

# REVISED FOURTH QUARTER 1993 MONITORING REPORT

UNION PACIFIC RAILROAD UNION PACIFIC MOTOR FREIGHT FACILITY OAKLAND, CALIFORNIA

#### PREPARED FOR:

UNION PACIFIC RAILROAD ENVIRONMENTAL MGMT. 1416 DODGE STREET, ROOM 930 OMAHA, NEBRASKA 68179

Prepared by:

USPCI
Consulting Services
5665 Flatiron Parkway
Boulder, Colorado 80301
Project No. 96120-844

May 4, 1994





#### **Consulting Services**



May 16, 1994

Mr. Harry Patterson Union Pacific Railroad 1416 Dodge Street, Room 930 Omaha, Nebraska 68179

RE: "Fourth Quarter 1993 Monitoring Report", UPMF Facility at 1750 Ferro Street, Oakland, California, USPCI Project No. 96120-844

Dear Mr. Patterson:

Enclosed is a copy of the "Revised Fourth Quarter 1993 Monitoring Report", dated May 4, 1994, for the Union Pacific Motor Freight (UPMF) Facility at the referenced site. As discussed in the cover letter for the "First Quarter 1994 Monitoring Report", dated April 29, 1994, a mathematical error had been made by the laboratory with the results for the "Fourth Quarter 1993 Monitoring Report." The error pertained to the total petroleum hydrocarbons as gasoline, and benzene, ethylbenzene, and xylenes concentration results in monitoring wells APL/UP-W1 and APL/UP-W2. Corresponding changes to the text, Table 1 Cumulative Analytical Results of Groundwater Samples", and Figure 4, "Dissolved BTXE Distribution Map" have been incorporated into the revised report. Please replace the enclosed report with the "Fourth Quarter 1993 Monitoring Report" dated January 1994.



Mr. Harry Patterson May 16, 1994 Page 2

If you have any questions, please call me at (303) 938-5539.

Sincerely,

Denton Mauldin Engineer III

cc:

Rick Pollard, USPCI
Steve Brinkman, USPCI
Jennifer Eberle, ACDEH
John Amdur, Port of Oakland
Philip Herden, APL
Ken Fossey, USPCI (cover letter)

Enclosures

DM/tjh

# REVISED FOURTH QUARTER 1993 MONITORING REPORT UNION PACIFIC RAILROAD UNION PACIFIC MOTOR FREIGHT FACILITY OAKLAND, CALIFORNIA USPCI Project No. 96120-844

#### Prepared for:

Union Pacific Railroad
Environmental Management - Room 930
1416 Dodge Street
Omaha, Nebraska 68179

for submittal to

Ms. Jennifer Eberle
Department of Environmental Health
Hazardous Materials Division
80 Swan Way, Room 200
Oakland, California 94621

Prepared by:

USPCI Consulting Services 24125 Aldine Westfield Road Spring, Texas 77373

Christopher S. Byerman

Geologist

Richard M. Pollard Geologist R.G. No. 4659

May 4, 1994

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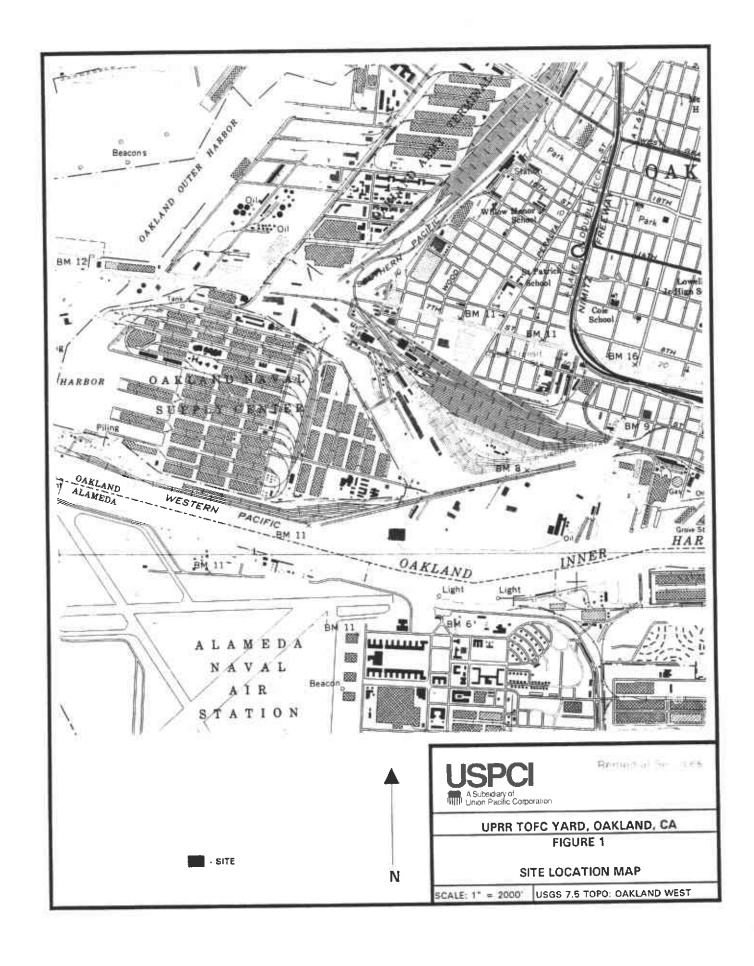
#### 1.0 INTRODUCTION

This Quarterly Monitoring Report has been prepared for Union Pacific Railroad (UPRR) by USPCI in response to a April 29, 1993, Alameda County Department of Environmental Health, Hazardous Materials Division (ACDEH) request for UPRR to begin a quarterly monitoring program at the Union Pacific Motor Freight (UPMF) Ferro Street facility in Oakland, California (See Figure 1). The facility was the site of a release of petroleum hydrocarbons from underground storage tanks (USTs).

The quarterly monitoring event involved:

- Measuring the static water levels, purging, and sampling the ten existing monitoring wells and determining the local groundwater gradient;
- Analyzing groundwater samples from the monitoring wells for total petroleum hydrocarbons (TPH, EPA Method 418.1); TPH diesel (TPH/D, EPA Method 8015 Modified, TPH gasoline (TPH/G, EPA Method 8015 Modified); benzene, toluene, ethylbenzene and xylenes (BTEX, EPA Method 8020); dissolved arsenic (As) and lead (Pb) by EPA Method 6000/7000; and for purgeable halocarbons (EPA Method 8010); and
- Preparation of a Quarterly Monitoring Report.

No phase-separated hydrocarbon (PSH) was detected in the monitoring wells in the Fourth Ouarter 1993 Monitoring Event. However, PSH was observed in at least one monitoring well during previous monitoring events. Groundwater samples were collected from the ten monitoring wells in November 1993. BTEX concentrations in groundwater samples ranged from below the Method Detection Limit (MDL) in monitoring well OKUS-W7 to 13,970 micrograms per liter (ug/L) in monitoring well OKUS-W2 which is located 15 feet south of the former fuel island next to the UPMF repair shop. Benzene concentrations exceeded the California Maximum Contaminant Level [MCL (Marshack, 1989)] in groundwater samples from eight of the ten monitoring wells. Ethylbenzene exceeded the MCL in samples from three of the ten monitoring TPH/G concentrations in groundwater samples ranged from below the MDL in monitoring well OKUS-W6 (450 feet southeast of one source area and 400 feet north-northwest of the another known source area) and OKUS-W7 (375 feet west-southwest of a known source area) to 24,000 ug/L in monitoring well OKUS-W2 (15 feet south of the former fuel island). Groundwater samples from all ten monitoring wells contained detectable concentrations (< 8,000 ug/L) of TPH diesel (TPH/D). Groundwater samples from three monitoring wells also contained minor concentrations [< 3.6 milligrams per liter (mg/L)] of TPH using EPA Method 418.1.



Analytical results indicated detectable concentrations of arsenic (As) in groundwater samples from three of the ten monitoring wells and minor concentrations of chlorobenzene in samples from five of the ten monitoring wells. There was no analytical evidence of chloroform which was present in elevated concentration in samples from three monitoring wells during the Preliminary Site Assessment (PSA). The presence of chloroform was most likely due to a laboratory error. Concentrations of As in groundwater samples ranged from 0.13 mg/L in OKUS-W2 to 0.53 mg/L in OKUS-W5. Chlorobenzene concentrations in groundwater samples ranged from 0.00089 mg/L in OKUS-W1 to 0.078 mg/L in OKUS-W2. The MCL for As is 0.050 mg/L, and the MCL for chlorobenzene is 0.100 mg/L (Marshack, 1989). Dissolved lead was not detected in any of the groundwater samples collected from the ten monitoring wells, which is consistent with previous sampling results.

The arsenic and chlorobenzene detected in groundwater samples are not believed to be related to the contents of the former USTs. The former USTs contained engine oil, waste oil, diesel fuel and gasoline.

Groundwater beneath the site flowed to the east-southeast at a gradient of 0.006 ft/ft on the date measured.

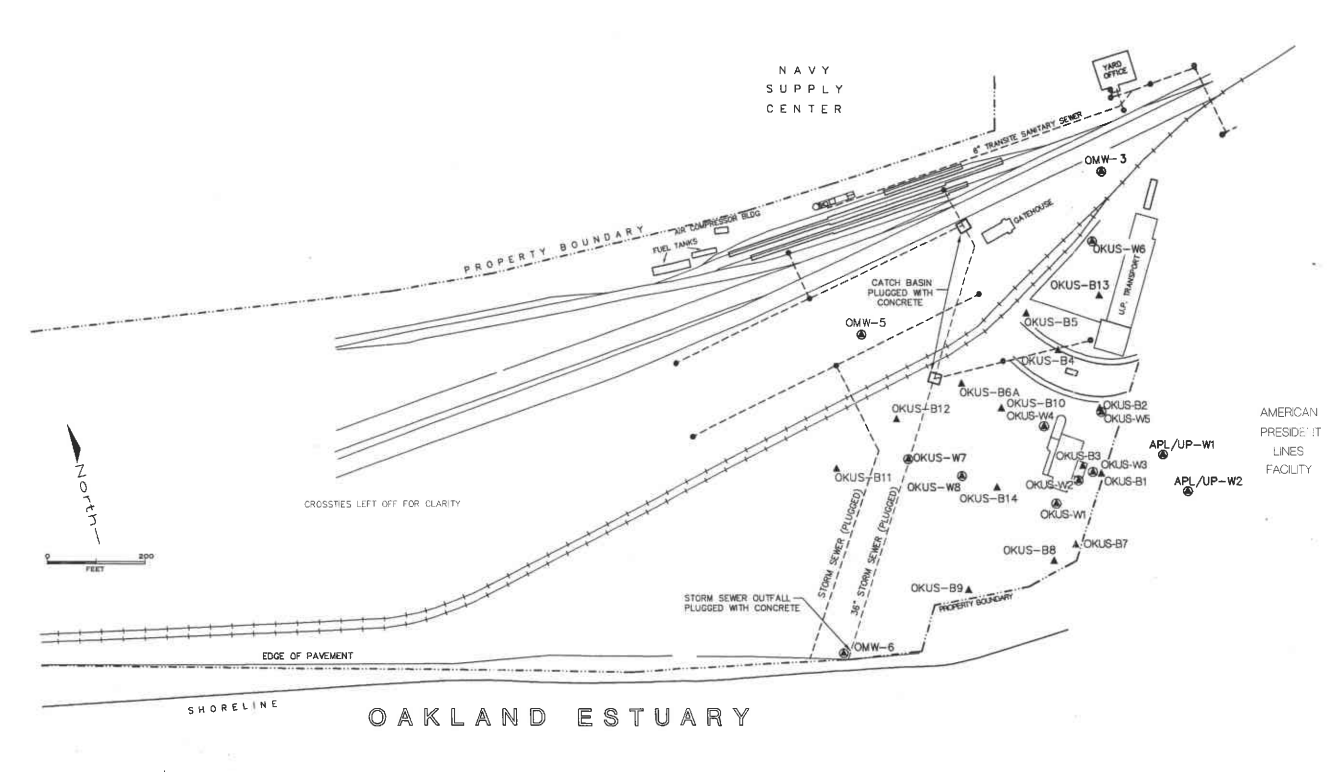
#### 1.1 Site Background

#### 1.1.1 General Description and Previous Activities

The site is located on the southeastern end of the UPRR TOFC Yard (Figure 2) located in the Port of Oakland. The area surrounding the site is used for heavy to light commerce, with residential areas being located inland to the east and west across the Oakland Estuary. Five USTs were removed from the UPMF site from 1987 to 1990 (Figure 3). The refueling portion of the TOFC yard, approximately 700 feet northwest of the truck repair shop, is currently undergoing groundwater remediation for recovery of diesel product. The limits of the diesel plume in that portion of the site have been adequately defined (USPCI, 1991), and the plume does not extend to the area of impacted groundwater at the truck repair facility in the TOFC Yard refueling area.

#### 1.1.2 Location and Access

The site is located in the UPRR TOFC Yard at 1750 Ferro Street in the Port of Oakland on east side of the Inner Harbor, Oakland, California. Access to the site is at the intersection of Middle Harbor Road and Ferro Street.



#### LEGEND

♠ OKUS-W1 MONITORING WELL LOCATION AND NUMBER

▲ OKUS-B1 BORING LOCATION AND NUMBER

CATCH BASIN FOR STORM SEWER

BY DATE
DEATH BILL 9/93
ORDORD
APPROVED
APPROVED
APPROVED
Union Pacific Corporation

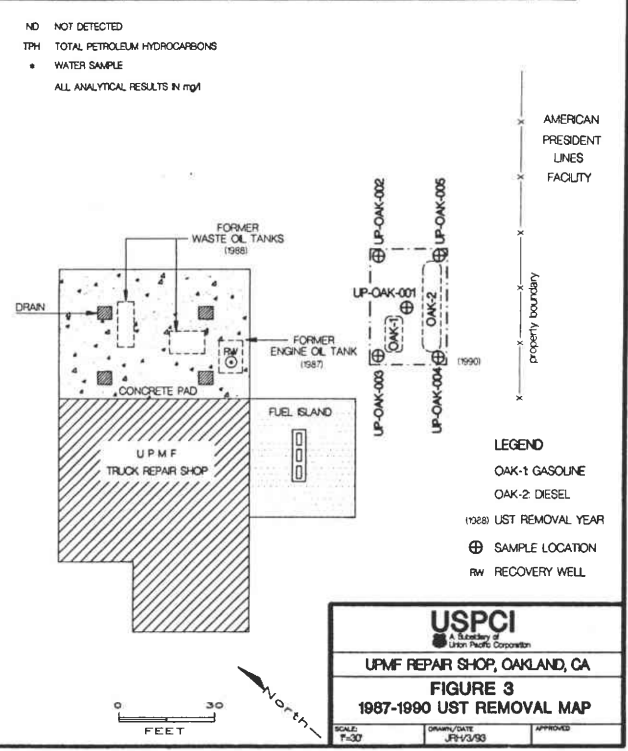
UPRR TOFC RALYARD
UPMF REPAIR SHOP, OAKLAND, CALIFORNIA

FIGURE 2 SITE VICINITY MAP

SCALE | DATE | DWG. NO. 96120-556 |

#### ANALYTICAL RESULTS FOR OAK-1 & OAK-2 REMOVAL IN 1990

SAMPLE LOCATION	6AMPLE ID	DATE SAMPLED	TPH CS-C20 mg/kg	TPH/D mg/kg	TPH/G mo/ka	BENZENE mo/kg	TOLLENE mo/kg	ETHML- BENZENE morks	XYLENE mo/kg	TOTAL BTEX mg/kg
CENTRAL *	UP-OAK-001	2/22/90	ND	NO	ND	0.063	0.023	ND	0.026	0.102
NW WALL	UP-OAK-002	2/22/90	ND	ND	ND	ND	ND	ND	ND	ND
SW WALL	UP-OAK-003	2/22/90	ND	ND	ND	ND	NO	ND	ND	NO
SE WALL	UP-OAK-004	2/22/90	ND	ND	0.032	NO	0.005	ND	0.025	0.025
NE WALL	UP-OAK-006	2/22/90	213	NO	ND	0.004	0.003	0.007	0.012	0.026



#### 1.2. Investigative Procedures

All USPCI field activities, including data recording procedures, decontamination methods, groundwater sample collection, and purge water disposal, were conducted in accordance with USPCI's Quality Assurance/Quality Control (QA/QC) Plan located in Appendix A.

The quarterly monitoring event was conducted by USPCI Remedial Services personnel from the Ontario, California office under the direct supervision of Christopher Byerman and Richard Pollard (California Registered Geologist #4659).

#### 2.0 CHRONOLOGY OF EVENTS

The following section presents a detailed chronology of 1992 and 1993 activities related to the site assessment, along with dates of relevant correspondence between the parties involved.

April 29, 1992	The Alameda County Department of Environmental Health (ACDEH) issued a letter to Mr. Andrew Clark-Clough of the Port of Oakland and
	Mr. John Seagle of UPRR requesting that a Preliminary Site Assessment
	(PSA) be performed at the UPRR facility at 1750 Ferro Street, Oakland,
	California.

June 10, 1992	UPRR contacted Mr. Paul Smith of the ACDEH regarding the ACDEH
<b>52.10</b> 23, 2112	letter dated April 29, 1992 concerning UPRR's UST removals at 1750
	Ferro Street, Oakland, California.

June 18, 1992	USPCI submitted a P\$A workplan to UPRR for review and submittal to
· ·	the ACDEH and the Port of Oakland presenting the proposed workscope,
	technical information, and methods used to conduct the assessment.

July 10, 1992	UPRR submitted the PSA workplan, along with analytical data from the
	December 1987 excavation and the 1990 UST removals, to Ms. Susan
	Hugo of the ACDEH and Ms. Michelle Heffes of the Port of Oakland.

- July 28, 1992 USPCI responded to a request from Ms. Jennifer Eberle of the ACDEH for supplemental information for the PSA workplan.
- August 6, 1992 The Port of Oakland responded to UPRR on USPCI's proposed PSA.

  Ms. Michelle Heffes of the Port of Oakland informed UPRR that permits would be required from the Port of Oakland, Bay Conservation and Development Commission (BCDC) and the Alameda County Flood

Control District (ACFCD) before proceeding with the PSA. A permit to install groundwater monitoring wells on Port property was also required.

- August 10, 1992 USPCI responded to a second request from Ms. Jennifer Eberle of the ACDEH for supplemental information for the PSA workplan.
- August 25, 1992 ACDEH approved USPCI's PSA workplan with comments. ACDEH also informed USPCI that Ms. Jennifer Eberle will be overseeing this case.
- November 10, 1992 USPCI requested a permit from the ACFCD and the Port of Oakland to install monitoring wells at the UPMF facility at 1750 Ferro Street, Oakland, California.
- November 16, 1992 USPCI received a permit application number (92580) from the ACFCD for the construction of the monitoring wells.
- December 6, 1992 USPCI received a permit application approval from the ACFCD for the construction of the monitoring wells.
- January 7, 1993 USPCI requested and received ticket numbers from UPRR Fiber Optics (# 672732) and Underground Service Alert (# 4997). Underground Service Alert notified all utilities that had underground lines in the area to mark lines by 8:00 AM, January 12, 1993. USPCI also informed Ms. Jennifer Eberle of the ACDEH that the PSA would begin on January 12, 1993.
- January 11, 1993 USPCI personnel arrived at the UPMF facility to collect site information and plan drilling activities.
- January 12-15, 1993 USPCI conducted a PSA on the UPMF facility at 1750 Ferro Street. Five monitoring wells and seven soil borings were installed. Soil and groundwater samples were collected and analyzed as part of the assessment.
- February 2, 1993 USPCI contacted Ms. Jennifer Eberle of the ACDEH to discuss report format and relay analytical information collected in the PSA.
- February 18, 1993 USPCI personnel collected additional groundwater samples due to elevated concentrations of metals and minor concentrations of semivolatiles and purgeable halocarbons that were found in several soil samples. Composite samples of drummed soil cuttings and purge/decon water were also collected for final treatment and/or disposal.

