



**SUPERFUND SUPPORT  
SAMPLING INSPECTION REPORT**

ANCHOR CHEMICAL SITE  
Hicksville, Long Island, New York

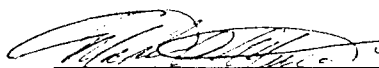
April 4, 1996

Participating Personnel:

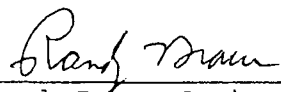
U.S. Environmental Protection Agency

Michael A. Mercado, Environmental Scientist  
Jennifer Fernada, Environmental Scientist  
Robert Morrell, Geologist  
Jennifer Snow-Ashbrook, Environmental Scientist

Report Prepared By:

 4/14/96  
Michael A. Mercado, Environmental Scientist

Approved for the Director By:

 6/14/96  
Randy Braun, Acting Chief,  
Surveillance and Monitoring Branch

REPORT  
ANCHOR CHEMICAL SITE

Sampling Dates:

March 29 thru. April 4, 1996

Objective:

The objective of this sampling event was to determine the effects, resulting from excavation on the site, to the underlying aquifer. The removal of subsurface soils may have had a positive impact by removing the contaminated soils but negative impacts (pollutant transport) may have resulted from subsurface soil disturbance. In order to determine the effects, five (5) monitoring wells were sampled. These samples were screened against Federal MCL standards for VOCs, TCL NVOAs and TAL metals.

Inspection Participants:

Michael Mercado, Environmental Scientist with EPA/ESD/SMB  
Jennifer Feranda, Environmental Scientist with EPA/ESD/SMB  
Jennifer Snow-Ashbrook, Environmental Scientist with EPA/ESD/SMB  
Robert Morrell, Geologist with EPA/ESD/SMB

Sampling Activities:

The ESD sampling team arrived at the Anchor Chemical Site, located on 500 West John Street, Hicksville, Long Island, NY at 6:45 am, March 29, 1996. At 7:00 am, we met with maintenance personnel at the site and started preparing to pull samples. At 8:05 am, we began preparing the equipment blanks and trip blanks. Weather was wet, snowy and icy, with strong winds and temperatures between 20°s and 30°s F.

Due to these weather conditions, the electronic water level indicator and the turbidity meter did not work properly. We measured the water column manually. The measurement was done by lowering the bailer into the well, until it touched the water. It was determined to touch the water when the weight of the bailer lessened. We then marked the cord to which the bailer was attached. Next we measured from the mark on the cord to end of the bailer. This measurement gives us the length from the top to the water column.

The team consisted of Michael Mercado, Jennifer Feranda and Robert Morrell. We sampled monitoring wells # MW-6D and MW-6S using bailers to purge and sample the wells. Due to the depth of the wells and the amount of water inside the wells, the wells were not stabilized until 2:00 pm. We sampled MW-6S at 2:04 pm and sampled MW-6D at 2:14 pm. After preserving and securing the samples in the cooler, we left the site and arrived at Edison approximately at 4:00 pm. Upon returning to the Edison Facility the samples were logged-in and turned over to the ESD Laboratory at 4:10 pm.

On April 1st, Michael Mercado had a telephone conversation with Thomas Taccome

in which they addressed the problem that only two of the five wells were sampled. It was decided that the sampling team would go back and finish sampling the other three wells.

On April 2nd, the sampling team consisting of Michael Mercado, Jennifer Feranda and Robert Morrell departed the Edison Facility at 8:30 am. The team arrived at 10:00 am at the site. At 10:15 am Trip Blank #2 was prepared. We then started to purge wells number, MW-5S and MW-4. We stabilized both wells by 1:00 pm. MW-5S was the first well sampled and at 1:20 pm, we then sampled MW-4. The duplicate sample, MW-4D, was taken from MW-4 along with Matrix Spike (MS)/Matrix Duplicate Spike (MDS) and Matrix Spike (MS)/Matrix Duplicate (MD) volumes. After preserving and securing the samples in the cooler, we left the site and arrived at Edison approximately at 4:00 pm. Upon returning to the Edison Facility the samples were logged-in and turned over to the ESD Laboratory at 4:10 pm.

On April 3rd, the sampling team consisting of Michael Mercado, Jennifer Feranda and Jennifer Snow-Askbrook departed the Edison Facility at 8:30 am. The team arrived at 10:00 am at the site. At 10:15 am Trip Blank #3 was prepared. We then started to purge well number, MW-5D. At 1:30 pm MW-5D was stabilized and samples were pulled. Acids to preserve both VOAs and metals were not available. Samples were preserved in ice and secured in a cooler. After preserving and securing the samples, we left the site and arrived at Edison approximately at 5:00 pm. Upon returning to the Edison Facility the samples for metals were open and HNO<sub>3</sub> was added to preserve the samples. Since the log-in-station was closed, the samples could not be logged-in. More ice was added to the cooler in which the samples were in. Custody seals were place on the four corners of the cooler and the cooler was placed inside the van. The van was locked-up and the key to the van was maintained by Michael Mercado.

On April 4th, at 8:00 am the cooler with the samples was checked. The van was still locked and the custody seals were not broken. The samples were logged-in and turned over to the ESD Laboratory at 8:30 am. The fact that the VOA samples and blanks were not preserved with HCl was identified at that time.

The well was considered stabilized and ready for sample collection when: pH, specific conductance and temperature, and turbidity stabilized. The samples were pulled using the same teflon bailers used to purge each well. Each well was sampled using a different bailer for the following constituents:

Volatile organic analytes (VOAs)  
Semi-Volatile organic analytes (SVOAs)  
Total metals (unfiltered)

All the above constituents were analyzed at the EPA ESD Laboratory in Edison NJ.

#### Observations & Findings:

During the four days of sampling, the weather ranged between the low 20°s and the mid 50°s, with rain, snow, ice and strong winds.

The objective was to determine the effects, resulting from excavation on the site, to the underlying aquifer, by sampling the monitoring wells on the site

and having the samples screened against Federal MCL standards for VOCs, TCL, NVOAs and TAL metals. A summary of the results from the analyses is presented below. Analyzed sample data are from the ESD laboratory, located in Edison, New Jersey and data sheets are attached as Appendix B.

RESULTS:

TRIP BLANKS:

#1

CHLOROFORM	1.9 ug/l
2-BUTANONE	0.9 ug/l (Estimate QM)

#2

CHLOROFORM	1.2 ug/l
TOLUENE	0.7 ug/l (Estimate QM)
ACETONE	8.0 ug/l (Estimate QF)
2-BUTANONE	1.0 ug/l (Estimate QM)

#3

CHLOROFORM	1.7 ug/l
TOLUENE	0.3 ug/l (Estimate QM)
NAPHTHALENE	0.3 ug/l (Estimate QM)
2-BUTANONE	0.6 ug/l (Estimate QM)

EQUIPMENT BLANKS:

BAILER:

CHLOROFORM	1.9 ug/l
ACETONE	7.5 ug/l (Estimate QF)
2-BUTANONE	0.9 ug/l (Estimate QM)
OLEYL ALCOHOL	13. ug/l (Estimate QT)

MONITORING WELLS:

MW-4:

1,1,1-TRICHLOROETHANE	0.8 ug/l (Estimate QM)
2-BUTANONE	0.4 ug/l (Estimate QM)
OCTANOIC ACID	3.7 ug/l (Estimate QT)
DODECANOIC ACID	12. ug/l (Estimate QT)
DIACETATE-1,13-TRIDECANEDIOL	5.5 ug/l (Estimate QT)
OLEIC ACID	4.8 ug/l (Estimate QT)
UNKNOWN COMPOUND #1	3.0 ug/l (Estimate QT)
ALUMINUM	2900 ug/l
CALCIUM	8000 ug/l
CHROMIUM	109 ug/l
IRON	5560 ug/l (Estimate QP)
MANGANESE	119 ug/l
SODIUM	32000 ug/l
LEAD	4.3 ug/l

MW-4D:

1,1-DICHLOROETHANE	0.3 ug/l (Estimate QM)
1,1,1-TRICHLOROETHANE	1.5 ug/l
OCTANOIC ACID	4.8 ug/l (Estimate QT)
DODECANOIC ACID	17. ug/l (Estimate QT)
DECANOIC ACID	2.4 ug/l (Estimate QT)
UNKNOWN COMPOUND #1	2.5 ug/l (Estimate QT)
UNKNOWN COMPOUND #2	5.9 ug/l (Estimate QT)
UNKNOWN COMPOUND #3	34. ug/l (Estimate QT)
UNKNOWN COMPOUND #4	5.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #5	38. ug/l (Estimate QT)
ALUMINUM	3280 ug/l
CALCIUM	8000 ug/l
CHROMIUM	91. ug/l
IRON	5190 ug/l (Estimate QP)
MANGANESE	125 ug/l
SODIUM	33000 ug/l
LEAD	4.3 ug/l

MW-5D:

OCTANOIC ACID	4.9 ug/l (Estimate QT)
DODECANOIC ACID	20. ug/l (Estimate QT)
TETRADECANOIC ACID	5.4 ug/l (Estimate QT)
OLEYL ALCOHOL	12. ug/l (Estimate QT)
METHYLESTER 9-HEXADECENOIC A	20. ug/l (Estimate QT)
OCTADECANOIC ACID	5.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #1	4.8 ug/l (Estimate QT)
UNKNOWN COMPOUND #2	4.4 ug/l (Estimate QT)
UNKNOWN COMPOUND #3	39. ug/l (Estimate QT)
BIS(2-ETHYLHEXYL) HEXANEDIOIC	5.4 ug/l (Estimate QT)
ALUMINUM	688 ug/l
CALCIUM	15000 ug/l
CHROMIUM	80. ug/l
COPPER	52. ug/l
IRON	793 ug/l (Estimate QP)
MANGANESE	23. ug/l
SODIUM	52000 ug/l
NICKEL	41. ug/l
LEAD	6.5 ug/l
ZINC	97. ug/l

MW-5S:

2-BUTANONE	0.4 ug/l (Estimate QM)
BENZOIC ACID	0.8 ug/l (Estimate QM)
FLUORANTHENE	0.2 ug/l (Estimate QM)
OCTANOIC ACID	8.0 ug/l (Estimate QT)
DODECANOIC ACID	30. ug/l (Estimate QT)
OLEIC ACID	13. ug/l (Estimate QT)
DECANOIC ACID	4.9 ug/l (Estimate QT)
NONANOIC ACID	15. ug/l (Estimate QT)
TETRADECANOIC ACID	6.9 ug/l (Estimate QT)

UNKNOWN COMPOUND #1	5.6 ug/l (Estimate QT)
UNKNOWN COMPOUND #2	38. ug/l (Estimate QT)
UNKNOWN COMPOUND #3	36. ug/l (Estimate QT)
UNKNOWN COMPOUND #4	3.1 ug/l (Estimate QT)
ALUMINUM	2020 ug/l
CALCIUM	15000 ug/l
CHROMIUM	108 ug/l
IRON	3150 ug/l (Estimate QP)
MANGANESE	61. ug/l
SODIUM	35000 ug/l
NICKEL	55. ug/l
LEAD	5.6 ug/l
ZINC	95 ug/l

MW-6D:

OCTANOIC ACID	2.7 ug/l (Estimate QT)
N,N-BIS(2 HYDROXYETHYL) DODEC	7.9 ug/l (Estimate QT)
(Z) 11-HEXADECEN-1-OL	2.8 ug/l (Estimate QT)
9-HEXADECANOIC ACID	6.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #1	2.6 ug/l (Estimate QT)
UNKNOWN COMPOUND #2	5.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #3	3.8 ug/l (Estimate QT)
UNKNOWN COMPOUND #4	35. ug/l (Estimate QT)
UNKNOWN COMPOUND #5	68. ug/l (Estimate QT)
UNKNOWN COMPOUND #6	7.0 ug/l (Estimate QT)
CALCIUM	13000 ug/l
CHROMIUM	139 ug/l
IRON	640 ug/l (Estimate QP)
MANGANESE	27. ug/l
SODIUM	46000 ug/l
NICKEL	149 ug/l

MW-6S:

CARBON DISULFIDE	0.3 ug/l (Estimate QM)
BENZYL ALCOHOL	0.2 ug/l (Estimate QM)
OCTANOIC ACID	8.2 ug/l (Estimate QT)
DODECANOIC ACID	33. ug/l (Estimate QT)
OLEIC ACID	12. ug/l (Estimate QT)
TETRADECANOIC ACID	7.1 ug/l (Estimate QT)
(Z) 11-HEXADECEN-1-OL	9.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #1	11. ug/l (Estimate QT)
UNKNOWN COMPOUND #2	9.5 ug/l (Estimate QT)
UNKNOWN COMPOUND #3	64. ug/l (Estimate QT)
UNKNOWN COMPOUND #4	6.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #5	8.0 ug/l (Estimate QT)
ALUMINUM	1910 ug/l
CALCIUM	16000 ug/l
CHROMIUM	126 ug/l
IRON	2840 ug/l (Estimate QP)
MANGANESE	63. ug/l
SODIUM	42000 ug/l
NICKEL	64. ug/l
LEAD	8.0 ug/l

ZINC

56 ug/l

DIFFERENCE BETWEEN DUPLICATE SAMPLES:

	MW-4	MW-4D (DUPL)
1,1-DICHLOROETHANE	NON-DETECT	0.3 ug/l (Estimate QM)
1,1,1-TRICHLOROETHANE	0.8 ug/l (Estimate QM)	1.5 ug/l
2-BUTANONE	0.4 ug/l (Estimate QM)	NON-DETECT
OCTANOIC ACID	3.7 ug/l (Estimate QT)	4.8 ug/l (Estimate QT)
DODECANOIC ACID	12. ug/l (Estimate QT)	17. ug/l (Estimate QT)
DECANOIC ACID	NON-DETECT	2.4 ug/l (Estimate QT)
DIACETATE-1,13-TRIDECANEDIOL	5.5 ug/l (Estimate QT)	NON-DETECT
OLEIC ACID	4.8 ug/l (Estimate QT)	NON-DETECT
UNKNOWN COMPOUND #1	3.0 ug/l (Estimate QT)	2.5 ug/l (Estimate QT)
UNKNOWN COMPOUND #2	NON-DETECT	5.9 ug/l (Estimate QT)
UNKNOWN COMPOUND #3	NON-DETECT	34. ug/l (Estimate QT)
UNKNOWN COMPOUND #4	NON-DETECT	5.2 ug/l (Estimate QT)
UNKNOWN COMPOUND #5	NON-DETECT	38. ug/l (Estimate QT)
ALUMINUM	2900 ug/l	3280 ug/l
CALCIUM	8000 ug/l	8000 ug/l
CHROMIUM	109 ug/l	91. ug/l
IRON	5560 ug/l (Estimate QP)	5190 ug/l (Estimate QP)
MANGANESE	119 ug/l	125 ug/l
SODIUM	32000 ug/l	33000 ug/l
LEAD	4.3 ug/l	4.3 ug/l

Attachment:

- Appendix A, Field Sampling Plan
- Appendix B, Sample Data Sheets for ESD laboratory
- Appendix C, Well Data Sheets
- Appendix D, Field Data Sheet for ESD laboratory
- Appendix E, Analysis Request for ESD laboratory
- Appendix F, Chain of Custody Record for ESD laboratory

Appendix A, Field Sampling Plan

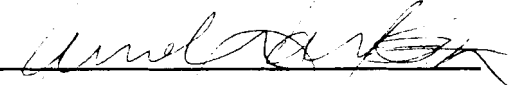


QA Plan Short Form  
Title Page

**WORK/QUALITY ASSURANCE PROJECT PLAN**  
**FOR GROUNDWATER SAMPLING**  
**AT ANCHOR CHEMICAL SUPERFUND SITE, HICKSVILLE, NEW YORK**

(Project Officer's Signature)  \_\_\_\_\_

(Project Officer's Name) Michael A. Mercado, Environmental Scientist  
Superfund Support Section

(Project Quality Assurance Officer's Signature)  \_\_\_\_\_

(Project Quality Assurance Officer's Name) Amelia Jackson, Chemist  
Toxic & Hazardous Waste Section

1. **Project Name:** Anchor Chemical, Hicksville, Long Island, NY
2. **Project Requested By:** US EPA - Region 2, ERRD, New York/Caribbean Superfund Branch II
3. **Date of Request received from RPM:** March 13, 1996
4. **Date of Project Initiation:** Same as date of request.
5. **EPA Project Officer:** Michael A. Mercado
6. **EPA Quality Assurance Officer:**
7. **Project Description:** This groundwater sampling event involves the collection of water samples from five (5) monitoring wells at the Anchor Chemical Superfund site in Hicksville, Long Island. The five (5) monitoring wells to be sampled are identified on the enclosure site map as in enclosure # 1. The five (5) monitoring wells are: MW-4, MW-5S, MW-5D, MW-6S and MW-6D. On 29 and 30 September 1995 the site's PRP removed approximately 24 cubic yards of contaminated sediments from four (4) on-site drywells. The drywells were designated DW2, DW3, DW6 and DW8.

**A. Objective and Scope Statement:**

The objective of this sampling event is to determine the effects, resulting from the excavation, to the underlying aquifer. The removal of subsurface soils may have had a positive impact by removing the contaminated soils but negative impacts (pollutant transport) may have resulted from subsurface soil disturbance. In order to determine the effects five (5) monitoring wells will be sampled. These samples will be screened against Federal MCL standards, for VOCs, TCL NVOAs and TAL metals. Since the groundwater in the area is a designated sole source aquifer, the purpose of the remedial activities is to ensure that the site does not contaminate the drinking water supply. Enclosure 3 is the Target Compound List (TCL) and the Target Analyte List (TAL).

**B. Data Usage:** Data and the interpretation of the against the MCL will be sent to ERRD for incorporation into the project file. ERRD will be responsible for informing affected residents and the local health department of pertinent results.

**C. Monitoring Network Design and Rationale:** Sample will be collected from each of the five monitoring wells as specified by ERRD (see enclosure 2). The only acceptable equipment available to sample for all requested parameter is the bailers, so the bailer will be used to do the sampling. All sampling tasks will conform with the Quality Assurance set forth in the current Region 2 CERCLA Quality Assurance Manual and the sampling procedures in EPA/540/P-91/007 dated Jan 91 for decontamination and sampling using a bailer. A copy of this Work/QA Short Form will be on site and available for reference during all sampling events.

The five monitoring wells to be sampled are MW-6D and MW-6S, both up gradient of the drywells and MW-4, MW-5D and MW-5S which are down gradient. All wells are located on the site (see enclosure 1, site map).

One VOC trip blank is required for each cooler containing VOC samples. Environmental duplicate, MS/MSD and MS/MD volume sample will be collected at a minimum of 1 in every 20 samples in each medium sampled. Field blanks are required, one per decontamination event, not to exceed one per day. The water used will be analyte-free from the ESD lab. This water is tested daily and any appearance of trace contaminants are noted and qualified.

Decontamination will conform to the following procedure used: equipment is scrubbed with Alconox and rinsed with water, then rinsed with nitric acid followed by another water rinse and then rinsed with methanol followed by another water rinse. As a minimum the final rinse will be deionized water.

We plan to have one trip blank, one environmental duplicate, a MS/MSD, a MS/MD and one field blank. The water used for the blanks will come from the ESD lab which is analyte-free and tested daily as discussed above.

**D. Monitoring Parameters and their Frequency of Collection:**

Monitoring well samples and blanks will be analyzed for drinking water levels of VOC's, TCL semivolatiles and TAL metals, except for trip blanks. Trip blanks will be analyzed for drinking water levels VOC's only. There will be six (6) samples including an environmental duplicate. The above parameters were chosen by ERRD. Monitoring well MW-4 has been chosen to supply the environmental duplicate. Each well will be monitored once during this sampling event. A follow-up survey will be recommended one year from this event as an additional check that pollutants are not migrating off-site through the groundwater.

**E. Parameter Table:** All analytical and quality assurance requirements of the Technical Support Branch will be followed. All samples parameter data will be incorporated into the table below:

PARAMETER TABLE

Parameter*1	Container Types	Analytical Method	Sample Preservation	Holding Times
MCL Organics VOC's	40ml VOC's Vial*2	TSB SOP C-49 (m.524.2)	(HCL to pH<2.0) Cool to 4 C	14 days
TCL Semivolatiles	1l amber glass bottle*3	TSB SOP C-3	Cool to 4 C	7 days
TAL Inorganic Metals	1l glass bottle*3	TSB SOP C-70	HNO3 to pH<2.0 Cool to 4 C	6 months (Hg 28 days)

\*1 For each parameter there will be five well samples and one duplicate, for a total of 6 samples.

\*2 Six, 40 ml vials for the first sample (no head space); three 40 ml vials for each additional sample.

\*3 Three, 1 liters bottles for the first sample, one 1 liter bottle for each additional sample.

