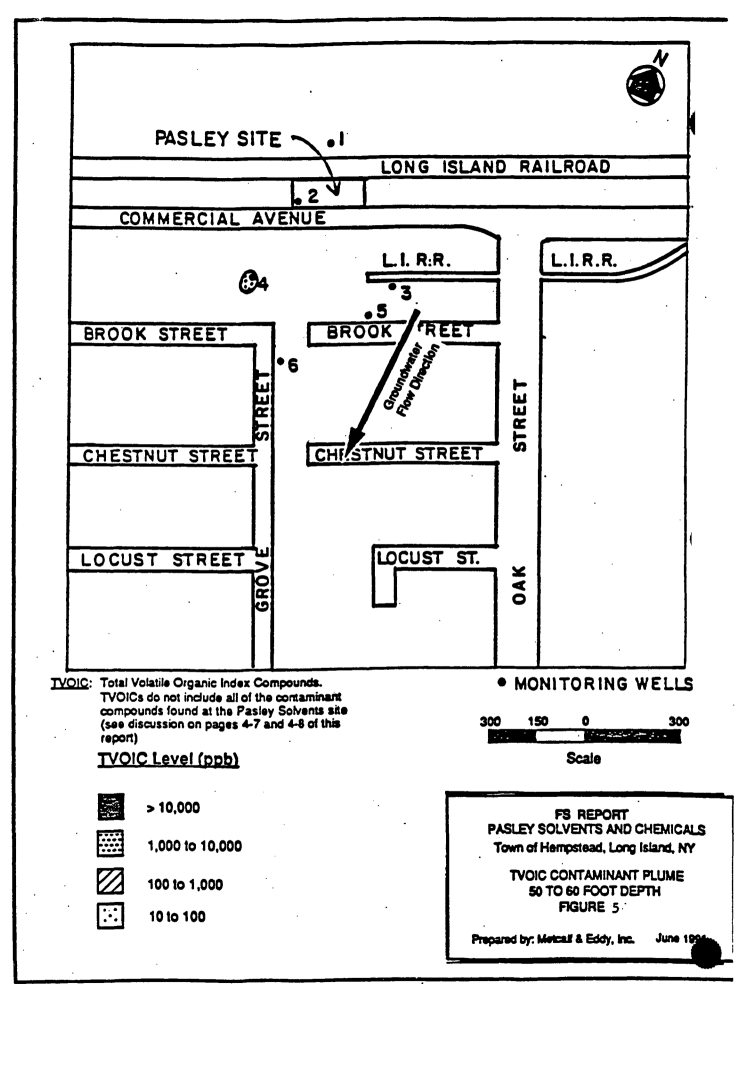


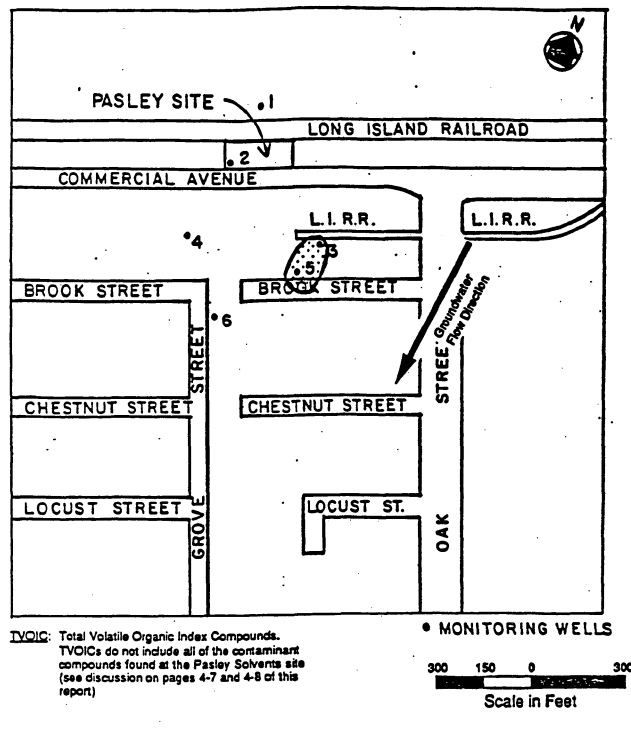
10 to 100

FS REPORT
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TVOIC CONTAMINANT PLUME 20 TO 30 FOOT DEPTH FIGURE § 4

Prepared by: Metcall & Eddy, Inc. June 1991





300

# TVOIC Level (ppb)

> 10,000

1,000 to 10,000

100 to 1,000

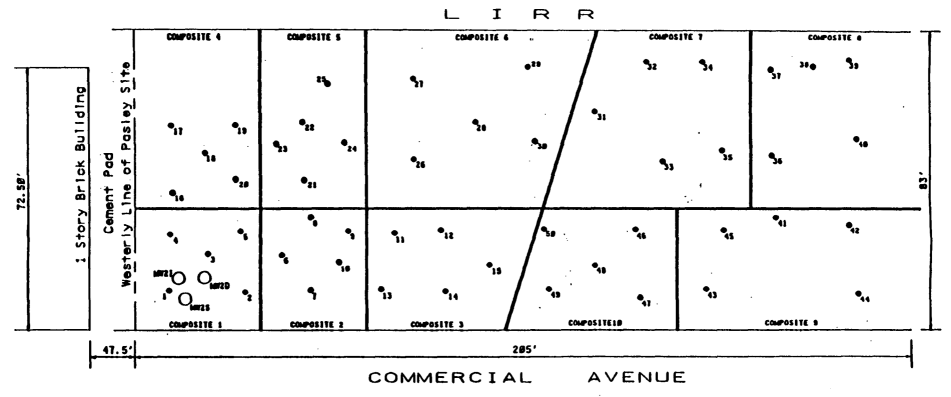
10 to 100

FS REPORT PASLEY SOLVENTS AND CHEMICALS Town of Hempstead, Long Island, NY

TVOIC CONTAMINANT PLUME 80 TO 90 FOOT DEPTH FIGURE 6:

Prepared by: Metcall & Eddy, Inc.

June 1991



Legend

- Surface Soll Samples
- O Monitoring Well

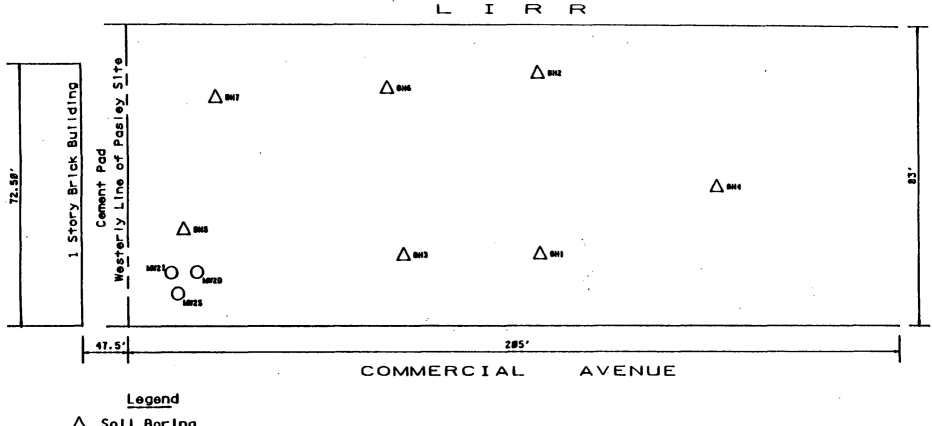


FINAL RI REPORT
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Town of Hempstead, Long Island, NY

ON-SITE SURFACE SOIL SAMPLING LOCATIONS Figure 7

Prepared by: Metcall & Eddy, Inc. 1

na 1991



△ Soll Boring Monitoring Well



**FINAL RI REPORT PASLEY SOLVENTS AND CHEMICALS** Town of Hempstead, Long Island, NY

**ON-SITE SUBSURFACE SOIL SAMPLING LOCATIONS** Figure 8

Prepared by: Metcall & Eddy, Inc. June 1991 **APPENDIX 2** 

PASILEY SOLVENTS AND CHEMICALS SITE FIRST ROUND GROUNDWATER SAMPLE RESULTS - MONITORING WELL

page P

SAMPLE NUMBERS	13	11	10	52	टा	21-NIP	20	33	31	30	43	41	10	54	51	50	63	61	60
UNITS	ug/l	ug/l	ug/l	ug/1	ug/l	ug/l	ug/l	ug/l	ug/l	ug/1	ug/l	ug/l	ug/1	ug/l	ug/l	ug/l	ug/1	ug/1	ug/I
MATRIX	Water	Weter	Water	Water	Water	Water	Water	Water			Water	Water	Water	Water	Water	Water	Water	Vator	. Vate
SAHPLE DATE (1990)	3-1	3-1	3-1	2-22	2-23	2-23	2-23	2-20	2-28	2-20	2-26	2-26	2-26	3-2	3-1	2-28	2-27	2-27	2-21
SAMPLE LOCATION		-LILCO			011-:	31TE			Lipp		(	REFINI	T	BA	00K 31	ľ		REDWI	17
VOLATILE ONCANTE COMPOUNDS															<del></del> -				
Methylene Chloride	-	-		-	-	R	-		-	_	-	•	-	-	•	-	•	-	•
Acetone	48J	130J	22J	1400J	1400J	580J	84J	510J	2200J	510J	TOUJ	3800J	130J	173	111	443	110J	260J	150J
Benzene	-	13	7	43J	10	11	3J	-	-	-	-	43	-	-	-	-	-	-	650
Chloroform	•	-	-	743	-	-	-	-	•	-	20	•	-	-	•	-	-	-	-
1,1-Diahloroethene	-	•	. •	641	-	-	W	-	-	-	-	•	-	•	•	-	-	. ••	-
1,1-Dichloroethane	•	•	-	630	-	•	IJ	-	-	-	21	-	-	-	•	-	-	·-	-
Trans 1,2-Dichlorosthess	•	•	- :	25000##	<b>2</b> 33	33	6	-	-	11	190	15	-	-	•	13	•	-	•
2-Butanone				•						•						· 🗑			
Ethy I benzene	-	•	•	510	-	-	-	-	-	-	42	•	-	-	•	-	-	-	-
Tetrechloroethene	21	•	•	160J	1,	ZJ	43	2	-	-	33	٩J	5	-	-	-	-	-	51
Tolwene	-	•	-	1100	1J	-	• .	-	-	•		-	•	-	•		-		-
Trichloroeth <b>ene</b>	11	•	15	140J	1,	IJ	8	-	<b>-</b>	50	<i>2</i> 50°,	1 15	10	•	2J	63	-	-	•1
1,1,1-Trichlo <del>roethans</del> .	32	-	•	3600	•	-	ZJ	2401	•	-	390		-	7	11	-	11	8	41
Chlorobenzene	-	•	-	510	-	-	-	-	-	-	•	-	•	. •	•	-	•	-	-
Xylenes (Total)	-	-	-	1100	-	-	-	-	-	. •	45	-	•	•	•	•	-	-	-
SEMINOLATELE ORGANIC COMPONIO	6								•										
bis(2-Ethylhoxyl) Phthelate		-		-	-	-	-	-	•	•	•		•	•	•	•	•	. 40	-
di-n-Butyl Phthelete	n			6.3	-		23		n		•		•	R	40		•	•	-
2-Methy Inaphthalone	-	-	-	110	-	-	-	-	-	-	16	-	-	-	•	•	•	-	•
Nephthalene	-	-	-	270	-	-	-	-	-	-	59	. •	-	-	•	•	•	-	•
Benzolo Aold	-	-	•	R '		•	-	-	-	-	1	-	-	-	-	•	•	-	-
Dibensofuran	-	•	•	5J	-	-		-	-	-	-	-	-	•	-	-	•	-	•
Phoneuthrone	•	-	-	5J	•	-	-	-	_	-	•	-	-	-	-		-	-	. •
41-n-Octvl Phthelate		2.1	-	•	•	-	-	-	-	-	**	•	-	-	-	-	-	-	-

MOTE: Only those compounds that are detected either as estimated, rejected or positive value is one or more sample are listed in this table.

J. Analyte present. Reported value may not be occurate or precise.

B. Unreliable Data Obtained. Data rejected by validator and is not usable.

Fire Fold diluted sample. See Appendix E for minimum detection limit attained.

Firey Fold diluted sample. Trans 1,2-Dichlorosethene concentration above the calibration range in this sample. See Appendix E for minimum detection limit attained.

(-) = Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

J. Shellow well; I. Intermediate well; D. a Deep well

BUP - Duplicate

H. \*\* Regated Compound not Present in Sample

E = estimated concentration due to interference.

\*\*\*U = Quantitation Limit is estimated.

June 19, 1991 Final RI Report PASILEY SOLVENTS AND CHEMICALS SITE TABLE | FIRST ROUND GROUNDWATER SAMPLE RESULTS - MONITORING WELL (Continued)

page Q	14000	1				. GAING						•			in we well
SAMPLE NUMBERS	E0-1	EB-2	EB-3	EB-4	EH-S	ED-6	E0-7	TO-1	TR-2	10-3	TB-1	18-5	TB-6	18-7	TR-B
UNITS	ug/l	un/l	ug/l	ug/l	ng/l	ug/l	ug/l	ug/1	ug/l	ug/l	ug/l	ug/l	Ug/1	UE/I	ug/1
MATRIX	Water	Water	Water	Veter	Valor	Water	Water	Vater	Water	Vator	Vater	Water	Veter	Water	Water
SAMPLE DATE (1990)	5-55	2-23	2-26	2-27	2-28	3-1	3-2	2-22	2-23	2-26	2-27	2-28	3-1	3-1	3-5
VOLATILE ORGANIC COMPOUNDS	<del></del>	<del>-</del>			<del> </del>				<del></del> -						
Chloromethane	3,3	•	•	-	-	· 🕳 '	•	•	-	-	-	-	_	-	•
Methylene Chloride	-		<b>.</b> .	-	-	-		, •	-	•	•	-	•	-	-
Acetone	•	110J	1200J	1800J	2300	83	R	_	25.J			-		-	•
Benzene	_ `	_	-	•	-	•	_	-		_	•	-	-	-	-
Chloroform	•	_	_	-	_		-	-	-	•	-	•	-	-	
1,1-Dichloroethene	-	-	-	-	-	•	-	-		-	•	_	<b>-</b> ·		-
1.1-Dichloroethane	•	_	-	_	_	-	-	-	•	-	-	•	•	-	•
Trans 1,2-Dighloroethens	•	_	-	-	-	-	-	-	•	•	-	•	•	-	•
2-Butanone							R								•
Ethylbenzene	-	•	-	-	-	-	_	-			-	-	•	-	•
Tetrechloroethene	23	23	5	21	-	-	~	-	-	-	_		-	-	•
Toluene		•	_	•	-	-	<b>-</b> '.	_	-	_	-	•	-	•	•
Trichloroethene	-		-	-	-	-	-	. 🕳	•	•	-	-	-	-	•
1,1,1-Trichloroethane	-	•		-	_	-	_	_	-		<b>-</b> ·	-	•	-	•
Xylenes	-	-	•	•		•	_		-	-	-	-	-	_	
SEMINOLATILE COGAMIC CONTOURS															
bis(2-Ethylhoxyl) Phthelate		-	_	-	-		-	•	•						•
di-n-Butyl Phthelate	-	-	61	-		Ü									
2-Hethy Inaphthalone	_	٠ .	-	-	-	-									
Naphthalone	-	•	•	•	-	-		•							
Benzolo Acid	•		•	-	-	-	-								
d1-n-Ootyl 'Phthelate	-	•	23	-	_	_	SJ								

NUTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Ammiyte present. Reported value may not be accourate or precise.

R = Unreliable Result Obtained. Data rejected by validator and is not weable.

<sup>(-)</sup> a Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory and field blanks.

