

Chem-Solv, Inc. Site
Cheswold, Kent County, Delaware

Remedial Investigation Report

Volume IV

May 1991



Engineers, Planners, Scientists
and Laboratory Services

REPORT

ARB 19026

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AR309027



IEA Assigned Number Index

Organic Analyses

Case No. 868-006

IEA
Assigned
"EPA"
Number

ECM Engineers Sample ID

BC001	41A
BC002	26AU
BC004	33ADU
BC006	33AU
BC008	Equipment Blank U (2/19/91)
BC010	Trip Blank U (2/19/91)

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BCM

APPENDIX M

GROUNDWATER ANALYTICAL RESULTS AND QUALITY ASSURANCE REVIEW
FEBRUARY 1991

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IEA, INC.
CASE 868-6
SDG EC001
CASE NARRATIVE & VOLATILES

1 OF 1

AR309030

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Industrial & Environmental Analysts, Inc.

P.O. Box 12848

Research Triangle Park, North Carolina 27709

(919) 877-0090

FAX (919) 877-0427

CASE: 868-006

SDG No.: BC001

SAMPLES: 6 Water Samples for CLP Volatile Analyse

The samples were received on February 20, 1991 in good condition. Upon receipt at IEA, each sample was assigned a 5-character "EPA" sample number for simplicity in forms generation. This package makes reference only to those 5-character ID's. Please consult the IEA Assigned Number Index for sample cross-identification. The letter "U" was added to the end of some of the client ID's to indicate that the sample was not filtered.

Some of the volatile RICs were not normalized to the largest non-solvent peak. The raw data for many volatile samples has Lab File ID that exceeds seven digits. This occurred when the autosampler was utilized, and we are in the process of obtaining software that will correct the filename.

I certify that this data package is in compliance with the terms and conditions of the contract, both technically and for completeness, for other than the conditions detailed above. Release of the data contained in this hardcopy data package has been authorized by the Laboratory Manager or his designee, as verified by the following signature.


Becky E. Linvill 4/4/91
CLP Organic Assistant Program Manager

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2A
WATER VOLATILE SURROGATE RECOVERY

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

	EPA SAMPLE NO.	S1 (TOL) #	S2 (BFB) #	S3 (DCE) #	OTHER	TOT OUT
1	VBLK01	99	99	100		0
2	BC006	100	103	96		0
3	BC004	99	100	99		0
4	BC008	99	92	96		0
5	BC010	106	98	97		0
6	BC001	108	97	100		0
7	BC001MSD	102	101	99		0
8	BC001MS	101	100	99		0
9	BC002	100	100	98		0
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QC LIMITS

S1 (TOL) = TOLUENE-d8 (88-110)
 S2 (BFB) = BROMOFLUOROBENZENE (86-115)
 S3 (DCE) = 1,2-DICHLOROCETHANE-D4 (76-114)

Column to be used to flag recovery values

* Values outside of contract required QC limits

D Surrogates diluted out

page 1 of 1

FORM II VOA-1

1 0001
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3A
WATER VOLATILE MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix Spike - EPA Sample No.: BC001

COMPOUND	SPIKE ADDED (UG/L)	SAMPLE CONCENTRATION (UG/L)	MS CONCENTRATION (UG/L)	MS % REC #	QC LIMITS REC.
1,1-DICHLOROETHENE	50.	0.	71.	143	61-145
TRICHLOROETHENE	50.	0.	59.	118	71-120
BENZENE	50.	0.	61.	122	76-127
TOLUENE	50.	0.	59.	118	76-125
CHLOROBENZENE	50.	0.	56.	113	75-130

COMPOUND	SPIKE ADDED (UG/L)	MSD CONCENTRATION (UG/L)	MSD % REC #	% RPD #	QC LIM RPD
1,1-DICHLOROETHENE	50.	72.	144	1	14 61-145
TRICHLOROETHENE	50.	60.	120	1	14 71-120
BENZENE	50.	59.	118	3	11 76-127
TOLUENE	50.	58.	117	1	13 76-125
CHLOROBENZENE	50.	57.	115	1	13 75-130

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits

RPD: 0 out of 5 outside limits

Spike Recovery: 0 out of 10 outside limits

COMMENTS:

FORM III VOA-1

1/87 Rev.

AR309035

1 0002

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4A
VOLATILE METHOD BLANK SUMMARY

Lab Name: IEA Contract: 68-W8-0045
 Lab Code: IEA Case No.: 868-6 SAS No.: SDG Nc.: BC001
 Lab File ID: 0301E02 Lab Sample ID:
 Date Analyzed: 3/ 1/91 Time Analyzed: 22:01
 Matrix: (soil/water) WATER Level: (low/med) LOW
 Instrument ID: msd5

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, MS AND MSD:

	EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	TIME ANALYZED
1	BC006		0301E03	22:36
2	BC004		0301E04	23:10
3	BC008		0301E05	23:50
4	BC010		0301E06	0:26
5	BC001		0301E07	1:05
6	BC001MSD		0301E09	2:29
7	BC001MS		0301E10	3:32
8	BC002		0301E11	4:07
9				
10				
11				
12				
13				
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COMMENTS:

page 1 of 1

FORM IV VOA

1 000009036 1/87 Rev.

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5A
VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: IEA Contract: 68-W8-0045
Lab Code: IEA Case No.: 868-6 SAS No.: SDG No.: BC001
Lab File ID: 02205BF BFB Injection Date: 2/20/91
Instrument ID.: msd5 BFB Injection Time: 0:11
Matrix:(soil/water) WATER Level:(low/med): LOW Column:(pack/cap) CAP

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	26.1
75	30.0 - 60.0% of mass 95	44.2
95	Base peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.6
173	Less than 2.0% of mass 174	.0 (.0)1
174	Greater than 50.0% of mass 95	63.9
175	5.0 - 9.0% of mass 174	5.6 (8.7)1
176	Greater than 95.0%, but less than 101.0% of mass 174	64.4 (100.8)1
177	5.0 - 9.0% of mass 176	5.6 (8.6)2

1-Value is % mass 174

2-Value is % mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDAF

	EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
1	VSTD50		0220502	2/20/91	1:18
2	VSTD100		0220503	2/20/91	2:08
3	VSTD150		0220504	2/20/91	2:42
4	VSTD200		0220505	2/20/91	3:17
5	VSTD20		0220506	2/20/91	3:52
6					
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FORM V VOA

1 0004

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5A
VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: IEA Contract: 68-W8-0045
Lab Code: IEA Case No.: 868-6 SAS No.: SDG No.: BC001
Lab File ID: 0301EBF BFB Injection Date: 3/ 1/91
Instrument ID.: msd5 BFB Injection Time: 20:50
Matrix:(soil/water) WATER Level:(low/med): LOW Column:(pack/cap) CAP

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	18.8
75	30.0 - 60.0% of mass 95	43.3
95	Base peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	7.4
173	Less than 2.0% of mass 174	.0 (.0)1
174	Greater than 50.0% of mass 95	63.6
175	5.0 - 9.0% of mass 174	4.0 (6.3)1
176	Greater than 95.0%, but less than 101.0% of mass 174	63.8 (100.3)1
177	5.0 - 9.0% of mass 176	5.5 (8.7)2

1-Value is % mass 174

2-Value is % mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

	EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
1	VSTD50		0301E01	3/ 1/91	21:24
2	VBLK01		0301E02	3/ 1/91	22:01
3	BC006		0301E03	3/ 1/91	22:36
4	BC004		0301E04	3/ 1/91	23:10
5	BC008		0301E05	3/ 1/91	23:50
6	BC010		0301E06	3/ 2/91	0:26
7	BC001		0301E07	3/ 2/91	1:05
8	BC001MSD		0301E09	3/ 2/91	2:29
9	BC001MS		0301E10	3/ 2/91	3:32
10	BC002		0301E11	3/ 2/91	4:07
11					
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22					

page 1 of 1

FORM V VOA

AR309038
1 0005 1/87 Rev.

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8A
VOLATILE INTERNAL STANDARD AREA SUMMARY

Lab Name: IEA Contract: 68-W8-0045
Lab Code: IEA Case No.: 868-6 SAS No.: SDG No.: BC001
Lab File ID (Standard): 0301E01 Date Analyzed: 3/ 1/91
Instrument ID: msd5 Time Analyzed: 21:24
Matrix:(soil/water) WATER Level:(low/med): LOW Column:(pack/cap) CAP

	IS1(BCM) AREA #	RT	IS2(DFB) AREA #	RT	IS3(CBZ) AREA #	RT
12 HOUR STD	192811.	11.04	864705.	12.91	767497.	18.33
UPPER LIMIT	385622.	11.54	1729410.	13.41	1534994.	18.83
LOWER LIMIT	96406.	10.54	432353.	12.41	383749.	17.83
EPA SAMPLE NO.						
1 VBLK01	191178.	11.05	854111.	12.91	765215.	18.32
2 BC006	188186.	11.06	820858.	12.91	726115.	18.33
3 BC004	191897.	11.06	852931.	12.92	756641.	18.33
4 BC008	199142.	11.06	855639.	12.92	745576.	18.33
5 BC010	189489.	11.05	832828.	12.91	678005.	18.33
6 BC001	182518.	11.06	793181.	12.92	655112.	18.33
7 BC001MSD	191500.	11.07	837886.	12.93	740878.	18.3
8 BC001MS	201213.	11.10	874366.	12.96	781565.	18.37
9 BC002	129449.	11.07	519782.	12.92	455633.	18.34
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						

IS1 (BCM) = BROMOCHLOROMETHANE
IS2 (DFB) = 1,4-DIFLUOROBENZENE
IS3 (CBZ) = CHLOROBENZENE-d5

UPPER LIMIT = + 100%
of internal standard area.
LOWER LIMIT = - 50%
of internal standard area.

Column used to flag internal standard area values with an asterisk

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FORM VIII VOA

1 0006

1/8

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D

SURROGATE AND INTERNAL STANDARD KEY SHEET
FOR VOLATILES
(AS THEY APPEAR ON THE RIC)

A = Bromochloromethane (Internal Standard)
B = 1,2-Dichloroethane-d4 (Surrogate)
C = 1,4-Difluorobenzene (Internal Standard)
D = Toluene-d8 (Surrogate)
E = Chlorobenzene-d5 (Internal Standard)
F = 4-Bromofluorobenzene (Surrogate)

AR309040
1 0007

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC001

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E07

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO.	COMPOUND	CONCENTRATION UNITS: (ug/L or ug/Kg) UG/L	Q
---------	----------	--	---

74-87-3	CHLOROMETHANE	10.	U
74-83-9	BROMOMETHANE	10.	U
75-01-4	VINYL CHLORIDE	10.	U
75-00-3	CHLOROETHANE	10.	U
75-09-2	METHYLENE CHLORIDE	5.	U
67-64-1	ACETONE	10.	U
75-15-0	CARBON DISULFIDE	5.	U
75-35-4	1,1-DICHLOROETHENE	5.	U
75-34-3	1,1-DICHLOROETHANE	5.	U
540-59-0	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3	CHLOROFORM	5.	U
107-06-2	1,2-DICHLOROETHANE	5.	U
78-93-3	2-BUTANONE	10.	U
71-55-6	1,1,1-TRICHLOROETHANE	5.	U
56-23-5	CARBON TETRACHLORIDE	5.	U
108-05-4	VINYL ACETATE	10.	U
75-27-4	BROMODICHLOROMETHANE	5.	U
78-87-5	1,2-DICHLOROPROPANE	5.	U
10061-01-5	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6	TRICHLOROETHENE	5.	U
124-48-1	DIBROMOCHLOROMETHANE	5.	U
79-00-5	1,1,2-TRICHLOROETHANE	5.	U
71-43-2	BENZENE	5.	U
10061-02-6	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2	BROMOFORM	5.	U
108-10-1	4-METHYL-2-PENTANONE	10.	U
591-78-6	2-HEXANONE	10.	U
127-18-4	TETRACHLOROETHENE	5.	U
79-34-5	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3	TOLUENE	5.	U
108-90-7	CHLOROBENZENE	5.	U
100-41-4	ETHYLBENZENE	5.	U
100-42-5	STYRENE	5.	U
1330-20-7	XYLENE (TOTAL)	5.	U

FORM I VOA

AR30904/97 Rev
1 0008

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC001

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E07

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 0

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
1.				
2.				
3.				
4.				
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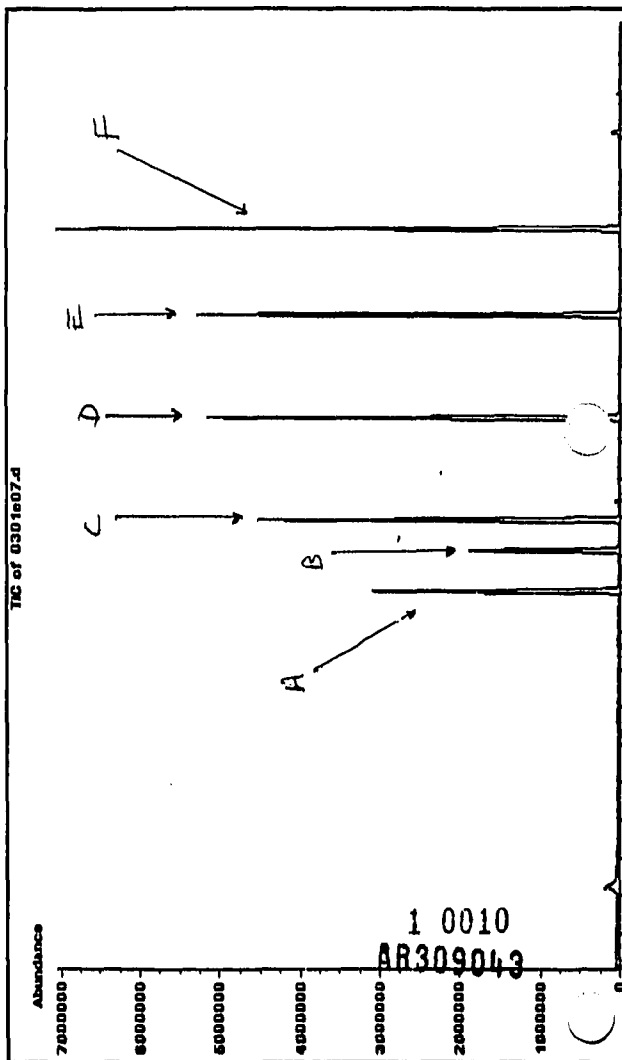
FORM I VOA-TIC

1/87 Rev.

1 00AR309042

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BCR 1
 3/2/91 1:0
 MSD
 12/2/91



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IEA INC CARY, NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.i/021991.b/0219505.d
 Date : 02 Mar 91 01:05 Inst ID: msd5
 Data file: 0301e07.d Tune Date: 23 Jan 91 10:10
 Name Info: BC001 868-006-1 41A 5ml undil ieA msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME		MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
MIN.	(REL)						
* 16 BROMOCHLOROMETHANE CAS #: 74-97-5							
11.05	(1.000)	128	182518	50.00		160.78 - 210.78	100.00
11.05		49	361369			106.54 - 156.54	197.99
11.05		130	238434				130.63
* 20 1,2-DICHLOROETHANE-D4 CAS #: 17060-07-0							
12.12	(1.097)	65	277691	50.23	100.46		100.00
12.12		102	63956			1.65 - 51.65	23.03
* 23 1,4-DIFLUOROBENZENE CAS #: 540-36-3							
12.92	(1.000)	114	793181	50.00		0.00 - 47.22	100.00
12.92		63	196519			0.00 - 45.47	24.77
12.92		88	178303				22.47
24 TRICHLOROETHENE CAS #: 79-01-6							
13.42	(1.040)	130	4647	0.67	BOL		100.00
27 2-CHLOROETHYL VINYL ETHER CAS #: 110-75-8							
14.42	(1.117)	63	537	0.12	BOL NT		100.00
* 37 CHLOROBENZENE-d5 CAS #: 3114-55-4 (OK 4/3/91)							
18.33	(1.000)	117	655112	50.00		37.16 - 87.16	100.00
18.33		82	419761			7.90 - 57.90	64.07
18.33		119	209066				31.91
* 30 TOLUENE-d8 CAS #: 2037-26-5							
15.63	(0.853)	98	745247	53.81	107.63		100.00
* 45 BROMOFIJIOROBENZENE CAS #: 460-00-4							
20.58	(1.122)	95	636176	48.46	96.92	41.39 - 91.39	100.00
20.58		174	423694			41.62 - 91.62	66.60
20.58		176	422036				66.33

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC002

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E11

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE	5.	U
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	5.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE	5.	U
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE	29.	U
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE	5.	U
108-90-7-----	CHLOROBENZENE	5.	U
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

AR309045

FORM I VOA

1 0012

1/87-REV

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC002

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E11

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 10

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
1. 78-78-4	Butane, 2-methyl-	5.49	100.	J
2. - -	UNKNOWN	5.66	100.	J
3. 96-14-0	Pentane, 3-methyl-	8.67	100.	J
4. 96-37-7	Cyclopentane, methyl-	10.43	100.	J
5. - -	UNKNOWN HYDROCARBON	11.72	100.	J
6. - -	UNKNOWN	12.44	100.	J
7. 108-87-2	Cyclohexane, methyl-	13.78	40.	J
8. - -	UNKNOWN HYDROCARBON	14.53	30.	J
9. 560-21-4	Pentane, 2,3,3-trimethyl-	14.78	40.	J
10. 767-58-8	1H-Indene, 2,3-dihydro-1-met	25.58	50.	J
11.				
12.				
13.				
14.				
15.				
16.				
17.				
18.				
19.				
20.				
21.				
22.				
23.				
24.				
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26.				
27.				
28.				
29.				
30.				

FORM I VOA-TIC

1/87 Rev.

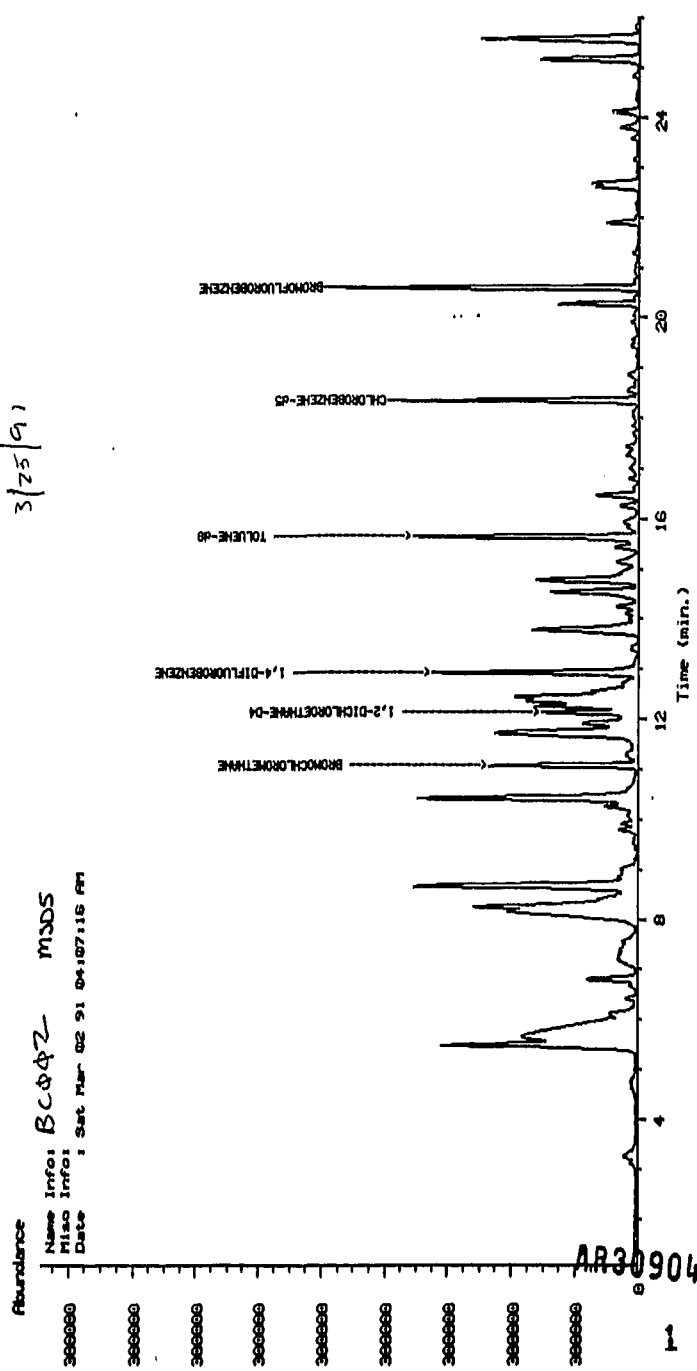
AR309046

1 0013

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CU
3/25/91

TIC of 03012101011.d



Name Info: BCO42 MDS
Misc Info:
Date : Sat Mar 02 91 04:07:16 PM

AR309047

1 0014

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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSDS

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/0219505.d
 Date : 02 Mar 91 04:07 Inst ID: msd5
 Data file: 0301e110111.d Tune Date: 23 Jan 91 10:10
 Name Info: BCLPZ
 Misc Info: *CKM 4/6/91*
 Operator : es Dilution Factor: 1.000
 Comment : voaCIP Method
 Sequence index : 1
 Als bottle num : 11
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE						
11.07 (1.000)	128	129449	50.00		CAS #: 74-97-5	100.00
11.06	49	257048			160.78 - 210.78	198.57
11.07	130	169067			106.54 - 156.54	130.60
7 ACETONE						
7.20 (0.652)	43	66493	145.42		CAS #: 67-64-1	100.00
					F.P	
9 METHYLENE CHLORIDE						
8.26 (0.748)	84	17191	5.00		CAS #: 75-09-2	100.00
					F.P	
17 CHLOROFORM						
11.23 (1.017)	83	11421	1.48		CAS #: 67-66-3	100.00
					BCL	
\$ 20 1,2-DICHLOROETHANE-D4						
12.13 (1.098)	65	191895	48.94		CAS #: 17060-07-0	100.00
12.13	102	45049			97.89	23.47
					1.65 - 51.65	
* 23 1,4-DIFLUOROBENZENE						
12.92 (1.000)	114	519782	50.00		CAS #: 540-36-3	100.00
12.92	63	133888			0.00 - 47.22	25.75
12.92	88	115986			0.00 - 45.47	22.31
21 BENZENE						
12.25 (0.949)	78	249973	29.27		CAS #: 71-43-2	100.00
33 1,1,2-TRICHLOROETHANE						
16.46 (1.275)	97	35678	10.02		CAS #: 79-00-5	100.00
					F.P	
* 37 CHLOROBENZENE-d5						
18.34 (1.000)	117	455633	50.00		CAS #: 3114-55-4	100.00
18.34	82	302103			37.16 - 87.16	66.30
18.34	119	143753			7.90 - 57.90	31.55
29 4-METHYL-2-PENTANONE						
15.47 (0.844)	43	7455	3.27		CAS #: 108-10-1	100.00
					BCL	
\$ 30 TOLUENE-d8						
15.64 (0.853)	98	480636	49.90		CAS #: 2037-26-5	100.00
					99.80	
34 TETRACHLOROETHENE						
16.79 (0.916)	164	6349	1.47		CAS #: 127-18-4	100.00
					BCL	
35 2-HEXANONE						
16.86 (0.919)	43	8217	5.26		CAS #: 591-78-6	100.00
					BCL	

AR309048
 18015

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\$ 45 BROMOFLUOROBENZENE

CAS #: 460-00-4

20.59 (1.123)	95	458815	50.25	100.50		100.00
20.59	174	298891			41.39 - 91.39	65.14
20.59	176	296522			41.62 - 91.62	64.62

TENTATIVELY IDENTIFIED COMPOUNDS

RET TIME	AREA	AMT(UG/L)	QUALITY	LIBRARY	LIB ENTRY
Butane, 2-methyl-			CAS #: 78-78-4		
5.49	189721300	97.95	87	WILEY.1	114154
UNKNOWN			CAS #:		
5.66	285280300	147.29	0		0
Pentane, 3-methyl-			CAS #: 96-14-0		
8.67	282444600	145.82	90	WILEY.1	1031
Cyclopentane, methyl-			CAS #: 96-37-7		
10.43	225870400	116.62	91	WILEY.1	114549
Cyclohexane(DOT			CAS #: 110-82-7		
11.72	202195000	104.39	72	WILEY.1	868
1-Butanol, 2-methyl-			CAS #: 137-32-6		
12.44	343224500	112.94	64	WILEY.1	114845
Cyclohexane, methyl-			CAS #: 108-87-2		
13.78	120067100	39.51	72	WILEY.1	1856
Pentane, 2,3,4-trimethyl-			CAS #: 565-75-3		
14.53	89133060	29.33	72	WILEY.1	4129
Pentane, 2,3,3-trimethyl-			CAS #: 560-21-4		
14.78	122557500	40.33	90	WILEY.1	116590
1H-Indene, 2,3-dihydro-1-methy			CAS #: 767-58-8		
25.58	143853700	49.00	87	WILEY.1	7898

BCΦΦZ
Φ3ΦIE11
MSP5

AR309049
1 001

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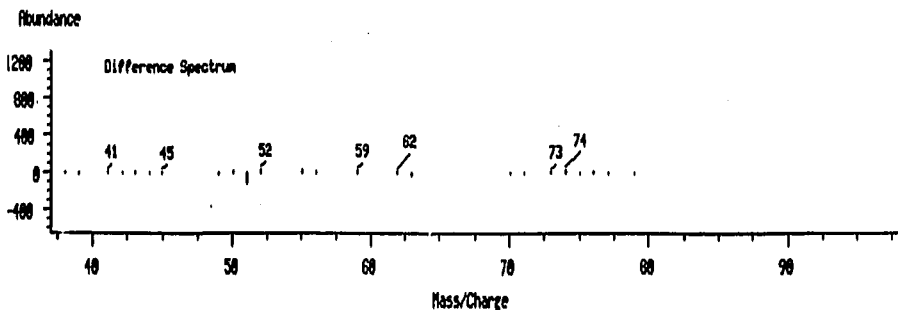
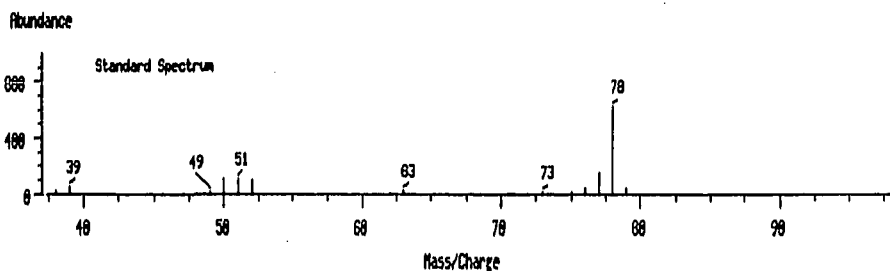
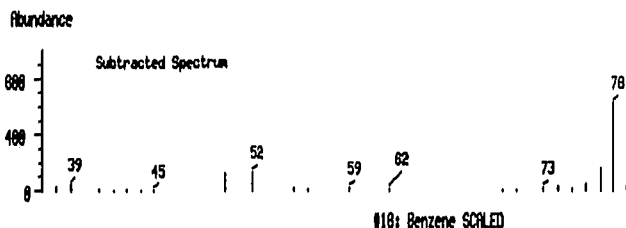
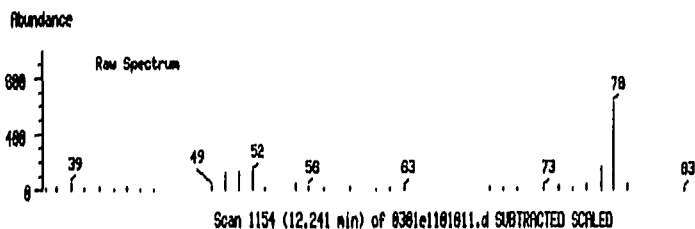
#21 BENZENE