8. Project Fiscal Information (Optional): Not included.

9. Schedule of Tasks and Products:

<u>Activity</u>	<u>Date</u>
Review and Background Information	March 19, 1996
Submit a QA plan	March 21, 1996
Book samples anticipated to be collected	March 20, 1996
Obtain Site Access	Prearranged by ERRD
Mobilize to Site	March 28, 1996
Complete Field Work	March 29, 1996
Package and ship samples to laboratory	Package at the time of sampling and will be delivered by samplers on March 29, 1996
Prepare Sampling Trip Report	Within one week of the sampling event
Prepare and submit data presentation to ERRD	Within two weeks of receipt of validated analytical data EPA Laboratory/TSB

**10. Project Organization and Responsibility:** The following is a list of key project personnel and their corresponding responsibilities:

Michael A Mercado, Superfund Support Section Project Officer	-sampling operations
Michael A Mercado, Superfund Support Section Project Officer	-sampling QC
TSB	-laboratory analysis
TSB	-laboratory QC
TSB	-data processing activities
TSB	-data processing QC
MMB or TSB QA/QC	-data quality review
N/A	-performance auditing
N/A	-systems auditing
MMB	-overall QA
Michael A Mercado, Superfund Support Section	-overall project
Jennifer Feranda, Superfund Support Section	-health and safety officer

**11. Data Quality Requirements and Assessments:** The data quality requirements for TSB are listed in the EPA-ESD laboratory for QA/QC Plan for GC/MS May 94 for organic analysis and Inorganic QA/QC Plan Nov 93. The method to be used, (524.2) -SOPs C-49, C-3, and C-70, will produce low level CRQLs within the ranges expected and needed for comparison to the drinking water MCLs.

**12. Sampling Procedures:** All monitoring well sampling will be in accordance with EPA/540/P-91/007 dated: Jan 91, Compendium of ERT Groundwater Sampling Procedures, for the sampling of groundwater and decontamination of equipment. Samples will be maintained in sealed cooler w/ice at 4 degrees C. All samples will be taken and delivered on the 29th of March.

**13. Sample Custody Procedures:** Sample custody seals will be placed on each cooler in which samples are contained. Chain of custody forms will accompany each cooler. Each time the seal is broken on the sample coolers, a new seal will be placed on. The custody seals will record the date and time of placement as well as the originator. It will also contain the date and time the seal was broken and the person responsible for this action. At the conclusion of the sampling event. all samples will be delivered personally to the EPA TSB for log-in.

**14. Calibration Procedures and Preventative Maintenance:** Laboratory will followed as specified under the EPA-ESD

Laboratory SOP's. The following field equipment will be used to check for stablization of the aquifer during purging: LaMotte Model 2008 Turbidity Meter, Orion Research Portable Meter 200 Series, and Cole-Parmer Conductively Meter, Model 1500. All of these will be calibrated and maintained IAW equipment operator's manual.

**15. Documentation, Data Reduction, and Reporting:**

**A. Documentation:** Data sheets, field logs, traffic reports, photographs and chain of custody forms will be kept by the project manager of each individual site.

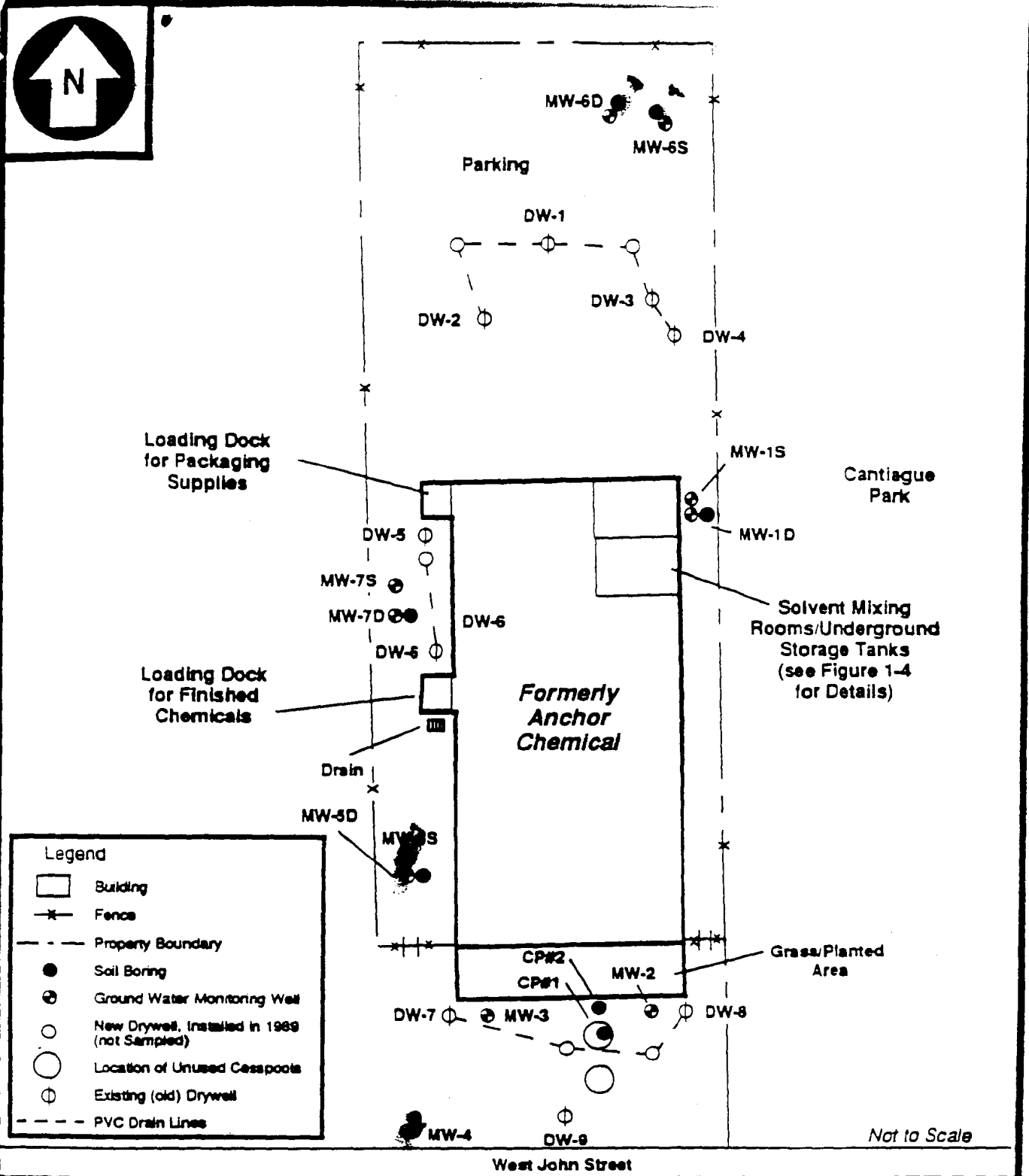
**B. Data Reduction and Reporting:** The laboratory performing the analysis will calculate and transfer data to ERRD-RPM for the site per ESD protocol.

**16. Data Validation:** The US EPA TSB will perform all data validation in house for all samples which it analyzes.

**17. Performance and Systems Audits:** As according to ESD-SMB and ESD-TSB SOP's.

**18. Corrective Action:** Corrective Action will be performed as required by the project manager in the field and by the audit report.

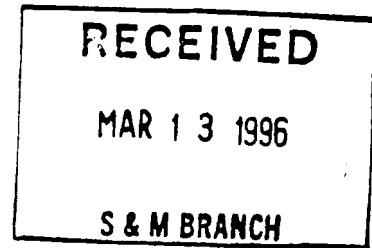
**19. Reports:** A data presentation will be prepared by the project manager and submitted to ERRD in New York. The report will include the data and will discuss whether or not the samples had exceeded the MCLs at any upgradient or downgradient locations. If any MCLs are exceeded, it will be up to the RPM to take any aditional action.




Encl 1

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY  
REGION II

DATE: MAR 1 1 1996



SUBJECT: Groundwater Samples at the Anchor Chemical Superfund Site

FROM: Carole Petersen, Chief   
NY/Caribbean Superfund Branch II

TO: Randy Braun, Acting Chief  
Surveillance and Analysis Branch

The purpose of this memo is to request the assistance of the Surveillance and Analysis Branch to collect and analyze five groundwater water samples from five groundwater monitoring wells, located at the Anchor Chemical Superfund Site. The Site is located at 500 West John Street in Hicksville, Long Island.

The sample data is needed to determine the effectiveness of a removal action, which resulted in the excavation of approximately 24 cubic yards of contaminated sediments from four on-Site drywells, designated DW2, 3, 6 and 8. The removal was performed on September 29 and September 30, 1995.

The wells, which need to be sampled, are designated MW-4, MW-5S, MW-5D, MW-6S and MW-6D. MW-6S and MW-6D are up gradient wells; MW-4, MW-5S and MW-5D are down gradient wells. The attached figure shows the drywell and well locations. Well construction logs for the five wells are also attached. All of the samples should be analyzed for TCL volatile and semi-volatile organic compounds and inorganic compounds. The groundwater sample data are needed by the middle of April 1996.

Please have your staff contact Tom Taccone, of my staff at (212) 637-4281, to arrange for the well sampling. If you have any questions, you may contact me at (212)637-4285.

We appreciate your assistance in this matter.

Attachments

cc: J. Greco, NYSDEC

Encl 2



Appendix B, Well Data Sheets

WELL DATA SHEET

SITE: ANCHOR CHEMICAL WELL#: MW-4

TYPE OF SAMPLE: GROUND WATER AIR MONITORING: (HNU) -0-

SAMPLING PERSONNEL: MERCADO/FERNADA/MORRELL

EVACUATION INFORMATION

DATE/TIME: 4/2/96 1320 HRS METHOD: BAILER

TOTAL DEPTH (FT): 80.61 WELL CSG. TYPE/DIAM.: 4" SS

TOP OF CSG. TO H2O LEVEL (FT): 66.57 WATER COLUMN HEIGHT (FT): 14.04

TOTAL VOLUME EVACUATED (GAL): 27 TURBIDITY TRUE READING: 4.98 NTU

pH METER CALIBRATION: 4(4.0) 7(7.0) 10(10.04) TIME: 1032HRS

SAMPLING INFORMATION

DATE/TIME OF SAMPLING: 4/2 - 1320 METHOD: BAILER

SAMPLE #: 200128

FIELD MEASUREMENT DATA

TIME	VOLUME EVACUATED (GAL)	TEMP (C)	*SPECIFIC CONDUCTANCE (U MHOS/CM)	pH (SU)	TURBIDITY (NTU)
1148HRS	8.0	14.3	153	6.19	173.7
1225	18.0	14.8	159	5.90	133.4
1248	28.0	14.6	160	5.89	107.6

\*W/O TEMPERATURE COMPENSATION - CONVERSIONS: 1 m=3.28'/1'=.305m

GENERAL INFORMATION

WEATHER CONDITIONS: STRONG WINDS TEMP IN THE 50s

SAMPLE CHARACTERISTICS: CLEAR WATER

COMMENTS AND OBSERVATIONS: \_\_\_\_\_

ANALYSIS/PRESERVATION: (SEE FIELD DATA SHEET)

CERTIFICATION: GLASSWARE EAGLE PICHER

WELL DATA SHEET

SITE: ANCHOR CHEMICAL WELL#: MW-4D  
TYPE OF SAMPLE: GROUND WATER AIR MONITORING: (HNU) -0-  
SAMPLING PERSONNEL: MERCADO/FERNADA/MORRELL

EVACUATION INFORMATION

DATE/TIME: 4/2/96 1320 HRS METHOD: BAILER  
TOTAL DEPTH (FT): 80.61 WELL CSG. TYPE/DIAM.: 4" SS  
TOP OF CSG. TO H2O LEVEL (FT): 66.57 WATER COLUMN HEIGHT (FT): 14.04  
TOTAL VOLUME EVACUATED (GAL): 27 TURBIDITY TRUE READING: 4.98 NTU  
pH METER CALIBRATION: 4(4.0) 7(7.0) 10(10.04) TIME: 1032HRS

SAMPLING INFORMATION

DATE/TIME OF SAMPLING: 4/2 - 1320 METHOD: BAILER  
SAMPLE #: 200129

FIELD MEASUREMENT DATA

TIME	VOLUME EVACUATED (GAL)	TEMP (C)	*SPECIFIC CONDUCTANCE (U MHOS/CM)	pH (SU)	TURBIDITY (NTU)
1148HRS	8.0	14.3	153	6.19	173.7
1225	18.0	14.8	159	5.90	133.4
1248	28.0	14.6	160	5.89	107.6

\*W/O TEMPERATURE COMPENSATION - CONVERSIONS: 1 m=3.28'/1'=.305m

GENERAL INFORMATION

WEATHER CONDITIONS: STRONG WINDS TEMP IN THE 50s  
SAMPLE CHARACTERISTICS: CLEAR WATER  
COMMENTS AND OBSERVATIONS: MW-4D IS THE DUPL OF MW-4  
ANALYSIS/PRESERVATION: (SEE FIELD DATA SHEET)  
CERTIFICATION: GLASSWARE EAGLE PICHER

WELL DATA SHEET

SITE: ANCHOR CHEMICAL WELL#: MW-5D  
TYPE OF SAMPLE: GROUND WATER AIR MONITORING: (HNU) -0-  
SAMPLING PERSONNEL: MERCADO/FERNADA/SNOW

EVACUATION INFORMATION

DATE/TIME: 4/3/96 1031 HRS METHOD: BAILER  
TOTAL DEPTH (FT): 122.6 WELL CSG. TYPE/DIAM.: 4" SS  
TOP OF CSG. TO H2O LEVEL (FT): 63.35 WATER COLUMN HEIGHT (FT): 59.25  
TOTAL VOLUME EVACUATED (GAL): 114 TURBIDITY TRUE READING: 4.97 NTU  
pH METER CALIBRATION: 4(4.0) 7(7.0) 10(10.05) TIME: 1020HRS

SAMPLING INFORMATION

DATE/TIME OF SAMPLING: 4/3 - 1408 METHOD: BAILER  
SAMPLE #: 200130

FIELD MEASUREMENT DATA

TIME	VOLUME EVACUATED (GAL)	TEMP (C)	*SPECIFIC CONDUCTANCE (U MHOS/CM)	pH (SU)	TURBIDITY (NTU)
1110HRS	19.0	14.0	N/A	6.60	N/A
1149HRS	38.0	14.2	N/A	6.55	N/A
1225HRS	55.0	14.4	N/A	6.64	N/A
1310HRS	74.0	14.2	N/A	6.46	N/A
1408HRS	94.0	14.4	N/A	6.40	N/A

\*W/O TEMPERATURE COMPENSATION - CONVERSIONS: 1 m=3.28' /1'=.305m

GENERAL INFORMATION

WEATHER CONDITIONS: IN THE HIGH 50s AND WINDY  
SAMPLE CHARACTERISTICS: CLEAR WATER  
COMMENTS AND OBSERVATIONS: \_\_\_\_\_  
ANALYSIS/PRESERVATION: (SEE FIELD DATA SHEET)  
CERTIFICATION: GLASSWARE EAGLE PICHER

WELL DATA SHEET

SITE: ANCHOR CHEMICAL WELL#: MW-5S

TYPE OF SAMPLE: GROUND WATER AIR MONITORING: (HNU) -0-

SAMPLING PERSONNEL: MERCADO/FERNADA/MORRELL

EVACUATION INFORMATION

DATE/TIME: 4/2/96 1100 HRS METHOD: BAILER

TOTAL DEPTH (FT): 78.35 WELL CSG. TYPE/DIAM.: 4" SS

TOP OF CSG. TO H2O LEVEL (FT): 63.16 WATER COLUMN HEIGHT (FT): 15.19

TOTAL VOLUME EVACUATED (GAL): 29 TURBIDITY TRUE READING: 4.98 NTU

pH METER CALIBRATION: 4(4.0) 7(7.0) 10(10.04) TIME: 1032HRS

SAMPLING INFORMATION

DATE/TIME OF SAMPLING: 4/2 - 1300 METHOD: BAILER

SAMPLE #: 200131

FIELD MEASUREMENT DATA

TIME	VOLUME EVACUATED (GAL)	TEMP (C)	*SPECIFIC CONDUCTANCE (U MHOS/CM)	pH (SU)	TURBIDITY (NTU)
1135HRS	10.0	13.8	296	6.26	108.5
1215HRS	20.0	14.1	316	5.96	129.6
1252HRS	30.0	14.1	316	6.06	89.0
1258HRS	32.0	14.1	318	6.06	94.3

\*W/O TEMPERATURE COMPENSATION - CONVERSIONS: 1 m=3.28'/1'=.305m

GENERAL INFORMATION

WEATHER CONDITIONS: STRONG WINDS TEMP IN THE 50s

SAMPLE CHARACTERISTICS: CLEAR WATER

COMMENTS AND OBSERVATIONS: \_\_\_\_\_

ANALYSIS/PRESERVATION: (SEE FIELD DATA SHEET)

CERTIFICATION: GLASSWARE EAGLE PICHER

WELL DATA SHEET

SITE: ANCHOR CHEMICAL WELL#: MW-6D  
TYPE OF SAMPLE: GROUND WATER AIR MONITORING: (HNU) -0-  
SAMPLING PERSONNEL: MERCADO/FERNADA/MORRELL

EVACUATION INFORMATION

DATE/TIME: 3/29/96 1320 HRS METHOD: BAILER  
TOTAL DEPTH (FT): 121 WELL CSG. TYPE/DIAM.: 4" SS  
TOP OF CSG. TO H2O LEVEL (FT): 70.0 WATER COLUMN HEIGHT (FT): 51.00  
TOTAL VOLUME EVACUATED (GAL): 105 TURBIDITY TRUE READING: 4.96 NTU  
pH METER CALIBRATION: 4(4.0) 7(7.0) 10(10.12) TIME: 0810HRS

SAMPLING INFORMATION

DATE/TIME OF SAMPLING: 3/29 - 1413 METHOD: BAILER  
SAMPLE #: 200132

FIELD MEASUREMENT DATA

TIME	VOLUME EVACUATED (GAL)	TEMP (C)	*SPECIFIC CONDUCTANCE (U MHOS/CM)	pH (SU)	TURBIDITY (NTU)
1013HRS	20.0	12.0	289	6.59	ICILY
1124HRS	40.0	12.3	280	6.56	ICILY
1333HRS	60.0	12.4	353	6.80	ICILY
1413HRS	80.0	12.5	360	6.80	ICILY

\*W/O TEMPERATURE COMPENSATION - CONVERSIONS: 1 m=3.28'/1'=.305m

GENERAL INFORMATION

WEATHER CONDITIONS: WET, SNOWY, AND ICILY. TEMP IN THE 20s  
SAMPLE CHARACTERISTICS: CLEAR WATER  
COMMENTS AND OBSERVATIONS: TURBIDITY METER NOT OPERABLE TO COLD  
ANALYSIS/PRESERVATION: (SEE FIELD DATA SHEET)  
CERTIFICATION: GLASSWARE EAGLE PICHER

WELL DATA SHEET

SITE: ANCHOR CHEMICAL WELL#: MW-6S

TYPE OF SAMPLE: GROUND WATER AIR MONITORING: (HNU) -0-

SAMPLING PERSONNEL: MERCADO/FERNADA/MORRELL

EVACUATION INFORMATION

DATE/TIME: 3/29/96 1045 HRS METHOD: BAILER

TOTAL DEPTH (FT): 81.5 WELL CSG. TYPE/DIAM.: 4" SS

TOP OF CSG. TO H2O LEVEL (FT): 63.00 WATER COLUMN HEIGHT (FT): 18.50

TOTAL VOLUME EVACUATED (GAL): 36 TURBIDITY TRUE READING: 4.96 NTU  
pH METER CALIBRATION: 4(4.0) 7(7.0) 10(10.12) TIME: 0810HRS

SAMPLING INFORMATION

DATE/TIME OF SAMPLING: 2/29 - 1404 METHOD: BAILER

SAMPLE #: 200133

FIELD MEASUREMENT DATA

TIME	VOLUME EVACUATED (GAL)	TEMP (C)	*SPECIFIC CONDUCTANCE (U MHOS/CM)	pH (SU)	TURBIDITY (NTU)
1118HRS	10.0	12.7	387	6.78	N/A
1303HRS	20.0	12.9	374	6.80	N/A
1349HRS	30.0	12.8	367	6.80	N/A
1404HRS	36.0	12.8	367	6.80	N/A

\*W/O TEMPERATURE COMPENSATION - CONVERSIONS: 1 m=3.28'/1'=.305m

GENERAL INFORMATION

WEATHER CONDITIONS: WET, SNOWY, AND ICILY. TEMP IN THE 20s

SAMPLE CHARACTERISTICS: CLEAR WATER

COMMENTS AND OBSERVATIONS: \_\_\_\_\_

ANALYSIS/PRESERVATION: (SEE FIELD DATA SHEET)

CERTIFICATION: GLASSWARE EAGLE PICHER

Appendix C, Sample Data Sheets for ESD laboratory



LAB DATA MANAGEMENT SYSTEM - REGION 11  
COMPLETED PROJECT APPROVAL

REPORT DATE 96/05/01

PROJECT NUMBER

235

PROJECT DATE

96/03/29

PROJECT NAME

ANCHOR CHEMICAL

APPROVED

*Barbara A. Piggott*  
5/2/96

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

## EXPLANATIONS OF REMARK CODES

REMARK CODE	EXPLANATION
B	RESULTS BASED UPON COLONY COUNTS OUTSIDE ACCEPTABLE RANGE
J	ESTIMATED VALUE
K	ACTUAL VALUE KNOWN TO BE LESS THAN VALUE GIVEN
L	ACTUAL VALUE KNOWN TO BE GREATER THAN VALUE GIVEN
N	NO OBSERVABLE EFFECT CONCENTRATION < 0.3%
O	SAMPLED BUT NOT ANALYZED DUE TO LAB ACCIDENT
T	REPORTED VALUE LESS THAN CRITERIA OF DETECTION
U	REPORTING LIMIT

## QA/QC REMARK CODES

CODE	EXPLANATION
QD	ACCURACY CHECK SAMPLE ABOVE UPPER ACCEPTANCE LIMIT
QE	ACCURACY CHECK SAMPLE BELOW LOWER ACCEPTANCE LIMIT
QF	PRECISION OF CALIBRATION CURVE LESS THAN ACCEPTANCE CRITERIA
QJ	ESTIMATED DETECTION LIMIT DUE TO INTERFERENCE
QG	CONTINUING CALIBRATION CHECK DOES NOT MEET ACCEPTANCE CRITERIA
QS	SPIKE RECOVERIES ABOVE UPPER ACCEPTANCE LIMIT
QR	SPIKE RECOVERIES BELOW LOWER ACCEPTANCE LIMIT
QP	SAMPLE REPLICATE PRECISION DOES NOT MEET ACCEPTANCE CRITERIA
QH	RECOMMENDED HOLDING TIMES EXCEEDED
QT	TENTATIVELY IDENTIFIED COMPOUND
QM	PRESENCE OF MATERIAL VERIFIED BUT NOT QUANTIFIED
QB	BLANK CONTAMINATED BY ANALYTE IN EXCESS OF ACCEPTANCE CRITERIA
QQ	SAMPLE IMPROPERLY PRESERVED

LOCATION CODES FOR IDENTIFICATION OF SAMPLING POINTS AT INDUSTRIAL /  
SANITARY FACILITIES, LANDFILLS, HAZARDOUS WASTE SITES.