TD - Trip Blanks enelysed for voletile organics only

ED - Equipment Blanks

PASILEY SOLVENTS AND CHEMICALS SITE June 14, 1991 TABLE 1 FIRST ROUND GROWNDWATER SAMPLE RESULTS - MONITORING WELL (Continued) Final RI Report

page N			_								- •	•	•		•		
SAMPLE NUMBERS	13	-11	10	28	51	21-009	SD	)3	31	3b	E0-1	ER-2	E0-3	EB-4	ED-5	En-6	E0-7
UNITS	ug/l	ug/l	ug/1	ug/l	ug/l	ug/l	ug/l	ug/1	ug/l	ug/1	ug/l	ug/l	ug/l	ug/l		ug/l	wg/L
MATRIZ	Water	Water	Water	Vator	Water	Vater	Water	Water	Water	Water	Water	Water	Water				Weter
SAMPLE DATE (1990)	3-1	3-1	3-1	2-22	2-23	2-23	2-23	2-28	2-28	2-28	2-22	2-23	2-26	2-21	2-28	3-1	3-2
SAMPLE LOCATION		-LILCO			011	-SITE			LIRA-								
HETALS				<del></del>			<del></del>				<del></del>						
Aluminum	<b>42100J</b>	4030J	155J	23400	. 1438	1360	416	15000	707	374	•	33.50	•		-	-	_
Ant Imony	-	-	-	-	-	_	-	-	-	-	•	-	•	-	-	•	-
Armento	-	-	-	-	-	-	-	-	•	•	•	-	-	-	-	•	•
Bartus	1113	68.8J	25.YJ	1128	51.1B	49.20	25.68	90.20	74.08	33.00	5.00	10. ID	•	18.50	20.00	19.6J	20.6J
Bory 1 tun	1.73	_	0.5	0.738	-	-	_	-	-	-		-	•	-	-	-	•
Cadetus	-	-	-	-	-	-	-	-	-	•	•	•	-	-	-	-	•
Coloium	36000J	18100J	14600J	13900	23800	23400	18100	<b>299</b> 00	18800	11400	10800	14900	-	15400	13200	13700J	15900J
Chrontus	23.1J	43.83	-	27.73	13.5J	17.6J	13.1J	90.6J	12.¶J	-	-	-	6.43	-	6.6B	9.63	•
Cobalt	10.9J	-	-	16J	-	-	_	_	-	-	-	-	-	-	•	-	-
Copper	1273	219J	28.5J	99.4	10.3B	8.7B	40.5	136	32.0	15.28	1440	3600	1.20	1320	4300	1880J	275J
Cyanide	70	•	-	15	-	_	-	20	-	-	-	10	10	-	-	-	-
Iron	28900J	4690J	368J	99100J	26 10 J	2450J	1570J	13500Ĵ	<b>21100J</b>	2290J	1713	120J	78.10	155J	331J	25.1J	99.2J
Lond				15.3	9.1	7.3	0.7	22.4	9.7	4.5		-,	-	-	-		
Hegnes I van	8330J	5510J	2760J	32808	48208	476B	27508	<b>46808</b>	26208	38608	15300	5610	-	5810	-	LOPOD	7390J
Hanganese	1060J	12200J	10.5J	1360	· 16100	15900	67.6	235	1760	221	-	0.9J	15.2	-	1.60	4.3J	1.8
Hereury	0.7	-	0.5	-	0.6	0.30	1.3	0.2 '	-	0.6	-	-	-	-	0.45	0.35	0.21
Miakel	53.61	129J	10.21	48.3	-	-	-	76.2	37.14	-	•	-	-	-	-	-	•
Potassium	4270J	31 10J	3290J	49608	16800	29308	6270	47 108	19208	-	•	•	-	-	-	1650J	1800
Selenium				-	-	-	-	-	•	-	•	-	-	•	-		
'81 lver	•	•	•	5. <b>6</b> J	-	•	-	•	-	-	•	•	•	-	•	•	-
Sodium	<b>4280</b> J	35300J	<b>28</b> 100J	390000J	<b>36700J</b>	34100J	26300J	18900J	33900J	2 <b>4</b> 500J	9250J	12300J	•	8060	1820J	7900J	12500
The Liture	-	-	•	•	•	-	•	-	•	•	•	•	. •	•	-	•	•
Vened1um	39.7J	-	5.93	27.60	•	-	7.00	10.18 -	•	•	•	•	6.30	•	•	•	•
21mg	1380	3200J	29.6J	859	67	51.2	917	1630	272	130	31.9	59.1	3.80	100	39.0	98.9J	29.1J

NOTE: Full Target Compound List metals are listed in this table.

J - Analyte present. Reported value may not be accurate or precise.

R - Unreliable Result Oblained. Data rejected by validator and is not meable.

(-) = Indicates compound was analyzed for but not detected.

3 - Shallow Well

I = Intermediate Well

D = Deep Well

EB = Equipment Blank

DUP = Duplicate

B = Trace levels (less than contract-required detection limits See Appendix F)

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page 3

F-43-4 -										
SAMPLE NUMBERS	<b>13</b>	41	40	55	51	50	63	61	60	-
UMITS	ter/1	ug/1	ug/1	ug/1	vg/1	ug/l	ug/1	ug/1	ug/l	
MATRIX	Water	Water	Vator	Water	Votor	**	Water	Votor	Water	
SAMPLE DATE (1990)	2-26	2-26	2-26	3-2	3-1	2-28	2-27	2-27	2-27	
SAMPLE LOCATION	****	-OREDINA	Y		ROOK STR	EET		-GREENVA		
NETAL3					<del></del>	<del></del>				-
Aluminum	97400	661	216	26400J	2390J	433	24000	734	747	
Ant Imony	39.938	-	_	-	-		-	-		
Arsento		-	-	_	. •	-	-	• .	-	
Bartum .	372	30.60	24.98	99.33	74.0J	30.1B	1190	70.69	66.5B	
Berylius	6.6	•	-	1.6J	-	_	2.138	_	139	
Codetus	-	•	-	-	•		-	9.5JB	•	
Calcium	22300	19500	10700	16600J	24900J	13300	24900	27900	17700	
Chrowing	63.6J	-	_	32.03	255J	-	25.83	22.03	16.5J	
Copper	194	40.8	41.5	76.6J	2523	85.2	71.2	30.2	64.1	
Cobalt	45.138	-	•	13.7J	19.6J	-		_	•	
Cyanida	to	-	to	-		, <b>-</b>		10	•	
Iron	152000J	3910J	903J	28300J	5050J	1180J	27600J	3180J	1860J	
Lead	39.6	9.5	0.2			5.0	17.0	11.0	11.1	
Hagnes ium	1130	31608	2770	4140J	12501	38208	40808	5050	31600	
Hanganese	9220	5630	119	659J	3380J	236	103	6610	1630	
Heroury	•	•	-	-	-	,	0.3	-		
Mickel	100	207	32.1JB	32.7J	310J	-	31.4.0	33.5JB	33.530	
Potessium	10200	26200	22008	5160J	6900J	-	25000	9550	3020B	
Selenius	•	-	•			-		•	•	
Silver	-	-	-	-	-	-	. 🍆	-		
Sodium	17000J	<b>L0078</b> 5	30400J	60601	37000J	<b>5</b> 5500J	13500J	306001	33700J	
Thelitus	-	-	•	5.73	•	-	•	•	•	
Vened Lus	94.0	-	-	40.7J	•	•	30.20	•		
Zino	1070	192	607	159J	2940	193	341	254	659	

MOTE: Full Target Compound List metals are listed in this table.

J a Analyte present. Reported value may not be accurate or precise.

R a Unreliable Result Obtained. Data rejected by validator and is not weable.

<sup>(-) .</sup> Indicates compound was analyzed for but not detected at a level eignificantly above the level reported in field and trip blanks.

<sup>5 .</sup> Shallow Well

I . Intermediate Well

<sup>9 .</sup> Doop Well

<sup>9 -</sup> Trace levels (less than contract required detection limit: See Appendix F)

# TABLE 2. SECOND ROUND GROUNDWATER SAMPLE RESULTS

page T

SAMPLE NUMBERS	25	21	21-DUP	2D	15	11	10	13		₹D	EB-1	EB-2	TB-1	10-2
UNITS	ug/l	ug/1	ug/1	ug/1	ug/l	ug/l	ug/l	ug/l	ug/l	ug/l	ug/l	ug/L	ug/l	ug/l
HATRIX	Water	Water	Water	Vator	Vater	•••		Water	Water	Water	Water	Weter	Weter	Water
SAMPLE DATE (1990)	4-19	4-18	4-10	9-18	9-19	1-19	9-18	4-18	4-18	4-18	4-18	9-19	4-10	4-19
LOCATION	371C-NO	OM-SITE	ON-SITE	OW-SITE	LILCO	LILCO	<b>LILCO</b>	CREENWAY	TAWEST	GREENMAY	V-,0	V-1,	V-10	4-13
VOLATILE ORGANICS		<del>*************************************</del>			<del></del>		<del></del>						<del></del>	<del></del>
Chloroethane	-	-	-	-	-	-	-	•	• .	•	77	-	-	_
Mothylene Chloride	163	-	-	•	-	•	-	•	• '	•	-	-	-	-
Acetone	-	-	-	-	-	200J	25 J	\$00°J	•	•	• .	373	_	-
1,1-Dichloroethene	<b>6</b> 2	-	-	• •	-	-	-	•	-	3.1	•	•	• -	-
1,1-Dichloroethane	300	-	-	-	•	•	•	20	-	-	•	•	-	-
Trans-1,2-Dichloroethene	37000 PB	•	-	3J	2J	•	3J	65			•	•	-	•
Chloroform	33	-	-	-	-	-	-	25	-	•	-	•	-	•
2 Butanone				R	•		R		n					
1,1,1-Trichloroethene	2800**	-	-	-	15	-	-	1BoJ*	-	•	•	-	-	-
Trichloroethene	320	-	-	11	12	-	12	150	- '	12	•	-	-	-
Benzene	20J	13	19	٩J	3.1	38	. 6	•	13	-	-	• •	-	-
Tet rechloroethese	91	-	-	5	24	-		29	23	7		•	-	-
Toluene .	750	-	23	_	-	-	-	-	2J		-	-	-	-
Ethylbenzene	340	-	-	-	-	-	-	18	-	-	-	-	-	-
Kylenes (Total)	210049	11	12	•	-	-		24	• .	-	-	-	-	_

HOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J - Analyte present. Reported value may not be accurate or precise.

H - Unreliable Result Obtained. Data rejected by validators and is not washis.

<sup>.</sup> Five fold diluted sample. See Appendix & for minimum detection limit attained.

<sup>\*\* - 250</sup> fold dilution. See Appendix & for winisum detection limit attained.

<sup>(-) -</sup> Indicates compound was analyzed for but not detected at a level significantly above the level report in inhoratory and field blanks.

S - Shallow Well

I . Intermediate Well

D . Doop Well

<sup>·</sup> ED . Equipment Blank

<sup>10 -</sup> Trip Blank

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#### PASLET SOLVENTS AND CHEMICALS SITE TABLE 2 SECOND ROUND GROUNDWATER SAMPLE RESULTS (Continued)

page U

SAMPLE NUMBERS	23	21	21-04P	20	13	11	10	43	41	ND.	EB-1	£0-2
MITS ·	ug/l	ug/l	ug/l	ug/1	ug/l	ug/l	ug/l	ug/l	ug/1	ug/l	ug/l	ug/l
MATRIX	Water	Water	Vater	Water	Water	Water	Water	Water	Vater	Water	Water	Water
SAMPLE DATK (1990)	4-19	4-18	9-18	9-18	4-19	4-19	4-19	4-16	4-16	4-14	4-18	4-19
LOCATION			9116			-LILCO-			-GREENVAY-			
SENT-VOLAYILE COULDICS		····	<del>.,,,,</del>	·		<del></del>			<del></del>	·		
Haphtha I one	180	-	•	-	-	-	-	23	-	-	-	•
2-Mothy imaphthalone	97	36	26J	-	-	-	-	9J	•	•	-	•
Acenaphthy lene	•	ŽI	16J	-	_	-	-	-	-	•	-	•
Acenaphthens	13	73	6.3	•	•	-	-	-	•	-	-	• .
Dibenzofurem	-	ŻJ	•	•		-	-	-	• '	-	-	•
Fluorene	3J	63	7,3	•		•	-	-	-	•	•	•
Phononthrone	-	3J	ŽĴ			•	•	•	-	•	•	-
Anthrocene	-	•	-	•	•	-	•	•	-	•	•	-
41-n-Butyl Phthalate	-					-	•				-	
Fluorenthene	-	•	•	•		_	•	•	• .	•	•	•
Pyrene	-	-	-	٠ ـ	•	_	•	•	•	•	•	•
bis(2-Ethylhoxyl) Phthalate	-	_	-	-	•	•	· •	•	•	-	•	R

MOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J. a. Analyte present. Reported value may not be accurate or precise.

<sup>#</sup> a Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-)</sup> a Indicates compound was analysed for but not detected at a level significantly above the level reported in laboratory and field blanks.

S . Shallow Well

I . Intermediate Well

D . Doop Well DBP . Duplicate

ES - Equipment Stank

# PASLEY SOLVENTS AND CHEMICALS SITE TABLE 3 MAY 1991 GROUNDWATER SAMPLE RESULTS - DEEP MONITORING WELL

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			<del>.</del>								00010140410
SAMPLE NUMBERS	MW-1D	MW-2D	MW-3D	MW-4D	MW-5D	MW-6D	TB-1	TB-2	EB-1	E8-2	MW-7D'
UNITS .	ug/l	ug/l	υg/l	ug/l	ug/f	บยู/เ	ug/l	ug/l	ug/I	ug/I	Ngu
MATRIX	Water	Water	Water	Water	Water	Water	Water	Water	Water	Water	Water
SAMPLE DATE	5/8/91	5/9/91	5/8/91	5/8/91	5/9/91	5/8/91	5/8/91	5/9/91	5/8/91	5/9/91	5/9/91
SAMPLE LOCATION	LILCO	On-Site	LIRR	Greenway	Brook St.	Greenway				'	DUP
VOLATILE ORGANIC COMPOU	NDS	<del></del>						<del></del>	<del></del>	•	····
Benzene	-	•	-	-	-	-	<u>.</u> .	•	-	•	0.9.
Bromochloromethane	A	•	•	A	-	-	R	R	R	R	-
Bromoform	A	R	R	R	R	R	R	P	R	. R	•
Chloromethane	•	•	•	-	<b>-</b> `	-	0.1	0.2	0.2	-	•
1,2-Dibromo-3-chloropropane	R	A	R	R .	R	A	R	R	R	A	A
Dichlorodifluoromethane	•	-	-	-	<b>-</b> ;	7.6	•	•	•	•	•
1,1-Dichloroethane	1.2J	5.1UJ	1.0UJ	1.9J	-	1.0UJ	•	1.0UJ	1.0UJ	1.0UJ	4.9
1,1-Dichloroethene	1.8	6.6UJ	-	2.9	-	-	•	•	•	•	4.3
Trans&Cis 1,2-Dichloroethene	2.2	87.8UJ	44.0	3.4	40.9	1.1	•	-	•	•	76.4J
Methylene Chloride	•	•	-	-	•	•	1.4J	2.5J	2.6J	2.23	•
Tetrachloroethene	7.2	7.6UJ	2.0	8.5	2.1	3.6	•	•	-	-	6.6
1,1,1-Trichloroethane	2.0	7.2UJ	-	2.9	-	5.4	•	•	•	•	6.7J
Trichloroethene	10.6	15UJ	99	16.3	91.0	9.1	•	-	-	-	14.5
trans-1,3 Dichloropropylene	A	A.	Я	A	R	R	A	R	A	Ħ	R
Carbon Disulfide	•	-	•	-	-	•	9.1J	•	-	-	•

Note: Only those compounds that are detected either as estimated, rejected, or positive values in one or more samples are listed in this table.

**UJ - Qualified Estimate** 

J = Analyte present. Reported value may not be accurate or precise.

R - Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) =</sup> Indicates compounds was analyzed for but not detected at a level significantly above the level reported in laboratory and field blanks.