Name Info:

Misc Info:

Date : Sat Mar 02 91 04:07:16 AM

Scan 1154 (12.241 min) of 8381e1101011.d SCALED



AR 30 900507

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Tentatively Identified Compound

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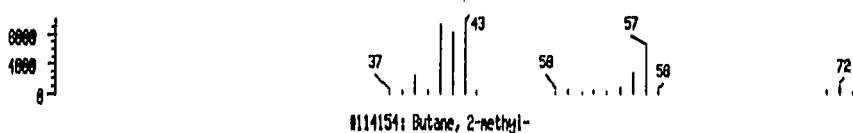
Date : Sat Mar 02 91 04:07:16 AM

PBM Library File: /chem/database/WILEY.1

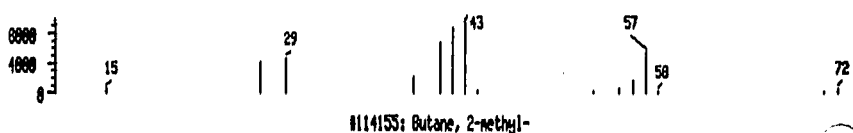
	CAS #	LIB INDEX	MATCH QUALITY
1. Butane, 2-methyl-	78-78-4	114154	87
2. Butane, 2-methyl-	78-78-4	114155	86
3. Butane, 2-methyl-	78-78-4	114153	86
4. Butane, 2-methyl-	78-78-4	424	80
5. Pentane	109-66-0	114156	64

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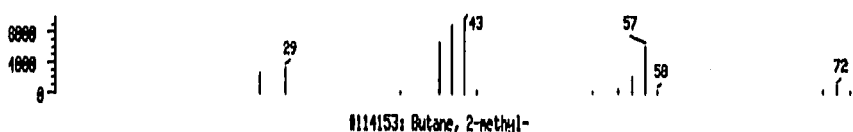
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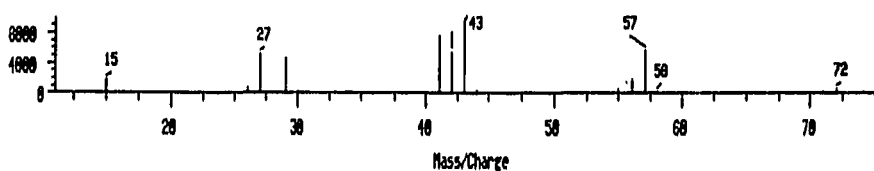
Abundance



Abundance



Abundance



AR309051

1 0018

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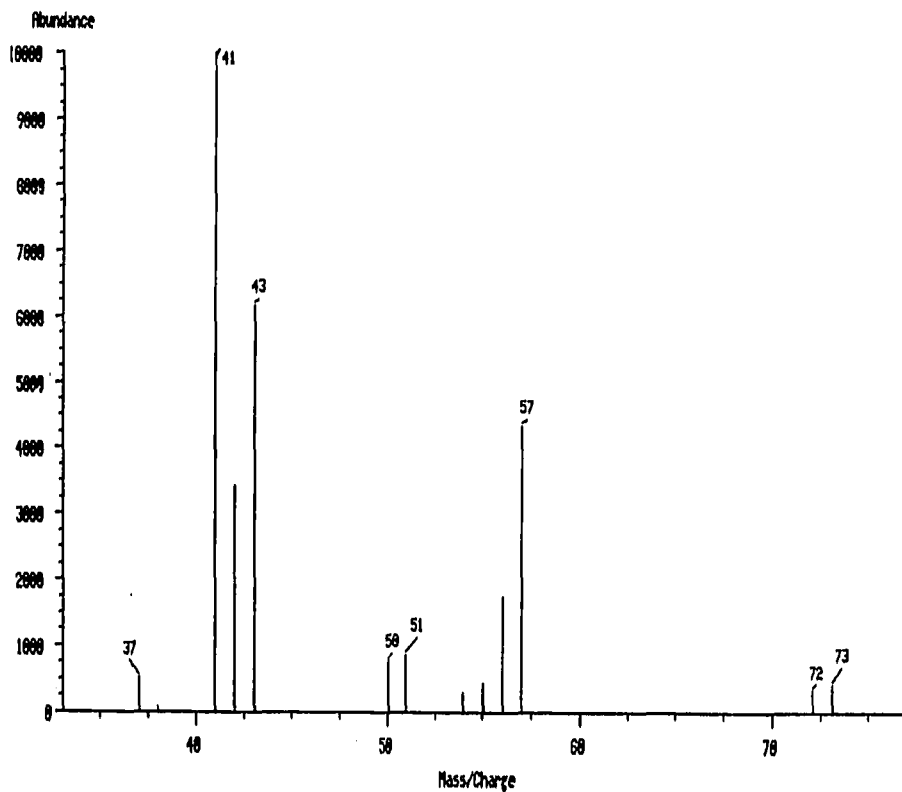
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Date : Sat Mar 02 91 04:07:16 AM

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AR309052
1 0019

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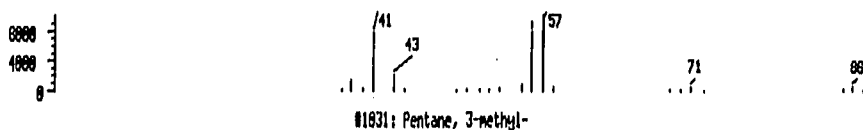
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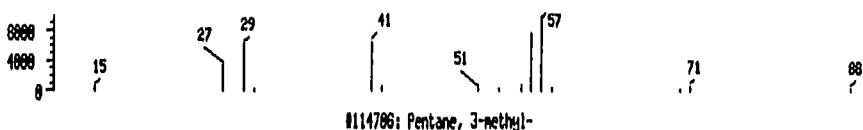
	CAS #	LIB INDEX	MATCH QUALITY
1. Pentane, 3-methyl-	96-14-0	1031	90
2. Pentane, 3-methyl-	96-14-0	114706	90
3. Pentane, 3-methyl-	96-14-0	114707	86

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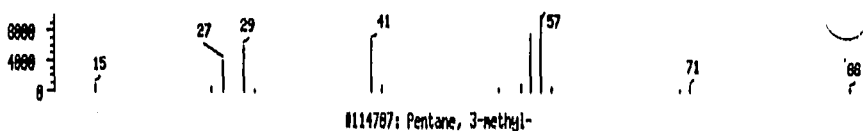
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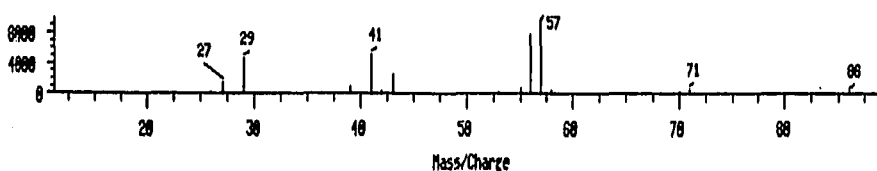
Abundance



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AR 0020
309053

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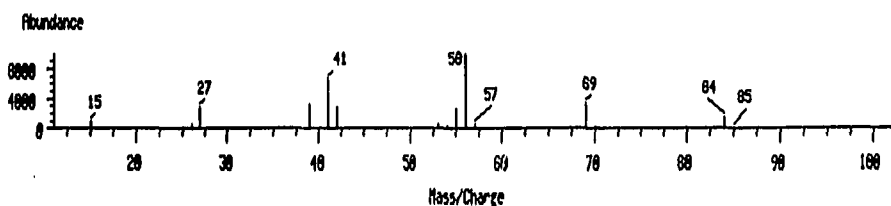
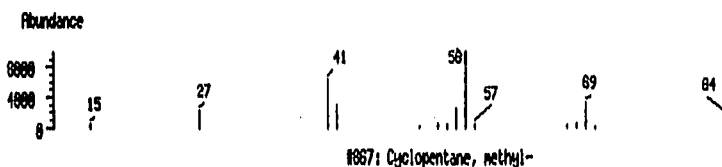
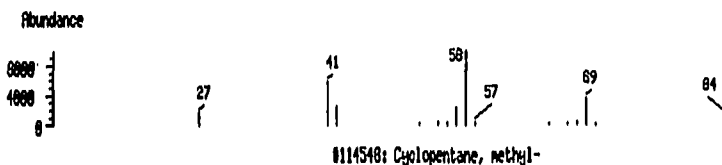
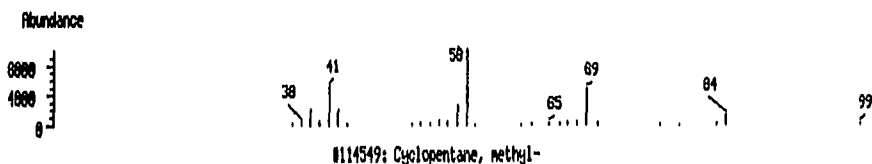
Misc Info:

Date : Sat Mar 02 91 04:07:16 AM

PBM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUALITY
1. Cyclopentane, methyl-	96-37-7	114549	91
2. Cyclopentane, methyl-	96-37-7	114548	91
3. Cyclopentane, methyl-	96-37-7	867	86
4. Cyclopentane, methyl-	96-37-7	114550	78
5. Cyclobutane, ethyl-	4806-61-5	866	72

Scan 967 (18.425 min) of 8381e1181811.d SUBTRACTED SCALED



AR300054

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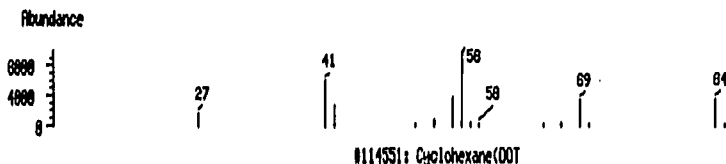
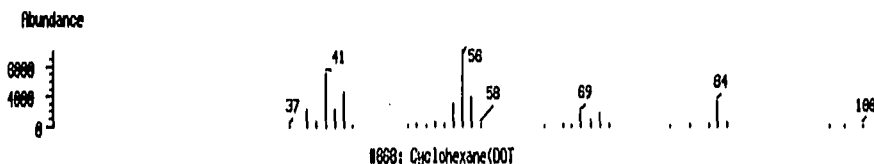
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Date : Sat Mar 02 91 04:07:16 AM

PFM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUAL
1. Cyclohexane(DOT)	110-82-7	868	72
2. 1-Pentene, 2-methyl-	763-29-1	114533	64
3. Cyclohexane(DOT)	110-82-7	114551	56
4. Cyclopentane, methyl-	96-37-7	114548	53

Scan 1100 (11.716 min) of 6301e1101011.d SUBTRACTED SCALED



AR30905
1 0022

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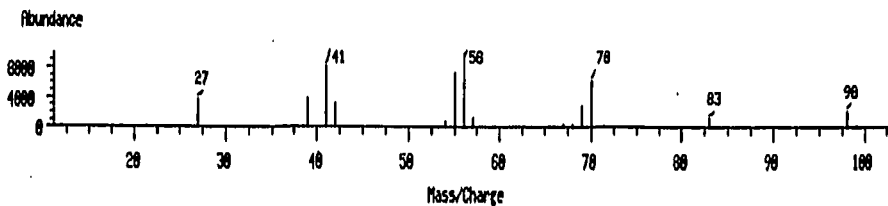
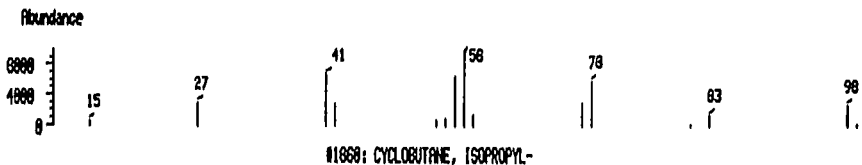
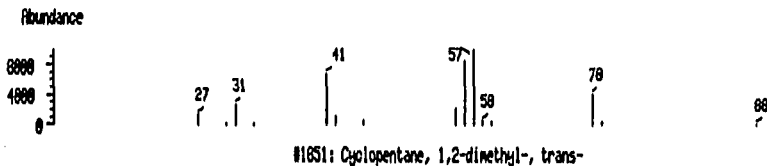
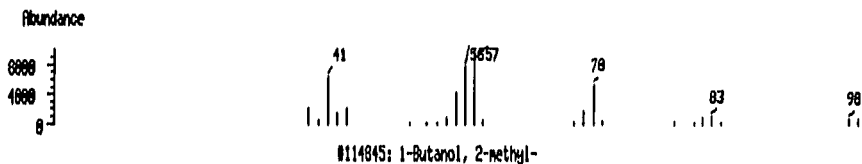
Misc Info:

Date : Sat Mar 02 91 04:07:16 AM

PEM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUALITY
1. 1-Butanol, 2-methyl-	137-32-6	114845	64
2. Cyclopentane, 1,2-dimethyl-, trans-	822-50-4	1851	59
3. CYCLOBUTANE, ISOPROPYL-		1860	59
4. 1-Butanol, 2-methyl-	137-32-6	114848	56
5. Cyclopentane, 1,3-dimethyl-, cis-	2532-58-3	1854	56

Scan 1174 (12.436 min) of 0301101011.d SUBTRACTED SCALED



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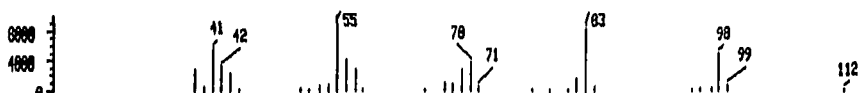
Date : Sat Mar 02 91 04:07:16 AM

PEM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUALITY
1. Cyclohexane, methyl-	108-87-2	1856	72
2. Cyclohexanone	108-94-1	115250	53
3. 3-Penten-2-one, 4-methyl-	141-79-7	115230	50

Scan 1313 (13.785 min) of 8301e1101011.d SUBTRACTED SCALED

Abundance



#1856: Cyclohexane, methyl-

Abundance



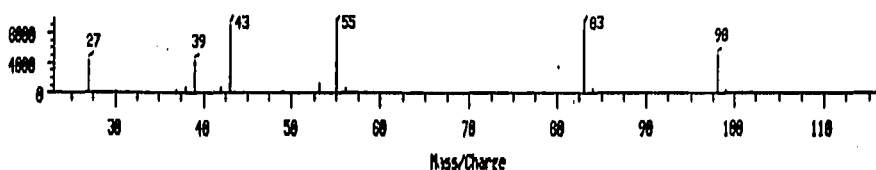
#115250: Cyclohexanone

Abundance



#115230: 3-Penten-2-one, 4-methyl-

Abundance



Mass/Charge

AR309C
1 0029C

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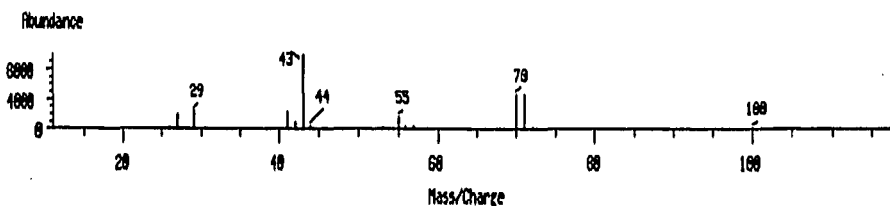
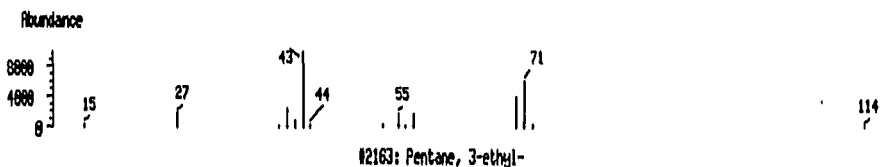
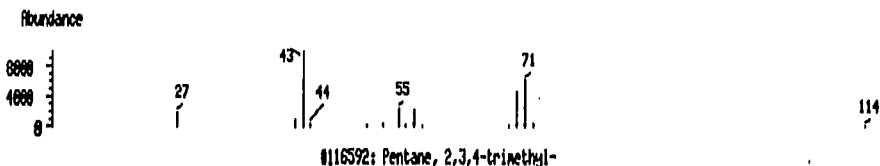
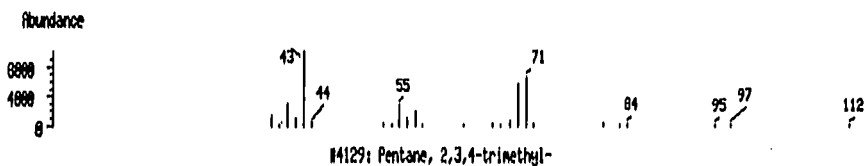
Misc Info:

Date : Sat Mar 02 91 04:07:16 AM

PEM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUALITY
1. Pentane, 2,3,4-trimethyl-	565-75-3	4129	72
2. Pentane, 2,3,4-trimethyl-	565-75-3	116592	72
3. Pentane, 3-ethyl-	617-78-7	2163	72
4. Propanoic acid, 2-methyl-, pentyl e	2445-72-9	120915	72
5. Pentane, 2,3,4-trimethyl-	565-75-3	116593	72

Scan 1370 (14.533 min) of 0301e11011.4 SUBTRACTED SCALED



AR309058
1 0025

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Tentatively Identified Compound

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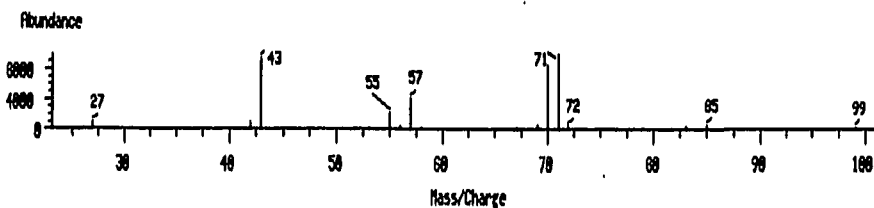
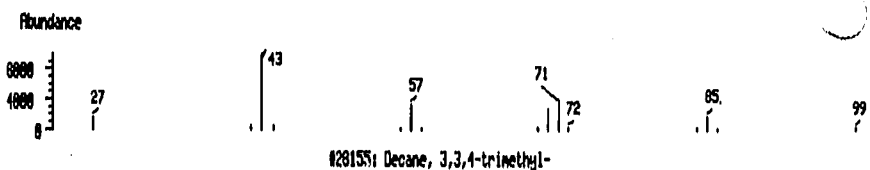
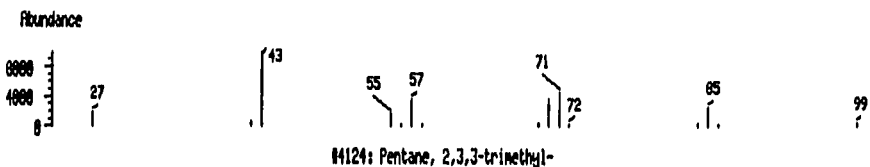
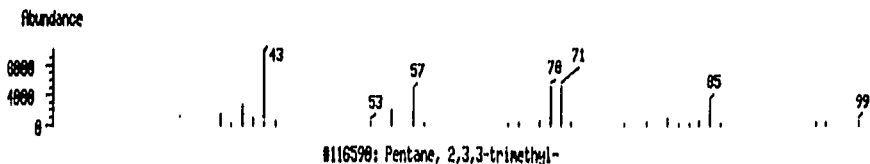
Misc Info:

Date : Sat Mar 02 91 04:07:16 AM

PEM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUALITY
1. Pentane, 2,3,3-trimethyl-	560-21-4	116590	90
2. Pentane, 2,3,3-trimethyl-	560-21-4	4124	83
3. Decane, 3,3,4-trimethyl-	49622-18-6	28155	59
4. Hexane, 2,3,4-trimethyl-	921-47-1	6966	59
5. Pentane, 2,3,4-trimethyl-	565-75-3	4129	59

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AR309059
1 0026

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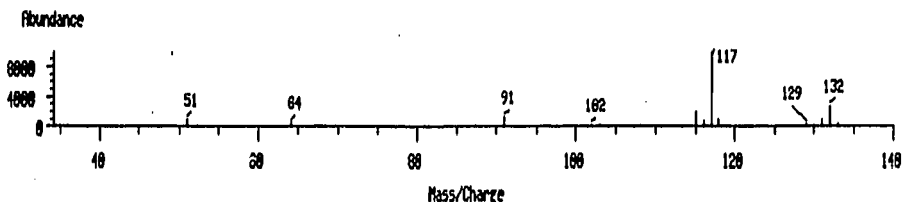
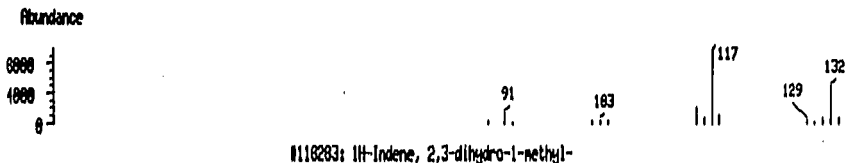
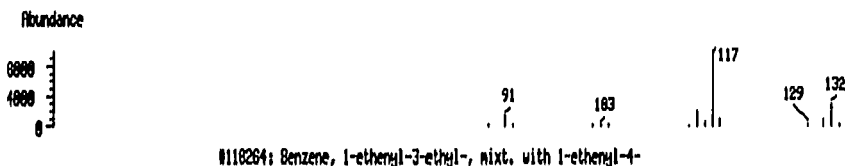
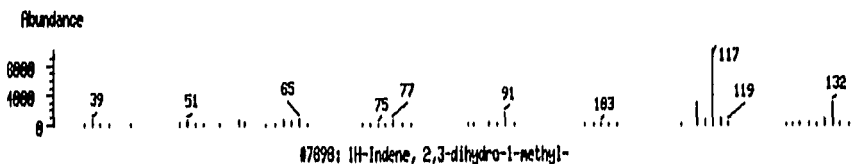
Misc Info:

Date : Sat Mar 02 91 04:07:16 AM

PBM Library File: /chem/database/WILEY.1

	CAS #	LIB INDEX	MATCH QUALITY
1. 1H-Indene, 2,3-dihydro-1-methyl-	767-58-8	7898	87
2. Benzene, 1-ethenyl-3-ethyl-, mixt.	55319-72-7	118264	87
3. 1H-Indene, 2,3-dihydro-1-methyl-	767-58-8	118283	87
4. Benzene, (2-methyl-2-propenyl)-	3290-53-7	7884	86
5. 1H-Indene, 2,3-dihydro-1-methyl-	767-58-8	118284	76

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC004

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E04

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE	5.	U
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	8.	
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE	110.	
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE	5.	U
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE	5.	U
108-90-7-----	CHLOROBENZENE	5.	U
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

AR309

FORM I VOA

1/87 Rev

1 0028

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC004

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 863-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E04

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 0

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
1.				
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FORM I VOA-TIC

1/87 Rev.

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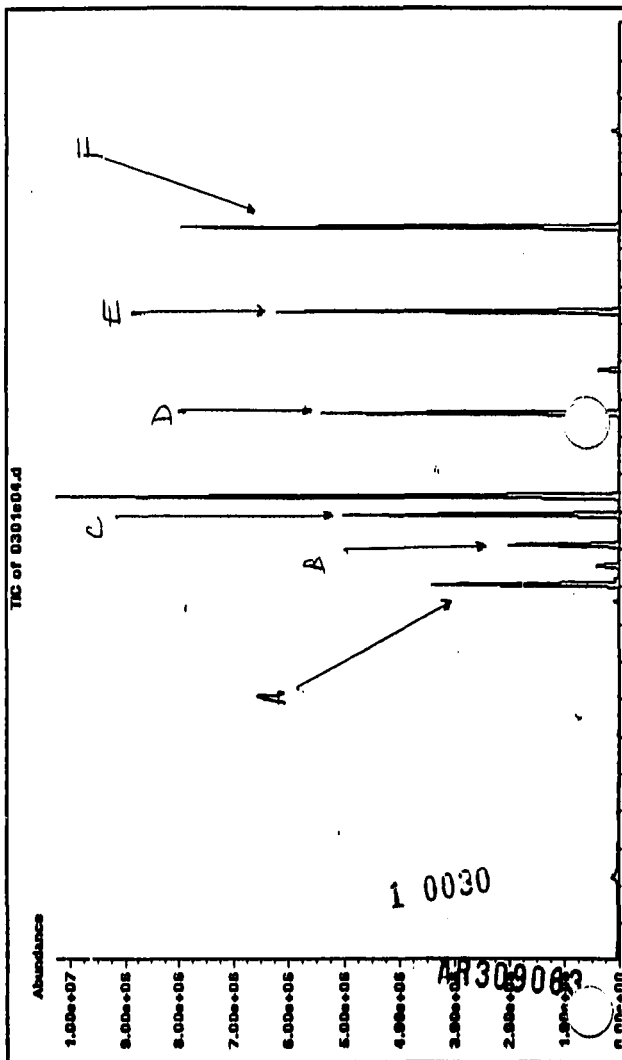
BCR4

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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/0219505.d
 Date : 01 Mar 91 23:10 Inst ID: msd5
 Data file: 0301e04.d Tune Date: 23 Jan 91 10:10
 Name Info: BC004 868-006-4 33ADU 5ml undil lea msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLIP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE					CAS #: 74-97-5	
11.06(1.000)	128	191897	50.00			100.00
11.06	49	397900			160.78 - 210.78	207.35
11.06	130	248107			106.54 - 156.54	129.29
13 cis-1,2-DICHLOROETHENE					CAS #: 156-59-2	
10.61(0.961)	96	6723	1.14	BQL		100.00
14 1,2-DICHLOROETHENE (TOTAL)					CAS #:	
10.61(0.961)	96	6723	1.14	BQL		100.00
17 CHLOROFORM					CAS #: 67-66-3	
11.21(1.015)	83	13915	1.22	BQL		100.00
\$ 20 1,2-DICHLOROETHANE-D4					CAS #: 17060-07-0	
12.12(1.097)	65	286724	49.33	98.66		100.00
12.12	102	67727			1.65 - 51.65	23.62
* 23 1,4-DIFLUOROBENZENE					CAS #: 540-36-3	
12.92(1.000)	114	852931	50.00			100.00
12.91	63	207356			0.00 - 47.22	24.31
12.92	88	184017			0.00 - 45.47	21.57
18 1,1,1-TRICHLOROETHANE					CAS #: 71-55-6	
11.56(0.896)	97	66047	8.11			100.00
19 CARBON TETRACHLORIDE					CAS #: 56-23-5	
11.56(0.896)	117	10082	1.24	BQL		100.00
24 TRICHLOROETHENE					CAS #: 79-01-6	
13.41(1.039)	130	803346	108.28			100.00
* 37 CHLOROBENZENE-d5					CAS #: 3114-55-4	
18.33(1.000)	117	756641	50.00			100.00
18.33	82	488385			37.16 - 87.16	64.54
18.33	119	249876			7.90 - 57.90	33.02
\$ 30 TOLUENE-d8					CAS #: 2037-26-5	
15.63(0.852)	98	791203	49.46	98.93		100.00
34 TETRACHLOROETHENE					CAS #: 127-18-4	
16.78(0.915)	164	20994	2.91	BQL		100.00
\$ 45 BROMOFLUOROBENZENE					CAS #: 460-00-4	
20.57(1.122)	95	756145	49.87	99.73		100.00