February 25, 1993	Mr. Craig Mayfield from the ACFCD contacted USPCI by letter and requested copies of the well construction diagram, boring logs and a site map showing the locations of the borings/monitoring wells. USPCI forwarded this information to Mr. Mayfield on February 26, 1993.
April 21, 1993	USPCI forwarded the completed Preliminary Site Assessment Report to UPRR, ACDEH, CRWQCB (San Francisco Bay) and the Port of Oakland.
April 29, 1993	Ms. Jennifer Eberle of the ACDEH informed UPRR that the county agrees with USPCI recommendations to begin a quarterly monitoring program and complete a Phase II Site Assessment.
May 12, 1993	USPCI personnel conducted a quarterly monitoring event as part of the agreed recommendations resulting from the PSA between the ACDEH, UPRR and USPCI. The five existing monitoring wells were purged and sampled. The analytical results were similar to the PSA groundwater results.
June 23, 1993	USPCI received a new permit number (93341) from Mr. Wyman Hong of the ACFCD for the Phase II Assessment.
July 9, 1993	USPCI received a verbal confirmation from Mr. Joe Marsh of the Port of Oakland that the previous permit number (3443) was still valid for the Phase II Assessment.
July 11-16, 1993	USPCI conducted a Phase II Site Assessment on the UPRR facility at 1750 Ferro Street. Five monitoring wells and eight soil borings were installed. Soil and groundwater samples were collected and analyzed as part of the assessment.
July 30, 1993	At the request of UPRR, USPCI forwarded the completed Second Quarter 1993 Monitoring Report to representatives of the ACDEH, CRWQCB (San Francisco Bay Region), and the Port of Oakland.
August 27, 1993	USPCI completed the Third Quarter 1993 Monitoring Event at the UPMF facility. The ten existing monitoring wells were purged and sampled. The analytical results were similar to the previous groundwater results.
October 29, 1993	At the request of UPRR, USPCI forwarded the completed Phase II Site Assessment Report to representatives of the ACDEH, CRWQCB (San Francisco Bay), the Port of Oakland and American President Lines.

November 12, 1993 USPCI completed the Fourth Quarter 1993 Monitoring Event at the UPMF facility. The ten existing monitoring wells were purged and sampled. The analytical results were similar to the previous groundwater results.

#### 3.0 FIELD INVESTIGATION RESULTS

#### 3.1 Groundwater Assessment Determinations

#### 3.1.1 Groundwater Characteristics

The UPRR Oakland TOFC Railyard and UPMF Facility (Figure 2) are immediately adjacent to the Oakland Estuary, which is located in the eastern portion of the San Francisco Bay. The close proximity of the Estuary to the site suggests that a direct hydrologic connection may exist between the Estuary and the groundwater beneath the site. Tidal influences from the Estuary may influence water levels in the monitoring wells at the site; however, previous studies in the San Francisco Bay Area suggests that tidal influences are generally minimal and are only detectable in monitoring wells in very close proximity to the Bay (usually within 200 feet; USPCI, 1991). The actual degree of influence is dependent on individual site characteristics.

#### 3.1.2 Results of Laboratory Analysis of Groundwater Samples

Analytical results indicate elevated TPH/G and BTEX concentrations in groundwater at the site (Figure 4 and Table 1). Total BTEX concentrations ranged from below analytical detection limits in the sample from monitoring well OKUS-W7 to 13,970 ug/L in sample OKUS-W2. Benzene concentrations were above the MCL (0.001 mg/L) in samples from eight of the ten monitoring wells. Ethylbenzene concentrations were above the MCL (680 mg/L) in samples from three of the ten monitoring wells. TPH/G concentrations ranged from below MDLs in samples OKUS-W6 and OKUS-W7 to 24,000 ug/L in sample OKUS-W2. TPH/D concentrations ranged from 160 ug/L in sample OKUS-W1 to 7700 ug/L in sample OKUS-W2. TPH 418.1 concentrations ranged from below the MDL in groundwater samples from wells OKUS-W1, -W4, -W6, -W7, -W8, APL/UP-W1 and APL/UP-W2 to 3.5 mg/L in a groundwater sample from well OKUS-W2.

TABLE 1: CUMULATIVE ANALYTICAL RESULTS OF GROUNDWATER SAMPLES AT THE UNION PACIFIC MOTOR FREIGHT FACILI OAKLAND, CALIFORNIA - USPCI PROJECT NO. 96120-844

SAMPLE LOCATION	SAMPLE	DATE SAMPLED	TPH/IR	TPH/D	TPH/G	BENZENE	TOLUENE	ETHYL- BENZENE	TOTAL XYLENES	TOTAL BTEX	As	Pb
			mg/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	mg/L	mg/L
OKUS-W1	OKUS-W1	01/14/93	ND	ND	410	20	4	220	ND	240	ND	N
	20000	05/12/93	80	120	ND	ND	ND	ND	ND	ND	ND	N
		08/25/83	ND	100	ND	ND	ND	ND	ND	ND	ND	N
		11/11/93	ND	160	91	1.1	0.88	21	1.6	25	ND	1
OKUS-W2	OKUS-W2	01/14/93	2.5	5400	14000	480	92	8500	ND	9100	0.036	1
0	н	05/12/93	ND	2800	8800	220	47	4600	100	5000	0.093	ı
		08/25/93	5,8	6500	22000	420	92	10000	210	11000	0.089	1
		11/11/93	3.5	7700	24000	540	150	13000	280	14000	ND	I
OKUS-W3	OKUS-W3	01/14/93	4.5	4200	4900	230	42	2600	44	2900	NA	ı
a	•	05/12/93	1.7	4400	4600	290	60	3500	72	3900	0.14	I
		08/25/93	1.5	2700	9400	280	55	4300	41	4700	0.08	I
		11/11/93	2.3	5000	9500	390	110	5100	130	5700	0.14	I
OKUS-W4	OKUS-W4	01/15/93	2.5	5400	8900	300	ND	4500	ND	4800	NA	
il i	0	05/12/93	1.3	2900	6000	320	110	4600	230	5300	0.16	
		08/26/93	ND	2200	6700	350	72	4800	130	5300	0.098	
		11/11/93		2400	5500	250	53	4600	140	5000	0.13	
OKUS-W5	OKUS-W5	01/15/93	ND	2900	550	53	11	180	20	260	NA	
		05/12/93	130	2100	550	81	14		37	380	0.56	
		08/25/93	PHASE SEP	ARATED HY	DROCARB	ONS - WEL	L NOT SAM			2000		
		11/11/93		1600	590		3.1	54	6.2	77	0.53	
OKUS-W6	OKUS-W6	07/16/93	BRK	BRK	ND	2.5				2.5	0.004	
TANKERS ME		08/25/93		590	ND	2.6	ND	4.9	1.3	8.8	0.013	
		11/12/93	ND	610	ND	3.6	ND	3.7	1.3	8.6	ND	
OKUS-W7	OKUS-W7	07/16/93	16	ND	ND		ND			2.1	0.009	
	velolitic (see (1876)	08/25/93	ND	930	56					4.1	ND	
		11/12/93	7 10 00 00	1100	ND	ND	ND	ND	ND	ND	ND	
OKUS-W8	OKUS-W8	07/16/93	15	ND	ND					ND	0.012	0.0
		08/27/93		1100	120					2.2	ND	0.0
		11/11/93	6.755.75.75	1300	190	3.5	1.3	46	4.9	56	ND	

ND - Not Detected

NA - Not Analyzed

BRK - Bottle broken during shipment

TPH - Total Petroleum Hydrocarbons

mg/L - milligram per liter

ug/L - microgram per liter

TPH/IR - analyzed using EPA Method 418.1

TPH/D - analyzed using EPA Method 8015 Mod.

TPH/G - analyzed using EPA Method 8015 Mod.

BTEX - analyzed using EPA Method 8020

As - analyzed using EPA Method 7060

Pb - analyzed using EPA Method 7421

TABLE 1: CUMULATIVE ANALYTICAL RESULTS OF GROUNDWATER SAMPLES AT THE UNION PACIFIC MOTOR FREIGHT FACILI OAKLAND, CALIFORNIA - USPCI PROJECT NO. 96120-844

SAMPLE LOCATION	SAMPLE	DATE SAMPLED	TPH/IR	TPH/D	TPH/G	BENZENE	TOLUENE	ETHYL- BENZENE	TOTAL XYLENES	TOTAL BTEX	As	Pb
			mg/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	ug/L	mg/L	mg/L
APL/UP-W1	APL/UPW1	07/16/93	1	700	300	25.4	1.7	ND	3.0	30	0.011	Nſ
		08/26/93	N	D 810	720	47	1.3	360	14.0	420	0.013	N
		11/11/93		ID 530	560	26	2 ND	220	11.0	260	ND	N
APL/UP-W2	APL/UP-W2	07/16/93	,	9 ND	ND	8.0	ND	ND	ND	8.0	0.016	N
		08/26/93	s N	ID 240	94	, ND	ND	35	2.4	37.0	0.023	N
		11/11/93	N N	ID 190	110	5.0	ND ND	38	2.6	46	ND	N
OKUS-W5	OKUS-W6	01/15/93	. N	ID 2800	510	50	10	170	19	250	NA	N
OKUS-W1	OKUS-W6	05/12/93		ID 140	ND	ND	ND	ND	ND	ND	ND	N
APL/UP-W1	QA/QC-1	07/16/93		2 ^ND	0.21	22.4	ND	ND	2.4	25	0.012	N
OKUS-W4	OKUS-W9	08/26/93	N	ID 2700	6200	340	78	4500	100	5000	0.1	N
OKUS-W8	OKUS-W9	11/11/93	ı	ID 1300	120	1.3	ND	4	1.4	6.7	2.4	NI
UPMF	OAK-FB 1	07/16/93	N	IA NA	NA		ND	ND		ND	NA	NA
UP <b>MF</b>	OAK-TB 2	07/16/93	1	IA NA	NA	ND ND	ND	ND		ND	NA	NA
UP <b>MF</b>	TB-1	08/27/93	1	IA NA	NA		ND	ND		ND	NA	NA
UPMF	TB-2	08/27/93	1	IA NA	NA		ND	ND		ND	NA	NA
UPMF	TB-1	11/12/93	l N	IA NA	NA	ND	ND	ND	ND	ND	NA	NA

ND - Not Detected

NA - Not Analyzed

BRK - Bottle broken during shipment

TPH - Total Petroleum Hydrocarbons

mg/L - milligram per liter

ug/L - microgram per liter

TPH/IR - analyzed using Method 418.1

TPH/D - analyzed using Method 8015 Mod. TPH/G - analyzed using Method 8015 Mod. BTEX - analyzed using Method 8020

As - analyzed using Method 7060

Pb - analyzed using Method 7421

TABLE 1. CUMULATIVE ANALYTICAL RESULTS OF GROUNDWATER SAMPLES AT THE UNION PACIFIC MOTOR FREIGHT FACILITY, OAKLAND, CALIFORNIA USPCI PROJECT NO. 96120-844

SAMPLE	SAMPLE ID	DATE SAMPLED	BROMO- FORM ug/L	CHLORO- BENZENE ug/L	CHLORO- FORM ug/L	-CHLOROETHY VINYL ETHER ug/L	DIBROMOCHLOR- OMETHANE ug/L	1,1-DICHLORO- ETHENE ug/L	cis-1,2-DICHLORO ETHENE ug/L	cle-1,3-DICHLORO- PROPANE ug/L	1,1,2,2-TETRA- CHLOROETHENE ug/L	1,1,1-TRICHL- OROETHANE ug/L
OKUS-W1	OKUS-W1	1/14/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
OKOS WI	OKOS WI	5/12/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
		8/25/83	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
		11/11/93	ND	0.89	ND	ND	ND	ND	ND	ND	ND	ND
OKIIE W/3	OKUS-W2	1/14/93	ND	14	290	15	ND	ND	ND	9	18	5
UKU3.442	0K03-412	5/12/93	ND	26	ND	ND	ND	ND	ND	ND	ND	ND
		8/25/93	ND	78	ND	20	18	21	20	10	20	ND
		11/11/93	ND	78	ND	ND	ND	ND	ND	ND	ND	ND
UNITE MIS	OKUS-W3	1/14/93	ND	15	140	ND	ND	ND	ND	ND	11	ND
H	-	5/12/93	ND	16	ND	ND	ND	ND	ND	ND	ND	ND
		8/25/93	ND	22	ND	5.4	3	3.2	ND	3.3	10	ND
		11/11/93	ND	11	ND	ND	ND	ND	ND	ND	ND	ND
OKUS.WA	OKUS-W4	1/15/93	ND GN	ND	75	ND	ND	ND	6.4	9.4	16	ND
0.000 114	N N	5/12/93	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
		8/26/93	ND	30	ND	ND	ND	ND	ND	ND	16	ND
		11/11/93	ND	32	ND	ND	ND	ND	ND	ND	ND	ND
OKUS.W5	OKUS-W5	1/15/93	5.9	ND	5.9	ND	ND	ND	ND	0.7	4.2	ND
- TOD-113	*	5/12/93	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
		8/25/93	''''				VELL NOT SAMPLED	)				
		11/11/93	ND	1.1	ND	ND	ND	ND	ND	ND	ND	ND
OKUS.W6	OKUS-W6	7/16/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
ONCO NO	01100 110	8/25/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
		11/12/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
OKUS.W7	OKUS-W7	7/16/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
0.100 117	2.100 117	8/25/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
		11/12/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
OKUS-W8	OKUS-W8	7/16/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
2.100 110	<b>,,,</b>	8/27/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
		11/11/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND

ND - Not Detected NA - Not Analyzed

BRK - Bottle broken during shipment

TPH - Total Petroleum Hydrocarbons

mg/L - milligram per liter ug/L - microgram per liter BROMOFORM - analyzed using EPA Method 8010
DIBROMOCHLOROETHANE - analyzed using EPA Method 8010
1,1-DICHLOROETHANE - analyzed using EPA Method 8010
cis-1,2-DICHLOROETHENE - analyzed using EPA Method 8010
cis-1,3-DICHLOROPROPANE - analyzed using EPA Method 8010
1,1,2,2-TETRACHLORETHANE - analyzed using EPA Method 8010
1,1,1-TRICHLOROETHANE -- analyzed using EPA Method 8010

TABLE 1. CUMULATIVE ANALYTICAL RESULTS OF GROUNDWATER SAMPLES AT THE UNION PACIFIC MOTOR FREIGHT FACILITY, OAKLAND, CALIFORNIA USPCI PROJECT NO. 96120-844

SAMPLE LOCATION	SAMPLE ID	DATE SAMPLED	BROMO- FORM ug/L	CHLORO- BENZENE ug/L	CHLORO- FORM ug/L	-CHLOROETHY VINYL ETHER ug/L	DIBROMOCHLORO METHANE ug/L	1,1-DICHLORO- ETHENE ug/L	1,2-DICHLORO- ETHENE ug/L	1,3-DICHLORO- PROPANE ug/L	1,1,2,2-TETRA- CHLOROETHENE ug/L	1,1,1-TRICHL- OROETHANE ug/L
A DI // ID 14/1	APL/UP-W1	7/16/93	ND	ND	5.4	ND	ND	ND	ND	ND	ND	4.2
AFL/OF-WI	AFE/OF-WI	8/26/93	ND	0.75	ND	ND	ND	ND	0.69	0.6	2.2	
		11/11/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND ND
APL/UP-W2 APL/UP-W2	7/16/93	ND	ND	2.5	ND	ND	ND	ND	ND	ND	3.7	
	8/26/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	
		11/11/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
QA/QC												
OKUS-W5	OKUS-W6	1/15/93	NA NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
OKUS-W1	OKUS-W6	5/12/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
APL/UP-W1		7/16/93	ND	ND	9.3	ND	ND	ND	ND	ND	ND	1.2
OKUS-W4	OKUS-W9	8/26/93	ND	22	ND	ND	ND	ND	ND	ND	15	ND
OKUS-W8	OKUS-W9	11/11/93	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
UPMF	OAK-FB 1	7/16/93	NA NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
UPMF	OAK-TB 2	7/16/93	NA NA	NA	NA	NA	NA:	NA	NA	NA	NA	NA
UPMF	TB-1	8/27/93	NA.	NA	NA	NA	NA	NA	NA	NA	NA	NA
UPMF	TB-2	8/27/93	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
UPMF	TB-1	11/12/93	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

ND - Not Detected NA - Not Analyzed

BRK - Bottle broken during shipment

TPH - Total Petroleum Hydrocarbons

mg/L - milligram per liter ug/L - microgram per liter BROMOFORM - analyzed using EPA Method 8010
CHLOROBENZENE - analyzed using Method 8010
CHLOROFORM - analyzed using Method 8010
2-CHLOROETHYLVINYL ETHER - analyzed using Method 8010
DIBROMOCHLOROETHANE - analyzed using Method 8010
1,1-DICHLOROETHANE - analyzed using Method 8010
cis-1,2-DICHLOROETHENE - analyzed using Method 8010
cis-1,3-DICHLOROPROPANE - analyzed using Method 8010
1,1,2,2-TETRACHLORETHANE - analyzed using Method 8010
1,1,1-TRICHLOROETHANE - analyzed using Method 8010

Groundwater samples were also analyzed for dissolved arsenic, lead, and purgeable halocarbons. The analytical results indicated dissolved arsenic in samples from three of the ten monitoring wells. The detected concentrations of dissolved arsenic in groundwater samples ranged from 0.13 mg/L in OKUS-W4 to 0.53 mg/L in OKUS-W5. Three groundwater samples contained concentrations of dissolved arsenic exceeding the MCL of 0.050 mg/L (Table 1). Dissolved lead was not detected above the MDL in any of the existing wells. Chlorobenzene was detected in groundwater water samples from five of the ten monitoring wells. Detected chlorobenzene concentrations in groundwater samples ranged from 0.00089 mg/L in OKUS-W1 to 0.078 mg/L in OKUS-W2. None of the groundwater samples contained chlorobenzene above the MCL of 0.100 mg/L. A complete listing of these analytical results are in Table 1.

#### 3.1.3 Groundwater Gradient

The static water levels measured in November 1993 in wells completed within the borings ranged from 6.43 to 10.30 feet below ground surface (BGS). The data from the water level measurements and mouth-of-casing elevations was used to calculate a local groundwater gradient of approximately 0.006 foot per foot to the east-southeast, toward the Oakland Estuary. The gradient data is consistent with the January 15, 1993 results. The data from the measurements and measuring point elevations (Table 2) were used to prepare the potentiometric surface map (Figure 5). Well stabilization and sampling reports are located in Appendix B.

#### 4.0 SUMMARY AND CONCLUSIONS

#### 4.1 Nature and Extent of Contamination

The analytical results from the 1993 site assessments and groundwater monitoring program indicate a dissolved plume of BTEX and TPH/G is not limited to the immediate area surrounding the UPMF facility. Other contaminants not clearly associated with the UST system were also detected in groundwater beneath the site. The downgradient lateral extent of groundwater impacted either by hydrocarbons or other contaminants has not yet been defined.

TABLE 2
CUMULATIVE FLUID LEVEL MEASUREMENT DATA
UNION PACIFIC RAILROAD MOTOR FREIGHT FACILITY
OAKLAND, CALIFORNIA
USPCI PROJECT NO. 96120-844

WELL NO.	ELEV.*	DATE	DEPTH TO PRODUCT	PRODUCT ELEVATION	PRODUCT THICKNESS	DEPTH TO WATER	WATER ELÉV. (UNCORRECTED)	WATER ELEV. (CORRECTED)
OKUS-W1	9.17	1/14/93	N/A	N/A	NP	8.42	0.75	0.75
	9.17	1/15/93	N/A	N/A	NP	8.45	0.72	0.72
	9,17	2/18/93	N/A	N/A	NP	7.79	1.38	1.38
	9.17	5/12/93	N/A	N/A	NP	8.04	1.13	1.13
	9.17	8/25/93	N/A	N/A	NP	8.61	0.56	0.56
	9.17	11/11/93	N/A	N/A	NP	9.24	-0.07	-0.07
OKUS-W2	9.71	1/14/93	N/A	N/A	NP	9.08	0.63	0.63
	9.71	1/15/93	N/A	N/A	NP	9.12	0.59	0.59
	9.71	2/18/93	N/A	N/A	NP	8.70	1.01	1.01
	9.71	5/12/93	N/A	N/A	NP	9.04	0.67	0.67
	9.71	8/25/93	N/A	N/A	NP	9.61	0.10	0.10
	9.71	11/11/93	N/A	N/A	NP	10.20	-0.49	-0.49
OKUS-W3	9.8	1/14/93	N/A	N/A	NP	9.39	0.41	0.41
	9.8	1/15/93	N/A	N/A	NP	9.33	0.47	0.47
	9.8	2/18/93	N/A	N/A	NP	8.85	0.95	0.95
	9,8	5/12/93	N/A	N/A	NP	9.23	0.57	0.54
	9.8	8/25/93	N/A	N/A	NP	9.82	-0.02	-0.02
	9.8	11/11/93	N/A	N/A	NP	10.30	-0.50	-0.5
OKUS-W4	7.35	1/14/93	N/A	N/A	NP	6.43	0.92	0.92
	7.35	1/15/93	N/A	N/A	NP	6.44	0.91	0.91
	7.35	2/18/93	N/A	N/A	NP	5.77	1.58	1.58
	7.35	5/12/93	6.39	0.96	0.01	6.40	0.95	0.96
	7.35	8/25/93	N/A	N/A	NP			
	7.35	11/11/93	N/A	N/A	N/A	7.10	0.25	0.25

<sup>\*</sup> All well casings measured to mean sea level (MSL).