TB - Trip Blanks analyzed for volatile organics only

EB = Equipment Blanks (Field Blanks)

<sup>\* -</sup> Sample MW-7D is a duplicate sample from well MW-2D

PASLET SOLVENTS AND CHEMICALS SITE
TABLE 4. ON-SITE SURFACE SOIL SAMPLE RESULTS

page

SAMPLE NUMBERS(Composites)	1-5	1-50UP	6-10	11-15	16-20	21-25	26-30	31-35	36-40	41-45	46-50	
UNITS	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	
MATRIX	Soll	Soil	Soll	3011	Soll	So11	Soll	3011	Soft	So11	Soti	
SAMPLE DATE (1989)	9-13	9-13	9-13	9-13	9-13	9-13	9-14	9-19	9-14	9-14	9-14	
SAMPLE DEPTH (In.)	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	
SPH-VOLATILE ORGANICS			<del></del>		<del></del>		·····		<del></del>			
1,2-Dichlorobenzene	-	-	-	-	-	-	2800J	1000J	-	890J	-	
Waphthalene	610J	_	3 <b>1</b> 00J	-	1000J	_	1300J	3803	<b>\$3000</b>	-	•	
2-Mothy lenaphthelene	1100J	•	10000	-	4600J	1100J	4000J	•	9800J	<b>460J</b>	2000J	
Fluorene	470J	-	•	-	-	-	-	-	•	•	- •	
Phononthrong	5700J	500J	1600J	1400J	2300J	1900J	900J	-	620J	370J	•	
Anthrocene	2600J		-	-	-	530J	•	•	-	•	•	
di-n-Butyl Phthalate	2500J	2000J	-	370J	68000	1700J	290J	4303	150000	<b>4800J</b>	•	
Fluoranthene	11000	700J	360J	400J	11003	1900J	370J	•	•	•	•	
Pyrono	£00#8	760J	680J	570J	4200J	1600J	620J	-	670J	• •	600J	
Benzo(a) Anthrecene	5000	_	-	-	-		-	•	-	•	•	
bis(2-Ethylhoxyl) Phthelate	-		-	-	-	` R	R	•		120000	Ħ	
Chrysene .	6000J	7903	-	280J	2900J	1100J	<b>360J</b>	-		-	-	
Benzo(b) Fluoranthene	LOOP	730J	-	-	1100J	990J	330J	•		-	-	
Benzo(k) Fluoranthene	1800J	-	-	-	-	450J	-	•	•	•	• ,	
Benzo(a) Pyrene	3300J	370J	-	-	1600J	` 750J	-	-	•	-	-	
Indeno (1,2,3-od) Pyrene	1600J	-	•	-	-	-	-	-	•	-	•	
Dibens (a,h) Anthrecens	-	-	-	-	-	-	•	•	-	-	-	
Benzo (g,h,1) Perylene	1500J	•	-	-	BAOJ	••	-	•	-	•	-	

NOTE: Only those compounds that are detected as either estimated, rejected or positive is one or more samples are listed in this table.

J - Analyte present. Reported value may not be accurate or precise.

R . Unreliable Result Obtained. Data rejected by validator, and is not usable.

<sup>.</sup> Medium level analysis with 20 fold dilution. See Appendig 8 for minimum detection limits attained.

<sup>\*\* -</sup> Medium level analysis with 15 fold dilution. See Appendix E for minimum detection limits attained.

<sup>\*\*\* .</sup> Medium level analysis with 30 fold dilution. See Appendix & for minimum detection limits attained.

<sup>(-) -</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported is laboratory or field blanks.

BUP - Duplicate

<sup>1-5 -</sup> Composite of samples 1-5

PASLEY SOLVENTS AND CHENTCALS SITE TABLE "44 SURFACE SOIL SAMPLE RESULTS (Continued)

page B

SAMPLE NUMBERS (Composites)	1-5	1-SMIP	6-10	11-15	16-20	21-25	2610	31-35	16-40	41-45	46-50	ER-01	
UNITS	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	ug/l	
MATRIX	Sett	3011	3011	Soil	3011	Soll	Soll	Soll	Soil	3011	5011	Water	
SAMPLE DATE (1989)	9-13	9-13	9-13	9-13	9-13	9-13	9-14	9-14	9-14	9-19	9-14	9-13	
SAMPLE DEPTH (In.)	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	,-,,	
ET4.3	<del>, . ,</del>				<del></del>		<del></del>				····	********	<del></del>
Atuninum ·	6330	5530	5030	3920	96 30	7450	7650	7970	4260	7620	2670	-	
Ant Leony	-	· -	-	LOS	16.7J	•	-	•	•	-	-	•	
Arseniq	11.5J	9.9J	16.73	14.25	4.3J	R	9.25	17.1J		1.43	2.88	* <b>=</b>	
Bartus	<b>94.6</b>	86.7	19.Z	49.5B	425	144	116	66.9	113	159	29.58	-	
Bory I tun	•	•		-	-	-	-	•	-	•	•	• .	
Code i um	•	-	-	-	-	-	-	-	-	-	-	-	
Caloium	442003	333003	180003	19000J	44300J	39800J	30800J	22700	5530J	18900J	19600J	217B	
Chromium	58.8	42.8	28.7	18.1	47.7	52.8	58.1	42.9	34.1	45.6	7.7	•	
Cobelt	4.58	4.38	4.98	2.98	6.58	7.6B	5.5B	6.90	3.80	6.90	-	•,	
Copper	51.2J	<b>40.4</b> J	<b>40</b> J	46.1J	72.3J	59.5J	53.5J	11.63	74.83	67.53	15.93	•	
Cyanide	-	_	-	-	0.28J	0. <b>4</b> 5J	-	_	1.13	0.343	0.39J	•	
l ron	23200	<b>22800</b>	17600	20700	41800	30700	23500	23100	20300	39900	11100	143	
Lead	335J	340J	19 <b>4</b> J	577J	1230 <b>J</b>	750J	` <b>416</b> J	22 <b>6</b> J	457J	709J	1343	•	
Magnes 1 up	22800	16400	8700	9430	22100	19300	15000	10900	2820	7460	7690		
Manganese	145	128	77	110	212	177	123	107	99.2	265	73.6		
Heroury	0.14J	0. 19J	-	· 0.14J	-	-		-	•	•	-	-	
Mickel	16.3	12.9	15.7	-	14.2	17.0	15.6	25.1	10.00	17	7.7B	•	
Pot <b>assius</b>	7120	4120	5118	6848	50 <del>98</del>	7018	5748	975B	3608	5680	3520	•	
Se l'entur		-	-		-	•	•	•	-	-	-	•	
Silver	•		-	-	-	-	-	•	•	•	` -	-	
Sodium	1468	1548	1390	66.88	1298	95.08	1658	2100	1080	1170	49.38	106B	
The ilium	1.9J	2.5J	3.1		3-3	2.50	2.09	-	-	-	-	•	
Yenedium	27.0	26.9	24.6	17.1	32.3	20.7	25.8	37.7	29.6	28.6	10.10	•	
Zino	215	202	131	179	455	308	292	133	555	658	712	20.3	

J - Analyte present. Reported value may not be accurate or precise.

R . Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) -</sup> Indicates compound was enalyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

DOP . Duplicate

ES . Equipment Blank; TB . Trip Blank

U . Trece level (less then contract required detection limit: See Appendix F)

UJ . Estimated detection limit

<sup>1-5 .</sup> Composite of samples 1-5.

PASLEY SOLVENTS AND CHEMICALS SITE TABLE 4 ON-SITE SURFACE SOIL SAMPLE RESULTS (Continued)

pege C

SAMPLE NUMBERS	IAB	IAR-DUP	248	3AB	PAR	SAB	6AB	748	BAB	9AH	10AB	TIAB
CTINU	wg/kg	ug/kg	ug/kg	ug/kg	vg/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg
MATRIE	Soil	3011	Soli	Soll	Soil	3011	Soll	Soil	<b>5</b> 011	3o11	3011	Soil
SAMPLE DATE (1989)	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13
SAWLE DEPTH (In.)	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12
VOLATILE ONGARIES										<del></del>		
Vinyl Chloride	•	130J	89J	77	-	250	68	-		•	210J	-
Chloroethans	•	-	-	-	-	-	-	•	•	-	-	•
Hethylene Chloride	100J	110J	<b>32J</b>	-	12J	13J	-	•	•	' <b>p</b>	790J	-
Agetone	46J	-	-	-	-	150	-	-	66J	•	•	-
1,1-Dichloroethene	90J	5001	52J	19J	30	49	28	•	160J	30	<b>36J</b>	-
Trans-1,2-Dichloroethene	600J	930J	7400°J	950	240	230	160	530J	240J	530	1100"	-
Chloroform	-	-	· -	•	10J		•	-	•	•	-	-
2-Butanone	BZJ	R	-		-	- '		,	•	_	-	•
1,1,1-Trichlorosthane	-	17J	_	_	-	•	•	•	11J	-	•	-
Trichloroethene	52J	97J	240J	57	49	12J	- 31	240J	32J	ZZJ	230J	-
Tetrachloroethene	213	110J	1403	130	81	41		220J	543	71	100J	32
Toluene	•	•	•	193	48	12J	•	10J	•	•	38J	-
Chlorobensone	-	_	•	,				-	113	-	•	-
Ethylbensene	_	-	•	•		-		-	•	•	-	-
lylene	-	_	-	65J	•	-	. =	_	19J	44	16J	16J

HOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R = Unrollable Result Obtained. Data rejected by validator and is not usable.

<sup>\*</sup> a Medium level analysis with ten fold dilution. See Appendix E for minimum detection limits attained.

(-) a Indicates compound was analysed for but not detected at a level significantly above the level reported in laboratory or field blanks.

DUP - Duplicate

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PASLET SOLVENTS AND CHEMICALS SITE TABLE - 4 ON-SITE SURFACE SOIL SAMPLE RESULTS (Continued)

pege D

SAMPLE NUMBERS	12AB	13AB	INAR	1548	16AFI	17AB	1848	19AB	20AB	ZIAD	22AB	23AB
UNITS	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg
MATRIX	<b>5</b> 011	So 1 1	3011	Soft	3011	3011	Soil	3o11	3011	Soft	3011	Soll
SAMPLE DATE (1989)	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9-13	9~13
SAMPLE DEPTH (In.)	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12
VOLATILE ONGANICS		<del></del>	<del></del>		<del></del>						<del>////</del>	· <del>····································</del>
Vinyl Chloride	670J	-	-		160J	-	_	-	-	•	-	-
Chloroethane	-	-	-	-	19J	-	-	•	•	•	•	-
Methylene Chloride	190J	15J			n				50J	59J	-	R
Agetone	· 720J	-	453	52	19000*	-	500	81	950J	290J	38J	•
1,1-Dichloroethene		-	· 🕳	-	•	-	-	-	-	-	-	467
1, 1-Dichloroethene	580J.	<b>9</b> J	•	-	270J	69	23J	79	120J	270J	•	73J
Trans-1,2-Dichloroethens	24000J	84	-	-	700J	20	160			25000J		n
Chloroform	-	· 🕳	-	•	-	-	-	-	-	-	•	_
2-But anone	-		-	R	110J		59	•	-	-	•	-
1,1,1-Trichloroethene	870J	•	-	-	-	6900	-	•	260J	470J	-	
Trans-1,3-Dichloropropens	-	•	-	-	-	:-	•	•	-	-	-	-
Trichloroethene	3500*	•	13J	28	190J	150	12J	517	24000*	19000*	3700*	R
Benzene	-	-	-	-	-	-	•	•	-	. 6J	-	-
Tet rechloroethens	370J	-	21J	•	493	<b>I</b>	19J	42	7600	84000*	87000°	90000°
Toluene	5403	•	-	8,3	12000°J	-	54000*	750	7603	2103	910J	•
Chlorobenzene	-	•	•	-	-		-	•	-	•	•	<b>48</b> J
Ethy I bensene	34J	-	-	-	35J	-	72	•	25.1	13J	380J	120J
Xylene (Total)	17000*	•	-	-	260J	-	290	•	453	15J		240J

NOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R . Unreliable Result Obtained. Data rejected by validator and is not weable.

<sup>.</sup> Medium level analysis with ten fold dilution. See Appendix & for minimum detection limit attained.

<sup>(-) .</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

VJ . Quantitation limit to estimated.

TABLE: 4: ON-SITE SURFACE SOIL SAMPLE RESULTS (Continued)

page E

SAMPLE NUMBERS	ZHAB	25AB	26AB	RATS	ZHAH	29AB	30AB	JIAR	32AB	33AH	34AB	35AP	
UNITS	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	
MATRIX	Soll	3011	3011	Soll	Soll	5011	Sofl	Soll	Soll	Sott	Soft	Sott	
SAMPLE DATE (1989)	9-13	9-13	9-14	9-14	9-14	9-14	9-14	9-14	9-14	9-14	9-14	9-14	
SAMPLE DEPTH (In.)	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	
VOLATILE ONGANICS					<del></del>				<del></del>				<del></del>
Vinyl Chloride	•	-		•	•.	-	-	-	•	•	-	-	
Chloroethene	•	-	-	•	-	-	•	•	•	•	•	-	
Hethylene Chloride	723	•	n	-	•	-	150J	243	35J	60	<b>NOJ</b>	170J -	
Acetone	•	-	•	-	-	· •	670J	-	-	-	-	6ZJ	
1, 1-Diahloroethene	39J	-	50	-	-	-	570 <b>J</b>	110J	-	•	-	120J	
Trans-1, 2-Diobioroethene		3400*	910	42	233	17J	82000*	16000*	-	31	30J	•	
Chloroform	-	_	-	-	-	193	-	350J	113	41	860J	• .	
2-Butanone			533	-	-	49J	-	_		69J	A	•	
1,1,1-Trichloroethene	140J	-	160	-	-	-	· •	LOS	121	-	-	59J	
Trans-1, 3-Dichloropropens	-	-	•	20	193	250	_	_	• ,	•	-	•	
Trichloroethene	5900ª	3300*	86	-	-	<b>-</b> √	7003	1200004	130J	71	790J	600J	
Benzene	•	-	-	•	-	•	87J	16J	•	•	-	-	
Tetrechloroethens	13000	48000®	140	86	ZZJ	150	21000	120000*	110	57	LOSP	34000*	
Toluene	21 J	170J	21	-	•	36	410000	9003	•	•	200J	170J	
Chlorobensens	•	-	•	-	-			-	•	•	•	•	
Ethy I benzene	•	140J	•	-	-	-	710	15J	•	-	-	300J	
Ivlene (Total)		5500*	100	_	-	-	27000°	613	•	-	-	21000	

NOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J - Analyte present. Reported value may not be accurate or precise.

<sup># =</sup> Unreliable Result Obtained. Data rejected by validator and is not unable.

<sup>.</sup> Medica level analysis with ten fold dilution. See Appendix E for minimum detection limit attained.

<sup>(-) .</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

#### PASILET SOLVENTS AND CHEMICALS SITE TABLE 4 ON-SITE SURFACE SOIL SAMPLE RESULTS (Continued)

page P

SAMPLE NUMBERS	36AB	37AR	38AB	39AD	HAOP	TIAB	42AB	TJAR	HAB	45AB	46AB	\$7AB	
et i mu	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	
MATRIE	3011	501 L	Soft	Soll	3011	Soft	3011	3011	3011	3011	Sofi	Soft	
Sample date (1989)	9-14	9-14	9-14	9-14	9-14	7-14	9-14	9-14	9-14	9-14	9-14	9-14	
SAMPLE DEPTH (In.)	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	6-12	
FOLATILE ORGANICS							<del></del>		<del></del>	<del></del>			
Vinyl Chloride	•	-	. •	-	-	-	-	•	-	-	•	-	
Chloroethene	•	-	-	-	•	-	-	-	•	•	-	•	
Methylene Chloride	55J	17.3	-	-	-	<b>45J</b>	1103		-	33UJ	1403	-	
Acetone	37.3	_	-	-	-	-	303	-		-	590J	-	
1,1-Diahlarosthans	LOP	63	-	•	•	-	72J	-	-	LOP	273	-	
Trans-1, 2-Dichloroethene	49J	LOPS	> 15J	•	-	LOPI	12000°J	-	-	160J	100 <i>J</i>	-	
Chloroform	12J	-	20J	-	50J	60J	690J	-	•	<b>410</b> J	17000*	ÐJ	
2-Butanone	. 51		59UJ	-			843	53J			<b>460J</b>	57J	
1,1,1-Trichloroethane	160J	360J	-	113	35.1	•	300J	-	-	<b>UI</b>	-	-	
Trans-1, 3-Dichloroethene	-	-	~	-	-		-			-	-	-	
Trichlaroethene	570J	2600°	LOOP	188	LOTP	<b>4016</b>	120000	72	11,	Locobs	67.3	-	
Benzene '	-	-	-	-	-	•	-	•	•	-	61J	-	
Tetrachloroethene	14000*	55000		640J	270000°	230000°	70000°	92	10	650J	25 J	-	
Tolvene	<b>4600</b>		-	-	34J		6700	•	•	27 J	<b>9</b> 7J	-	
Chlorobenzene	-	_	-	-	-	<b>~</b> `	-	•	•	•	-	-	
Ethylbenzene	3000ª	2700	-	-	-	- '	620	•	-	•	-	-	
Tylene (Total)	1100	35000°	_	-	-	73J	20000®	•	-	LOI	-	•	

HOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

B = Unreliable Result Obtained. Data rejected by validator and is not weable.