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41.62 - 91.62 65.06

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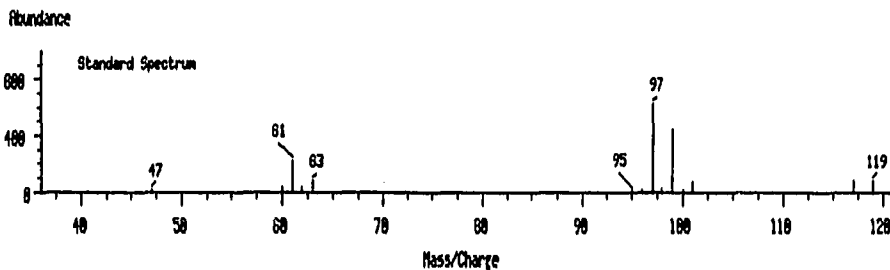
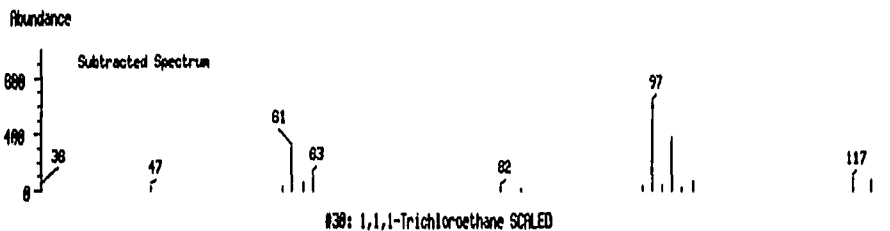
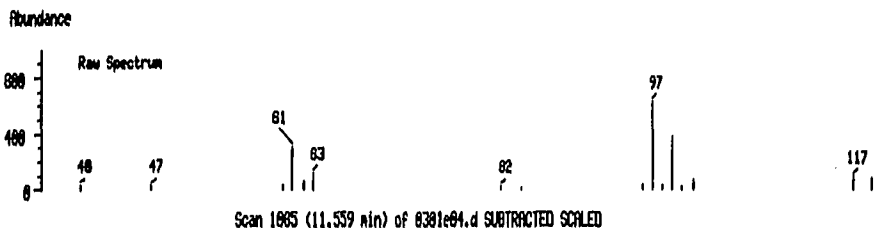
#18 1,1,1-TRICHLOROETHANE

Name Info: BC004 868-006-4 33ADU 5ml undil lea msd5

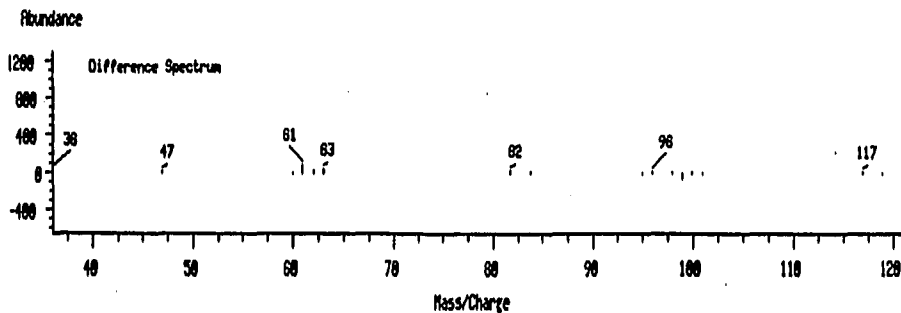
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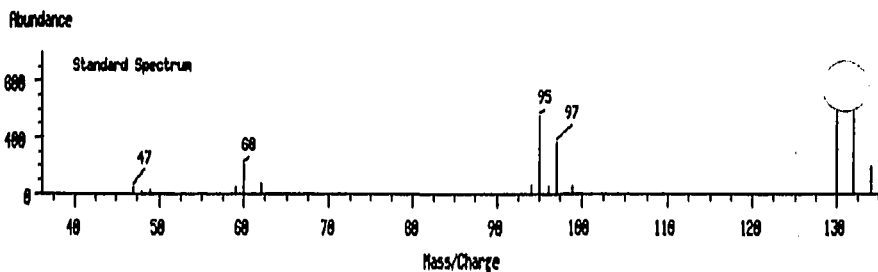
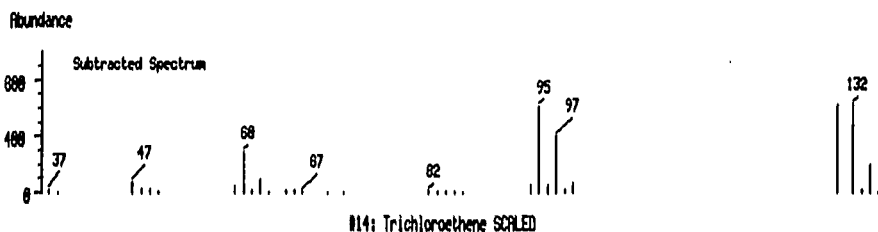
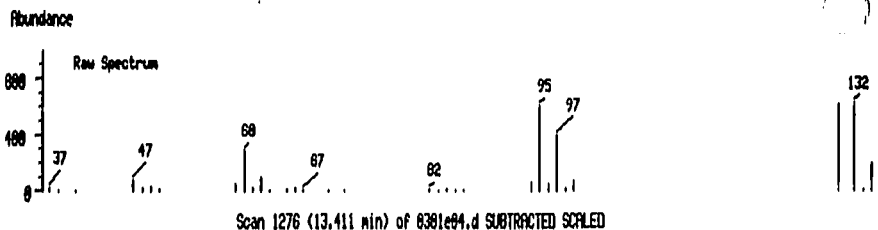
#24 TRICHLOROETHENE

Name Info: BC004 868-006-4 33ADU 5ml undil lea msd5

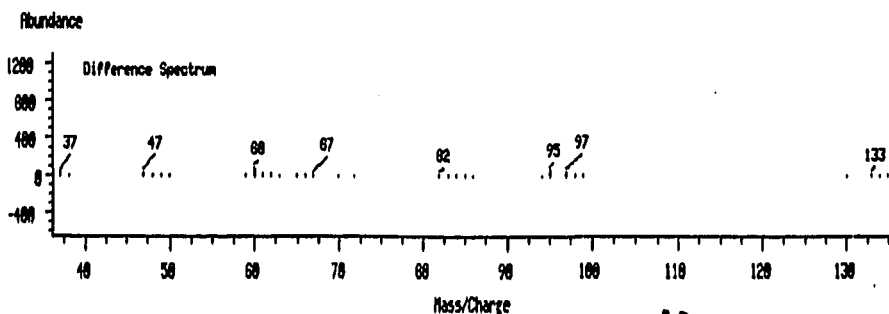
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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC006

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E03

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE	5.	U
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	10.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE	120.	U
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE	5.	U
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	4.	J
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE	5.	U
108-90-7-----	CHLOROBENZENE	5.	U
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

FORM I VOA

1 08399068 Rev.

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC006

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 863-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E03

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 0

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
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FORM I VOA-TIC

1/87 Rev

1 0036

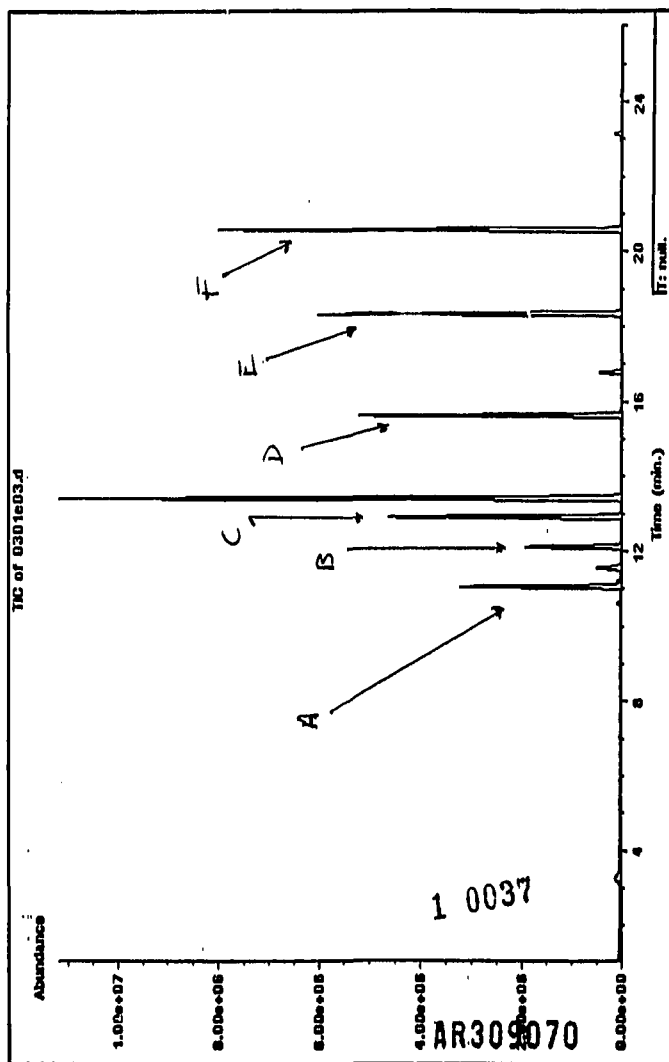
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IEA INC CARY, NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/02195
 Date : 01 Mar 91 22:36 Inst ID: msd5
 Data file: 0301e03.d Tune Date: 23 Jan 91 10:10
 Name Info: BC006 33AU 868-006-6 5ml undil lea msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16	BROMOCHLOROMETHANE				CAS #: 74-97-5	
11.06(1.000)	128	188186	50.00			100.00
11.05	49	380558			160.78 - 210.78	202.22
11.06	130	239864			106.54 - 156.54	127.46
13	cis-1,2-DICHLOROETHENE				CAS #: 156-59-2	
10.60(0.960)	96	8326	1.44	B2L		100.00
14	1,2-DICHLOROETHENE (TOTAL)				CAS #:	
10.60(0.960)	96	8326	1.44	B2L		100.00
17	CHLOROFORM				CAS #: 67-66-3	
11.21(1.015)	83	15294	1.36	B2L		100.00
\$ 20	1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.11(1.097)	65	273099	47.91	95.83		100.00
12.11	102	68468			1.65 - 51.65	25.07
* 23	1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.91(1.000)	114	820858	50.00			100.00
12.91	63	200590			0.00 - 47.22	24.43
12.91	88	174305			0.00 - 45.47	21.23
18	1,1,1-TRICHLOROETHANE				CAS #: 71-55-6	
11.55(0.895)	97	76477	9.76			100.00
19	CARBON TETRACHLORIDE				CAS #: 56-23-5	
11.56(0.896)	117	11210	1.43	B2L		100.00
24	TRICHLOROETHENE				CAS #: 79-01-6	
13.41(1.039)	130	869828	121.83			100.00
25	1,2-DICHLOROPROPANE				CAS #: 78-87-5	
13.42(1.040)	63	1637	0.24	B2L		100.00
* 37	CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.33(1.000)	117	726115	50.00			100.00
18.33	82	471053			37.16 - 87.16	64.87
18.33	119	248055			7.90 - 57.90	34.16
29	4-METHYL-2-PENTANONE				CAS #: 108-10-1	
15.62(0.852)	43	4696	1.29	B2L		100.00
\$ 30	TOLUENE-d8				CAS #: 2037-26-5	
15.63(0.852)	98	768478	50.06	100.13		100.00

If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

34	TETRACHLOROETHENE		CAS #: 127-18-4	
16.78 (0.915)	164	27478	4.00	100.00
\$ 45	BROMOFLUOROBENZENE		CAS #: 460-00-4	
20.57 (1.122)	95	747235	51.35 102.70	100.00
20.57	174	486529	41.39 - 91.39	65.11
20.57	176	481510	41.62 - 91.62	64.43

BCAΦ6
 Φ3Φ1EΦ3
 MSD5

AR3090032

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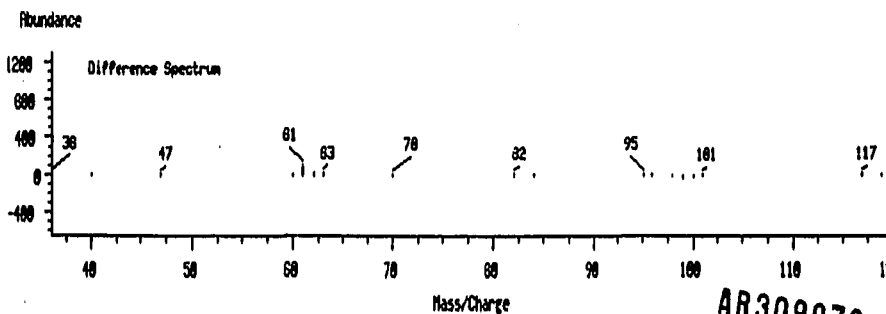
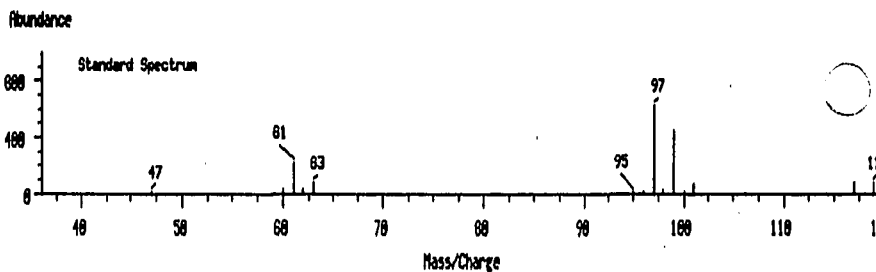
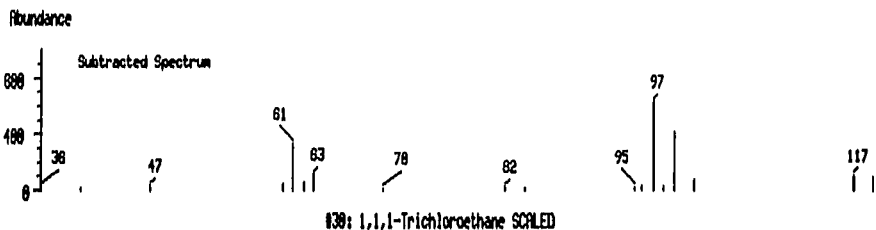
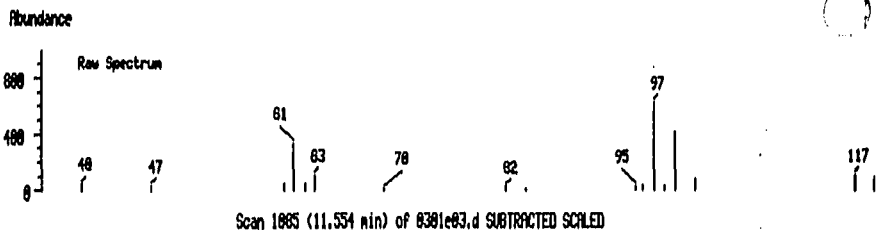
#18 1,1,1-TRICHLOROETHANE

Name Info: BC006 33AU 868-006-6 5ml undil lea msd5

Misc Info:

Date : Fri Mar 01 91 10:36:02 PM

Scan 1865 (11.554 min) of 8301e63.d SCALED



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#24 TRICHLOROETHENE

Name Info: BC006 33AU 868-006-6 5ml undil lea msd5

Misc Info:

Date : Fri Mar 01 91 10:36:02 PM

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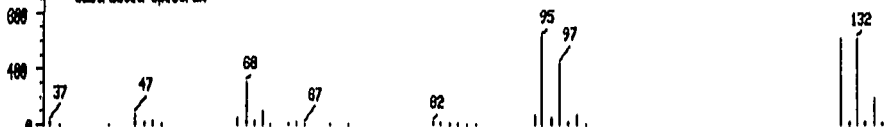
Abundance

Raw Spectrum



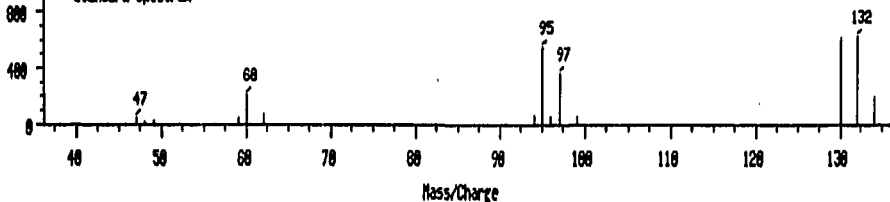
Abundance

Subtracted Spectrum



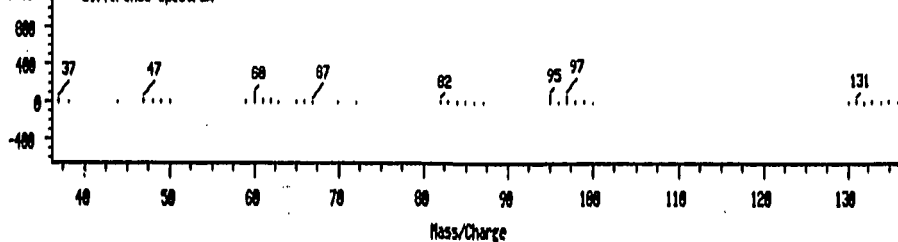
Abundance

Standard Spectrum



Abundance

Difference Spectrum



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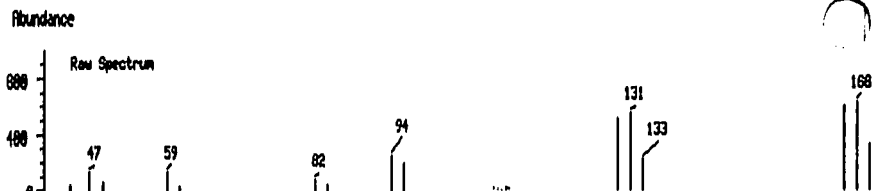
#34 TETRACHLOROETHENE

Name Info: BC006 33AU 868-006-6 5ml undil lea msd5

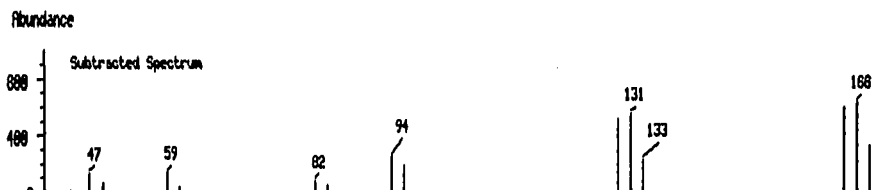
MISC Info:

Date : Fri Mar 01 91 10:36:02 PM

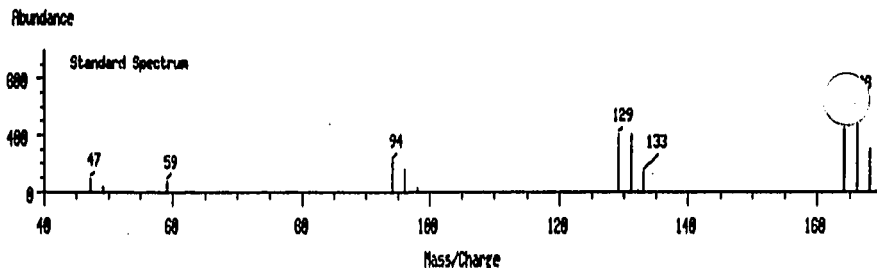
Scan 1624 (16.776 min) of 8301e83.d SCALED



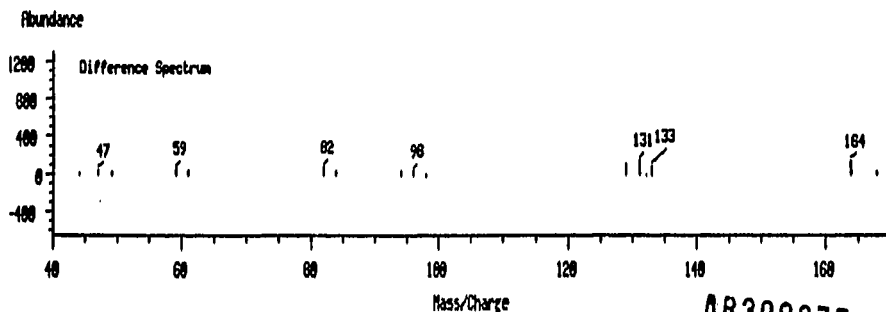
Scan 1624 (16.776 min) of 8301e83.d SUBTRACTED SCALED



#17: Tetrachloroethene SCALED



Scan 1624 (16.776 min) of 8301e83.d SUBTRACTED SCALED



AR309075

1 0042

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC008

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E05

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE	5.	U
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	5.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE	5.	U
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE	5.	U
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE	5.	U
108-90-7-----	CHLOROBENZENE	5.	U
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

AR389076

FORM I VOA

1 0043

1/87 Rev.

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC008

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E05

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 0

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
1.				
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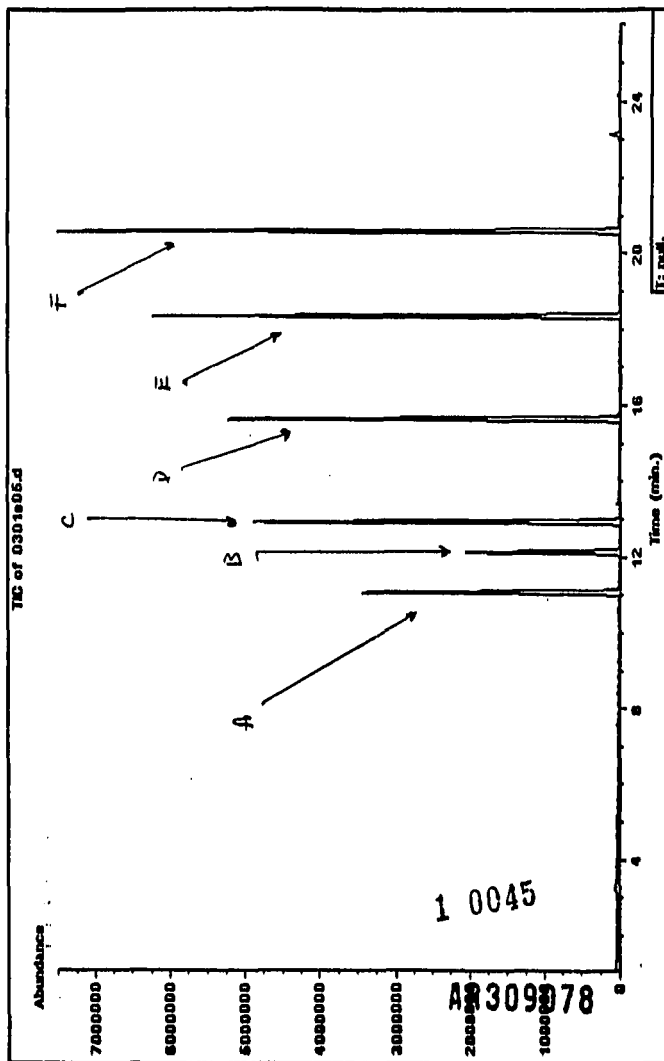
FORM I VOA-TIC

1/87 Rev.
AR309077

1 0044

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BC&B
3/1/91 23:51
MSDS
CK
3/25/91



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IEA INC CARY, NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/02195
 Date : 01 Mar 91 23:50 Inst ID: msd5
 Data file: 0301e05.d Tune Date: 23 Jan 91 10:10
 Name Info: BC008 868-006-8 EQUIPMENT BLANK U 5ml undil ie a msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voacLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE						
11.06(1.000)	128	199142	50.00		CAS #: 74-97-5	100.00
11.06	49	401057			160.78 - 210.78	201.39
11.06	130	264064			106.54 - 156.54	132.60
\$ 20 1,2-DICHLOROETHANE-D4						
12.12(1.098)	65	290628	48.18	96.37	CAS #: 17060-07-0	100.00
12.12	102	72545			1.65 - 51.65	24.96
* 23 1,4-DIFLUOROBENZENE						
12.92(1.000)	114	855639	50.00		CAS #: 540-36-3	100.00
12.92	63	205218			0.00 - 47.22	23.98
12.92	88	180391			0.00 - 45.47	21.08
* 37 CHLOROBENZENE-d5						
18.33(1.000)	117	745576	50.00		CAS #: 3114-55-4	100.00
18.33	82	488271			37.16 - 87.16	65.48
18.34	119	239286			7.90 - 57.90	32.09
\$ 30 TOLUENE-d8						
15.63(0.853)	98	782129	49.62	99.25	CAS #: 2037-26-5	100.00
\$ 45 BROMOFLUOROBENZENE						
20.58(1.122)	95	684760	45.83	91.66	CAS #: 460-00-4	100.00
20.58	174	431224			41.39 - 91.39	62.97
20.58	176	431962			41.62 - 91.62	63.08

1 00A90

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC010

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301206

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE	5.	U
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	5.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE	5.	U
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE	5.	U
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE	5.	U
108-90-7-----	CHLORO BENZENE	5.	U
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

FORM 1 VOA

AR309080
1/87 REV.