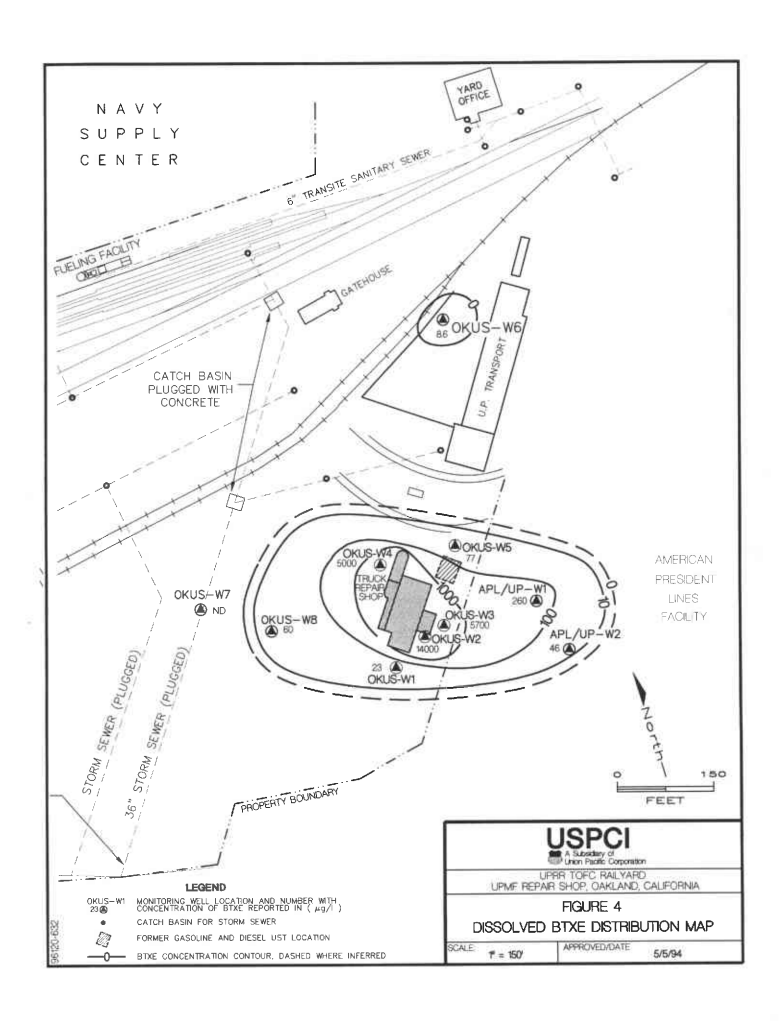
N/A Non Applicable NP - No Product

# TABLE 2 cont. CUMULATIVE FLUID LEVEL MEASUREMENT DATA UNION PACIFIC RAILROAD MOTOR FREIGHT FACILITY OAKLAND, CALIFORNIA USPCI PROJECT NO. 96120-844

WELL NO.	ELEV.*	DATE	DEPTH TO PRODUCT	PRODUCT ELEVATION	PRODUCT THICKNESS	DEPTH TO WATER	WATER ELEV. (UNCORRECTED)	(CORRECTED)
OKUS-W5	9.25	1/14/93	N/A	N/A	NP	9,13	0.12	0.12
	9.25	1/15/93	N/A	N/A	NP	9.15	0.10	0.10
	9.25	2/18/93	N/A	N/A	NP	8.85	0.40	0,40
	9.25	5/12/93	9.18	0.07	0.02	9.20	0.05	0.07
	9.25	8/25/93	8.82	0.43	0.02	8.84	0.41	0.16
	9.25	11/11/93	N/A	N/A	NP	10.15	-0.80	-0.80
OKUS-W6	7.02	7/16/93	N/A	N/A	NP	6.20	0.82	0.82
0702-110	7.02	8/25/93	N/A	N/A	NP	6.52	0.50	0.50
	7.02	11/12/93	N/A	N/A	NP	7.22	-0.20	-0.20
	7.02	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,						
OKUS-W7	6.91	7/16/93	N/A	N/A	NP	5.72	1.19	1.19
01100-117	6.91	8/25/93	N/A	N/A	NP	5.94	0.97	0.97
	6.91	11/12/93	N/A	N/A	NP	6.50	0.39	0.39
OKUS-W8	6.75	7/16/93	N/A	N/A	ΝP	5.56	1.19	1.19
	6.75	8/27/93	N/A	N/A	NP	5.88	0.97	0.87
	6.75	11/11/93	N/A	N/A	NP	6.43	0.33	0.33
					110	10.00	-1.90	-1.90
APL/UP-W1	8.12	7/16/93	N/A	N/A	NP	10.02	-1.81	-1.81
	8.12	8/26/93	N/A	N/A	NP	9.93		
	8.12	11/11/93	N/A	N/A	NP	10.25	-2.13	-2.13
APL/UP-W2	7,31	7/16/93	N/A	N/A	NP	9.38	-2.07	-2.07
	7.31	8/26/93	N/A	N/A	NP	9.20	-1.89	-1.89
	7.31	11/11/93	N/A	N/A	NP	9.65	-2.34	-2.34

N/A Non Applicable NP - No Product

<sup>\*</sup> All well casings measured to mean sea level (MSL).



#### 4.2 Conclusions

TPH/G and BTEX were detected in groundwater samples from wells in the vicinity of the truck repair shop/UST system as well as up and downgradient from the former UST system. Benzene concentrations were above the MCL in samples from nine of the ten existing monitoring wells. Ethylbenzene concentrations were above the MCL in samples from five of the ten monitoring wells. TPH/G concentrations in groundwater samples were as high as 24,000 ug/L. The BTEX and TPH/G concentrations in samples from the downgradient off-site monitoring wells has essentially remained the same as previous monitoring events.

The concentrations of dissolved arsenic exceeded the MCL of 0.050 mg/L in groundwater samples from three of ten wells. Chlorobenzene was detected in groundwater water samples in five of the ten monitoring wells. The chlorobenzene concentrations did not exceed the MCL of 0.100 mg/L. No source for the detectable concentrations of dissolved arsenic and chlorobenzene has been determined.

#### 5.0 RECOMMENDATIONS

USPCI's recommendations for this site were listed in the Phase II Site Assessment Report, and include:

- Continue the quarterly monitoring program throughout 1994 to increase the data base on groundwater characteristics.
- Begin extracting the PSH from the recovery well and determine if PSH exists in any of the monitoring well.
- Conduct an aquifer pump test to determine the hydraulic conductivity of the impacted area.

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#### USPCI

### QUALITY ASSURANCE / QUALITY CONTROL PLAN

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### USPCI Ouality Assurance /Quality Control Plan

#### INTRODUCTION

The USPCI quality assurance/quality control program (QA/QC) is intended to facilitate the acquisition of accurate and reliable data for environmental assessments.

The Quality Assurance Program is a totally integrated program for assuring the reliability of laboratory data, including quality planning, quality assessment and quality improvement efforts to meet project requirements at an economical level. Quality Assurance incorporates procedures for field sampling, sample handling and storage, analytical quality control and document preparation and review.

The Quality Control Program is a routine application of procedures such as blanks, spikes and spike duplicates for obtaining prescribed standards of performance in the measuring process. Quality Control is an audit of the overall Quality Assurance Program. Both programs are necessary to provide accurate data and documentation for investigations and laboratory analyses. The following personnel requirements and field and laboratory procedures will be implemented to ensure that QA/QC objectives are met on all USPCI projects.

#### 1.0 FIELD PERSONNEL

All USPCI Project management personnel are formally educated and, at minimum, hold bachelor's degrees in their fields of technical expertise. In addition, many have advanced degrees in their technical disciplines. Where applicable, USPCI professionals are State Registered or Certified in their fields of expertise. Experience levels will vary depending on job responsibilities. Project Managers have at least five years experience in conducting and managing environmental drilling and sampling programs. Field personnel are experienced in conducting field activities involving monitoring well installation, water sampling, aquifer testing, core description, field program management, and data analysis.

Due to the importance of protecting the health of USPCI employees, subcontractor personnel and others, all on-site workers involved in USPCI projects must have Occupational Safety and Health Administration (OSHA) 40-hour Health and Safety Training Certification. Additionally, USPCI personnel receive periodic training in the use of special equipment for air monitoring and contaminant detection, excavation and shoring, and computerized project management systems.

#### 2.0 FIELD TECHNIQUES

#### 2.1 Recording of Field Data

All information pertinent to the field investigation will be documented on field forms. Information to be documented includes at least the following:

- O Sample numbers
- O Locations of sample collection
- O Soil boring or well numbers, as applicable
- O Depths at which samples were obtained
- O Names of sample collectors
- O Dates and times of collection
- O Purpose of sample
- O Sample distribution (e.g., laboratory, archive, etc.)
- Field observations
- o Field measurements (e.g., PID readings, Ph, conductivity, water levels).
- Other data records (e.g., development log, soil sampling report, well log, etc.)

#### 2.2 Field Equipment Calibration and Maintenance

The following measurement or monitoring equipment may be used during environmental assessments. Equipment is grouped by field activity. Calibration procedures and frequency are listed for each piece.

Soil Borings and Well dimensions- Steel and coated cloth tape. Calibration: none.

<u>Water Level Measurements in Wells- Steel surveyors tape.</u> Calibration: manufacturer supplied temperature correction will be applied as applicable for field conditions. Electrical well sounders. Calibration: check against steel surveyor's tape.

Organic Vapors- Photoionization detector. Calibration: daily field calibration using an isobutylene standard as per manufacturers instructions.

Groundwater pH Measurement- Digital pH meter. Calibration: standard pH solutions of 4, 7, and 10 will be utilized for daily field calibration according to manufacturers instructions.

<u>Electrical Conductivity</u>- Electrical conductivity meter. Calibration: factory-calibrated annually and periodically calibrated against laboratory prepared standard calibration solution.

<u>Water Temperature</u>- Mercury or digital thermometers. Calibration: factory-calibrated once.

<u>Combustible Gas/Oxygen</u>- Combustible gas/oxygen meter calibration: Factory calibrated, field calibrated monthly, zeroed daily according to manufacturers instructions.

<u>Miscellaneous Measuring Devices</u>- Calibration procedures for any other measuring device used will be documented at the request of the regulatory authority.

All equipment will be checked daily and replaced as necessary. Instrument manuals and an instrument log book will accompany all equipment into the field. Any calibration, repairs or related information will be recorded in the log book.

#### 3.0 SOIL SAMPLING METHODOLOGY

3.1 Soil Sampling and Drilling Equipment Decontamination

All equipment used for drilling and sampling during USPCI environmental assessments will be decontaminated using a steam cleaner or high pressure washer prior to use. In addition, the equipment will be decontaminated subsequent to completion of each boring/monitoring well. All equipment used for collection of more than one sample, such as split-spoon soil samplers, will be decontaminated between each use to prevent cross contamination between samples. The sampling equipment decontamination procedure for pesticides and organic analysis will consist of a low phosphate detergent (Alconox or equivalent) bath followed by tap water, and deionized water rinses. Nylon brushes will be used to scrub sediment from the

equipment. If the equipment is used to sample for metals, the initial rinse will be conducted with 0.1 N nitric acid followed by tap water and deionized water. Clean equipment will be placed on a rack and allowed to air dry. Bath and rinse water will be replaced as necessary to ensure adequate cleaning. The water used in the decontamination procedure will be stored in containers certified for hazardous materials storage by U.S. Department of Transportation (DOT). The drums will be secured on-site.

#### 3.2 Soil Sample Collection During Drilling Activities

A proposal will be submitted to the lead Regulatory Authority with proposed boring/sampling locations. The exact location and number of borings at each site will be determined in the field by the Project Geologist/Engineer.

Drilling will be conducted by a State Certified Well Driller under the supervision of the USPCI field representative. Soil borings will be advanced with a continuous flight, hollow stem auger drill rig. No petroleum hydrocarbon based lubricants will be allowed on the drill string and associated connections. Soil cores will be collected with either a split-spoon sampler or a continuous coring device.

#### 3.2.1 Split-Spoon Sampler

A California modified split-spoon sampler consists of a thin-walled steel cylinder, held together on each end by threaded steel end pieces, which separates longitudinally into two halves allowing the removal of brass or stainless steel liners which are used to contain the sampled soil interval. The sampler is 18 to 24 inches long and typically contains 3 to 4, six inch long, 2 to 2.5 inch diameter liners. The sampler will be driven ahead of the hollow stem auger by a 140 pound hammer with a 30 inch drop in accordance with the American Society for Testing and Materials (ASTM) Methods D 1586-84 for split barrel sampling of soil and D 1587-83 for thin-walled tube sampling of soils. The blows required to drive the sampler each six inch interval will be recorded on the boring log. The sampler will be removed from the boring and opened to reveal the liners. Latex gloves will be worn to prevent cross-contamination with other samples. The disposable gloves will be discarded after collection of samples from each sample drive.

Whenever possible, the bottom liner will be selected for laboratory analysis. The liner will be sealed on each end with aluminum foil, plastic end caps and duct tape. Samples selected for laboratory analysis will be preserved, stored and transported in accordance with USPCI sample processing protocol (see Section 6).

Soil in the other liners and sampler shoe will be described by the USPCI field representative according to ASTM Standard Practice for Description and Identification of Soils, Visual-Manual Procedure (ASTM D-2488-90). Stratigraphic,

genetic and other data/interpretations will also be recorded. Alternatively, one of the other sample liners may be used for the preparation of a duplicate sample. Field observations and selected sample intervals for laboratory analysis will be noted on the log prepared for each soil boring/ monitoring well. An explanation of the ASTM soil classification system will be included with the soil boring/well logs in an appendix of the assessment report.

#### 3.2.2 Continuous Coring Devices

A variety of continuous coring devices may be employed for core collection. During coring operations, samples selected for laboratory analysis will be contained in glass jars and processed in accordance with the above mentioned USPCI sample processing protocol.

At a minimum, soil samples will be collected at five foot intervals, at significant changes in lithology and at intervals of obvious contamination in order to develop a complete profile of soil contamination.

#### 3.3 Soil Sampling by Hand Auger and Coring Hammer

Hand tools will be utilized to collect soil samples from areas which are inaccessible to drilling rigs or do not require one. A hand auger will be used to advance the soil boring to the interval of interest. A hand held sliding hammer soil coring device will be utilized to drive a steel liner to obtain a undisturbed sample. Latex gloves will be worn to prevent cross-contamination with other samples. The disposable gloves will be discarded after sample collection from each interval. The steel liner containing the collected sample will be sealed on each end with aluminum foil, plastic end caps and duct tape. Samples selected for laboratory analysis will be preserved, stored and transported in accordance with USPCI sample processing protocol (See Section 6).

Soil description and sample collection intervals will follow methods discussed in Section 3.2.

#### 3.4 Sampling from Soil Piles or Shallow Soil Pits

Soil samples will be collected and transported from excavated material or shallow pits in the manner described in the previous section except that a backhoe will not be utilized. If composite samples are collected, four metal liners (brass or stainless steel) will be filled for every 50 cubic yards of material to be sampled unless otherwise specified by the regulatory agency. The samples will be composited in a

State Certified laboratory prior to analysis.

#### 3.5 Sample Collection During Underground Tank Removal

Soil samples will be collected as soon as possible after removal of the tank. Where feasible, all preparations for soil sampling will be made prior to tank removal. Soil samples collected from a backhoe bucket or directly from the excavation floor will be collected in thin-walled stainless steel or brass liners at least three inches long by one inch in diameter. From 3 to 24 inches of soil will be removed from the immediate surface area where the sample is to be taken and the cylinder then pounded into the soil with a wooden mallet, bulk density driver, or other decontaminated driving device. No head space will be present in the cylinder once the sample is collected. Care will be taken to avoid contamination of both the inside and outside of the cylinder as well as its contents. During sampling, latex gloves will be worn to prevent cross contamination with other samples. The disposable gloves will be discarded after collection of each sample.

Once the sample is collected, the liner will be sealed on each end with aluminum foil or teflon tape, polyethylene lids, and duct tape. The sample will be stored and transported to the laboratory in accordance with USPCI Sample Processing Protocol (Section 6).

#### 4.0 SOIL BORING ABANDONMENT AND DRILL CUTTINGS DISPOSAL

#### 4.1 Soil Boring Abandonment

Upon completion of sampling activities, all USPCI soil borings will be abandoned with neat cement in order to prevent development of any preferential pathways from the surface to subsurface. The neat cement shall be composed of one sack of Portland cement (94 pounds or 43 kilograms) to 4.5 to 6.5 (depending on cement type and additives used) gallons (17 to 25 liters) of clean water. The borings will be backfilled in one continuous operation from the bottom up either through the drilling augers or via tremie pipe.

#### 4.2 Disposal of Drill Cuttings

All soil cuttings generated during drilling activities will be contained in DOT approved, labeled steel drums certified for the storage of hazardous materials. The drums will be secured on-site.

## 5.0 GROUNDWATER MONITORING WELLS/ INSTALLATION, DEVELOPMENT, SURVEY, MONITORING, AND SAMPLING

#### 5.1 Monitoring Well Installation

If a soil boring is converted to a groundwater monitoring well, all well screen and casing, centralizers and casing handling equipment will be decontaminated with a steam cleaner or high pressure, hot water washer utilizing potable water immediately prior to installation. Well construction material decontamination will be conducted on impermeable surfaces and all decontamination effluent will be contained and transferred to DOT approved plastic or steel drums. The drums will be secured on-site.

Well casing will be selected based on the chemical compounds targeted for laboratory analysis, anticipated lifetime of the monitoring program, well depth and geochemistry. In most cases, polyvinyl chloride (PVC) well casing and screen will be utilized. Site specific conditions may, in some cases, require the use of other well construction materials. The casing/screen will be flush threaded. Unless site-specific conditions warrant otherwise, 0.020 inch slotted screen will be installed. All appropriate measures will be taken to ensure that the well casing is centrally located in the boring. The screened interval will extend up to 15 feet below the water table. Five feet of screen will extend above the saturated zone in unconfined conditions in order to allow for monitoring of free product under conditions of a rising water table. Screened intervals completed in confined aquifers will not extend above the saturated zone. In order to prevent potential dilution of target chemical compounds in water samples, no more than 20 feet of screen will be installed in any monitoring well.

A coarse-grained sand filter pack (e.g. #2/12 Lonestar, #3 Monterey) will usually be utilized to mitigate siltation of the well by fine-grained sediments in the surrounding aquifer (grain size of the filter pack will be of appropriate size to ensure hydraulic connection between the well bore and the adjacent water-bearing formation). The sand will be introduced through the drilling augers in order to ensure the integrity of the filter pack. A minimum 3 inch differential between the outer diameter of the well screen and the inner diameter of the augers will be maintained in order to ensure effective placement of filter pack. In some instances, saturated fine-grained sand (flowing sand) may enter the drill string during well completion. Although every effort will be made to prevent entry of native materials into the drill string

during well completion (e.g. loading the augers with water), it may sometimes be necessary to utilize native material for filter pack. Information regarding filter pack condition will be included on the well log. The filter pack will extend to at least one foot but no more than two feet above the top of the screened interval to allow for filter pack settling during well development.

Subsequent to introduction of the filter pack, the surface sanitary seal will be completed. At least a two foot thick interval of sodium bentonite pellets will be deposited directly above the filter pack. The pellets will then be hydrated with potable water. A neat cement grout seal will be placed via tremie pipe from the bentonite pellet seal to just below the frost line. The neat cement grout seal will be composed of one sack of Portland cement (94 pounds or 43 kilograms) to 4.5 to 6.5 (depending on cement type and additives used) gallons (17 to 25 liters) of clean water.

Soundings will be made by the USPCI field representative during all stages of well construction to ensure proper placement of filter pack and sealant materials. Moreover, the volume of filter pack and sealant required will be calculated to establish the correct subsurface distribution of the materials. The actual volume of materials used will be recorded during well construction. Discrepancies between calculated volumes and actual volumes be noted and explained on the monitoring well construction log.

A subgrade traffic-rated well box, or aboveground steel casing imbedded in concrete will be installed to protect the wellhead. The concrete cap will extend from below the frost line to the surface and blend into a four-inch thick apron at least two feet in diameter. The annulus between the well casing and the steel casing will be filled with bentonite pellets or chips from below the frost line to the surface. The bentonite sealant material will then be hydrated with potable water. This non-bonding surface seal will serve to protect the well casing from damage during periods of frost heaving. The wellhead will be locked to provide monitoring well security.