<sup>. -</sup> Medium level analysis with ten fold dilution. See Appendix E for minimum detection limit attained.

<sup>(-)</sup> a Indicates compound use analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

UJ - Quantitation limit is estimated.

PASLEY SOLVENTS AND CHEMICALS SITE TABLE 4 " ON-SITE SURFACE SOIL SAMPLE RESULTS (Continued)

page G

SAMPLE NUMBERS	HAD	49AB	5040	18-1	BLK-1	
UNITS	ug/kg	ug/kg		ug/l	ug/l	
MATRIX			3 0	•	•	
SAMPLE DATE (1989)	7-14	9-14	9-14	9-14	9-14	
SAMPLE DEPTH (In.)	6-12	6-12				
VOLATILE ORGANICS	***************************************				<del></del>	_
Vinyl Chloride	-	-	370J	•	-	
Chloroethane	-	-	-	-	-	
Methylene Chloride	-	-	L0#1	-	-	
Acetone	-	-	750J	-	-	
1,1-Dichloroethane	-	-	L091	-	-	
Trans-1, Z-Dichloroethone	-	-	20000*	-	-	
Chloroform	•	-	-	-	-	
Z-Butanone	52J	391	320J			
1,1,1-Trichloroethene	•	-	52J	•	~	
Trichloroethene	•	•	120J	•	-	
Tetrachloroethene	50	-	130J	•	-	
Toluens	-	-	330J	•	-	
Ethylbenxene .	•	•	•	-	-	
Iylone (Total)	-	•	220J	-	-	

MOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J's Analyte present. Reported value may not be accurate or precise.

R - Unreliable Result Obtained. Date rejected by validator and is not usable.

- Medium level analysis with ten fold dilution. See Appendix E for minimum detection limit attained.

<sup>(-)</sup> a Indicates compound use enalysed for but not detected at a level significantly above the level reported is laboratory or field blanks.

TB . Trip Blank

M.E-1 - Soutpoont Blank

#### PASLET SOLVENTS AND CHEMICALS SITE TABLE 5. ON-SITE SOIL BORING SAMPLE RESULTS

page H

SAMPLE NUMBERS	BH-1A	BH-18	BH-2A	BH-28	BH-3A	60 16	3A-DUP	39 040	600 BA	BH-48	DIL CA	60.50		-50-75	
DANTE NUMBRS BNITS					_		-	-			BH-5A	BH-50			
MATRIX	ug/kg Soll	ug/kg So11	ug/kg Soll	ug/kg Soll	ug/kg Soll	ug/kg Soll	ug/kg 3o11	ug/kg Soll	ug/kg Soil	ug/kg Soll	ug/kg	ug/kg	ug/kg	ug/kg	
SAMPLE DATE (1989)	9-19	9-19	9~20	9-20	9~22	9-22	9-22	9-21	9-20		3011	3011	Soll	Soll	
SAMPLE DEPTH (Ft.)	12-14	24-26	12-14	24-26	-	24-26		24-26	12~14	9-21 . 24-26	9-21 12-14	9-21 22-24	9-25 12-14	9- <i>2</i> 5 22-24	
SERI-VOLATILE ONGLATICS															
Maphthalene	437	450	5500	2900	1200J	290J	1700	-	-	-	-	-	-	-	
2-Hethy Insphthalene	681	2500	5500	3000	13000	2400	15000	-	-		-	-	790	• •	
Acenephthene		_	-	-	760J	-	-	-	-	-	-	-	-	-	
Dibenzofuran	-	-	-	-	-	. 220J	1100J	-	-	-	-	-	-	-	
Fluorene	-	-	280J	160J	-	3403	1400J	-	-	-	-	-	190J	-	
Phenanthrene	-	340J	390J	220J	2500	880	2300	-	-	693	~	-	2603	-	
Anthrecene	-	-	-	-	-	86J	240J	-	_	-	-	_	-	-	
d1-n-Butyl Phthalate	310J	270J	390J	490	150J	130J	120J	77J	680	120J	120J	76J	140J	120J	
Fluoranthene	-	33J	100J	59J	200J	321	230J	-	-	-	-	-	•	-	
Pyrene '	100J	873	1903	100J	240J	<b>4</b> 2J	280J	-	-	-	-	-	213	-	
bis(2-Ethylhexyl) Phthelate		R	R	4900	_	-	-			R		R	-	-	
Chrysene	-	-	160J	55J	-	-	`. <del></del>	-	-	-	-	-	-	• •	
di-n-Octyl Phthalate	87J	. 170J	_	131	, -	-	-	-	-	-	-	-	-	-	

NOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R . Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) =</sup> Indicates compound was analysed for but not detected at a level significantly above the level reported in laboratory or field blanks.

BH . Borehole

DUP - Duplicate

TABLE 5 ON-SITE SOIL BORING SAMPLE RESULTS (Continued)

page I

SAMPLE NUMBER	BH-7A	BH-78	BH-8A	8H-8B	BH-EB-1	BH-EB-2	BH-EB-3	BH-EB-4	8H-EB-5	BH-EB-6	
mi t9	ug/kg	ug/kg	ug/kg	ug/kg	ug/l	ug/l	ug/l	ug/l	ug/l	ug/l	
IATRIZ	3011	<b>5</b> 011	Soil	So11	Soll	Vator	Water	Water	Water	Water	
IAMPLE DATE (1989)	9-25	9-25	10-23	10-23	9-19	9-20	9-21	9-22	9-25	10-23	
SAMPLE DEPTH (Ft.)	12-14	22-24	12-14	22-24							
MOT-VOLATILE ORGANICS											<del></del>
Maphthalene	650	COOL	•	3600	-	<b>-</b> ,	-	-	-	-	
2-Mothy Inaphthalone	2600	9100	7500	4800	-	_	-	-	-	-	
Aconaphthono	-	-	-	_	-	-	-	-	-	-	
Dibenzofuren	280J	550J	<b>480</b> J	_	-	-	-	-	-	-	•
Diethyl Phthelate	-	-	-	-	-	-	-	. 93	-	~	
Fluorene	420	820J	~	-	-	-	-	-	-	-	
Phenenthrene	910	1000J	560J	490	-	-	-	-	-	-	
Anthrecene	170J	-	-	-	-	-	-		-	-	
Di-n-Butyl Phthelete	130J	210J	120J	42J	110	130	828	65B	7J	IJ	
Fluoranthene	71J		47J	<b>42</b> J	-	-	-	-	-	•	•
Pyrene	270J	150J	59J	39J	-	-	-	-	-	-	
bis (2-Ethylhexyl) Phthalate			410J	1000	••		-	-	-	-	
Chrysene	120J	-	-	193	-	_	-	•	-	_	
di-n-octyl Phthalate		-	_	1103		_	_	_	_	_	

MOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R = Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) =</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

BH = Borehole

EB . Equipment Blank

B = Trace level (less than contract required detection limit: See Appendix F)

TABLE 5 PASLET SOLVENTS AND CHEMICALS SITE ON-SITE SOIL BORING SAMPLE RESULTS (Continued)

page J

							******			THE TEST		~				
SAMPLE NUMBER	BH-14	BH-18		BH-28	BH-3A		_	3B-DUP		BH-48	BH-5A	-	BH-6A	BH-6B	BH-7A	BH-78
UNITS _	ug/kg	ug/kg		ug/kg	ug/kg		ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg	ug/kg
MATRIX	Soll	3011	Soil	Sott	Soil	Soll	Soll	Soil	Soll	Soil	Soll	Soll	Soil	Soll	Sofi	Soil
SAMPLE DATE (1989)	9-19	9-19	9-20	9-20	9-22	9-22	9-22	9-21	9- <i>2</i> 0	9-21	9-21	9-21	9-25	9-25	9-25	9-25
SAMPLE DEPTH (Ft.)	12-14	24-26	12-14	24-26	12-14	24-26	12-14	24-26	12-14	24-26	12-14	22-24	12-14	22-24	12-14	22-24
VOLATILE ONGANICS					<del></del>						·			<del></del>	*****	<del></del>
Methylene Chloride	-	1200J	790J	-	710J	-	-	-	•	-	-	-	17J	-	-	360J
Acetone	130J	-	-	1 10J	-	11J	-	34J	19J	•	-	24J	18J	-	-	-
1,1-Dichloroethane	12J	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Trans-1, 2-Dichloroethene	99	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Chloroform	<b>9</b> J	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
2-Butanone	R	1	R	R	R	R	R		#	R	Ħ	•		R	R	R
1,1,1-Trichloroethane	-	-	-	-	-	-	-	-	-	-	•	-	-	-	-	320J
Trichloroethene	100J	-	-	-	-	160J	-	-	-	-	-	-	_	-	-	2800
4 Methyl - 2-pentanone	-	-	-	-	-	-	-	-	-	-	-	-	-	3600	-	-
Tetrachloroethene	21J	-	12000	-	480J	55J	680J	-	21J	-	-	-	-	9700	520	21000
Toluene	-	200	1200	-	4400J	<b>4</b> 60J	6900J	-	7,3	-	-	-	-	230J	21J	590J
Ethy1benzene	-	-	500J	-	1600J	510J	2200J	-	-	•	-	-	-	540J	23J	330J
Total Tylene	•	390	1000	-	6000J		8300J	-	-	12J	-	-	_	2000	71	750
1,1,2-Trichloroethane	-	-	-	-	•	-	-	. 🕶	-	-		-	_		-	-

MOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R = Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) =</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

BH - Borehole

DUP . Duplicate

PASILEY SOLVENTS AND CHEMICALS SITE TABLE 5 . ON-SETE SOIL BORING SAMPLE RESULTS (Continued)

page K

SAMPLE NUMBERS	BH-8A	BH-8B	BH-EB1	BH-EB2	BH-EB3	BH-EB4	BH-EB5	вн-ев6	BH-TB1	BH-TB2	BH-TB3
UNITS	ug/kg	ug/kg	ug/1 .	ug/1	ug/l	ug/l	ug/l	ug/1	ug/l	ug/l	ug/1
MATRIX	Soil	Soil	Vator	Weter	Water	Water	Water	Water	Water	Water	Water
SAMPLE DATE (1989)	10-23	10-23	9-19	9-20	9-21	9-22	9-25	10-23	9-19	9-20	9-21
SAMPLE DEPTH (Ft.)	12-14	22-21	•	•	•	•				, 25	,
VOLATILE ORGANICS						<del></del>		<del></del>		<del></del>	
Hethylene Chloride	-	-	2J	-	-	-	-	-	_	-	
Agetone	-	27 J	6J	•	2 <b>4</b> J	20J	-	R	-	•	-
2-Butenone		R	n	R	•	R	n	R		R	
1,1,1-Trichloroethane	-	70	-	•	-	-	_	-	_	-	-
Trichloroethene		300	-	-	-	-	-	-	-	-	-
4 Hothyl - 2-pentanone	-	-	-	- '	-	-	_	-	-	4	-
Tetrachloroethene	110	180	-	-	-	-	-	-	-	-	-
Toluene	243	310	-	-	-	-	-	-	-	-	-
Ethylbenzene	15	330	-	-	-	-	-	-	-	•	-
Total Tylene	210	1200	-	-	-	-			-		-
Total 1,2-Dichloroethane	-	98	-	-	-	-	_	-	-		-

MOTE: Only those compounds that are detected either as estimated, rejected or positive value in one or more sample are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R = Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) =</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

EB = Equipment Blank

TB = Trip Blank analyzed for volatile organics only

TABLE 5 ON-SITE SOIL BORING SAMPLE RESULTS (Continued) page L

SAMPLE MUMBER	BH-1A	BH-18	BH-2A	BH-28	BH-3A	BH-3B	3B-DUP	3A-DUP	BH-4A	BH-48	BH-54	BH-58	BH-6A	BH-68
Units	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	es/ks	mg/kg
MATRIX	3011	3011	Soil	3011	Soil	3011	Soil	3011	Soil	Sott	3011	3011	Soti	Soll
SAMPLE DATE (1989)	9-19	9-19	9-20	9-20	9-22	9-22	9-22	9-22	9-20	9-21	9-21	9-21	9-25	9-25
SAMPLE DEPTH (Ft.)	12-14	24-26	12-14	24-26	12-14	24-26	24-26	12-14	12-14	24-26	12-14	22-24	12-14	22-24
METALS								<del></del>						
Aluminum	13300	2010	14600	8130	3150	1130	1700	3220	5820	1890	4210	2640	4710	1280
Antimony	<b>-</b> '	•	-	-	-	-	-	-	-	-	-	-	-	-
Arsenio	_	-	-	-	-	-	-	-	<b>-</b> .	-	-	1.68	-	-
Berlum	30.8B	-	29.78	19.30	-	-	-	-	-	-	-	-	-	-
Bery 11um	-	-	-	•	-	•	-	-	<b>-</b> ,	-	-	-	-	-
Cadmium	-	-	-	-	-	-	•	-	-	-	-	-	-	-
Cobalt	6.1B	-	5.28	3.38	3.4B	-	-	3. <i>2</i> 8	8.70	-	-	-	4.4B	-
Calcium	14408	9640	9448	837B	R	R	R	R	8598	R		R	6038	784B
Chrosius	R		29.6		5.8	3.4	5	4.5	R	R	3.7	20.1	7.5	2.23
Copper	31.2	16	25.7	13.6	23.2	20.7	33.9	19.6	16.3	24.0	20.5	27.2	12.3	10.7
Cyanide	-	-	-	-	-	-	- '	3.1	- •	-	•	-	•	-
Iron		R			4520	2090	2170	4660	R	2670	9100	11300	4110	1970
Lead	59J	•	-	12.6J	-	6.9J	6.4J	-	12.2J	13.7J	13.4J	7.5J	n	17.3J
Magnes i un	1800	4188	<b>84</b> 0B	595B	406B	2918	304B	· 409B	512B	27 3B	293B	323B	367B	242B
Hanganése	R	n	R		<i>2</i> 6.9	11.4	7.6	, 30.4	R	1.9	190	23.3	22.1	8.7
Hercury	0.19	-	-	0.48	-	-	1.1	-	-	0.94	-	-	-	0.28
Wickel	12.1J	-	9.8J	•	to.3	6.3J	-	6.1J	7.1J	6.5J	•	-	5. <b>8</b> J	-
Potassiu <b>a</b>	4448	153B	695B	130B	145B	-	2458	191B	-	-	-	207B	-	-
Selentum '	-	-	-	-	-	•	-	=	-	-		-	-	-
Silver	-	-	•	-			-	-	-	3.33	-	~	-	3.2J
Sod Lum	75.6D	62B	10 1B	78Ð	118B	1430	1528	1328	91.9B	1118	95.7B	1480	R	R
Thellium	-	-	-	-	-	-		•		-	-	- •		~
Venedium	24.6	4. 3B	24.4	12.48	6.0B	3.50	4.6B	5.18	7.78	3.5B	3.88	17.2	6.6B	-
Zino	71.8J	R	83.91	61.7J	31J	30.1J	58.1J	31.5J	ħ	49.7J	48.3J	39.6J	39.BJ	A

MOTE: Full Target Compound list metals are listed in this table.

J = Analyte present. Reported value may not be accurate or precise.

R = Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-)</sup> x Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

B = Trace level (less than contract required detection limit: See Appendix F)

DUP . Duplicate

BH = Borehole

TABLE 5 PASILET SOLVENTS AND CHEMICALS SITE ON-SITE SOIL BORING SAMPLING DATA (Continued)

page H

page n										
SAMPLE NUMBER	BH-7Å	BH-78	BH-8A	BH-88	EB-1	EB-2	EB-3	EB-4	EB-5	EB-6
UNITS	mg/kg	mg/kg	mg/kg	mg/kg	ug/l	սց/1	ug/l	ug/l	ug/l	ug/1
MATRIX	Soil	Soi 1	Soll	Soil	Water	Water	Water	Water	Water	Vater
SAMPLE DATE (1989)	9-25	9-25	10-23	10~23	9-19	9~20	9-21	9-22	9-25	10-23
SAMPLE DEPTH (Ft.)	12-14	22-24	12-14	22-24						•
HETALS	····	<del></del>					·····			<del></del>
Aluminum	13700	1610	10800	2200J	-	-	-	-	-	-
ânt isony	-	-	•	-	-	-	-	-	-	-
Arsenlo	1.78	-	-	-	-	-	-	•	-	-
Bertue	34.38	-	278	-	-	-	-	-	•	-
Bery lium	-	-	-	-	-	-	-	-	-	~
Codelus	-	-	-	-	-	-	-	-	-	-
Cobalt	4.58	•	4.50	-	-	-	-	. •	-	-
Calcium	<b>0</b> 758	<b>8</b> 658	8081 f	1 38B	-	1010	15 IB	92.4B	3128	_ =
Chrontun	12.8	•	16.4J	5. IJ	13.0	7.7B	-	-	6.2J	6.83
Copper	25.8	14.2	29.4	16.9	_	-	-	-	-	-
Cyantde	. •	-	•	-	-	10.6		-	-	-
Iron	14000	2120	11400J	34703	12500	24100	62.98	279	218	-
Lead	-	#	22.2J	-	6.2	6.4	_	6.0	11.8	
Hagnes i um	5118	264B	4778	192B	•	•	•	•	138B	52.88
Manganese	46.7	9.2	37.9	32.9	83.7	205	, <b>-</b>	•	-	-
Heroury	-	0.27	0.43J	~	-	-		-	-	-
Nickel	-	-	•	-	-	-		-	-	-
Potessium	-	-	4170	500B	-	-	-	-	-	1230B
Selenium	•	-	~	-	-	-	-	-	-	~
Silver	-	-	-	5.3	~				-	<del>_</del>
Sodium		R	55.08	34.78	-	55.6B	61.68	116B	186B	80.18
Thellius	•	-	<b>-</b> '	-	-	-	-	•	-	-
Venedium	10.2	` •	17.28	5.8J	-	-			-	-
Zino	<b>2</b> 02J	••	43.6J	49.5J	-	23.2	20.1	26.8	-	•

MOTE: Full Target Compound list metals are listed in this table

J = Analyte present. Reported value may not be accurate or precise.

R . Unreliable Result Obtained. Data rejected by validator and is not usable.

<sup>(-) =</sup> Indicates compound was analyzed for but not detected at a level significantly above the level reported in laboratory or field blanks.

B. = Trace level (less than contract required detection limit: See Appendix F)

BH = Borehole

<sup>·</sup> EB = Equipment Slank

### TABLE 6 ON SITE SOIL SAMPLE DATA - TOTAL VOCS GREATER THAN 1 PPM

# PASLEY SOLVENTS AND CHEMICALS SITE HEMPSTEAD, NEW YORK

SURFACE SOIL	TOTAL VOC				TOTAL VOC	
SAMPLE	CONCENTRATION	PRIMARY	BORING	DEPTH	CONCENTRATION	PRIMARY
LOCATION	(PPB)	CONTAMINANTS (")	LOCATION	Fī	(PPB)	CONTAMINANTS (*)
1	1594	trans-1,2-dichloroethene				
2	7953	trans-1,2-dichloroethene				
3	1312	trans-1.2-dichloroethene				
7	1000	trans-1.2-dichloroethene				
		trichloroethane				
		tetrachloroethene				
10	5770	trans-1,2-dichloroethene	3	12	19300	ethylbenzene,xylene
		*	_	22	1200	toluene,ethylbenzene
12	47691	trans-1,2-dichloroethene				
		xylene	,			
16	28153	toluene		<del></del> -	<del></del>	
	35755					
		acetone				
17	7147 55100	1,1.1-trichloroethane		22	26000	tetrachloroethene
19	980					<del></del>
20	33830	trichioroethene			<del></del>	<del> </del>
	2222					
		ethylbenzene				
21	129000	tetrachioroethene				
22	92000	tetrachioroethene			<del></del>	<del></del>
23	90481	tetrachiorcethene	11	22	1800	methylene chloride, xylen
24	79180	tetrachioroethene	8	22	16000	tetrachioroethene
						4-methyl-2-pentanone
25	60500	tetrachioroethene				
28	1550	trans-1,2-dichloroethene				
30	903000	toluene	2	12	15500	tetrachioroethene
31	258000	trichloroethene			•	
		tetrachioroethene				•
34	2300	trichloroethene			<del></del>	
•						
		tetrachioroethene	•	•		
<del></del>		chloroform				
35	56500	tetrachloroethene				
		xylene				
38	23700	tetrachloroethene			<del></del>	
37	96000	tetrachiorosthene				
	<del></del>	xylene			·	
40		tetrachioroethene				
41	231228	tetrachloroethene				
45		trichioroethene			•	
46	18800	chloroform	<del></del>			
50	22800	trans-1,2-dichloroethene			<u> </u>	

<sup>\*</sup> Each primary contaminant accounts for at least 20% of the total VOC concentration

TABLE 7 SUMMARY OF CHEMICALS OF POTENTIAL CONCERN FOR THE PASLEY SOLVENTS AND CHEMICALS SITE

Chemical	On Site Surface Soil	On Site Subsurface Soil	Upper Glacial Aquifer	Upper Magothy Aquifer
Organic Chemicals:				
A south the second				
Acenaphthene	•	ž	•	•
Anthracene .	•	x	x	•
Senzene Chloroform	x		Ŷ	_
Chrysene	•	X X	•	•
Dibenzofuran	-	•	-	•
	_	â	•	X
1,1-Dichloroethane 1,2-Dichloroethane (total)	-	â	•	•
		•	×	x
1,1-Dichloroethene trans-1,2-Dichloroethene	X	X	i	â
Pine how lebeled	•	•	-	•
Di-n-butylphthalate Di-n-octylphthalate		. •	-	•
Ethylbenzene		X	· <b>X</b>	•
bis(2-Ethylhexyl)phthalate	X	Ç	-	-
Fluoranthene	•	Ş	•	•
Fluorene	•	Ç	•	•
4-Methyl-2-pentanone	•	Ş	•	<u>.</u>
Methylene chloride	•	Ç	•	• .
2-Methylnaphthalene	X	Ş	X	•
Naphtha Lene	â	* * * * * * * * * * * * * * * * * * *	â	•
Phenanthrene	:	Ş	:	•
Pyrene	•	Ş	•	•
Tetrachloroethene	×	Ÿ	¥	•
Toluene	x	Ÿ	X X	•
1,1,1-Trichlorgethane	<b>?</b> ,	Ŷ	Ÿ	x
Trichioroethene	x	x		:
Vinyl chloride	x	<b>.</b>	•	•
Xylenes (total)	X	<b>X</b> '	X	•
Inorganic Chemicals:				
Aluminum	•	~	•	x
Antimony	×	Ŷ	•	:
renic	Ŷ	Ŷ	•	•
Sarium	X X	Ÿ	•	•
Servilium	:	X X X X	x	x
Camium	•	Ï	:	-
Chronium	×	•	x	•
Ephait .	•	X .	•	x
Cyanide	•	•	•	×
Leas	×	•	X	X
Kanganese	•	•	:	Ÿ
Nickel	•	•	•	x
Silver	•	×	•	•
Thallium	X	•	•	•
Vanadium	X	×	•	×
Zinc	X	•	•	X

<sup>\* #</sup> Not selected as a chemical of potential concern. X \* Selected as a chemical of potential concern.

TABLE 8 SUMMARY OF CHEMICALS DETECTED IN ON-SITE SURFACE SOIL SAMPLES PASLEY SOLVENTS AND CHEMICALS (a)

(Concentrations reported in mg/kg)

Chemical	Frequency of Detection (b)	Range of Detected Concentrations
Organic Chemicals:		
Acetone	18 / 50	0.026 - 14
Anthracene	2 / 10	0.53 - 1.4
Benzene	4 / 36	0.006 - 0.087
Benzo(a)anthracene	4 / 36 1 / 10 4 / 10	2.6
Senzo(b)fluoranthene	4 / 10	\$.33 · 2.6
Benzo(k)fluoranthene	2 / 10	0.45 - 0.98
Benzo(g,h,i)perylene Benzo(a)pyrene	2 / 10	0.83 - 0.84 0.75 - 1.8
2-Butanone	2 / 10 2 / 10 3 / 10 13 / 34	0.039 - 0.46
Chioropenzene	6/91	0.011 - 0.048
Chloroethane	1 / 50	0.079
Chloroform	14 / 50	0.008 - 17
Chrysene Di-n-butylphthalate	5 / 10	9.28 • 3.4 9.29 • 150
1,2-Dichlorobenzene	3 / 10	0.89 - 2.8
1,1-Dichloroethane	5 / 10 8 / 10 3 / 10 29 / 50 1 / 12	0.006 • 0.58
1.1-Dichic:pethene	1 / 12	0.046
trans-1,2-Dichloroethene trans-1,3-Dichloropropene	33 / 44	0.015 - <b>8</b> 2
trans-1,3-Dichloropropene	3/36	0.005
Ethylbenzene	14 / 50	0.013 · 3 6.8 · 120
bis(2-Ethylhexyl)phthalate Fluoranthene	2 / 6 6 / 10	0.36 - 5.9
Indeno(1,2,3-c,d)pyrene	1 / 10	0.88
Methylene Chloride	22 / 40	0.012 - 0.74
2-Methylnaphthalene	8 / 10 6 / 10	0.46 - 10
Naphthalene	6 / 10	0.38 - 43
Phenanthrene	8 / 10 8 / 10 43 / 48	0.37 - 3.1
Pyrene - Tetrachloroethene	6 / 10	0.57 • 4.6 0.01 • 270
Toluene	26 / 47	0.008 - 470
1,1,1-Trichloroethane	17 / 48 41 / 49	0.0098 - 6.9
Trichloroethene	41 / 49	0.004 - 120
Vinyl Chloride	9 / 49	0.068 - 0.67
Xylenes	21 / 49	0.01 - 35
norganic Chemicals:		
Aluminum	10 / 10	2,670 - 9,630
Antimony	2 / 10	16.7 • 48
Arsenic	8 / 8	2.8 - 17.1
Berium	10 / 10 10 / 10	29.5 - 430
Calcium	10 / 10	5,530 - 44,300
Chromium Cobelt	0 / 10	9.9 - 58.1 2.9 - 7.6
Copper	10 / 10	15.9 - 74.8
Cyanide	/ / 40	0.3 - 4.4
Iron	10 / 10 10 / 10 10 / 10	11,100 - 41,800 130 - 1,230
Lead	10 / 10	130 - 1,230
Kagnesium	70 / 10	2,820 - 22,100
Kanganese	10 / 10 1 / 10 9 / 10	73.6 - 270
Mercury Nickel	0 / 10	0.1 7.7 - 25.1
Potessium	10 / 10	350 - 980
Thellium	5/0	2 - 3.3
Vanadium	10 / 10	2 - 3.3 10.1 - 37.7
Zinc	10 / 10	130 - 710

 <sup>(</sup>a) Samples 1-10 (composite samples) and TAB-SDAB.
 (b) The number of samples in which the chemical was detected divided by the total number of samples analyzed for that chemical.

<sup>\* =</sup> Selected as chemical of potential concern.

### TABLE 8 (continued)

## SLAMARY. OF CHEMICALS DETECTED IN UPPER GLACIAL AGUIFER GROUNDWATER PASLEY SOLVENTS AND CHEMICALS (8)

(Concentrations reported in ug/L)

	Range of Detected Concentrations									
Chemical	Upgradient (b)	On Site (c)	Downgradient (d)							
Organic Chemicals:										
• • • • • • • • • • • • • • • • • • •		40 54 5								
* Benzene	2.75 - 25.5	12 - 31.5	8.5							
Acenaphthene	10	1 - 6.5	<b>8D</b>							
Acenaphthylene	<b>10</b>	10 - 18.5	MD							
Chlorobenzene	AID.	. 20	<b>#</b> D							
• Chloroform	<b>ED</b>	53.5	25							
Dibenzofuran	. 🛍	2 • 5	<b>ID</b>							
1,1-Dichloroethane	<b>MD</b>	465	<b>23.</b> 5							
• 1,1-Dichloroethene	_ <b>MD</b>	_ 73	MD .							
<ul><li>trans-1,2-Dichloroethene</li></ul>	2.25	2.75 - 31,000	8 - 102.5							
<ul> <li>Ethylbenzene</li> </ul>	<b>MD</b>	425	30							
fluorene	MD	3 • 6.5	· MD							
<ul> <li>2-Methylnaphthalene</li> </ul>	10	<b>36 - 103.5</b>	12.5							
* Naphthalene	<b>10</b>	<b>22</b> 5 "	41							
di-n-Octylphthalate	2	<b>***</b>	<b>90</b>							
Phenanthrene	<b>8</b> TD	2.5 - 5	<b>AD</b>							
* Tetrachioroethene	25.5	1.5 - 125.5	3 - 31							
* Toluene	ND	1.5 - 1,100	2							
* 1,1,1-Trichloroethane	. MD	3,200	5.25 - 285							
* Trichioroethene	11.5	1 - 230	15 - 205							
• Xylenes		11.5 - 1,600	34.5							
Inorganic Chemicals:	. •	·								
**************************************		4/4 52 /00	0/4 . 02 /00							
Aluninum	4,030 - 42,100	141 - 23,400	<b>8</b> 61 - <b>97,40</b> 0							
Barium	68.8 · 111	50.2 - 112	38.6 - 372							
• Beryllium	1.7	0.73	6.6							
Colcium	18,100 - 36,000	13,900 - 23,600	19,500 - 22,300							
• Chromium	23.1 - 43.8	15.6 - 27.7	63.6 - 258							
Cobalt	10 <u>.9</u>	16	40.8 - 194							
Cyanide	70	15	ND							
Iron	4,690 - 28,900	2,530 <b>- 9</b> 9,100	3,410 - 152,000							
• Lead	ND	8.2 - 15.3	9.5 - 34.6							
Manganese	1,060 - 12,200	1,360 - 16,000	4,220 - 5,630							
Nickel	53.8 - 129	48.3	100 - 207							
Silver	ND	5.6	SED.							
Sodium	4,280 - 35,300	35,400 - 390,000	17,000 - 28,700							
Vanadium	34.7	27.6	94.8							
2ins	1,380 - 3,200	<b>59.1 - 8</b> 59.	192 - 1,070							

 <sup>(</sup>a) The reported range represents concentrations found at shallow and intermediate depths within the well.
 (b) Well 1.
 (c) Well 2.
 (d) Well 4.

ND = Not detected.

<sup>\* =</sup> Selected as chemical of potential concern.