1 0047

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC010

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6 - SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E06

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 0

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
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FORM I VOA-TIC

AR309081
1/87

1 0048

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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/021950...d
 Date : 02 Mar 91 00:26 Inst ID: msd5
 Data file: 0301a06.d Tune Date: 23 Jan 91 10:10
 Name Info: BC 010 868-006-10 TRIP BLANK U 5ml undil lea msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE						
11.05(1.000)	128	189489	50.00		CAS #: 74-97-5	100.00
11.05	49	382007			160.78 - 210.78	201.59
11.05	130	241506			106.54 - 156.54	127.45
\$ 20 1,2-DICHLOROETHANE-D4						
12.11(1.097)	65	277808	48.40	96.81	CAS #: 17060-07-0	100.00
12.11	102	65805			1.65 - 51.65	23.68
* 23 1,4-DIFLUOROBENZENE						
12.91(1.000)	114	832828	50.00		CAS #: 540-36-3	100.00
12.91	63	204739			0.00 - 47.22	24.58
12.91	88	184184			0.00 - 45.47	22.11
* 37 CHLOROBENZENE-d5						
18.33(1.000)	117	678005	50.00		CAS #: 3114-55-4	100.00
18.33	82	460971			37.16 - 87.16	67.98
18.33	119	236993			7.90 - 57.90	34.95
\$ 30 TOLUENE-d8						
15.63(0.852)	98	761306	53.12	106.23	CAS #: 2037-26-5	100.00
\$ 45 BROMOFIUFOROBENZENE						
20.57(1.122)	95	668534	49.20	98.41	CAS #: 460-00-4	100.00
20.58	174	433906			41.39 - 91.39	64.90
20.58	176	435011			41.62 - 91.62	65.06

AR309083

1 0050

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6A
VOLATILE ORGANICS INITIAL CALIBRATION DATA

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Instrument ID: msd5

Calibration Date(s): 2/20/91

2/20/91

Matrix: (noil/water) WATER Level: (low/med): LOW Column: (pack/cap) CAP

Min RRF for SPCC(%) = .300 (0.250 for Bromoform) Max %RSD for CCC(*) = 30.0%

LAB FILE ID:		RRF20 = 0220506	RRF50 = 0220502				
RRF100= 0220503		RRF150= 0220504	RRF200= 0220505				
COMPOUND	RRF20	RRF50	RRF100	RRF150	RRF200	RRF	% RSD
CHLOROMETHANE	.826	.860	.853	.826	.845	.842	1.9*
BROMOMETHANE	1.001	1.091	1.109	.906	.849	.991	11.4
VINYL CHLORIDE	* 1.001	1.073	1.078	.989	1.022	1.032	4.0*
CHLOROETHANE	.690	.732	.725	.622	.552	.664	11.4
METHYLENE CHLORIDE	1.197	1.214	1.265	1.157	1.178	1.202	3.4
ACETONE	.183	.132	.104	.124	.147	.138	21.6
CARBON DISULFIDE	2.709	2.588	2.803	2.667	2.619	2.677	3.1
1,1-DICHLOROETHENE	* 1.025	1.017	1.067	.979	.923	1.002	5.4*
1,1-DICHLOROETHANE	* 2.540	2.451	2.603	2.388	2.460	2.488	3.4*
1,2-DICHLOROETHENE (TOTAL)	1.300	1.313	1.367	1.234	1.225	1.288	4.6
CHLOROFORM	* 2.706	2.641	2.722	2.498	2.421	2.598	5.1*
1,2-DICHLOROETHANE	1.530	1.445	1.400	1.321	1.309	1.401	6.5
2-BUTANONE	.072	.068	.053	.066	.066	.065	11.3
1,1,1-TRICHLOROETHANE	.447	.430	.430	.418	.407	.426	3.6
CARBON TETRACHLORIDE	.470	.411	.419	.418	.405	.424	6.1
VINYL ACETATE	.408	.372	.302	.374	.403	.372	11.4
BROMODICHLOROMETHANE	.652	.661	.613	.589	.569	.617	6.4
1,2-DICHLOROPROPANE	* .398	.391	.384	.373	.377	.385	2.7*
CIS-1,3-DICHLOROPROPENE	.578	.579	.537	.543	.526	.553	4.4
TRICHLOROETHENE	.458	.409	.392	.367	.351	.395	10.5
DIBROMOCHLOROMETHANE	.567	.555	.492	.502	.469	.517	8.1
1,1,2-TRICHLOROETHANE	.342	.317	.270	.282	.278	.298	10.3
BENZENE	.815	.750	.744	.705	.699	.743	6.3
TRANS-1,3-DICHLOROPROPENE	.467	.448	.411	.426	.414	.433	5.5
BROMOFORM	* .435	.420	.348	.371	.361	.387	9.9*
4-METHYL-2-PENTANONE	.238	.203	.136	.214	.230	.204	19.7
2-HEXANONE	.157	.133	.106	.139	.154	.138	14.7
TETRACHLOROETHENE	.524	.441	.418	.401	.363	.429	14.0
1,1,2,2-TETRACHLOROETHANE	* .649	.581	.493	.564	.573	.572	9.7*
TOLUENE	* .651	.602	.604	.596	.584	.607	4.2*
CHLOROBENZENE	* .961	.894	.888	.856	.828	.885	5.6*
ETHYLBENZENE	* .410	.370	.378	.377	.371	.381	4.4*
STYRENE	.868	.781	.726	.683	.651	.742	11.6
XYLENE (TOTAL)	.492	.451	.453	.426	.420	.448	6.3
TOLUENE-d8	1.263	1.061	1.061	1.036	.978	1.080	10.0
BROMOFLUOROBENZENE	1.142	.962	.910	.872	.834	.944	12.8
1,2-DICHLOROETHANE-D4	1.473	1.269	1.349	1.289	1.296	1.335	6.2

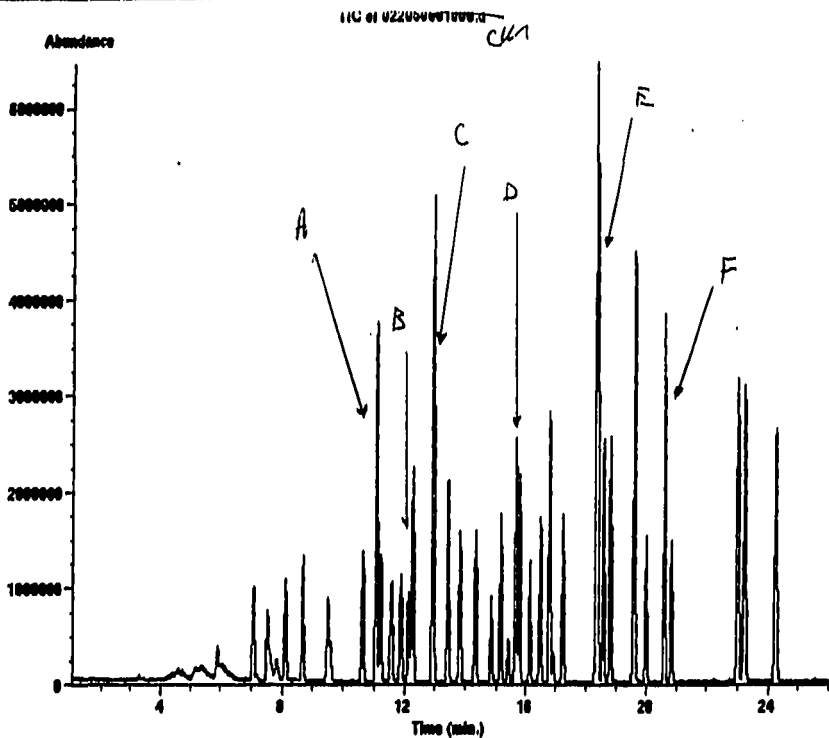
FORM VI VOA

AR309084

1/87 Rev.

1 0051

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VSTD20

2/20/91 03:52

MSD5

AR309085

1 0052

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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date: 20 Feb 91 03:52 Cal Operator: es
 Cal Date: 20 Feb 91 08:06 INSTRUMENT Cal File: d5.1/022091.b/022050601006.d
 Data: 20 Feb 91 03:52 Inst ID: msd5
 Data file: 022050601006.d Tune Date: 23 Jan 91 10:10
 Name Info: *clm*
 Misc Info: *3/25/91*
 VSTD 20
 Operator: es
 Comment: voaCLP Method
 Sequence index: 4
 Als bottle num: 6
 Replicate num: 1
 Dilution Factor: 1.000
 Calibration Sample, Level: 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT (UG/L)	TARGET RANGE	RATIO
* 16	BROMOCHLOROMETHANE			CAS #: 74-97-5	
11.05 (1.000)	128	215696	50.00		100.00
11.05	49	443494		180.61 - 230.61	205.61
11.05	130	281734		105.61 - 155.61	130.61
1	CHLOROMETHANE			CAS #: 74-87-3	
4.38 (0.396)	50	71245	19.62		100.00 (M)
2	VINYL CHLORIDE			CAS #: 75-01-4	
4.67 (0.423)	62	86367	19.39		100.00 (M)
3	BROMOMETHANE			CAS #: 74-83-9	
5.19 (0.470)	94	86402	20.20		100.00
4	CHLOROETHANE			CAS #: 75-00-3	
5.41 (0.490)	64	59559	20.78		100.00
5	TRICHLOROFLUOROMETHANE			CAS #: 75-69-4	
5.87 (0.531)	101	205772	21.78		100.00
6	1,1-DICHLOROETHENE			CAS #: 75-35-4	
7.03 (0.637)	96	88411	20.45		100.00
7	ACETONE			CAS #: 67-64-1	
7.16 (0.648)	43	15811	26.58		100.00
8	CARBON DISULFIDE			CAS #: 75-15-0	
7.49 (0.678)	76	233730	20.24		100.00
9	METHYLENE CHLORIDE			CAS #: 75-09-2	
8.08 (0.73)	84	103274	19.91		100.00
10	trans-1,2-DICHLOROETHENE			CAS #: 540-59-0	
8.65 (0.783)	96	105103	20.13		100.00
11	1,1-DICHLOROETHANE			CAS #: 75-34-3	
9.48 (0.858)	63	219160	20.42		100.00
13	cis-1,2-DICHLOROETHENE			CAS #: 156-59-2	
10.61 (0.960)	96	119257	20.25		100.00
14	1,2-DICHLOROETHENE (TOTAL)			CAS #: AR309086	
10.61 (0.960)	96	119257 224,360	20.25	40.00	100.00
15	2-BUTANONE			CAS #: 78-93-3	

1 0053

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10.62(0.961)	72	6246	22.26		100.00
17 CHLOROFORM				CAS #: 67-66-3	
11.20(1.014)	83	233442	20.83		100.00
\$ 20 1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.12(1.096)	65	127083	22.06		100.00
12.12	102	31963		0.15 - 50.15	25.15
22 1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.26(1.109)	62	132007	21.84		100.00
* 23 1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.92(1.000)	114	905772	50.00		100.00
12.92	63	220761		0.00 - 49.37	24.37
12.92	88	197404		0.00 - 46.79	21.79
12 VINYL ACETATE				CAS #: 108-05-4	
9.57(0.741)	43	147872	21.96		100.00
18 1,1,1-TRICHLOROETHANE				CAS #: 71-55-6	
11.55(0.894)	97	162107	20.99		100.00
19 CARBON TETRACHLORIDE				CAS #: 56-23-5	
11.88(0.919)	117	170153	22.13		100.00
21 BENZENE				CAS #: 71-43-2	
12.24(0.948)	78	295399	21.96		100.00
24 TRICHLOROETHENE				CAS #: 79-01-6	
13.42(1.038)	130	165949	23.17		100.00
25 1,2-DICHLOROPROPANE				CAS #: 78-87-5	
13.82(1.070)	63	144347	20.71		100.00
26 BROMODICHLOROMETHANE				CAS #: 75-27-4	
14.31(1.108)	83	236074	21.13		100.00
27 2-CHLOROETHYL VINYL ETHER				CAS #: 110-75-8	
14.83(1.148)	63	97625	23.36		100.00
28 CIS-1,3-DICHLOROPROPENE				CAS #: 10061-01-5	
15.13(1.171)	75	221943	22.16		100.00
32 TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
16.13(1.249)	75	155691	19.84		100.00
33 1,1,2-TRICHLOROETHANE				CAS #: 79-00-5	
16.48(1.276)	97	124019	22.97		100.00
36 DIBROMOCHLOROMETHANE				CAS #: 124-48-1	
17.24(1.334)	129	205424	21.94		100.00
44 BROMOFORM				CAS #: 75-25-2	
19.97(1.546)	173	157732	22.50		100.00
* 37 CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.33(1.000)	117	758547	50.00		100.00
18.33	82	480807		38.38 - 88.38	63.38
18.33	119	248564		74.38 - 90.38	32.76
\$ 29 TOLUENE-d8				CAS #: 2037-26-5	
15.63(0.853)	98	383296	23.40		100.00
30 TOLUENE				CAS #: 108-88-3	
15.76(0.859)	92	197652	21.45		100.00

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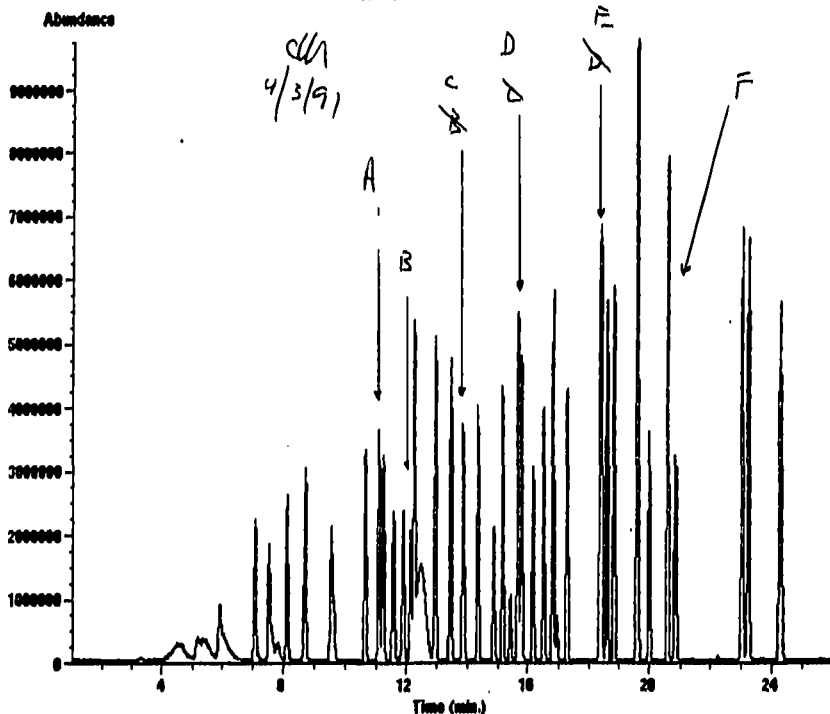
31 4-METHYL-2-PENTANONE			CAS #: 108-10-1	
15.39(0.839) 43	72312	23.33		100.00
34 TETRACHLOROETHENE			CAS #: 127-18-4	
16.78(0.915) 164	159119	24.42		100.00
35 2-HEXANONE			CAS #: 591-78-6	
16.92(0.923) 43	47780	22.83		100.00
38 CHLOROBENZENE			CAS #: 108-90-7	
18.39(1.003) 112	291523	21.71		100.00
39 ETHYLBENZENE			CAS #: 100-41-4	
18.57(1.013) 106	124465	21.53		100.00
40 m-, p-XYLENE			CAS #: 108-38-3	
18.80(1.025) 106	154201	21.73		100.00
41 o-XYLENE			CAS #: 95-47-6	
19.56(1.067) 106	144244	22.17		100.00
42 XYLENE (TOTAL)	298 845	40	CAS #: 1330-20-7	
19.56(1.067) 106	144264	22.17		100.00
43 STYRENE			CAS #: 100-42-5	
19.58(1.068) 104	263486	23.42		100.00
\$ 45 BROMOFLUOROBENZENE			CAS #: 460-00-4	
20.58(1.122) 95	346462	24.19		100.00
20.58 174	232076		41.98 - 91.98	66.98
20.58 176	226442		40.35 - 90.35	65.35
46 1,1,2,2-TETRACHLOROETHANE			CAS #: 79-34-5	
20.82(1.135) 83	196827	22.69		100.00
47 1,3-DICHLOROBENZENE			CAS #: 541-73-1	
23.20(1.266) 146	338704	23.52		100.00
48 1,4-DICHLOROBENZENE			CAS #: 106-46-7	
24.22(1.321) 146	320845	23.29		100.00
49 1,2-DICHLOROBENZENE			CAS #: 95-50-1	
24.22(1.321) 146	320845	23.29		100.00

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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 20 Feb 91 01:18 Cal Operator : es
 Cal Date : 20 Feb 91 07:37 INSTRUMENT Cal File: em/msd5.1/022091.b/0220502.d
 Date : 20 Feb 91 01:18 Inst ID: msd5
 Data file: 0220502.d Tune Date: 23 Jan 91 10:10
 Name Info: vstd50 lea msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method Calibration Sample, Level: 2
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	TARGET RANGE	RATIO
* 16	BROMOCHLOROMETHANE			CAS #: 74-97-5	
11.07(1.000)	128	214697	50.00		100.00
11.07	49	416544		169.01 - 219.01	194.01
11.07	130	283341		106.97 - 156.97	131.97
1	CHLOROMETHANE			CAS #: 74-87-3	
4.42(0.399)	50	184693	61.90		100.00
2	VINYL CHLORIDE			CAS #: 75-01-4	
4.59(0.414)	62	230303	59.09		100.00 (M)
3	BROMOMETHANE			CAS #: 74-83-9	
5.17(0.467)	94	234232	55.09		100.00
4	CHLOROETHANE			CAS #: 75-00-3	
5.42(0.489)	64	157072	55.07		100.00
5	TRICHLOROFLUOROMETHANE			CAS #: 75-69-4	
5.90(0.533)	101	483641	58.21		100.00 (M)
6	1,1-DICHLOROETHENE			CAS #: 75-35-4	
7.06(0.638)	96	218430	50.76		100.00
7	ACETONE			CAS #: 67-64-1	
7.18(0.648)	43	28256	47.73		100.00
8	CARBON DISULFIDE			CAS #: 75-15-0	
7.52(0.679)	76	555682	48.34		100.00
9	METHYLENE CHLORIDE			CAS #: 75-09-2	
8.10(0.732)	84	260736	50.51		100.00
10	trans-1,2-DICHLOROETHENE			CAS #: 540-59-0	
8.67(0.783)	96	264551	50.91		100.00
11	1,1-DICHLOROETHANE			CAS #: 75-34-3	
9.51(0.858)	63	526264	49.25		100.00
13	cis-1,2-DICHLOROETHENE			CAS #: 156-59-2	
10.63(0.960)	96	299334	51.06		100.00
14	1,2-DICHLOROETHENE (TOTAL)			CAS #:	
10.63(0.960)	96	299334	51.06		100.00
15	2-BUTANONE			CAS #: 78-93-3	

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10.64(0.961)	72	14691	52.59		100.00
17 CHLOROFORM				CAS #: 67-66-3	
11.22(1.013)	83	566972	50.83		100.00
\$ 20 1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.14(1.096)	65	272545	47.53		100.00
12.14	102	73521		1.97 - 51.97	26.97
22 1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.28(1.109)	62	310147	51.56		100.00
* 23 1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.94(1.000)	114	906608	50.00		100.00
12.93	63	203520		0.00 - 47.44	22.44
12.93	88	192577		0.00 - 46.24	21.24
12 VINYL ACETATE				CAS #: 108-05-4	
9.60(0.742)	43	337147	50.02		100.00
18 1,1,1-TRICHLOROETHANE				CAS #: 71-55-6	
11.58(0.895)	97	389745	50.41		100.00
19 CARBON TETRACHLORIDE				CAS #: 56-23-5	
11.90(0.920)	117	372323	48.39		100.00
21 BENZENE				CAS #: 71-43-2	
12.26(0.948)	78	679931	50.50		100.00
24 TRICHLOROETHENE				CAS #: 79-01-6	
13.43(1.038)	130	370729	51.72		100.00
25 1,2-DICHLOROPROPANE				CAS #: 78-87-5	
13.83(1.069)	63	354775	50.86		100.00
26 BROMODICHLOROMETHANE				CAS #: 75-27-4	
14.33(1.108)	83	599437	53.62		100.00
27 2-CHLOROETHYL VINYL ETHER				CAS #: 110-75-8	
14.85(1.148)	63	228378	54.59		100.00
28 CIS-1,3-DICHLOROPROPENE				CAS #: 10061-01-5	
15.14(1.171)	75	556783	55.55		100.00
32 TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
16.15(1.248)	75	373267	47.53		100.00
33 1,1,2-TRICHLOROETHANE				CAS #: 79-00-5	
16.50(1.275)	97	287835	53.26		100.00
36 DIBROMOCHLOROMETHANE				CAS #: 124-48-1	
17.25(1.334)	129	503024	53.67		100.00
44 BROMOFORM				CAS #: 75-25-2	
19.98(1.545)	173	380382	54.22		100.00
* 37 CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.35(1.000)	117	769054	50.00		100.00
18.35	82	474179		36.65 - 86.65	61.65
18.35	119	260311		8.84 - 58.84	33.84
\$ 29 TOLUENE-d8				CAS #: 2037-26-5	
15.65(0.853)	98	815608	49.11		100.00
30 TOLUENE				CAS #: 108-88-3	
15.77(0.860)	92	463095	49.57		100.00

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31 4-METHYL-2-PENTANONE			CAS #: 108-10-1	
15.40(0.839) 43	155976	107.24		100.00
34 TETRACHLOROETHENE			CAS #: 127-18-4	
16.80(0.916) 164	339338	51.37		100.00
35 2-HEXANONE			CAS #: 591-78-6	
16.93(0.923) 43	102245	48.19		100.00
38 CHLOROBENZENE			CAS #: 108-90-7	
18.41(1.003) 112	687606	50.50		100.00
39 ETHYLBENZENE			CAS #: 100-41-4	
18.59(1.013) 106	284372	48.51		100.00
40 m-, p-XYLENE			CAS #: 108-38-3	
18.81(1.025) 106	353186	49.09		100.00
41 o-XYLENE			CAS #: 95-47-6	
19.57(1.067) 106	340593	51.64		100.00
42 XYLENE (TOTAL)			CAS #: 1330-20-7	
19.57(1.067) 106	340631 693,779	51.64 100	ck 4/2/91	100.00
43 STYRENE			CAS #: 100-42-5	
19.59(1.068) 104	600579	52.64		100.00
\$ 45 BROMOFLUOROBENZENE			CAS #: 460-00-4	
20.59(1.122) 95	740052	50.97		100.00
20.59 174	489949		41.20 - 91.20	66.20
20.59 176	479546		39.79 - 89.79	64.79
46 1,1,2,2-TETRACHLOROETHANE			CAS #: 79-34-5	
20.83(1.135) 83	446527	50.77		100.00
47 1,3-DICHLOROBENZENE			CAS #: 541-73-1	
23.23(1.266) 146	752766	51.55		100.00
48 1,4-DICHLOROBENZENE			CAS #: 106-46-7	
24.25(1.321) 146	717135	51.34		100.00
49 1,2-DICHLOROBENZENE			CAS #: 95-50-1	
24.25(1.321) 146	717135	51.34		100.00

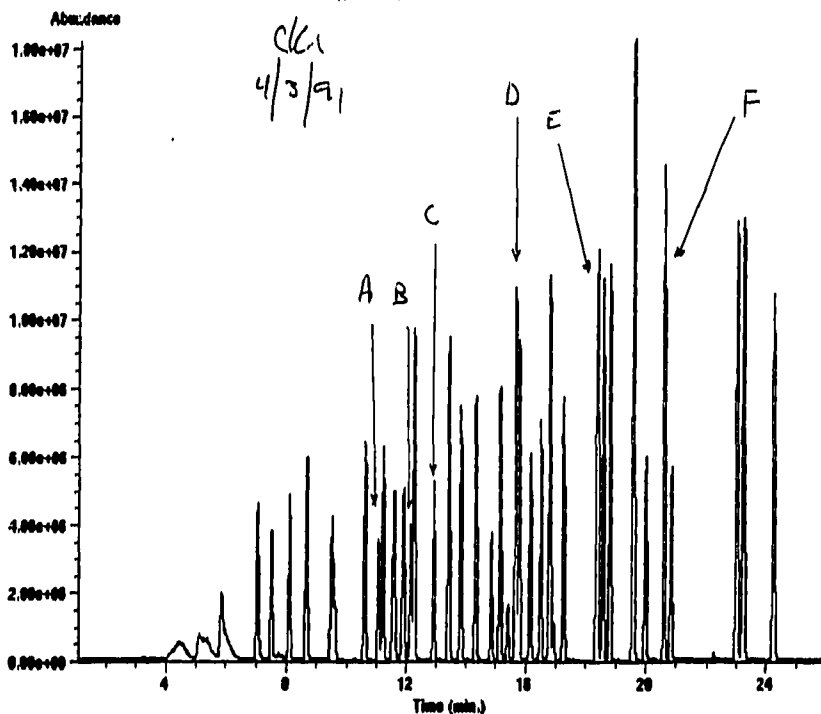
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VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 20 Feb 91 02:08 Cal Operator : es
 Cal Date : 20 Feb 91 07:44 INSTRUMENT Cal File: d5.i/022091.b/022050301003.d
 Date : 20 Feb 91 02:08 Inst ID: msd5
 Data file: 022050301003.d Tune Date: 23 Jan 91 10:10
 Name Info: VSTD 100 3/25/91
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voacLP Method Calibration Sample, Level: 3
 Sequence index : 1
 Als bottle num : 3
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE				CAS #: 74-97-5	
11.06(1.000)	128	205957	50.00		100.00
11.06	49	402758		170.55 - 220.55	195.55
11.06	130	266878		104.57 - 154.57	129.57
1 CHLOROMETHANE				CAS #: 74-87-3	
4.30(0.389)	50	351478	112.52		100.00 (M)
2 VINYL CHLORIDE				CAS #: 75-01-4	
4.51(0.408)	62	444098	118.77		100.00
3 BROMOMETHANE				CAS #: 74-83-9	
5.13(0.464)	94	457004	112.04		100.00
4 CHLOROETHANE				CAS #: 75-00-3	
5.36(0.485)	64	298471	109.08		100.00
5 TRICHLOROFLUOROMETHANE				CAS #: 75-69-4	
5.87(0.531)	101	1003328	125.88		100.00
6 1,1-DICHLOROETHENE				CAS #: 75-35-4	
7.05(0.637)	96	439510	106.48		100.00
7 ACETONE				CAS #: 67-64-1	
7.16(0.647)	43	42754	75.29		100.00
8 CARBON DISULFIDE				CAS #: 75-15-0	
7.50(0.678)	76	1154464	104.69		100.00
9 METHYLENE CHLORIDE				CAS #: 75-09-2	
8.09(0.731)	84	520929	105.20		100.00
10 trans-1,2-DICHLOROETHENE				CAS #: 540-59-0	
8.66(0.783)	96	525961	105.52		100.00
11 1,1-DICHLOROETHANE				CAS #: 75-34-3	
9.49(0.858)	63	1072208	104.60		100.00
13 cis-1,2-DICHLOROETHENE				CAS #: 156-59-2	
10.61(0.959)	96	599914	106.68		100.00
14 1,2-DICHLOROETHENE (TOTAL)			2.00	CAS #:	
10.61(0.959)	96	599914	106.68		
15 2-BUTANONE		1125875	100	CAS #: 78-93-3	1.0061

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10.63(0.961)	72	21778	81.27		100.00
17 CHLOROFORM				CAS #: 67-66-3	
11.21(1.014)	83	1121346	104.80		100.00
\$ 20 1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.12(1.096)	65	555685	101.02		100.00
12.12	102	145123		1.11 - 51.11	26.11
22 1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.26(1.109)	62	576870	99.97		100.00
* 23 1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.92(1.000)	114	947948	50.00		100.00
12.92	63	223008		0.00 - 48.52	23.52
12.92	88	200184		0.00 - 46.11	21.11
12 VINYL ACETATE				CAS #: 108-05-4	
9.58(0.742)	43	572184	81.19		100.00
18 1,1,1-TRICHLOROETHANE				CAS #: 71-55-6	
11.56(0.895)	97	814948	100.81		100.00
19 CARBON TETRACHLORIDE				CAS #: 56-23-5	
11.88(0.919)	117	794425	98.74		100.00
21 BENZENE				CAS #: 71-43-2	
12.24(0.947)	78	1410180	100.16		100.00
24 TRICHLOROETHENE				CAS #: 79-01-6	
13.42(1.038)	130	742320	99.04		100.00
25 1,2-DICHLOROPROPANE				CAS #: 78-87-5	
13.82(1.070)	63	727660	99.76		100.00
26 BROMODICHLOROMETHANE				CAS #: 75-27-4	
14.32(1.108)	83	1161322	99.34		100.00
27 2-CHLOROETHYL VINYL ETHER				CAS #: 110-75-8	
14.84(1.148)	63	405831	92.79		100.00
28 CIS-1,3-DICHLOROPROPENE				CAS #: 10061-01-5	
15.13(1.171)	75	1080181	103.07		100.00
32 TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
16.13(1.248)	75	716256	87.23		100.00
33 1,1,2-TRICHLOROETHANE				CAS #: 79-00-5	
16.48(1.275)	97	512470	90.69		100.00
36 DIBROMOCHLOROMETHANE				CAS #: 124-48-1	
17.24(1.334)	129	932167	95.12		100.00
44 BROMOFORM				CAS #: 75-25-2	
19.97(1.546)	173	659236	89.86		100.00
* 37 CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.34(1.000)	117	783711	50.00		100.00
18.34	82	485687		36.97 - 86.97	61.97
18.34	119	258326		7.96 - 57.96	32.96
\$ 29 TOLUENE-d8				CAS #: 2037-26-5	
15.64(0.853)	98	1663412	98.29		100.00
30 TOLUENE				CAS #: 108-88-3	
15.76(0.860)	92	946425	99.40		100.00

