

# QUARTERLY MONITORING REPORT

UNOCAL Service Station No. 3690 14999 Farnsworth Street San Leandro, CA



2140 WEST WINTON AVENUE HAYWARD, CALIFORNIA 94545

(510) 352-4800

April 15, 1992

Unocal Corporation P.O. Box 5155 San Ramon, California 94583

Attn:

Mr. Robert A. Boust

Re:

QUARTERLY MONITORING REPORT

Unocal Service Station No. 3690

14999 Farnsworth Street San Leandro, California

#### Gentlemen:

This Quarterly Monitoring Report has been prepared by GeoStrategies Inc. (GSI) and presents the results of the 1992 first quarter sampling for the above referenced site (Plate 1).

There are currently three monitoring wells at the site; Wells U-1, U-2 and U-3 (Plate 2). These wells were installed in 1991 by GSI.

#### CURRENT QUARTER SAMPLING RESULTS

Depth to water-level measurements were obtained in each monitoring well on February 12, 1992. Static ground-water levels were measured from the surveyed top of the well box and recorded to the nearest  $\pm 0.01$  foot. Water-level elevations were referenced to Mean Sea Level (MSL) datum and are presented in Table 1. Water-level data were used to construct a quarterly potentiometric map (Plate 3). Shallow ground-water flow direction is to the south with an approximate hydraulic gradient of 0.004.

Each well was checked for the presence of floating product. Floating product was not observed in the wells this quarter.

Unocal Corporation April 15, 1992 Page 2

Ground-water samples were collected on February 12, 1992. Samples were analyzed for Total Petroleum Hydrocarbons calculated as Gasoline (TPH-Gasoline), according to EPA Method 8015 (Modified), for Benzene, Toluene, Ethylbenzene, Xylenes (BTEX) according to EPA Method 8020. The ground-water samples were analyzed by Sequoia Analytical, a California State-certified laboratory located in Redwood City, California. The laboratory analytical report and Chain-of-Custory form is included in Appendix A. These data are summarized and included with the historical chemical analytical data presented in Table 2. A chemical concentration map for benzene is presented on Plate 4. Groundwater sampling field methods and procedures are included in Appendix B.

If you have any questions, please call.

GeoStrategies Inc., by,

Ellen C. Fostersmith

Geologist

Diane M. Lundquist, P.E.

Senior Engineer

C 46725

ECF/DML/dls

Plate 1. Vicinity Map Plate 2. Site Plan

Plate 3. Potentiometric Map

Plate 4. Benzene Concentration Map

Appendix A: Laboratory Analytical Report and Chain-of-Custody form

NO. C46725

Appendix B: Field Methods and Procedures

QC Review:

**TABLES** 

TABLE 1

#### FIELD MONITORING DATA

							·					
WELL NO.	MONITORING DATE	CASING DIA.	TOTAL WELL DEPTH (FT)	WELL ELEV.	DEPTH TO WATER (FT)	PRODUCT THICKNESS (FT)	STATIC WATER ELEV. (FT)	PURGED WELL VOLUMES	pH	TEMPERATURE (F)	CONDUCTIVITY (u MHOS/CM)	
U-1	12-Feb-92	2	30.2	17.24	8.97		8.27	5	7.29	66.0	692	
U-2	12-Feb-92	2	30.6	16.85	8.49		8.36	3	7.29	65.7	506	
U-3	12-Feb-92	2	30.3	17.76	9.27		8.49	5	7.25	66.6	832	

Notes: 1. Static water elevations referenced to Mean Sea Level (MSL).

<sup>2.</sup> Physical parameter measurements represent stabilized values.

SAMPLE Date	SAMPLE POINT	TPH-G (PPB)	BENZENE (PPB)	TOLUENE (PPB)	ETHYLBENZENE (PPB)	XYLENES (PPB)	OIL & GREAS (PPM)
\$ <b>22</b> =========	=========				=======================================	=======	=========
30-Sep-91	U-1	<30	<0.30	<0.30	<0.30	<0.30	N/A
12-Feb-92	U-1	<30	<0.30	<0.30	<0.30	<0.30	N/A
30-Sep-91	U-2	<30	<0.30	<0.30	<0.30	<0.30	N/A
12-Feb-92	U-2	<30	<0.30	<0.30	<0.30	<0.30	N/A
30-Sep-91	u-3	<30	<0.30	<0.30	<0.30	<0.30	<5.0
12-Feb-92	U-3	<30	1.7	<0.30	<0.30	<0.30	N/A

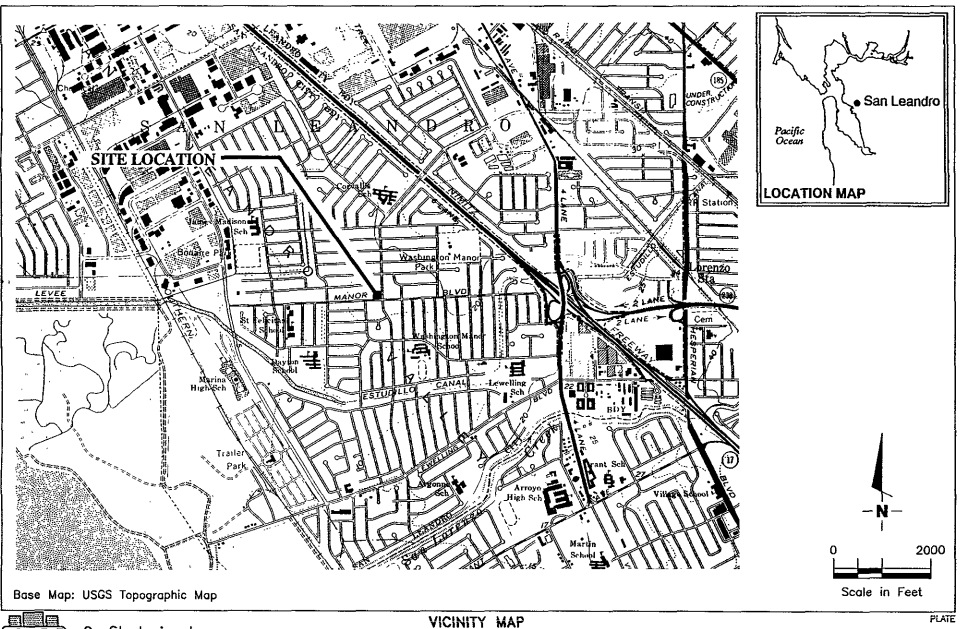
TPH-G = Total Petroleum Hydrocarbons calculated as Gasoline

PPB = Parts Per Billion

NOTE 1. All data shown as <X are reported as ND (none detected).

HVOC?