CODE NUMBERS	SAMPLING POINTS
1001 - 1050	EFFLUENT PIPE NUMBER 001 TO 050
1051 - 1099	OTHER EFFLUENTS SUCH AS COOLING TOWER DISCHARGE, DISCHARGE FROM HOLDING PONDS, ETC...
1100 - 1249	IN PLANT SAMPLES
1435 - 1454	SEPARATE INFLUENT POINTS/WATER SOURCES
15XX	INFLUENT ASSOCIATED WITH EFFLUENT 10XX
2000	BLANK FOR VOLATILE ORGANICS
3000 - 3099	GROUND WATER FROM WELL 01 TO 99
3100 - 3199	SEDIMENT SAMPLE (WATER BOTTOM)
3200 - 3299	SOIL SAMPLE
3300 - 3399	STREAM WATER SAMPLE
3400 - 3499	LAGOON SAMPLE
3500 - 3599	STORAGE TANK SAMPLE
3600 - 3699	LEACHATE SAMPLE
3700 - 3799	OTHER TYPE SAMPLE

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
NONE	96/03/29	0803							
DEPTH: 0000 SUBSTRATE: AQUEOUS									
DESCRIPTION: TRIP BLANK									
			200126	99999	CHLOROMETHANE	UG/L		1 U	
				99999	BROMOMETHANE	UG/L		1 U	
				39175	VINYL CHLORIDE	UG/L		1 U	
				34311	CHLOROETHANE	UG/L		1 U	
				34423	METHYLENE CHLORIDE	UG/L		1 U	
				34488	TRICHLOROFLUOROMETHANE	UG/L		1 U	
				34501	1,1-DICHLOROETHYLENE	UG/L		1 U	
				34496	1,1-DICHLOROETHANE	UG/L		1 U	
				99964	CARBON DISULFIDE	UG/L		1 U	
				34546	TRANS 1,2 DICHLOROETHYLENE	UG/L		1 U	
				99999	CIS 1,2- DICHLOROETHYLENE	UG/L		1 U	
				99999	2,2 DICHLOROPROPANE	UG/L		1 U	
				32106	CHLOROFORM	UG/L		1.9	
				99999	DIBROMOMETHANE	UG/L		1 U	
				34506	1,1,1-TRICHLOROETHANE	UG/L		1 U	
				32102	CARBON TETRACHLORIDE	UG/L		1 U	
				32101	DICHLOROBROMOMETHANE	UG/L		1 U	
				99999	1,1-DICHLOROPROPENE	UG/L		1 U	
				34541	1,2-DICHLOROPROPANE	UG/L		1 U	
				99999	CIS-1,3-DICHLOROPROPENE	UG/L		1 U	
				39180	TRICHLOROETHYLENE	UG/L		1 U	
				34030	BENZENE	UG/L		1 U	
				32103	1,2-DICHLOROETHANE	UG/L		1 U	
				99999	1,3 DICHLOROPROPANE	UG/L		1 U	
				99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		1 U	
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200126	99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SECUBYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		7 U	
				99999	2-BUTANONE	UG/L		0.9 J	QM
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
			200127	99999	CHLOROMETHANE	UG/L		1 U	
				99999	BROMOMETHANE	UG/L		1 U	
				39175	VINYL CHLORIDE	UG/L		1 U	
				34311	CHLOROETHANE	UG/L		1 U	
				34423	METHYLENE CHLORIDE	UG/L		1 U	
				34488	TRICHLOROFLUOROMETHANE	UG/L		1 U	
				34501	1,1-DICHLOROETHYLENE	UG/L		1 U	
				34496	1,1-DICHLOROETHANE	UG/L		1 U	
				99964	CARBON DISULFIDE	UG/L		1 U	
				34546	TRANS 1,2 DICHLOROETHYLENE	UG/L		1 U	
				99999	CIS 1,2- DICHLOROETHYLENE	UG/L		1 U	
				99999	2,2 DICHLOROPROPANE	UG/L		1 U	
				32106	CHLOROFORM	UG/L		1.9	
				99999	DIBROMOMETHANE	UG/L		1 U	

NONE 96/03/29 0805  
 DEPTH: 0000 SUBSTRATE: AQUEOUS  
 DESCRIPTION: EQUIPMENT BLANKS

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200127	34506	1,1,1-TRICHLOROETHANE	UG/L		1 U	
				32102	CARBON TETRACHLORIDE	UG/L		1 U	
				32101	DICHLOROBROMOMETHANE	UG/L		1 U	
				99999	1,1-DICHLOROPROPENE	UG/L		1 U	
				34541	1,2-DICHLOROPROPANE	UG/L		1 U	
				99999	CIS-1,3-DICHLOROPROPENE	UG/L		1 U	
				39180	TRICHLOROETHYLENE	UG/L		1 U	
				34030	BENZENE	UG/L		1 U	
				32103	1,2-DICHLOROETHANE	UG/L		1 U	
				99999	1,3 DICHLOROPROPANE	UG/L		1 U	
				99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		1 U	
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SEC BUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200127	34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		7.5 J	QF
				99999	2-BUTANONE	UG/L		0.9 J	QM
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
				34273	BIS(2-CHLOROETHYL) ET.	UG/L		4.5 U	
				34694	PHENOL	UG/L		4.5 U	
				34586	2-CHLOROPHENOL	UG/L		4.5 U	
				34566	1,3-DICHLOROBENZENE	UG/L		4.5 U	
				34571	1,4-DICHLOROBENZENE	UG/L		4.5 U	
				34536	1,2-DICHLOROBENZENE	UG/L		4.5 U	
				99999	BENZYL ALCOHOL	UG/L		4.5 U	
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		4.5 U	
				99999	2-METHYL PHENOL	UG/L		4.5 U	
				99999	4-METHYL PHENOL	UG/L		4.5 U	
				34396	HEXACHLOROETHANE	UG/L		4.5 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		4.5 U	
				34447	NITROBENZENE	UG/L		4.5 U	
				34408	ISOPHORONE	UG/L		4.5 U	
				34591	2-NITROPHENOL	UG/L		4.5 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		4.5 U	
				99999	BENZOIC ACID	UG/L		4.5 U	
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		4.5 U	
				34601	2,4-DICHLOROPHENOL	UG/L		4.5 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		4.5 U	
				34696	NAPHTHALENE	UG/L		4.5 U	
				99999	4-CHLOROANILINE	UG/L		4.5 U	
				39702	HEXACHLOROBUTADIENE	UG/L		4.5 U	
				34452	P-CHLORO-M-CRESOL	UG/L		4.5 U	
				99999	2-METHYL NAPHTHALENE	UG/L		4.5 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		36 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		4.5 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		4.5 U	
				34581	2-CHLORONAPHTHALENE	UG/L		4.5 U	
				99999	2-NITROANILINE	UG/L		4.5 U	
				34200	ACENAPHTHYLENE	UG/L		4.5 U	
				34341	DIMETHYL PHTHALATE	UG/L		4.5 U	
				34626	2,6-DINITROTOLUENE	UG/L		4.5 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200127	99999	3-NITROANILINE	UG/L		4.5 U	
				34205	ACENAPHTHENE	UG/L		4.5 U	
				34616	2,4-DINITROPHENOL	UG/L		36 U	
				99999	DIBENZOFURAN	UG/L		4.5 U	
				34646	4-NITROPHENOL	UG/L		4.5 U	
				34611	2,4-DINITROTOLUENE	UG/L		4.5 U	
				34381	FLUORENE	UG/L		4.5 U	
				34641	4-CHLOROPHENYL PHENYL ET.	UG/L		4.5 U	
				99999	4-NITROANILINE	UG/L		4.5 U	
				34336	DIETHYL PHTHALATE	UG/L		4.5 U	
				34657	4,6-DINITRO-O-CRESOL	UG/L		36 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		4.5 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		4.5 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		4.5 U	
				39700	HEXACHLOROBENZENE	UG/L		4.5 U	
				39032	PENTACHLOROPHENOL	UG/L		36 U	
				34461	PHENANTHRENE	UG/L		4.5 U	
				34220	ANTHRACENE	UG/L		4.5 U	
				34376	FLUORANTHENE	UG/L		4.5 U	
				39110	DI-N-BUTYLPHTHALATE	UG/L		4.5 U	
				34469	PYRENE	UG/L		4.5 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		4.5 U	
				34526	1,2-BENZANTHRACENE	UG/L		4.5 U	
				34320	CHRYSENE	UG/L		4.5 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		4.5 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		4.5 U	
				34230	3,4-BENZOFUORANTHENE	UG/L		4.5 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		4.5 U	
				34247	BENZO(A)PYRENE	UG/L		4.5 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		4.5 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		4.5 U	
				34521	1,12-BENZOPERYLENE	UG/L		4.5 U	
				99999	OLEYL ALCOHOL	UG/L		13 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINUM	UG/L		200 U	
				01002	ARSENIC	UG/L		10 U	
				01007	BARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	
				00916	CALCIUM	MG/L		5 U	
				01027	CADMIUM	UG/L		5 U	
				01037	COBALT	UG/L		50 U	

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PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE		TIME	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE &		
	FROM	TO							REMARK	QA/QC	
			OF							REMARK	
			DAY								
				200127	01034	CHROMIUM	UG/L		10	U	
					01042	COPPER	UG/L		25	U	
					01045	IRON	UG/L		100	U	QP
					71900	MERCURY	UG/L		0.2	U	
					00937	POTASSIUM	MG/L		5	U	
					00927	MAGNESIUM	MG/L		5	U	
					01055	MANGANESE	UG/L		15	U	
					00929	SODIUM	MG/L		5	U	
					01067	NICKEL	UG/L		40	U	
					01051	LEAD	UG/L		3	U	
					01097	ANTIMONY	UG/L		60	U	
					01147	SELENIUM	UG/L		5	U	
					01059	THALLIUM	UG/L		10	U	
					01087	VANADIUM	UG/L		50	U	
					01092	ZINC	UG/L		20	U	

NONE 96/04/02 1320  
 DEPTH: 67.0 SUBSTRATE: AQUEOUS  
 DESCRIPTION: MW-4

				200128	99999	CHLOROMETHANE	UG/L		1	U	
					99999	BROMOMETHANE	UG/L		1	U	
					39175	VINYL CHLORIDE	UG/L		1	U	
					34311	CHLOROETHANE	UG/L		1	U	
					34423	METHYLENE CHLORIDE	UG/L		1	U	
					34488	TRICHLOROFLUOROMETHANE	UG/L		1	U	
					34501	1,1-DICHLOROETHYLENE	UG/L		1	U	
					34496	1,1-DICHLOROETHANE	UG/L		1	U	
					99964	CARBON DISULFIDE	UG/L		1	U	
					34546	TRANS 1,2 DICHLOROETHYLENE	UG/L		1	U	
					99999	CIS 1,2- DICHLOROETHYLENE	UG/L		1	U	
					99999	2,2 DICHLOROPROPANE	UG/L		1	U	
					32106	CHLOROFORM	UG/L		1	U	
					99999	DIBROMOMETHANE	UG/L		1	U	
					34506	1,1,1-TRICHLOROETHANE	UG/L		0.8	J	QM
					32102	CARBON TETRACHLORIDE	UG/L		1	U	
					32101	DICHLOROBROMOMETHANE	UG/L		1	U	
					99999	1,1-DICHLOROPROPENE	UG/L		1	U	
					34541	1,2-DICHLOROPROPANE	UG/L		1	U	
					99999	CIS-1,3-DICHLOROPROPENE	UG/L		1	U	
					39180	TRICHLOROETHYLENE	UG/L		1	U	



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PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE		TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE &		QA/QC REMARK	
	FROM	TO							REMARK	REMARK		
				200128	34030	BENZENE	UG/L			1	U	
					32103	1,2-DICHLOROETHANE	UG/L			1	U	
					99999	1,3-DICHLOROPROPANE	UG/L			1	U	
					99999	1,1,1,2-TETRACHLOROETHANE	UG/L			1	U	
					34010	TOLUENE	UG/L			1	U	
					34475	TETRACHLOROETHYLENE	UG/L			1	U	
					99999	1,2,3-TRICHLOROPROPANE	UG/L			1	U	
					34516	1,1,2,2-TETRACHLOROETHANE	UG/L			1	U	
					99999	TRANS-1,3-DICHLOROPROPENE	UG/L			1	U	
					34511	1,1,2-TRICHLOROETHANE	UG/L			1	U	
					32105	CHLORODIBROMOMETHANE	UG/L			1	U	
					34301	CHLOROBENZENE	UG/L			1	U	
					34371	ETHYLBENZENE	UG/L			1	U	
					32104	BROMOFORM	UG/L			1	U	
					99999	BROMOBENZENE	UG/L			1	U	
					99999	ISOPROPYLBENZENE	UG/L			1	U	
					99921	STYRENE	UG/L			1	U	
					99902	O-XYLENE	UG/L			1	U	
					99999	P+M XYLENE	UG/L			1	U	
					99905	N-PROPYLBENZENE	UG/L			1	U	
					99999	1,2-DIBROMOETHANE	UG/L			1	U	
					99999	2-HEXANONE	UG/L			1	U	
					99999	2-CHLOROTOLUENE	UG/L			1	U	
					99999	4-CHLOROTOLUENE	UG/L			1	U	
					99999	TERTBUTYLBENZENE	UG/L			1	U	
					34566	1,3-DICHLOROBENZENE	UG/L			1	U	
					99999	SEC-BUTYLBENZENE	UG/L			1	U	
					34536	1,2-DICHLOROBENZENE	UG/L			1	U	
					34571	1,4-DICHLOROBENZENE	UG/L			1	U	
					99909	N-BUTYLBENZENE	UG/L			1	U	
					99999	1,2,4-TRIMETHYLBENZENE	UG/L			1	U	
					99907	1,3,5-TRIMETHYLBENZENE	UG/L			1	U	
					99999	4-ISOPROPYLTOLUENE	UG/L			1	U	
					34551	1,2,4-TRICHLOROBENZENE	UG/L			1	U	
					34696	NAPHTHALENE	UG/L			1	U	
					39702	HEXACHLOROBUTADIENE	UG/L			1	U	
					99999	1,2,3-TRICHLOROBENZENE	UG/L			1	U	
					99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L			1	U	
					99930	ACETONE	UG/L			7	U	
					99999	2-BUTANONE	UG/L			0.4	J	QM
					99999	BROMOCHLOROMETHANE	UG/L			1	U	

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STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200128	99999	4-METHYL-2-PENTANONE	UG/L		1 U	
				34273	BIS(2-CHLOROETHYL) ET.	UG/L		4.1 U	
				34694	PHENOL	UG/L		4.1 U	
				34586	2-CHLOROPHENOL	UG/L		4.1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		4.1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		4.1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		4.1 U	
				99999	BENZYL ALCOHOL	UG/L		4.1 U	
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		4.1 U	
				99999	2-METHYL PHENOL	UG/L		4.1 U	
				99999	4-METHYL PHENOL	UG/L		4.1 U	
				34396	HEXACHLOROETHANE	UG/L		4.1 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		4.1 U	
				34447	NITROBENZENE	UG/L		4.1 U	
				34408	ISOPHORONE	UG/L		4.1 U	
				34591	2-NITROPHENOL	UG/L		4.1 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		4.1 U	
				99999	BENZOIC ACID	UG/L		4.1 U	
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		4.1 U	
				34601	2,4-DICHLOROPHENOL	UG/L		4.1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		4.1 U	
				34696	NAPHTHALENE	UG/L		4.1 U	
				99999	4-CHLOROANILINE	UG/L		4.1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		4.1 U	
				34452	P-CHLORO-M-CRESOL	UG/L		4.1 U	
				99999	2-METHYL NAPHTHALENE	UG/L		4.1 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		33 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		4.1 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		4.1 U	
				34581	2-CHLORONAPHTHALENE	UG/L		4.1 U	
				99999	2-NITROANILINE	UG/L		4.1 U	
				34200	ACENAPHTHYLENE	UG/L		4.1 U	
				34341	DIMETHYL PHTHALATE	UG/L		4.1 U	
				34626	2,6-DINITROTOLUENE	UG/L		4.1 U	
				99999	3-NITROANILINE	UG/L		4.1 U	
				34205	ACENAPHTHENE	UG/L		4.1 U	
				34616	2,4-DINITROPHENOL	UG/L		33 U	
				99999	DIBENZOFURAN	UG/L		4.1 U	
				34646	4-NITROPHENOL	UG/L		4.1 U	
				34611	2,4-DINITROTOLUENE	UG/L		4.1 U	
				34381	FLUORENE	UG/L		4.1 U	

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STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200128	34641	4-CHLOROPHENYL PHENYL ET.	UG/L		4.1 U	
				99999	4-NITROANILINE	UG/L		4.1 U	
				34336	DIETHYL PHTHALATE	UG/L		4.1 U	
				34657	4,6-DINITRO-O-CRESOL	UG/L		33 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		4.1 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		4.1 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		4.1 U	
				39700	HEXACHLOROBENZENE	UG/L		4.1 U	
				39032	PENTACHLOROPHENOL	UG/L		33 U	
				34461	PHENANTHRENE	UG/L		4.1 U	
				34220	ANTHRACENE	UG/L		4.1 U	
				34376	FLUORANTHENE	UG/L		4.1 U	
				39110	DI-N-BUTYLPHTHALATE	UG/L		4.1 U	
				34469	PYRENE	UG/L		4.1 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		4.1 U	
				34526	1,2-BENZANTHRACENE	UG/L		4.1 U	
				34320	CHRYSENE	UG/L		4.1 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		4.1 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		4.1 U	
				34230	3,4-BENZOFUORANTHENE	UG/L		4.1 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		4.1 U	
				34247	BENZO(A)PYRENE	UG/L		4.1 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		4.1 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		4.1 U	
				34521	1,12-BENZOPERYLENE	UG/L		4.1 U	
				34524	OCTANOIC ACID	UG/L		3.7 J	QT
				34521	DODECANOIC ACID	UG/L		12 J	QT
				99999	DIACETATE-1,13-TRIDECANEDIOL	UG/L		5.5 J	QT
				99999	OLEIC ACID	UG/L		4.8 J	QT
				99999	UNKNOWN COMPOUND #1	UG/L		3.0 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINUM	UG/L		2900	
				01002	ARSENIC	UG/L		10 U	QR
				01007	BARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	
				00916	CALCIUM	MG/L		8	
				01027	CADMIUM	UG/L		5 U	
				01037	COBALT	UG/L		50 U	
				01034	CHROMIUM	UG/L		109	
				01042	COPPER	UG/L		25 U	
				01045	IRON	UG/L		5560 J	QP

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STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200128	71900	MERCURY	UG/L		0.2 U	
				00937	POTASSIUM	MG/L		5 U	
				00927	MAGNESIUM	MG/L		5 U	
				01055	MANGANESE	UG/L		119	
				00929	SODIUM	MG/L		32	
				01067	NICKEL	UG/L		40 U	
				01051	LEAD	UG/L		4.3	
				01097	ANTIMONY	UG/L		60 U	
				01147	SELENIUM	UG/L		5 U	QR
				01059	THALLIUM	UG/L		10 U	
				01087	VANADIUM	UG/L		50 U	
				01092	ZINC	UG/L		20 U	

NONE 96/04/02 1330  
 DEPTH: 67.0 SUBSTRATE: AQUEOUS  
 DESCRIPTION: MW-4D

200129	99999	CHLOROMETHANE	UG/L		1 U		
	99999	BROMOMETHANE	UG/L		1 U		
	39175	VINYL CHLORIDE	UG/L		1 U		
	34311	CHLOROETHANE	UG/L		1 U		
	34423	METHYLENE CHLORIDE	UG/L		1 U		
	34488	TRICHLOROFLUOROMETHANE	UG/L		1 U		
	34501	1,1-DICHLOROETHYLENE	UG/L		1 U		
	34496	1,1-DICHLOROETHANE	UG/L		0.3 J	QM	
	99964	CARBON DISULFIDE	UG/L		1 U		
	34546	TRANS 1,2 DICHLOROETHYLENE	UG/L		1 U		
	99999	CIS 1,2- DICHLOROETHYLENE	UG/L		1 U		
	99999	2,2 DICHLOROPROPANE	UG/L		1 U		
	32106	CHLOROFORM	UG/L		1 U		
	99999	DIBROMOMETHANE	UG/L		1 U		
	34506	1,1,1-TRICHLOROETHANE	UG/L		1.5		
	32102	CARBON TETRACHLORIDE	UG/L		1 U		
	32101	DICHLOROBROMOMETHANE	UG/L		1 U		
	99999	1,1-DICHLOROPROPENE	UG/L		1 U		
	34541	1,2-DICHLOROPROPANE	UG/L		1 U		
	99999	CIS-1,3-DICHLOROPROPENE	UG/L		1 U		
	39180	TRICHLOROETHYLENE	UG/L		1 U		
	34030	BENZENE	UG/L		1 U		
	32103	1,2-DICHLOROETHANE	UG/L		1 U		
	99999	1,3 DICHLOROPROPANE	UG/L		1 U		