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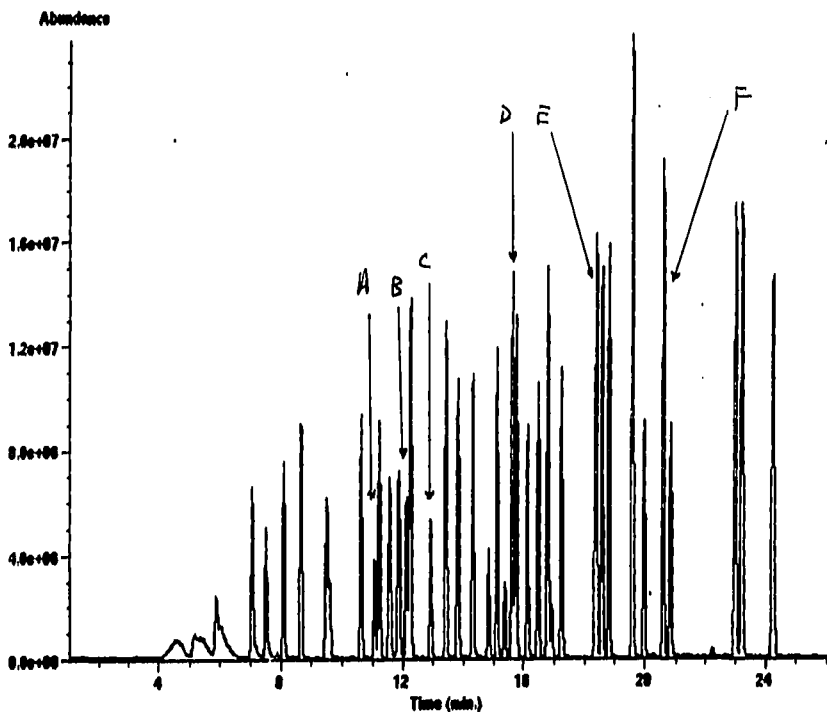
31 4-METHYL-2-PENTANONE			CAS #: 108-10-1	
15.39 (0.839) 43	213878	113.91		100.00 (M)
34 TETRACHLOROETHENE			CAS #: 127-18-4	
16.79 (0.916) 164	654525	97.23		100.00
35 2-HEXANONE			CAS #: 591-78-6	
16.92 (0.923) 43	166906	77.19		100.00
38 CHLOROBENZENE			CAS #: 108-90-7	
18.40 (1.003) 112	1391828	100.31		100.00
39 ETHYLBENZENE			CAS #: 100-41-4	
18.58 (1.013) 106	592033	99.11		100.00
40 m-, p-XYLENE			CAS #: 108-38-3	
18.80 (1.025) 106	745831	101.72		100.00
41 o-XYLENE			CAS #: 95-47-6	
19.57 (1.067) 106	673310	100.17		100.00
42 XYLENE (TOTAL)	14/9/41	200	CAS #: 1330-20-7	
19.57 (1.067) 106	673384	100.18	100-18 20-3/28/91	100.00
43 STYRENE			CAS #: 100-42-5	
19.59 (1.068) 104	1137585	97.85		100.00
\$ 45 BROMOFLUOROBENZENE			CAS #: 460-00-4	
20.58 (1.122) 95	1426061	96.39		100.00
20.58 174	922433		39.68 - 89.68	64.68
20.58 176	930419		40.24 - 90.24	65.24
46 1,1,2,2-TETRACHLOROETHANE			CAS #: 79-34-5	
20.82 (1.136) 83	773369	86.28		100.00
47 1,3-DICHLOROBENZENE			CAS #: 541-73-1	
23.21 (1.266) 146	1442547	96.94		100.00
48 1,4-DICHLOROBENZENE			CAS #: 106-46-7	
24.23 (1.321) 146	1371209	96.33		100.00
49 1,2-DICHLOROBENZENE			CAS #: 95-50-1	
24.23 (1.321) 146	1371209	96.33		100.00

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0220503
VSD50
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1 0063
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VSTD150

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IEA INC CARY, NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 20 Feb 91 02:42 Cal Operator : es
 Cal Date : 20 Feb 91 07:51 INSTRUMENT Cal File: d5.1/022091.b/022050401004.d
 Date : 20 Feb 91 02:42 Inst ID: msd5
 Data file: 022050401004.d Tune Date: 23 Jan 91 10:10
 Name Info: *15-TO 150* *3/25/91*
 Misc Info: *15-TO 150*
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method Calibration Sample, Level: 4
 Sequence index : 2
 Als bottle num : 4
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE				CAS #: 74-97-5	
11.04(1.000)	128	218777	50.00		100.00
11.04	49	432245		172.57 - 222.57	197.57
11.04	130	285560		105.52 - 155.52	130.52
1 CHLOROMETHANE				CAS #: 74-87-3	
4.41(0.400)	50	541892	163.31		100.00
2 VINYL CHLORIDE				CAS #: 75-01-4	
4.61(0.417)	62	648871	163.37		100.00
3 BROMOMETHANE				CAS #: 74-83-9	
5.15(0.466)	94	594947	137.33		100.00 (M)
4 CHLOROETHANE				CAS #: 75-00-3	
5.39(0.488)	64	408565	140.56		100.00
5 TRICHLOROFUOROMETHANE				CAS #: 75-69-4	
5.86(0.531)	101	1491124	155.58		100.00 (M)
6 1,1-DICHLOROETHENE				CAS #: 75-35-4	
7.02(0.636)	96	642255	146.48		100.00
7 ACETONE				CAS #: 67-64-1	
7.15(0.648)	43	81163	134.55		100.00
8 CARBON DISULFIDE				CAS #: 75-15-0	
7.48(0.677)	76	1750675	149.45		100.00
9 METHYLENE CHLORIDE				CAS #: 75-09-2	
8.07(0.731)	84	759284	144.35		100.00
10 trans-1,2-DICHLOROETHENE				CAS #: 540-59-0	
8.64(0.782)	96	761002	143.72		100.00
11 1,1-DICHLOROETHANE				CAS #: 75-34-3	
9.47(0.857)	63	1567485	143.96		100.00
13 cis-1,2-DICHLOROETHENE				CAS #: 156-59-2	
10.59(0.959)	96	858257	143.67		100.00
14 1,2-DICHLOROETHENE (TOTAL)				CAS #: <i>300</i>	
10.59(0.959)	96	858257 <i>1619259</i>	143.67 <i>150</i>	<i>AR309098</i>	100.00
15 2-BUTANONE				CAS #: 78-93-3	1 0065

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10.61(0.961)	72	43047	151.23		100.00
17 CHLOROFORM				CAS #: 67-66-3	
11.19(1.014)	83	1639374	144.24		100.00
\$ 20 1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.11(1.096)	65	846273	144.84		100.00
12.11	102	223324		1.38 - 51.38	26.38
22 1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.25(1.109)	62	866698	141.39		100.00
* 23 1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.91(1.000)	114	932511	50.00		100.00
12.91	63	222405		0.00 - 48.85	23.85
12.91	88	201389		0.00 - 46.59	21.59
12 VINYL ACETATE				CAS #: 108-05-4	
9.56(0.741)	43	1044909	150.73		100.00
18 1,1,1-TRICHLOROETHANE				CAS #: 71-55-6	
11.55(0.895)	97	1169819	147.11		100.00
19 CARBON TETRACHLORIDE				CAS #: 56-23-5	
11.86(0.919)	117	1168790	147.67		100.00
21 BENZENE				CAS #: 71-43-2	
12.23(0.947)	78	1972369	142.41		100.00
24 TRICHLOROETHENE				CAS #: 79-01-6	
13.40(1.038)	130	1026782	139.26		100.00
25 1,2-DICHLOROPROPANE				CAS #: 78-87-5	
13.81(1.070)	63	1044431	145.55		100.00
26 BROMODICHLOROMETHANE				CAS #: 75-27-4	
14.30(1.108)	83	1646381	143.17		100.00
27 2-CHLOROETHYL VINYL ETHER				CAS #: 110-75-8	
14.82(1.148)	63	478395	111.19		100.00 (M)
28 CIS-1,3-DICHLOROPROPENE				CAS #: 10061-01-5	
15.12(1.171)	75	1610183	156.18		100.00
32 TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
15.12(1.249)	75	1096667	135.77		100.00
33 1,1,2-TRICHLOROETHANE				CAS #: 79-00-5	
16.47(1.276)	97	788247	141.80		100.00
36 DIBROMOCHLOROMETHANE				CAS #: 124-48-1	
17.23(1.335)	129	1403208	145.55		100.00
44 BROMOFORM				CAS #: 75-25-2	
19.97(1.547)	173	1037165	143.72		100.00
* 37 CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.33(1.000)	117	746504	50.00		100.00
18.33	82	467837		37.67 - 87.67	62.67
18.33	119	245225		7.84 - 118.09	98.99
\$ 29 TOLUENE-d8				CAS #: 2037-26-5	
15.63(0.852)	90	2319591	143.89		100.00
30 TOLUENE				CAS #: 108-88-3	
15.75(0.859)	92	1334069	147.10		100.00

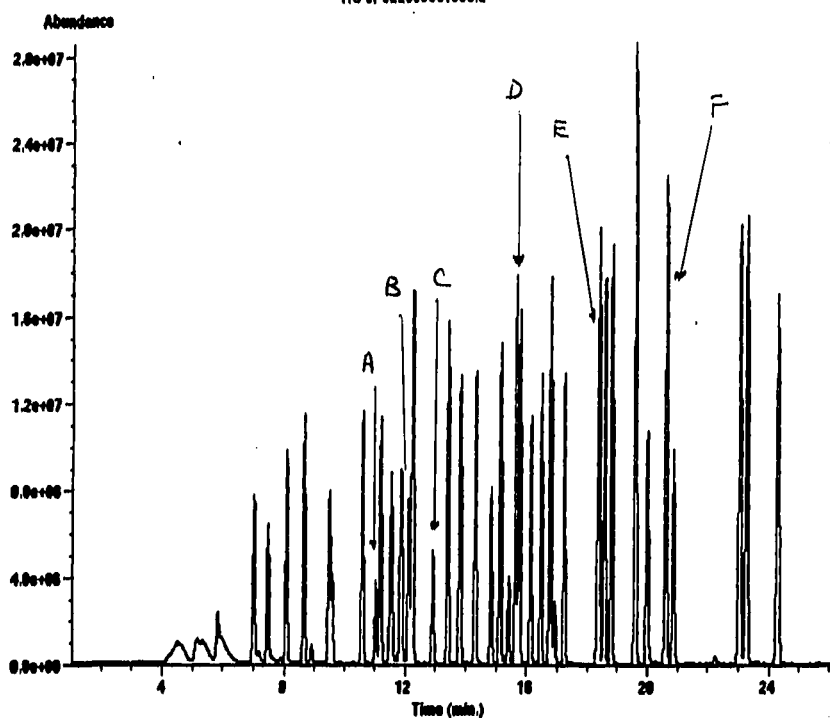
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31 4-METHYL-2-PENTANONE			CAS #: 108-10-1	
15.38 (0.839) 43	479621	200.19		100.00 (M)
34 TETRACHLOROETHENE			CAS #: 127-18-4	
16.78 (0.915) 164	897505	139.97		100.00
35 2-HEXANONE			CAS #: 591-78-6	
16.91 (0.922) 43	311162	151.07		100.00
38 CHLOROBENZENE			CAS #: 108-90-7	
18.39 (1.003) 112	1916353	144.99		100.00
39 ETHYLBENZENE			CAS #: 100-41-4	
18.57 (1.013) 106	844680	148.45		100.00
40 m-, p-XYLENE			CAS #: 108-38-3	
18.80 (1.025) 106	1004320	143.80		100.00
41 o-XYLENE			CAS #: 95-47-6	
19.56 (1.067) 106	903444	141.11		100.00
42 XYLENE (TOTAL)			CAS #: 1330-20-7	
19.56 (1.067) 106	903444	141.10	³⁰⁰ 141.10 _{8/27/91}	100.00
43 STYRENE			CAS #: 100-42-5	
19.59 (1.068) 104	1528541	138.03		100.00
\$ 45 BROMOFUOROBENZENE			CAS #: 460-00-4	
20.58 (1.123) 95	1953180	138.59		100.00
20.58 174	1214784		37.19 - 87.19	62.19
20.58 176	1226101		37.77 - 87.77	62.77
46 1,1,2,2-TETRACHLOROETHANE			CAS #: 79-34-5	
20.82 (1.136) 83	1263113	147.94		100.00
47 1,3-DICHLOROBENZENE			CAS #: 541-73-1	
23.21 (1.266) 146	1974387	139.30		100.00
48 1,4-DICHLOROBENZENE			CAS #: 106-46-7	
24.23 (1.322) 146	1911880	141.01		100.00
49 1,2-DICHLOROBENZENE			CAS #: 95-50-1	
24.23 (1.322) 146	1911880	141.01		100.00

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 VSTD 150
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VOLATILE QUANTITATION REPORT MSD5

Cal Inj Data : 20 Feb 91 03:17 Cal Operator : es
 Cal Date : 20 Feb 91 08:02 INSTRUMENT Cal File: d5.i/022091.b/022050501005.d
 Date : 20 Feb 91 03:17 Inst ID: msd5
 Data file: 022050501005.d *CKC 3/2/91* Tune Date: 23 Jan 91 10:10
 Name Info: *VSTD 200*
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voac/CP Method
 Sequence index : 3 Calibration Sample, Level: 5
 Als bottle num : 5
 Replicate num : 1

RETENTION TIME	MIN. (REL)	MASS	AREA	AMT (UG/L)	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE					CAS #: 74-97-5	
11.03 (1.000)	128	208629	50.00			100.00
11.03	49	446319			188.92 - 238.92	213.92
11.04	130	275713			107.15 - 157.15	132.15
1 CHLOROMETHANE					CAS #: 74-87-3	
4.39 (0.397)	50	704924	222.78			100.00
2 VINYL CHLORIDE					CAS #: 75-01-4	
4.61 (0.418)	62	852644	225.12			100.00
3 BROMOMETHANE					CAS #: 74-83-9	
5.14 (0.466)	94	708510	171.26			100.00 (M)
4 CHLOROETHANE					CAS #: 75-00-3	
5.36 (0.486)	64	461007	166.32			100.00
5 TRICHLOROFLUOROMETHANE					CAS #: 75-69-4	
5.82 (0.528)	101	1340620	146.68			100.00
6 1,1-DICHLOROETHENE					CAS #: 75-35-4	
6.98 (0.633)	96	770107	184.18			100.00
7 ACETONE					CAS #: 67-64-1	
7.15 (0.648)	43	122679	213.26			100.00
8 CARBON DISULFIDE					CAS #: 75-15-0	
7.43 (0.674)	76	2185186	195.62			100.00
9 METHYLENE CHLORIDE					CAS #: 75-09-2	
8.04 (0.729)	84	982948	195.96			100.00
10 trans-1,2-DICHLOROETHENE					CAS #: 540-59-0	
8.61 (0.781)	96	971305	192.36			100.00
11 1,1-DICHLOROETHANE					CAS #: 75-34-3	
9.45 (0.857)	63	2052693	197.69			100.00
13 cis-1,2-DICHLOROETHENE					CAS #: 156-59-2	
10.58 (0.959)	96	1072875	188.34			100.00
14 1,2-DICHLOROETHENE (TOTAL)					CAS #:	
10.58 (0.959)	96	1072875	188.34	<i>400</i>	<i>MS/MS</i>	<i>AR309102</i>
15 2-BUTANONE		<i>2044100</i>	<i>200</i>		CAS #: 78-93-3	1 0069

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10.61(0.962)	72	55084	202.92	100.00
17 CHLOROFORM			CAS #: 67-66-3	
11.19(1.014)	83	2020722	186.44	100.00
\$ 20 1,2-DICHLOROETHANE-D4			CAS #: 17060-07-0	
12.10(1.097)	65	1081511	194.10	100.00
12.10	102	280500	0.93 - 50.93	25.93
22 1,2-DICHLOROETHANE			CAS #: 107-06-2	
12.24(1.110)	62	1092393	186.88	100.00
* 23 1,4-DIFLUOROBENZENE			CAS #: 540-36-3	
12.90(1.000)	114	906061	50.00	100.00
12.90	63	219406	0.00 - 49.21	24.21
12.90	88	196816	0.00 - 46.72	21.72
12 VINYL ACETATE			CAS #: 108-05-4	
9.56(0.741)	43	1461334	216.95	100.00
18 1,1,1-TRICHLOROETHANE			CAS #: 71-55-6	
11.54(0.894)	97	1473457	190.70	100.00
19 CARBON TETRACHLORIDE			CAS #: 56-23-5	
11.85(0.919)	117	1467063	190.77	100.00
21 BENZENE			CAS #: 71-43-2	
12.22(0.947)	78	2532970	188.23	100.00
24 TRICHLOROETHENE			CAS #: 79-01-6	
13.40(1.038)	130	1272464	177.63	100.00
25 1,2-DICHLOROPROPANE			CAS #: 78-87-5	
13.80(1.070)	63	1365714	195.89	100.00
26 BROMODICHLOROMETHANE			CAS #: 75-27-4	
14.30(1.108)	83	2062778	184.61	100.00
27 2-CHLOROETHYL VINYL ETHER			CAS #: 110-75-8	
14.82(1.149)	63	895532	214.21	100.00
28 CIS-1,3-DICHLOROPROPENE			CAS #: 10061-01-5	
15.11(1.172)	75	2021486	201.80	100.00
32 TRANS-1,3-DICHLOROPROPENE			CAS #: 10061-02-6	
16.12(1.250)	75	1380702	175.93	100.00
33 1,1,2-TRICHLOROETHANE			CAS #: 79-00-5	
16.47(1.277)	97	1009007	186.82	100.00
36 DIBROMOCHLOROMETHANE			CAS #: 124-48-1	
17.22(1.335)	129	1701476	181.64	100.00
44 BROMOFORM			CAS #: 75-25-2	
19.97(1.548)	173	1309406	186.75	100.00
* 37 CHLOROBENZENE-d5			CAS #: 3114-55-4	
18.32(1.000)	117	726087	50.00	100.00
18.32	82	476612	40.64 - 90.64	65.64
18.32	119	243045	8.47 - 58.47	33.47
\$ 29 TOLUENE-d8			CAS #: 2037-26	AR 309100
15.62(0.853)	98	2839840	181.12	100.00
30 TOLUENE			CAS #: 108-88-3	1
15.75(0.860)	92	1696553	192.33	100.00

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31 4-METHYL-2-PENTANONE			CAS #: 108-10-1	
15.38(0.840) 43	666851	224.80		100.00 (M)
34 TETRACHLOROETHENE			CAS #: 127-18-4	
16.77(0.915) 164	1055461	169.23		100.00
35 2-HEXANONE			CAS #: 591-78-6	
16.90(0.922) 43	447039	223.15		100.00
38 CHLOROBENZENE			CAS #: 108-90-7	
18.38(1.003) 112	2403916	187.00		100.00
39 ETHYLBENZENE			CAS #: 100-41-4	
18.56(1.013) 106	1076747	194.55		100.00
40 m-, p-XYLENE			CAS #: 108-38-3	
18.79(1.025) 106	1298754	191.19		100.00
41 o-XYLENE			CAS #: 95-47-6	
19.55(1.067) 106	1141101	183.24		100.00
42 XYLENE (TOTAL)	2437755	400	CAS #: 1330-20-7	
19.55(1.067) 106	1141101	183.23	04/25/91	100.00
43 STYRENE			CAS #: 100-42-5	
19.58(1.069) 104	1890491	175.52		100.00
\$ 45 BROMOFLUOROBENZENE			CAS #: 460-00-4	
20.58(1.123) 95	2420866	176.61		100.00
20.58	174		36.15 - 86.15	61.15
20.58	176		37.17 - 87.17	62.17
46 1,1,2,2-TETRACHLOROETHANE			CAS #: 79-34-5	
20.83(1.137) 83	1662956	200.25		100.00
47 1,3-DICHLOROBENZENE			CAS #: 541-73-1	
23.23(1.268) 146	2467573	178.99		100.00
48 1,4-DICHLOROBENZENE			CAS #: 106-46-7	
24.25(1.324) 146	2388404	181.10		100.00
49 1,2-DICHLOROBENZENE			CAS #: 95-50-1	
24.25(1.324) 146	2388404	181.10		100.00

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7A
VOLATILE CONTINUING CALIBRATION CHECK

Lab Name: IEA Contract: 68-W8-0045
Lab Code: IEA Case No.: 868-6 SAS No.: SDG No.: BC001
Instrument ID: msd5 Calibration Date: 3/ 1/91 Time: 21:24
Lab File ID: 0301E01 Init. Calib. Date(s): 2/20/91 2/20/91
Matrix:(soil/water) WATER Level:(low/med): LOW Column:(pack/cap) CAP
Min RRF50 for SPCC(#) = .300 (0.250 for Bromoform) Max %D for CCC(*) = 25.0

COMPOUND	RRF	RRF50	%D
CHLOROMETHANE	.842	.672	20.2
BROMOMETHANE	.991	.540	45.5
VINYL CHLORIDE	1.032	.889	13.9
CHLOROETHANE	.664	.692	4.2
METHYLENE CHLORIDE	1.202	1.329	10.5
ACETONE	.138	.177	28.1
CARBON DISULFIDE	2.677	2.668	.4
1,1-DICHLOROETHENE	1.002	1.031	2.9
1,1-DICHLOROETHANE	2.488	2.730	9.7
1,2-DICHLOROETHENE (TOTAL)	1.288	1.437	11.6
CHLOROFORM	2.598	2.982	14.8
1,2-DICHLOROETHANE	1.401	1.716	22.5
2-BUTANONE	.065	.092	42.0
1,1,1-TRICHLOROETHANE	.426	.477	12.0
CARBON TETRACHLORIDE	.424	.476	12.3
VINYL ACETATE	.372	.433	16.4
BROMODICHLOROMETHANE	.617	.715	15.9
1,2-DICHLOROPROPANE	.385	.421	9.3
CIS-1,3-DICHLOROPROPENE	.553	.608	10.0
TRICHLOROETHENE	.395	.435	10.0
DIBROMOCHLOROMETHANE	.517	.612	18.5
1,1,2-TRICHLOROETHANE	.298	.343	14.9
BENZENE	.743	.822	10.6
TRANS-1,3-DICHLOROPROPENE	.433	.466	7.7
BROMOFORM	.387	.465	20.1
4-METHYL-2-PENTANONE	.204	.250	22.5
2-HEXANONE	.138	.168	21.9
TETRACHLOROETHENE	.429	.473	10.1
1,1,2,2-TETRACHLOROETHANE	.572	.642	12.3
TOLUENE	.607	.618	1.8
CHLOROBENZENE	.885	.901	1.7
ETHYLBENZENE	.381	.384	.8
STYRENE	.742	.789	6.4
XYLENE (TOTAL)	.448	.466	3.9
TOLUENE-d8	1.080	1.057	2.1
BROMOFLUOROBENZENE	.944	1.002	6.2
1,2-DICHLOROETHANE-D4	1.335	1.514	13.4

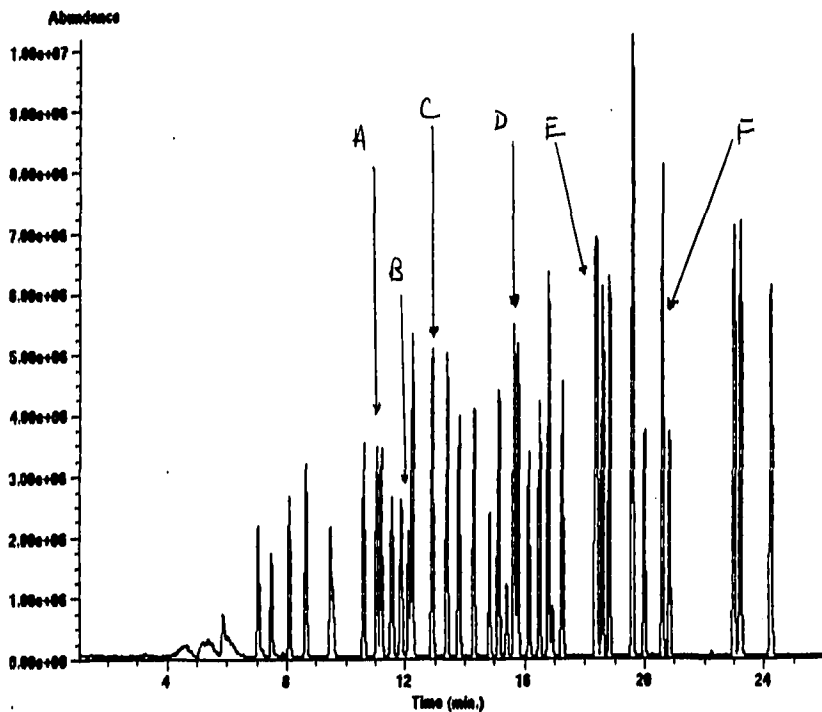
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FORM VII VOA

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VSTD54
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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/02195.
 Date : 01 Mar 91 21:24 Inst ID: msd5
 Data file: 0301e01.d Tune Date: 23 Jan 91 10:10
 Name Info: vstd50 iea msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME	MIN. (REL)	MASS	AREA	AMT (UG/L)	TARGET RANGE	RATIO
* 16	BROMOCHLOROMETHANE				CAS #: 74-97-5	
11.04(1.000)	128	192811	50.00			100.00
11.04	49	397189			160.78 - 210.78	205.99
11.05	130	254254			106.54 - 156.54	131.86
1	CHLOROMETHANE				CAS #: 74-87-3	
4.42(0.400)	50	129570	50.00			100.00
2	VINYL CHLORIDE				CAS #: 75-01-4	
4.64(0.420)	62	171377	50.00			100.00
3	BROMOMETHANE				CAS #: 74-83-9	
5.16(0.468)	94	104211	50.00			100.00
4	CHLOROETHANE				CAS #: 75-00-3	
5.37(0.486)	64	133421	50.00			100.00
5	TRICHLOROFLUOROMETHANE				CAS #: 75-69-4	
6.00(0.543)	101	231786	50.00			100.00
6	1,1-DICHLOROETHENE				CAS #: 75-35-4	
7.04(0.638)	96	198799	50.00			100.00
7	ACETONE				CAS #: 67-64-1	
7.16(0.648)	43	34054	50.00			100.00
8	CARBON DISULFIDE				CAS #: 75-15-0	
7.49(0.678)	76	514376	50.00			100.00
9	METHYLENE CHLORIDE				CAS #: 75-09-2	
8.08(0.732)	84	256215	50.00			100.00
10	trans-1,2-DICHLOROETHENE				CAS #: 540-59-0	
8.65(0.783)	96	258178	50.00			100.00
11	1,1-DICHLOROETHANE				CAS #: 75-34-3	
9.48(0.858)	63	526365	50.00			100.00
13	cis-1,2-DICHLOROETHENE				CAS #: 156-59-2	
10.60(0.960)	96	296121	50.00			100.00
14	1,2-DICHLOROETHENE (TOTAL)				CAS #: AR309107	100.00
10.60(0.960)	96	206121	50.00			100.00
15	2-BUTANONE				CAS #: 78-93-3	1 0074