## TABLE 8 (continued)

## SUPPLIES OF CHEMICALS DETECTED IN ON-SITE SUBSURFACE SOIL PASLEY SOLVENTS AND CHEMICALS (a)

(Concentrations reported in mg/kg)

Chemical	Frequency of Detection (b)	Range of Detected Concentrations
Organic Chemicals:		
Acenaphthene		0.31
Anthracene	1/82/8	8.16 - 0.17
Chloroform	177	0.0058
Chrysene	3 / B	0.019 - 0.12
Dibenzofuran	3/8	0.32 - 0.42
1,1-Dichloroethane	1/7	0.0073
1,2-Dichloroethane	1/1	0.05
trans-1,2-Dichloroethene	1/7	0.05
Di-n-octylphthalate	2 / 8	0.13 - 0.14
P Ethylbenzene P his/7-Fthulherullmhthalete	5 / 8 2 / 5	0.18 - 1.1 0.71 - 4.9
<pre>bis(2-Ethylhexyl)phthalate Fluoranthene</pre>	5/8	0.045 - 0.16
* Fluorene	4/8	0.18 - 0.62
4-Methyl-2-pentanone	1/8	1.8
Methylene Chloride	5 / 8	0.0098 - 0.60
2-Methylmaphthalene	6/8	0.48 - 7.6
Naphthalene	5/8	0.25 - 4.2
Phenanthrene	7/8	D.12 - 1.5
Pyrene	6 / 8	0.049 - 0.21
Tetrachloroethene	7/8	0.12 - 11 0.0048 - 2.9
'Toluene '1,1,1-Trichloroethane	2 / 8	0.036 - 0.16
Trichloroethene	4/8	0.042 - 1.4
Xylenes (total)	7/8	0.0073 - 4.8
norganic Chemicals:		
Aluminum	8 / 8	2,300 - 11,400
Antimony	1 / 8	4.5
Arsenic	2 / 8	1.1 - 1.2
' Barium	3/8	22.2 - 110
'Beryllium	1/8	0.4
Cegnium	1 / 8	0.4
Cobalt	6 / 8	2.9 - 5.6
Copper	8 / 8 8 / 8	11.5 - 24.4
Magnesium Mercury	7 / 8	304 - 1,110 0.1 - 0.5
Nickel	. 5/8	3.9 - 7.1
Potassium	5/8	210 - 660
Silver	3/8	1.9 - 2.9
Sodium	6/6	44.9 - 136
Venedium	8/8	4.6 - 18.4

 <sup>(</sup>a) Samples 8H-1 - 8H-8.
 (b) The number of samples in which the chemical was detected divided by the total number of samples analyzed for that chemical.

<sup>\* \*</sup> Selected as chemical of potential concern.

POTENTIAL EXPOSURE PATHWAYS FOR THE PASLEY SOLVENTS AND CHEMICALS SITE CURRENT LAND USE CONDITIONS

Exposure Medium	Source and Mechanism of Release	Potential Receptor	Exposure Route	Potential for Significant Exposure	Method of Evaluation
Surface Soll	Post spills and direct discharge from on site tanks to surface soil	Тгезраметв	Incidental Ingestion, dermal absorption	None. Site soil has been covered by gravel and access is restricted. No potential for direct contact or incidental ingestion exists.	None. Incomplete pathway.
Subsurface Soll	Direct discharge from on site tasks or leaching from surface soils	Trespessors	Incidental Ingestion, dermal absorption	None. Persons do not come into contact with subsurface soil.	None. Incomplete pathway.
Groundwater	Leaching to groundwater from soil; Migration to public water supply wells	Résidents	Ingestion, inhalstion, and dermal absorption of chemicals during home use	None. Site related contamination has not migrated to public supply wells, and no private residential wells exist near or downgradient of the site.	None. Incomplete pathway.
Air (vapors)	Volatilization of chemicals from soli	Тгезрамета	Inhalation	None. Although volatilization of chemicals may occur, access to the site is restricted.	None. Incomplete pathway.
Air (vapora)	Volatilization of chemicals from soil to sir	Nearby residents	Inhatation	Low. Dispersion in ambient sir significantly decreases concentrations of chemicals.	Quantitative. Emission estimates will be based on measured surface soil concentrations.
Air (dust)	Dust released from surface soil to air	Trespossers/ Nearby residents	Inhalation	Negligible. Dust generation is unlikely because site soil has been covered by gravel. Further, site access is restricted and trespassers are not expected.	None. Incomplete pathway.

### TABLE 9 (continued)

## POTENTIAL EXPOSURE PATHWAYS FOR THE PASLEY SOLVENTS AND CHEMICALS SITE FUTURE LAND USE CONDITIONS

Exposure Medium	Source and Mechanism of Release	Potential Receptor	Exposure Route	Potential for Significant Exposure	Method of Evaluation
Surface Soll	Past spills and direct discharge from on site tanks to surface soil	Workers	Incidental ingestion, definal absorption	Low to high depending on nature of activities at the site and the degree of vegetation or pavement at the site.	Quantitative. Estimates will be based on current surface soll concentrations.
Subsurface Soll	Direct discharge from on alte tanks or leaching from surface soils	Workers	incidental ingestion, dermal absorption	Low. Contact with subsurface soils is likely to be infrequent and of short duration.	Nanc.
Orcundwater	Leaching to groundwater from soil; Pumping from an on site well	V forkers	Ingestion of drinking water	Moderate to high if use occurs.	Quantitative.
Oroundwater	Migration to residential well adjacent to site	Nearly Resident	Ingestion and inhalation and dermal absorption of chemicals during home use	Moderate to high for ingestion and inhalation if we occurs. Negligible for dermal absorption compared to ingestion and inhalation.	Quantitative. Estimates will be based on measured groundwater concentrations.
Air (vapors)	Volatilization of chemicals from soil to air.	Workers	Inhalation	Moderate. Volatilization of chemicals from on site soil will occur.	Quantitative, Emission estimates will be hased on measured current soil concentrations.

TABLE 10

ORAL CRITICAL TOXICITY VALUES FOR CHEMICALS OF POTENTIAL CONCERN
PASLEY SOLVENTS AND CHEMICALS

Chemical	Chronic RfD (mg/kg-day)	Uncertainty factor (a)	Target Organ (b)	RfD Source	Stope factor (SF) (mg/kg-day)-1	EPA Weight of Evidence Classification (c)	SF Source
Organics;				· ,			
Acenaphthena	6.00E-02	3,000	Liver	IRIS	•••	•••	•••
Anthrocene	3.00E-01	3,000	None observed	IRIS	•••	. <b>D</b>	IRIS
lenzene .	•••	••••		IRIS	2.90E-02	A	. IRIS
chtoroform	1.00E-02	1,000	Liver	IRIS	6.10E-03	, 02	IRIS
thrysene	•••	•••	***	•••	•••	B2	IRIS
l benzofuran		•••	***	MEAST	•••	D	IRIS
l, 1-Dichloroeth <b>ene</b>	1.006-01	1,000	Kidney	REAST	•••	С	IRIS
, 2-Dichloroethana	•••	•••	***	1815	9.10E-02	92	IRIS
, 1-Dichloroethene	9.00E-03	1,000	Liver	IRIS	6.00E-01	Ç	1815
:ls-1,2-Dichloroethene	•••	•••		'	•••	Ď	IRIS
rans-1,2-Dichloroethane	2.00E-02	1,000	Liver	IRIS	•••	•-•	•••
i-n-butylphthalate	1.00E-01	1,000	Mortal fty	IRIS	•••	•••	•••
i-n-octylphthalate	2.00E-02	1,000	Liver/Kidney	MEAST	•••	•••	•••
thyl Benzene	1.006-01	1,000	Liver/Kidney	IRIS	•••	D	IRIS
lis(2-ethylhexyl)phthalate	2.006-02	1,000	Liver	IRIS	1.40E-02	<b>0</b> 2	IRIS
luoranthene	4.00E-02	3,000	Kidney/Liver	IRIS	•••	•••	•••
luorene	4.00E-02	3,000	Hemotology	IRIS	•••	D	IRIS
lethylene Chloride	6.00€∙02	100	Liver	IRIS	7.50E-03	92	IRIS
2-Hethylnophthalene			•-•	• • •	***	,	
laph tha lene	4.00E-03	10,000	<pre><body pre="" weight<=""></body></pre>	MEAST	•••	Ð	IRIS
henonthrene	• • •	***	•••	HEAST		D	1215
'yrene	3.00E-02	3,000	Kidney	IRIS		D	IRIS
etrachloroethene	1.00E-02	1,000	Liver	IRIS	5.10E-02	92	MEAST
oluene	2.00€-01	1,000	Liver/Kidney	IRIS	•••	D	IRIS
, 1, 1-Trichloroethane	9.00€-02	1,000	Liver	IRES		D	1813
richloroethene	7.35E-03	1,000	tiver	HA	1.106-02	<b>D</b> 2	NEAST
/inyl Chloride	•••	****			1.90E+00	A	HEAST
lylenes (total)	2.00E+00	100	CNS, Mortality	IRIS		D	IRIS

### TABLE 10 (continued)

### IMMALATION CRITICAL TOXICITY VALUES FOR CHEMICALS OF POTENTIAL CONCERN PASLEY SOLVENIS AND CHEMICALS

Chemical	Chronic RfD (mg/kg-dny).		Target Organ (b)	RfD Source	Unit Risk (UR) (ug/m3)-1	EPA Weight of Evidence Classification (c)	UR Source
Organics:							
Benzene	•••	•••	•••	IRIS	8.30E-06	<b>A</b> .	1819
Ehloroform	•••	•••	•••	IRIS	2.30£ ·05	92	IRIS
1.1-Dichtgroethane	1.00E-01	1,000	Kidney	MEASI	2.302 07	Ĉ	1813
1, 1-Dichtor bethene		1,000	NIGHT.	1815	5.00E-05	. č	1815
trans · 1 · 2 · Dichloroethene	•••	•••	•••	IRIS	J.002-03	•••	1413
Di-n-butylphthalate	•••	•••	•••	1815		4-4	•••
Ethyl Benzene	£1.00£+001 *	100				0	
	(1.005*00) -		Devel apment	IRIS	•••		IRIS
bis(2-Ethylhenyl)phthalate		•••	•••	IRIS		. 05	IRIS
letrachtoroethene	40 400 400 4	***	***	IRIS	5.20E-07	03	MEAST
loluene	12.006+001 *	100	- CHS, Irritation	MEAST	•••	P	IRIS
1,1,1-trichloroethane	3.00E-01	1,000	Liver	HEAST		_0	1813
Irichloroethene	•••	•••	•••	IRIS	1.70E-06	92	MEAST
Vinyl Chloride	•••	•••	_		8.40E · 05	A	MEAST
Kylenes (total)	(3.006-01) •	100	CMS, respiratory	MEAST	•••	0	IRIS
Inorganica:						•	
Atunima		•••	•••	MEAST	•••	•••	
Ant Improv		•••	Concer	IRIS	•••	• •••	•••
Arsenic			Concer	1815	4.00E-03 (d)	· A	1812
Par lum	15.00E-041 *	1.000	Fetotoxicity	MEAST	••••	,	••••
Deryllum	***	•,•••		IRIS	2.40E-03	92	lats
Chromium III and Compounds	[2.00E-06] *	100	lesel Mucose	WEAST	2.402-03	•••	***
Chromium VI and Compounds	(2.00E-06) *	300	lesel Mucose	WEAST	1.206-02	<b>A</b>	IRIS
Cobelt	(6.000 00)		, 10301 1100000	***	***************************************	***	••••
Evenide	•••	•••		2181	•••	4.0	•••
lead	•••	•••	CWS	1815	•••	12	1012
lenganese	(4.00E-04) *	900		MEASI	•••		
lickel refinery dust	[4,002,04]	700	CHS, Respiratory		2.40E-04	•	IRIS
		=	***	IRIS		Ą	IRIS
tickei subsulfide Lilver	•••	•••	•••	1815	4.80E-04 (e)		IRIS
	•••	• • •	•••	1015	•••	•••	
thattium (In soluble salts)	•••	•••	•••	MEAST	•••	•••	•••
Venedium	•••	•••	•••	MEAST	• • •	•••	• •••
linc and compounds	•••	•••	•••	MEAST	•••	•	tris

<sup>(</sup>a) Uncertainty factors are a measure of the uncertainty in the data available. A higher uncertainty factor represents a greater amount of uncertainty in the date.

(d) An absorption factor of 30% was used to calculate the unit risk from the stope factor.

### . Integrated fisk information System.

MEAST - Health Effects Assessment Summery Tables.

<sup>(</sup>b) A target organ is the organ most sensitive to a chemical's toxic effect. Afts are based on toxic affects in the target organ, If an RID was based on a study in which a target organ was not identified, an organ or system known to be affected by the chemical in listed.

<sup>(</sup>c) EPA Weight of Evidence for Carcinogenic Effects:

<sup>[</sup>A] = Human carcinogen based on adequate evidence from human atudies;
[B2] = Probable human carcinogen based on inadequate evidence from human atudies and adequate evidence from atudies;
[C] = Possible human carcinogen based on ilmited evidence from animal atudies in the absence of human atudies;

<sup>(</sup>D) - Not classified as to human carcinogenicity; and

<sup>(</sup>e) The conser unit risk for nicket subsuifide was conservatively used to calculate the risks associated with Inhelation of nicket.

<sup>.</sup> No information evaluate.

<sup>&</sup>quot; Value is a unit risk in mo/ml.

TABLE 11 SUMMARY OF POTENTIAL HEALTH RISKS ASSOCIATED WITH THE PASLEY SOLVENTS AND CHEMICALS SITE

Exposure Pathway	Upper Bound Excess Lifetime Cancer Risk <sup>a</sup>	Hazard Index for Noncarcinogenic Effects <sup>6</sup>
CURRENT LAND USE:	•	
Inhalation 0-30 Year Old Residents Adult Residents  FUTURE LAND USE:	18x10 <sup>-7</sup> 8x10 <sup>-7</sup>	<b>&lt;1</b> <b>&lt;1</b>
Soil Ingestion Workers	2d0 <sup>4</sup>	<b>&lt;</b> 1
Dermal Absorption from Soil Matrix Workers	- <b>2</b> 10 <sup>6</sup>	- <b>-≪1</b> -
Inhalation Workers	7x10 <sup>-5</sup>	<1
Ingestion of Upgradient Upper Glacial Groundwater Workers 0-30 Year Old Residents Adult Residents	3x10 <sup>-5</sup> 2x10 <sup>-4</sup> 1x10 <sup>-4</sup>	<1 <1 <1
Ingestion of On Site Upper Glacial Groundwater Workers 0-30 Year Old Residents Adult Residents	2x10 <sup>-4</sup> 9x10 <sup>-4</sup> 7x10 <sup>-4</sup>	>1 >1 >1
Ingestion of Downgradient Upper Glacial Aquifer Groundwater 0-30 Year Old Residents Adult Residents	5x10 <sup>-4</sup> 4x10 <sup>-4</sup>	>1 >1
Ingestion of Upgradient Upper Magothy Aquifer Groundwater Workers		: <b>&lt;1</b>
0-30 Year Old Residents Adult Residents	-	<1 <1

### TABLE 11 (Continued)

# SUMMARY OF POTENTIAL HEALTH RISKS ASSOCIATED WITH THE PASLEY SOLVENTS AND CHEMICALS SITE

Exposure Pathway	Upper Bound Excess Lifetime Cancer Risk <sup>a</sup>	Hazard Index for Noncarcinogenic Effects <sup>5</sup>
FUTURE LAND USE (CORL.):		
Ingestion of On Site Upper Magothy Aquifer Groundwater Workers 0-30 Year Old Residents Adult Residents	3x10 <sup>-5</sup> 1x10 <sup>-4</sup> 1x10 <sup>-4</sup>	<1 1 <1
Ingestion of Downgradient Upper Magothy Aquifer Groundwater 0-30 Year Old Residents Adult Residents	2x10 <sup>-5</sup> 3x10 <sup>-5</sup>	<1 <1
Inhalation While Showering with Upgradient Upper Glacial Groundwater Adult Residents	7x10 <sup>-6</sup>	-
Inhalation While Showering with On Site Upper Glacia! Groundwater Adult Residents	2x10 <sup>-4</sup>	<1
Inhalation While Showering with Downgradient Upper Glacial Groundwater Adult Residents	3x10 <sup>-5</sup>	<1
Inhalation While Showering with Upgradient Upper Magothy Groundwater Adult Residents	NE	NE
Inhalation While Showering with On Site Upper Magothy Groundwater Adult Residents	4x10 <sup>-6</sup>	<1
Inhalation While Showering with Downgradient Upper Magothy Groundwater Adult Residents	4x10 <sup>-6</sup>	

The upperbound individual excess lifetime cancer risk represents the additional probability that an individual may develop cancer over a 70-year lifetime as a result of exposure conditions evaluated.

NE = Not evaluated. Pathway only evaluated for chemicals of concern which volatilize.

The hazard index indicates whether or not exposure to mixtures of noncarcinogenic chemicals may result in adverse health effects. A hazard index less than one indicates that adverse human health effects are unlikely to occur.

<sup>--- =</sup> Not applicable. Chemicals of potential concern for this pathway do not exhibit carcinogenic (or noncarcinogenic) effects.