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STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200129	99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		1 U	
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SECBUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		7 U	
				99999	2-BUTANONE	UG/L		1 U	
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
				34273	BIS(2-CHLOROETHYL) ET.	UG/L		3.8 U	
				34694	PHENOL	UG/L		3.8 U	

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PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200129	34586	2-CHLOROPHENOL	UG/L		3.8 U	
				34566	1,3-DICHLOROBENZENE	UG/L		3.8 U	
				34571	1,4-DICHLOROBENZENE	UG/L		3.8 U	
				34536	1,2-DICHLOROBENZENE	UG/L		3.8 U	
				99999	BENZYL ALCOHOL	UG/L		3.8 U	
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		3.8 U	
				99999	2-METHYL PHENOL	UG/L		3.8 U	
				99999	4-METHYL PHENOL	UG/L		3.8 U	
				34396	HEXACHLOROETHANE	UG/L		3.8 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		3.8 U	
				34447	NITROBENZENE	UG/L		3.8 U	
				34408	ISOPHORONE	UG/L		3.8 U	
				34591	2-NITROPHENOL	UG/L		3.8 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		3.8 U	
				99999	BENZOIC ACID	UG/L		3.8 U	
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		3.8 U	
				34601	2,4-DICHLOROPHENOL	UG/L		3.8 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		3.8 U	
				34696	NAPHTHALENE	UG/L		3.8 U	
				99999	4-CHLOROANILINE	UG/L		3.8 U	
				39702	HEXACHLOROBUTADIENE	UG/L		3.8 U	
				34452	P-CHLORO-M-CRESOL	UG/L		3.8 U	
				99999	2-METHYL NAPHTHALENE	UG/L		3.8 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		30 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		3.8 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		3.8 U	
				34581	2-CHLORONAPHTHALENE	UG/L		3.8 U	
				99999	2-NITROANILINE	UG/L		3.8 U	
				34200	ACENAPHTHYLENE	UG/L		3.8 U	
				34341	DIMETHYL PHTHALATE	UG/L		3.8 U	
				34626	2,6-DINITROTOLUENE	UG/L		3.8 U	
				99999	3-NITROANILINE	UG/L		3.8 U	
				34205	ACENAPHTHENE	UG/L		3.8 U	
				34616	2,4-DINITROPHENOL	UG/L		30 U	
				99999	DIBENZOFURAN	UG/L		3.8 U	
				34646	4-NITROPHENOL	UG/L		3.8 U	
				34611	2,4-DINITROTOLUENE	UG/L		3.8 U	
				34381	FLUORENE	UG/L		3.8 U	
				34641	4-CHLOROPHENYL PHENYL ET.	UG/L		3.8 U	
				99999	4-NITROANILINE	UG/L		3.8 U	
				34336	DIETHYL PHTHALATE	UG/L		3.8 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200129	34657	4,6-DINITRO-O-CRESOL	UG/L		30 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		3.8 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		3.8 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		3.8 U	
				39700	HEXACHLOROBENZENE	UG/L		3.8 U	
				39032	PENTACHLOROPHENOL	UG/L		30 U	
				34461	PHENANTHRENE	UG/L		3.8 U	
				34220	ANTHRACENE	UG/L		3.8 U	
				34376	FLUORANTHENE	UG/L		3.8 U	
				39110	DI-N-BUTYLPHTHALATE	UG/L		3.8 U	
				34469	PYRENE	UG/L		3.8 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		3.8 U	
				34526	1,2-BENZANTHRACENE	UG/L		3.8 U	
				34320	CHRYSENE	UG/L		3.8 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		3.8 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		3.8 U	
				34230	3,4-BENZOFLUORANTHENE	UG/L		3.8 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		3.8 U	
				34247	BENZO(A)PYRENE	UG/L		3.8 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		3.8 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		3.8 U	
				34521	1,12-BENZOPERYLENE	UG/L		3.8 U	
				34524	OCTANOIC ACID	UG/L		4.8 J	QT
				34521	DODECANOIC ACID	UG/L		17 J	QT
				99999	DECANOIC ACID	UG/L		2.4 J	QT
				99999	UNKNOWN COMPOUND #1	UG/L		2.5 J	QT
				99999	UNKNOWN COMPOUND #2	UG/L		5.9 J	QT
				99999	UNKNOWN COMPOUND #3	UG/L		34 J	QT
				99999	UNKNOWN COMPOUND #4	UG/L		5.2 J	QT
				99999	UNKNOWN COMPOUND #5	UG/L		38 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINUM	UG/L		3280	
				01002	ARSENIC	UG/L		10 U	
				01007	BARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	
				00916	CALCIUM	MG/L		8	
				01027	CADMIUM	UG/L		5 U	
				01037	COBALT	UG/L		50 U	
				01034	CHROMIUM	UG/L		91	
				01042	COPPER	UG/L		25 U	
				01045	IRON	UG/L		5190 J	QP

109.8

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200129	71900	MERCURY	UG/L		0.2 U	
				00937	POTASSIUM	MG/L		5 U	
				00927	MAGNESIUM	MG/L		5 U	
				01055	MANGANESE	UG/L		125	
				00929	SODIUM	MG/L		33	
				01067	NICKEL	UG/L		40 U	
				01051	LEAD	UG/L		4.3	
				01097	ANTIMONY	UG/L		60 U	
				01147	SELENIUM	UG/L		5 U	QR
				01059	THALLIUM	UG/L		10 U	
				01087	VANADIUM	UG/L		50 U	
				01092	ZINC	UG/L		20 U	

NONE 96/04/03 1408  
 DEPTH: 63.4 SUBSTRATE: AQUEOUS  
 DESCRIPTION: MW-5D

200130	99999	CHLOROMETHANE	UG/L	1 U
	99999	BROMOMETHANE	UG/L	1 U
	39175	VINYL CHLORIDE	UG/L	1 U
	34311	CHLOROETHANE	UG/L	1 U
	34423	METHYLENE CHLORIDE	UG/L	1 U
	34488	TRICHLOROFLUOROMETHANE	UG/L	1 U
	34501	1,1-DICHLOROETHYLENE	UG/L	1 U
	34496	1,1-DICHLOROETHANE	UG/L	1 U
	99964	CARBON DISULFIDE	UG/L	1 U
	34546	TRANS 1,2 DICHLOROETHYLENE	UG/L	1 U
	99999	CIS 1,2- DICHLOROETHYLENE	UG/L	1 U
	99999	2,2 DICHLOROPROPANE	UG/L	1 U
	32106	CHLOROFORM	UG/L	1 U
	99999	DIBROMOMETHANE	UG/L	1 U
	34506	1,1,1-TRICHLOROETHANE	UG/L	1 U
	32102	CARBON TETRACHLORIDE	UG/L	1 U
	32101	DICHLOROBROMOMETHANE	UG/L	1 U
	99999	1,1-DICHLOROPROPENE	UG/L	1 U
	34541	1,2-DICHLOROPROPANE	UG/L	1 U
	99999	CIS-1,3-DICHLOROPROPENE	UG/L	1 U
	39180	TRICHLOROETHYLENE	UG/L	1 U
	34030	BENZENE	UG/L	1 U
	32103	1,2-DICHLOROETHANE	UG/L	1 U
	99999	1,3 DICHLOROPROPANE	UG/L	1 U



## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200130	99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		1 U	
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORO DIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SECBUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		6 U	
				99999	2-BUTANONE	UG/L		1 U	
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
				34273	BIS(2-CHLOROETHYL) ET.	UG/L		4.1 U	
				34694	PHENOL	UG/L		4.1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200130	34586	2-CHLOROPHENOL	UG/L		4.1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		4.1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		4.1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		4.1 U	
				99999	BENZYL ALCOHOL	UG/L		4.1 U	
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		4.1 U	
				99999	2-METHYL PHENOL	UG/L		4.1 U	
				99999	4-METHYL PHENOL	UG/L		4.1 U	
				34396	HEXACHLOROETHANE	UG/L		4.1 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		4.1 U	
				34447	NITROBENZENE	UG/L		4.1 U	
				34408	ISOPHORONE	UG/L		4.1 U	
				34591	2-NITROPHENOL	UG/L		4.1 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		4.1 U	
				99999	BENZOIC ACID	UG/L		4.1 U	
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		4.1 U	
				34601	2,4-DICHLOROPHENOL	UG/L		4.1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		4.1 U	
				34696	NAPHTHALENE	UG/L		4.1 U	
				99999	4-CHLOROANILINE	UG/L		4.1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		4.1 U	
				34452	P-CHLORO-M-CRESOL	UG/L		4.1 U	
				99999	2-METHYL NAPHTHALENE	UG/L		4.1 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		33 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		4.1 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		4.1 U	
				34581	2-CHLORONAPHTHALENE	UG/L		4.1 U	
				99999	2-NITROANILINE	UG/L		4.1 U	
				34200	ACENAPHTHYLENE	UG/L		4.1 U	
				34341	DIMETHYL PHTHALATE	UG/L		4.1 U	
				34626	2,6-DINITROTOLUENE	UG/L		4.1 U	
				99999	3-NITROANILINE	UG/L		4.1 U	
				34205	ACENAPHTHENE	UG/L		4.1 U	
				34616	2,4-DINITROPHENOL	UG/L		33 U	
				99999	DIBENZOFURAN	UG/L		4.1 U	
				34646	4-NITROPHENOL	UG/L		4.1 U	
				34611	2,4-DINITROTOLUENE	UG/L		4.1 U	
				34381	FLUORENE	UG/L		4.1 U	
				34641	4-CHLOROPHENYL PHENYL ET.	UG/L		4.1 U	
				99999	4-NITROANILINE	UG/L		4.1 U	
				34336	DIETHYL PHTHALATE	UG/L		4.1 U	

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REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200130	34657	4,6-DINITRO-O-CRESOL	UG/L		33 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		4.1 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		4.1 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		4.1 U	
				39700	HEXACHLOROBENZENE	UG/L		4.1 U	
				39032	PENTACHLOROPHENOL	UG/L		33 U	
				34461	PHENANTHRENE	UG/L		4.1 U	
				34220	ANTHRACENE	UG/L		4.1 U	
				34376	FLUORANTHENE	UG/L		4.1 U	
				39110	DI-N-BUTYLPHthalate	UG/L		4.1 U	
				34469	PYRENE	UG/L		4.1 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		4.1 U	
				34526	1,2-BENZANTHRACENE	UG/L		4.1 U	
				34320	CHRYSENE	UG/L		4.1 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		4.1 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		4.1 U	
				34230	3,4-BENZOFUORANTHENE	UG/L		4.1 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		4.1 U	
				34247	BENZO(A)PYRENE	UG/L		4.1 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		4.1 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		4.1 U	
				34521	1,12-BENZOPERYLENE	UG/L		4.1 U	
				34524	OCTANOIC ACID	UG/L		4.9 J	QT
				34521	DODECANOIC ACID	UG/L		20 J	QT
				34524	TETRADECANOIC ACID	UG/L		5.4 J	QT
				99999	OLEYL ALCOHOL	UG/L		12 J	QT
				99999	METHYLESTER 9-HEXADECENOIC A	UG/L		20 J	QT
				99999	OCTADECANOIC ACID	UG/L		5.2 J	QT
				99999	UNKNOWN COMPOUND #1	UG/L		4.8 J	QT
				99999	UNKNOWN COMPOUND #2	UG/L		4.4 J	QT
				99999	UNKNOWN COMPOUND #3	UG/L		39 J	QT
				99999	BIS(2-ETHYLHEXYL)HEXANEDIOIC	UG/L		5.4 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINUM	UG/L		688	
				01002	ARSENIC	UG/L		10 U	
				01007	BIARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	
				00916	CALCIUM	MG/L		15	
				01027	CADMIUM	UG/L		5 U	
				01037	COBALT	UG/L		50 U	
				01034	CHROMIUM	UG/L		80	

121.1

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200130	01042	COPPER	UG/L		52	
				01045	IRON	UG/L		793 J	QP
				71900	MERCURY	UG/L		0.2 U	
				00937	POTASSIUM	MG/L		5 U	
				00927	MAGNESIUM	MG/L		5 U	
				01055	MANGANESE	UG/L		23	
				00929	SODIUM	MG/L		52	
				01067	NICKEL	UG/L		41	
				01051	LEAD	UG/L		6.5	
				01097	ANTIMONY	UG/L		60 U	
				01147	SELENIUM	UG/L		5 U	
				01059	THALLIUM	UG/L		10 U	
				01087	VANADIUM	UG/L		50 U	
				01092	ZINC	UG/L		97	
			200131	99999	CHLOROMETHANE	UG/L		1 U	
				99999	BROMOMETHANE	UG/L		1 U	
				39175	VINYL CHLORIDE	UG/L		1 U	
				34311	CHLOROETHANE	UG/L		1 U	
				34423	METHYLENE CHLORIDE	UG/L		1 U	
				34488	TRICHLOROFLUOROMETHANE	UG/L		1 U	
				34501	1,1-DICHLOROETHYLENE	UG/L		1 U	
				34496	1,1-DICHLOROETHANE	UG/L		1 U	
				99964	CARBON DISULFIDE	UG/L		1 U	
				34546	TRANS 1,2 DICHLOROETHYLENE	UG/L		1 U	
				99999	CIS 1,2- DICHLOROETHYLENE	UG/L		1 U	
				99999	2,2 DICHLOROPROPANE	UG/L		1 U	
				32106	CHLOROFORM	UG/L		1 U	
				99999	DIBROMOMETHANE	UG/L		1 U	
				34506	1,1,1-TRICHLOROETHANE	UG/L		1 U	
				32102	CARBON TETRACHLORIDE	UG/L		1 U	
				32101	DICHLOROBROMOMETHANE	UG/L		1 U	
				99999	1,1-DICHLOROPROPENE	UG/L		1 U	
				34541	1,2-DICHLOROPROPANE	UG/L		1 U	
				99999	CIS-1,3-DICHLOROPROPENE	UG/L		1 U	
				39180	TRICHLOROETHYLENE	UG/L		1 U	
				34030	BENZENE	UG/L		1 U	

NONE 96/04/02 1300  
 DEPTH: 63.4 SUBSTRATE: AQUEOUS  
 DESCRIPTION: MW-5S

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200131	32103	1,2-DICHLOROETHANE	UG/L		1 U	
				99999	1,3 DICHLOROPROPANE	UG/L		1 U	
				99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		1 U	
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SECBUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		7 U	
				99999	2-BUTANONE	UG/L		0.4 J	QM
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200131	34273	BIS(2-CHLOROETHYL) ET.	UG/L		3.8 U	
				34694	PHENOL	UG/L		3.8 U	
				34586	2-CHLOROPHENOL	UG/L		3.8 U	
				34566	1,3-DICHLOROBENZENE	UG/L		3.8 U	
				34571	1,4-DICHLOROBENZENE	UG/L		3.8 U	
				34536	1,2-DICHLOROBENZENE	UG/L		3.8 U	
				99999	BENZYL ALCOHOL	UG/L		3.8 U	
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		3.8 U	
				99999	2-METHYL PHENOL	UG/L		3.8 U	
				99999	4-METHYL PHENOL	UG/L		3.8 U	
				34396	HEXACHLOROETHANE	UG/L		3.8 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		3.8 U	
				34447	NITROBENZENE	UG/L		3.8 U	
				34408	ISOPHORONE	UG/L		3.8 U	
				34591	2-NITROPHENOL	UG/L		3.8 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		3.8 U	
				99999	BENZOIC ACID	UG/L		0.8 J	QM
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		3.8 U	
				34601	2,4-DICHLOROPHENOL	UG/L		3.8 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		3.8 U	
				34696	NAPHTHALENE	UG/L		3.8 U	
				99999	4-CHLOROANILINE	UG/L		3.8 U	
				39702	HEXACHLOROBUTADIENE	UG/L		3.8 U	
				34452	P-CHLORO-M-CRESOL	UG/L		3.8 U	
				99999	2-METHYL NAPHTHALENE	UG/L		3.8 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		30 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		3.8 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		3.8 U	
				34581	2-CHLORONAPHTHALENE	UG/L		3.8 U	
				99999	2-NITROANILINE	UG/L		3.8 U	
				34200	ACENAPHTHYLENE	UG/L		3.8 U	
				34341	DIMETHYL PHTHALATE	UG/L		3.8 U	
				34626	2,6-DINITROTOLUENE	UG/L		3.8 U	
				99999	3-NITROANILINE	UG/L		3.8 U	
				34205	ACENAPHTHENE	UG/L		3.8 U	
				34616	2,4-DINITROPHENOL	UG/L		30 U	
				99999	DIBENZOFURAN	UG/L		3.8 U	
				34646	4-NITROPHENOL	UG/L		3.8 U	
				34611	2,4-DINITROTOLUENE	UG/L		3.8 U	
				34381	FLUORENE	UG/L		3.8 U	
				34641	4-CHLOROPHENYL PHENYL ET.	UG/L		3.8 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200131	99999	4-NITROANILINE	UG/L		3.8 U	
				34336	DIETHYL PHTHALATE	UG/L		3.8 U	
				34657	4,6-DINITRO-O-CRESOL	UG/L		30 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		3.8 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		3.8 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		3.8 U	
				39700	HEXACHLOROBENZENE	UG/L		3.8 U	
				39032	PENTACHLOROPHENOL	UG/L		30 U	
				34461	PHENANTHRENE	UG/L		3.8 U	
				34220	ANTHRACENE	UG/L		3.8 U	
				34376	FLUORANTHENE	UG/L		0.2 J	QM
				39110	DI-N-BUTYLPHTHALATE	UG/L		3.8 U	
				34469	PYRENE	UG/L		3.8 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		3.8 U	
				34526	1,2-BENZANTHRACENE	UG/L		3.8 U	
				34320	CHRYSENE	UG/L		3.8 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		3.8 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		3.8 U	
				34230	3,4-BENZOFUORANTHENE	UG/L		3.8 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		3.8 U	
				34247	BENZO(A)PYRENE	UG/L		3.8 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		3.8 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		3.8 U	
				34521	1,12-BENZOPERYLENE	UG/L		3.8 U	
				34524	OCTANOIC ACID	UG/L		8.0 J	QT
				34521	DODECAHOIC ACID	UG/L		30 J	QT
				99999	OLEIC ACID	UG/L		13 J	QT
				99999	DECAHOIC ACID	UG/L		4.9 J	QT
				00929	NONANOIC ACID	UG/L		15 J	QT
				34524	TETRADECAHOIC ACID	UG/L		6.9 J	QT
				99999	UNKNOWN COMPOUND #1	UG/L		5.6 J	QT
				99999	UNKNOWN COMPOUND #2	UG/L		38 J	QT
				99999	UNKNOWN COMPOUND #3	UG/L		36 J	QT
				99999	UNKNOWN COMPOUND #4	UG/L		3.1 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINUM	UG/L		2020	
				01002	ARSENIC	UG/L		10 U	
				01007	BARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	
				00916	CALCIUM	MG/L		15	
				01027	CADMIUM	UG/L		5 U	

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## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200131	01037	COBALT	UG/L		50 U	
				01034	CHROMIUM	UG/L		108	
				01042	COPPER	UG/L		25 U	
				01045	IRON	UG/L		3150 J	QP
				71900	MERCURY	UG/L		0.2 U	
				00937	POTASSIUM	MG/L		5 U	
				00927	MAGNESIUM	MG/L		5 U	
				01055	MANGANESE	UG/L		61	
				00929	SODIUM	MG/L		35	
				01067	NICKEL	UG/L		55	
				01051	LEAD	UG/L		5.6	
				01097	ANTIMONY	UG/L		60 U	
				01147	SELENIUM	UG/L		5 U	
				01059	THALLIUM	UG/L		10 U	
				01087	VANADIUM	UG/L		50 U	
				01092	ZINC	UG/L		95	

NONE 96/03/29 1413  
 DEPTH: 0070 SUBSTRATE: AQUEOUS  
 DESCRIPTION: MW-6D

200132	99999	CHLOROMETHANE	UG/L	1 U
	99999	BROMOMETHANE	UG/L	1 U
	39175	VINYL CHLORIDE	UG/L	1 U
	34311	CHLOROETHANE	UG/L	1 U
	34423	METHYLENE CHLORIDE	UG/L	1 U
	34488	TRICHLOROFLUOROMETHANE	UG/L	1 U
	34501	1,1-DICHLOROETHYLENE	UG/L	1 U
	34496	1,1-DICHLOROETHANE	UG/L	1 U
	99964	CARBON DISULFIDE	UG/L	1 U
	34546	TRANS 1,2 DICHLOROETHYLENE	UG/L	1 U
	99999	CIS 1,2- DICHLOROETHYLENE	UG/L	1 U
	99999	2,2 DICHLOROPROPANE	UG/L	1 U
	32106	CHLOROFORM	UG/L	1 U
	99999	DIBROMOMETHANE	UG/L	1 U
	34506	1,1,1-TRICHLOROETHANE	UG/L	1 U
	32102	CARBON TETRACHLORIDE	UG/L	1 U
	32101	DICHLOROBROMOMETHANE	UG/L	1 U
	99999	1,1-DICHLOROPROPENE	UG/L	1 U
	34541	1,2-DICHLOROPROPANE	UG/L	1 U
	99999	CIS-1,3-DICHLOROPROPENE	UG/L	1 U



## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200132	39180	TRICHLOROETHYLENE	UG/L		1 U	
				34030	BENZENE	UG/L		1 U	
				32103	1,2-DICHLOROETHANE	UG/L		1 U	
				99999	1,3-DICHLOROPROPANE	UG/L		1 U	
				99999	1,1,1,2-TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		1 U	
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SEC-BUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		7 U	
				99999	2-BUTANONE	UG/L		1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200132	99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
				34273	BIS(2-CHLOROETHYL) ET.	UG/L		3.8 U	
				34694	PHENOL	UG/L		3.8 U	
				34586	2-CHLOROPHENOL	UG/L		3.8 U	
				34566	1,3-DICHLOROBENZENE	UG/L		3.8 U	
				34571	1,4-DICHLOROBENZENE	UG/L		3.8 U	
				34536	1,2-DICHLOROBENZENE	UG/L		3.8 U	
				99999	BENZYL ALCOHOL	UG/L		3.8 U	
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		3.8 U	
				99999	2-METHYL PHENOL	UG/L		3.8 U	
				99999	4-METHYL PHENOL	UG/L		3.8 U	
				34396	HEXACHLOROETHANE	UG/L		3.8 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		3.8 U	
				34447	NITROBENZENE	UG/L		3.8 U	
				34408	ISOPHORONE	UG/L		3.8 U	
				34591	2-NITROPHENOL	UG/L		3.8 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		3.8 U	
				99999	BENZOIC ACID	UG/L		3.8 U	
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		3.8 U	
				34601	2,4-DICHLOROPHENOL	UG/L		3.8 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		3.8 U	
				34696	NAPHTHALENE	UG/L		3.8 U	
				99999	4-CHLOROANILINE	UG/L		3.8 U	
				39702	HEXACHLOROBUTADIENE	UG/L		3.8 U	
				34452	P-CHLORO-M-CRESOL	UG/L		3.8 U	
				99999	2-METHYL NAPHTHALENE	UG/L		3.8 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		30 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		3.8 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		3.8 U	
				34581	2-CHLORONAPHTHALENE	UG/L		3.8 U	
				99999	2-NITROANILINE	UG/L		3.8 U	
				34200	ACENAPHTHYLENE	UG/L		3.8 U	
				34341	DIMETHYL PHTHALATE	UG/L		3.8 U	
				34626	2,6-DINITROTOLUENE	UG/L		3.8 U	
				99999	3-NITROANILINE	UG/L		3.8 U	
				34205	ACENAPHTHENE	UG/L		3.8 U	
				34616	2,4-DINITROPHENOL	UG/L		30 U	
				99999	DIBENZOFURAN	UG/L		3.8 U	
				34646	4-NITROPHENOL	UG/L		3.8 U	
				34611	2,4-DINITROTOLUENE	UG/L		3.8 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200132	34381	FLUORENE	UG/L		3.8 U	
				34641	4-CHLOROPHENYL PHENYL ET.	UG/L		3.8 U	
				99999	4-NITROANILINE	UG/L		3.8 U	
				34336	DIETHYL PHTHALATE	UG/L		3.8 U	
				34657	4,6-DINITRO-O-CRESOL	UG/L		30 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		3.8 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		3.8 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		3.8 U	
				39700	HEXACHLOROBENZENE	UG/L		3.8 U	
				39032	PENTACHLOROPHENOL	UG/L		30 U	
				34461	PHENANTHRENE	UG/L		3.8 U	
				34220	ANTHRACENE	UG/L		3.8 U	
				34376	FLUORANTHENE	UG/L		3.8 U	
				39110	DI-N-BUTYLPHthalate	UG/L		3.8 U	
				34469	PYRENE	UG/L		3.8 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		3.8 U	
				34526	1,2-BENZANTHRACENE	UG/L		3.8 U	
				34320	CHRYSENE	UG/L		3.8 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		3.8 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		3.8 U	
				34230	3,4-BENZOFUORANTHENE	UG/L		3.8 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		3.8 U	
				34247	BENZO(A)PYRENE	UG/L		3.8 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		3.8 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		3.8 U	
				34521	1,12-BENZOPERYLENE	UG/L		3.8 U	
				34524	OCTANOIC ACID	UG/L		2.7 J	QT
				99999	N,N-BIS(2-HYDROXYETHYL)DODEC	UG/L		7.9 J	QT
				99999	(Z)11-HEXADECEN-1-OL	UG/L		2.8 J	QT
				99999	9-HEXADECANOIC ACID	UG/L		6.2 J	QT
				99999	UNKNOWN COMPOUND #1	UG/L		2.6 J	QT
				99999	UNKNOWN COMPOUND #2	UG/L		5.2 J	QT
				99999	UNKNOWN COMPOUND #3	UG/L		3.8 J	QT
				99999	UNKNOWN COMPOUND #4	UG/L		35 J	QT
				99999	UNKNOWN COMPOUND #5	UG/L		68 J	QT
				99999	UNKNOWN COMPOUND #6	UG/L		7.0 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINUM	UG/L		200 U	
				01002	ARSENIC	UG/L		10 U	
				01007	BARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200132	00916	CALCIUM	MG/L		13	
				01027	CADMIUM	UG/L		5 U	
				01037	COBALT	UG/L		50 U	
				01034	CHROMIUM	UG/L		139	
				01042	COPPER	UG/L		25 U	
				01045	IRON	UG/L		640 J	QP
				71900	MERCURY	UG/L		0.2 U	
				00937	POTASSIUM	MG/L		5 U	
				00927	MAGNESIUM	MG/L		5 U	
				01055	MANGANESE	UG/L		27	
				00929	SODIUM	MG/L		46	
				01067	NICKEL	UG/L		149	
				01051	LEAD	UG/L		3 U	
				01097	ANTIMONY	UG/L		60 U	
				01147	SELENIUM	UG/L		5 U	
				01059	THALLIUM	UG/L		10 U	
				01087	VANADIUM	UG/L		50 U	
				01092	ZINC	UG/L		20 U	

NONE 96/03/29 1404  
 DEPTH: 0063 SUBSTRATE: AQUEOUS  
 DESCRIPTION: MW-6S

200133	99999	CHLOROMETHANE	UG/L	1 U	
	99999	BROMOMETHANE	UG/L	1 U	
	39175	VINYL CHLORIDE	UG/L	1 U	
	34311	CHLOROETHANE	UG/L	1 U	
	34423	METHYLENE CHLORIDE	UG/L	1 U	
	34488	TRICHLOROFLUOROMETHANE	UG/L	1 U	
	34501	1,1-DICHLOROETHYLENE	UG/L	1 U	
	34496	1,1-DICHLOROETHANE	UG/L	1 U	
	99964	CARBON DISULFIDE	UG/L	0.3 J	QM
	34546	TRANS 1,2 DICHLOROETHYLENE	UG/L	1 U	
	99999	CIS 1,2- DICHLOROETHYLENE	UG/L	1 U	
	99999	2,2 DICHLOROPROPANE	UG/L	1 U	
	32106	CHLOROFORM	UG/L	1 U	
	99999	DIBROMOMETHANE	UG/L	1 U	
	34506	1,1,1-TRICHLOROETHANE	UG/L	1 U	
	32102	CARBON TETRACHLORIDE	UG/L	1 U	
	32101	DICHLOROBROMOMETHANE	UG/L	1 U	
	99999	1,1-DICHLOROPROPENE	UG/L	1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE		TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE &		QA/QC
	FROM	TO							REMARK	REMARK	
				200133	34541	1,2-DICHLOROPROPANE	UG/L				1 U
					99999	CIS-1,3-DICHLOROPROPENE	UG/L				1 U
					39180	TRICHLOROETHYLENE	UG/L				1 U
					34030	BENZENE	UG/L				1 U
					32103	1,2-DICHLOROETHANE	UG/L				1 U
					99999	1,3 DICHLOROPROPANE	UG/L				1 U
					99999	1,1,1,2 TETRACHLOROETHANE	UG/L				1 U
					34010	TOLUENE	UG/L				1 U
					34475	TETRACHLOROETHYLENE	UG/L				1 U
					99999	1,2,3 TRICHLOROPROPANE	UG/L				1 U
					34516	1,1,2,2-TETRACHLOROETHANE	UG/L				1 U
					99999	TRANS-1,3-DICHLOROPROPENE	UG/L				1 U
					34511	1,1,2-TRICHLOROETHANE	UG/L				1 U
					32105	CHLORODIBROMOMETHANE	UG/L				1 U
					34301	CHLOROBENZENE	UG/L				1 U
					34371	ETHYLBENZENE	UG/L				1 U
					32104	BROMOFORM	UG/L				1 U
					99999	BROMOBENZENE	UG/L				1 U
					99999	ISOPROPYLBENZENE	UG/L				1 U
					99921	STYRENE	UG/L				1 U
					99902	O-XYLENE	UG/L				1 U
					99999	P+M XYLENE	UG/L				1 U
					99905	N-PROPYLBENZENE	UG/L				1 U
					99999	1,2-DIBROMOETHANE	UG/L				1 U
					99999	2-HEXANONE	UG/L				1 U
					99999	2-CHLOROTOLUENE	UG/L				1 U
					99999	4-CHLOROTOLUENE	UG/L				1 U
					99999	TERTBUTYLBENZENE	UG/L				1 U
					34566	1,3-DICHLOROBENZENE	UG/L				1 U
					99999	SECBUTYLBENZENE	UG/L				1 U
					34536	1,2-DICHLOROBENZENE	UG/L				1 U
					34571	1,4-DICHLOROBENZENE	UG/L				1 U
					99909	N-BUTYLBENZENE	UG/L				1 U
					99999	1,2,4-TRIMETHYLBENZENE	UG/L				1 U
					99907	1,3,5-TRIMETHYLBENZENE	UG/L				1 U
					99999	4-ISOPROPYLTOLUENE	UG/L				1 U
					34551	1,2,4-TRICHLOROBENZENE	UG/L				1 U
					34696	NAPHTHALENE	UG/L				1 U
					39702	HEXACHLOROBUTADIENE	UG/L				1 U
					99999	1,2,3-TRICHLOROBENZENE	UG/L				1 U
					99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L				1 U

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200133	99930	ACETONE	UG/L		7 U	
				99999	2-BUTANONE	UG/L		1 U	
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
				34273	BIS(2-CHLOROETHYL) ET.	UG/L		3.8 U	
				34694	PHENOL	UG/L		3.8 U	
				34586	2-CHLOROPHENOL	UG/L		3.8 U	
				34566	1,3-DICHLOROBENZENE	UG/L		3.8 U	
				34571	1,4-DICHLOROBENZENE	UG/L		3.8 U	
				34536	1,2-DICHLOROBENZENE	UG/L		3.8 U	
				99999	BENZYL ALCOHOL	UG/L		0.2 J	QM
				34283	BIS(2-CHLOROISOPROPYL) ET.	UG/L		3.8 U	
				99999	2-METHYL PHENOL	UG/L		3.8 U	
				99999	4-METHYL PHENOL	UG/L		3.8 U	
				34396	HEXACHLOROETHANE	UG/L		3.8 U	
				34428	N-NITROSODI-N-PROPYLAMINE	UG/L		3.8 U	
				34447	NITROBENZENE	UG/L		3.8 U	
				34408	ISOPHORONE	UG/L		3.8 U	
				34591	2-NITROPHENOL	UG/L		3.8 U	
				34606	2,4-DIMETHYLPHENOL	UG/L		3.8 U	
				99999	BENZOIC ACID	UG/L		3.8 U	
				34278	BIS(2-CHLOROETHOXY) METH.	UG/L		3.8 U	
				34601	2,4-DICHLOROPHENOL	UG/L		3.8 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		3.8 U	
				34696	NAPHTHALENE	UG/L		3.8 U	
				99999	4-CHLOROANILINE	UG/L		3.8 U	
				39702	HEXACHLOROBUTADIENE	UG/L		3.8 U	
				34452	P-CHLORO-M-CRESOL	UG/L		3.8 U	
				99999	2-METHYL NAPHTHALENE	UG/L		3.8 U	
				34386	HEXACHLOROCYCLOPENTADIENE	UG/L		30 U	
				34621	2,4,6-TRICHLOROPHENOL	UG/L		3.8 U	
				88894	2,4,5-TRICHLOROPHENOL	UG/L		3.8 U	
				34581	2-CHLORONAPHTHALENE	UG/L		3.8 U	
				99999	2-NITROANILINE	UG/L		3.8 U	
				34200	ACENAPHTHYLENE	UG/L		3.8 U	
				34341	DIMETHYL PHTHALATE	UG/L		3.8 U	
				34626	2,6-DINITROTOLUENE	UG/L		3.8 U	
				99999	3-NITROANILINE	UG/L		3.8 U	
				34205	ACENAPHTHENE	UG/L		3.8 U	
				34616	2,4-DINITROPHENOL	UG/L		30 U	
				99999	DIBENZOFURAN	UG/L		3.8 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200133	34646	4-NITROPHENOL	UG/L		3.8 U	
				34611	2,4-DINITROTOLUENE	UG/L		3.8 U	
				34381	FLUORENE	UG/L		3.8 U	
				34641	4-CHLOROPHENYL PHENYL ET.	UG/L		3.8 U	
				99999	4-NITROANILINE	UG/L		3.8 U	
				34336	DIETHYL PHTHALATE	UG/L		3.8 U	
				34657	4,6-DINITRO-O-CRESOL	UG/L		30 U	
				34433	N-NITROSODIPHENYLAMINE	UG/L		3.8 U	
				34346	1,2-DIPHENYLHYDRAZINE	UG/L		3.8 U	
				34636	4-BROMOPHENYL PHENYL ET.	UG/L		3.8 U	
				39700	HEXACHLOROBENZENE	UG/L		3.8 U	
				39032	PENTACHLOROPHENOL	UG/L		30 U	
				34461	PHENANTHRENE	UG/L		3.8 U	
				34220	ANTHRACENE	UG/L		3.8 U	
				34376	FLUORANTHENE	UG/L		3.8 U	
				39110	DI-N-BUTYLPHTHALATE	UG/L		3.8 U	
				34469	PYRENE	UG/L		3.8 U	
				34292	BUTYL BENZYL PHTHALATE	UG/L		3.8 U	
				34526	1,2-BENZANTHRACENE	UG/L		3.8 U	
				34320	CHRYSENE	UG/L		3.8 U	
				39100	BIS(2-ETHYLHEXYL) PHTHAL.	UG/L		3.8 U	
				34596	DI-N-OCTYL PHTHALATE	UG/L		3.8 U	
				34230	3,4-BENZOFUORANTHENE	UG/L		3.8 U	
				34242	11,12-BENZOFUORANTHENE	UG/L		3.8 U	
				34247	BENZO(A)PYRENE	UG/L		3.8 U	
				34403	INDENO(1,2,3-C,D) PYRENE	UG/L		3.8 U	
				34556	1,2:5,6-DIBENZANTHRACENE	UG/L		3.8 U	
				34521	1,12-BENZOPERYLENE	UG/L		3.8 U	
				34524	OCTANOIC ACID	UG/L		8.2 J	QT
				34521	DODECANOIC ACID	UG/L		33 J	QT
				99999	OLEIC ACID	UG/L		12 J	QT
				34524	TETRADECANOIC ACID	UG/L		7.1 J	QT
				99999	(Z)11-HEXADECEN-1-OL	UG/L		9.2 J	QT
				99999	UNKNOWN COMPOUND #1	UG/L		11 J	QT
				99999	UNKNOWN COMPOUND #2	UG/L		9.5 J	QT
				99999	UNKNOWN COMPOUND #3	UG/L		64 J	QT
				99999	UNKNOWN COMPOUND #4	UG/L		6.2 J	QT
				99999	UNKNOWN COMPOUND #5	UG/L		8.0 J	QT
				01077	SILVER	UG/L		10 U	
				01105	ALUMINIUM	UG/L		1910	
				01002	ARSENIC	UG/L		10 U	

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## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200133	01007	BARIUM	UG/L		200 U	
				01012	BERYLLIUM	UG/L		5 U	
				00916	CALCIUM	MG/L		16	
				01027	CADMIUM	UG/L		5 U	
				01037	COBALT	UG/L		50 U	
				01034	CHROMIUM	UG/L		126	
				01042	COPPER	UG/L		25 U	
				01045	IRON	UG/L		2840 J	QP
				71900	MERCURY	UG/L		0.2 U	
				00937	POTASSIUM	MG/L		5 U	
				00927	MAGNESIUM	MG/L		5 U	
				01055	MANGANESE	UG/L		63	
				00929	SODIUM	MG/L		42	
				01067	NICKEL	UG/L		64	
				01051	LEAD	UG/L		8.0	
				01097	ANTIMONY	UG/L		60 U	
				01147	SELENIUM	UG/L		5 U	
				01059	THALLIUM	UG/L		10 U	
				01087	VANADIUM	UG/L		50 U	
				01092	ZINC	UG/L		56	

NONE 96/04/02 1015  
 DEPTH: 0000 SUBSTRATE: AQUEOUS  
 DESCRIPTION: TRIP BLANK #2

200134	99999	CHLOROMETHANE	UG/L	1 U
	99999	BROMOMETHANE	UG/L	1 U
	39175	VINYL CHLORIDE	UG/L	1 U
	34311	CHLOROETHANE	UG/L	1 U
	34423	METHYLENE CHLORIDE	UG/L	1 U
	34488	TRICHLOROFUOROMETHANE	UG/L	1 U
	34501	1,1-DICHLOROETHYLENE	UG/L	1 U
	34496	1,1-DICHLOROETHANE	UG/L	1 U
	99964	CARBON DISULFIDE	UG/L	1 U
	34546	TRANS 1,2 DICHLOROETHYLENE	UG/L	1 U
	99999	CIS 1,2- DICHLOROETHYLENE	UG/L	1 U
	99999	2,2 DICHLOROPROPANE	UG/L	1 U
	32106	CHLOROFORM	UG/L	1.2
	99999	DIBROMOMETHANE	UG/L	1 U
	34506	1,1,1-TRICHLOROETHANE	UG/L	1 U
	32102	CARBON TETRACHLORIDE	UG/L	1 U



## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200134	32101	DICHLOROBROMOMETHANE	UG/L		1 U	
				99999	1,1-DICHLOROPROPENE	UG/L		1 U	
				34541	1,2-DICHLOROPROPANE	UG/L		1 U	
				99999	CIS-1,3-DICHLOROPROPENE	UG/L		1 U	
				39180	TRICHLOROETHYLENE	UG/L		1 U	
				34030	BENZENE	UG/L		1 U	
				32103	1,2-DICHLOROETHANE	UG/L		1 U	
				99999	1,3 DICHLOROPROPANE	UG/L		1 U	
				99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		0.7 J	QM
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	
				34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SECBUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		1 U	
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
NONE	96/04/03	0800	200134	99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		8 J	QF
				99999	2-BUTANONE	UG/L		1 J	QM
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	
DEPTH: 0000	SUBSTRATE: AQUEOUS	DESCRIPTION: TRIP BLANK #3	200135	99999	CHLOROMETHANE	UG/L		1 U	
				99999	BROMOMETHANE	UG/L		1 U	
				39175	VINYL CHLORIDE	UG/L		1 U	
				34311	CHLOROETHANE	UG/L		1 U	
				34423	METHYLENE CHLORIDE	UG/L		1 U	
				34488	TRICHLOROFUOROMETHANE	UG/L		1 U	
				34501	1,1-DICHLOROETHYLENE	UG/L		1 U	
				34496	1,1-DICHLOROETHANE	UG/L		1 U	
				99964	CARBON DISULFIDE	UG/L		1 U	
				34546	TRANS 1,2 DICHLOROETHYLENE	UG/L		1 U	
				99999	CIS 1,2- DICHLOROETHYLENE	UG/L		1 U	
				99999	2,2 DICHLOROPROPANE	UG/L		1 U	
				32106	CHLOROFORM	UG/L		1.7	
				99999	DIBROMOMETHANE	UG/L		1 U	
				34506	1,1,1-TRICHLOROETHANE	UG/L		1 U	
				32102	CARBON TETRACHLORIDE	UG/L		1 U	
				32101	DICHLOROBROMOMETHANE	UG/L		1 U	
				99999	1,1-DICHLOROPROPENE	UG/L		1 U	
				34541	1,2-DICHLOROPROPANE	UG/L		1 U	
				99999	CIS-1,3-DICHLOROPROPENE	UG/L		1 U	
				39180	TRICHLOROETHYLENE	UG/L		1 U	
				34030	BENZENE	UG/L		1 U	
				32103	1,2-DICHLOROETHANE	UG/L		1 U	
				99999	1,3 DICHLOROPROPANE	UG/L		1 U	
				99999	1,1,1,2 TETRACHLOROETHANE	UG/L		1 U	
				34010	TOLUENE	UG/L		0.3 J	QM
				34475	TETRACHLOROETHYLENE	UG/L		1 U	
				99999	1,2,3 TRICHLOROPROPANE	UG/L		1 U	
				34516	1,1,2,2-TETRACHLOROETHANE	UG/L		1 U	
				99999	TRANS-1,3-DICHLOROPROPENE	UG/L		1 U	