If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

10.61(0.961)	72	17815	50.00		100.00
17 CHLOROFORM				CAS #: 67-66-3	
11.19(1.014)	83	575049	50.00		100.00
\$ 20 1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.11(1.096)	65	292000	50.00		100.00
12.11	102	71558		1.65 - 51.65	24.50
22 1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.24(1.109)	62	330788	50.00		100.00
* 23 1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.91(1.000)	114	864705	50.00		100.00
12.90	63	215262		0.00 - 47.22	24.89
12.91	88	191796		0.00 - 45.47	22.18
12 VINYL ACETATE				CAS #: 108-05-4	
9.57(0.742)	43	374137	50.00		100.00
18 1,1,1-TRICHLOROETHANE				CAS #: 71-55-6	
11.54(0.894)	97	412762	50.00		100.00
19 CARBON TETRACHLORIDE				CAS #: 56-23-5	
11.86(0.919)	117	411981	50.00		100.00
21 BENZENE				CAS #: 71-43-2	
12.23(0.947)	78	710410	50.00		100.00
24 TRICHLOROETHENE				CAS #: 79-01-6	
13.41(1.039)	130	376067	50.00		100.00
25 1,2-DICHLOROPROPANE				CAS #: 78-87-5	
13.81(1.070)	63	363687	50.00		100.00
26 BROMODICHLOROMETHANE				CAS #: 75-27-4	
14.31(1.108)	83	618016	50.00		100.00
27 2-CHLOROETHYL VINYL ETHER				CAS #: 110-75-8	
14.82(1.148)	63	244690	50.00		100.00
28 CIS-1,3-DICHLOROPROPENE				CAS #: 10061-01-5	
15.12(1.171)	75	557362	53.00		100.00
32 TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
16.13(1.249)	75	370894	46.00		100.00
33 1,1,2-TRICHLOROETHANE				CAS #: 79-00-5	
16.48(1.277)	97	296241	50.00		100.00
36 DIBROMOCHLOROMETHANE				CAS #: 124-48-1	
17.24(1.336)	129	529617	50.00		100.00
44 BROMOFORM				CAS #: 75-25-2	
19.97(1.547)	173	401870	50.00		100.00
* 37 CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.33(1.000)	117	767497	50.00		100.00
18.33	82	486897		37.16 - 87.16	63.43
18.33	119	251673		7.90 - 57.90	32.79
29 4-METHYL-2-PENTANONE				CAS #: 108-10-1	
15.38(0.839)	43	192107	50.00	AR309108	100.00 (M)
\$ 30 TOLUENE-d8				CAS #: 2037-26-5	1 0075
15.63(0.852)	98	811237	50.00		100.00

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msds

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31 TOLUENE			CAS #: 108-88-3	
15.75(0.859)	92	474642	50.00	100.00
34 TETRACHLOROETHENE			CAS #: 127-18-4	
16.78(0.915)	164	362867	50.00	100.00
35 2-HEXANONE			CAS #: 591-78-6	
16.91(0.922)	43	129036	50.00	100.00
38 CHLOROBENZENE			CAS #: 108-90-7	
18.39(1.003)	112	691169	50.00	100.00
39 ETHYLBENZENE			CAS #: 100-41-4	
18.57(1.013)	106	294730	50.00	100.00
40 m-, p-XYLENE			CAS #: 108-38-3	
18.80(1.025)	106	365972	50.00	100.00
41 o-XYLENE			CAS #: 95-47-6	
19.56(1.067)	106	348954	50.00	100.00
42 XYLENE (TOTAL)		714926	100	CAS #: 1330-20-7
19.56(1.067)	106	348954	50.00	100.00
43 STYRENE			CAS #: 100-42-5	
19.58(1.068)	104	605695	50.00	100.00
\$ 45 BROMOFLUOROBENZENE			CAS #: 460-00-4	
20.57(1.122)	95	769032	50.00	100.00
20.58	174	495033		41.39 - 91.39 64.37
20.58	176	487000		41.62 - 91.62 63.32
46 1,1,2,2-TETRACHLOROETHANE			CAS #: 79-34-5	
20.82(1.135)	83	492999	50.00	100.00
47 1,3-DICHLOROBENZENE			CAS #: 541-73-1	
23.19(1.265)	146	792560	50.00	100.00
48 1,4-DICHLOROBENZENE			CAS #: 106-46-7	
24.21(1.320)	146	754777	50.00	100.00
49 1,2-DICHLOROBENZENE			CAS #: 95-50-1	
24.21(1.320)	146	754835	50.00	100.00

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4/2/91

0301E01
VSTD50
MSD5

AR309109
1 0076

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8A
VOLATILE INTERNAL STANDARD AREA SUMMARY

Lab Name: IEA Contract: 68-W8-0045
 Lab Code: IEA Case No.: 868-6 SAS No.: SDG No.: BC001
 Lab File ID (Standard): 0301E01 Date Analyzed: 3/ 1/91
 Instrument ID: msd5 Time Analyzed: 21:24
 Matrix:(soil/water) WATER Level:(low/med): LOW Column:(pack/cap) CAP

	IS1(BCM) AREA #	RT	IS2(DFB) AREA #	RT	IS3(CBZ) AREA #	RT
=====	=====	=====	=====	=====	=====	=====
12 HOUR STD	192811.	11.04	864705.	12.91	767497.	18.33
=====	=====	=====	=====	=====	=====	=====
UPPER LIMIT	385622.	11.54	1729410.	13.41	1534994.	18.83
=====	=====	=====	=====	=====	=====	=====
LOWER LIMIT	96406.	10.54	432353.	12.41	383749.	17.83
=====	=====	=====	=====	=====	=====	=====
EPA SAMPLE NO.						
=====	=====	=====	=====	=====	=====	=====
1 VBLK01	191178.	11.05	854111.	12.91	765215.	18.32
2 BC006	188186.	11.06	820858.	12.91	726115.	18.33
3 BC004	191897.	11.06	852931.	12.92	756641.	18.33
4 BC008	199142.	11.06	855639.	12.92	745576.	18.33
5 BC010	189489.	11.05	832828.	12.91	678005.	18.33
6 BC001	182518.	11.06	793181.	12.92	655112.	18.33
7 BC001MS	191500.	11.07	837886.	12.93	740878.	18.34
8 BC001MSD	201213.	11.10	874366.	12.96	781565.	18.37
9 BC002	129449.	11.07	519782.	12.92	455633.	18.34
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						

IS1 (BCM) = BROMOCHLOROMETHANE
 IS2 (DFB) = 1,4-DIFLUOROBENZENE
 IS3 (CBZ) = CHLOROBENZENE-d5

UPPER LIMIT = + 100%
 of internal standard area.
 LOWER LIMIT = - 50%
 of internal standard area.

Column used to flag internal standard area values with an asterisk

page 1 of 1

FORM VIII VOA

AR309110

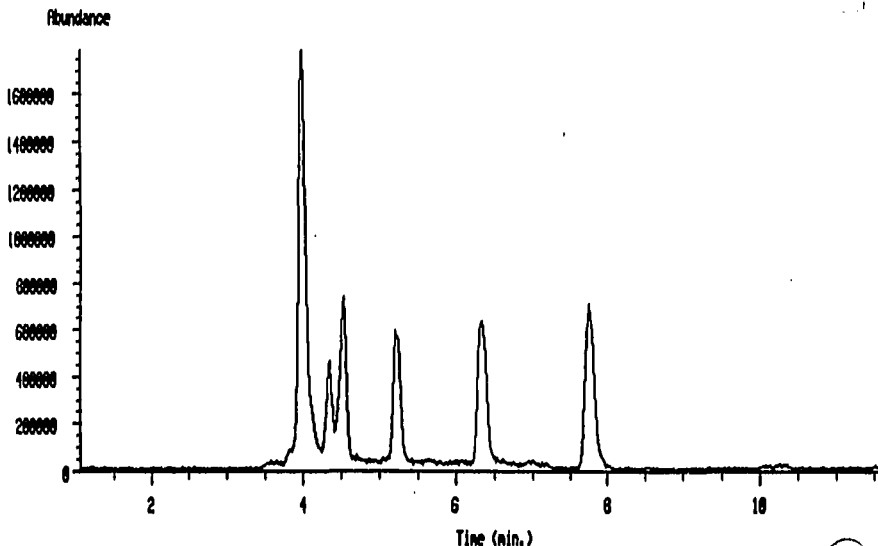
1 0077

1/87 Rev.

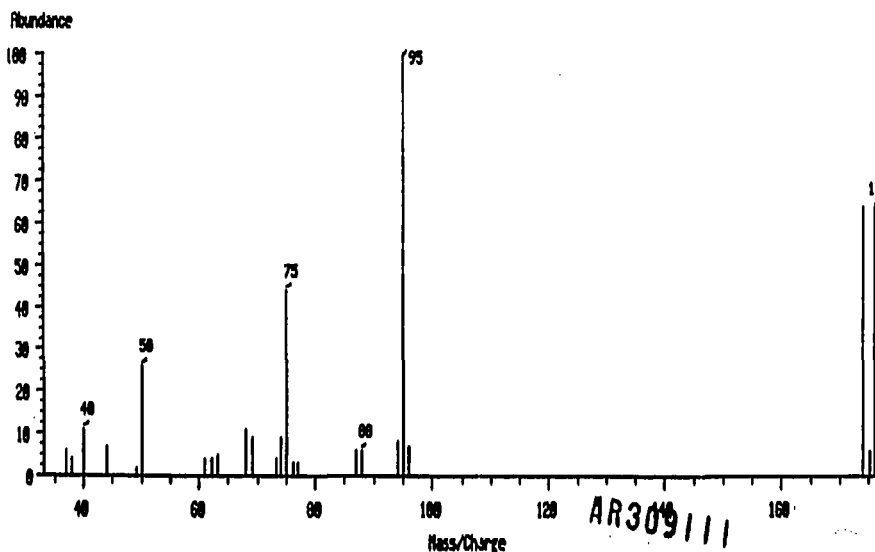
If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

Name Info: 50 ng bfb lea msd5
Misc Info:
Date : Wed Feb 20 91 12:11:04 AM

TIC of 022856.d



Scan 414 (7.672 min) of 022856.d SCALED



AR309111

1 0078

If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

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50 ng bfb iea msd5

Modified: scaled

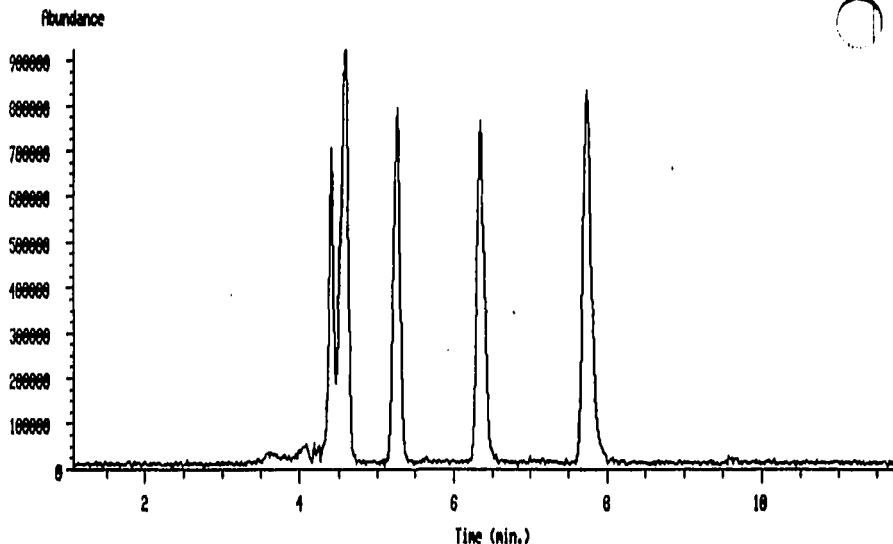
m/z	abund.	m/z	abund.	m/z	abund.	m/z	abund.
37.00	6	62.10	4	75.05	44	95.05	100
38.10	4	63.10	5	76.25	3	96.05	7
40.00	11	67.90	11	77.05	3	173.95	64
44.00	7	69.00	9	86.95	6	175.05	6
49.10	2	73.25	4	87.85	6	175.95	64
50.10	26	74.05	9	94.05	8	176.95	6
60.90	4						

AR3099929

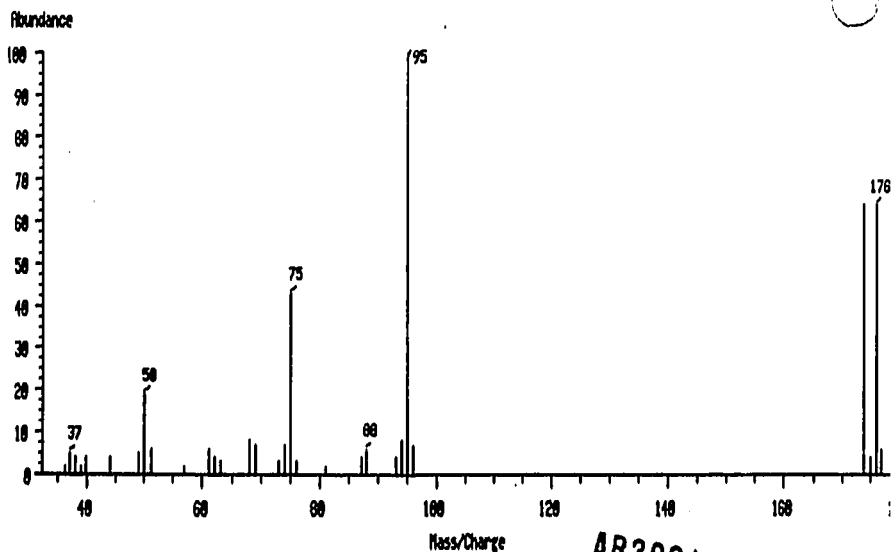
If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

Name Info: 50 ng bfb iea msd5
Misc Info:
Date : Fri Mar 01 91 08:50:41 PM

TIC of 8381ebf.d



Scan 418 (7.756 min) of 8381ebf.d SCALED



AR309113 0050

If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

Scan 418 (7.756 min) of 0301ebf.d

50 ng bfb lea msd5

Modified: scaled

m/z	abund.	m/z	abund.	m/z	abund.	m/z	abund.
36.20	2	51.10	6	74.05	7	94.05	8
37.10	5	57.00	2	75.05	43	95.05	100
38.00	4	61.10	6	75.95	3	96.05	7
39.10	2	62.00	4	80.95	2	173.85	64
39.90	4	63.00	3	87.05	4	174.95	4
44.00	4	68.00	8	87.85	6	175.95	64
49.00	5	69.00	7	93.05	4	176.85	6
50.00	19	72.95	3				

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1AR989114

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

VBLK01

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E02

Level: (low/med) LOW

Date Received: 0/ 0/ 0

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE	5.	U
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	5.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE	5.	U
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE	5.	U
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE	5.	U
108-90-7-----	CHLOROBENZENE	5.	U
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

FORM I VOA

AR809715

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

VBLK01

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SD3 No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E02

Level: (low/med) LOW

Date Received: 0/ 0/ 0

% Moisture: not dec. 100.

Date Analyzed: 3/ 1/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

Number TICs found: 0

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q
1.				
2.				
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FORM I VOA-TIC

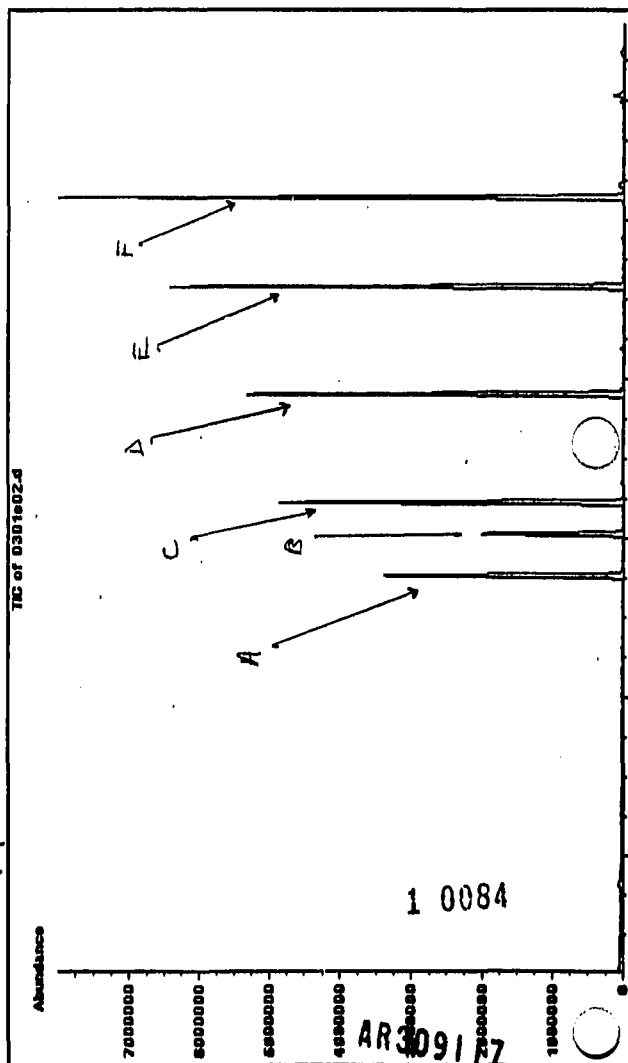
1/87 Rev.

AR309196

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clm
3/25/91

VB1K91
3/1/91 22:01 MSDS



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IEA INC CARY, NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.i/021991.b/0219505.d
 Date : 01 Mar 91 22:01 Inst ID: msd5
 Data file: 0301e02.d Tune Date: 23 Jan 91 10:10
 Name Info: ~~vik~~ lea msd5 VBLKQ1
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16 BROMOCHLOROMETHANE CAS #: 74-97-5						
11.05(1.000)	128	191178	50.00			100.00
11.05	49	398816			160.78 - 210.78	208.60
11.05	130	252171			106.54 - 156.54	131.90
\$ 20 1,2-DICHLOROETHANE-D4 CAS #: 17060-07-0						
12.11(1.097)	65	289764	50.04	100.08		100.00
12.11	102	71236			1.65 - 51.65	24.58
* 23 1,4-DIFLUOROBENZENE CAS #: 540-36-3						
12.91(1.000)	114	854111	50.00			100.00
12.91	63	211487			0.00 - 47.22	24.76
12.91	88	194166			0.00 - 45.47	22.73
* 37 CHLOROBENZENE-d5 CAS #: 3114-55-4						
18.32(1.000)	117	765215	50.00			100.00
18.32	82	494267			37.16 - 87.16	64.59
18.32	119	247048			7.90 - 57.90	32.28
\$ 30 TOLUENE-d8 CAS #: 2037-26-5						
15.62(0.852)	98	799714	49.44	98.87		100.00
31 TOLUENE CAS #: 108-88-3						
15.75(0.859)	92	3155	0.39	B-L		100.00
\$ 45 BROMOFLUOROBENZENE CAS #: 460-00-4						
20.57(1.122)	95	757623	49.41	98.81		100.00
20.57	174	487557			41.39 - 91.39	64.35
20.57	176	484367			41.62 - 91.62	63.93
47 1,3-DICHLOROBENZENE CAS #: 541-73-1						
23.19(1.265)	146	6220	0.32	NT		100.00
48 1,4-DICHLOROBENZENE CAS #: 106-46-7						
24.20(1.320)	146	6024	0.40	NT		100.00
49 1,2-DICHLOROBENZENE CAS #: 95-50-1						
24.20(1.320)	146	6024	0.40	NT		100.00

AR309118
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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC001MS

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301R10

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO.	COMPOUND	CONCENTRATION UNITS: (ug/L or ug/Kg) UG/L	Q
---------	----------	--	---

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE		
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	2-BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	5.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE		
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE		
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE		
108-90-7-----	CHLOROBENZENE		
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

FORM I VOA

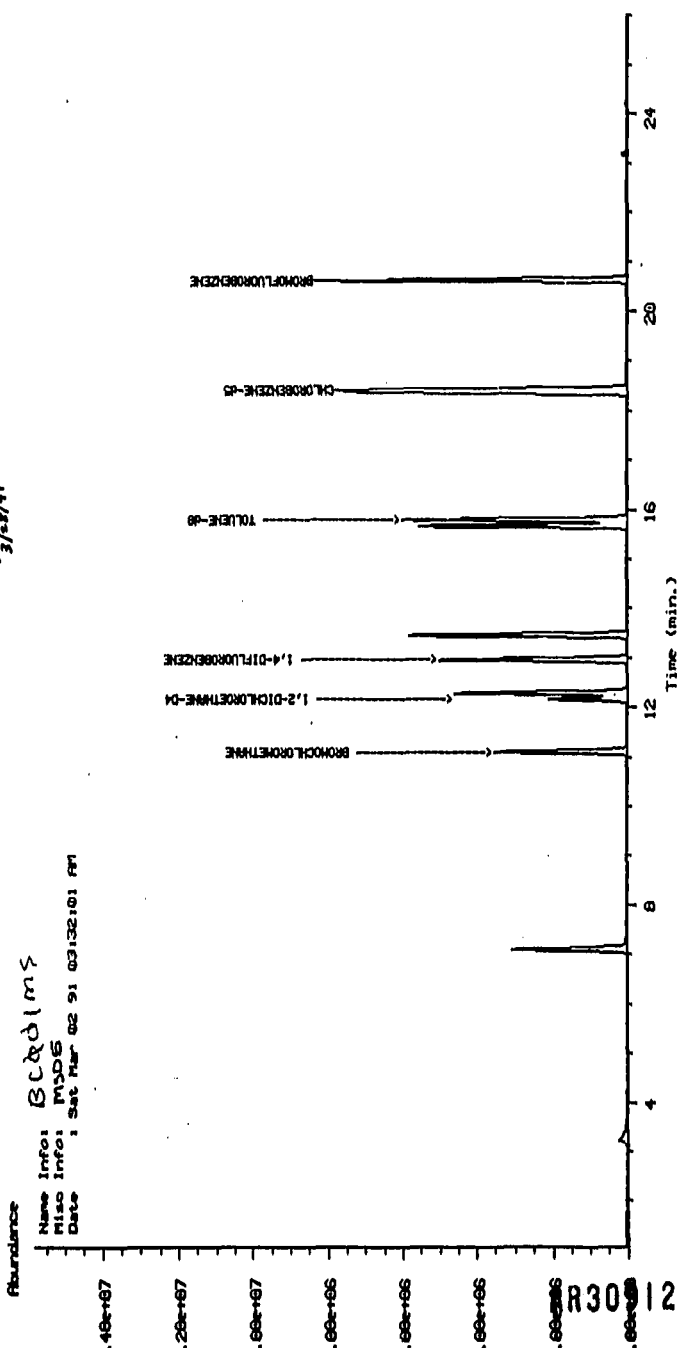
AR30911986

1/87 Rev

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TIC of 0301e1001e10-d-
3/2/91

Name Info: BCD1m3
Misc Info: M305
Date: Sat Mar 02 91 03:32:01 PM



R3001201 0087

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IEA INC CARY, NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29
Cal Date : 1 Mar 91 23:03 chemist

Cal Operator : es

Date : 02 Mar 91 03:32

Cal File: em/msd5.1/021991.b/02195

Data file: 0301e10e1010.d

Inst ID: msd5

Tune Date: 23 Jan 91 10:10

Name Info:

Misc Info:

Operator : es

Dilution Factor: 1.000

Comment : voaCLP Method

Sequence index :

Als bottle num :

Replicate num :

1

10

1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT (UG/L)	%REC	TARGET RANGE	RATIO
* 16	BROMOCHLOROMETHANE				CAS #: 74-97-5	
11.10 (1.000)	128	201213	50.00			100.00
11.10	49	408103			160.78 - 210.78	202.82
11.10	130	265350			106.54 - 156.54	131.87
6	1,1-DICHLOROETHENE				CAS #: 75-35-4	
7.09 (0.642)	96	296042	71.35	143%		100.00
8	CARBON DISULFIDE				CAS #: 75-15-0	
7.08 (0.641)	76	432	0.04	BQL		100.00
\$ 20	1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.16 (1.101)	65	301539	49.48	98.96		100.00
12.16	102	74608			1.65 - 51.65	24.74
22	1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.30 (1.113)	62	9031	1.31	BQL		100.00
* 23	1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.96 (1.000)	114	874366	50.00			100.00
12.96	63	211230			0.00 - 47.22	24.15
12.96	88	189794			0.00 - 45.47	21.70
21	BENZENE				CAS #: 71-43-2	
12.28 (0.952)	78	875778	60.96	122%		100.00
24	TRICHLOROETHENE				CAS #: 79-01-6	
13.46 (1.043)	130	449788	59.14	118%		100.00
32	TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
15.79 (1.224)	75	5632	0.69	BQL		100.00
* 37	CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.37 (1.000)	117	781565	50.00			100.00
18.37	82	504453			37.16 - 87.16	64.28
18.37	119	258660			7.90 - 57.90	33.09
29	4-METHYL-2-PENTANONE				CAS #: 108-10-1	
15.80 (0.862)	43	6966	1.78	BQL		100.00
\$ 30	TOLUENE-d8				CAS #: 2037-26-5	
15.67 (0.855)	98	834130	50.49	100.97		100.00
31	TOLUENE				CAS #: 108-88-3	
15.79 (0.861)	92	571994	59.17	118%		100.00

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38 CHLOROBENZENE			CAS #: 108-90-7	
18.42 (1.005)	112	794188	56.42	100.00
			1137	
\$ 45 BROMOFLUOROBENZENE			CAS #: 460-00-4	
20.61 (1.124)	95	780641	49.84	100.00
20.61	174	516507		41.39 - 91.39 66.16
20.61	176	516593		41.62 - 91.62 66.17

TENTATIVELY IDENTIFIED COMPOUNDS

RET TIME	AREA	AMT(UG/L)	QUALITY	LIBRARY	LIB ENTRY
TOLUOL, PERDEUTERO-			CAS #:		
15.67	221854600	52.58	52	WILEY.1	1352

Q331E10
BC001MS
MSD5

ckr
3/25/91

AR309122 0089

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1A
VOLATILE ORGANICS ANALYSIS DATA SHEET

EPA SAMPLE NO.