**ILLUSTRATIONS** 



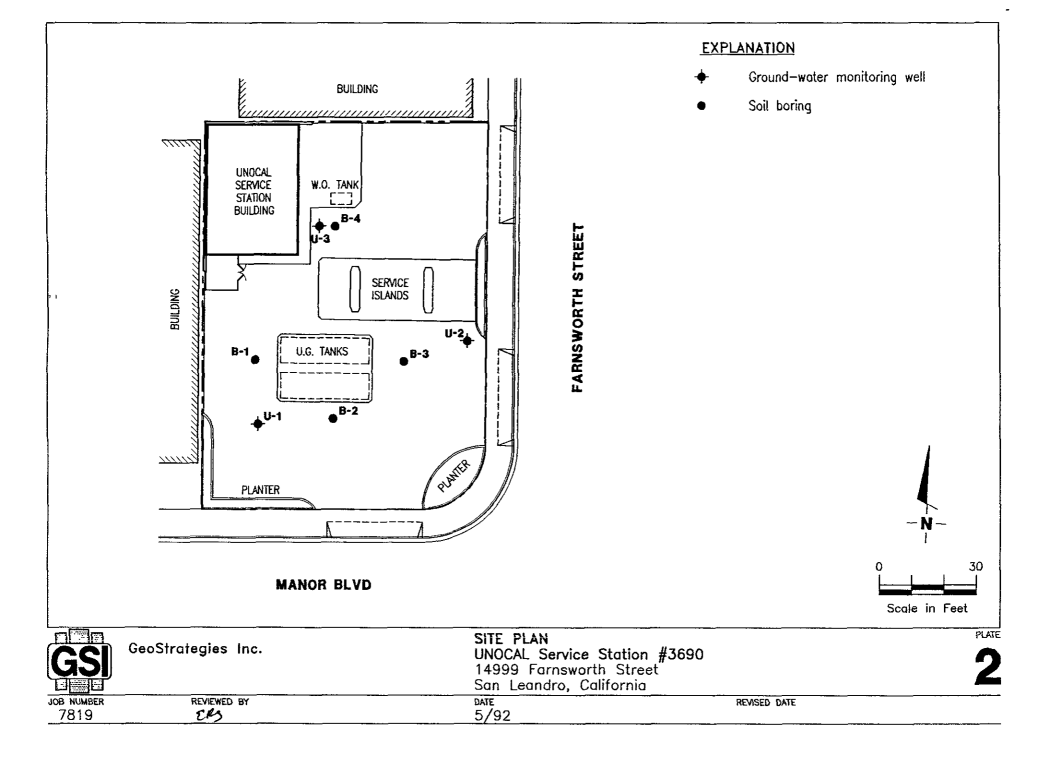
UNOCAL Service Station #3690 14999 Farnsworth Street San Leandro, California

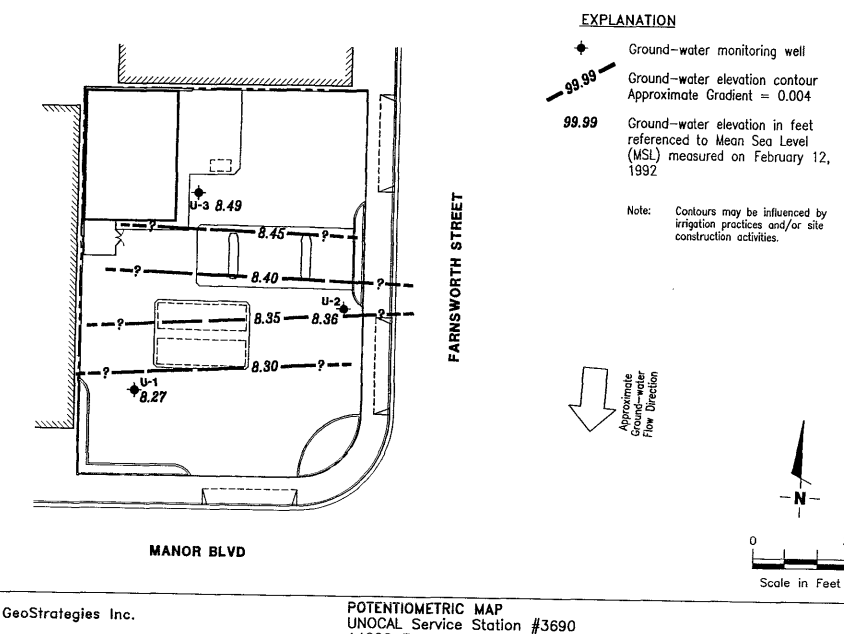
DATE 5/91

JOB NUMBER 7819

REVIEWED BY EUS

REVISED DATE





GSI

JOB NUMBER REVIEWED BY 781902-3 CFS

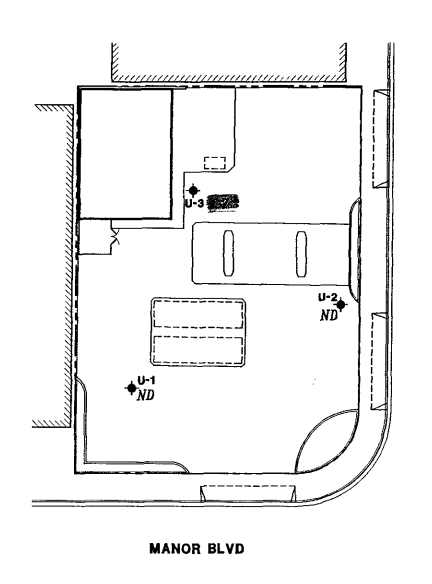
POTENTIOMETRIC MAP UNOCAL Service Station #3690 14999 Farnsworth Street San Leandro, California

DATE 5/92

REVISED DATE

PLATE

30



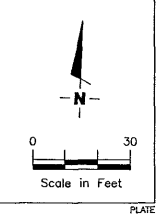
# **EXPLANATION**

Ground-water monitoring well

0.05 concentration in ppb sampled on February 12, 1992

ND Not Detected (See laboratory reports for detection limits)

FARNSWORTH STREET



GeoStrategies Inc.

BENZENE CONCENTRATION MAP UNOCAL Service Station #3690 14999 Farnsworth Street San Leandro, California

DATE

REVISED DATE

JOB NUMBER

REVIEWED BY EKS

5/92

781902-3

# APPENDIX A LABORATORY ANALYTICAL REPORT AND CHAIN-OF-CUSTODY FORM





# GETTLER-RYAN INC.

#### GENERAL CONTRACTORS

Gettler Ryan 2150 W. Winton Avenue Hayward, CA 94545 : Attention: John Zwierzycki adologija is is manistra in 1965. Gasaci

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

#3819.01, Unocal, #3690, San Leandro Water, U-1 **EPA 8240** 202-2385

Sampled: Received: Analyzed: Reported:

Feb 12, 1992 Feb 13, 1992 Feb 25, 1992 Feb 27, 1992

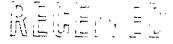
# **VOLATILE ORGANICS by GC/MS (EPA 8240)**

Analyte	Detection Limit µg/L		Sample Results µg/L
Acetone	10	***************************************	N.D.
Benzene	2.0	***************************************	N.D.
Bromodichloromethane	2.0	***************************************	N.D.
Bromoform	2.0		N.D.
Bromomethane	2.0		N.D.
2-Butanone	10	***************************************	N.D.
Carbon disulfide	2.0	***************************************	N.D.
Carbon tetrachloride	2.0	***************************************	N.D.
Chlorobenzene	2.0	***************************************	N.D.
Chloroethane	2.0	***************************************	N.D.
2-Chloroethyl vinyl ether	10	****************************	N.D.
Chloroform	2.0	•••••	N.D.
Chloromethane	2.0		N.D.
Dibromochloromethane	2.0	***************************************	N.D.
1,1-Dichloroethane	2.0	***************************************	N.D.
1,2-Dichloroethane	2.0	••••	N.D.
1,1-Dichloroethene	2.0	•••••	N.D.
cis-1,2-Dichloroethene	2.0	***************************************	N.D.
trans-1,2-Dichloroethene	2.0	***************************************	N.D.
1,2-Dichloropropane	2.0	••••	N.D.
cis-1,3-Dichloropropene	2.0	***************************************	N.D.
trans-1,3-Dichloropropene	2.0	***************************************	N.D.
Ethylbenzene	2.0		N.D.
2-Hexanone	10		N.D.
Methylene chloride	5.0	***************************************	N.D.
4-Methyi-2-pentanone	10	***************************************	N.D.
Styrene	2.0		N.D.
1,1,2,2-Tetrachloroethane	2.0	***************************************	N.D.
Tetrachioroethene	2.0		N.D.
Toluene	2.0		N.D.
1,1,1-Trichloroethane	2.0	***************************************	N.D.
1,1,2-Trichloroethane	2.0	***************************************	N.D.
Trichloroethene	2.0	***************************************	N.D.
Trichlorofluoromethane	2.0	***************************************	N.D.
Vinyl acetate	2.0	***************************************	N.D.
Vinyl chloride	2.0	***************************************	N.D.
Total Xylenes	2.0	***************************************	N.D.