## COMPLETED ANALYSIS REPORT

REPORT DATE: 96/05/01

PROJECT NO: 235

PROJECT NAME: ANCHOR CHEMICAL

STATION NO	DATE FROM TO	TIME OF DAY	LABNO	PARNO	PARAMETER NAME	UNITS	CHEMISTRY	VALUE & REMARK	QA/QC REMARK
			200135	34511	1,1,2-TRICHLOROETHANE	UG/L		1 U	
				32105	CHLORODIBROMOMETHANE	UG/L		1 U	
				34301	CHLOROBENZENE	UG/L		1 U	
				34371	ETHYLBENZENE	UG/L		1 U	
				32104	BROMOFORM	UG/L		1 U	
				99999	BROMOBENZENE	UG/L		1 U	
				99999	ISOPROPYLBENZENE	UG/L		1 U	
				99921	STYRENE	UG/L		1 U	
				99902	O-XYLENE	UG/L		1 U	
				99999	P+M XYLENE	UG/L		1 U	
				99905	N-PROPYLBENZENE	UG/L		1 U	
				99999	1,2-DIBROMOETHANE	UG/L		1 U	
				99999	2-HEXANONE	UG/L		1 U	
				99999	2-CHLOROTOLUENE	UG/L		1 U	
				99999	4-CHLOROTOLUENE	UG/L		1 U	
				99999	TERTBUTYLBENZENE	UG/L		1 U	
				34566	1,3-DICHLOROBENZENE	UG/L		1 U	
				99999	SEC-BUTYLBENZENE	UG/L		1 U	
				34536	1,2-DICHLOROBENZENE	UG/L		1 U	
				34571	1,4-DICHLOROBENZENE	UG/L		1 U	
				99909	N-BUTYLBENZENE	UG/L		1 U	
				99999	1,2,4-TRIMETHYLBENZENE	UG/L		1 U	
				99907	1,3,5-TRIMETHYLBENZENE	UG/L		1 U	
				99999	4-ISOPROPYLTOLUENE	UG/L		1 U	
				34551	1,2,4-TRICHLOROBENZENE	UG/L		1 U	
				34696	NAPHTHALENE	UG/L		0.3 J	QM
				39702	HEXACHLOROBUTADIENE	UG/L		1 U	
				99999	1,2,3-TRICHLOROBENZENE	UG/L		1 U	
				99999	1,2-DIBROMO-3-CHLOROPROPANE	UG/L		1 U	
				99930	ACETONE	UG/L		6 U	
				99999	2-BUTANONE	UG/L		0.6 J	QM
				99999	BROMOCHLOROMETHANE	UG/L		1 U	
				99999	4-METHYL-2-PENTANONE	UG/L		1 U	

\*\*\*\*\* END OF PROJECT \*\*\*\*\*

Appendix D, Field Data Sheet for ESD laboratory

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name Franklin Township  
Collector(s) [Signature] Affiliation USEPA

SAMPLING METHOD (Circle)  
 Kemmerer Dredge Ponar Manual  
 Niskin Net Seine Trawl Bucket  
 Trowel Cream Dipper  
 Automatic  
 Other [Signature]

LDMS CODE \_\_\_\_\_  
 DATA BASE CODE \_\_\_\_\_  
 STA. TYPE CODE \_\_\_\_\_

Samples to:  
 Bact Bio Chem  Other

Station No.  

--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--

Sample Depth (Ft.)/Fac. Loc. Code  

--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--

SUBSTRATE TYPE (Circle)  Aqueous Sediment Sludge Oil Biological  
 Solvent Extract Other ( )

Lab Number  

200126

BOD — Seed Supplied  Yes  No

Source: \_\_\_\_\_

Sample Preparation (Circle)		Sample Source Type (Circle)	
Container	Cleaning Procedure	Landfill	Industrial
Glass Jar	Detergent Wash	Leachate	Effluent
Plastic Jar	Water Rinse	Drum	Process Stream
Metal	Acid Rinse	Test Well	Holding Pond
<u>POA Vial</u>	Solvent Rinse:	Depth:	Drum
Cubitainer	Acetone	Other: _____	Waste Pile
Acetate Core	Hexane		Municipal Treatment
Paper Cap	Methylene Chloride	Storage Tank	Influent
<u>Teflon Cap</u>	Other (Specify):	Top	Effluent-CI
Foil Cap	<u>Et Glassware</u>	Middle	Effluent-Non CI
Other _____		Bottom	Sludge
		Truck	Ambient
		Drum	Lake
Preservation		Tank	Stream
Acid <u>HCl</u>		Other _____	Pond
Solvent _____			Ocean
Chemical _____		Wells	Estuary
<u>Wet Ice</u>		Monitoring	
Dry Ice		Production	
Ambient		Drinking	
Other _____		Private	

Type of Sample  
 Grab  Composite  
 Time Space

Collection (Ending) Date  

Yr	Mo	Day
96	03	27

Ending Time (24 Hr)  

08	03
----	----

Beginning Date  

Yr	Mo	Day

Beginning Time (24 Hr)  

--	--	--	--

pH  

<	2	.	
---	---	---	--

Sample Temp. (°C)  

--	--	--	--

DO (mg/l)  

--	--	--	--

Cond. (µMHOS/CM)  

--	--	--	--	--	--

Salinity(‰)  

--	--	--	--	--	--

Sample Split  
 Yes  No

If Yes With Whom? \_\_\_\_\_

Receipt  Yes  No

Sample Location Description:

Trip Blank

Remarks:

VOCs - Drinking Water Standards - MCL - 3, 40 µg/l  
 Vials pres HCl pH < 2. Cool to 4°C

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name Procter Chemical  
Collector(s) Elizabeth Williams Affiliation USEPA

SAMPLING METHOD (Circle)  
Kemmerer Dredge Ponar Manual  
Niskin Net Seine Trawl Bucket  
Trowel Cream Dipper  
Automatic  
Other Amber

LDMS CODE \_\_\_\_\_  
DATA BASE CODE \_\_\_\_\_  
STA. TYPE CODE \_\_\_\_\_

SUBSTRATE TYPE (Circle) Aqueous Sediment Sludge Oil Biological  
Solvent Extract Other ( )

BOD — Seed Supplied  Yes  No Source:

Sample Preparation (Circle)		Sample Source Type (Circle)	
Container	Cleaning Procedure	Landfill	Industrial
<u>Glass Jar</u>	Detergent Wash	Leachate	Effluent
Plastic Jar	Water Rinse	Drum	Process Stream
<u>Metal</u>	Acid Rinse	Test Well	Holding Pond
<u>POA Vial</u>	Solvent Rinse:	Depth:	Drum
Cubitainer	Acetone	Other:	Waste Pile
Acetate Core	Hexane		Municipal Treatment
Paper Cap	Methylene Chloride	Storage Tank	Influent
<u>Tetlon Cap</u>	Other (Specify):	Top	Effluent-CI
Foil Cap	<u>EP Glassware</u>	Middle	Effluent-Non CI
Other _____		Bottom	Sludge
		Truck	Ambient
		Drum	Lake
		Tank	Stream
		Other _____	Pond
			Ocean
		Wells	Estuary
		Monitoring	
		Production	
		Drinking	
		Private	

Sample Location Description:

Equipment Blanks

Remarks:

VOL. DRINKING WATER (Std-NCl): 3, 40ml Vials pres  
HCl pH < 2 Cool to 4°C  
NOOCs: 1, 1 Liter Amber Glass Bottle Cool to 4°C  
TAL METAL: 1, 1 Liter Glass Bottle Cool to 4°C pres HNO<sub>3</sub>  
pH < 2

Samples to:

Bact Bio Chem Other

Station No.

\_\_\_\_\_

Sample Depth (Ft.)/Fac. Loc. Code

0000

Lab Number

200127

Type of Sample

Grab Composite  
Time Space

Collection (Ending) Date

7/6 Yr 03 Mo 27 Day

Ending Time (24 Hr)

0825

Beginning Date

Yr Mo Day

Beginning Time (24 Hr)

\_\_\_\_\_

pH

\_\_\_\_\_

Sample Temp. (°C)

\_\_\_\_\_

DO (mg/l)

\_\_\_\_\_

Cond. (uMHOS/CM)

\_\_\_\_\_

Salinity(‰)

\_\_\_\_\_

Sample Split

Yes  No

If Yes With Whom?

Receipt  Yes  No

## FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name Anchor Chemical  
 Collector(s) M. MERCADO/R. NORRILL/S. FERRERA Affiliation USEPA

SAMPLING METHOD (Circle)  
 Kemmerer Dredge Ponar Manual  
 Niskin Net Seine Trawl Bucket  
 Trowel Cream Dipper  
 Automatic  
 Other Bailer

LDMS CODE \_\_\_\_\_  
 DATA BASE CODE \_\_\_\_\_  
 STA. TYPE CODE \_\_\_\_\_

SUBSTRATE TYPE (Circle) Aqueous Sediment Sludge Oil Biological  
 Solvent Extract Other ( )

BOD - Seed Supplied  Yes  No

Source:

Sample Preparation (Circle)

Sample Source Type (Circle)

Container  
Glass Jar  
 Plastic Jar  
 Metal  
~~POA Vial~~  
 Cubitainer  
 Acetate Core  
 Paper Cap  
Teflon Cap  
 Foil Cap  
 Other \_\_\_\_\_

Cleaning Procedure  
 Detergent Wash  
 Water Rinse  
 Acid Rinse  
 Solvent Rinse:  
 Acetone  
 Hexane  
 Methylene Chloride  
 Other (Specify):  
E/P Glassware

Landfill  
 Leachate  
 Drum  
 Test Well  
 Depth:  
 Other: \_\_\_\_\_

Industrial  
 Effluent  
 Process Stream  
 Holding Pond  
 Drum  
 Waste Pile  
 Municipal Treatment

Storage Tank  
 Top  
 Middle  
 Bottom  
 Truck  
 Drum  
 Tank  
 Other: \_\_\_\_\_

Influent  
 Effluent-CI  
 Effluent-Non CI  
 Sludge  
 Ambient  
 Lake  
 Stream  
 Pond  
 Ocean  
 Estuary

Wells  
Monitoring  
 Production  
 Drinking  
 Private

Preservation  
 Acid HCl, HNO<sub>3</sub>  
 Solvent  
 Chemical  
Wet Ice  
 Dry Ice  
 Ambient  
 Other \_\_\_\_\_

Sample Location Description:

MW-4

Remarks:

VOC's (DRINKING WATER 5+d-MCL) : 6, 40ml Vials pres HCl  
pH < 2 Cool to 4°C  
NVOC's : 3, 1 Liter Amber Glass Bottle, Cool to 4°C  
TAL METALS : 3, 1 Liter Glass Bottle, Cool to 4°C pres. HNO<sub>3</sub>  
pH < 2

Samples to:

Bact  Bio  Chem  Other

Station No.

--	--	--	--	--	--	--	--	--	--	--	--

Sample Depth (Ft.)/Fac. Loc. Code

--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--

Lab Number

200128

Type of Sample

Grab  Composite

--	--	--	--	--	--	--	--

Collection (Ending) Date

Yr	Mo	Day
95	04	02

Ending Time (24 Hr)

--	--	--	--

Beginning Date

Yr	Mo	Day

Beginning Time (24 Hr)

--	--	--	--

pH

--	--	--	--

Sample Temp. (°C)

--	--	--	--

DO (mg/l)

--	--	--	--

Cond. (uMHOS/CM)

--	--	--	--	--	--	--	--

Salinity(‰)

--	--	--	--	--	--	--	--

Sample Split

Yes  No

If Yes With Whom?

Receipt  Yes  No

FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name Anchor Chemical  
Collector(s) A. H. Garcia / R. Morrell / J. P. ... Affiliation USEPA

SAMPLING METHOD (Circle)  
Kemmerer Dredge Ponar Manual  
Niskin Net Seine Trawl Bucket  
Trowel Cream Dipper  
Automatic  
Other Barker

LDMS CODE \_\_\_\_\_  
DATA BASE CODE \_\_\_\_\_  
STA. TYPE CODE \_\_\_\_\_

Samples to:  
Bact  Bio  Chem  Other

Station No. \_\_\_\_\_

Sample Depth (Ft.)/Fac. Loc. Code  
670

SUBSTRATE TYPE (Circle) Aqueous Sediment Sludge Oil Biological  
Solvent Extract Other ( )

Lab Number  
200129

BOD - Seed Supplied  Yes  No Source:

Sample Preparation (Circle)  
Cleaning Procedure  
Glass Jar  
Plastic Jar  
Metal  
POA Vial  
Cubitainer  
Acetate Core  
Paper Cap  
Teflon Cap  
Foil Cap  
Other \_\_\_\_\_  
Preservation  
Acid HCl/HNO3  
Solvent \_\_\_\_\_  
Chemical \_\_\_\_\_  
Wet Ice  
Dry Ice  
Ambient  
Other \_\_\_\_\_  
E/P  
Glassware

Sample Source Type (Circle)  
Landfill  
Leachate  
Drum  
Test Well  
Depth:  
Other: \_\_\_\_\_  
Storage Tank  
Top  
Middle  
Bottom  
Truck  
Drum  
Tank  
Other \_\_\_\_\_  
Wells  
Monitoring  
Production  
Drinking  
Private  
Industrial  
Process Stream  
Holding Pond  
Drum  
Waste Pile  
Municipal Treatment  
Influent  
Effluent-CI  
Effluent-Non CI  
Sludge  
Ambient  
Lake  
Stream  
Pond  
Ocean  
Estuary

Type of Sample  
Grab  Composite  
Time  Space

Collection (Ending) Date  
Yr Mo Day  
9 6 04  
01 2

Ending Time (24 Hr)  
1330

Beginning Date  
Yr Mo Day

Beginning Time (24 Hr)

pH  
5.89

Sample Temp. (°C)  
14.6

DO (mg/l)

Cond. (uMHOS/CM)  
160

Salinity(‰)

Sample Split  
 Yes  No

If Yes With Whom?  
Receipt  Yes  No

Sample Location Description:  
MW-4D

Remarks:  
VOCs (Drinking Water Std - MCL): 3, 40ml Vials pres HCl  
pH < 2 Cool to 4°C  
TCV NUOCs: 1, 1 Liter Amber Glass bottle Cool to 4°C  
TAL Metals: 1, 1 Liter Glass bottle Cool to 4°C pres HNO3  
pH < 2



# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name <u>Anchor Chemical</u>		Samples to:																																																																																					
Collector(s) <u>M. Hernandez / J. Fernandez</u> Affiliation <u>USEPA</u>		<input type="checkbox"/> Bact <input type="checkbox"/> Bio <input checked="" type="checkbox"/> Chem <input type="checkbox"/> Other																																																																																					
SAMPLING METHOD (Circle) Kemmerer   Dredge   Ponar <u>Manual</u> Niskin   Net   Seine   Trawl   Bucket Trowel   Cream   Dipper Automatic Other <u>Bailer</u>		LDMS CODE _____ DATA BASE CODE _____ STA. TYPE CODE _____																																																																																					
SUBSTRATE TYPE (Circle) <u>Aqueous</u> Sediment    Sludge    Oil    Biological Solvent    Extract    Other (     )		Station No. <table border="1" style="width: 100%; height: 20px;"><tr><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td></tr></table> Sample Depth (Ft.)/Fac. Loc. Code <table border="1" style="width: 100%; height: 20px;"><tr><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td><td> </td></tr></table>																																																																																					
BOD — Seed Supplied <input type="checkbox"/> Yes <input type="checkbox"/> No    Source: _____ Sample Preparation (Circle)     Sample Source Type (Circle)		Lab Number <div style="border: 1px solid black; padding: 5px; display: inline-block; font-size: 1.2em;">200130</div>																																																																																					
<table border="1" style="width: 100%; border-collapse: collapse;"> <tr> <th style="width: 20%;">Container</th> <th style="width: 20%;">Cleaning Procedure</th> <th style="width: 20%;">Landfill</th> <th style="width: 20%;">Industrial</th> </tr> <tr> <td><u>Glass Jar</u></td> <td>Detergent Wash</td> <td>Leachate</td> <td>Effluent</td> </tr> <tr> <td>Plastic Jar</td> <td>Water Rinse</td> <td>Drum</td> <td>Process Stream</td> </tr> <tr> <td>Metal</td> <td>Acid Rinse</td> <td>Test Well</td> <td>Holding Pond</td> </tr> <tr> <td><u>POA Vial</u></td> <td>Solvent Rinse:</td> <td>Depth:</td> <td>Drum</td> </tr> <tr> <td>Cubitainer</td> <td>Acetone</td> <td>Other: _____</td> <td>Waste Pile</td> </tr> <tr> <td>Acetate Core</td> <td>Hexane</td> <td></td> <td>Municipal Treatment</td> </tr> <tr> <td>Paper Cap</td> <td>Methylene Chloride</td> <td>Storage Tank</td> <td>Influent</td> </tr> <tr> <td><u>Teflon Cap</u></td> <td>Other (Specify):</td> <td>Top</td> <td>Effluent-CI</td> </tr> <tr> <td>Foil Cap</td> <td><u>EP</u></td> <td>Middle</td> <td>Effluent-Non CI</td> </tr> <tr> <td>Other _____</td> <td><u>Glassware</u></td> <td>Bottom</td> <td>Sludge</td> </tr> <tr> <td></td> <td></td> <td>Truck</td> <td>Ambient</td> </tr> <tr> <td></td> <td></td> <td>Drum</td> <td>Lake</td> </tr> <tr> <td></td> <td></td> <td>Tank</td> <td>Stream</td> </tr> <tr> <td></td> <td></td> <td>Other _____</td> <td>Pond</td> </tr> <tr> <td></td> <td></td> <td></td> <td>Ocean</td> </tr> <tr> <td></td> <td></td> <td>Wells</td> <td>Estuary</td> </tr> <tr> <td></td> <td></td> <td><u>Monitoring</u></td> <td></td> </tr> <tr> <td></td> <td></td> <td>Production</td> <td></td> </tr> <tr> <td></td> <td></td> <td>Drinking</td> <td></td> </tr> <tr> <td></td> <td></td> <td>Private</td> <td></td> </tr> </table>		Container	Cleaning Procedure	Landfill	Industrial	<u>Glass Jar</u>	Detergent Wash	Leachate	Effluent	Plastic Jar	Water Rinse	Drum	Process Stream	Metal	Acid Rinse	Test Well	Holding Pond	<u>POA Vial</u>	Solvent Rinse:	Depth:	Drum	Cubitainer	Acetone	Other: _____	Waste Pile	Acetate Core	Hexane		Municipal Treatment	Paper Cap	Methylene Chloride	Storage Tank	Influent	<u>Teflon Cap</u>	Other (Specify):	Top	Effluent-CI	Foil Cap	<u>EP</u>	Middle	Effluent-Non CI	Other _____	<u>Glassware</u>	Bottom	Sludge			Truck	Ambient			Drum	Lake			Tank	Stream			Other _____	Pond				Ocean			Wells	Estuary			<u>Monitoring</u>				Production				Drinking				Private		Type of Sample <input checked="" type="checkbox"/> Grab <input type="checkbox"/> Composite Time    Space	
Container	Cleaning Procedure	Landfill	Industrial																																																																																				
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Yr	Mo	Day																																																																																					
Sample Location Description:  <div style="font-size: 1.5em; text-align: center;">MW-5D</div>		pH <div style="border: 1px solid black; padding: 5px; display: inline-block; font-size: 1.2em;">6.40</div>																																																																																					
Remarks: <u>VOCs (Drinking Water Std - NCL): 3, 40ml Vials pres HCl</u> <u>pH 2 Cool to 4°C</u> <u>TU, NO<sub>3</sub> 1, 1 Liter Amber Glass Bottle, Cool to 4°C</u> <u>TAL METALS: 1, 1 Liter Glass Bottle, Cool to 4°C, pres HNO<sub>3</sub></u> <u>pH 2</u>		Sample Temp. (°C) <table border="1" style="width: 100%; text-align: center;"> <tr> <td> </td> <td> </td> <td> </td> <td> </td> </tr> <tr> <td>1</td> <td>7</td> <td>4</td> <td> </td> </tr> </table>						1	7	4																																																																													
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Form: FTB RPD-11-82-2		DO (mg/l) <table border="1" style="width: 100%; text-align: center;"> <tr> <td> </td> <td> </td> <td> </td> <td> </td> </tr> </table>																																																																																					
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Sample Split <input type="checkbox"/> Yes <input checked="" type="checkbox"/> No		If Yes With Whom? <input type="checkbox"/> Yes <input type="checkbox"/> No																																																																																					

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name Anchor Chemical

Collector(s) M. MERCADO, B. MORENO, J. FERRERA Affiliation USEPA

SAMPLING METHOD (Circle)

Kemmerer Dredge Ponar  Manual  
Niskin Net Seine Trawl Bucket  
Trowel Cream Dipper  
Automatic  
Other Boiler

LDMS CODE \_\_\_\_\_

DATA BASE CODE \_\_\_\_\_

STA. TYPE CODE \_\_\_\_\_

Samples to:

Bact	Bio	<input checked="" type="checkbox"/> Chem	Other
------	-----	--	-------

Station No.