A typical monitoring well completion is diagramed in Figure A. All well completion information will be included in the well log.

## 5.2 Well Development Protocol

Groundwater monitoring wells will be surged and developed subsequent to well completion. Flow reversals or surges will be created by using surge blocks, bailers or pumps. Formation water will be used to surge the well. In low yielding water bearing formations, an outside source of water may be introduced into the well to facilitate development. In such cases this water will be chemically analyzed

will

beforehand to evaluate its potential impact on in-situ water quality. At no time will air be used to develop a well. Approximately 4 to 10 times the volume of water in the casing and pores of the filter pack will be withdrawn, if possible. Development volumes will be calculated in the following manner:

Volume of Schedule 40 PVC Pipe

Diameter (inches)	I.D. (inches)	Volume Gal/linear ft.
2	2.067	0.17
4	4.026	0.66

## Volume of Open Borehole and Annular Space Between Casing and Hole

Hole Diameter (inches)	Volum ft. of	e/linear hole	Normal Casing Diameter		linear ft. of* lar Space
(======)	Gal.	Cu. Ft.	(inches)	Gal.	Cu. ft.
7.25	2.14	0.29	2	1.91	0.26
8.25	2.78	0.37	2	2.55	0.34
10.25	4.29	0.57	2	4.06	0.54
10.25	4.29	0.57	4	3.46	0.46
12.25	6.13	0.82	4	5.30	0.71

\*Note: Annular volumes will be multiplied by 30% to account for porosity of filter pack.

If the aquifer is slow to recharge, development will continue until recharge is too slow to practically continue. The volume of water produced versus time will be recorded on the well log.

All withdrawn groundwater will be stored on-site in DOT approved containers for hazardous material storage unless prior permission is granted by the appropriate regulatory agency to discharge the water to the ground surface or sanitary sewer. Contained water will be labeled with the source of the water to help ensure appropriate disposal based on contamination levels.

## 5.3 Elevation Survey of Monitoring Wells

All monitoring wells at USPCI project sites will be surveyed to a common datum by a qualified surveyor. Where required by regulatory agencies, the wells will be surveyed to mean sea level datum (MSLD) by a Registered land surveyor to an accuracy of 0.01 foot. The surveyor's report will be included as an appendix to the report. For consistency, the wells will be surveyed from the north side of the top of the monitoring well casing.

## 5.4 Documentation of Well Design, Construction and Development

The following well design and construction details for each monitoring well will be included on the boring log, well construction log, purge log, or surveyor' report:

- 1) Date/time of construction
- 2) Drilling method and drilling fluid used
- 3) Well location (within 0.5 ft.)
- 4) Bore hole diameter and well casing diameter
- 5) Well depth (within 0.1 ft.)
- 6) Drilling and lithologic logs
- 7) Casing materials
- 8) Screen materials and design
- 9) Casing and screen joint type
- 10) Screen slot size /length
- 11) Filter pack material/size
- 12) Filter pack volume calculations
- 13) Filter pack placement method
- 14) Sealant materials (percent bentonite)
- 15) Sealant volume (lbs/gallon of cement)
- 16) Sealant placement method
- 17) Surface seal design/construction
- 18) Well development procedure

19) Type of protective well cap

20) Ground surface elevation (within 0.01 ft.)

21) Top of monitoring well casing elevation (within 0.01 ft.)

22) Detailed drawing of well (including dimensions)

## 5.5 Groundwater Monitoring Protocol

During a sampling event the depth to standing water and total depth of the well (bottom of screened interval) well be measured to an accuracy of 0.01 foot. For consistency, all measurements will be taken from the north side of the wellhead at the survey mark. These measurements are required to calculate the volume of stagnant water in the well and provide a check of the integrity of the well (e.g., identify siltation problems). The devices used to detect the water level surface and calibration methods have been discussed previously (Section 2.2).

To reduce the potential for cross contamination between wells, well monitorings will proceed in order from the least to most contaminated wells, if known. Wells containing free product will be monitored last. Between each well monitoring the equipment will be decontaminated following the procedure detailed in Section 3.1.

Water elevations will be collected during each subsequent sampling event in order to determine if horizontal and vertical flow gradients have changed since initial site characterization. A change in hydrologic conditions may necessitate modification to the design of the site groundwater monitoring system.

## 5.5.1 Detection of Immiscible Layers

The thickness of immiscible layers (i.e., "floaters" and/or "sinkers") within a monitoring well, if present, will be determined during each sampling event. "Floaters" are those relatively insoluble organic liquids that are less dense than water and which spread across the potentiometric surface. "Sinkers" are those relatively insoluble organic liquids that are more dense than water and tend to migrate vertically through sand and gravel aquifers to the underlying confining layer.

The following procedures will be utilized for detecting the presence of light and/or dense phase immiscible organic layers. These procedures will be conducted prior to well evacuation for conventional sampling:

- 1) Remove the locking and protective well caps.
- 2) Sample the air in the wellhead for organic vapors using either a photoionization analyzer or an organic vapor analyzer, and record

measurements. The air above the wellhead will be monitored in order to determine the potential for fire, explosion, and/or toxic effects on workers.

- 3) Determine, using an interface probe, the static liquid level and thickness, if present, of any floating immiscible organic layers.
- 4) Determine the presence of dense phase immiscible layers by lowering an interface probe to the bottom of the well.

## 5.5.2 Collection of immiscible solutions

The approach to collecting light phase immiscibles is dependent upon the depth to the surface of the floating layer and the thickness of that layer. If the thickness of the phase is 2 feet or greater, a bottom valve bailer will be used. The bailer will be lowered slowly until contact is made with the surface of the immiscible phase, then lowered to a depth less than that of the immiscible/water interface depth as determined by preliminary measure with the interface probe.

A double check valve bailer will be used to collect dense phase immiscibles. The bailer will be slowly lowered and raised for sample collection.

Floating product thickness is calculated by subtracting the depth to product from the depth to water. In addition, water elevations are adjusted for the presence of fuel with the following calculation:

(Product Thickness) (.8)+(Water Elevation)
= Corrected Water Elevation

Note: The factor of 0.8 accounts for the density difference between water and petroleum hydrocarbons.

Newly installed wells will be allowed to stabilize for 24 hours after development prior to free product inspection.

A transparent surface sampler will be used for visual inspection of the groundwater in order to note sheens (difficult to detect with an Interface Probe), odors, microbial action and sediments.

If free product is detected, laboratory analysis of groundwater at the interface for dissolved product will not be conducted. A product sample will be collected for source identification.

## 5.6 Groundwater Sampling Protocol

## 5.6.1 Sampling Equipment Decontamination

Prior to arriving at the sampling site, all groundwater sampling equipment except pre-cleaned disposable materials, and laboratory prepared sampling containers will be washed with a low phosphate detergent (Alconox or equivalent), rinsed twice with tap water, and once with deionized water. If more than one monitoring well is on-site, this procedure will be carried out prior to sampling of each of the other monitoring wells.

Latex gloves will be worn at each sampling location to prevent cross-contamination with other samples. The disposable gloves will be discarded after collection of samples from each well.

## 5.6.2 Well Purging

The water standing in a well prior to sample collection may not be representative of in-situ ground-water quality. Prior to sample collection, the well will be purged with a bailer, WaTerra pump, or positive-gas-displacement pump until indicator parameters (temperature, conductivity and pH) stabilize. This generally requires the removal of at least three well casing volumes by bailing or pumping. The water will be drawn from the uppermost part of the water column in high-yield formations to ensure that fresh water from the formation will move upward in the screen. In low-yield formations, water will be purged so that it is removed from the bottom of the screened interval.

The criteria for determining well casing volumes and disposition of purged water is outlined in Section 5.3 (Well Development Protocol). The indicator parameter measurements will be taken both before and after purging of each well casing volume. Once indicator parameters have stabilized, a sample will be collected after the water level approaches 80 percent of its initial elevation. Where water level recovery is slow (exceeding 2 hours), the sample will be collected after stabilization is achieved and enough water is present to collect an adequate amount of sample for analysis. At no time will a well be pumped dry if the recharge rate causes the formation water to vigorously cascade down the sides of the screen and cause an accelerated loss of volatiles. All well development and purging information will be noted on purge logs and included as an appendix of the report.

## 5.6.4 Sample Collection

Observations made during to groundwater sample collection will include a description of the area surrounding the well, possible impacts by surface-water runoff, ambient weather conditions and other factors which could affect the final data analysis. This

documentation will be recorded on the well purge log.

Sampling will proceed from the least contaminated to the most contaminated well, if that information is available before sample collection, or if such information can be determined by field evidence. Where several types of analysis will be performed for a given well, individual samples will be collected in order of decreasing volatility as follows:

- 1. Volatile organics
- 2. Purgeable organic carbon
- 3. Purgeable organic halogens
- 4. Total organic carbon
- 5. Total organic halogens
- 6. Extractable organics
- 7. Total metals
- 8. Dissolved metals
- 9. Phenols
- 10. Cyanide
- 11. Sulfate and chloride
- 12. Turbidity
- 13. Nitrate and ammonia

The specific analytical methods to be utilized for the various analyses are shown on Table 2.

All sampling procedures will conform with the following:

- 1) Water samples will be collected with a teflon bailer equipped with a bottom emptying device, a WaTerra pump, or a positive gas displacement bladder pump.
- 2) All sampling equipment introduced to the well will be constructed of inert materials (i.e. teflon or stainless steel).
- 3) Positive gas displacement bladder pumps will be operated in a continuous manner so that they do not produce pulsating samples that are aerated in the return tube or upon discharge.

- 4) Check valves will be designed and inspected to assure that fouling problems do not reduce delivery capabilities or result in aeration of the sample.
- 5) Sampling equipment (e.g., especially bailers) will never be dropped into the well, which causes degassing of the water upon impact. When using a bailer, the device will be lowered slowly into the well to mitigate agitation (causing turbidity) of the water. The bailer will be retrieved gently but quickly and the sample then poured slowly into the sample container in order to minimize agitation, turbulence and exposure to air.
- 6) Clean sampling equipment will not be placed directly on the ground or other contaminated surfaces prior to insertion into the well.
- 7) Duplicate samples will be transferred to vials or containers that meet Regulatory specifications (Table 1). When filling 40 ml vials, groundwater will be transferred from the sampling device to the sample container by allowing the fluid to flow slowly along the sides of the vessel. All containers will be filled above the top of the opening to form a positive meniscus. No head space should be present in the sample container once it is sealed. After the vial is capped, it will be inverted to check for air bubbles. If bubbles are present, the sample will be discarded and replaced. If it is not possible to collect a sample without head space, the problem will be noted on the field technician's sampling log.
- 8) Immediately following sample collection, sample containers will be stored and transferred to the laboratory in accordance with USPCI sample processing protocol (Section 6).
  - 9) If a positive gas displacement bladder pump is used for sample collection, pumping rates will not exceed 100 milliliters/minute. Higher rates can increase the loss of volatile constituents and can cause fluctuation in pH and pH-sensitive analytes. Once the portions of the sample reserved for the analysis of volatile components have been collected, higher pumping rates may be utilized for sample collection for other analyses. However, the sampling flow rate will not exceed the flow rate used while purging.

#### 6.0 SAMPLE PROCESSING

## 6.1 Sample Containers

Soil and Groundwater samples will be placed in the proper containers for the desired analysis. Table 1 summarizes the required sample containers.

All sample containers will be verified clean in the laboratory prior to

shipment to a sampling site.

## 6.2 Sample Preservation

Samples will be preserved in order to: 1) retard biological activity, 2) retard hydrolysis, and 3) reduce sorption effects. Soil and groundwater samples will be preserved as indicated on Table 1 and placed in an ice chest immediately after collection. Chemical ice (blue ice), dry ice, or, where allowed, regular ice, sealed in plastic bags will be used to cool and maintain samples at a temperature of 4°C.

Samples requiring analysis for organics will not be filtered. Samples will not be transferred from one container to another which could result in loss of organic material onto the walls of the container or through aeration.

Metallic ions that migrate through the unsaturated (vadose) and saturated zones and arrive at a ground-water monitoring well may be present in the well. Particles (e.g., silt, clay), which may be present in the well even after well evacuation procedures, may absorb or adsorb various ionic species to effectively lower the dissolved metal content in the well water. Ground-water samples on which metals analysis will be conducted will be split into two portions. One portion will be filtered through a 0.45 micron membrane filter, transferred to a bottle, preserved with nitric acid to a pH less than 2 (Table 1), and analyzed for dissolved metals. Dissolved metals content is utilized to determine hydrochemical conditions. The remaining portion will be transferred to a bottle, preserved with nitric acid, and analyzed for total metals. Total metals content is used to determine worst case contaminant concentrations. Any difference in concentration between the total and dissolved fractions may be attributed to the original metallic ion content of the particles and any sorption of ions to the particles. Disposable filters will be dedicated to individual wells to prevent cross-contamination of other samples.

## 6.3 Sample Labeling

Each sample container will be labeled to prevent misidentification. The label will contain at least the following information:

- O Sample number which uniquely identifies the sample
- O Project title or number
- Location of sample collection
- Soil boring or well number, as applicable
- Name of collector
- O Date and time of collection
- O Type of analysis requested.

Parameter	Matrix	Holding Container	Time	Preservation
Total Petrol.	Soil	3" stainless steel	14 days¹/	4°C
Hydrocarbons		or brass cylinder	40 days <sup>2</sup> /	
(Light Fractions)	Water	40 ml glass vial, teflon-faced silicon septum	14 days <sup>1</sup> / 20 days <sup>2</sup> /	4°C HC1 to pH * 2 (except CaCO,
water)				
(Heavy fractions)	Water	<pre>1 amber bottles, teflon seal/silicon septum</pre>	14 days <sup>1</sup> / 40 days <sup>2</sup> /	4°C
Benzene Toluene	Soil	3" stainless steel or brass cylinder	14 days¹/	4 • C
Xylene Ethylbenzene	Water	40 ml glass vial,	7 days1/	4°C HC1
-		teflon seal/silicon septum	14 days <sup>3</sup> /	to pH * 2 (except CaCO;
water)			14	44.0
Purgeable Halocarbons	Soil	3" stainless steel or brass cylinder	14 days¹/	4°C
	Water	500 ml glass vial, teflon seal/silicon septum	14 days¹/	
Organic lead	Soil	3" stainless steel or brass cylinder	14 days1/	4°C
	Water	40 ml glass vial, teflon seal/silicon septum	14 days¹/	4°C
Ethylene Dibromide	Soil	<pre>3" stainless steel or brass cylinder</pre>	14 days²/	40C
	Water	40 ml glass vial, teflon faced silicon septum	14 days¹/	4°C
Polynuclear Aromatic Hydrocarbons	Soil	8 oz. wide mouth glass with teflon seal	14 days¹/ 40 days²/	4 <b>°</b> C
	Water	1000 m. amber glass with teflon seal	7 days¹/ 40 days²/	4°C

		Holdin		. •
Parameter	Matrix	Container	Time	Preservation
Poly- Chlorinated Biphenyls	Soil	8 oz. wide mouth glass with teflon seal	7 days¹/ 40 days²/	4°C
	Water	1000 ml amber glass with teflon seal	7 days¹/ 40 days²/	4°C
Total Metals	Soil	3" stainless steel or brass cylinder	6 months	
	Water	1000 ml plastic	6 months	pH<2
Dissolved Metals	Water	1000 ml plastic	6 months	pH<2 0.45 micron filtration
Pesticides	Soil	3" stainless steel or brass cylinder	14 days²/	4°C
	Water	1000 ml glass	7 days²/	4 <b>0</b> C

Note:

- / Maximum holding time for sample (extract within this time or analyze if extraction is not required).
- $^2/$  Maximum holding time for extract (analyze within this time).
- 3/ Maximum holding time for sample when pH adjusted with HCl.

## 6.4 Chain-of-Custody Record and Sample Analysis Request Form

A chain-of-custody record for each container or sample will be used to track possession of the samples from collection in the field until arrival at the laboratory.

The chain-of-custody record will contain the following information:

- 1. Site name
- 2. Signature of collector
- 3. Date and time of collection
- 4. Sample identification number(s)
- 5. Number of containers in sample set
- Description of sample and container(s)
- 7. Name and signature of persons, and the companies or agencies they represent, who are involved in the chain of possession
- 8. Inclusive dates and times of possession
- 9. Requested analysis for each sample

## 6.5 Delivery of Samples to Laboratory

Samples will be delivered to the laboratory within 48 hours when possible. Delivered samples will be accompanied by a chain-of-custody record. The laboratory shall note sample condition on the chain-of-custody (e.g. chilled, presence or absence of head space) upon arrival. Samples will be transported either by USPCI personnel or by private carrier. Analytical holding times will be considered in determining sampling and shipping schedules. Friday shipment/ Saturday laboratory receipt of samples will be coordinated in advance with the laboratory.

## 6.6 Quality Control Field Samples

A QC program independent from the laboratory's program will be instituted. The program entails "blind" submittals to the laboratory of blank and duplicate samples. No spiked samples will be supplied from the field for these investigations. All QC samples will be assigned independent sample numbers and made indistinguishable from non QC samples.

When submitting groundwater samples, travel blanks will be used to detect the introduction of contaminants during sample handling or transportation from the field to the laboratory. The travel blanks, provided by the analytical laboratory, will remain sealed in the field and accompany the collected groundwater samples to the laboratory for analysis. The blanks will consist of deionized, analytically confirmed organic-free water. The blanks will be numbered, packaged, and sealed in the same manner as the other samples. Each blank will carry the appropriate preservative for the analytes of

concern. A minimum of one trip blank per sampling event will be collected.

Field or Equipment blanks will be collected in order to detect introduction (if any) of cross-contamination into environmental matrices from nondedicated sampling devices that have been cleaned in the laboratory or field. Laboratory prepared analyte-free water, brought to the field in sealed containers, will be poured over decontaminated sampling equipment, collected in basins and transferred to appropriate sample jars for shipment to the laboratory. Each Equipment blank will carry the appropriate preservative for the analyte of concern. These blanks will be numbered, packaged, and sealed in the same manner as the groundwater samples. A minimum of one equipment blank will be processed during each day of well sampling activity.

Analytical results for travel and equipment blanks will not be used to correct groundwater data. If contaminants are found in the blanks, the source of the contamination will be identified and corrective action, including resampling, will be initiated.

Ten percent of groundwater samples submitted to the laboratory for analysis will be duplicates. Water sample duplicates will be collected by filling two sample bottles from the one bailer volume. If more than one bailer volume is required, each bailer volume will be split between containers.

## 6.7 Laboratory QA/QC Plan

Soil and groundwater samples will be submitted to a State Certified Hazardous Waste Laboratory for chemical analysis of hazardous constituents. Established QA/QC procedures for analytical laboratory operations will include sample custody procedures, standards of analytical accuracy, analysis of matrix spikes and method blanks, data reduction, verification of raw analytical data, and maintenance of control charts to monitor analytical performance. These QA/QC procedures are outlined in the laboratory QA/QC Plan which is available upon request. Chemical analyses will be performed in accordance with standard procedures established by the United States Environmental Protection Agency (EPA) in "Guidelines Establishing Test Procedures for the Analysis of Pollutants Under the Clean Water Act" (40 CFR Part 136, October 1984). Analytical laboratories are periodically evaluated through external performance audits conducted by EPA and State agencies through government QC labs. The specific analytical methods to be utilized for purgeable and semivolatile hydrocarbons analyses are shown on Table 2.

Provided the data base is of sufficient size, statistical techniques may be employed for data validation.