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

SEQUOIA ANALYTICAL

Vickie Tague Project Manager





# GETTLER-RYAN INC

GENERAL CONTRACTORS

Gettler Ryan 2150 W. Winton Avenue Hayward, CA 94545 Attention: John Zwierzycki

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Client Project ID: Sample Descript:

#3819.01, Unocal, #3690, San Leandro Water, U-2

Sampled: Received:

Feb 12, 1992 Feb 13, 1992 Feb 25, 1992

Analysis Method: Lab Number:

**EPA 8240** 202-2386

Analyzed: Reported:

Feb 27, 1992

# **VOLATILE ORGANICS by GC/MS (EPA 8240)**

Analyte	Detection Limit µg/L		Sample Results µg/L
Acetone	10	******************************	N.D.
Benzene	2.0		N.D.
Bromodichloromethane	2.0	****************************	N.D.
Bromoform	2.0	*******************************	N.D.
Bromomethane	2.0	*******************************	N.D.
2-Butanone	10	*****************************	N.D.
Carbon disulfide	2.0	*******************************	N.D.
Carbon tetrachloride	2.0	****************************	N.D.
Chlorobenzene	2.0		N.D.
Chioroethane	2.0	*************************	N.D.
2-Chloroethyl vinyl ether	10	************************	N.D.
Chloroform	2.0	*******************	N.D.
Chloromethane	2.0	*******************************	N.D.
Dibromochloromethane	2.0	***************************************	N.D.
1,1-Dichloroethane	2.0	********************	N.D.
1,2-Dichloroethane	2.0	*************************	N.D.
1,1-Dichloroethene	2.0	*************************	N.D.
cis-1,2-Dichloroethene	2.0	****************************	N.D.
trans-1,2-Dichloroethene	2.0	*******************************	N.D.
1,2-Dichloropropane	2.0	***************************************	N.D.
cis-1,3-Dichloropropene	2.0	***************************************	N.D.
trans-1,3-Dichloropropene	2.0	*******************************	N.D.
Ethylbenzene	2.0	***************************************	N.D.
2-Hexanone	10	***************************************	N.D.
Methylene chloride	5.0	*******************************	N.D.
4-Methyl-2-pentanone	10	*******************************	N.D.
Styrene	2.0	***************************************	N.D.
1,1,2,2-Tetrachioroethane	2.0	***************************************	N.D.
Tetrachloroethene	2.0	***************************************	N.D.
Toluene	2.0	*************************	N.D.
1,1,1-Trichloroethane	2.0	*********************	N.D.
1,1,2-Trichloroethane	2.0	*****************	N.D.
Trichloroethene	2.0	***************	N.D.
Trichlorofluoromethane	2.0	***************************************	N.D.
Vinyl acetate	2.0	***************************************	N.D.
Vinyl chloride	2.0		N.D.
Total Xylenes	2.0	***************************************	N.D.

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

<del>Vi</del>ckie Tague Project Manager



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# GETTLER-RYAN INC

GENERAL CO. That is

Gettler Ryan 2150 W. Winton Avenue Hayward, CA 94545 & Attention: John Zwierzycki

Client Project ID: Sample Descript: Analysis Method:

Lab Number:

#3819.01, Unocal, #3690, San Leandro Water, U-3 **EPA 8240** 202-2387

Sampled: Feb 12, 1992 Received: Feb 13, 1992 Analyzed: Feb 25, 1992

Reported: Feb 27, 1992

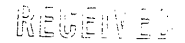
# **VOLATILE ORGANICS by GC/MS (EPA 8240)**

Analyte	Detection Limit μg/L		Sample Results
Acetone	10	•••••	N.D.
Benzene	2.0	***************************************	N.D.
Bromodichloromethane	2.0	*******************************	N.D.
Bromoform	2.0	***************************************	N.D.
Bromomethane	2.0	***************************************	N.D.
2-Butanone	10	***************************************	N.D.
Carbon disulfide	2.0		N.D. N.D.
Carbon tetrachloride	2.0	*****************************	
Chlorobenzene	2.0	******************************	N.D.
Chioroethane	2.0 2.0	******************************	N.D.
2-Chloroethyl vinyl ether	2.0 10	***************************************	N.D.
Chioroform	2.0	*****************************	N.D.
Chloromethane		****************************	N.D.
Dibromochloromethane	2.0	•	N.D.
1/4-Dichloroethane	2.0	***************************************	N.D.
1,2-Dichloroethane			
1.2-Dichloroothone	2.0	***************************************	N.D.
1,1-Dichloroethene	2.0		N.D.
cis-1,2-Dichloroethene	2.0	*******************************	N.D.
trans-1,2-Dichloroethene	2.0	**************************	N.D.
1,2-Dichloropropane	2.0	**************	N.D.
cis-1,3-Dichloropropene	2.0	***************************************	N.D.
trans-1,3-Dichloropropene	2.0	******************************	N.D.
Ethylbenzene	2.0	***************************************	N.D.
2-Hexanone	10	•	N.D.
Methylene chloride	5.0	***************************************	N.D.
4-Methyl-2-pentanone	10	***************************************	N.D.
Styrene	2.0	***************************************	N.D.
1,1,2,2-Tetrachioroethane	2.0	******************************	N.D.
Tetrachloroethene	2.0	***************************************	N.D.
Toluene	2.0	*************	N.D.
1/1,1-Trichloroethane	2.0	*****	
1,1,2-Trichloroethane	2.0	***************************************	N.D.
Trichloroethene	2.0	*************************	N.D.
Trichlorofluoromethane	2.0	************************	N.D.
Vinyl acetate	2.0	••••••	N.D.
Vinyl chloride	2.0	***************************************	N.D.
Total Xylenes	2.0	***************************************	N.D.

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

SEQUOIA ANALYTICAL

√Vickie Tague Project Manager





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# GETTLER-RYAN INC.