--	--	--	--	--	--	--	--	--	--	--	--

Sample Depth (Ft.)/Fac. Loc. Code

6	3	4	

Lab Number

2	0	0	1	3	1
---	---	---	---	---	---

SUBSTRATE TYPE (Circle)

Aqueous    Sediment    Sludge    Oil    Biological  
Solvent    Extract    Other (                )

BOD — Seed Supplied     Yes     No    Source:

Sample Preparation (Circle)

Sample Source Type (Circle)

Container  
 Glass Jar  
Plastic Jar  
Metal  
 POA Vial  
Cubittainer  
Acetate Core  
Paper Cap  
 Teflon Cap  
Foil Cap  
Other \_\_\_\_\_

Cleaning Procedure  
Detergent Wash  
Water Rinse  
Acid Rinse  
Solvent Rinse:

Acetone  
Hexane  
Methylene Chloride  
Other (Specify):

E/P Glassware

Preservation  
Acid HCl/HNO<sub>3</sub>  
Solvent  
Chemical  
 Wet Ice  
Dry Ice  
Ambient  
Other \_\_\_\_\_

Landfill  
Leachate  
Drum  
Test Well  
Depth: \_\_\_\_\_  
Other: \_\_\_\_\_

Storage Tank  
Top  
Middle  
Bottom

Truck  
Drum  
Tank  
Other \_\_\_\_\_  
Wells  
 Monitoring  
Production  
Drinking  
Private

Industrial  
Effluent  
Process Stream  
Holding Pond  
Drum  
Waste Pile  
Municipal Treatment  
Influent  
Effluent-CI  
Effluent-Non CI  
Sludge  
Ambient  
Lake  
Stream  
Pond  
Ocean  
Estuary

Type of Sample

Grab    Composite

<input checked="" type="checkbox"/> Grab	Time	Space
--	------	-------

Collection (Ending) Date

Yr	Mo	Day
9	6	04
		012

Ending Time (24 Hr)

1	3	0	0
---	---	---	---

Beginning Date

Yr	Mo	Day

Beginning Time (24 Hr)

--	--	--	--

pH

6	0	6
---	---	---

Sample Temp. (°C)

1	4	1
---	---	---

DO (mg/l)

--	--	--	--

Cond. (µMHOS/CM)

		3	1	8

Salinity(‰)

--	--	--	--

Sample Split

Yes     No

If Yes With Whom?

Receipt     Yes     No

Sample Location Description:

MW-55

Remarks:

VOCs (Drinking Water Std - HCL): 3, ~~4~~ <sup>40 ml</sup> Vials pres HCl  
pH < 2 Cool to 4°C  
TCL NVOCS: 1, 1 Liter Amber Glass Bottle, Cool to 4°C  
TAL METALS: 1, 1 Liter Glass Bottle, Cool to 4°C pres HNO<sub>3</sub>  
pH < 2

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name: Anchor Chemical  
Collector(s): W. P. ... Affiliation: ES&E

Samples to:

Bact	Bio	Chem <input checked="" type="checkbox"/>	Other
------	-----	--	-------

SAMPLING METHOD (Circle)

Kemmerer  Dredge  Ponar  Manual  
Niskin  Net  Seine  Trawl  Bucket  
Trowel  Cream  Dipper  
Automatic   
Other: Anchor

LDMs CODE \_\_\_\_\_  
DATA BASE CODE \_\_\_\_\_  
STA. TYPE CODE \_\_\_\_\_

Station No.

--	--	--	--	--	--	--	--	--	--	--

Sample Depth (Ft.)/Fac. Loc. Code

70	-
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SUBSTRATE TYPE (Circle) Aqueous Sediment Sludge Oil Biological  
Solvent Extract Other ( )

Lab Number

200132
--------

BOD — Seed Supplied  Yes  No Source: \_\_\_\_\_

Sample Preparation (Circle)		Sample Source Type (Circle)	
<u>Glass Jar</u>	Detergent Wash	Landfill	Industrial
Plastic Jar	Water Rinse	Leachate	Effluent
Metal	Acid Rinse	Drum	Process Stream
<u>POA Vial</u>	Solvent Rinse: Acetone Hexane Methylene Chloride Other (Specify): <u>E/C Glassware</u>	Test Well	Holding Pond
Cubitainer		Depth:	Drum
Acetate Core		Other:	Waste Pile
<u>Paper Cap</u>		Storage Tank	Municipal Treatment
<u>Teflon Cap</u>		Top	Influent
Foil Cap	Middle	Effluent-Cl	
Other _____	Bottom	Effluent-Non Cl	
Preservation	Truck	Sludge	
Acid <u>HCl, HNO3</u>	Drum	Ambient	
Solvent _____	Tank	Lake	
Chemical _____	Other _____	Stream	
Wet Ice	<u>Wells</u>	Pond	
<u>Dry Ice</u>	<u>Monitoring</u>	Ocean	
Ambient	Production	Estuary	
Other _____	Drinking		
	Private		

Type of Sample

Grab <input checked="" type="checkbox"/>	Composite
Time	Space

Collection (Ending) Date

Yr: <u>96</u>	Mo: <u>03</u>	Day: <u>27</u>
---------------	---------------	----------------

Ending Time (24 Hr)

14	13
----	----

Beginning Date

Yr	Mo	Day

Beginning Time (24 Hr)

--	--

pH

6	8	0
---	---	---

Sample Temp. (°C)

1	2	5
---	---	---

DO (mg/l)

--	--	--

Cond. (uMHOS/CM)

		3	6	0
--	--	---	---	---

Salinity(‰)

--	--	--	--

Sample Split

Yes  No

If Yes With Whom?

Receipt  Yes  No

Sample Location Description:

MW-6D

Remarks:

VOCs (Drinking Water Std - MCL): 3, 40 ml Vials pres  
HCl pH=2 Cool to 4°C  
NVOCs: 1, 1 Liter Amber Glass Bottle Cool to 4°C  
TLV Metals: 1, 1 Liter Glass Bottle Cool to 4°C. pres HNO<sub>3</sub>  
pH=2

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey

## ENVIRONMENTAL SERVICES DIVISION

Project Name Asbestos Chemical  
 Collector(s) Michael J. Murphy Affiliation USEPA

**SAMPLING METHOD (Circle)**  
 Kemmerer Dredge Ponar  Manual  
 Niskin Net Seine Trawl Bucket  
 Trowel Cream Dipper  
 Automatic  
 Other Buiter

LDMS CODE \_\_\_\_\_  
 DATA BASE CODE \_\_\_\_\_  
 STA. TYPE CODE \_\_\_\_\_

**SUBSTRATE TYPE (Circle)**  Aqueous Sediment Sludge Oil Biological  
 Solvent Extract Other ( )

BOD - Seed Supplied  Yes  No Source: \_\_\_\_\_

**Sample Preparation (Circle)**

Container	Cleaning Procedure
<input checked="" type="radio"/> <u>Glass Jar</u>	Detergent Wash
Plastic Jar	Water Rinse
Metal	Acid Rinse
<input checked="" type="radio"/> <u>POA Vial</u>	Solvent Rinse:
Cubitainer	Acetone
Acetate Core	Hexane
Paper Cap	Methylene Chloride
<input checked="" type="radio"/> <u>Teflon Cap</u>	Other (Specify):
Foil Cap	<u>EP</u> <u>Glassware</u>
Other _____	

**Preservation**  
 Acid HCl, HNO<sub>3</sub>  
 Solvent \_\_\_\_\_  
 Chemical \_\_\_\_\_  
 Wet Ice  
 Dry Ice  
 Ambient  
 Other \_\_\_\_\_

**Sample Source Type (Circle)**

Landfill	Industrial
Leachate	Effluent
Drum	Process Stream
Test Well	Holding Pond
Depth:	Drum
Other:	Waste Pile
	Municipal Treatment
Storage Tank	Influent
Top	Effluent-CI
Middle	Effluent-Non CI
Bottom	Sludge
Truck	Ambient
Drum	Lake
Tank	Stream
Other _____	Pond
	Ocean
Wells	Estuary
<input checked="" type="radio"/> <u>Monitoring</u>	
Production	
Drinking	
Private	

Sample Location Description:

MW-65

**Remarks:**

VOCs (Drinking Water Standard - MCL): 3, 40ml Vials Pres  
HCl, H<sub>2</sub>SO<sub>4</sub> Cool to 4°C  
MVOCs: 1, 1 liter Amber Glass Bottle Cool to 4°C  
TRE METALS: 1, 1 liter Glass Bottle Cool to 4°C, Pres  
HNO<sub>3</sub> pH < 2

**Samples to:**

Bact	Bio	Chem <input checked="" type="checkbox"/>	Other
------	-----	--	-------

**Station No.**

--	--	--	--	--	--	--	--	--	--

**Sample Depth (Ft.)/Fac. Loc. Code**


**Lab Number**

200133
--------

**Type of Sample**

Grab <input checked="" type="checkbox"/>	Composite
Time	Space

**Collection (Ending) Date**

Yr	Mo	Day
96	03	29

**Ending Time (24 Hr)**

1404
------

**Beginning Date**

Yr	Mo	Day

**Beginning Time (24 Hr)**

--	--	--

**pH**

680
-----

**Sample Temp. (°C)**

128
-----

**DO (mg/l)**

--	--	--

**Cond. (µMHOS/CM)**


**Salinity(‰)**

--	--	--

**Sample Split**

Yes  No

**If Yes With Whom?**

Receipt  Yes  No

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey  
ENVIRONMENTAL SERVICES DIVISION

Project Name Anchor Chemical  
 Collector(s) M. MERCADO & H. HERRERA Affiliation USEPA

SAMPLING METHOD (Circle)  
 Kemmerer Dredge Ponar Manual  
 Niskin Net Seine Trawl Bucket  
 Trowel Cream Dipper  
 Automatic  
 Other Bowler

LDMS CODE \_\_\_\_\_  
 DATA BASE CODE \_\_\_\_\_  
 STA. TYPE CODE \_\_\_\_\_

SUBSTRATE TYPE (Circle) Aqueous Sediment Sludge Oil Biological  
 Solvent Extract Other ( )

BOD - Seed Supplied <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No		Source:	
Sample Preparation (Circle)		Sample Source Type (Circle)	
Container	Cleaning Procedure	Landfill	Industrial
Glass Jar	Detergent Wash	Leachate	Effluent
Plastic Jar	Water Rinse	Drum	Process Stream
Metal	Acid Rinse	Test Well	Holding Pond
<u>POA Vial</u>	Solvent Rinse:	Depth:	Drum
Cubitainer	Acetone	Other: _____	Waste Pile
Acetate Core	Hexane		Municipal Treatment
Paper Cap	Methylene Chloride	Storage Tank	Influent
<u>Teflon Cap</u>	Other (Specify):	Top	Effluent-CI
Foil Cap	<u>E/P</u>	Middle	Effluent-Non CI
Other _____	<u>Glassware</u>	Bottom	Sludge
		Truck	Ambient
Preservation		Drum	Lake
Acid <u>HCl</u>		Tank	Stream
Solvent		Other _____	Pond
Chemical			Ocean
<u>Wet Ice</u>		Wells	Estuary
Dry Ice		Monitoring	
Ambient		Production	
Other _____		Drinking	
		Private	

Sample Location Description:  
  
Trip Blank #2

Remarks:  
VOCs (Drinking Water Std - HCL). 3, 40 ml vials pres HCl  
pH < 2, Cool to 4°C

Samples to:  

Bact	Bio	Chem <input checked="" type="checkbox"/>	Other
------	-----	--	-------

Station No.  

--	--	--	--	--	--	--	--	--	--	--	--

Sample Depth (Ft.)/Fac. Loc. Code  

--	--	--	--	--	--	--	--

00

Lab Number  
200134

Type of Sample  

Grab		Composite	
Time	Space		

  
 Time  Space

Collection (Ending) Date  

Yr	Mo	Day
96	04	02

Ending Time (24 Hr)  

--	--	--	--

1015

Beginning Date  

Yr	Mo	Day

Beginning Time (24 Hr)  

--	--	--	--

pH  

--	--	--	--

Sample Temp. (°C)  

--	--	--	--

DO (mg/l)  

--	--	--	--

Cond. (uMHOS/CM)  

--	--	--	--	--	--

Salinity (‰)  

--	--	--	--

Sample Split  
 Yes  No

If Yes With Whom?  
 Receipt  Yes  No

# FIELD DATA SHEET

ENVIRONMENTAL PROTECTION AGENCY - Region II, Edison, New Jersey

ENVIRONMENTAL SERVICES DIVISION

Project Name Anchoe Chemical  
Collector(s) M. HERAZO / J. FERANDA / J. SNOW Affiliation US EPA

SAMPLING METHOD (Circle)

Kemmerer Dredge Ponar Manual  
Niskin Net Seine Trawl Bucket  
Trowel Cream Dipper  
Automatic Direct Pour  
Other Direct Pour

LDMS CODE \_\_\_\_\_

DATA BASE CODE \_\_\_\_\_

STA. TYPE CODE \_\_\_\_\_

Samples to:

Bact Bio  Chem Other

Station No.

\_\_\_\_\_

Sample Depth (Ft.)/Fac. Loc. Code

00

SUBSTRATE TYPE (Circle)

Aqueous Sediment Sludge Oil Biological  
Solvent Extract Other ( )

Lab Number

200135

BOD - Seed Supplied  Yes  No Source: \_\_\_\_\_

Sample Preparation (Circle)

Sample Source Type (Circle)

Container  
Glass Jar  
Plastic Jar  
Metal  
POA Vial  
Cubitainer  
Acetate Core  
Paper Cap  
Teflon Cap  
Foil Cap  
Other \_\_\_\_\_  
Preservation  
Acid \_\_\_\_\_  
Solvent \_\_\_\_\_  
Chemical \_\_\_\_\_  
Wet Ice  
Dry Ice  
Ambient  
Other \_\_\_\_\_

Cleaning Procedure  
Detergent Wash  
Water Rinse  
Acid Rinse  
Solvent Rinse:  
Acetone  
Hexane  
Methylene Chloride  
Other (Specify): EP Glassware

Landfill  
Leachate  
Drum  
Test Well  
Depth:  
Other: \_\_\_\_\_  
Storage Tank  
Top  
Middle  
Bottom  
Truck  
Drum  
Tank  
Other \_\_\_\_\_  
Wells  
Monitoring  
Production  
Drinking  
Private

Industrial  
Effluent  
Process Stream  
Holding Pond  
Drum  
Waste Pile  
Municipal Treatment  
Influent  
Effluent-Cl  
Effluent-Non Cl  
Sludge  
Ambient  
Lake  
Stream  
Pond  
Ocean  
Estuary

Type of Sample

Grab  Composite  
Time Space

Collection (Ending) Date

Yr Mo Day  
9 6 04 0 3

Ending Time (24 Hr)

0800

Beginning Date

Yr Mo Day

Beginning Time (24 Hr)

\_\_\_\_\_

pH

\_\_\_\_\_

Sample Temp. (°C)

\_\_\_\_\_

DO (mg/l)

\_\_\_\_\_

Cond. (uMHOS/CM)

\_\_\_\_\_

Salinity(‰)

\_\_\_\_\_

Sample Split

Yes  No

If Yes With Whom?

Receipt  Yes  No

Sample Location Description:

TRIP Blank #3

Remarks:

VOCs (Drinking Water Std-MCL): 3, 40 ml Vials pres  
HCL pH 2.2 Cool to 4°C

Appendix E, Analysis Request for ESD laboratory

# ANALYSIS REQUEST

CHEM <input checked="" type="checkbox"/>	BIO. <input type="checkbox"/>	BACT <input type="checkbox"/>	OTHER <input type="checkbox"/>
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ENVIRONMENTAL PROTECTION AGENCY  
Environmental Services Division

EDISON, N.J.

Date of Request 3/24/10 Priority  Immediate  Normal  Deferred  
Source of Sample(s) Industrial Chemical  
Sample Number(s) 200126, 200127, 200132 & 200133  
Type of Sample  Water  Sediment  Oil  Air  Other (Specify)

## PHYSICAL CHARACTERISTICS

- |  |  |  |   |
|--|--|--|---|
| <input type="checkbox"/> Turbidity                         | <input type="checkbox"/> Color             | <input type="checkbox"/> Specific Gravity    | <input type="checkbox"/> Corrosivity (RCRA) |
| <input type="checkbox"/> Volatile Solids                   | <input type="checkbox"/> Total Solids      | <input type="checkbox"/> Viscosity           | <input type="checkbox"/> Other _____        |
| <input checked="" type="checkbox"/> Total Suspended Solids | <input type="checkbox"/> Dissolved Solids  | <input type="checkbox"/> % Solids            | _____                                       |
| <input type="checkbox"/> Volatile Suspended Solids         | <input type="checkbox"/> Settleable Solids | <input type="checkbox"/> Ignitability (RCRA) | _____                                       |

## ORGANIC/DEMAND ANALYSES

- |  |   |   |  |
|--|---|---|--|
| <input type="checkbox"/> _____ Day BOD     | <input type="checkbox"/> Phenol   | <input checked="" type="checkbox"/> Priority Pollutants | <input type="checkbox"/> Specific Compound |
| <input type="checkbox"/> COD               | <input type="checkbox"/> Pesticides   | <input checked="" type="checkbox"/> POA <u>_____</u>    | <input type="checkbox"/> Identify _____    |
| <input type="checkbox"/> TOC               | <input type="checkbox"/> Herbicides   | <input checked="" type="checkbox"/> NVOA                | _____                                      |
| <input type="checkbox"/> TOD               | <input type="checkbox"/> Long-term O <sub>2</sub> Demand (Carbon)           | <input type="checkbox"/> Other Major Peaks              | _____                                      |
| <input type="checkbox"/> PCB's             | <input checked="" type="checkbox"/> Long-term O <sub>2</sub> Demand (Total) | <input type="checkbox"/> EP Toxicity                    | <input type="checkbox"/> Quantitate _____  |
| <input type="checkbox"/> Total             | <input type="checkbox"/> Volatile Acids                                     | <input type="checkbox"/> Pesticides                     | _____                                      |
| <input type="checkbox"/> Specific Aroclors | <input type="checkbox"/> Oil (Identify)                                     | <input type="checkbox"/> Herbicides                     | _____                                      |
|  |   | <input type="checkbox"/> Oil & Grease (Quantitate)      |  |

## INORGANIC ANALYSES

- |  |  |  |                              |  |
|--|--|--|------------------------------|--|
| <input type="checkbox"/> pH                  | <input type="checkbox"/> Alkalinity        | <input type="checkbox"/> TKN                                     | <input type="checkbox"/> Cd  | <input type="checkbox"/> Ba            |
| <input type="checkbox"/> Conductivity        | <input type="checkbox"/> CO <sub>3</sub>   | <input type="checkbox"/> Org N                                   | <input type="checkbox"/> Co  | <input checked="" type="checkbox"/> Se |
| <input checked="" type="checkbox"/> Salinity | <input type="checkbox"/> Total             | <input type="checkbox"/> NH <sub>3</sub> -N                      | <input type="checkbox"/> Cu  | <input checked="" type="checkbox"/> Ag |
| <input type="checkbox"/> Chloride            | <input type="checkbox"/> HCO <sub>3</sub>  | <input type="checkbox"/> NO <sub>2</sub> -N                      | <input type="checkbox"/> Pb  | <input type="checkbox"/> Asbestos      |
| <input type="checkbox"/> SO <sub>4</sub>     | <input type="checkbox"/> Chlorine Demand   | <input type="checkbox"/> NO <sub>3</sub> -N                      | <input type="checkbox"/> Zn  | <input type="checkbox"/> Hexavalent Cr |
| <input type="checkbox"/> SO <sub>3</sub>     | <input type="checkbox"/> Chlorine Residual | <input type="checkbox"/> Total P                                 | <input type="checkbox"/> Fe  |  |
| <input type="checkbox"/> Dissolved S         | <input type="checkbox"/> Free              | <input type="checkbox"/> AHP                                     | <input type="checkbox"/> Cr  |  |
| <input type="checkbox"/> Hardness            | <input type="checkbox"/> Total             | <input type="checkbox"/> Ortho-P                                 | <input type="checkbox"/> As  |  |
| <input type="checkbox"/> Ca                  | <input type="checkbox"/> Acidity           | <input checked="" type="checkbox"/> Metal Scan <u>TAL METALS</u> | <input type="checkbox"/> CN- |  |
| <input type="checkbox"/> Mg                  | <input type="checkbox"/> Free              | <input type="checkbox"/> EP Toxicity (Metals)                    | <input type="checkbox"/> F-  |  |
| <input type="checkbox"/> Total/METHOD        | <input type="checkbox"/> Total             | <input type="checkbox"/> Hg                                      | <input type="checkbox"/> NI  |  |

## SENSITIVITY / METHOD

- |   |                                      |  |   |
|---|--------------------------------------|--|---|
| <input type="checkbox"/> COD                    | <input type="checkbox"/> Phosphorous | <input type="checkbox"/> Phenol          | <input type="checkbox"/> Metals           |
| <input type="checkbox"/> High Level (> 50 mg/L) | <input type="checkbox"/> Total       | <input type="checkbox"/> 0-1,000 ppb     | <input type="checkbox"/> Total            |
| <input type="checkbox"/> Low Level (< 50 mg/L)  | <input type="checkbox"/> Dissolved   | <input type="checkbox"/> Above 1,000 ppb | <input type="checkbox"/> Dissolved        |
|   |                                      |  | <input type="checkbox"/> Low Sensitivity  |
|   |                                      |  | <input type="checkbox"/> High Sensitivity |

## MICROBIOLOGY

- |                             |                                     |            |  |
|-----------------------------|-------------------------------------|------------|--|
| MF                          | MPN                                 | Est. Range |  |
| TC <input type="checkbox"/> | <input type="checkbox"/>            | _____      | <input type="checkbox"/> Clostridium perfringens |
| FC <input type="checkbox"/> | <input checked="" type="checkbox"/> | _____      | <input type="checkbox"/> Mutagenicity Tests      |
| FS <input type="checkbox"/> | <input type="checkbox"/>            | _____      | <input type="checkbox"/> Ampe Test               |
|                             | <input type="checkbox"/> Pathogens  |            | <input type="checkbox"/> Viral Enhancement       |
|                             | <input type="checkbox"/> Bacterial  |            | <input type="checkbox"/> Other (Specify)         |
|                             | <input type="checkbox"/> Viral      |            | <input type="checkbox"/> ATP                     |

## BIOLOGY

- |   |   |
|---|---|
| <input type="checkbox"/> 24 Hour Bioassay | <input type="checkbox"/> Static             |
| <input type="checkbox"/> 48 Hour Bioassay | <input type="checkbox"/> Flow-Through       |
| <input type="checkbox"/> 96 Hour Bioassay | <input type="checkbox"/> Static Replacement |
| <input type="checkbox"/> Chronic Bioassay | <input type="checkbox"/> Laboratory         |
| <input type="checkbox"/> Benthos ID       | <input type="checkbox"/> On Site            |
| <input type="checkbox"/> Fish ID          | <input type="checkbox"/> Identify           |
|   | <input type="checkbox"/> Quantitate         |

Requested by [Signature] Date 3/24/10 Approved by \_\_\_\_\_ Date \_\_\_\_\_

Remarks

Form ETS 898-11-92



# ANALYSIS REQUEST

CHEM <input checked="" type="checkbox"/>	BIO. <input type="checkbox"/>	BACT <input type="checkbox"/>	OTHER <input type="checkbox"/>
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ENVIRONMENTAL PROTECTION AGENCY  
Environmental Services Division

EDISON, N.J.