# Table 2 Laboratory Test Methodology Underground Tank Sites

Hydrocarbon Leak	Soil Anal	ysis	Water An	nalysis
Unknown Fuel	TPH G	GCFID(5030)	TPH G	GCFID(5030)
Olikilowii Fuei	TPH D	GCFID(3550)	TPH D	GCFID(3510)
	BTX&E	8020 or 8240	BTX&E	602 or 624
	DIAGE	0020 01 02 10		
Leaded Gas	TPH G	GCFID(5030)	TPH G	GCFID(5030)
	BTX&E	8020 or 8240	BTX&E	602 or 624
	Option	al	TEL	DHS-LUFT
	TEL	DHS-LUFT	EDB	DHS-AB1803
	EDB	DHS-AB1803		
III.I. adad Caa	TPH G	GCFID(5030)	TPH G	GCFID(5030)
Unleaded Gas	BTX&E	8020 or 8240	BTX&E	602 or 624
	DIVOE	8020 OF 8240	DIXAL	002 0, 02
Diesel TPH D	GCFID(3	550)	TPH D	GCFID(3510)
Diesei IIII D	BTX&E	8020 or 8240	BTX&E	602 or 624
Jet Fuel	TPH D	GCFID(3550)	TPH D	GCFID(3510)
001.00.	BTX&E	8020 or 8240	BTX&E	602 OR 624
Kerosene	TPH D	GCFID(3550)	TPH D	GCFID(3510)
	BTX&E	8020 or 8240	BTX&E	602 or 624
Fuel Oil	TPH D	GCFID(3550)	TPH D	GCFID(3510)
rue: Oil	BTX&E	8020 or 8240	BTX&E	602 or 624
	BIAGE	0020 5. 02.10		
Chlorinated Solvents	CL HC	8010 or 8240	CL HC	601 or 624
Gindinated Dollaria	BTX&E	8020 or 8240	BTX&E	602 or 624
				0.0510 (0540)
Non Chlorinated Solvents	TPH D	GCFID(3550)	TPH D	GCFID(3510)
	BTX&E	8020 or 8240	BTX&E	602 or 624
Waste Oil or Unknown	TPH G	GCFID(5030)	TPH G	GCFID(5030)
waste on or orknown	TPH D	GCFID(3550)	TPH D	GCFID(3510)
	0 & G	503D&E	O & G	503A&E
	BTX&E	8020 or 8240	BTX&E	602 or 624
	CL HC	8010 or 8240	CL HC	601 or 624
	CL HC	5010 or 6240	CETTO	001 0. 02 .
	ICAP or	AA to Detect Metals:	Cd, Cr, Pb, Zi	n
	Method	8270 for Soil or Wate	er to Detect:	
	PCB*		PCB*	
	PCP*		PCP*	
	PNA		PNA	
		o Crenente	111/5	
	Creosot	e Creosote		

<sup>\*</sup> If found, analyze for dibenzofurans (PCBs) or dioxins (PCP)

us	PCI Project N	lame:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
Me	asuring Point	(MP)	Top of casing	-		Well No.	OKUS-W1
We	ell Depth: (Be	olow MP):	22.05	Feet			
Ca	sing diameter		2	Inches	<u></u>	Sampling Date:	11/11/93
De	pth To Groun	d Water (Below N	MP): 9.24	Feet		Sample ID No.	OKUS-W1
Me	thod Of Well	Development			Time:	1150	
L)	·	[ ] Submersible P		[X] Inertia Pump		Riser Elevation (MP):	9.17
	Bailer	[] Centrifugal Pu	imp	[] Other	Top of Screen Elevation:	6.85	Feet
1	mpling Collec		· · · · · · · · · · · · · · · · · · ·		Sample Appearance:	turbid	<del>.</del>
	Тар	[ ] Submersible P		[ ] Inertia Pump:	Odor:	modorate petroleum odo	<u> </u>
[X]	Bailer	Tγpe:	[] Teflon	[] Steinless Steel	Sampling Problems (if any	):	
┛	<del></del>	[X] HDPE Plastic		[] PVC	[X] Disposable		
		Bailer Set At:	3.0	Feet Below MP	Decontamination Performs	ed:	
Tu	bing Type (if	Used):	HDPE				
Tul	bing Used for	r:	[]SampleCollection	[X] Well Developme	nt/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1,
_				<u> </u>		T	As, Pb, 8010
			Temperature			Cumulative Volume	Pumping Rate
	Time	рН	Corrected	Temperature	Water Level	of Water	in
		(Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
-	4450	7.0	(umho/cm)	20.5		(Gallons) 5.0	(GPM)
	1158 1201	7.3 7.2	900 1200	20.5 21.0		10.0	
4-	1207	7.2	1300	20.6		15.0	
_ saı	mples collecte	<del></del>					
						<u> </u>	
	<del> </del>	<u> </u>	-				
1—	***					<del></del>	
				<u> </u>	<del>                                     </del>		
	<del> </del>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	<u> </u>	
At	Least	3	Well Bore Volumes We	re Purged Before Samp	I Discharge Rate =	GPM x 0.00223 =	cfs
_	mments:						
			<u></u>			<u></u>	
<u>.</u>					MCanagad Pro	Ara Mardirosain	
Fo	rm Complete	d By:	Don Ostrand		Witnessed Bγ:	Ara Warun Obalii	

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JSPCI Project N	lame:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
Measuring Point		Top of casing			Well No.	OKUS-W2
Vell Depth: (Be		22.50	Feet			
asing diameter		2	inches		Sampling Date:	11/11/93
٠	nd Water (Below I	MP): 10.20	Feet		Sample ID No.	OKUS-W2
epth To Produ	ct (Below MP):	N/A				
Method Of Well	Development			Time	: 1048	
] Tap	[ ] Submersible I	Pump	[X] Inertia Pump		Riser Elevation (MP):	9.71
] Bailer	[] Centrifugal Po	ımp	[] Other	Top of Screen Elevation:	7.05	Feet
ampling Collec	tion Method:			Sample Appearance:	slightly turbid	
] Tap	[] Submersible [	Pump	[] Inertia Pump:	Odor:	modorate petroleum odo	<u>r</u>
X] Bailer	Type:	[] Teflon	[] Stainless Steel	Sampling Problems (if any	n:	
	[X] HDPE Plastic	;	[] PVC	[X] Disposable		
ump Intake Or		3.0	Feet Below MP	Decontamination Perform	ed:	
Tubing Type (if		HDPE				
		[]SampleCollection	[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1,
Tubing Used for	1-	r Togrupia-consection	ive trail retainbille			As, Pb, 8010
	1	T	l		Cumulative Volume	Pumping Rate
		Temperature				in
Time	pН	Corrected	Temperature	Water Level	of Water Removed From Well	Gallons/Minut
	(Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	***************************************	(GPM)
		(umho/cm)		<u> </u>	(Gallons)	/GPIN)
1056	7.0	2000	21.1		10.0	
1102	7.0	2300	20.9		15.0	
1108	6.9	2500	20.9	<u> </u>	10.0	
amples collect	eu at iiii					
****					<u> </u>	
<del></del>						
						<u> </u>
						<b></b>
	<del></del>				<del> </del>	+
		-				
·	<u></u>	1	<u> </u>		<u> </u>	<u>L</u>
					0014 0 00000	ofo
At Least	3	Well Bore Volumes We		of Discharge Rate =	GPM x 0.00223 =	cfs
Comments:	15 gallons of w	ater removed during p	urging.			
		. <u> </u>	<del></del>			
					· ·	

JSPCI Project N	lame:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
		Top of casing			Well No.	OKUS-W3
Measuring Point		22.38	Feet		1	
Vell Depth: (Be		2	Inches		Sampling Date:	11/11/93
asing diameter	r: nd Water (Below I		Feet		Sample ID No.	OKUS-W3
-	ict (Below MP): 1		7000		,	
Method Of Well		3/A		Time	: 0951	
Tap	[] Submersible l	Pump	[X] Inertia Pump		Riser Elevation (MP):	9.80
) Bailer	[] Centrifugal Pu		[] Other	Top of Screen Elevation:	6.55	Feet
ampling Collec				Sample Appearance:	turbid	
	[] Submersible I	Dump	[ ] Inertia Pump:	Odor:	modorate petroleum odo	r
) Tap			[] Stainless Steel	Sampling Problems (if an	v):	
X] Bailer	Түре:	[] Teflon			<i></i>	
. <u> </u>	[X] HDPE Plastic		[] PVC	[X] Disposable		
ump Intake Or	Bailer Set At:	3.0	Feet Below MP	Decontamination Perform	led:	
ubing Type (if	Used):	HDPE		<u> </u>	<del></del>	· · ·
ubing Used fo	r;	[ ]SampleCollection	[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX.TPH/G/D/418.1,
				···		As, Pb, 8010
······································		Temperature			Cumulative Volume	Pumping Rate
Time	рН	Corrected	Temperature	Water Level	of Water	in
	(Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
		(umho/cm)			(Gallons)	(GPM)
1003	7.1	1800	22.8		5.0	
1009	7.0	1900	22.4		10.0	
1016	7.0	1900	22.8		15.0	
samples collect	ted at 1018		<u> </u>	<u> </u>		
			ļ			
······································			ļ			
. <u>.</u>		<u> </u>				
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	<del> </del>		1			
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	-		<del>                                     </del>			
·	<del></del>		<del>                                     </del>			
	1	1				
	2	Wall Bore Volumes W	ere Purged Before Sam	of Discharge Rate =	GPM x 0.00223 =	cfs
At Least	15 gallons of w	vater removed during				
Comments:	19 gallons of W	rater removed donning	8" '8'			
		<del></del>				

		UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
SPCI Project N	lame:		acinty Quarterly	14.010143	Well No.	OKUS-W4
leasuring Point		Top of casing			175,110.	
ell Depth: (Be	elow MP):	20.92	Feet		Sampling Date:	11/11/93
asing diameter		22	Inches		Sample ID No.	OKUS-W4
•	nd Water (Below N		Feet		Complete the control of the control	
	ct (Below MP): N	/A		Time:	1416	
	Development	·	[X] Inertia Pump		Riser Elevation (MP):	7.36
Тар	[] Submersible P		[] Other	Top of Screen Elevation:	6.08	Feet
Bailer		mp	[] 0.1.0.	Sample Appearance:	turbid	
ampling Collec	ction Method:		E 3 I will Burnet	Odor:	strong petroleum odor	
] Тар	[ ] Submersible F		[] Inertia Pump:			
() Bailer	Type:	[] Teflon	[] Stainless Steel	Sampling Problems (if any		
	[X] HDPE Plastic		[] PVC	[X] Disposable		
ump Intake O	Bailer Set At:	3.0	Feet Below MP	Decontamination Perform	ed:	
ubing Type (if	Used):	HDPE				
ubing Used fo		[ ]SampleCollection	[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1,
						As, Pb, 8010
· · · · · · · · · · · · · · · · · · ·		Temperature			Cumulative Volume	Pumping Rate
<b>T</b>	u	Corrected	Temperature	Water Level	of Water	in
Time	pH (Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
	(Omits)	(umho/cm)			(Gallons)	(GPM)
1422	7.2	2400	19.9		5.0	<u> </u>
1422	7.1	2500	19.8		10.0	
1428	7.1	2600	19.8		15.0	
samples collec		1				
amples conce						
	<del>                                     </del>					
					<del> </del>	
					<del> </del>	<del>                                     </del>
						<del> </del>
				_	<del>- </del>	
						<del>-</del>
				<del></del>		
At Least	3		Vere Purged Before San	npl Discharge Rate =	GPM x 0.00223 =	cfs
Comments:	15 gallons of v	vater removed during	har Auria.			
	<u> </u>					
		Don Ostrand		Witnessed By:	Ara Mardirosain	
Form Comple				**************************************		

USPCI Project N	ama:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
<u> </u>		Top of casing		- · · · · · · · · · · · · · · · · · · ·	Well No.	OKUS-W5
Measuring Point Well Depth: (Be		20.68	Feet		1	_ <del>_</del>
Casing diameter		2	Inches		Sampling Date:	11/11/93
	d Water (Below M		Feet		Sample ID No.	OKUS-W5
•	ct (Below MP): N					
Method Of Well				Time:	0821	
	[ ] Submersible P	ump	[X] Inertia Pump		Riser Elevation (MP):	9.25
	[] Centrifugal Pur		[] Other	Top of Screen Elevation:	5.95	Feet
Sampling Collect		····		Sample Appearance:	turbid	
	[ ] Submersible P	ump	[] Inertia Pump:	Odor:	petroleum odor	
	Type:	[] Teflon	[ ] Stainless Steel	Sampling Problems (if any	):	
	[X] HDPE Plastic		[] PVC	[X] Disposable		
Pump Intake Or		3.0	Feet Below MP	Decontamination Performe	ed:	
Tubing Type (if	Used):	HDPE				
Tubing Used for		[]SampleCollection	(X) Well Developme	nt/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1,
						Ав, Рь, 9010
		Temperature			Cumulative Volume	Pumping Rate
Time	рН	Corrected	Temperature	Water Level	of Water	in
	(Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
		(umho/cm)			(Gallons)	(GPM)
0829	7.3	4100	20.9	<del></del>	10.0	
0837 0849	7.5 7.4	3300 2900	20.6		15.0	
samples collecte		2300				
adilipico dollecti						
				<u>,</u>	<u> </u>	
		<u> </u>				
	<u></u>	N-1		-		
	-		<del> </del>			
		1	1	<u></u>	<u></u>	:el :::::::::::::::::::::::::::::::::::
At Least	3	Well Bore Volumes We	ere Purged Before Samp	l Discharge Rate =	GPM x 0.00223 =	cfs
Comments:				was removed from the wel	1.	
						<u></u>
·		<u> </u>		Witnessed Pro	Ara Mardirosain	
Form Completed	d By:	Don Ostrand		Witnessed By:	ALS INGLESSORS	

JSPCI Project N	ame:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
Measuring Point		Top of casing			Well No.	OKUS-W6
Veli Depth: (Be		16.30	Feet			. <u> </u>
Casing diameter		2	Inches		Sampling Date:	11/12/93
	d Water (Below N		Feet		Sample ID No.	OKUS-W6
•	ct (Below MP): N					
Method Of Well		· · · · · · · · · · · · · · · · · · ·		Time:	1010	
	[ ] Submersible P	ump	[X] Inertia Pump		Riser Elevation (MP):	7.29
1 Bailer	[] Centrifugal Pu		[] Other	Top of Screen Elevation:	2.29	Feet
Sampling Collec	<del></del>	•		Sample Appearance:	slightly turbid	
	[] Submersible P	ump	[] Inertia Pump:	Odor:	petroleum odor	
X) Bailer	Type:	[] Teflon	[] Stainless Steel	Sampling Problems (if any	n:	
	(X) HDPE Plastic		[] PVC	[X] Disposable		
Pump Intake Or		3.0	Feet Below MP	Decontamination Perform	ed:	
Tubing Type (if		HDPE				
Tubing Used for			[X] Well Developme	nt/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1,
Toma Cook lot	<del>-</del>		,			As, Pb, 8010
		Temperature			Cumulative Volume	Pumping Rate
Time	pH	Corrected	   Temperature	Water Level	of Water	in
) ime	(Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minut
	(0,1,10)	(umho/cm)	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		(Gallons)	(GPM)
1025	7.0	3100	22.7		5.0	
1031	6.9	3100	21.1		10.0	
1037	6.9	3000	21.9_		15.0	
amples collect	ed at 1042					<del>                                     </del>
<del></del>			<u> </u>			
			<del>                                     </del>		-	
			<u> </u>			
		<u> </u>	<u> </u>		<del> </del>	
		1				<del>-</del>
	<del>                                     </del>	<del>                                     </del>		<del> </del>	-	
			<del> </del>			
<del></del>	1	-				
	<u> </u>	<u> </u>	<u> </u>			
At Least	3		ere Purged Before Samp		GPM x 0.00223 =	cfs
Comments:	Petroleum produ	ect noted on the side	or the bailer when it	was removed from the we	D	
<del> </del>		<u> </u>				
	<del></del>	<u></u>				
					Ara Mardirosain	

JSPCI Project N	Name:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
Measuring Poin		Top of casing			Well No.	OKUS-W7
Well Depth: (Bo	·	20.50	Feet		1	
		2	Inches		Sampling Date:	11/12/93
asing diamete	r: nd Water (Below N		Feet	· · · · · · · · · · · · · · · · · · ·	Sample ID No.	OKUS-W7
•	ict (Below MP): N		1000			<u></u>
	l Development	<u> </u>		Time	: 0909	
<u>Леппоц от туек</u> ] Тар	[ ] Submersible F	Pump	[X] Inertia Pump		Riser Elevation (MP):	7.4
] Bailer	[ ] Centrifugal Pu		[] Other	Top of Screen Elevation:	2.4	Feet
Sampling Collec				Sample Appearance:	slightly turbid	_
		Ouron.	[] Inertia Pump:	Odor:	slight petroleum odor	······································
] Tap	[ ] Submersible F			Sampling Problems (if an		
X) Bailer	Type:	[] Teflon	[] Stainless Steel		y).	
	[X] HDPE Plastic		[] PVC	[X] Disposable		
<sup>2</sup> ump Intake Or	r Bailer Set At:	3.0	Feet Below MP	Decontamination Perform	ned:	
Tubing Type (if	Used):	HDPE				
Tubing Used fo	or:	[ ]SampleCollection	[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1.
						As, Pb, 8010
		Temperature			Cumulative Volume	Pumping Rate
Tires	pН	Corrected	Temperature	Water Level	of Water	in
Time	pri (Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
	(Gille)	(umho/cm)	(acitigrade)	,	(Gallons)	(GPM)
0922	7.5	2800	19.0	<del></del>	5.0	
0925	7.4	2800	19.5		10.0	
0927	7.4	2900	19.9		15.0	
samples collect		<u> </u>				
	1	1				<u> </u>
						<u> </u>
_						<del></del>
			<u> </u>			<del> </del>
			<u> </u>			<del> </del>
				<u> </u>	<u></u>	
At Least	3		ere Purged Before Samp	pl Discharge Rate =	GPM x 0.00223 =	cts
Comments:	15 gallons of w	ater removed during p	ourging.			
						<del></del>
				146	Ara Mardinassia	
Form Complete	ed By:	Don Ostrand		Witnessed By:	Ara Mardirosain	

JSPCi Project N	vame:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
Measuring Poin		Top of casing			Well No.	OKUS-W8
Well Depth: (B		15.30	Feet		1	
		2	Inches		Sampling Date:	11/11/93
Casing diamete	r: nd Water (Below N		Feet		Sample ID No.	OKUS-W8
-	ict (Below MP): N		1000		•	
	Development	<u> </u>		Time	: 1525	
] Tap	[ ] Submersible F	Pump	[X] Inertia Pump		Riser Elevation (MP):	7.11
] Bailer	[] Centrifugal Pu		[] Other	Top of Screen Elevation:	2.11	Feet
ampling Collec				Sample Appearance:	turbid	
	[] Submersible F	Duran	[] Inertia Pump:	Odor:	modorate petroleum odo	
] Tap				Sampling Problems (if any	· · · · · · · · · · · · · · · · · · ·	
X] Bailer	Туре:	[] Teflon	[ ] Stainless Steel		γι.	
	[X] HDPE Plastic		[] PVC	[X] Disposable	<u> </u>	
ump Intake O	r Bailer Set At:	3.0	Feet Below MP	Decontamination Perform	ed:	
ubing Type (if	Used):	HDPE				
ubing Used fo	r:	[ ]SampleCollection	[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX,TPH/G/D/418.1,
						As, Pb, 8010
		Temperature			Cumulative Volume	Pumping Rate
Time	pH	Corrected	Temperature	Water Level	of Water	in
IIIIPS	(Unite)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
	(0,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	(umho/cm)	,====		(Gallons)	(GPM)
1546	7.1	3700	20.8		5.0	
1552	7.2	3900	20.4		10.0	,
1558	7.2	4000	20.8		15.0	
amples collec	ted at 1617					
						<u> </u>
						<u> </u>
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			<del> </del>		<del> </del>	
			<del>                                      </del>		<del>                                     </del>	
<u> </u>	<u> </u>	<b></b>	<del> </del>		<del></del>	
	1		<u> </u>			<u></u>
						_
At Least	3		ere Purged Before Sam	ol Discharge Rate =	GPM x 0.00223 =	cfs
Comments:	15 gallons of w	ater removed during p	purging.			
				<u> </u>		
	<u> </u>				A B	
Form Complete	ed By:	Don Ostrand		Witnessed By:	Ara Mardirosain	