GENERAL CO HUAGIORI

Gettier Ryan \$2150 W. Winton Avenue Hayward, CA 94545 Attention: John Zwierzycki

Client Project ID: Sample Descript: Analysis Method:

Lab Number:

#3819.01, Unocal, #3690, San Leandro Water, Trip Blank **EPA 8240** 

202-2388

Sampled

Feb 12, 1992 Received: Feb 13, 1992 Feb 26, 1992

Analyzed: Reported: Feb 27, 1992 

# **VOLATILE ORGANICS by GC/MS (EPA 8240)**

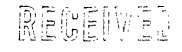
Analyte	Detection Limit µg/L		Sample Results  µg/L
Acetone	10	*************	N.D.
Benzene	2.0	*************************	N.D.
Bromodichloromethane	2.0	************	N.D.
Bromoform	2.0	***************************************	N.D.
Bromomethane	2.0	***********	N.D.
2-Butanone	10	***************************************	N.D.
Carbon disulfide	2.0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N.D.
Carbon tetrachloride	2.0	***************************************	N.D.
Chlorobenzene	2.0	***************************************	N.D.
Chloroethane	2.0	*******************************	N.D.
2-Chloroethyl vinyl ether	10	***************************************	N.D.
Chloroform	2.0	***************************************	N.D.
Chioromethane	2.0	***************************************	N.D.
Dibromochioromethane	2.0	***************************************	N.D.
1,1-Dichloroethane	2.0	***************************************	N.D.
1,2-Dichloroethane	2.0	***************************************	N.D.
1,1-Dichloroethene	2.0	***************************************	N.D.
cis-1,2-Dichioroethene	2.0		N.D. N.D.
trans-1,2-Dichloroethene	2.0	*************************	N.D. N.D.
1,2-Dichloropropane	2.0	******************************	N.D. N.D.
cis-1,3-Dichioropropene	2.0	**************************	N.D.
trans-1,3-Dichloropropene	2.0	***********************	
Ethylbenzene	2.0	***************************************	N.D.
2-Hexanone	10	*******************************	N.D.
		1-1	N.D.
Methylene chloride	10	***************************************	N.D.
4-Methyl-2-pentanone	10	***************************************	N.D.
Styrene	2.0	***************************************	N.D.
1,1,2,2-Tetrachloroethane	2.0	***************************************	N.D.
Tetrachloroethene	2.0	***************************************	N.D.
Toluene	2.0	***************************************	N.D.
1,1,1-Trichloroethane	2.0	***************************************	N.D.
1,1,2-Trichloroethane	2.0	*************************	N.D.
Trichloroethene	2.0	***************************************	N.D.
Trichlorofluoromethane	2.0	***************************************	N.D.
Vinyl acetate	2.0	***************************************	N.D.
Vinyl chloride	2.0	***************************************	N.D.
Total Xylenes	2.0	***************************************	N.D.

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

SEQUOIA ANALYTICAL

**Project Manager** 





FTD 77, 1992

# GETTLER-RYAN INC.

GENERAL CONTILAC OT

Gettler Ryan

Client Project ID:

#3819.01, Unocal, #3690, San Lean

Sampled:

Feb 12, 1992

2150 W. Winton Avenue :Hayward, CA 94545

Matrix Descript:

Water

Received:

Feb 13, 1992 Feb 20, 1992

Attention: John Zwierzycki

Analysis Method:

EPA 5030/8015/8020

Analyzed:

First Sample #: ara era de l'accesso l'experi 202-2385

Reported: 201486918752751882674894878866667666666111119667613888

Feb 27, 1992

# TOTAL PETROLEUM FUEL HYDROCARBONS with BTEX DISTINCTION (EPA 8015/8020)

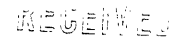
Sample Number	Sample Description	Low/Medium B.P. Hydrocarbons μg/L (ppb)	Benzene μg/L (ppb)	Toluene μg/L (ppb)	Ethyl Benzene µg/L (ppb)	<b>Xylenes</b> μg/L (ppb)
202-2385	U-1	N.D.	N.D.	N.D.	N.D.	N.D.
202-2386	U-2	N.D.	N.D.	N.D.	N.D.	N.D.
202-2387	บ-3	N.D.	1.7	N.D.	N.D.	N.D.
202-2388	Trip Blank	N.D.	N.D.	N.D.	N.D.	N.D.

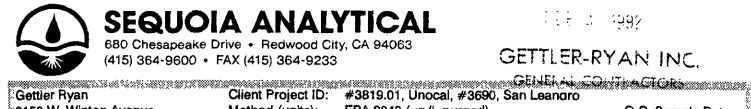
Detection Limits:	30	0.30	0.30	0.30	0.30

Low to Medium Boiling Point Hydrocarbons are quantitated against a gasoline standard. Analytes reported as N.D. were not present above the stated limit of detection.

SEQUOIA ANALYTICAL

Project Manager





1 EF 1 1992 GETTLER-RYAN INC.

Gettier Ryan

2150 W. Winton Avenue

Hayward, CA 94545 Attention: John Zwierzycki

Method (units): EPA 8240 (µg/L purged)

Analyst(s): QC Sample #:

M.Williams

202-3334

Q.C. Sample Dates

Analyzed: Feb 25, 1992

Reported: Feb 27, 1992 Portograman (LT). Literato especies lucas el com la les especies de la compositació de la compositació de la c

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

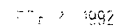
Analyte	Sample Conc.	Spike Conc. Added	Conc. Matrix Spike	Matrix Spike % Recovery	Conc. Matrix Spike Duplicate	Matrix Spike Duplicate % Recovery	Relative % Difference	
1,1-Dichloro- ethene	N.D.	50	43	86	44	88	2.3	_
Trichloroethene	N.D.	50	46	92	47	94	2.2	
Benzene	N.D.	50	45	90	47	94	4.3	
Toluene	N.D.	50	45	90	47	94	4.3	
Chlorobenzene	N.D.	50	49	98	49	98	0.0	

SEQUOIA ANALYTICAL

Vickie Tague Project Manager

% Recovery:	Conc. of M.S Conc. of Sample	x 100	
	Spike Conc. Added		
Relative % Difference:	Conc. of M.S Conc. of M.S.D.	x 100	
	(Conc. of M.S. + Conc. of M.S.D.) / 2		





# GETTLER-RYAN INC.

GENERAL CONTRACTORS

Gettler Ryan

Client Project ID: #3819.01, Unocal, #3690, San Leandro

2150 W. Winton Avenue Hayward, CA 94545

Attention: John Zwierzycki

QC Sample Group: 2022385 - 88

Reported:

Feb 27, 1992 

# **QUALITY CONTROL DATA REPORT**

ANALYTE			Ethyl-	
	Benzene	Toluene	Benzene	Xylenes
Method: Analyst: Reporting Units: Date Analyzed: QC Sample #:	EPA 8020 M.Nipp μg/L Feb 20, 1992 GBLK022092	EPA 8020 M.Nipp μg/L Feb 20, 1992 GBLK022092	EPA 8020 M.Nipp µg/L Feb 20, 1992 GBLK022092	EPA 8020 M.Nipp µg/L Feb 20, 1992 GBLK022092
Sample Conc.:	N.D.	N.D.	N.D.	N.D.
Spike Conc. Added:	10	10	10	30
Conc. Matrix Splke:	9.0	8.9	8.9	27
Matrix Spike % Recovery:	90	89	89	90
Conc. Matrix Spike Dup.:	8.5	8.6	ઇ.4	26
Matrix Spike Duplicate % Recovery:	85	86	84	87
Relative % Difference:	5.1	3.4	5.8	3.8

vickie Tague Project Manager % Recovery: Conc. of M.S. - Conc. of Sample x 100 Spike Conc. Added Relative % Difference: Conc. of M.S. - Conc. of M.S.D. x 100

(Conc. of M.S. + Conc. of M.S.D.) / 2

2022385.GET <7>

COMPANY	Unocal	? ss #	3690 1	2007	\ /7 JOB	Chain of Custo
JOB LOCATION		Tarns wor	1		Ÿ,	
CITY		candro		· ; · · · · · · · · · · · · · · · · · ·	PHONE NO.	(510) 783-70
AUTHORIZED		v.erzyck:	DATE	2-12-92	P.O. NO	3818.01
SAMPLE ID	NO. OF CONTAINERS	SAMPLE MATRIX	DATE/TIME SAMPLED	ANALYSIS REQ		SAMPLE CONDITION LAB ID
U-1	4	Liquid	2.12-52/0905	EPA 80		2022385
U-2		6/	1,1000			86
U-3		1	10944	'		87
trip		1	- /-	V		2 88
<del></del>						
· · · · · · · · · · · · · · · · · · ·					<u></u>	
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# APPENDIX B

FIELD METHODS AND PROCEDURES

#### FIELD METHODS AND PROCEDURES

#### **EXPLORATION DRILLING**

#### Mobilization

Prior to any drilling activities, GeoStrategies Inc. (GSI) will verify that necessary drilling permits have been secured.