TABLE 12 POTENTIAL ARARS FOR GROUNDWATER CONTAMINANTS
PASLEY SOLVENTS AND CHEMICAL SITE

					ARARS				GOAL TO	BE CONSIDERED	)	·	
	BAANIBANA.	1	MOST			101		PROPOSED	M AMBERT	EPA DRIBOUNG		REFERENCE	
	CONCENTRATION	į	STRINGENT	FEDERAL	NV AMBENT	CROLINDWATER	rederal.	FEDERAL	WATER	WATER		CONCENTRATION	
	DETECTED	MOST	COAL	COMA	WATER	CHARLEA	SDWA.	SDWA	CHALFFY	HEALTH	EPA AWOC	FOR POTEMBAL	
	MON-SITE	STRUCCENT	TORE	MCL	<b>CILMINITY</b>	STANDARDS	MOLO	MCL	GLIDANCE	ADVISORES	DW ONLY	CARCINOCENS	
	WELLS 23 & 28	WELLS 23 & 28	ARAR	CONSIDERED	(+)	STANDANDS (M)	(c)			WILES (H)	₩	•	•
VOLATLE ORGANICS COMPOUNDS	ugA .	ugh	144	فيب	Nat		ugA	ugA	ايون	ugh .	legel .	اليب	
Methylene Chicride	161		•	NS	564	NS .	OP	5	MS	NS	0(0.19)	4.7	
Bentene	431	MD		5	ND()#4	ND	0	NS	<b>CM</b>	NS	0(0.67)	1.2	
Acetone	30001	50	NS	NS	NS	NS	NS	NS	50G	NS	MS ,	MS	
Chlarotarm	743	100	0	NS	100()(1)	100	NS	NS	NS	NS	0(0.19)	6.7	
1,1 - Dichlaraethene	943	5	0	1	564	NS	7	NS	NS	· NS	0(0.33)	.06	
1,1 - Dichlaroethene	630	5	NS	NS	504	. NS	NS	N9	NS	NS	NS	NS	
Trans - 1,2 - Dichloroethene	37,000		100	100	5(1)	N9	100	NS	N9	350	NS	NS	
Ethythenzene	510	5	700	700	5(4)	NS	700	NS	NS	3,400	2400	NS	
Teb achteroethene	160.1	5	0	5	5#4	NS	0	NS	NS	NS	0(0 <b>88)</b>	.7	
Toksene	1100		1000	1000	504	, en	1000	N3	NS	10,000	15,000	NO	
Trichtgroethene	320	5	•	5	564	10	0	NS	NS	N9	0(2.4)	32	
1,1,1 - Trichtoroethene	3000	5	200	200	5(14)	NS ·	200	NS	NS	1.000	19,000	NS	
Chlorobenzene	510	6	100	100	5(14)	NS .	100	PN PN	NS	3.150	400	HS	
Kylene (Total)	817.3		2,200	10,000	50(4)	NS	10,000	NS	MS	2,200	NS	MS	
SEMI-VOLATLE GROANC COMPOUNDS													
# - n - buly! phtralete	40	50	44,000	NS .	NS	N9	NS	NS	50G(1)	N9	44,000	NS	
2 - Methylnephthelene	110	50	NS	N3	NS	NS	NS	en	50G	HG	MO	. NS	
Vaphthelene	270	10	NS	NS	NS	N9	NS	NS	10G <b>(1)</b>	NS	· NS	NS	
Obenzokran	5.1	50	MS	NS	NS	NS	NS	N9	50G	N3	NB	NS	
herantivane	เม	50	NS	N3	NS	NS	NS	NS	50G <b>(4)</b>	NS	NS	NS	
i -n-Octyl phtheiste	S1	50	NS	N3	NS	, NS	NS	NS	20GM	N3	NS	N9	
comphiliylano	21	50	NS	N9	NS	ен	NS	N9	50G	NS	N9	NS	
cemphihene	73 .	20	20	N3	NS	NS	NS	NS	30GM	N3	20	NS	
Lorene	73	50	NS	NS	NS	N9	NS	RM	50G#)	NS	NS	NS	
is (2 - othythosyl)phthalato	40	50	25	NS	NS	N9	NS	NS	503	NS	NS	2.5	

PG. 1 OF 3

## TABLE 12 Cont'd POTENTIAL ARARS: WOUNDWATER CONTAMINANTS PASLEY SOLVENTS AND CHEMICAL SITE

		·			ARARS		[	·	GOAL TO	BE CONSIDERED	)	
	MAXIMA		MOST			W		PROPOSED	W MBEN	EPA DROVING		REFERENCE
•	CONCENTRATION	1	STRACENT	PEDERAL	NY AMBENI	GROLINDWATER	/EDERAL	FEDERAL	WATER	WATER		CONCENTRATION
•	DETECTED	MOST	QOAL	SDWA	WATER	CITACULA (1)	<b>SDWA</b>	SDWA	CHAPILA	HEALTH	EPA AWQC	POR POTEMBAL
	MON-SITE	STRACENT	10 <b>9</b> 6	MOL	CHARLITY	STANDARDS	MOLG	MOL	GUIDANCE	ADMISORES	DWONLY	CARCINIDIENS
	WELLS 29 4 28	AAAA	CONSIDERED		STANDARDS (+)	<b>(c)</b>	14	14	VALLES A		•	<b></b>
METALS	Agu	ugA	Los	ug/l	ugA	ug/l	ugil	ugA	Agu	ugil '	lg/	ug/l
Akardram	97,400	100	50	NS	100@	NS	50 - 200(A)	NS	МЭ	NS	NS	NS.
Artimony	39.9	3	3	NS	3	NS	36	10/5(m)	N3	NS	146	NS
Areenic	-	25	20	50	25	25	50P	NS	NS	60	(25 ng/l)	20
Backen .	372	1,000	· 1,800	1,000	1,000	1,000	5,000P	NS	NS	1,000	NS	N9
Berytium	••	•		NS	. 30	NS	OP	•	N9	NS	(3.9 ng/)	.000
Cochrision	4.5	5	5	5	10	10	5	NS	· N3	10	10	NS
Calchen	36,000	NS.	NS	NS	NS.	N9	NS	NS	NS	NS	NS	NS
Chromhan	255	50	50	100	50	50	100	NS	N9	170	60	NS
Cobalt .	45.1	5	NS	NS	5 <b>(4)</b>	· NS	NS	NS	NS,	NS	NS	NS
Соррег	270	200	1,000	PM PM	200	1,000	1300P	1300	. NS	NS	1000	NS
Cyanide	70	100	200	N9	100	200	200P	200	en <sup>°</sup>	750	200	NS
ton	152,000	300	NS	en	300 (n)	300(n)	300(4)	NS	NS	N3	MS .	N3
Lead	34.6	15	0	15	25	25	: OP	NS	NS	20 up/day	60	N9
Magneeken	6330	35,000	NS	NS	<b>35,000 (G)</b>	NS	NS	MS	NS	NS	NS	N9
Manganese	16,100	50	NS	NS	300(4)	200(4	50 <b>(</b> )	MS	NS	MS	NS	, NS
Macay	<u> </u>	2	2	2	2	2	2	NS	NS	5.5	10	NS
Mickel	310	100	15.4	NS	(N)A	NS	100P	100	NS	360	15.4	NS
Potestan	10,200	NG	NS	149	N3 .	NS	MS	NS	NS	NS	NS	N3
Selerium	-	10	10	50	10	50	50	N9	NS	NS	10	NS
Béver .	560	50	. 50	NS	50	50	1006)	NS.	N3	NS	50	N9
Sadken	1000,000	20,000	NS ·	NS	20,000	N9	N9	NS.	HS	NS	NS	N9
Theilium	5.7	•	17.0	NS	40	NS	NS	2/1(m)	NS	NS	17.0	N9
Versethen	94.8	14	MS	-MS	140	MS	PM SM	NS	N3	NO	· NS	NS
Zine	3,200	300	5,000	N9	300	500	5,000(A)	NS	NS	NS	5000	NS

PG 2 OF 3

#### NOTES:

- J ANALYTE PRESENT. REPORTED VALUES MAY NOT BE ACCURATE OR PRECISE.
- P PROPOSED VALUE
- NS NO STANDARD OR GUDGLINE EXISTS
- Q GUIDANCE VALUES
- NO NOT DETECTABLE
- (a) SAFE DRINKING WATER ACT MAXIMUM CONTAMINANT LEVEL
- (b) 6 MYCRR PARTS 701 AND 702 AND 10 MYCRR PARTS 170 AND 5 AS SLIMMARZED IN MYSDEC DIVISION OF WATER TECHNICAL AND OPERATIONALS GLIDANCE SERIES
  - (1.1.1) GEPT. 25, 1990.
- (c) 6 MYCRR PART 703
- (d) SAFE DRINKING WATER ACT MAXIMUM CONTAMINANT LEVEL GOALS
- (o) EPA DRINKING WATER HEALTH ADVISORIES, SUPERFLIND PUBLIC HEALTH EVALUATION MANUAL, 1986
- IN EPA AMBIENT WATER CLIALITY CRITERIA FOR PROTECTION OF HUMAN HEALTH ADJUSTED FOR DRINKING WATER ONLY (CONCENTRATIONS IN PARENTHESES CORRESPOND TO MIDPOINT OF RISK FRANCE FOR POTENTIAL CARCINGGINS ONLY)
- (a) CORRESPONDS TO AN INCREASED LIFETIME CANCER RISK OF 1E 8, CALCULATED FROM SLOPE FACTORS PUBLISHED IN THE HEALTH EFFECTS ASSESSMENT SUMMARY
  TABLES (1991) AS FOLLOWS: REFERENCE CONCENTRATION (1E-8 X 70 KG/(SLOPE FACTOR IN BIGHG/DAY) X 2L/DAY)
- IN TOTAL ORGANIC CHEMICALS CANNOT EXCEED 100 UGA.
- **(A PROPOSED FOR REVISION**
- (A APPLIES TO EACH ISOMER INDIVIDUALLY
- (A) SECONDARY MOL
- (I) NO HUMAN HEALTH STANDARDS. THIS STANDARD IS FOR PROTECTION OF AQUATIC LIFE.
- IN) TWO OPTIONS PROPOSED BY EPA RESILTING IN DIFFERENT STANDARDS.
- IN IF IRON & MANGANESE ARE PRESENT, THE TOTAL CONCENTRATION OF BOTH SHOULD NOT EXCEED 500 MGA.
- (1) FORMALA TO DETERMINE STANDARD EXPID. 76(In (PPM HAPDNESS)) + 1.08

Recommended soil cleanup objectives (mg/kg or ppm)
Pesley Solvents Site, 8 1-30-016

Contaminant	Solubility	Partition -	Grounduater	Allouable	b MM Soil Cleanup	USEPA Heal (ppn)			. NAN
	ng/l or ppn S	caefficient Kac	Standards/ Criteria Cu ug/l or ppb.	Soil conc. ppn. Cs	objectives to Protect GH Quality (ppm)	Carcinogens	Systemic Tenicants	CRQL (ppb)	Roc.soil Cloup Objet. (ppn)
Kylenes	198	210	S	0.012	1.2	H/A	200,000		1.2
Ethylbenzono	152	1,100	. 5	0.055	5.5	H/A	6,000	5	5.5
Toluene	\$35	300	5	0.015	1.5	H/A	2,000	5	1.5
Yelrechloroethene ·	150	277	5	0.014	- 1.4	84	800	5	1.5
Tri chilaraethene	1,100	126	5	0.007	0.70	61	N/A	5	1.0
1,1,1-frichloroethene	1,500	152	5	0.0076	0.76	MA	7,000	5	1.6
1,2-0i chl croelhone(trans)	6,300	59	5 '	0.003	0.3	N/A	N/A	5	0.5
Chloroforn	0,200	31 <sup>-</sup>	*	0.002	0.2	114	800	5	0.2
1,2-Dichlerobenzene	100	1,700	4.7	0.079	7.9	MA	NA	330	9.0
Phenanthrene	1.0	4,365	50	2.20	220.0	N/A	M/A	330	50.0
Fluoranthene	0.206	38,000	50	19	1900.0	N/A	3,000	330	50.0
Haphthal one	31.70	1,300	10	0.130	13.0	N/B	300	230	13.0
2-nethyl nephthal ene	26.00	727	50	0.365	36.0	K/A .	N/R	330	36.0
Di-n-butyl phthelete	<b>1</b> 00	162	50	0-08	8.0	H/A	4,000	330	0.0

MIN is Nothed Detection Limit

MAN As per proposed FAGM, Total VOCs & 10 ppm., Total Seni-VOCs & 500 ppm. and Individual Seni-VOCs & 50 ppm.

Note: Soil cloup objectives are developed for soil organic carbon content (f) of 12, and should be adjusted for the actual soil organic carbon content if it is known.

a. Allowable Soil Concentration Cs = f x Cu x Koc

b. Soil cleanup objective = Cs x Correction Factor (CF)

M Partition coefficient is calculated by using the following equation: log Koc  $\approx$  -0.55 log S + 3.64. Other values are experimental values.

NX Correction Factor (CF) of 100 is used as per proposed FRGH

**APPENDIX 3** 

# New York State Department of Environmental Conservation io Wolf Road, Albany, New York 12233 - 7010



MAR 1 8 1992

Ms. Carole Petersen
Chief
NY/Caribbean Superfund Branch II
U.S. Environmental Protection Agency
Region II
26 Federal Plaza
New York, NY 10278

Dear Ms. Petersen:

Re: Pasley Solvents & Chemicals Site ID No. 130016
Draft Record of Decision

The New York State Department of Environmental Conservation (NYSDEC) has reviewed the March 13, 1992 draft Record of Decision (ROD) for the Pasley Solvents and Chemicals site, as telexed to us on March 16, 1992.

The remedy presented in the draft ROD includes treating contaminated soil via soil vacuuming followed by soil flushing, if necessary, and treating groundwater via metals precipitation/air stripping with vapor phase granular activated carbon/GAS polishing.

As per conversations between our respective staff, this March 13 draft reflects the several changes made to the March 5, 1992 draft. Consequently, the NYSDEC concurs with the draft ROD for the Pasley Solvents and Chemicals site.

Sincerely,

Edward O. Sullivan Deputy Commissioner

cc: M. Hauptmann, USEPA-Region II

S. Henry, USEPA-Region II

#3/25/92

### Index Document Number Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 1

Document Number: PAI-881-8881 To 8112 Date: 88/38/88

Title: Final Field Operations Plan for Remedial Investigation/Feasibility Study - Pasley Solvents

and Chemicals Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Blanar, Edward W: ICF Technology

Recipient: none: US EPA

Document Number: PAI-001-0113 To 0275 Date: 68/39/88

Title: Final Work Plan for Remedial Investigation/Feasibility Study - Pasley Solvents and Chemicals

Site, Town of Heapstead, Long Island NY

Type: PLAN

Author: Blanar, Edward W: ICF Technology

Recipient: none: US EPA

Document Number: PAI-001-0276 To 0341 Date: 09/01/88

Title: Final Work Plan for Tank Demolition and Removal at the Pasley Solvents and Chemicals Site,

Town of Heapstead, Long Island NY

Type: PLAN

Author: Russell, William 6: EA Engineering Science & Technology

Recipient: none: Commander Dil Corporation

Date: \$3/\$1/89

Document Number: PAI-001-0342 To 0616

Title: Soil Vapor Contaminant Assessment for Remedial Investigation/Feasibility Study - Pasley Solvents

and Chemicals Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Schultz, James A: EA Engineering Science & Technology

Recipient: none: Commander Oil Corporation

Document Number: PAI-881-8617 To 8762 Date: 18/81/91

Title: Remedial Investigation Report - Pasley Solvents & Chemicals Site, Town of Hempstead, Long

Island NY

Type: REPORT

Author: none: Metcalf & Eddy

Recipient: none: Commander Oil Corporation

Document Number: PAI-801-8763 To 8783 Date: 11/88/98

Title: (Letter forwarding attached EPA comments on the Draft Remedial Investigation Report for the

site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation .

Document Number: PAI-801-8784 To 1889 Date: 83/14/91

Title: (Letter forwarding data, received from the Nassau County Department of Public Works for the

Mitchel Field site, to be incorporated into the Pasley Remedial Investigation Report, and transmitting

attached Monitoring Program Sampling Report)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-001-1010 To 1013 . Date: \$3/21/91

Title: (Letter forwarding attached analytical results of groundwater samples from existing wells

at the former Texaco service station, Garden City NY)

Type: CORRESPONDENCE

Author: Brooker, Lauren J: Star Enterprise

Recipient: Mirza, Misbahuddin K: MY Dept of Environmental Conservation

Page: 3

Document Number: PAI-881-1814 To 1817 Date: 85/38/91

Title: (Letter forwarding attached comments from EPA about Metcalf & Eddy's Remedial Investigation

Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-801-1018 To 1018 Date: 07/19/91

Title: (Letter requesting information about any hazardous waste site located near Stewart Avenue

which may be upgradient of the Pasley Solvents & Chemicals site)

. Type: CORRESPONDENCE

Author: Henry, Sherrel D: US EPA

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Document Number: PAI-861-1619 To 1631 Date: 18/84/91

Title: (Letter forwarding attached EPA comments on the third revision of the June 1991 Remedial Investigation

Report)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-001-1032 To 1032 Date: 12/05/91

Title: (Letter approving the revised Remedial Investigation Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Dil Corporation

£3/25/92

## Index Document Number Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 4

Document Number: PAI-801-1833 To 1326 Date: 82/81/92

Title: Feasibility Study Report - Pasley Solvents and Chemicals Site, Town of Hempstead, Nassau County

NY

Type: REPORT

Author: Roth, Robert J: Metcalf & Eddy

Recipient: none: US EPA

Document Number: PAI-881-1327 To 1346 Date: 82/81/92

Dutument Rumber: FRI-801-132/ IU 1340

Title: Superfund Proposed Plan - Pasley Solvents and Chemicals Site, Town of Hempstead NY

Type: PLAN

Author: none: US EPA Recipient: none: none

Document Number: PAI-881-1347 To 1357 Date: 18/24/91

Title: (Letter forwarding attached EPA comments on the Draft Feasibility Study Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-001-1358 To 1368 Date: 12/18/91

Title: (Letter forwarding attached comments on the Feasibility Study Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-881-1361 To 1362 Date: 12/27/91

Title: (Letter containing NYSDEC and NYSDOH comments on the EPA Proposed Plan for the site)

Type: CORRESPONDENCE

Author: O'Toole, Michael J Jr: NY Dept of Environmental Conservation

Recipient: Hauptman, Mel: US EPA

**B3/25/92** 

### Index Document Number Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 5

Document Number: PAI-881-1363 To 1364 Date: #5/31/91

Title: (Letter stating what has to be done to stop the dissolved product plume from moving onto the

property of the Texaco service station)

Type: CORRESPONDENCE

Author: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Recipient: Brooker, Lauren J: Star Enterprise

Document Number: PAI-001-1365 To 1366 Date: 86/17/91

Title: (Letter containing response to NYSDEC correspondence regarding the former Texaco service station

at the site)

Type: CORRESPONDENCE

Author: Brooker, Lauren J: Star Enterprise

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Document Number: PAI-801-1367 To 1384 Date: \$8/19/88

Title: Administrative Order on Consent in the Matter of Commander Dil Corporation

Type: LEGAL DOCUMENT

Author: Muszynski, William J: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Date: 67/11/91

Document Number: PAI-881-1385 To 1385

Title: (Letter regarding the Mitchel Field facility that Purex has constructed pursuant to a consent

judgaent)

Type: CORRESPONDENCE

Author: Smith, Jeffrey M: Purex Industries Inc

Recipient: Henry, Sherrel D: US EPA

## Index Document Number Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 6

Bocusent Number: PAI-891-1386 To 1395 Bate: \$5/81/91

Title: Engineering Bulletin: In Situ Soil Vapor Extraction Treatment

Type: CORRESPONDENCE Author: none: US EPA Recipient: none: none

Document Number: PAI-801-1396 To 1437 Date: 83/18/92

Title: (Transcript of the 83/85/92 Public Meeting for the Pasley Solvents & Chemicals site)

Type: LEGAL DOCUMENT

Author: Lewis, Virginia E: court reporter

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Page: 1

Document Number: PAI-801-1367 To 1384 Date: 88/19/88

Title: Administrative Order on Consent in the Matter of Commander Oil Corporation

Type: LEGAL DOCUMENT

**83/25/92** 

Author: Muszynski, William J: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

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Document Number: PAI-881-8881 To 8112 Date: 88/38/88

Title: Final Field Operations Plan for Remedial Investigation/Feasibility Study - Pasley Solvents

and Chemicals Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Blanar, Edward W: ICF Technology

Recipient: none: US EPA

Document Number: PAI-881-8113 To 8275 Date: 88/38/88

Title: Final Mork Plan for Remedial Investigation/Feasibility Study - Pasley Solvents and Chemicals

Site, Town of Heapstead, Long Island NY

Type: PLAN

Author: Blanar, Edward W: ICF Technology

Recipient: none: US EPA

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Document Number: PAI-601-8276 To 8341 Date: 69/81/88

Title: Final Work Plan for Tank Desolition and Resoval at the Pasley Solvents and Chemicals Site,

Town of Hempstead, Long Island NY

Type: PLAN

Author: Russell, William 6: EA Engineering Science & Technology

Recipient: none: Commander Oil Corporation

Page: 2

Document Number: PAI-881-8342 To 8616 Date: 83/81/89

Title: Soil Vapor Contaminant Assessment for Remedial Investigation/Feasibility Study - Pasley Solvents

and Chemicals Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Schultz, James A: EA Engineering Science & Technology

Recipient: none: Commander Oil Corporation

Document Number: PAI-001-0763 To 0783 Date: 11/88/98

Title: {Letter forwarding attached EPA comments on the Draft Remedial Investigation Report for the

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

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Document Number: PAI-001-0784 To 1809 Date: 83/14/91

mitle: (Letter forwarding data, received from the Nassau County Department of Public Works for the

Mitchel Field site, to be incorporated into the Pasley Remedial Investigation Report, and transmitting

attached Monitoring Program Sampling Report)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-881-1818 To 1813

Title: (Letter forwarding attached analytical results of groundwater samples from existing wells

at the former Texaco service station, Garden City NY)

Type: CORRESPONDENCE

Author: Brooker, Lauren J: Star Enterprise

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

## Index Chronological Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 3

Document Number: PAI-001-1386 To 1395 Date: 05/01/91

Title: Engineering Bulletin: In Situ Soil Vapor Extraction Treatment

Type: CORRESPONDENCE
Author: none: US EPA
Recipient: none: none

Document Number: PAI-881-1814 To 1817 Date: 85/38/91

Title: (Letter forwarding attached comments from EPA about Metcalf & Eddy's Remedial Investigation

Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Dil Corporation

Document Number: PAI-881-1363 To 1364 Date: 85/31/91

Title: (Letter stating what has to be done to stop the dissolved product plume from moving onto the

property of the Texaco service station)

Type: CORRESPONDENCE

Author: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Recipient: Brooker, Lauren J: Star Enterprise

Document Number: PAI-881-1365 To 1366 Date: 86/17/91

Title: (Letter containing response to NYSDEC correspondence regarding the former Texaco service station

at the site)

Type: CORRESPONDENCE

Author: Brooker, Lauren J: Star Enterprise

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

### Index Chronological Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 4

Document Number: PAI-881-1385 To 1385

Date: 87/11/91

Title: (Letter regarding the Mitchel Field facility that Purex has constructed pursuant to a consent

judgment)

Type: CORRESPONDENCE

Author: Smith, Jeffrey M: Purex Industries Inc

Recipient: Henry, Sherrel D: US EPA

Document Number: PAI-881-1818 To 1818

. Date: 87/19/91

Title: (Letter requesting information about any hazardous waste site located near Stewart Avenue

which may be upgradient of the Pasley Solvents & Chemicals site)

Type: CORRESPONDENCE

Author: Henry, Sherrel D: US EPA

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Date: 18/81/91

Document Number: PAI-801-8617 To 8762

Title: Remedial Investigation Report - Pasley Solvents & Chemicals Site, Town of Hempstead, Long

Island NY

Type: REPORT

Author: none: Metcalf & Eddy

Recipient: none: Commander Oil Corporation

Document Number: PAI-001-1019 To 1031

Date: 18/84/91

Title: (Letter forwarding attached EPA comments on the third revision of the June 1991 Remedial Investigation

Report)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

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Page: 5

Document Number: PAI-881-1347 To 1357 Date: 18/24/91

Title: (Letter forwarding attached EPA comments on the Draft Feasibility Study Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-881-1832 To 1832 Date: 12/85/91

Title: (Letter approving the revised Remedial Investigation Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Dil Corporation

Document Number: PAI-801-1358 To 1360 Date: 12/18/91

Title: (Letter forwarding attached comments on the Feasibility Study Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: PAI-801-1361 To 1362 Date: 12/27/91

Title: (Letter containing NYSDEC and NYSDOH comments on the EPA Proposed Plan for the site)

Type: CORRESPONDENCE

Author: O'Toole, Michael J Jr: NY Dept of Environmental Conservation

Recipient: Hauptman, Hel: US EPA

Document Number: PAI-881-1833 To 1326 Date: 02/81/92

Title: Feasibility Study Report - Pasley Solvents and Chemicals Site, Town of Hempstead, Nassau County

MY

Type: REPORT

Author: Roth, Robert J: Metcalf & Eddy

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Page: 6

Document Number: PAI-881-1327 To 1346 Date: 82/81/92

Title: Superfund Proposed Plan - Pasley Solvents and Chemicals Site, Town of Hempstead NY

Type: PLAN

Author: none: US EPA Recipient: none: none

Document Number: PAI-881-1396 To 1437 Date: 03/10/92

Title: (Transcript of the 83/85/92 Public Meeting for the Pasley Solvents & Chemicals site)

Type: LEGAL DOCUMENT

Author: Lewis, Virginia E: court reporter

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### Index Author Name Order PASLEY SOLVENTS & CHEMICALS Documents

Document Number: PAI-881-8617 To 8762 Date: 18/81/91

Title: Remedial Investigation Report - Pasley Solvents & Chemicals Site, Town of Hempstead, Long

Type: REPORT

Author: none: Metcalf & Eddy

Recipient: none: Commander Oil Corporation

Document Number: PAI-801-1327 To 1346 Date: 82/81/92

Title: Superfund Proposed Plan - Pasley Solvents and Chemicals Site, Town of Hempstead NY

Type: PLAN

Author: none: US EPA Recipient: none: none

Document Number: PAI-881-1386 To 1395 Date: 85/81/91

Title: Engineering Bulletin: In Situ Soil Vapor Extraction Treatment

Type: CORRESPONDENCE Author: none: US EPA Recipient: none: none

Document Number: PAI-081-0881 To 0112 Date: 08/30/88

Title: Final Field Operations Plan for Remedial Investigation/Feasibility Study - Pasley Solvents

and Chemicals Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Blanar, Edward W: ICF Technology

Recipient: none: US EPA

Document Number: PAI-801-8113 To 8275 Date: 68/38/88

Title: Final Work Plan for Remedial Investigation/Feasibility Study - Pasley Solvents and Chemicals

Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Blanar, Edward W: ICF Technology

Document Number: PAI-801-1010 To 1013 Date: 03/21/91

Title: (Letter forwarding attached analytical results of groundwater samples from existing wells

at the former Texaco service station, Garden City NY)

Type: CORRESPONDENCE

Author: Brooker, Lauren J: Star Enterprise

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Document Number: PAI-801-1365 To 1366 Date: 86/17/91

Title: (Letter containing response to NYSDEC correspondence regarding the former Texaco service station

at the site)

Type: CORRESPONDENCE

Author: Brooker, Lauren J: Star Enterprise

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Document Number: PAI-881-1818 To 1818 Date: 87/19/91

Title: (Letter requesting information about any hazardous waste site located near Stewart Avenue

which may be upgradient of the Pasley Solvents & Chemicals site)

Type: CORRESPONDENCE

Author: Henry, Sherrel D: US EPA

Recipient: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Document Number: PAI-081-1396 To 1437 Date: 83/18/92

Title: (Transcript of the 03/05/92 Public Meeting for the Pasley Solvents & Chemicals site)

Type: LEGAL DOCUMENT

Author: Lewis, Virginia E: court reporter

## Index Author Name Order PASLEY SOLVENTS & CHEMICALS Documents

Page: 3

Document Number: PAI-001-1363 To 1364 Date: 05/31/91

Title: (Letter stating what has to be done to stop the dissolved product plume from moving onto the

property of the Texaco service station)

Type: CORRESPONDENCE

Author: Mirza, Misbahuddin K: NY Dept of Environmental Conservation

Recipient: Brooker, Lauren J: Star Enterprise

Document Number: PAI-801-1367 To 1384 Date: 88/19/88

Title: Administrative Order on Consent in the Matter of Commander Oil Corporation

Type: LEGAL DOCUMENT

Author: Muszynski, William J: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Document Number: FAI-001-1361 To 1362 Date: 12/27/91

Title: (Letter containing NYSDEC and NYSDOH comments on the EPA Proposed Plan for the site)

Type: CORRESPONDENCE

Author: O'Toole, Michael J Jr: NY Dept of Environmental Conservation

Recipient: Hauptman, Mel: US EPA

Document Number: PAI-881-8763 To 8783 Date: 11/88/98

Title: (Letter forwarding attached EPA comments on the Draft Remedial Investigation Report for the

site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Bil Corporation

Document Number: PAI-881-8784 To 1889 Date: 83/14/91

Title: (Letter forwarding data, received from the Nassau County Department of Public Works for the

Mitchel Field site, to be incorporated into the Pasley Remedial Investigation Report, and transmitting

attached Monitoring Program Sampling Report)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

E3/25/92

Document Number: PAI-BB1-1814 To 1817 Date: 85/38/91 Title: (Letter forwarding attached comments from EPA about Metcalf & Eddy's Remedial Investigation Report for the site) Type: CORRESPONDENCE Author: Petersen, Carole: US EPA Recipient: Shapiro, Joseph 6: Commander Oil Corporation Document Number: PAI-801-1819 To 1831 Date: 10/04/91 Title: {Letter forwarding attached EPA comments on the third revision of the June 1991 Remedial Investigation Report) Type: CORRESPONDENCE Author: Petersen, Carole: US EPA Recipient: Shapiro, Joseph 6: Commander Oil Corporation Document Number: PAI-981-1832 To 1832 Date: 12/85/91 Title: (Letter approving the revised Remedial Investigation Report for the site) Type: CORRESPONDENCE Author: Petersen, Carole: US EPA Recipient: Shapiro, Joseph 6: Commander Oil Corporation Document Number: PAI-881-1347 To 1357 Date: 18/24/91 Title: (Letter forwarding attached EPA comments on the Braft Feasibility Study Report for the site) Type: CORRESPONDENCE Author: Petersen, Carole: US EPA Recipient: Shapiro, Joseph 6: Commander Oil Corporation Document Number: PAI-801-1358 To 1368 Date: 12/18/91

Title: (Letter forwarding attached comments on the Feasibility Study Report for the site)

Type: CORRESPONDENCE

Author: Petersen, Carole: US EPA

Recipient: Shapiro, Joseph 6: Commander Oil Corporation

Borugent Number: PAI-881-1833 To 1326

Date: 82/81/92

Title: Feasibility Study Report - Pasley Solvents and Chemicals Site, Town of Hempstead, Nassau County

Type: REPORT

Author: Roth, Robert J: Metcalf & Eddy

Recipient: none: US EPA

Document Number: PAI-881-8276 To 8341 Date: 89/81/88

Title: Final Work Plan for Tank Demolition and Removal at the Pasley Solvents and Chemicals Site,

Town of Heapstead, Long Island NY

Type: PLAN

Author: Russell, William 6: EA Engineering Science & Technology

Recipient: none: Commander Oil Corporation

Date: 83/81/89

Document Number: PAI-001-0342 To 8616

Title: Soil Vapor Contaminant Assessment for Remedial Investigation/Feasibility Study - Pasley Solvents

and Chemicals Site, Town of Hempstead, Long Island NY

Type: PLAN

Author: Schultz, James A: EA Engineering Science & Technology

Recipient: none: Commander Oil Corporation

Document Number: PAI-081-1385 To 1385 Date: 87/11/91

Title: (Letter regarding the Mitchel Field facility that Purex has constructed pursuant to a consent

judgment)

Type: CORRESPONDENCE

Author: Smith, Jeffrey M: Purex Industries Inc

Recipient: Henry, Sherrel D: US EPA