JSPCI Project N	lame:	UPMF Oakland F	acility Quarterly	Monitoring	USPCI Project Number:	96120-844
Measuring Point	t (MP)	Top of casing			Well No.	APL/UP-W1
Vell Depth: (Be		22.00	Feet			
asing diameter		2	inches		Sampling Date:	11/11/93
<del>T</del>	nd Water (Below I	MP): 10.25	Feet	<del>.</del>	Sample ID No.	APL/UP-W1
•	ct (Below MP): N					
	Development			Time	: 1714	
] Tap	[] Submersible [	Pump	[X] Inertia Pump		Riser Elevation (MP):	7.11
] Bailer	[] Centrifugal Pu	ımp	[] Other	Top of Screen Elevation:	2.11	Feet
ampling Collec	tion Method:			Sample Appearance:	turbid	
] Tap	[] Submersible	Pump	[] Inertia Pump:	Odor:	modorate petroleum odo	<u> </u>
() Bailer	Type:	[] Teflon	[] Stainless Steel	Sampling Problems (if an	y):	
	[X] HDPE Plastic	<u> </u>	[] PVC	[X] Disposable	<del> </del>	
ump Intake Or	Bailer Set At:	3.0	Feet Below MP	Decontamination Perform	ed:	
ubing Type (if	Used):	HDPE				
ubing Used fo	r:	[ ]SampleCollection	[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX.TPH/G/D/418.1,
•						As. Pb, 8010
		Temperature			Cumulative Volume	Pumping Rate
Time	pН	Corrected	Temperature	Water Level	of Water	in
	(Units)	Conductance	(Centigrade)	(Nearest 0.01 Ft.)	Removed From Well	Gallons/Minut
		(umho/cm)			(Gallons)	(GPM)
1735	7.2	1700	18.6		5.0	ļ
1737	7.2	1800	18.8		10.0	
1741	7.3	1700	18.6		15.0	<u> </u>
amples collect	ted at 1748					<u></u>
	ļ					
	<u> </u>					
·····			<del> </del>			
	ļ		<del> </del>		<del>-  </del>	
	<u> </u>	<u> </u>	-			· · · · · · · · · · · · · · · · · · ·
	-	<del>- </del>	<del> </del>			1
	-	<u> </u>	<del> </del>		<del>                                     </del>	
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At Least	3	Well Bore Volumes W	ere Purged Before Sam	pl Dischar <u>ge</u> Rate ≃	GPM x 0.00223 =	cfs
Comments:		rater removed during				
<u> </u>						
Form Complete		Don Ostrand		Witnessed By:	Ara Mardirosain	

SPCI Project i	Name:	UPMF Oakland	Facility Quarterly	Monitoring	USPCI Project Number:	96120-844
leasuring Poin	nt (MP)	Top of casing	<u> </u>		Well No.	APL/UP-W2
/ell Depth: (B	Below MP):	17.30	Feet		<u></u>	
asing diamete	er;	2	Inches		Sampling Date:	11/11/93
	nd Water (Below	MP): 9.20	Feet		Sample ID No.	APL/UP-W2
lethod Of We	il Development			Time	: 1810	
] Tap	[] Submersible	Pump	[X] Inertia Pump		Riser Elevation (MP):	7.62
] Bailer	[] Centrifugal P	ump	[] Other	Top of Screen Elevation:	2.62	. Feet
ampling Colle	ction Method:			Sample Appearance:	turbid	
] Тар	[] Submersible	Pump	[] Inertia Pump:	Odor:	modorate petroleum ode	or
X] Bailer	Type:	[] Teflon	[] Stainless Steel	Sampling Problems (if an	y):	
<u> </u>	[X] HDPE Plastic	<u> </u>	[] PVC	[X] Disposable		
ump Intake O	r Bailer Set At:	3.0	Feet Below MP	Decontamination Perform	ned:	
ubing Type (it		HDPE	W. C.			
Tubing Used fo			[X] Well Developme	ent/Field Tests	Samples Collected:	BTEX,TPH/Q/D/418.1,
uning Usea IC	<u> </u>	1 Journal of the Control of the Cont	. 424 11011 E010/0PH	<u></u>		As, Pb. 8010
		T			Cumulative Volume	Pumping Rate
		Temperature	-	Maran I amal	of Water	in
Time	pH	Corrected	Temperature	Water Level (Nearest 0.01 Ft.)	Removed From Well	Gallons/Minute
	(Units)	Conductance	(Centigrade)	(Nearest O.O. Pt.)	(Gallons)	(GPM)
1824	7.5	(umho/cm) 1800	17.9		5.0	
1830	7.6	1900	18.0		10.0	
1834	7.6	1900	18.1	-	15.0	
samples collec						
						<u> </u>
	<u> </u>				ļ	<del></del>
	<del> </del>					
			<del> </del>		<del> </del>	
			-			
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	- <b>_</b>		<u>, , , , , , , , , , , , , , , , , , , </u>			
At Least	3	Well Bore Volumes V	Vere Purged Before Sam	pl Discharge Rate =	GPM x 0.00223 =	cfs
Comments:						
						<del></del>
Form Complet	ted By:	Don Ostrand		Witnessed By:	Ara Mardirosain	



U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID: Sample Matrix:

Analysis Method:

First Sample #:

96120-844/UPRR MF Yard - Oakland

on power and the second se

Water

EPA 5030/8015/8020

311-0913

Sample

Receive Report

## TOTAL PURGEABLE PETROLEUM HYDROCARBONS with BTEX DISTINCTION

Analyte	Reporting Limit μg/L	Sample I.D. 311-0913 OKUS-W5	Sample I.D. 311-0914 OKUS-W3	Sample I.D. 311-0915 OKUS-W2	Sample I.D. 311-0916 OKUS-W1	Sample I.D. 311-0917 OKUS-W4	
Purgeable Hydrocarbons	50	590	9,500	24,000	91	5,500	
Benzene	0.5	14	390	540	1.1	250	
Toluene	0.5	3.1	110	150	0.88	53	
Ethyl Benzene	0.5	54	5,100	13,000	21	4,600	
Total Xylenes	0.5	6.2	130	280	1.6	140	
Chromatogram Pat	tern:	Gasoline	Gasoline	Gasoline	Gasolin <b>e</b>	Gasoline	

**Quality Control Data** 

Report Limit Multiplication Factor:	5.0	100	100	1.0	20
Date Analyzed:	11/19/93	11/21/93	11/21/93	11/22/93	11/22/93
Instrument Identification:	HP-5	HP-5	HP-5	HP-2	HP-2
Surrogate Recovery, %: (QC Limits = 70-130%)	115	106	106	101	111

Purgeable Hydrocarbons are quantitated against a fresh gasoline standard.

Analytes reported as N.D. were not detected above the stated reporting limit.

SEQUOIA ANALYTICAL



## SEQUOIA ANALYTICA

1900 Bates Avenue • Suite LM • Concord, California 94520 (510) 686-9600 • FAX (510) 686-9689

U.S.P.C.I.

24125 Aldine Westfield

Attention: Chris Byerman

Client Project ID:

96120-844/UPRR MF Yard - Oakland

Sampled:

Nov 10, 1993

Spring, TX 77373

CONTO PO RESERVADO AL SE CENSE ATO A

Sample Matrix: Analysis Method:

Water EPA 5030/8015 Received: Reported: Nov 11, 1993 Dec 8, 1993

First Sample #:

311-0918

## TOTAL PURGEABLE PETROLEUM HYDROCARBONS

Analyte	Reporting Limit μg/L	Sample I.D. 311-0918 Trip Blank*	
Purgeable Hydrocarbons	50	92	
Chromatogram Patte	ern:	Discrete	

**Quality Control Data** 

Report Limit Multiplication Factor:

1.0

Peak

Date Analyzed:

11/18/93

Instrument Identification:

HP-4

Surrogate Recovery:

(QC Limits = 70-130%)

95

Purgeable Hydrocarbons are quantitated against a fresh gasoline standard. Analytes reported as N.D. were not detected above the stated reporting limit.

**SEQUOIA ANALYTICAL** 

Karen L. Enstrom Project Manager

Please Note:

\*Purgeable Hydrocarbons are due to a discrete peak in the MTBE range. This peak was also found in all five OKUS samples.



U.S.P.C.I.

24125 Aldine Westfield

Spring, TX 77373 Attention: Chris Byerman Client Project ID: Sample Matrix:

First Sample #:

96120-844/UPRR MF Yard - Oakland

Water EPA 3510/3520/8015

Analysis Method: 311-0913 

Sampled:

Nov 11, 1993 Nov 11, 1993

Received: Reported:

Dec 8, 1993

## TOTAL EXTRACTABLE PETROLEUM HYDROCARBONS

Analyte	Reporting Limit μg/L	Sample I.D. 311-0913 OKUS-W5	Sample I.D. 311-0914 OKUS-W3	Sample I.D. 311-0915 OKUS-W2	Sample I.D. 311-0916 OKUS-W1	Sample I.D. 311-0917 OKUS-W4	
Extractable Hydrocarbons	50	1,600	5,000	7,700	160	2,400	
Chromatogram Pa	attern:	Diesel & Non-Diesel Mixture ( <c14;>C18)</c14;>	Diesel & Non-Diesel Mixture ( <c14)< td=""><td>Diesel &amp; Non-Diesel Mixture (<c14)< td=""><td>Diesel &amp; Non-Diesel Mixture (&gt;C20)</td><td>Diesel &amp; Non-Diesel Mixture (<c14;>C20)</c14;></td><td></td></c14)<></td></c14)<>	Diesel & Non-Diesel Mixture ( <c14)< td=""><td>Diesel &amp; Non-Diesel Mixture (&gt;C20)</td><td>Diesel &amp; Non-Diesel Mixture (<c14;>C20)</c14;></td><td></td></c14)<>	Diesel & Non-Diesel Mixture (>C20)	Diesel & Non-Diesel Mixture ( <c14;>C20)</c14;>	

**Quality Control Data** 

Report Limit Multiplication Factor:	1.0	10	10	1.0	1.0
Date Extracted:	11/17/93	11/17/93	11/17/93	11/17/93	11/17/93
Date Analyzed:	11/22/93	11/23/93	11/23/93	11/22/93	11/22/93
Instrument Identification:	HP-3B	НР-ЗА	HP-3A	HP-3A	HP-3A

Extractable Hydrocarbons are quantitated against a fresh diesel standard. Analytes reported as N.D. were not detected above the stated reporting limit.

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID: Matrix Descript:

96120-844/UPRR MF Yard - Oakland

Water

EPA 418.1 (I.R. with clean-up)

Analysis Method: First Sample #: 311-0913 Sampled:

Nov 11, 1993

Received: Nov 11, 1993 Nov 17, 1993 Extracted:

Analyzed: Nov 18, 1993 Reported: Dec 8, 1993

## TOTAL RECOVERABLE PETROLEUM HYDROCARBONS

Sample Number	Sample Description	Petroleum Oil mg/L (ppm)
311-0913	OKUS-W5	. 2.7
311-0914	OKUS-W3	2.3
311-0915	OKUS-W2	3.5
311-0916	OKUS-W1	N.D.
311-0917	OKUS-W4	N.D.

**Detection Limits:** 

1.0

Analytes reported as N.D. were not present above the stated limit of detection.

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## SEQUOIA ANALYTICAL

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman Client Project ID: Sample Descript:

Analysis for:

First Sample #:

96120-844/UPRR MF Yard - Oakland

Water

**Total Dissolved Arsenic** 

311-0913

Sampled:

Nov 11, 1993

Received: Nov 11, 1993 Extracted: Nov 21, 1993

Analyzed: Dec 3, 1993 Reported: Dec 8, 1993

## LABORATORY ANALYSIS FOR:

## **Total Dissolved Arsenic**

Sample Number	Sample Description	Detection Limit mg/L	Sample Result mg/L
311-0913	OKUS-W5	0.10	0.53
311-0914	OKUS-W3	0.10	0.14
311-0915	OKUS-W2	0.10	N.D.
311-0916	OKUS-W1	0.10	N.D.
311-0917	OKUS-W4	0.10	0.13

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



## SEQUOIA ANALYTICAL

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

Client Project ID: estfield Sample Descript:

96120-844/UPRR MF Yard - Oakland

Water

Analysis for: Total Dissolved Lead First Sample #: 311-0913

Sampled: Nov

Nov 11, 1993

Received: Nov 11, 1993 Extracted: Nov 21, 1993

Analyzed: Dec 3, 1993 Reported: Dec 8, 1993

#### LABORATORY ANALYSIS FOR:

## **Total Dissolved Lead**

Sample Number	Sample Description	Detection Limit mg/L	Sample Result mg/L
311-0913	OKUS-W5	0.020	N.D.
311-0914	OKUS-W3	0.020	N.D.
311-0915	OKUS-W2	0.020	N.D.
311-0916	OKUS-W1	0.020	N.D.
311-0917	OKUS-W4	0.020	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID:

Lab Number:

96120-844/UPRR MF Yard - Oakland

Sample Descript: Water, OKUS-W5

Analysis Method: EPA 5030/8010 311-0913

Sampled:

Nov 11, 1993 Nov 11, 1993 Received:

Analyzed: Reported: Nov 22, 1993 Dec 8, 1993

## **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit μg/L		Sample Results μg/L
Bromodichloromethane	0.50		N.D.
Bromoform	0.50		N.D.
Bromomethane	1.0	***************************************	N.D.
Carbon tetrachloride	0.50	***************************************	N.D.
Chlorobenzene	0.50	*************************	2, 20, 20, 20, 20, 20, 20, 20, 20, 20, 2
Chloroethane	1.0		N.D.
2-Chloroethylvinyl ether	1.0		N.D.
Chloroform	0.50		N.D.
Chioromethane	1.0		N.D.
Dibromochloromethane	0.50		N.D.
1,3-Dichlorobenzene	0.50	***************************************	N.D.
1,4-Dichlorobenzene	0.50		N.D.
1,2-Dichlorobenzene	0.50		N.D.
1,1-Dichloroethane	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1,2-Dichloroethane	0.50		N.D.
1,1-Dichloroethene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
cis-1,2-Dichloroethene	0.50		N.D.
trans-1,2-Dichloroethene	0.50		N.D.
1,2-Dichloropropane	0.50	***************************************	N.D.
cis-1,3-Dichloropropene	0.50		N.D.
trans-1,3-Dichloropropene	0.50	2442444444444	N.D.
Methylene chloride	5.0	***********	N.D.
1,1,2,2-Tetrachloroethane	0.50	***********	N.D.
Tetrachloroethene	0.50		N.D.
1,1,1-Trichloroethane	0.50	***************************************	N.D.
1,1,2-Trichloroethane	0.50	***************************************	N.D.
Trichloroethene	0.50	***************************************	N.D.
Trichlorofluoromethane	0.50	4114414111499999999999	N.D.
Vinyl chloride	1.0	***************************************	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID: Sample Descript:

Lab Number:

96120-844/UPRR MF Yard - Oakland

Water, OKUS-W3

Analysis Method: EPA 5030/8010 311-0914

Sampled:

Nov 11, 1993 Nov 11, 1993

Received: Analyzed:

Nov 22, 1993

Reported:

Dec 8, 1993

## **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit µg/L		Sample Results µg/L
Bromodichloromethane	0.50		N.D.
Bromoform	0.50		N.D.
Bromomethane	1.0		N.D.
Carbon tetrachloride	0.50		N.D.
Chlorobenzene	0.50	**************************************	ennagnigagnoods societys yr. h. <u>h. dynager yn, grennege</u> yr.
Chloroethane	1.0	***************************************	N.D.
2-Chloroethylvinyl ether	1.0	*************	N.D.
Chloroform	0.50		N.D.
Chloromethane	1.0	***************************************	N.D.
Dibromochloromethane	0.50		N.D.
1,3-Dichlorobenzene	0.50		N.D.
1,4-Dichlorobenzene	0.50		N.D.
1,2-Dichlorobenzene	0.50		N.D.
1,1-Dichloroethane	0.50		N.D.
1,2-Dichloroethane	0.50	***************************************	N.D.
1,1-Dichloroethene	0.50		N.D.
cis-1,2-Dichloroethene	0.50	***************************************	N.D.
trans-1,2-Dichloroethene	0.50		N.D.
1,2-Dichloropropane	0.50		N.D.
cis-1,3-Dichloropropene	0.50		N.D.
trans-1,3-Dichloropropene	0.50	17718	N.D.
Methylene chloride	5.0		N.D.
1,1,2,2-Tetrachloroethane	0.50		N.D.
Tetrachloroethene	0.50		N.D.
1,1,1-Trichloroethane	0.50	,	N.D.
1,1,2-Trichloroethane	0.50		N.D.
Trichloroethene	0.50		N.D.
Trichlorofluoromethane	0.50		N.D.
Vinyl chloride	1.0		N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



(510) 686-9600 • FAX (510) 686-9689

DELERGINGEREER DOR DE KOSELLES U.S.P.C.I. 24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

Client Project ID: Sample Descript: Water, OKUS-W2 Analysis Method:

Lab Number:

96120-844/UPRR MF Yard - Oakland

EPA 5030/8010 311-0915

Sampled: Received: Nov 11, 1993 Nov 11, 1993

Nov 22, 1993 Analyzed: Dec 8, 1993 Reported:

## **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit µg/L		Sample Results μg/L
Bromodichloromethane	5.0		N.D.
Bromoform	5.0		N.D.
Bromomethane	10	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Carbon tetrachloride	5.0		N.D.
Chlorobenzene	5.0		. 78
Chloroethane	10		N.D.
2-Chloroethylvinyl ether	10	***************************************	N.D.
Chloroform	5.0	***************************************	N.D.
Chloromethane	10	***************************************	N.D.
Dibromochloromethane	5.0	***************************************	N.D.
1,3-Dichlorobenzene	5.0	***************************************	N.D.
1,4-Dichlorobenzene	5.0	***************************************	N.D.
1,2-Dichlorobenzene	5.0	***************************************	N.D.
1,1-Dichloroethane	5.0	**!!**!!*!!*!	N.D.
1,2-Dichloroethane	5.0	***************************************	N.D.
1,1-Dichloroethene	5.0	***************************************	N.D.
cis-1,2-Dichloroethene	5.0		N.D.
trans-1,2-Dichloroethene	5.0	***************************************	N.D.
1,2-Dichloropropane	5.0		N.D.
cis-1,3-Dichloropropene	5.0	***************************************	N.D.
trans-1,3-Dichloropropene	5.0	***************************************	N.D.
Methylene chloride	50	******************************	N.D.
1,1,2,2-Tetrachloroethane	5.0	***************************************	N.D.
Tetrachloroethene	5.0		N.D.
1,1,1-Trichloroethane	5.0	***********	N.D.
1,1,2-Trichloroethane	5.0	,,.,.,.,	N.D.
Trichloroethene	5.0		N.D.
Trichloroflyoromethane	5.0	******	N.D.
Vinyl chloride	10		N.D.

Analytes reported as N.D. were not present above the stated limit of detection. Because matrix effects and/or other factors required additional sample dilution, detection limits for this sample have been raised.