Utility locations will be located and drilling will be conducted so as not to disrupt activities at a project site. GSI will obtain and review available public data on subsurface geology and if warranted, the location of wells within a half-mile of the project site will be identified. Drillers will be notified in advance so that drilling equipment can be inspected prior to performing work.

#### **Drilling**

The subsurface investigations are typically performed to assess the lateral and vertical extent of petroleum hydrocarbons present in soils and groundwater. Drilling methods will be selected to optimize field data requirements as well as be compatible with known or suspected subsurface geologic conditions.

Monitoring wells are installed using a truck-mounted hollow-stem auger drill rig or mud-rotary drill rig. Typically, the hollow-stem rig is used for wells up to 100 feet, if subsurface conditions are Wells greater than 100-feet deep are typically drilled favorable. When mud rotary drilling is used, an using mud-rotary techniques. electric log will be performed for additional lithological Also during mud rotary drilling, precautions will be information. taken to prevent mud from circulating contaminants by using a conductor casing to seal off contaminated zones. Samples will be collected for lithologic logging by continuous chip, and where needed by drive sample or core as specified by the supervising geologist.

#### Soil Sampling

Shallow soil borings will be drilled using a truck-mounted hollow-stem auger drilling rig, unless site conditions favor a different drilling method. Drilling and sampling methods will be consistent with ASTM Method D-1452-80. The auger size will be a minimum 6-inch nominal outside-diameter (O.D). No drilling fluids will be used during this drilling method. The augers and other tools used in the bore hole will be steam cleaned before use and between borings to minimize the possibilities of cross-contamination between borings.

Soil samples are typically collected at 5-foot intervals as a minimum from ground surface to total depth of boring. Additional soil samples will be collected based on significant lithologic changes and/or potential chemical content. Soil samples from each sampling interval will be lithologically described by a GSI geologist (Figure 1). Soil colors will be described using the Munsell Color Chart. Rock units will be logged using appropriate lithologic terms, and colors described by the G.S.A. Rock Color Chart.

Head-space analyses will be performed to check for the evidence of volatile organic compounds. Head-space analyses will be performed using an organic vapor analyzer; either an OVA, HNU, or OVM. Organic vapor concentrations will be recorded on the GSI field log of boring (Figure 1). The selection of soil samples for chemical analysis are typically based on the following criteria:

- 1) Soil discoloration
- 2) Soil odors
- 3) Visual confirmation of chemical in soil
- 4) Depth with respect to underground tanks (or existing grade)
- 5) Depth with respect to ground water
- 6) OVA reading

Soil samples (full brass liners) selected for chemical analysis are immediately covered with aluminum foil and the liner ends are capped to prevent volatilization. The samples are labeled and entered onto a Chain-of-Custody form, and placed in a cooler on blue ice for transport to a State-certified analytical laboratory.

Soil cuttings are stockpiled on-site. Soils are sampled and analyzed for site-specific chemical parameters. Disposition of soils is dependent of chemical analytical results of the samples.

#### Soil Sampling - cont.

Soil borings not converted to monitoring wells will be backfilled (sealed) to ground surface using either a neat cement or cement-bentonite grout mixture. Backfilling will be tremied by continuously pumping grout from the bottom to the top of the boring where depth exceeds 20' or as required by local permit requirements.

All field and office work, including exploratory boring logs, are prepared under the direction of a registered geologist.

#### Monitoring Well Installation

Monitoring well casing and screen will be constructed of Schedule 40, flush-joint threaded polyvinylchloride (PVC). The well screen will be factory mill-slotted unless additional open area is required (eg. conversion to an extraction well in a low-yield aquifer). The screen length will be placed adjacent to the aquifer material to a minimum of 2-feet above encountered water. No screen shall be placed in a borehole that potentially creates hydraulic interconnection of two or more aquifer units. Screen slot size and well sand pack will be compatible with encountered aquifer materials, as confirmed by sieve analysis.

Monitoring wells will be completed below grade (Figure 2) unless special conditions exist that require above-grade completion design. In the event a monitoring well is required in an aquifer unit beneath an existing aquifer, the upper aquifer will be sealed off by installing a steel conductor casing with an annular neat cement or cement-bentonite grout seal. This seal will be continuously tremie pumped from the bottom of the annulus to ground surface.

The monitoring well sand pack will be placed adjacent to the entire screened interval and will extend a recommended minimum distance of 2-feet above the top of the screen. No sand pack will be placed that interconnects two or more aquifer units. A minimum 2-foot bentonite pellet or bentonite slurry seal will be placed above the sand pack. Sand pack, bentonite, and cement seal levels will be confirmed by sounding the annulus with a calibrated weighted tape. The remaining annular space above the bentonite seal will be grouted with a bentonite-cement mixture and will be tremie-pumped from the bottom of the annular space to the ground surface. The bentonite content of the grout will not exceed 5 percent by weight. A field log of boring and a field well completion form will be prepared by GSI for each well installed.

Decontamination of drilling equipment before drilling and between wells will consist of steam cleaning, and/or Alconox wash.

#### Well Development

All newly installed wells will be properly developed within 48 hours of completion. No well will be developed until the well seal has set a minimum of 12 hours. Development procedures will include one or more of the methods described below:

#### Bailing

Bailing will be used to remove suspended sediments and drilling fluids from the well, where applicable. The bailer will be raised and lowered through the column of water in the well so as to create a gentle surging action in the screened interval. This technique may be used in conjunction with other techniques, such as pumping, and may be used alone if the well is of low yield.

#### Pumping

Pumping will be used in conjunction with bailing or surging. The pump will be operated in such a manner as to gently surge the entire screened interval of the well. This may involve operating the pump with a packer type mechanism attached and slowly raising and lowering the pump, or by cycling the pump off and on to allow water to move in and out of the screened interval. Care will be used not to overpump a well.

#### Surging

Surging will be performed on wells that are screened in known or suspected high yield formations and/or on larger diameter (recovery) wells. A surge block will be raised and lowered through the entire screened interval, forcing water in and out of the well screen and sand pack. Pumping or air lifting will be used in conjunction with this method of development to remove any sediment brought into the well during surging.

#### Air Lifting

Air lifting will be used to remove sediment from wells as an alternative to pumping under certain conditions. When appropriate, a surge block designed for use with air lifting will be used to agitate the entire screened interval and water will be lifted out of the well using forced air. When air lifting is performed, the air source will be either nitrogen or filtered air and the procedure will be performed gently to prevent any damage to the well screen or casing and to insure that discharged water is contained.