BC001MSD

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E09

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CAS NO. COMPOUND CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L Q

74-87-3-----	CHLOROMETHANE	10.	U
74-83-9-----	BROMOMETHANE	10.	U
75-01-4-----	VINYL CHLORIDE	10.	U
75-00-3-----	CHLOROETHANE	10.	U
75-09-2-----	METHYLENE CHLORIDE	5.	U
67-64-1-----	ACETONE	10.	U
75-15-0-----	CARBON DISULFIDE	5.	U
75-35-4-----	1,1-DICHLOROETHENE		
75-34-3-----	1,1-DICHLOROETHANE	5.	U
540-59-0-----	1,2-DICHLOROETHENE (TOTAL)	5.	U
67-66-3-----	CHLOROFORM	5.	U
107-06-2-----	1,2-DICHLOROETHANE	5.	U
78-93-3-----	BUTANONE	10.	U
71-55-6-----	1,1,1-TRICHLOROETHANE	5.	U
56-23-5-----	CARBON TETRACHLORIDE	5.	U
108-05-4-----	VINYL ACETATE	10.	U
75-27-4-----	BROMODICHLOROMETHANE	5.	U
78-87-5-----	1,2-DICHLOROPROPANE	5.	U
10061-01-5-----	CIS-1,3-DICHLOROPROPENE	5.	U
79-01-6-----	TRICHLOROETHENE		
124-48-1-----	DIBROMOCHLOROMETHANE	5.	U
79-00-5-----	1,1,2-TRICHLOROETHANE	5.	U
71-43-2-----	BENZENE		
10061-02-6-----	TRANS-1,3-DICHLOROPROPENE	5.	U
75-25-2-----	BROMOFORM	5.	U
108-10-1-----	4-METHYL-2-PENTANONE	10.	U
591-78-6-----	2-HEXANONE	10.	U
127-18-4-----	TETRACHLOROETHENE	5.	U
79-34-5-----	1,1,2,2-TETRACHLOROETHANE	5.	U
108-88-3-----	TOLUENE		
108-90-7-----	CHLOROBENZENE		
100-41-4-----	ETHYLBENZENE	5.	U
100-42-5-----	STYRENE	5.	U
1330-20-7-----	XYLENE (TOTAL)	5.	U

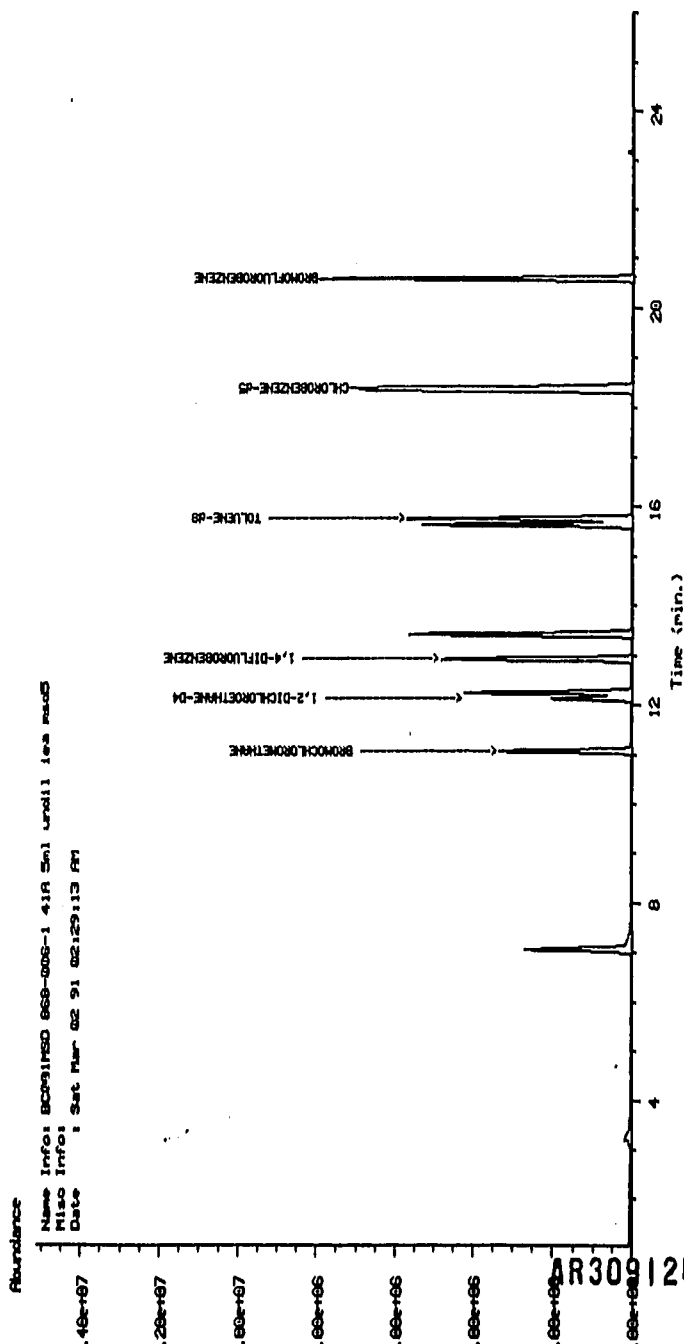
AR309 1630

FORM I VOA

1/87 Rev

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TIC of 0301e09.d



AR3091241 0091

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IEA INC CARY,NC

VOLATILE QUANTITATION REPORT MSD5

Cal Inj Date : 19 Feb 91 16:29 Cal Operator : es
 Cal Date : 1 Mar 91 23:03 chemist Cal File: em/msd5.1/021991.b/021950
 Date : 02 Mar 91 02:29 Inst ID: msd5
 Data file: 0301a09.d Tune Date: 23 Jan 91 10:10
 Name Info: BC001MSD 868-006-1 41A 5ml undil ie a msd5
 Misc Info:
 Operator : es Dilution Factor: 1.000
 Comment : voaCLP Method
 Sequence index : 0
 Als bottle num : 0
 Replicate num : 1

RETENTION TIME MIN. (REL)	MASS	AREA	AMT(UG/L)	%REC	TARGET RANGE	RATIO
* 16	BROMOCHLOROMETHANE				CAS #: 74-97-5	
11.07(1.000)	128	191500	50.00			100.00
11.06	49	390607			160.78 - 210.78	203.97
11.07	130	254728			106.54 - 156.54	133.01
6	1,1-DICHLOROETHENE				CAS #: 75-35-4	
7.06(0.639)	96	284927	72.15	144%		100.00
\$ 20	1,2-DICHLOROETHANE-D4				CAS #: 17060-07-0	
12.13(1.098)	65	288205	49.69	99.38		100.00
12.13	102	70941			1.65 - 51.65	24.61
22	1,2-DICHLOROETHANE				CAS #: 107-06-2	
12.24(1.108)	62	9579	1.46	BOL		100.00
* 23	1,4-DIFLUOROBENZENE				CAS #: 540-36-3	
12.93(1.000)	114	837886	50.00			100.00
12.92	63	204942			0.00 - 47.22	24.45
12.93	88	187755			0.00 - 45.47	22.40
21	BENZENE				CAS #: 71-43-2	
12.25(0.949)	78	812230	59.00	113%		100.00
24	TRICHLOROETHENE				CAS #: 79-01-6	
13.42(1.040)	130	436419	59.88	120%		100.00
32	TRANS-1,3-DICHLOROPROPENE				CAS #: 10061-02-6	
15.76(1.221)	75	5123	0.86	BOL		100.00
* 37	CHLOROBENZENE-d5				CAS #: 3114-55-4	
18.34(1.000)	117	740878	50.00			100.00
18.34	82	477550			37.16 - 87.16	64.45
18.34	119	241609			7.90 - 57.90	32.61
29	4-METHYL-2-PENTANONE				CAS #: 108-10-1	
15.75(0.859)	43	9429	2.54	BOL		100.00
\$ 30	TOLUENE-d8				CAS #: 2037-26-5	
15.64(0.853)	98	797716	50.93	101.87		100.00
31	TOLUENE				CAS #: 108-88-3	
15.76(0.860)	92	535756	58.47	111%		100.00
38	CHLOROBENZENE				CAS #: 108-90-7	
18.40(1.004)	112	764153	57.27	115%		100.00

1 0092

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\$ 45 BROMOFLUOROBENZENE

CAS #: 460-00-4

20.59 (1.123)	95	753013	50.72 101.44	100.00
20.59	174	483734	41.39 - 91.39	64.23
20.59	176	484182	41.62 - 91.62	64.29

TENTATIVELY IDENTIFIED COMPOUNDS

RET TIME	AREA	AMT (UG/L)	QUALITY	LIBRARY	LIB ENTRY
15.64	211638900	52.03	78	WILEY.1	1352

BCD 1 MSD
 0301E09
 MSD5

CLL
 3/25/91

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AR309127

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IEA, INC.
CASE 868-006
SDG BC_001
METALS DATA

1 OF 1

AR309128

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Industrial & Environmental Analysts, Inc.

P.O. Box 12848
Research Triangle Park, North Carolina 27709
(919) 677-0090
FAX (919) 677-0427

CASE: 868-006

SDG No: BC_001

SAMPLES: Twenty-six (26) filtered and un-filtered Water Samples for the following analyses: Four Samples for manganese, seven samples for manganese and zinc, six samples for mercury, six samples for inorganic mercury, and three samples for mercury and manganese.

This case was closed on February 21, 1991. Samples BC_013 and BC_025 were received in broken bottles. Samples BC_018 and BC_021 were received without lids. Although samples had leaked out into the cooler during transition, the client requested that the analyses proceed as normal. The remaining samples were received intact. Upon receipt at IEA, each sample was assigned a 6-character "EPA" sample number for simplicity in forms generation. This package makes reference only to those 6-character IDs. Please consult the IEA Assigned Number Index for sample cross-identification.

On samples analyzed for mercury only, the color after and clarity after are noted as not applicable (NA) on Forms I. Due to procedural requirements, the samples are normally purple following mercury digestion. No significant changes occurred during sample preparation.

Due to Plasma2 instrument problems during the first quarter (January - March), the Linear Range has not been determined for manganese and zinc. The Instrument Detection Limits (IDL), however, were determined. The IDL for the inorganic form of mercury has not been determined.

Sample BC_006 was initially prepared on 03/05/91. Software limitations would not allow this preparation to be recorded on Form XIII.

The remaining analyses proceeded with no notable difficulties.


Anna Feather
Assistant CLP Manager
April 1, 1991

AR309129

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IEA Assigned Number Index

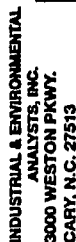
Inorganic Analyses

Case No. 868-006

IEA Assigned "EPA" Number	BCM Sample ID
BC_002	26AU
BC_003	26AF
BC_004	33ADU
BC_005	33ADF
BC_006	33AU
BC_007	33AF
BC_008	EQUIPMENT BLANK U (02/19/91)
BC_009	EQUIPMENT BLANK F (02/19/91)
BC_010	TRIP BLANK U (02/19/91)
BC_011	39AU
BC_012	39AF
BC_013	9ADU
BC_014	9ADF
BC_015	9ADIU
BC_016	9ADIF
BC_017	9BU
BC_018	9BF
BC_019	9BIU
BC_020	9BIF
BC_021	9AU
BC_022	9AF
BC_023	9AIU
BC_024	9AIF
BC_025	EQUIPMENT BLANK U (02/20/91)
BC_026	EQUIPMENT BLANK F (02/20/91)
BC_027	TRIP BLANK U (02/20/91)

AR309130

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CHAIN OF CUSTODY RECORD

REGULATORY CLASSIFICATION - PLEASE SPECIFY

☐ NPDES ☐ DRINKING WATER ☐ RCRA ☒ OTHER CERCLA / SARA

NO: 12133

[illegible]

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ENVIROFORMS/CLP 788

COVER PAGE - INORGANIC ANALYSES DATA PACKAGE

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

SOW No.: 7/88

Sample No.	Lab Sample ID.
BC 002	868-006-2
BC 003	868-006-3
BC 004	868-006-4
BC 004D	868-006-4D
BC 004S	868-006-4S
BC 005	868-006-5
BC 005D	868-006-5D
BC 005S	868-006-5S
BC 006	868-006-6
BC 006D	868-006-6D
BC 006S	868-006-6S
BC 007	868-006-7
BC 008	868-006-8
BC 009	868-006-9
BC 010	868-006-10
BC 011	868-006-11
BC 012	868-006-12
BC 013	868-006-13
BC 014	868-006-14
BC 015	868-006-15

Were ICP interelement corrections applied?

Yes/No NO

Were ICP background corrections applied?

Yes/No YES

If yes, were raw data generated before application of background corrections?

Yes/No NO

Comments:

I certify that this data package is in compliance with the terms, and conditions of the contract, both technically and for completeness, for other than the conditions detailed in the case narrative. Release of the data contained in this hardcopy data package has been authorized by the Laboratory Manager or the Manager's designee, as verified by the following signature.

Signature: Mark R. Bent

Name: Mark R. Bent AR309132

Date: 7-1-91

Title: CLP Manager

COVER PAGE - IN

1: 001 REV 6/89

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ENVIROFORMS/CLP 788

COVER PAGE - INORGANIC ANALYSES DATA PACKAGE

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

SOW No.: 7/88

Sample No.	Lab Sample ID.
BC 016	868-006-16
BC 016D	868-006-16D
BC 016S	868-006-16S
BC 017	868-006-17
BC 018	868-006-18
BC 019	868-006-19
BC 019D	868-006-19D
BC 019S	868-006-19S
BC 020	868-006-20
BC 021	868-006-21
BC 021D	868-006-21D
BC 021S	868-006-21S
BC 022	868-006-22
BC 022D	868-006-22D
BC 022S	868-006-22S
BC 023	868-006-23
BC 023D	868-006-23D
BC 023S	868-006-23S
BC 024	868-006-24
BC 025	868-006-25

Were ICP interelement corrections applied? Yes/No NO

Were ICP background corrections applied? Yes/No YES

If yes, were raw data generated before application of background corrections? Yes/No NO

Comments:

I certify that this data package is in compliance with the terms, and conditions of the contract, both technically and for completeness, for other than the conditions detailed in the case narrative. Release of the data contained in this hardcopy data package has been authorized by the Laboratory Manager or the Manager's designee, as verified by the following signature.

Signature:

Mark R. Barr

Name:

Mark R. Barr

Date:

4-1-91

Title:

CLP Manager

COVER PAGE - IN

1: 002 REV 6/85

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ENVIROFORMS/CLP 788

COVER PAGE - INORGANIC ANALYSES DATA PACKAGE

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: 8C_001

SOW No.: 7/88

Sample No.

8C 026

8C 027

Lab Sample ID.

868-006-26

868-006-27

Were ICP interelement corrections applied?

Yes/No NO

Were ICP background corrections applied?

Yes/No YES

If yes, were raw data generated before application of background corrections?

Yes/No NO

Comments:

I certify that this data package is in compliance with the terms, and conditions of the contract, both technically and for completeness, for other than the conditions detailed in the case narrative. Release of the data contained in this hardcopy data package has been authorized by the Laboratory Manager or the Manager's designee, as verified by the following signature.

Signature:

Mark R. Bero

Name:

Mark R. Bero

Date:

4-1-91

Title:

CLP Manager

AR309134

COVER PAGE - IN

1: 003

REV 6/89

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_004

Lab Name: IEA, INC.

Case No.: 86B-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 26AU

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	23400			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: YELLOW

Clarity After: CLEAR

44009135

Comments:

26AU

FORM I - IN

1. 004

7.

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ENVIROFORMS/CLP 78B

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_003

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 26AF

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	20300			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: YELLOW

Clarity After: CLEAR

Artifacts:

Comments:

AR309136

26AF

FORM I - IN

1 005

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_001

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 33ADU

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	27.3			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc	3280			P
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Comments:

Artifacts:
AR309137

33ADU

FORM I - IN

1 006

7

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_005

Lab Name: IEA, INC.

Case No.: 86B-006

Lab Code: IEA

SDG No.: 8C_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 33ADF

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum		-		NR
7440-36-0	Antimony		-		NR
7440-38-2	Arsenic		-		NR
7440-39-3	Barium		-		NR
7440-41-7	Beryllium		-		NR
7440-43-9	Cadmium		-		NR
7440-70-2	Calcium		-		NR
7440-47-3	Chromium		-		NR
7440-48-4	Cobalt		-		NR
7440-50-8	Copper		-		NR
7439-89-6	Iron		-		NR
7439-92-1	Lead		-		NR
7439-95-4	Magnesium		-		NR
7439-96-5	Manganese	30.1	-		P
7439-97-6	Mercury		-		NR
7440-02-0	Nickel		-		NR
7440-09-7	Potassium		-		NR
7782-49-2	Selenium		-		NR
7440-22-4	Silver		-		NR
7440-23-5	Sodium		-		NR
7440-28-0	Thallium		-		NR
7440-62-2	Vanadium		-		NR
7440-66-6	Zinc	2180	-	E	P
	Cyanide		-		NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309138

33ADF

FORM I - IN

7/88

1 007

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ENVIROFORMS/CLP 788

SAMPLE

INORGANIC ANALYSIS DATA SHEET

BC_006

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 33AU

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	30.8			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc	3340			P
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

AR309139

Comments:

33AU

FORM I - IN

1 008

7/

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_007

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 33AF

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	31.6			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc	2240		E	P
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309140

33AF

FORM I - IN

1 009

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_008

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: EQUIPMENT BLANK U

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	2.0	U		P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc	10.5	B		P
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309141

EQUIPMENT BLANK U

FORM I - IN

1 010

7/

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_009

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: EQUIPMENT BLANK F

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum		-		NR
7440-36-0	Antimony		-		NR
7440-38-2	Arsenic		-		NR
7440-39-3	Barium		-		NR
7440-41-7	Beryllium		-		NR
7440-43-9	Cadmium		-		NR
7440-70-2	Calcium		-		NR
7440-47-3	Chromium		-		NR
7440-48-4	Cobalt		-		NR
7440-50-8	Copper		-		NR
7439-89-6	Iron		-		NR
7439-92-1	Lead		-		NR
7439-95-4	Magnesium		-		NR
7439-96-5	Manganese	2.0	U		P
7439-97-6	Mercury		-		NR
7440-02-0	Nickel		-		NR
7440-09-7	Potassium		-		NR
7782-49-2	Selenium		-		NR
7440-22-4	Silver		-		NR
7440-23-5	Sodium		-		NR
7440-28-0	Thallium		-		NR
7440-62-2	Vanadium		-		NR
7440-66-6	Zinc	7.0	U	F	P
	Cyanide		-		NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309142

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FORM I - IN

1 011

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_010

Lab Name: IEA, INC.

Case No.: 868-000

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: TRIP BLANK U

Level (low/med): LOW

Date Received: 02/20/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	2.0	U		P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc	16.4	B		P
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309143

TRIP BLANK U

FORM I - IN

1 012

7/

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_011

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 39AU

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	768			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: YELLOW

Clarity After: CLEAR

Artifacts:

Comments:

AR309144

39AU

FORM I - IN

1 013

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_012

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 39AF

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	647			P
7439-97-6	Mercury				NR
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLOUDY

Texture:

Color After: YELLOW

Clarity After: CLEAR

AR 809145

Comments:

394F

FORM I - IN

1 014

7

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_013

Lab Name: IEA, INC.

Case No.: 868-000

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9ADU

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309146

9ADU

FORM I - IN

1 015

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_014

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9ADF

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

AR309747

Comments:

9ADF

FORM I - IN

1 016

7

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_015

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9ADIU

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum		-		NR
7440-36-0	Antimony		-		NR
7440-38-2	Arsenic		-		NR
7440-39-3	Barium		-		NR
7440-41-7	Beryllium		-		NR
7440-43-9	Cadmium		-		NR
7440-70-2	Calcium		-		NR
7440-47-3	Chromium		-		NR
7440-48-4	Cobalt		-		NR
7440-50-8	Copper		-		NR
7439-89-6	Iron		-		NR
7439-92-1	Lead		-		NR
7439-95-4	Magnesium		-		NR
7439-96-5	Manganese		-		NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel		-		NR
7440-09-7	Potassium		-		NR
7782-49-2	Selenium		-		NR
7440-22-4	Silver		-		NR
7440-23-5	Sodium		-		NR
7440-28-0	Thallium		-		NR
7440-62-2	Vanadium		-		NR
7440-66-6	Zinc		-		NR
	Cyanide		-		NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309148

9ADIU

FORM I - IN

1 017

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

INORGANIC ANALYSIS DATA SHEET

BC_016

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9ADIF

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

AR308449

Comments:

9ADIF

FORM 1 - IN

1 018

7/

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ENVIROFORMS/CLP 78B

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_017

Lab Name: IEA, INC.

Case No.: 86B-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 98U

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum		-		NR
7440-36-0	Antimony		-		NR
7440-38-2	Arsenic		-		NR
7440-39-3	Barium		-		NR
7440-41-7	Beryllium		-		NR
7440-43-9	Cadmium		-		NR
7440-70-2	Calcium		-		NR
7440-47-3	Chromium		-		NR
7440-48-4	Cobalt		-		NR
7440-50-8	Copper		-		NR
7439-89-6	Iron		-		NR
7439-92-1	Lead		-		NR
7439-95-4	Magnesium		-		NR
7439-96-5	Manganese		-		NR
7439-97-6	Mercury	2.2	-		CV
7440-02-0	Nickel		-		NR
7440-09-7	Potassium		-		NR
7782-49-2	Selenium		-		NR
7440-22-4	Silver		-		NR
7440-23-5	Sodium		-		NR
7440-28-0	Thallium		-		NR
7440-62-2	Vanadium		-		NR
7440-66-6	Zinc		-		NR
	Cyanide		-		NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309150

98U

FORM I - IN

1 019

7/88

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ENVIROFORMS/CLP 788

1
INORGANIC ANALYSIS DATA SHEET

SAMPLE

BC_018

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 98F

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	2.1			CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

AR089151

Comments:

98F

FORM I - IN

1 020

7.

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_019

Lab Name: IEA, INC.

Case No.: 868-000

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9BIU

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum		-		NR
7440-36-0	Antimony		-		NR
7440-38-2	Arsenic		-		NR
7440-39-3	Barium		-		NR
7440-41-7	Beryllium		-		NR
7440-43-9	Cadmium		-		NR
7440-70-2	Calcium		-		NR
7440-47-3	Chromium		-		NR
7440-48-4	Cobalt		-		NR
7440-50-8	Copper		-		NR
7439-89-6	Iron		-		NR
7439-92-1	Lead		-		NR
7439-95-4	Magnesium		-		NR
7439-96-5	Manganese		-		NR
7439-97-6	Mercury	2.6	-		CV
7440-02-0	Nickel		-		NR
7440-09-7	Potassium		-		NR
7782-49-2	Selenium		-		NR
7440-22-4	Silver		-		NR
7440-23-5	Sodium		-		NR
7440-28-0	Thallium		-		NR
7440-62-2	Vanadium		-		NR
7440-66-6	Zinc		-		NR
	Cyanide		-		NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309152

9BIU

FORM I - IN

1 021 7/88

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ENVIROFORMS/CLP 788

1
INORGANIC ANALYSIS DATA SHEET

SAMPLE NO

BC_020

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 98IF

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.27			CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

98IF

FORM I - IN

AR309153

1 022

71

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

8C_021

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9AU

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum		-		NR
7440-36-0	Antimony		-		NR
7440-38-2	Arsenic		-		NR
7440-39-3	Barium		-		NR
7440-41-7	Beryllium		-		NR
7440-43-9	Cadmium		-		NR
7440-70-2	Calcium		-		NR
7440-47-3	Chromium		-		NR
7440-48-4	Cobalt		-		NR
7440-50-8	Copper		-		NR
7439-89-6	Iron		-		NR
7439-92-1	Lead		-		NR
7439-95-4	Magnesium		-		NR
7439-96-5	Manganese		-		NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel		-		NR
7440-09-7	Potassium		-		NR
7782-49-2	Selenium		-		NR
7440-22-4	Silver		-		NR
7440-23-5	Sodium		-		NR
7440-28-0	Thallium		-		NR
7440-62-2	Vanadium		-		NR
7440-66-6	Zinc		-		NR
	Cyanide		-		NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309154

9AU

FORM I - IN

1 023

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_O.

Lab Name: IEA, INC.

Case No.: B68-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9AF

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309155

9AF

FORM I - IN

1 024

7/

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_023

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9AIU

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

Artifacts:

Comments:

AR309156

9AIU

FORM I - IN

1 025

7/88

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_024

Lab Name: IEA, INC.

Case No.: B6B-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: 9AIF

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese				NR
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: NA

Clarity After: NA

AR309157

Comments:

9AIF

FORM I - IN

1 026

7,

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ENVIROFORMS/CLP 708

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_025

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: EQUIPMENT BLANK U

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	2.0	U		P
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLOUDY

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309158

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FORM I - IN

1 027

7/88

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ENVIROFORMS/CLP 788

1
INORGANIC ANALYSIS DATA SHEET

SAMPLE NO.

BC_026

Lab Name: IEA, INC.

Case No.: 868-000

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: EQUIPMENT BLANK F

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	2.0	U		P
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLOUDY

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309159

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FORM I - IN

1 028

7/

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ENVIROFORMS/CLP 788

SAMPLE NO.

1
INORGANIC ANALYSIS DATA SHEET

BC_027

Lab Name: IEA, INC.

Case No.: 868-006

Lab Code: IEA

SDG No.: BC_001

Lab Sample ID:

Matrix (soil/water): WATER

Client ID: TRIP BLANK U

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7429-90-5	Aluminum				NR
7440-36-0	Antimony				NR
7440-38-2	Arsenic				NR
7440-39-3	Barium				NR
7440-41-7	Beryllium				NR
7440-43-9	Cadmium				NR
7440-70-2	Calcium				NR
7440-47-3	Chromium				NR
7440-48-4	Cobalt				NR
7440-50-8	Copper				NR
7439-89-6	Iron				NR
7439-92-1	Lead				NR
7439-95-4	Magnesium				NR
7439-96-5	Manganese	2.0	U		P
7439-97-6	Mercury	0.20	U		CV
7440-02-0	Nickel				NR
7440-09-7	Potassium				NR
7782-49-2	Selenium				NR
7440-22-4	Silver				NR
7440-23-5	Sodium				NR
7440-28-0	Thallium				NR
7440-62-2	Vanadium				NR
7440-66-6	Zinc				NR
	Cyanide				NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309160

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FORM I - IN --

1 029

7/88

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ENVIROFORMS/CLP 788

2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: 8C_001

Initial Calibration Source: SQL.PLUS

Continuing Calibration Source: SQL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0	593.80	99.0	1000.0	1005.48	100.5	999.92	100.0	P
Mercury	4.0	3.92	99.8	6.0	5.89	98.2	5.75	95.8	CV
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0	769.17	96.1	1000.0	959.65	96.0	971.64	97.2	P
Cyanide	100.0			200.0					

AR309161

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 030

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ENVIROFORMS/CLP 788

2A
INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0	980.46	98.0	978.95	97.9	P
Mercury	4.0			6.0	5.75	95.8	5.82	97.0	CV
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0	971.74	97.2	967.03	96.7	P
Cyanide	100.0			200.0					

AR309162

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 031

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ENVIROFORMS/CLP 788

2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0			6.0	5.78	96.3	5.72	95.3	Q
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 032

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ENVIROFORMS/CLP 788

2A
INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Initial Calibration Source: SQL.PLUS

Continuing Calibration Source: SQL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0	3.96	99.0	6.0	6.16	102.7	6.20	103.3	CV
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

AR309164

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 033

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ENVIROFORMS/CLP 788

2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-CC6

SAS No.:

SDG No.: 8C_00.

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0	4.07	101.8	6.0	5.79	96.5	5.63	93.8	0
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

AR309165

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 034

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ENVIROFORMS/CLP 788

2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0			6.0	5.67	94.5	5.71	95.2	CV
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

AR309166

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/1.3

1 035

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ENVIROFORMS/CLP 788

2A
INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Initial Calibration Source: SQL.PLUS

Continuing Calibration Source: SQL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration				
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)
Aluminum	4000.0			10000.0				
Antimony	2400.0			4000.0				
Arsenic	24.0			40.0				
Barium	4000.0			10000.0				
Beryllium	320.0			200.0				
Cadmium	400.0			2000.0				
Calcium	4000.0			10000.0				
Chromium	400.0			500.0				
Cobalt	2000.0			2000.0				
Copper	1000.0			1000.0				
Iron	4000.0			4000.0				
Lead	4000.0			4000.0				
Magnesium	4000.0			10000.0				
Manganese	600.0			1000.0				
Mercury	4.0			6.0	5.67	94.5	5.48	91.3
Nickel	1600.0			2000.0				
Potassium	9000.0			10000.0				
Selenium	15.0			25.0				
Silver	800.0			500.0				
Sodium	4000.0			10000.0				
Thallium	25.0			40.0				
Vanadium	2000.0			2000.0				
Zinc	800.0			1000.0				
Cyanide	100.0			200.0				

AR309167

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 036

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2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0			6.0	5.60	93.3	5.71	95.2	CV
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

AR309168

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

7/88

1 037

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2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00.

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0	3.86	96.5	6.0	5.63	93.8	5.17	86.2	C
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

AR309169

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

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1 038

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2A

INITIAL AND CONTINUING CALIBRATION VERIFICATION

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: B6B-006

SAS No.:

SDG No.: BC_001

Initial Calibration Source: SOL.PLUS

Continuing Calibration Source: SOL.PLUS

Concentration Units: ug/L

Analyte	Initial Calibration			Continuing Calibration					M
	True	Found	%R(1)	True	Found	%R(1)	Found	%R(1)	
Aluminum	4000.0			10000.0					
Antimony	2400.0			4000.0					
Arsenic	24.0			40.0					
Barium	4000.0			10000.0					
Beryllium	320.0			200.0					
Cadmium	400.0			2000.0					
Calcium	4000.0			10000.0					
Chromium	400.0			500.0					
Cobalt	2000.0			2000.0					
Copper	1000.0			1000.0					
Iron	4000.0			4000.0					
Lead	4000.0			4000.0					
Magnesium	4000.0			10000.0					
Manganese	600.0			1000.0					
Mercury	4.0	3.51	87.8	6.0	6.31	105.2	6.22	103.7	CV
Nickel	1600.0			2000.0					
Potassium	9000.0			10000.0					
Selenium	15.0			25.0					
Silver	800.0			500.0					
Sodium	4000.0			10000.0					
Thallium	25.0			40.0					
Vanadium	2000.0			2000.0					
Zinc	800.0			1000.0					
Cyanide	100.0			200.0					

AR309170

(1) Control Limits: Mercury 80-120; Other Metals 90-110;
Cyanide 85-115

FORM II (PART 1) - IN

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28

CRDL STANDARD FOR AA AND ICP

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

AA CRDL Standard Source: INORG. VENT.