SEQUOIA ANALYTICAL

U.S.P.C.I. 24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman Client Project ID: Sample Descript: Analysis Method:

Lab Number:

96120-844/UPRR MF Yard - Oakland

Water, OKUS-W1

EPA 5030/8010 311-0916 Sampled: N Received: N

Analyzed:

Reported:

Nov 11, 1993 Nov 11, 1993

Nov 22, 1993 Dec 8, 1993

## **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit µg/L		Sample Results μg/L
Bromodichloromethane	0.50	***************************************	N.D.
Bromoform	0.50	***************************************	N.D.
Bromomethane	1.0		N.D.
Carbon tetrachloride	0.50	•••••	N.D.
Chlorobenzene	0.50	<b>16140709070807080708070807080</b>	. 0.83
Chloroethane	1.0		N.D.
2-Chloroethylvinyl ether	1.0		N.D.
Chloroform	0.50		N.D.
Chloromethane	1.0		N.D.
Dibromochloromethane	0.50		N.D.
1,3-Dichlorobenzene	0.50		N.D.
1,4-Dichlorobenzene	0.50		N.D.
1,2-Dichlorobenzene	0.50		N.D.
1,1-Dichloroethane	0.50	***************************************	N.D.
1,2-Dichloroethane	0.50		N.D.
1,1-Dichloroethene	0.50		N.D.
cis-1,2-Dichloroethene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
trans-1,2-Dichloroethene	0.50	***************************************	N.D.
1,2-Dichloropropane	0.50	**********	N.D.
cis-1,3-Dichloropropene	0.50	,	N.D.
trans-1,3-Dichloropropene	0.50	***************************************	N.D.
Methylene chloride	5.0		N.D.
1,1,2,2-Tetrachloroethane	0.50	4	N.D.
Tetrachloroethene	0.50	*******************************	N.D.
1,1,1-Trichloroethane	0.50	*********	N.D.
1,1,2-Trichloroethane	0.50	465448446466444644	N.D.
Trichloroethene	0.50	************	N.D.
Trichlorofluoromethane	0.50	,	N.D.
Vinyl chloride	1.0		N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman Client Project ID: 96120-844/UPRR Sample Descript: Water, OKUS-W4

Lab Number:

96120-844/UPRR MF Yard - Oakland

Sample Descript: Water, OKUS-We Analysis Method: EPA 5030/8010

311-0917

Sampled: Received: Nov 11, 1993 Nov 11, 1993

Analyzed: Reported:

Nov 22, 1993 Dec 8, 1993

## **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

ŀ	Analyte	Detection Limit µg/L		Sample Results µg/L
	Bromodichloromethane	0.50	***************************************	N.D.
	Bromoform	0.50	*********	N.D.
	Bromomethane	1.0	*********	N.D.
	Carbon tetrachloride	0.50		N.D.
•	Chlorobenzene	0.50	#*****************************	
	Chloroethane	1.0	***************************************	N.D.
	2-Chloroethylvinyl ether	1.0	***************************************	N.D.
	Chloroform	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
ľ	Chloromethane	1.0		N.D.
ļ	Dibromochloromethane	0.50		N.D.
	1,3-Dichlorobenzene	0.50	***************************************	N.D.
•	1,4-Dichlorobenzene	0.50	,	N.D.
ı	1,2-Dichlorobenzene	0.50		N.D.
•	1,1-Dichloroethane	0.50	***************************************	N.D.
	1,2-Dichloroethane	0.50	***************************************	N.D.
Ì	1,1-Dichloroethene	0.50		N.D.
ŀ	cis-1,2-Dichloroethene	0.50		N.D.
	trans-1,2-Dichloroethene	0.50	141,447,44444444444444444	N.D.
	1,2-Dichloropropane	0.50	1**************************************	N.D.
l	cis-1,3-Dichloropropene	0.50	141744744444444444444444444444444444444	N.D.
,	trans-1,3-Dichloropropene	0.50	.44	N.D.
	Methylene chloride	5.0	,	N.D.
ŀ	1,1,2,2-Tetrachloroethane	0.50	***************************************	N.D.
l,	Tetrachloroethene	0.50		N.D.
	1,1,1-Trichloroethane	0.50	***************************************	N.D.
	1,1,2-Trichloroethane	0.50	*******************************	N.D.
Ī	Trichloroethene	0.50	************************	N.D.
ľ	Trich orofluoromethane	0.50	***************************************	N.D.
	Vinyl chloride	1.0	***************************************	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

**SEQUOIA ANALYTICAL** 



## **SEQUOIA ANALYTICAL**

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

SECRETERAL ESPOS O LA MISE

Client Project ID:

96120-844/UPRR MF Yard - Oakland

Matrix: Water

QC Sample Group: 3110913-917

Reported:

Dec 8, 1993

#### QUALITY CONTROL DATA REPORT

ANALYTE	Lead	Arsenic	
Method:	EPA 200.7	EPA 200.7	
Analyst:	K.A.	K.A.	
MS/MSD			
Batch#:	3111663	3111663	
Date Prepared:	12/1/93	12/1/93	
Date Analyzed:	12/3/93	12/3/93	
Instrument I.D.#:	Liberty 100	Liberty 100	
Conc. Spiked:	0.50 mg/L	0.50 mg/L	
Matrix Spike			
% Recovery:	92	90	
Matrix Spike			
Duplicate %			
Recovery:	92	92	
Relative %			

LCS Batch#:

Difference:

BLK112193

0.0

BLK112193

2.2

Date Prepared: Date Analyzed: 11/21/93 12/3/93 11/21/93

Instrument I.D.#:

12/3/93 Liberty 100 12/3/93 Liberty 100

LCS %

Recovery:

92

91

% Recovery Control Limits:

75-125

75-125

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Karen L. Enstrom Project Manager Please Note:

The LCS is a control sample of known, interferent free matrix that is analyzed using the same reagents, preparation, and analytical methods employed for the samples. The matrix spike is an aliquot of sample fortified with known quantities of specific compounds and subjected to the entire analytical procedure. If the recovery of analytes from the matrix spike does not fall within specified control limits due to matrix interference, the LCS recovery is to be used to validate the batch.



## SEQUOIA ANALYTICA

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U.S.P.C.I.

24125 Aldine Westfield

Spring, TX 77373 Attention: Chris Byerman Client Project ID:

96120-844/UPRR MF Yard - Oakland

Liquid Matrix:

QC Sample Group: 3110913-918

Reported:

Dec 8, 1993

#### QUALITY CONTROL DATA REPORT

ANALYTE	Benzene	Toluene	Ethyl Benzene	Xylenes	Diesel	Oil & Grease	
Method: Analyst:	EPA 8020 A.T.	EPA 8020 A.T.	EPA 8020 A.T.	EPA 8020 A.T.	EPA 8015 K.W.	EPA 418.1 S.L.	
MS/MSD Batch#:	3111144	3111144	3111144	3111144	BLK111793	BLK111793	
Date Prepared: Date Analyzed: Instrument I.D.#: Conc. Spiked:	11/18/93 11/18/93 HP-4 20 µg/L	11/18/93 11/18/93 HP-4 20 µg/L	11/18/93 11/18/93 HP-4 20 µg/L	11/18/93 11/18/93 HP-4 60 µg/L	11/17/93 11/22/93 HP-3B 300 µg/L	11/17/93 11/17/93 Miran IFF 5.0 mg/L	
Matrix Spike % Recovery:	100	95	100	100 98 9		76	
Matrix Spike Duplicate % Recovery:	100	100	105	103	101	74	
Relative % Difference:	0.0	5.1	4.9	4.9	6.8	2.7	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,

	LCS Batch#:	2LC\$111893	2LCS111893	2LCS111893	2LCS111893	BLK111793	LCS111793	
!r	Date Prepared: Date Analyzed: nstrument I.D.#:	11/18/93 11/18/93 HP-4	11/18/93 11/18/93 HP-4	11/18/93 11/18/93 HP-4	11/18/93 11/18/93 HP-4	11/17/93 11/22/93 HP-3B	11/17/93 11/17/93 Miran IFF	
j	LCS % Recovery:	98	97	98	97	95	76	
	% Recovery Control Limits:	71-133	72-128	72-130	71-120	80-120	70-130	

Karen L. Enstrom Project Manager

Please Note:

The LCS is a control sample of known, interferent free matrix that is analyzed using the same reagents, preparation, and analytical methods employed for the samples. The matrix spike is an aliquot of sample fortified with known quantities of specific compounds and subjected to the entire analytical procedure. If the recovery of analytes from the matrix spike does not fall within specified control limits due to matrix interference, the LCS recovery is to be used to validate the batch.



## SEQUOIA ANALYTICAL

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Para Parakanan salah Julya Sarah

Client Project ID:

96120-844/UPRR MF Yard - Oakland

Matrix:

Liquid

QC Sample Group: 3110913-917

Reported:

Dec 8, 1993

#### **QUALITY CONTROL DATA REPORT**

ANALYTE	1,1-Dichloro-	Triables	OH-
ANALITE		Trichloro-	Chloro-
	ethene	ethene	benzene
Method:	EPA 8010	EPA 8010	EPA 8010
Analyst:	K.N.	K.N.	K.N.
MS/MSD			
Batch#:	3110916	3110916	3110916
			0110010
Date Prepared:	11/22/93	11/22/93	11/22/93
Date Analyzed:	11/22/93	11/22/93	11/22/93
Instrument I.D.#:	HP-5890/6	HP-5890/6	HP-5890/6
Conc. Spiked:	10 µg/L	10 μg/L	10 μg/L
Matrix Spike			
% Recovery:	140	97	120
•			
Matrix Spike			
Duplicate %			
Recovery:	140	89	110
Relative %			
Difference:	0.0	0.6	
Dillelence.	U,U	8.6	8.7

LCS Batch#:	LCS112293	LCS112293	LCS112293
Date Prepared: Date Analyzed: estrument I.D.#:	11/22/93 11/22/93 HP-5890/6	11/22/93 11/22/93 HP-5890/6	11/22/93 11/22/93 HP-5890/6
LCS % Recovery:	140	110	110

% Recovery			
Control Limits:	28-167	35-146	38-150

**SEQUOIA ANALYTICAL** 

Karen L. Enstrom Project Manager Please Note:

The LCS is a control sample of known, interferent free matrix that is analyzed using the same reagents, preparation, and analytical methods employed for the samples. The matrix spike is an aliquot of sample fortified with known quantities of specific compounds and subjected to the entire analytical procedure. If the recovery of analytes from the matrix spike does not fall within specified control limits due to matrix interference, the LCS recovery is to be used to validate the batch.

#### CONTACT Chris Byerman CONTACT Same COMPANY Same COMPANY USPCIT A Subsidiary of Union Pacific Corporation ADDRESS 24125 Aldre West Field ADDRESS \_\_\_\_\_ Ship To: USPCI Remedial Services CITY Spring ST. TX ZIP 77313 T CITY \_\_\_\_\_\_ ST.\_\_ ZIP\_\_\_\_ 24125 Aldine Westfield PHONE (713) 350-7265 FAX Spring, TX 77373 PHONE \_\_\_\_\_\_ PO # (713) 350-7240 **CHAIN OF CUSTODY RECORD** PROJ. NO. 96120 - 844 STANDARD TURNAROUND. PROJECT NAME LIPRR MF Yard - Ochstand N **RUSH TURNAROUND** (specify required date) BTEX 0108 (SIGNATURE) ₩ 1.80 CUSTOMER LABORATORY DATE TIME MATRIX REMARKS SAMPLE I.D. SAMPLE I.D. 3110913 AG Wenter OKU8-WS 11/11/93 0901 H 11 H 11 4 u 11 2 11 11 " il. 10 u ıl 11 Filtered in Œ U 11 t ( Field 13110914 AG-OKUS-W3 11/11/93 1018 Water 11 11 11 ١ T. 11 11 u 2 11 11 L u 4 ( 11 11 11 u Filtered in 11 10 11 BECEIVED BY COURIER DATE / TIME RELINQUISHED BY

HECEIVED BY

DATE / TIME

AIRBILL NO

# **USPCI**A Substitions of

A Subsidiary of Union Pacific Corporation Ship To: USPCI Remedial Services

24125 Aldine Westfield Spring, TX 77373 (713) 350-7240

R	CONTACT Chris Byerman
E	COMPANY USPCT
O R	ADDRESS 24125 Aldine West Field
Ť	CITY Spring ST.TX ZIP 77313
ŏ	PHONE (7/3) 350-7265 FAX

CONTACT Surve	Sory	_Nº	000166
COMPANY Sume	<b>.</b>		· ·
ADDRESS			
CITY	ST	_ ZIP_	•
PHONE	PO#		

						СНА	IN OI	F CU	STO	DY R	ECOR	)					
	16120-844 HE UPRR ME	Yard-0	akternel	#CON+4-NE	BTEX	TPH-6	0/08	TPH-D		Notal Divolved					RUSH	DARD TU	
CUSTOMER SAMPLE I.D.	DATE	TIME	MATRIX	RS	187	11	80	7	1h	401						RATORY PLE I.D.	REMARKS
OKUS-WZ	11/1/93	701	water	1	/										3110	19151	16
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U	11	4	11	2													
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4	11	"	((	1					/								<u> </u>
1/	11	11	11	1						-					7	/	Filtered in Field
OKUS-WI	1/11/93	1233	water	1											311	0 <sup>0</sup> 116	Filtered in Field AG
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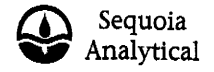
A Subsidiary of Union Pacific Corporation

Ship To: USPC1 Remedial Services 24125 Aldine Westfield Spring, TX 77373 (713) 350-7240

R	CONTACT Chris Byerman
Ē	COMPANY USPCI
OR	ADDRESS 24125 Aldine Westfield
T	CITY Sering ST. TX ZIP 77373
T	CITY Spring ST. TX ZIP 77373  PHONE (713) 350-7265 FAX

1	CONTACT Sum		Νº	0001.67
8	COMPANY Sume			•
L	ADDRESS			
Ţ	CITY	ST	ZIP_	•
٦	PHONE	PO#		

						CHA	IN O	F CU	STO	DY R	ECOF	RD						
	EUPRR MF		Oakland	<b>JEZ-&gt;</b> 4200#	8TEX	TPH-G	0/08	TPH-D	1.8.1	Total Dussolved Ariquic + Leach			is the state of th			FI (s	USH TURNA pecily require	ed date)
CUSTOMER SAMPLE I.D.	DATE	TIME	MATRIX	S	8	7	28	1	7/	70 th							BORATORY	REMARKS
OKUS-WY	11/1/93	1442	water	1	<b>/</b>											3	11091	TAG
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U	ધ	ιι	Ιį	2_			<b>'</b>											
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11	l l	l l	11	l					~	<u> </u>								
11	11	11	(1	1						V							<u> </u>	Filtered in Field
Trip Blank	11/10/93	0810	abter	1		/										3	11091	
RELINOUISHED BY			11/11/9	DATE / T	1615	-	ECEIVED		<u> </u>					14/	DATE:		COURIEF AIRBILL N	



680 Chesapeake Drive 1900 Bates Avenue, Suite L 819 Striker Avenue, Suite 8

Redwood City, CA 94063 Concord, CA 94520 Sacramento, CA 95834

(415) 364-9600 (510) 686-9600 (916) 921-9600 FAX (415) 364-9233 FAX (510) 686-9689 FAX (916) 921-0100

24125 Aldine Westfield Spring, TX 77373

U.S.P.C.I, Client Project ID: Sample Matrix:

D: #96120-844/UPRR MF Yard - Oakland

Water

Sampled: Received:

Nov 11, 1993 Nov 12, 1993

Attention: Chris Byerman

Analysis Method: First Sample #:

EPA 5030/8015/8020

Reported:

Dec 8, 1993

311-1019

#### TOTAL PURGEABLE PETROLEUM HYDROCARBONS with BTEX DISTINCTION

Analyte	Reporting Limit µg/L	Sample I.D. 311-1019 APL-W1	ا 811ھر	mple .D. -1020 L-W2 i	Sample I.D. 311-1021 OKUS-W8	Sample I.D. 311-1022 OKUS-W9	<b>Sample</b> I.D. 311-1023 OKUS-W7	Sample I.D. 311-1024 OKUS-W6
Purgeable Hydrocarbons	50	560	•	110	190	120	N.D.	N.D.
Benzene	0.5	26	!	5.0 🗸	3.5	1.3	N.D.	3.6
Toluene	0.5	N.D.	٨	I.D.	1.3	N.D.	N.D.	N.D.
Ethyl Benzene	0.5	220	1	38	46	4.0	N,D.	3.7
Total Xylenes	0.5	11	;	2.6	4.9	1.4	N.D.	1.3
Chromatogram Patt	ern:	Gasoline	Ga	oline	Gasoline	Gasoline	••	<del></del>

Quality Control Data

Report Limit Multiplication Factor:	4.0	1.0	1.0	1.0	1.0	1.0
Date Analyzed:	11/19/93	11/19/93	11/19/93	11/19/93	11/18/93	11/19/93
Instrument identification:	HP-4	HP-4	HP-2	HP-2	HP-4	HP-4
Surrogate Recovery, %: (QC Limits = 70-130%)	94	91	114	106	90	96

Purgeable Hydroparbons are quantitated against a fresh gasoline standard. Analytes reported as N.D. were not detected above the stated reporting limit.

**SEQUOIA ANALYTICAL** 

Please Note:

Revised Report 4/28/94

rolect Manage



U.S.P.C.I.

24125 Aldine Westfield

Spring, TX 77373

Client Project ID:

#96120-844/UPRR MF Yard - Oakland Water

Nov 10, 1993

Spring, TX 77373

Sample Matrix: Analysis Method:

EPA 5030/8015/8020

Received: Reported:

Sampled:

Nov 12, 1993 Dec 8, 1993

Attention: Chris Byerman

First Sample #:

311-1029

ious kuus, populis (1990 agga paraki pet patatina tii 1 teeteti keete. 2007 2009)

#### TOTAL PURGEABLE PETROLEUM HYDROCARBONS with BTEX DISTINCTION

Analyte	Reporting Limit μg/L	Sample I.D. 311-1029 Trip Blank	
Purgeable Hydrocarbons	50	N.D.	
Benzene	0.5	N.D.	
Toluene	0.5	N.D.	
Ethyl Benzene	0.5	N.D.	
Total Xylenes	0.5	N.D.	
Chromatogram Patt	ern:		

#### **Quality Control Data**

Report Limit Multiplication Factor:

1.0

Date Analyzed:

11/18/93

Instrument Identification:

HP-5

Surrogate Recovery, %:

100

(QC Limits = 70-130%)

Purgeable Hydrocarbons are quantitated against a fresh gasoline standard. Analytes reported as N.D. were not detected above the stated reporting limit.

SEQUOIA ANALYTICAL



U.S.P.C.L

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID:

#96120-844/UPRR MF Yard - Oakland

Sampled:

Nov 11, 1993

Sample Matrix: Analysis Method:

EPA 3510/3520/8015

Received:

Nov 12, 1993 Dec 9, 1993

First Sample #:

311-1019

Water

Reported:

#### TOTAL EXTRACTABLE PETROLEUM HYDROCARBONS

Analyte	Reporting Limit μg/L	Sample 1.D. 311-1019 APL-W1	Sample I.D. 311-1020 PAL-W2	Sample I.D. 311-1021 OKUS-W8	Sample I.D. 311-1022 OKUS-W9	Sample I.D. 311-1023 OKUS-W7	Sample I.D. 311-1024 OKUS-W6
Extractable Hydrocarbons	50	530	190	1,300	1,300	1,100	610
Chromatogram Pa	ttern:	Diesel & Non-Diesel Mixture (< C14; > C20)	Diesel & Non-Diesel Mixture ( <c14;>C20)</c14;>	Diesel & Non-Diesel Mixture ( <c14;>C20)</c14;>	Diesel & Non-Diesel Mixture ( <c14;>C20)</c14;>	Diesel & Non-Diesel Mixture ( > C20)	Diesel & Non-Diesel Mixture ( > C20)

**Quality Control Data** 

Report Limit Multiplication Factor:	1.0	1.0	1.0	1.0	1.0	1.0
Date Extracted:	11/17/93	11/17/93	11/17/93	11/17/93	11/17/93	11/17/93
Date Analyzed:	11/22/93	11/22/93	11/22/93	11/22/93	11/22/93	11/22/93
Instrument Identification:	HP-3A	HP-3A	HP-3A	HP-3A	HP-3A	HP-3B

Extractable Hydrocarbons are quantitated against a fresh diesel standard. Analytes reported as N.D. were not detected above the stated reporting limit.

SEQUOIA ANALYTICAL



U.S.P.C.I. 24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Sample Descript: Water; APL-W1

Lab Number:

Client Project ID: #96120-844/UPRR MF Yard - Oakland

Analysis Method: EPA 5030/8010 311-1019

Sampled: Received: Analyzed:

Nov 11, 1993 Nov 12, 1993

Nov 23, 1993 Reported: Dec 9, 1993

#### **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

ļ	Analyte	Detection Limit µg/L		Sample Results µg/L
E	Bromodichloromethane	0.50	***************************************	N.D.
E	Bromoform	0.50	••••••	N.D.
E	Bromomethane	1.0	***************************************	N.D.
(	Carbon tetrachloride	0.50		N.D.
(	Chlorobenzene	0.50	*******************************	N.D.
(	Chloroethane	1.0		N.D.
2	2-Chloroethylvinyl ether	1.0		N.D.
	Chloroform	0.50		N.D.
(	Chloromethane	1.0		N.D.
. [	Dibromochloromethane	0.50	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1	I,3-Dichlorobenzene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1	I,4-Dichlorobenzene	0.50	*********	N.D.
	1,2-Dichlorobenzene	0.50		N.D.
1	1-Dichloroethane	0.50		N.D.
1	2-Dichloroethane	0.50	,,,	N.D.
1	1,1-Dichloroethene	0.50		N.D.
	cis-1,2-Dichloroethene	0.50		N.D.
	rans-1,2-Dichloroethene	0.50		N.D.
	1,2-Dichloropropane	0.50		N.D.
	cis-1,3-Dichloropropene	0.50		N.D.
	rans-1,3-Dichloropropene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
	Methylene chloride	5.0	,	N.D.
	1,1,2,2-Tetrachloroethane	0.50		N.D.
Ī	Tetrachloroethene	0.50		N.D.
1	1,1,1-Trichloroethane	0.50	***************************************	N.D.
. 1	1,1,2-Trichloroethane	0.50	p.,	N.D.
	Frichloroethene	0.50		N.D.
	Frichlorofluoromethane	0.50	.,,,,,	N.D.
١	/inyl chloride	1.0	***************************************	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

**SEQUOIA ANALYTICAL** 

#### SEQUOIA ANALYTICAL 1900 Bates Avenue • Suite LM • Concord, California 94520 (510) 686-9600 • FAX (510) 686-9689

U.S.P.C.I. 24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

Sample Descript: Water; APL-W2 Analysis Method:

Lab Number:

Client Project ID: #96120-844/UPRR MF Yard - Oakland

EPA 5030/8010 311-1020

Sampled: Received: Analyzed: Nov 11, 1993. Nov 12, 1993 Nov 23, 1993,

Reported: Dec 9, 1993

#### **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

	Analyte	Detection Limit µg/L		Sample Result <b>s</b> µg/L
	Bromodichloromethane	0.50	***************************************	N.D.
	Bromoform	0.50		N.D.
	Bromomethane	1.0		N.D.
	Carbon tetrachloride	0.50	***************************************	N.D.
	Chlorobenzene	0.50	***************************************	N.D.
	Chloroethane	1.0	***************************************	N.D.
	2-Chloroethylvinyl ether	1.0	***************************************	N.D.
	Chloroform	0.50	***************************************	N.D.
	Chloromethane	1.0	***************************************	N.D.
	Dibromochloromethane	0.50	***************************************	N.D.
	1,3-Dichlorobenzene	0.50	***************************************	N.D.
	1,4-Dichlorobenzene	0.50	***************************************	N.D.
	1,2-Dichlorobenzene	0.50		N.D.
	1,1-Dichloroethane	0.50		N.D.
	1,2-Dichloroethane	0.50	<pre><pre></pre></pre>	N.D.
	1.1-Dichloroethene	0.50	***************************************	N.D.
r	cis-1,2-Dichloroethene	0.50	***************************************	N.D.
	trans-1,2-Dichloroethene	0.50	***************************************	N.D.
	1,2-Dichloropropane	0.50		N.D.
	cis-1,3-Dichloropropene	0.50		N.D.
	trans-1,3-Dichloropropene	0.50	,	N.D.
	Methylene chloride	5.0	14.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
u	1,1,2,2-Tetrachloroethane	0.50	1465445441854548855444444444444444444444	N.D.
	Tetrachloroethene	0.50		N.D.
	1,1,1-Trichloroethane	0.50		N.D.
ļ	1,1,2-Trichloroethane	0.50		N.D.
ĺ	Trichloroethene	0.50	***************************************	N.D.
	Trichlorofluoromethane	0.50	*  *	N.D.
	Vinvl chloride	1.0	\$14\$44\$474\$4P4P4P4P4P4	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID: Sample Descript:

#96120-844/UPRR MF Yard - Oakland

Water, OKUS-W8 Analysis Method: EPA 5030/8010

Lab Number: 311-1021

Sampled: Received:

Nov 11, 1993 Nov 12, 1993.