#### Well Development - cont.

All well developing equipment will be thoroughly decontaminated prior to development using a steam cleaner and/or Alconox detergent wash and clean water rinse. During development procedures, field parameters (temperature, specific conductance and pH) will be monitored and recorded on well development forms (Figure 3). Equilibration requirements consist of a minimum of three readings with the following accuracy standards:

pH ± 0.1 pH units
Specific Conductance ± 10% of full scale reading
Temperature ± 0.5 degrees Celsius

The wells will be developed until water is visibly clear and free of sediment, and well purging parameters stabilized. A minimum of 8 to 10 well volumes will be purged from each well, if feasible. If well purging parameters have not stabilized before 10 casing volumes have been removed, well development will continue until purging parameters have stabilized and formation water is being drawn into the well. The adequacy of well development will be judged by the field technician performing the well development and based on known formation conditions.

#### Well Surveying

Monitoring wells will be surveyed to obtain top of box elevations to the nearest  $\pm 0.01$  foot. Water level measurements will be recorded to the nearest  $\pm 0.01$  foot and referenced to Mean Sea Level (MSL). If additional wells are required, then existing and newly installed wells are surveyed relative to MSL.

#### GROUND-WATER SAMPLING AND ANALYSIS

#### Quality Assurance/Quality Control Objectives

The sampling and analysis procedures employed by Gettler-Ryan Inc. (G-R) for ground-water sampling and monitoring follow specific Quality Assurance/Quality Control (QA/QC) guidelines. Quality Assurance objectives have been established by G-R to develop and implement procedures for obtaining and evaluating water quality and field data in an accurate, precise, and complete manner so that sampling procedures and field measurements provide information that is comparable and representative of actual field conditions. Quality Control (QC) is maintained by G-R by using specific field protocols and requiring the analytical laboratory to perform internal and external QC checks. It is the goal of G-R to provide data that are accurate, precise, complete, comparable, and representative. The definitions for accuracy, precision, completeness, comparability, and representativeness are as follows:

- Accuracy the degree of agreement of a measurement with an accepted referenced or true value.
- <u>Precision</u> a measure of agreement among individual measurements under similar conditions. Usually expressed in terms of the standard deviation.
- <u>Completeness</u> the amount of valid data obtained from a measurement system compared to the amount that was expected to meet the project data goals.
- <u>Comparability</u> expresses the confidence with which one data set can be compared to another.
- Representativeness a sample or group of samples that reflects the characteristics of the media at the sampling point. It also includes how well the sampling point represents the actual parameter variations which are under study.

As part of the G-R QA/QC program, applicable federal, state, and local reference guidance documents are followed. The procedures outlined in these regulations, manuals, handbooks, guidance documents, and journals are incorporated into the G-R sampling procedures to assure that; (1) ground-water samples are properly collected, (2) ground-water samples are identified, preserved, and transported in a manner such that they are representative of field conditions, and (3) chemical analysis of samples are accurate and reproducible.

# Guidance and Reference Documents Used to Collect Groundwater Samples

These documents are used to verify G-R sampling procedures and are consistent with current regulatory guidance. If site specific work and sampling plans are required, those plans will be developed from these documents, and newly received applicable documents.

U.S.E.P.A 330/9-51-002	NEIC Manual for Groundwater/Subsurface Investigation at Hazardous Waste Sites
U.S.E.P.A 530/SW611	Procedures Manual for Groundwater Monitoring at Solid Waste Disposal Facilities (August, 1977)
U.S.E.P.A 600/4-79-020	Methods for Chemical Analysis of Water and Wastes (1983)
U.S.E.P.A 600/4-82-029	Handbook for Sampling and Sample Preservation of Water and Wastewater (1982)
U.S.E.P.A 600/4-82-057	Test Methods for Organic Chemical Analysis of Municipal and Industrial Wastewater (July, 1982)
U.S.E.P.A SW-846#, 3rd Edition	Test Methods for Evaluating Solid Waste - Physical/Chemical Methods (November, 1986)
40 CFR 136.3e, Table II (Code of Federal Regulations)	Required Containers, Preservation Techniques, and Holding Times
Resources Conservation and Recover Act (OSWER 9950.1)	Groundwater Monitoring Technical Enforcement Guidance Document (September, 1986)
California Regional Water Quality Control Board (Central Valley Region)	A Compilation of Water Quality Goals (September, 1988); Updates (October, 1988)
California Regional Water Quality Control Board (North Coast, San Francisco Bay, and Central Valley)	Regional Board Staff Recommendations for Initial Evaluations and Investigation of Underground Tanks: Tri-Regional Recommendations (June,

1988)

#### Guidance and Reference Documents Used to Collect Groundwater Samples (cont.)

Regional Water Quality Control Board (Central Valley Region)

Memorandum: Disposal, Treatment, and Refuse of Soils Contaminated with Petroleum Fractions (August, 1986)

State of California Department of Health Services

Hazardous Waste Testing Laboratory Certification List (March, 1987)

State of California Water Resources Control Board Leaking Underground Fuel Tank (LUFT) Field Manual (May, 1988), and LUFT Field Manual Revision (April, 1989)

State of California Water Resources Control Board Title 23. (Register #85.#33-8-17-85), 16: Underground Subchapter Regulations; Article 3, Sections 2632 and 2634; Article 4, Sections 2645, 2647, and 2648: Article 2646. Sections 2670, 2671. and 2672 1988 (October, 1986: including Amendments)

Alameda County Water District

Groundwater Protection Program: Guidelines for Groundwater and Soil Investigations at Leaking Underground Fuel Tank Sites (November, 1988)

American Public Health Association

Standard Methods for the Examination of Water and Wastewaters, 16th Edition

Analytical Chemistry (journal)

Principles of Environmental Analysis, Volume 55, Pages 2212-2218 (December, 1983)

Napa County

Napa County Underground Storage Tank Program: Guidelines for Site Investigations; February 1989.

Santa Clara Valley Water District

Guidelines for Preparing or Reviewing Sampling Plans for Soil and Groundwater Investigation of Fuel Contamination Sites (January, 1989)

# Guidance and Reference Documents Used to Collect Groundwater Samples (cont.)

Santa Clara Valley Water District

Investigation and Remediation at Fuel Leak sites: Guidelines for Investigation and Technical Report

Preparation (March 1989)

Santa Clara Valley Water District

American Petroleum Institute

Well Standards for Santa Revised Clara County (July 18, 1989) Groundwater Monitoring & Sample API Publication 4367, Bias: Environmental Affairs Department,

June 1983

American Petroleum Institute

American Petroleum Institute

A Guide Assessment and to the Remediation of Underground Petroleum Releases; API Publication 1628, February 1989

Summary: Hydrocarbon Literature Attenuations Solubilities and API Publication 4414. Mechanisms,

August 1985

Site Specific (as needed)

specific regulatory General and documents as required.

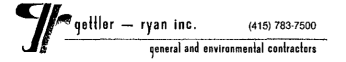
Because ground-water samples collected by G-R are analyzed to the parts per billion (ppb) range for many compounds, extreme care is exercised to prevent contamination of samples. When volatile or semi-volatile organic compounds are included for analysis, G-R sampling crew members will adhere to the following precautions in the field:

- 1. A clean pair of new, disposable gloves are worn for each well being sampled.
- 2. When possible, samples are collected from known or suspected wells that are least contaminated (i.e. background) followed by wells in increasing order of contamination.
- 3. Ambient conditions are continually monitored to maintain sample integrity.