Date of Request 4/2/96 Priority  Immediate  Normal  Deferred

Source of Sample(s) Anchor Chemical

Sample Number(s) 200125, 200129, 200131 & 200134

Type of Sample  Water  Sediment  Oil  Air  Other (Specify)

### PHYSICAL CHARACTERISTICS

- |  |  |  |   |
|--|--|--|---|
| <input type="checkbox"/> Turbidity                 | <input type="checkbox"/> Color                       | <input type="checkbox"/> Specific Gravity    | <input type="checkbox"/> Corrosivity (RCRA) |
| <input type="checkbox"/> Volatile Solids           | <input checked="" type="checkbox"/> Total Solids     | <input type="checkbox"/> Viscosity           | <input type="checkbox"/> Other _____        |
| <input type="checkbox"/> Total Suspended Solids    | <input checked="" type="checkbox"/> Dissolved Solids | <input type="checkbox"/> % Solids            | _____                                       |
| <input type="checkbox"/> Volatile Suspended Solids | <input type="checkbox"/> Settleable Solids           | <input type="checkbox"/> Ignitability (RCRA) | _____                                       |

### ORGANIC/DEMAND ANALYSES

- |  |   |   |  |
|--|---|---|--|
| <input type="checkbox"/> _____ Day BOD     | <input type="checkbox"/> Phenol                                   | <input checked="" type="checkbox"/> Priority Pollutants | <input type="checkbox"/> Specific Compound |
| <input type="checkbox"/> COD               | <input type="checkbox"/> Pesticides                               | <input checked="" type="checkbox"/> POA (HCL)           | <input type="checkbox"/> Identify _____    |
| <input type="checkbox"/> TOC               | <input type="checkbox"/> Herbicides                               | <input checked="" type="checkbox"/> NVOA (TCL)          | _____                                      |
| <input type="checkbox"/> TOD               | <input type="checkbox"/> Long-term O <sub>2</sub> Demand (Carbon) | <input type="checkbox"/> Other Major Peaks              | <input type="checkbox"/> Quantitate _____  |
| <input type="checkbox"/> PCB's             | <input type="checkbox"/> Long-term O <sub>2</sub> Demand (Total)  | <input type="checkbox"/> EP Toxicity                    | _____                                      |
| <input type="checkbox"/> Total             | <input type="checkbox"/> Volatile Acids                           | <input type="checkbox"/> Pesticides                     | _____                                      |
| <input type="checkbox"/> Specific Aroclors | <input type="checkbox"/> Oil (Identify)                           | <input type="checkbox"/> Herbicides                     | _____                                      |
|  |   | <input type="checkbox"/> Oil & Grease (Quantitate)      | _____                                      |

### INORGANIC ANALYSES

- |  |  |   |                              |  |
|--|--|---|------------------------------|--|
| <input type="checkbox"/> pH              | <input type="checkbox"/> Alkalinity        | <input type="checkbox"/> TKN                                | <input type="checkbox"/> Cd  | <input type="checkbox"/> Ba            |
| <input type="checkbox"/> Conductivity    | <input type="checkbox"/> CO <sub>3</sub>   | <input type="checkbox"/> Org N                              | <input type="checkbox"/> Co  | <input type="checkbox"/> Se            |
| <input type="checkbox"/> Salinity        | <input type="checkbox"/> Total             | <input type="checkbox"/> NH <sub>3</sub> -N                 | <input type="checkbox"/> Cu  | <input type="checkbox"/> Ag            |
| <input type="checkbox"/> Chloride        | <input type="checkbox"/> HCO <sub>3</sub>  | <input type="checkbox"/> NO <sub>2</sub> -N                 | <input type="checkbox"/> Pb  | <input type="checkbox"/> Asbestos      |
| <input type="checkbox"/> SO <sub>4</sub> | <input type="checkbox"/> Chlorine Demand   | <input type="checkbox"/> NO <sub>3</sub> -N                 | <input type="checkbox"/> Zn  | <input type="checkbox"/> Hexavalent Cr |
| <input type="checkbox"/> SO <sub>2</sub> | <input type="checkbox"/> Chlorine Residual | <input type="checkbox"/> Total P                            | <input type="checkbox"/> Fe  |  |
| <input type="checkbox"/> Dissolved S     | <input type="checkbox"/> Free              | <input type="checkbox"/> AH-P                               | <input type="checkbox"/> Cr  |  |
| <input type="checkbox"/> Hardness        | <input type="checkbox"/> Total             | <input type="checkbox"/> Ortho-P                            | <input type="checkbox"/> As  |  |
| <input type="checkbox"/> Ca              | <input type="checkbox"/> Acidity           | <input checked="" type="checkbox"/> Metal Scan (TAL METALS) | <input type="checkbox"/> CN- |  |
| <input type="checkbox"/> Mg              | <input type="checkbox"/> Free              | <input type="checkbox"/> EP Toxicity (Metals)               | <input type="checkbox"/> F-  |  |
| <input type="checkbox"/> Total/METHOD    | <input type="checkbox"/> Total             | <input type="checkbox"/> Hg                                 | <input type="checkbox"/> Ni  |  |

### SENSITIVITY / METHOD

- |   |   |   |   |
|---|---|---|---|
| <input type="checkbox"/> COD                              | <input type="checkbox"/> Phosphorous      | <input type="checkbox"/> Phenol                     | <input type="checkbox"/> Metals           |
| <input type="checkbox"/> High Level (> 50 mg/l)           | <input checked="" type="checkbox"/> Total | <input type="checkbox"/> 0-1,000 ppb                | <input type="checkbox"/> Total            |
| <input checked="" type="checkbox"/> Low Level (< 50 mg/l) | <input type="checkbox"/> Dissolved        | <input checked="" type="checkbox"/> Above 1,000 ppb | <input type="checkbox"/> Dissolved        |
|   |   |   | <input type="checkbox"/> Low Sensitivity  |
|   |   |   | <input type="checkbox"/> High Sensitivity |

### MICROBIOLOGY

- |    | MF                       | MPN                      | Est. Range |  |
|----|--------------------------|--------------------------|------------|--|
| TC | <input type="checkbox"/> | <input type="checkbox"/> | _____      | <input type="checkbox"/> Clostridium perfringens |
| FC | <input type="checkbox"/> | <input type="checkbox"/> | _____      | <input type="checkbox"/> Mutagenicity Tests      |
| FS | <input type="checkbox"/> | <input type="checkbox"/> | _____      | <input type="checkbox"/> Ames Test               |
|    |                          |                          |            | <input type="checkbox"/> Viral Enhancement       |
|    |                          |                          |            | <input type="checkbox"/> Other (Specify)         |
|    |                          |                          |            | <input type="checkbox"/> ATP                     |
|    |                          |                          |            | <input type="checkbox"/> Pathogens               |
|    |                          |                          |            | <input type="checkbox"/> Bacterial               |
|    |                          |                          |            | <input type="checkbox"/> Viral                   |

### BIOLOGY

- |  |  |
|--|--|
| <input checked="" type="checkbox"/> 24 Hour Bioassay | <input type="checkbox"/> Static              |
| <input type="checkbox"/> 48 Hour Bioassay            | <input type="checkbox"/> Flow-Through        |
| <input type="checkbox"/> 96 Hour Bioassay            | <input type="checkbox"/> Static Replacement  |
| <input type="checkbox"/> Chronic Bioassay            | <input type="checkbox"/> Laboratory          |
| <input type="checkbox"/> Benthos ID                  | <input checked="" type="checkbox"/> On Site  |
| <input type="checkbox"/> Fish ID                     | <input checked="" type="checkbox"/> Identify |
|  | <input type="checkbox"/> Quantitate          |

Requested by [Signature] Date 4/2/96 Approved by \_\_\_\_\_ Date \_\_\_\_\_

Remarks

# ANALYSIS REQUEST

CHEM <input checked="" type="checkbox"/>	BIO. <input type="checkbox"/>	BACT <input type="checkbox"/>	OTHER <input type="checkbox"/>
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ENVIRONMENTAL PROTECTION AGENCY  
Environmental Services Division

EDISON, N.J.

Date of Request 4/3/96 Priority  Immediate  Normal  Deferred  
 Source of Sample(s) Anchor Chemical, Sticksville, NY  
 Sample Number(s) 200130 & 200135  
 Type of Sample  Water  Sediment  Oil  Air  Other (Specify)

**PHYSICAL CHARACTERISTICS**

- |  |   |  |   |
|--|---|--|---|
| <input type="checkbox"/> Turbidity                 | <input type="checkbox"/> Color                        | <input type="checkbox"/> Specific Gravity    | <input type="checkbox"/> Corrosivity (RCRA) |
| <input type="checkbox"/> Volatile Solids           | <input type="checkbox"/> Total Solids                 | <input type="checkbox"/> Viscosity           | <input type="checkbox"/> Other _____        |
| <input type="checkbox"/> Total Suspended Solids    | <input checked="" type="checkbox"/> Dissolved Solids  | <input checked="" type="checkbox"/> % Solids | _____                                       |
| <input type="checkbox"/> Volatile Suspended Solids | <input checked="" type="checkbox"/> Settleable Solids | <input type="checkbox"/> Ignitability (RCRA) | _____                                       |

**ORGANIC/DEMAND ANALYSES**

- |  |   |  |   |
|--|---|--|---|
| <input type="checkbox"/> Day BOD           | <input type="checkbox"/> Phenol                                   | <input type="checkbox"/> Priority Pollutants       | <input checked="" type="checkbox"/> Specific Compound |
| <input type="checkbox"/> COD               | <input type="checkbox"/> Pesticides                               | <input checked="" type="checkbox"/> POA - MCL      | <input type="checkbox"/> Identify _____               |
| <input checked="" type="checkbox"/> TOC    | <input type="checkbox"/> Herbicides                               | <input checked="" type="checkbox"/> NVOA - TCL     | _____   |
| <input type="checkbox"/> TOD               | <input type="checkbox"/> Long-term O <sub>2</sub> Demand (Carbon) | <input type="checkbox"/> Other Major Peaks         | <input type="checkbox"/> Quantitate _____             |
| <input type="checkbox"/> PCB's             | <input type="checkbox"/> Long-term O <sub>2</sub> Demand (Total)  | <input type="checkbox"/> EP Toxicity               | _____   |
| <input type="checkbox"/> Total             | <input type="checkbox"/> Volatile Acids                           | <input type="checkbox"/> Pesticides                | _____   |
| <input type="checkbox"/> Specific Aroclors | <input type="checkbox"/> Oil (Identify)                           | <input type="checkbox"/> Herbicides                | _____   |
|  |   | <input type="checkbox"/> Oil & Grease (Quantitate) |   |

**INORGANIC ANALYSES**

- |  |  |  |                              |  |
|--|--|--|------------------------------|--|
| <input type="checkbox"/> pH              | <input type="checkbox"/> Alkalinity        | <input type="checkbox"/> TKN                                     | <input type="checkbox"/> Cd  | <input type="checkbox"/> Ba            |
| <input type="checkbox"/> Conductivity    | <input type="checkbox"/> CO <sub>3</sub>   | <input type="checkbox"/> Org N                                   | <input type="checkbox"/> Co  | <input type="checkbox"/> Se            |
| <input type="checkbox"/> Salinity        | <input type="checkbox"/> Total             | <input type="checkbox"/> NH <sub>3</sub> -N                      | <input type="checkbox"/> Cu  | <input checked="" type="checkbox"/> Ag |
| <input type="checkbox"/> Chloride        | <input type="checkbox"/> HCO <sub>3</sub>  | <input type="checkbox"/> NO <sub>2</sub> -N                      | <input type="checkbox"/> Pb  | <input type="checkbox"/> Asbestos      |
| <input type="checkbox"/> SO <sub>4</sub> | <input type="checkbox"/> Chlorine Demand   | <input type="checkbox"/> NO <sub>3</sub> -N                      | <input type="checkbox"/> Zn  | <input type="checkbox"/> Hexavalent Cr |
| <input type="checkbox"/> SO <sub>2</sub> | <input type="checkbox"/> Chlorine Residual | <input type="checkbox"/> Total P                                 | <input type="checkbox"/> Fe  |  |
| <input type="checkbox"/> Dissolved S     | <input type="checkbox"/> Free              | <input type="checkbox"/> AH-P                                    | <input type="checkbox"/> Cr  |  |
| <input type="checkbox"/> Hardness        | <input type="checkbox"/> Total             | <input type="checkbox"/> Ortho-P                                 | <input type="checkbox"/> As  |  |
| <input type="checkbox"/> Ca              | <input type="checkbox"/> Acidity           | <input checked="" type="checkbox"/> Metal Scan <u>TAL METALS</u> | <input type="checkbox"/> CN- |  |
| <input checked="" type="checkbox"/> Mg   | <input type="checkbox"/> Free              | <input type="checkbox"/> EP Toxicity (Metals)                    | <input type="checkbox"/> F-  |  |
| <input type="checkbox"/> Total/METHOD    | <input type="checkbox"/> Total             | <input type="checkbox"/> Hg                                      | <input type="checkbox"/> Ni  |  |

**SENSITIVITY / METHOD**

- |   |                                      |  |   |
|---|--------------------------------------|--|---|
| <input type="checkbox"/> COD                    | <input type="checkbox"/> Phosphorous | <input type="checkbox"/> Phenol          | <input type="checkbox"/> Metals           |
| <input type="checkbox"/> High Level (> 50 mg/l) | <input type="checkbox"/> Total       | <input type="checkbox"/> 0-1,000 ppb     | <input type="checkbox"/> Total            |
| <input type="checkbox"/> Low Level (< 50 mg/l)  | <input type="checkbox"/> Dissolved   | <input type="checkbox"/> Above 1,000 ppb | <input type="checkbox"/> Dissolved        |
|   |                                      |  | <input type="checkbox"/> Low Sensitivity  |
|   |                                      |  | <input type="checkbox"/> High Sensitivity |

**MICROBIOLOGY**

- | <table style="width: 100%;"> <tr> <th style="width: 10%;">MF</th> <th style="width: 10%;">MPN</th> <th style="width: 10%;">Est. Range</th> </tr> <tr> <td>TC</td> <td><input type="checkbox"/></td> <td>_____</td> </tr> <tr> <td>FC</td> <td><input type="checkbox"/></td> <td>_____</td> </tr> <tr> <td>FS</td> <td><input type="checkbox"/></td> <td>_____</td> </tr> </table> | MF                       | MPN        | Est. Range | TC | <input type="checkbox"/> | _____ | FC | <input type="checkbox"/> | _____ | FS | <input type="checkbox"/> | _____ | <input type="checkbox"/> Clostridium perfringes<br><input type="checkbox"/> Mutagenicity Tests<br><input type="checkbox"/> Ames Test<br><input type="checkbox"/> Viral Enhancement<br><input type="checkbox"/> Other (Specify)<br><input type="checkbox"/> ATP |
|---|--------------------------|------------|------------|----|--------------------------|-------|----|--------------------------|-------|----|--------------------------|-------|--|
| MF  | MPN                      | Est. Range |            |    |                          |       |    |                          |       |    |                          |       |  |
| TC  | <input type="checkbox"/> | _____      |            |    |                          |       |    |                          |       |    |                          |       |  |
| FC  | <input type="checkbox"/> | _____      |            |    |                          |       |    |                          |       |    |                          |       |  |
| FS  | <input type="checkbox"/> | _____      |            |    |                          |       |    |                          |       |    |                          |       |  |
| <input type="checkbox"/> Pathogens<br><input type="checkbox"/> Bacterial<br><input type="checkbox"/> Viral  |                          |            |            |    |                          |       |    |                          |       |    |                          |       |  |

**BIOLOGY**

- |  |   |
|--|---|
| <input type="checkbox"/> Static                      | <input type="checkbox"/> Flow-Through       |
| <input checked="" type="checkbox"/> 24 Hour Bioassay | <input type="checkbox"/> Static Replacement |
| <input type="checkbox"/> 48 Hour Bioassay            | <input type="checkbox"/> Laboratory         |
| <input type="checkbox"/> 96 Hour Bioassay            | <input type="checkbox"/> On Site            |
| <input checked="" type="checkbox"/> Chronic Bioassay | <input type="checkbox"/> Identify           |
| <input type="checkbox"/> Benthos ID                  | <input type="checkbox"/> Quantitate         |
| <input checked="" type="checkbox"/> Fish ID          |   |

Requested by [Signature] Date 4/3/96 Approved by \_\_\_\_\_ Date \_\_\_\_\_

Remarks

Appendix F, Chain of Custody Record for ESD laboratory

# CHAIN OF CUSTODY RECORD

ENVIRONMENTAL PROTECTION AGENCY - REGION II  
Environmental Services Division  
EDISON, NEW JERSEY 08817

Name of Unit and Address:  
*1001 ...*

Sample Number	Number of Containers	Description of Samples
200124	3	<i>...</i>
200127	5	<i>...</i>
200128	5	<i>...</i>
200133	5	<i>...</i>

Person Assuming Responsibility for Sample: *[Signature]* Time: *1420* Date: *3/25/96*

Sample Number	Relinquished By:	Received By:	Time	Date	Reason for Change of Custody
	<i>[Signature]</i>	<i>[Signature]</i>			<i>...</i>

# CHAIN OF CUSTODY RECORD

ENVIRONMENTAL PROTECTION AGENCY - REGION II  
Environmental Services Division  
EDISON, NEW JERSEY 08817

Name of Unit and Address: *Edison Electric Institute  
20010*

Sample Number	Number of Containers	Description of Samples
20010	3	<p><u>300 (Dioxin) - UCL</u>: 3, 40ml Vial Cool to 4°C</p> <p><u>700 (Dioxin)</u>: 1, 100ml Amber Glass bottle Cool to 4°C</p> <p><u>TPH (Dioxin)</u>: 1, 100ml Amber Glass bottle Cool to 4°C per HPLC, pH=2</p> <p><u>200 (Dioxin)</u>: 3, 40ml Vial Cool to 4°C per HPLC, pH=2</p>

Person Assuming Responsibility for Sample: \_\_\_\_\_ Time \_\_\_\_\_ Date *4/2/82*

Sample Number	Relinquished By:	Received By:	Time	Date	Reason for Change of Custody
	<i>[Signature]</i>	<i>[Signature]</i>	<i>11:15</i>	<i>4/2</i>	<i>L. J. [Signature]</i>
	<i>[Signature]</i>				

# CHAIN OF CUSTODY RECORD

ENVIRONMENTAL PROTECTION AGENCY - REGION II  
Environmental Services Division  
EDISON, NEW JERSEY 08817

Name of Unit and Address: *Anchor Chemical Site  
Hicksville NY*

Sample Number	Number of Containers	Description of Samples
200128	12	VOCs (Drinking Water Std - MCL) : 6, 40ml Vials pres HCl pH < 2 Cool to 4°C TCL NUOCs : 3, 1 Liter Amber Glass Bottle, Cool to 4°C <u>TAL METALS</u> : 3, 1 Liter Glass Bottle, Cool to 4°C, HNO <sub>3</sub> pH < 2
200129	5	VOCs (Drinking Water Std MCL) : 3, 40ml Vials pres HCl, pH < 2 Cool to 4°C TCL NUOCs : 1, 1 Liter Amber Glass Bottle, Cool to 4°C <u>TAL METALS</u> : 1, 1 Liter Glass Bottle Cool to 4°C pres HNO <sub>3</sub> , pH < 2
200131	5	SAME AS 200129
200134	3	VOCs (Drinking Water Std) : 3, 40ml Vials pres HCl pH < 2 Cool to 4°C

Person Assuming Responsibility for Sample: *[Signature]* Time: 1400 Date: 4/2/96

Sample Number	Relinquished By:	Received By:	Time	Date	Reason for Change of Custody
All	<i>[Signature]</i>				