Analyzed: Reported:

Nov 23, 1993 Dec 9, 1993

#### **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

	Analyte	Detection Limit µg/L		Sample Results μg/L
	Bromodichloromethane	0.50		N.D.
i	Bromoform	0.50		N.D.
	Bromomethane	1.0		N.D.
	Carbon tetrachloride	0.50	,	N.D.
	Chlorobenzene	0.50	***************************************	N.D.
	Chloroethane	1.0		N.D.
	2-Chloroethylvinyl ether	1.0	********************************	N.D.
	Chloroform	0.50		N.D.
•	Chloromethane	1.0	**!**!**!	N.D.
,	Dibromochloromethane	0.50	***************	N.D.
	1,3-Dichlorobenzene	0.50		N.D.
	1,4-Dichlorobenzene	0.50		N.D.
	1,2-Dichlorobenzene	0.50		N.D.
	1,1-Dichloroethane	0.50		N.D.
	1,2-Dichloroethane	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
	1,1-Dichloroethene	0.50	491104494444444444	N.D.
)	cis-1,2-Dichloroethene	0.50		N.D.
	trans-1,2-Dichloroethene	0.50	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
	1,2-Dichloropropane	0.50		N.D.
	cis-1,3-Dichloropropene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
	trans-1,3-Dichloropropene	0.50	145455455555577777777777777777777777777	N.D.
	Methylene chloride	5.0		N.D.
ŀ	1,1,2,2-Tetrachloroethane	0.50		N.D.
ļ	Tetrachloroethene	0.50	,.,,,,	N.D.
	1,1,1-Trichloroethane	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1	1,1,2-Trichloroethane	0.50	***************************************	N.D.
	Trichloroethene	0.50	}\$\$14\$494949;*******************************	N.D.
ı	Trichlorofluoromethane	0.50	***************************************	N.D.
	Vinyl chloride	1.0	***************************************	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



1900 Bates Avenue • Suite LM • Concord, California 94520 (510) 686-9600 • FAX (510) 686-9689

U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID: Sample Descript: Analysis Method:

Lab Number:

#96120-844/UPRR MF Yard - Oakland

Water; OKUS-W9 EPA 5030/8010

311-1022

Sampled: Received:

Nov 11, 1993 Nov 12, 1993

Analyzed: Reported:

Nov 23, 1993 Dec 9, 1993

#### **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit µg/L		Sample Results µg/L
Bromodichloromethane	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Bromoform	0.50	4,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Bromomethane	1.0	************	N.D.
Carbon tetrachloride	0.50	***********	N.D.
Chlorobenzene	0.50		N.D.
Chloroethane	1.0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
2-Chloroethylvinyl ether	1.0	***************************************	N.D.
Chloroform	0.50	1001400	N.D.
Chloromethane	1.0		N.D.
Dibromochloromethane	0.50	************	N.D.
1,3-Dichlorobenzene	0.50		N.D.
1,4-Dichlorobenzene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1,2-Dichlorobenzene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1,1-Dichloroethane	0.50		N.D.
1,2-Dichloroethane	0.50		N.D.
1,1-Dichloroethene	0.50		N.D.
cis-1,2-Dichloroethene	0.50	***************************************	N.D.
trans-1,2-Dichloroethene	0.50	*14************************************	N.D.
1,2-Dichloropropane	0.50	***************************************	N.D.
cis-1,3-Dichloropropene	0.50	214471437777777777	N.D.
trans-1,3-Dichloropropene	0.50	************	N.D.
Methylene chloride	5.0		N.D.
1,1,2,2-Tetrachloroethane	0.50		N.D.
Tetrachloroethene	0.50		N.D.
1,1,1-Trichloroethane	0.50		N.D.
1,1,2-Trichloroethane	0.50		N.D.
Trichloroethene	0.50	******************************	N.D.
Trichlorofluoromethane	0.50	***************************************	N.D.
Vinyl chloride	1.0		N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

**SEQUOIA ANALYTICAL** 

U.S.P.C.I. 24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

Client Project ID: Sample Descript: Analysis Method: Lab Number:

#96120-844/UPRR MF Yard - Oakland Water; OKUS-W7 EPA 5030/8010 311-1023

Sampled: Nov 11, 1993 Received: Analyzed: Reported:

Nov 12, 1993 Nov 23, 1993 orted: Dec 9, 1993

#### **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit µg/L		Sample Results µg/L
Bromodichloromethane	0.50		N.D.
Bromoform	0.50	*!*************************************	N.D.
Bromomethane	1.0	24427491498498498478787878787878	N.D.
Carbon tetrachloride	0.50	***************************************	N.D.
Chlorobenzene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Chloroethane	1.0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
2-Chloroethylvinyl ether	1.0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Chloroform	0.50		N.D.
Chloromethane	1.0	14014014444011114	N.D.
Dibromochloromethane	0.50		N.D.
1,3-Dichlorobenzene	0.50	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1,4-Dichlorobenzene	0.50		N.D.
1,2-Dichlorobenzene	0.50	,	N.D.
1,1-Dichloroethane	0.50	1,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
1,2-Dichloroethane	0.50	***********	N.D.
1,1-Dichloroethene	0.50	**********	N.D.
cis-1,2-Dichloroethene	0.50	<pre></pre>	N.D.
trans-1,2-Dichloroethene	0.50		N.D.
1,2-Dichloropropane	0.50		N.D.
cis-1,3-Dichloropropene	0.50	4,,44,,11,41,41,41,41,41,41,41,41,41,41,	N.D.
trans-1,3-Dichloropropene	0.50	4	N.D.
Methylene chloride	5.0	***************************************	N.D.
1,1,2,2-Tetrachloroethane	0.50	***************************************	N.D.
Tetrachloroethene	0.50	***************************************	N.D.
1,1,1-Trichloroethane	0.50		N.D.
1,1,2-Trichloroethane	0.50	,,	N.D.
Trichloroethene	0.50	(*)(*)(*)	N.D.
Trichlorofluoromethane	0.50	***************************************	N.D.
Vinyl chloride	1.0	***************************************	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



U.S.P.C.I. 24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

Lab Number:

Client Project ID: #96120-844/UPRR MF Yard - Öakland

Sample Descript: Water; OKUS-W6 Analysis Method: EPA 5030/8010 311-1024

Sampled: Received: Analyzed:

Nov 11, 1993 Nov 12, 1993

Nov 23, 1993 Reported: Dec 9, 1993

#### **HALOGENATED VOLATILE ORGANICS (EPA 8010)**

Analyte	Detection Limit µg/L		Sample Results µg/L
Bromodichloromethane	0.50	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Bromoform	0.50	***************************************	N.D.
Bromomethane	1.0		N.D.
Carbon tetrachloride	0.50	***************************************	N.D.
Chlorobenzene	0.50		N.D.
Chloroethane	1.0		N.D.
2-Chloroethylvinyl ether	1.0		N.D.
Chloroform	0.50		N.D.
Chloromethane	1.0	******	N.D.
Dibromochloromethane	0.50		N.D.
1,3-Dichlorobenzene	0.50	***************************************	N.D.
1,4-Dichlorobenzene	0.50	***************************************	N.D.
1,2-Dichlorobenzene	0.50	******	N.D.
1,1-Dichloroethane	0.50	***************************************	N.D.
1,2-Dichloroethane	0.50		N.D.
1,1-Dichloroethene	0.50	***************************************	N.D.
cis-1,2-Dichloroethene	0.50		N.D.
trans-1,2-Dichloroethene	0.50		N.D.
1,2-Dichloropropane	0.50	***************************************	N.D.
cis-1,3-Dichloropropene	0.50	***************************************	N.D.
trans-1,3-Dichloropropene	0.50	***************************************	N.D.
Methylene chloride	5.0	***************************************	N.D.
1,1,2,2-Tetrachloroethane	0.50		N.D.
Tetrachloroethene	0.50		N.D.
1,1,1-Trichloroethane	0.50	***************************************	N.D.
1,1,2-Trichloroethane	0.50		N.D.
Trichloroethene	0.50	491499149171777777777	N.D.
Trichlorofluoromethane	0.50	***************************************	N.D.
Vinyl chloride	1.0		N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



1900 Bates Avenue • Suite LM • Concord, California 94520 (510) 686-9600 • FAX (510) 686-9689

U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

. Paguadetis silijudije jastajsgasjes

Client Project ID: Sample Descript:

Analysis for:

First Sample #:

#96120-844/UPRR MF Yard - Oakland

Water; APL-W1 Dissolved Arsenic

311-1019

Sampled:

Nov 11, 1993

Received: Extracted: Nov 12, 1993 Nov 30, 1993

Analyzed: Reported: 

Dec 5, 1993 Dec 9, 1993

#### LABORATORY ANALYSIS FOR:

#### **Dissolved Arsenic**

Sample Number	Sample Description	Detection Limit mg/L	Sample Result mg/L
311-1019	APL-W1	0.10	N.D.
311-1020	APL-W2	0.10	N.D.
311-1021	OKUS-W8	0.10	N.D.
311-1022	OKUS-W9	0.10	N.D.
311-1023	OKUS-W7	0.10	N.D.
311-1024	OKUS-W6	0.10	N.D.
311-1025	OMW-1	0.10	N.D.
311-1026	OMW-8	0.10	N.D.
311-1027	OMW-4	0.10	N.D.
311-1028	OMW-9	0.10	2.4

Analytes reported as N.D. were not present above the stated limit of detection.

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#### SEQUOIA ANALYTICAL

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373 Attention: Chris Byerman

Client Project ID:

#96120-844/UPRR MF Yard - Oakland

Sample Descript: Analysis for: First Sample #: Water; APL-W1 Dissolved Lead 311-1019 Sampled: Received: Nov 11, 1993 Nov 12, 1993

Extracted: Analyzed:

Reported:

Nov 30, 1993 Dec 5, 1993 Dec 9, 1993

#### LABORATORY ANALYSIS FOR:

#### **Dissolved Lead**

Sample Number	Sample Description	Detection Limit mg/L	Sample Result mg/L
311-1019	APL-W1	0.020	N.D.
311-1020	APL-W2	0.020	N.D.
311-1021	OKUS-W8	0.020	N.D.
311-1022	OKUS-W9	0.020	N.D.
311-1023	OKUS-W7	0.020	N.D.
311-1024	OKUS-W6	0.020	N.D.

Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL



U.S.P.C.I.

24125 Aldine Westfield

Spring, TX 77373

Attention: Chris Byerman

Client Project ID:

#96120-844/UPRR MF Yard - Oakland

Matrix:

QC Sample Group: 3111019-028

Reported:

Dec 9, 1993

#### **QUALITY CONTROL DATA REPORT**

l								
ANALYTE	Lead	Arsenic						
Method:	EPA 200.7	EPA 200.7						
Analyst:	K.A.	K.A.	 					
MS/MSD								
Batch#:	3111019	3111019						
<b>D</b> 410,, .	0111010							
Date Prepared:	11/30/93	11/30/93						
Date Analyzed:	12/5/93	12/5/93						
Instrument I.D.#:	Liberty 100	Liberty 100						
Conc. Spiked:	1.0 mg/L	1.0 mg/L						
Matrix Spike								
% Recovery:	96	111						
•								
Matrix Spike								
Duplicate %								
Recovery:	98	113						
Relative %								
Difference:	2.1	1.8						

LCS Batch#:

BLK113093

BLK113093

**Date Prepared:** Date Analyzed: 11/30/93 12/5/93

11/30/93

Instrument I.D.#:

Liberty 100

12/5/93 Liberty 100

LCS %

Recovery:

82

88

% Recovery

**Control Limits:** 

75-125

75-125

SEQUOIA ANALYTICAL

(aren L. Enstrom Project Manager

Please Note:

The LCS is a control sample of known, interferent free matrix that is analyzed using the same reagents, preparation, and analytical methods employed for the samples. The matrix spike is an aliquot of sample fortified with known quantities of specific compounds and subjected to the entire analytical procedure. If the recovery of analytes from the matrix spike does not fall within specified control limits due to matrix interference, the LCS recovery is to be used to validate the batch.



## SEQUOIA ANALYTICAL

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U.S.P.C.I.

24125 Aldine Westfield Spring, TX 77373

Attention: Chris Byerman

Client Project ID:

#96120-844/UPRR MF Yard - Oakland

Matrix:

Liquid

QC Sample Group: 3111019-024

Reported:

Dec 9, 1993

#### QUALITY CONTROL DATA REPORT

ANALYTE	1,1-Dichloro- ethene	Trichloro- ethene	Chloro- benzene	
Method: Analyst:	EPA 8010 K.N.	EPA 8010 K.N.	EPA 8010 K.N.	
Analysti	K.N.	13.14.	IXIIV.	
MS/MSD				
Batch#:	3111805	3111805	3111805	
Date Prepared:	11/23/93	11/23/93	11/23/93	
Date Analyzed:	11/23/93	11/23/93	11/23/93	
Instrument I.D.#:	HP-5890/6	HP-5890/6	HP-5890/6	
Conc. Spiked:	10 μg/L	10 µg/L	10 μg/L	
Matrix Spike				
% Recovery:	140	100	110	
Matrix Spike			,	
Duplicate %				
Recovery:	140	98	110	
Relative %				
Difference:	0.0	2.0	0.0	

LUS Batch#:	LCS112393	LCS112393	LCS112393	
Date Prepared:	11/23/93	11/23/93	11/23/93	
Date Analyzed:	11/23/93	11/23/93	11/23/93	
Instrument I.D.#:	HP-5890/6	HP-5890/6	HP-5890/6	
LCS %				
Recovery:	140	100	110	
% Recovery		<del> </del>		
Control Limits:	28-167	35-146	38-150	

1.00110000

SEQUOIA ANALYTICAL

Karen L. Enstrom Project Manager Please Note:

The LCS is a control sample of known, interferent free matrix that is analyzed using the same reagents, preparation, and analytical methods employed for the samples. The matrix spike is an aliquot of sample fortified with known quantities of specific compounds and subjected to the entire analytical procedure. If the recovery of analytes from the matrix spike does not fall within specified control limits due to matrix interference, the LCS recovery is to be used to validate the batch.

## **USPCI**

A Subsidiary of Union Pacific Corporation Ship To: USPCI Remedial Services 24125 Aldine Westfield Spring, TX 77373 (713) 350-7240

R	CONTACT Chris Byerman
E	COMPANY_USPCT
OR	ADDRESS 24125 Aldine Westfield
T	CITY Spring ST. TX ZIP 77373  PHONE (7/3) 150-7265 FAX
ò	PHONE (7/3) 150-7265 FAX

CONTACT Scine		15	000155
COMPANY Same			
ADDRESS			· · · · · ·
CITY	ST Z	<u>'</u> IP	
PHONE	PO#		

					(	CHAI	N OF	CU	STO	DY R	ECOR	D								
	16120-844 E UPRR ME		Hand	#CONTA-NE	BTEX	TPH-6	8010	TOH-D	माउत्त	Total Dissulved							RUSH <sup>1</sup> (specify	TURNARe y required		
CUSTOMER SAMPLE I.D.	DATE	TIME	MATRIX	R S	3	7	Ø	7	7	Te							LABOR/ SAMPI	ATORY LE I.D.	REMARK	5
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# **USPCI**

A Subsidiary of Union Pacific Corporation

Ship To: USPCI Remedial Services
24125 Aldine Westfield
Spring, TX. 77373
(713) 350-7240

	CONTACT Chris Byerman		CONTACT Same		Nº 00015€
E	COMPANY USPCI	B	COMPANY Sware		
P	ADDRESS 24125 Aldine Westfield	L	ADDRESS		
Ť	CITY Spring ST. TX ZIP 77373  PHONE (213) 350-7265 FAX	Ţ	CITY	ST	_ ZIP
o	PHONE (713) 350-7265 FAX		PHONE	PO #	
	OHAN OF OUOTOBY BEOO	<u> </u>			

	CHAIN OF CUSTODY RECORD																	
	6120-84L ELLPRE ME		bal-land	#CONTA-NERS	RTEX	TPH-6	&O (O	TPH-D	41811	Total Dissolved Arsente + Lend				F 18			STANDARD TO RUSH TURNA (specify require	ROUND od date)
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A Subsidiary of Union Pacific Corporation

Ship To: USPCI Remedial Services 24125 Aldine Westfield Spring, TX 77373 (713) 350-7240

R	CONTACT Chris Byerman
R E P O	ADDRESS 24125 Aldine Westfield
Τ	CITY 5 pring ST. TX ZIP 77373
o	PHONE (713) 350-7265 FAX

	CONTACT Same	Nº	000168
В	COMPANY Surve		
L L	ADDRESS		
Ţ	CITY	ST ZIP_	
O	PHONE	PO#	<u>.</u>

	<del></del>				(	CHA	IN OF	CUS	STOE	Y RI	COR	D	·							
PROJECT NAM SAMPLERS — (SIGNATURE)	6120-84 E UPRR ME		aldand	#CONTAINER	Brex	TPH-G	0108	TPH-D	-	Total Disserved Arsenic + Lead							RUSH (specify	TURNAR TURNAR y required	i date)	<b>\</b>
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# A Subsidiary of Union Pacific Corporation Ship To: USPCI Remedial Services 24125 Aldine Westfield Spring, TX 77373 (713) 350-7240

R E P O R T T O	CONTACT Chris Byernan
	COMPANY USPCT
	ADDRESS 24125 Aldine Westfield  CITY Spring SITX ZIP 77171
	CITY _ Spring ST.TX ZIP ST.TX ZIP ST.TX ZIP

-	CONTACT Same		N2	0001	5.7
	COMPANY Suna				· 
	ADDRESS				
	CITY	ST	_ ZIP_	•	
	PHONE	PO #			

						CHA	IN OF	CUS	STOE	Y RI	CO	RD		_						
PROJ. NO. 9 PROJECT NAM SAMPLERS (SIGNATURE)	#CONTAINERS	A T A T A T A T A T A T A T A T A T A T								STANDARD TURNAROUND  RUSH TURNAROUND (specify required date)										
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6mw-1	11/12/93	1214	Water	1	1										3	111025	Fig	ered eld	<u> </u>	
omw-8		1235	water	1	V											1026				
H-mmo	11/12/93	1254	water	1	/											1027				
		1320	Water	1							=					1028				
OMW-9 Trip Nank	11/10/91	0830	water	1		<b>\</b>									ما	1029				
THELINOUISHED BY	(bt.	-0	11/12	DATE /	TIME Z	Y7#	ECEIVED E	3Y		<u> </u>			(1/1	DATE / T	IME	COURIER	,	and define the desired state of the second sta		
RELINQUISHED BY	1. o sour		10/12/	DATE/	TIME	RI	ECEIVED	CIAA):	<b>U</b>	<u></u>	KYT <b>()</b>	<i>L/-</i> -k_	 + ' + '	DATE/T	CX	AIRBILL NO	)			