When known or potential organic compounds are being sampled for, the following additional precautions are taken:

- 1. All sample bottles and equipment are kept away from fuels and solvents. When possible, gasoline (used in generators) is stored away from bailers, sample bottles, purging pumps, etc.
- 2. Bailers are made of Teflon or Stainless Steel. Other materials such as plastic may contaminate samples with phthalate esters which interfere with many Gas Chromatography (GC) analyses.
- 3. Volatile organic ground-water samples are collected so that air passage through the sample does not occur or is minimal (to prevent volatiles from being stripped from the samples): sample bottles are filled by slowly running the sample down the side of the bottle until there is a positive convex meniscus over the neck of the bottle; the Teflon side of the septum (in cap) is positioned against the meniscus, and the cap screwed on tightly; the sample is inverted and the bottle lightly tapped. The absence of an air bubble indicates a successful seal; if a bubble is evident, the cap is removed, more sample is added, and the bottle is resealed.
- 4. Extra Teflon seals are brought into the field in case seals are difficult to handle and/or are dropped. Dropped seals are considered contaminated and are not used. When replacing seals or if seals become flipped, care is taken to assure that the Teflon seal faces down.

Sample analysis methods, containers, preservatives and holding times are shown on Table 1.



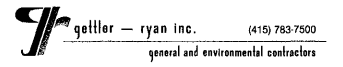
Laboratory and field handling procedures of samples are monitored by including QC samples for analysis with every submitted sample lot from a project site. QC samples may include any combination of the following:

- A. <u>Trip Blank</u>: Used for purgeable organic compounds only; QC samples are collected in 40 milliliter (ml) sample vials filled in the analytical laboratory with organic-free water. Trip blanks are sent to the project site, and travel with project site samples. Trip blanks are not opened, and are returned from a project site with the project site samples for analysis.
- B. <u>Field Blank</u>: Prepared in the field using organic-free water. These QC samples accompany project site samples to the laboratory and are analyzed for specific chemical parameters unique to the project site where they were prepared.
- C. <u>Duplicates</u>: Duplicated samples are collected "second samples" from a selected well and project site. They are collected as either split samples or second-run samples collected from the same well.
- D. <u>Equipment Blank</u>: Periodic QC sample collected from field equipment rinsate to verify decontamination procedures.

The number and types of QC samples are determined as follows:

- A. Up to 2 wells Trip Blank Only
- B. 2 to 5 Wells 1 Field Blank and 1 Trip Blank
- C. 5 to 10 Wells 1 Field blank, 1 Trip Blank, and 1 Duplicate
- D. More than 10 Wells 1 Field Blank, 1 Trip Blank, and 1 Duplicate per each 12 wells
- E. If sampling extends beyond one day, quality control samples will be collected for each day.

Additional QC is performed through ongoing and random reviews of duplicate samples to evaluate the precision of the field sampling procedures and analytical laboratory. Precision of QC data is accomplished by calculating the Relative Percent Difference (RPD). The RPD is evaluated to assess whether values are within an acceptable range (typically ± 20% of duplicate sample).



#### SAMPLE COLLECTION

This section describes the routine procedures followed by G-R while collecting ground-water samples for chemical analysis. These procedures include decontamination, water-level measurements, well purging, physical parameter measurements, sample collection, sample preservation, sample handling, and sample documentation. Critical sampling objectives for G-R are to:

- 1. Collect ground-water samples that are representative of the sampled matrix and,
- 2. Maintain sample integrity from the time of sample collection to receipt by the analytical laboratory.

Sample analyses methods, containers, preservation, and holding times are presented in Table 1.

#### Decontamination Procedures

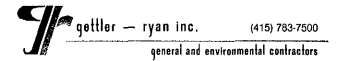
All physical parameter measuring and sampling equipment are decontaminated prior to sample collection using Alconox or equivalent detergent followed by steam cleaning with deionized water. Any sampling equipment surfaces or parts that might absorb specific contaminants, such as plastic pump valves, impellers, etc., are cleaned in the same manner.

Sample bottles, bottle caps, and septa used for sampling volatile organics are thoroughly cleaned and prepared in the laboratory. Sample bottles, bottle caps, and septa are protected from all potential chemical contact before actual usage at a sample location.

During field sampling, equipment placed in a well are decontaminated before purging or sampling the next well. The equipment are decontaminated by cleaning with Alconox or equivalent detergent followed by steam cleaning with deionized water.

#### Water-Level Measurements

Prior to purging and sampling a well, the static-water levels are measured in all wells at a project site using an electric sounder and/or calibrated portable oil-water interface probe (Figure 4). Both static water-level and separate-phase product thickness are measured to the nearest  $\pm 0.01$  foot. The presence of separate-phase product is confirmed using a clean, acrylic or polyvinylchloride (PVC) bailer, measured to the nearest  $\pm 0.01$  foot with a decimal scale tape.



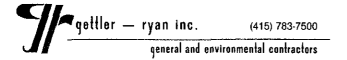
#### Water-Level Measurements (continued)

The monofilament line used to lower the bailer is replaced between possibility new line to preclude the wells with Field observations (e.g. well integrity, product cross-contamination. color, turbidity, water color, odors, etc.) are noted on the G-R Well Sampling Field Data Sheet shown in Figure 4. Before and after each sounder, interface and bailer probe electric decontaminated by washing with Alconox or equivalent detergent rinsing with deionized water followed by cross-contamination.

As mentioned previously, water-levels are measured in wells with known or suspected lowest dissolved chemical concentrations to the highest dissolved concentrations.

#### Well Purging

Before sampling occurs, well casing storage water and interstitial water in the artificial sand pack will be purged using (1) a positive displacement bladder pump constructed of inert, non-wetting, Teflon and stainless steel, (2) a pneumatic-airlift pumping system, (3) a centrifigal pumping system, or (4) a Teflon or Stainless steel bailer (Figure 5). Methods of purging will be assessed based on well size. location, accessibility, and known chemical conditions. well purge volumes are calculated from borehole volumes which take into account the sand packed interval in the well annular space. As a general rule, a minimum of 3 and a maximum of 10 borehole volumes will be purged. Wells which dewater or demonstrate slow recharge periods (i.e. low-yield wells) during purging activities may be sampled after fewer purging cycles. If a low-yield (low recovery) well is to be sampled, sampling will not take place until at least 80 percent of the previously measured water column has been replaced by recharge, or as per local requirements. Physical parameter measurements (temperature, pH, and specific conductance) are closely monitored throughout the well purging process and are used by the G-R sampling crew as indicators for assessing sufficient purging. Purging is continued physical parameters have stabilized. Specific until all three read to the nearest conductance (conductivity) meters are umhos/cm, and are calibrated daily. pH meters are read to the nearest ±0.1 pH units and are calibrated daily. Temperature is read to the nearest 0.1 degree F. Calibration of physical parameter meters will follow manufacturers specifications. Monitoring wells will be purged according to the protocol presented in Figure 5. Collected field data during purging activities will be entered on the G-R Well Sampling Field Data Sheet shown in Figure 4. Copies of the G-R Field Data Sheets will be reviewed by the G-R Sampling Manager for accuracy and completeness.



#### **DOCUMENTATION**

#### Sample Container Labels

Each sample container will be labeled by an adhesive label, noted in permanent ink immediately after the sample is collected. Label information will include:

Sample point designation (i.e. well number or code)

Sampler's identification

Project number

Date and time of collection

Type of preservation used

#### Well Sampling Data Forms

In the field, the G-R sampling crew will record the following information on the Well Sampling Data Sheet for each sample collected:

Project number

Client

Location

Source (i.e. well number)

Time and date

Well accessibility and integrity

Pertinent well data (e.g. depth, product thickness, static water-level, pH, specific conductance, temperature)

Calculated and actual purge volumes

#### Chain-of-Custody

A Chain-of-Custody record (Figure 6) shall be completed and accompany every sample and every shipment of samples to the analytical laboratory in order to establish the documentation necessary to trace sample possession from time of collections. The record will contain the following information:

- Sample or station number or sample identification (ID)
- Signature of collector, sampler, or recorder
- Date and time of collection
- Place of collection
- Sample type
- Signatures of persons involved in chain of possession
- Inclusive dates of possession

Samples shall <u>always</u> be accompanied by a Chain-of-Custody record. When transferring the samples, the individual relinquishing and receiving the samples will sign, date, and note the time on the Chain-of-Custody record. G-R will be responsible for notifying the laboratory coordinator when and how many samples will be sent to the laboratory for analysis, and what types of analyses shall be performed.

TABLE 1
SAMPLE ANALYSIS METHOOS, CONTAINERS, PRESERVATIONS, AND HOLDING TIMES

	Analytical	Reporting		_	Maximum Holding
Parameter	Method	Units	Container	Preservation	Time
Total Petroleum	EPA 8015	l\gm	40 ml. viel	cool, 4 C	14 days (maximum)
Hydrocarbons (Gasoline)	(modified)	ug/l	glass, Teflon	HCl to pH<2	
Benzene	EPA 8020	mg/l	50 ml. viat	cool, 4 C	7 days (w/o preservative)
Toluene		ug/l	glass, Tefion	HCl to pH<2	14 days (w preservative)
Ethylbenzene Xylenes (BTEX			lined septum		
Oil & Grease	SM 503E	mg/l	1 l glass, Teflon	H2SO4 or HCl	28 days (maximum)
		ug/l	lined septum	to pH<2	
Total Petroleum	EPA 8015	mg/l	40 ml. vial	cool, 4 C	14 days (maximum)
Hydrocarbons	(modified)	ug/l	glass, Teflon		
(Diesel)			lined septum		
Halogented	2010	mg/l	40 ml. vial	cool, 4 C	14 days (maximum)
Volatile Organics		∪g/l	glass, Teflon		
(chlorinated solvents)			lined septum		
Non chlerinated	8020	mg/l	40 ml. vial	cool, 4 C	14 days (maximum)
solvents		ug/l	glass, Teflon lined septum	HCl to pH<2	
Volatile Organics	8240	mg/l	40 ml. vial	cool, 4 C	14 days (maximum)
		ug/l	glass, Teflon	HCl to pH<2	
•			lined septum	75	
Semi-Volatile	8270	mg/l	1 l amber	cool, 4 C	7 days extract
Organics		ug/l	glass, Teflon lined septum		40 days (maximum to analyze)
Specific		umhos/cm			
Conductance					
(Field test)					
pH (Field test)		pH units			
Temperature		Deg F			
(Field test)					

# GETTLER-RYAN INC.

# General and Environmental Contractors

# WELL SAMPLING FIELD DATA SHEET

FIGURE 4

COMPANY	JOB #				
·					
Well ID.	Well Condition				
Well Diameter	in. Hydrocarbon Thickness	ft.			
	Tt. Volume 2" = 0.17 6" = 1.5	50 12" = 5.80 80 10			
(# of casing volumes) x	(Fatiment and	gal.			
Purging Equipment					
Sampling Equipment					
Starting Time	Purging Flow Rate				
Æstimated\	/Purging\ _ /Anticipated	gpm min.			
(Purge yolume) gal.	Rate / Time	)			
Time p	H Conductivity Temperature	Volume			
Did well dewater?	Volum	e			
Sampling Time	Weather Conditions	and a desired fill a control of the			
Analysis	Bottles Used				
Chain of Custody Number_					
COMMENTS					

#### Sampling Crew Reviews Project Sampling Requirements/Schedule Field Decontamination and Instrumentation Calibration Check Integrity of Well (Inspect for Well Damage) Measure and Record Depth to Water and Total Well Depth (Electric Well Sounder) Check for Floating Product (Oil/Water Interface Probe) Floating Product Present Floating Product Not Present Purge Volume Calculation Confirm Product Thickness (Acrylic or PVC Bailer) $V = \pi (r/12)^{2} h(_{x} \text{ vol})(7.48) = ___/gallons$ V = Purge volume (gailons) Collect Free-Product Sample $\tau = 3.14159$ h = Height of Water Column (feet) Dissolved Product Sample Not r = Borehole radius (inches) Required Record Data on Field Data Form Evacuate water from well equal to the calculated purge volume while monitoring groundwater stabilization indicator parameters (pH, conductivity, temperature) at intervals of one casing volume. Well Readily Recovers Well Dewaters after One Purge Volume (Low yield well) Well Recharges to 80% of Initial Record Groundwater Stability Indicator Parameters from each Additional Purge Volume Measured Water Column Reight in Feet within 24 hrs. of Evacuation. Stability indicated when the following Criteria are met: pH : + 0.1 pH units Measure Groundwater Stability Indicator Parameters (pH, Temperature, Conductivity) Conductivity: ± 10% Temperature: 1.0 degrees F Groundwater Stability Achieved Groundwater Stability Not Achieved Collect Sample and Complete Chain-of-Custody Collect Sample and Complete Continue Purging Until Stability Chain-of-Custody is Achieved Preserve Sample According to Required Preserve Sample According Collect Sample and complete Chain-of-Custody Chemical Analysis to Required Chemical Analysis Preserve Sample According to Required Chemical Analysis Transport to Analytical Laboratory Transport to Analytical Laboratory Transport to Analytical Laboratory

Monitoring Well Sampling Protocol Schematic

Gettler - R	yan inc	E N	VIRONMENTAL DI	/ISION	Chain of Custody FIGURE 6
COMPANY				J	
JOB LOCATION					
CITY		· · · · · · · · · · · · · · · · · · ·		PHONE N	0
AUTHORIZED			DATE _	P.O. NO.	
SAMPLE ID	NO. OF CONTAINERS	SAMPLE MATRIX	DATE/TIME SAMPLED	ANALYSIS REQUIRED	SAMPLE CONDITION LAB ID
				•	
			,		
RELINQUISHED BY	<u>':</u>	1	RECE	EIVED BY:	
RELINQUISHED BY	<b>:</b>			EIVED BY:	
RELINQUISHED BY			RECE	EIVED BY LAB:	
DESIGNATED LABO	ORATORY;		·····	DHS #:	············
REMARKS:					
DATE COMPLETED			FORE	MAN	
				•	

2140 WEST WINTON AVENUE HAYWARD, CALIFORNIA 94545 2312218 23198

(510) 352-4800

April 15, 1992

Alameda County Health Agency Division of Hazardous Materials Department of Environmental Health 80 Swan Way, Room 200 Oakland, California 94521

Attention:

Ms. Pamela Evans

Reference:

UNOCAL Service Station No. 3690

14999 Farnsworth Street San Leandro, California

Ms. Evans:

As requested by Mr. Robert Boust of UNOCAL Corporation, we are forwarding a copy of the Quarterly Monitoring Report for the above referenced location. This report presents the results of the 1992 first quarter groundwater sampling conducted at this site.

If you have any questions or comments, please call.

Sincerely,

David J. Vossler Senior Geologist

Jan Wasalu

DJV/cmg

Enclosure

cc:

Mr. Robert Boust, UNOCAL Corporation

Mr. Richard Hiett, Regional Water Quality Control Board