ICP CRDL Standard Source: SPEX

Concentration Units: ug/L

Analyte	CRDL Standard for AA			CRDL Standard for ICP				
	True	Found	%R	True	Initial Found	%R	Found Found	%R
Aluminum								
Antimony				120.0				
Arsenic	10.0							
Barium								
Beryllium				10.0				
Cadmium				10.0				
Calcium								
Chromium				20.0				
Cobalt				100.0				
Copper				50.0				
Iron								
Lead	3.0			60.0				
Magnesium								
Manganese				30.0	28.12	93.7	28.94	96.
Mercury	0.2							
Nickel				80.0				
Potassium								
Selenium	5.0							
Silver				20.0				
Sodium								
Thallium	10.0							
Vanadium				100.0				
Zinc				40.0	39.35	98.4	36.79	92.

AR309171

1 040

FORM II (PART 2) - IN

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ENVIROFORMS/CLP 788

3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)	C	Continuing Calibration Blank (ug/L)						Prepa- ration Blank	C	M
			1	C	2	C	3	C			
Aluminum											
Antimony											
Arsenic											
Barium											
Beryllium											
Cadmium											
Calcium											
Chromium											
Cobalt											
Copper											
Iron											
Lead											
Magnesium											
Manganese	2.0	U	2.0	U	2.0	U	2.0	U	2.0	U	P
Mercury	0.2	U	0.2	U	0.2	U	0.2	U	0.2	U	CV
Nickel											
Potassium											
Selenium											
Silver											
Sodium											
Thallium											
Vanadium											
Zinc	7.0	U	7.0	U	7.0	U	7.0	U	7.0	U	P
Cyanide											

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FORM III - IN

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1 041

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3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_00

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)	Continuing Calibration Blank (ug/L)						Prepa- ration Blank		
		1	C	2	C	3	C		C	M
Aluminum										
Antimony										
Arsenic										
Barium										
Beryllium										
Cadmium										
Calcium										
Chromium										
Cobalt										
Copper										
Iron										
Lead										
Magnesium										
Manganese		2.0	U					2.0	U	P
Mercury		0.2	U	0.2	U	0.2	U	0.2	U	CV
Nickel										
Potassium										
Selenium										
Silver										
Sodium										
Thallium										
Vanadium										
Zinc		7.0	U					7.0	U	P
Cyanide										

AR309173

FORM III - III

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1 04:

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ENVIROFORMS/CLP 788

3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_001

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)		Continuing Calibration Blank (ug/L)						Preparation Blank	
	C		1	C	2	C	3	C	C	M
Aluminum										
Antimony										
Arsenic										
Barium										
Beryllium										
Cadmium										
Calcium										
Chromium										
Cobalt										
Copper										
Iron										
Lead										
Magnesium										
Manganese										
Mercury	0.2	U	0.2	U	0.2	U			0.2	U CV
Nickel										
Potassium										
Selenium										
Silver										
Sodium										
Thallium										
Vanadium										
Zinc										
Cyanide										

AR309174

FORM III - IN

7/88

1 043

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ENVIROFORMS/CLP 788

3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_OC

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)	C	Continuing Calibration Blank (ug/L)						Prepa- ration Blank	C	M
			1	C	2	C	3	C			
Aluminum											
Antimony											
Arsenic											
Barium											
Beryllium											
Cadmium											
Calcium											
Chromium											
Cobalt											
Copper											
Iron											
Lead											
Magnesium											
Manganese											
Mercury	0.2	U	0.2	U	0.2	U	0.2	U	0.2	U	CV
Nickel											
Potassium											
Selenium											
Silver											
Sodium											
Thallium											
Vanadium											
Zinc											
Cyanide											

AR309175

FORM III - IN

7/88 1 044

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3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: 8C_001

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)	C	Continuing Calibration Blank (ug/L)						Prepa- ration Blank	C	M
			1	C	2	C	3	C			
Aluminum											
Antimony											
Arsenic											
Barium											
Beryllium											
Cadmium											
Calcium											
Chromium											
Cobalt											
Copper											
Iron											
Lead											
Magnesium											
Manganese											
Mercury			0.2	U	0.2	U	0.2	U	0.2	U	CV
Nickel											
Potassium											
Selenium											
Silver											
Sodium											
Thallium											
Vanadium											
Zinc											
Cyanide											

AR309176

FORM III - IN

7/88 1 045

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ENVIROFORMS/CLP 788

3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: B68-006 SAS No.:

SDG No.: BC_00

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)	C	Continuing Calibration Blank (ug/L)						Prepa- ration Blank	C	M
			1	C	2	C	3	C			
Aluminum											
Antimony											
Arsenic											
Barium											
Beryllium											
Cadmium											
Calcium											
Chromium											
Cobalt											
Copper											
Iron											
Lead											
Magnesium											
Manganese											
Mercury			0.2	U	0.2	U					CV
Nickel											
Potassium											
Selenium											
Silver											
Sodium											
Thallium											
Vanadium											
Zinc											
Cyanide											

AR309177

FORM III - IN

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1 046

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ENVIROFORMS/CLP 788

3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: B68-006

SAS No.:

SDG No.: BC_001

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)		Continuing Calibration Blank (ug/L)						Preparation Blank	
	C		1	C	2	C	3	C	C	M
Aluminum										
Antimony										
Arsenic										
Barium										
Beryllium										
Cadmium										
Calcium										
Chromium										
Cobalt										
Copper										
Iron										
Lead										
Magnesium										
Manganese										
Mercury	0.2	U	0.2	U	0.2	U				CV
Nickel										
Potassium										
Selenium										
Silver										
Sodium										
Thallium										
Vanadium										
Zinc										
Cyanide										

AR309178

FORM III - IN

7/88

1 047

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ENVIROFORMS/CLP 788

3
BLANKS

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_00.

Preparation Blank Matrix (soil/water): WATER

Preparation Blank Concentration Units (ug/L or mg/kg): UG/L

Analyte	Initial Calib. Blank (ug/L)		Continuing Calibration Blank (ug/L)						Preparation Blank	
	C		1	C	2	C	3	C	C	M
Aluminum										
Antimony										
Arsenic										
Barium										
Beryllium										
Cadmium										
Calcium										
Chromium										
Cobalt										
Copper										
Iron										
Lead										
Magnesium										
Manganese										
Mercury	0.2	U	0.2	U	0.2	U				CV
Nickel										
Potassium										
Selenium										
Silver										
Sodium										
Thallium										
Vanadium										
Zinc										
Cyanide										

AR309179

FORM III - IN

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1 048

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4

ICP INTERFERENCE CHECK SAMPLE

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

ICP ID Number: PLAMSA2

ICS Source: SOL.PLUS

Concentration Units: ug/L

Analyte	True		Initial Found			Final Found		
	Sol. A	Sol. AB	Sol. A	Sol. AB	%R	Sol. A	Sol. AB	%R
Aluminum	500000	500000						
Antimony								
Arsenic								
Barium		500						
Beryllium		500						
Cadmium		1000						
Calcium	500000	500000						
Chromium		500						
Cobalt		500						
Copper		500						
Iron	200000	200000						
Lead		1000						
Magnesium	500000	500000						
Manganese		500	19	478.6	95.7	20	469.6	93.9
Mercury								
Nickel		1000						
Potassium								
Selenium								
Silver		1000						
Sodium								
Thallium								
Vanadium		500						
Zinc		1000	73	954.2	95.4	72	990.0	99.0

AR309180

FORM IV - IN

7/88 1 049

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ENVIROFORMS/CLP 788

SA
SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_0045

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	C	Spike Added (SA)	%R	Q	M
Aluminum									
Antimony									
Arsenic									
Barium									
Beryllium									
Cadmium									
Calcium									
Chromium									
Cobalt									
Copper									
Iron									
Lead									
Magnesium									
Manganese	75-125	452.4405		27.3382		500.00	85.0		
Mercury									
Nickel									
Potassium									
Selenium									
Silver									
Sodium									
Thallium									
Vanadium									
Zinc		3558.7433		3284.0897		500.00	54.9		
Cyanide									

Comments:

AR30918

1 050

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ENVIROFORMS/CLP 788

5A
SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_005S

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	C	Spike Added (SA)	%R	Q	M
Aluminum									NR
Antimony									NR
Arsenic									NR
Barium									NR
Beryllium									NR
Cadmium									NR
Calcium									NR
Chromium									NR
Cobalt									NR
Copper									NR
Iron									NR
Lead									NR
Magnesium									NR
Manganese	75-125	522.4696		30.1433		500.00	98.5		P
Mercury									NR
Nickel									NR
Potassium									NR
Selenium									NR
Silver									NR
Sodium									NR
Thallium									NR
Vanadium									NR
Zinc		2904.3584		2182.3537		500.00	144.4		P
Cyanide									NR

AR309182

Comments:

1 051

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ENVIROFORMS/CLP 788

5A
SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_006S

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR) C	Sample Result (SR) C	Spike Added (SA)	%R	Q1
Aluminum						
Antimony						
Arsenic						
Barium						
Beryllium						
Cadmium						
Calcium						
Chromium						
Cobalt						
Copper						
Iron						
Lead						
Magnesium						
Manganese	75-125	529.9856	30.7628	500.00	99.8	
Mercury						
Nickel						
Potassium						
Selenium						
Silver						
Sodium						
Thallium						
Vanadium						
Zinc		3700.2090	3344.1034	500.00	71.2	
Cyanide						

AR309183

Comments:

1 052

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ENVIROFORMS/CLP 788

SA
SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_016S

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	C	Spike Added (SA)	%R	Q	M
Aluminum			-		-				NR
Antimony			-		-				NR
Arsenic			-		-				NR
Barium			-		-				NR
Beryllium			-		-				NR
Cadmium			-		-				NR
Calcium			-		-				NR
Chromium			-		-				NR
Cobalt			-		-				NR
Copper			-		-				NR
Iron			-		-				NR
Lead			-		-				NR
Magnesium			-		-				NR
Manganese			-		-				NR
Mercury	75-125	1.0300	-	0.2000	U	1.00	103.0		CV
Nickel			-		-				NR
Potassium			-		-				NR
Selenium			-		-				NR
Silver			-		-				NR
Sodium			-		-				NR
Thallium			-		-				NR
Vanadium			-		-				NR
Zinc			-		-				NR
Cyanide			-		-				NR

AR309184

Comments:

1 053

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ENVIROFORMS/CLP 788

SA
SPIKE SAMPLE RECOVERY

SAMPLE

BC_019S

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR) C	Sample Result (SR) C	Spike Added (SA)	%R	Q M
Aluminum						
Antimony						
Arsenic						
Barium						
Beryllium						
Cadmium						
Calcium						
Chromium						
Cobalt						
Copper						
Iron						
Lead						
Magnesium						
Manganese						
Mercury	75-125	3.7800	2.6000	1.00	118.0	
Nickel						
Potassium						
Selenium						
Silver						
Sodium						
Thallium						
Vanadium						
Zinc						
Cyanide						

AR309185

Comments:

1 054

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ENVIROFORMS/CLP 783

5A
SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_021S

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	C	Spike Added (SA)	%R	Q	M
Aluminum									NR
Antimony									NR
Arsenic									NR
Barium									NR
Beryllium									NR
Cadmium									NR
Calcium									NR
Chromium									NR
Cobalt									NR
Copper									NR
Iron									NR
Lead									NR
Magnesium									NR
Manganese									NR
Mercury	75-125	0.9500		0.2000	U	1.00	95.0		CV
Nickel									NR
Potassium									NR
Selenium									NR
Silver									NR
Sodium									NR
Thallium									NR
Vanadium									NR
Zinc									NR
Cyanide									NR

Comments:

AR309186

1 055

FORM V (PART 1) - TN

7/88

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ENVIROFORMS/CLP 788

5A
SPIKE SAMPLE RECOVERY

SAMPLE

BC_0225

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	Sample Result (SR)	Spike Added (SA)	%R	Q/M
Aluminum						
Antimony						
Arsenic						
Barium						
Beryllium						
Cadmium						
Calcium						
Chromium						
Cobalt						
Copper						
Iron						
Lead						
Magnesium						
Manganese						
Mercury	75-125	0.9300	0.2000	1.00	93.0	
Nickel						
Potassium						
Selenium						
Silver						
Sodium						
Thallium						
Vanadium						
Zinc						
Cyanide						

AR30918

Comments:

1 056

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ENVIROFORMS/CLP 788

5A
SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_023S

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	C	Spike Added (SA)	%R	Q	M
Aluminum									NR
Antimony									NR
Arsenic									NR
Barium									NR
Beryllium									NR
Cadmium									NR
Calcium									NR
Chromium									NR
Cobalt									NR
Copper									NR
Iron									NR
Lead									NR
Magnesium									NR
Manganese									NR
Mercury	75-125	1.0200		0.2000	U	1.00	102.0		CV
Nickel									NR
Potassium									NR
Selenium									NR
Silver									NR
Sodium									NR
Thallium									NR
Vanadium									NR
Zinc									NR
Cyanide									NR

AR309188

Comments:

1 057

FORM V (PART 1) - TN

7/88

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ENVIROFORMS/CLP 788

5B
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO

BC_004A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	Spike Added (SA)	%R	Q	M
Aluminum								NE
Antimony								NE
Arsenic								NE
Barium								NE
Beryllium								NE
Cadmium								NE
Calcium								NE
Chromium								NE
Cobalt								NE
Copper								NE
Iron								NE
Lead								NE
Magnesium								NE
Manganese								NE
Mercury								NE
Nickel								NE
Potassium								NE
Selenium								NE
Silver								NE
Sodium								NE
Thallium								NE
Vanadium								NE
Zinc								NE
Cyanide								NE

Comments:

AR30918

1 058

FORM 11 (PART 2) - 7/81

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ENVIROFORMS/CLP 788

58
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_005A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR) C	Sample Result (SR)	Spike Added (SA)	%R	Q M
Aluminum						NR
Antimony						NR
Arsenic						NR
Barium						NR
Beryllium						NR
Cadmium						NR
Calcium						NR
Chromium						NR
Cobalt						NR
Copper						NR
Iron						NR
Lead						NR
Magnesium						NR
Manganese						NR
Mercury						NR
Nickel						NR
Potassium						NR
Selenium						NR
Silver						NR
Sodium						NR
Thallium						NR
Vanadium						NR
Zinc						NR
Cyanide						NR

AR309190

Comments:

1 059

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ENVIROFORMS/CLP 788

5B
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO

BC_006A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	Sample Result (SR)	Spike Added (SA)	%R	Q	M
Aluminum							
Antimony							
Arsenic							
Barium							
Beryllium							
Cadmium							
Calcium							
Chromium							
Cobalt							
Copper							
Iron							
Lead							
Magnesium							
Manganese							
Mercury							
Nickel							
Potassium							
Selenium							
Silver							
Sodium							
Thallium							
Vanadium							
Zinc							
Cyanide							

Comments:

AR309191

1 060

FORM V (PART 2) - IN

REV 9/

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ENVIROFORMS/CLP 788

5B
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_016A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	Spike Added (SA)	%R	Q	M
Aluminum			-					NR
Antimony			-					NR
Arsenic			-					NR
Barium			-					NR
Beryllium			-					NR
Cadmium			-					NR
Calcium			-					NR
Chromium			-					NR
Cobalt			-					NR
Copper			-					NR
Iron			-					NR
Lead			-					NR
Magnesium			-					NR
Manganese			-					NR
Mercury			-					NR
Nickel			-					NR
Potassium			-					NR
Selenium			-					NR
Silver			-					NR
Sodium			-					NR
Thallium			-					NR
Vanadium			-					NR
Zinc			-					NR
Cyanide			-					NR

AR309192

Comments:

1 061

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ENVIROFORMS/CLP 788

58
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO

BC_019A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR) C	Sample Result (SR)	Spike Added (SA)	%R	Q M
Aluminum						
Antimony						
Arsenic						
Barium						
Beryllium						
Cadmium						
Calcium						
Chromium						
Cobalt						
Copper						
Iron						
Lead						
Magnesium						
Manganese						
Mercury						
Nickel						
Potassium						
Selenium						
Silver						
Sodium						
Thallium						
Vanadium						
Zinc						
Cyanide						

AR309193

Comments:

1 062

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ENVIROFORMS/CLP 788

SB
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_021A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	Spike Added (SA)	%R	Q/M
Aluminum			-				NR
Antimony			-				NR
Arsenic			-				NR
Barium			-				NR
Beryllium			-				NR
Cadmium			-				NR
Calcium			-				NR
Chromium			-				NR
Cobalt			-				NR
Copper			-				NR
Iron			-				NR
Lead			-				NR
Magnesium			-				NR
Manganese			-				NR
Mercury			-				NR
Nickel			-				NR
Potassium			-				NR
Selenium			-				NR
Silver			-				NR
Sodium			-				NR
Thallium			-				NR
Vanadium			-				NR
Zinc			-				NR
Cyanide			-				NR

AR309194

Comments:

1 063

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ENVIROFORMS/CLP 788

58
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE N

BC_022A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR) C	Sample Result (SR)	Spike Added (SA)	%R	Q M
Aluminum						
Antimony						
Arsenic						
Barium						
Beryllium						
Cadmium						
Calcium						
Chromium						
Cobalt						
Copper						
Iron						
Lead						
Magnesium						
Manganese						
Mercury						
Nickel						
Potassium						
Selenium						
Silver						
Sodium						
Thallium						
Vanadium						
Zinc						
Cyanide						

AR309195

Comments:

1 064

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ENVIROFORMS/CLP 788

58
POST DIGEST SPIKE SAMPLE RECOVERY

SAMPLE NO.

BC_023A

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Control Limit %R	Spiked Sample Result (SSR)	C	Sample Result (SR)	Spike Added (SA)	%R	Q	M
Aluminum			-					NR
Antimony			-					NR
Arsenic			-					NR
Barium			-					NR
Beryllium			-					NR
Cadmium			-					NR
Calcium			-					NR
Chromium			-					NR
Cobalt			-					NR
Copper			-					NR
Iron			-					NR
Lead			-					NR
Magnesium			-					NR
Manganese			-					NR
Mercury			-					NR
Nickel			-					NR
Potassium			-					NR
Selenium			-					NR
Silver			-					NR
Sodium			-					NR
Thallium			-					NR
Vanadium			-					NR
Zinc			-					NR
Cyanide			-					NR

AR309196

Comments:

1 065

FORM V (PART 2) - IN

REV 8/88

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO

BC_004D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q/M
Aluminum							
Antimony							
Arsenic							
Barium							
Beryllium							
Cadmium							
Calcium							
Chromium							
Cobalt							
Copper							
Iron							
Lead							
Magnesium							
Manganese	15.0	27.3382		25.4323		7.2	P
Mercury							
Nickel							
Potassium							
Selenium							
Silver							
Sodium							
Thallium							
Vanadium							
Zinc		3284.0897		3258.4606		0.8	P
Cyanide							

AR309197

FORM VI - IN

7/88 1 066

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO.

BC_005D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum								
Antimony								
Arsenic								
Barium								
Beryllium								
Cadmium								
Calcium								
Chromium								
Cobalt								
Copper								
Iron								
Lead								
Magnesium								
Manganese	15.0	30.1433		34.3376		13.0		P
Mercury								
Nickel								
Potassium								
Selenium								
Silver								
Sodium								
Thallium								
Vanadium								
Zinc		2182.3537		2355.3897		7.6		P
Cyanide								

AR309198

FORM VI - I'

7/88

1 067

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ENVIROFORMS/CLP 788

6
 DUPLICATES

SAMPLE N

BC_006D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: B68-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum								
Antimony								
Arsenic								
Barium								
Beryllium								
Cadmium								
Calcium								
Chromium								
Cobalt								
Copper								
Iron								
Lead								
Magnesium								
Manganese	15.0	30.7628		32.0664		4.1		P
Mercury								
Nickel								
Potassium								
Selenium								
Silver								
Sodium								
Thallium								
Vanadium								
Zinc		3344.1034		3338.9517		0.2		P
Cyanide								

AR309102

FCRM VI - IN

7/89

1 068

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO.

BC_016D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q/M
Aluminum							
Antimony							
Arsenic							
Barium							
Beryllium							
Cadmium							
Calcium							
Chromium							
Cobalt							
Copper							
Iron							
Lead							
Magnesium							
Manganese							
Mercury		0.2000	U	0.2000	U		CV
Nickel							
Potassium							
Selenium							
Silver							
Sodium							
Thallium							
Vanadium							
Zinc							
Cyanide							

AR309200

1 069

FORM VI - IN

7/88

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO.

BC_019D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q/M
Aluminum							
Antimony							
Arsenic							
Barium							
Beryllium							
Cadmium							
Calcium							
Chromium							
Cobalt							
Copper							
Iron							
Lead							
Magnesium							
Manganese							
Mercury		2.6000		2.5500		1.9	CV
Nickel							
Potassium							
Selenium							
Silver							
Sodium							
Thallium							
Vanadium							
Zinc							
Cyanide							

AR309201

FORM VI - 1N

7/88

1 070

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO.

BC_021D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum								
Antimony								
Arsenic								
Barium								
Beryllium								
Cadmium								
Calcium								
Chromium								
Cobalt								
Copper								
Iron								
Lead								
Magnesium								
Manganese								
Mercury		0.2000	U	0.2000	U			CV
Nickel								
Potassium								
Selenium								
Silver								
Sodium								
Thallium								
Vanadium								
Zinc								
Cyanide								

AR309202

FORM VI - IN

7/88

1 071

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO.

BC_022D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00:

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q/M
Aluminum							
Antimony							
Arsenic							
Barium							
Beryllium							
Cadmium							
Calcium							
Chromium							
Cobalt							
Copper							
Iron							
Lead							
Magnesium							
Manganese							
Mercury		0.2000	U	0.2000	U		CV
Nickel							
Potassium							
Selenium							
Silver							
Sodium							
Thallium							
Vanadium							
Zinc							
Cyanide							

AR309200

1 072

FORM VI - IN

7/89

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ENVIROFORMS/CLP 788

6
DUPLICATES

SAMPLE NO.

BC_023D

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

% Solids for Sample: 0.0

% Solids for Duplicate:

Concentration Units (ug/L or mg/kg dry weight): UG/L

Analyte	Control Limit	Sample (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum								
Antimony								
Arsenic								
Barium								
Beryllium								
Cadmium								
Calcium								
Chromium								
Cobalt								
Copper								
Iron								
Lead								
Magnesium								
Manganese								
Mercury		0.2000	U	0.2000	U			CV
Nickel								
Potassium								
Selenium								
Silver								
Sodium								
Thallium								
Vanadium								
Zinc								
Cyanide								

AR309204

FORM VI - IN

7/88

1 073

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ENVIROFORMS/CLP 788

7

LABORATORY CONTROL SAMPLE

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006 SAS No.:

SDG No.: BC_001

Solid LCS Source: ERA

Aqueous LCS Source: SOL.PLUS

Analyte	Aqueous (ug/L)			Solid (mg/kg)			
	True	Found	%R	True	Found	C	Limits
Aluminum	4000.0			11400.0		8600.0	14200.0
Antimony	2400.0			32.0		0.0	48.0
Arsenic	40.0			40.0		15.0	65.0
Barium	4000.0			372.0		311.0	481.0
Beryllium	320.0			13.0		9.6	16.0
Cadmium	400.0			52.0		33.0	64.0
Calcium	4000.0			8310.0		6940.0	10100.0
Chromium	400.0			58.0		44.0	70.0
Cobalt	2000.0			27.0		21.0	32.0
Copper	1000.0			201.0		158.0	245.0
Iron	4000.0			16400.0		12800.0	20100.0
Lead	20.0			38.0		19.0	46.0
Magnesium	4000.0			4090.0		3050.0	5110.0
Manganese	600.0	573.59	95.6	326.0		244.0	392.0
Mercury				1.4		0.7	2.1
Nickel	1600.0			160.0		122.0	201.0
Potassium	9000.0			3680.0		3060.0	4440.0
Selenium	10.0			38.0		19.0	48.0
Silver	800.0			74.0		0.0	94.0
Sodium	4000.0			524.0		262.0	786.0
Thallium	50.0			10.0		5.0	15.0
Vanadium	2000.0			57.0		45.0	69.0
Zinc	800.0	746.20	93.3	325.0		276.0	421.0
Cyanide				98.0		39.0	157.0

AR309205

FORM VII - IN

7/88

1 07

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ENVIROFORMS/CLP 788

7

LABORATORY CONTROL SAMPLE

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Solid LCS Source: ERA

Aqueous LCS Source: SQL.PLUS

Analyte	Aqueous (ug/L)			Solid (mg/kg)				%R
	True	Found	%R	True	Found	C	Limits	
Aluminum	4000.0			11400.0			8600.0 14200.0	
Antimony	2400.0			32.0			0.0 48.0	
Arsenic	40.0			40.0			15.0 65.0	
Barium	4000.0			372.0			311.0 481.0	
Beryllium	320.0			13.0			9.6 16.0	
Cadmium	400.0			52.0			33.0 64.0	
Calcium	4000.0			8310.0			6940.0 10100.0	
Chromium	400.0			58.0			44.0 70.0	
Cobalt	2000.0			27.0			21.0 32.0	
Copper	1000.0			201.0			158.0 245.0	
Iron	4000.0			16400.0			12800.0 20100.0	
Lead	20.0			38.0			19.0 46.0	
Magnesium	4000.0			4090.0			3050.0 5110.0	
Manganese	600.0	578.36	96.4	326.0			244.0 392.0	
Mercury				1.4			0.7 2.1	
Nickel	1600.0			160.0			122.0 201.0	
Potassium	9000.0			3680.0			3060.0 4440.0	
Selenium	10.0			38.0			19.0 48.0	
Silver	800.0			74.0			0.0 94.0	
Sodium	4000.0			524.0			262.0 786.0	
Thallium	50.0			10.0			5.0 15.0	
Vanadium	2000.0			57.0			45.0 69.0	
Zinc	800.0	749.77	93.7	325.0			276.0 421.0	
Cyanide				98.0			39.0 157.0	

AR309206

FORM VII - IN

7/88

1 075

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8 STANDARD ADDITION RESULTS

SDG No.: BC_001

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7/8t

1 076

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ENVIROFORMS/CLP 788

9
ICP SERIAL DILUTIONS

SAMPLE NO.

BC_004L

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Differ- ence	Q	M
Aluminum							
Antimony							
Arsenic							
Barium							
Beryllium							
Cadmium							
Calcium							
Chromium							
Cobalt							
Copper							
Iron							
Lead							
Magnesium							
Manganese	27.34		19.43	8	28.9		P
Mercury							
Nickel							
Potassium							
Selenium							
Silver							
Sodium							
Thallium							
Vanadium							
Zinc	3284.09		3148.28		4.1		P

AR309208

1 077

FORM IX - IN

7/8L

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ENVIROFORMS/CLP 788

9
ICP SERIAL DILUTIONS

SAMPLE N

BC_005L

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Initial Sample Result (I) C	Serial Dilution Result (S) C	% Differ- ence	Q	M
Aluminum					
Antimony					
Arsenic					
Barium					
Beryllium					
Cadmium					
Calcium					
Chromium					
Cobalt					
Copper					
Iron					
Lead					
Magnesium					
Manganese	30.14	21.25	29.5		P
Mercury					
Nickel					
Potassium					
Selenium					
Silver					
Sodium					
Thallium					
Vanadium					
Zinc	2182.35	1922.85	11.9	E	P

AR309209

1 078

FORM IX - IN

7/88

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ENVIROFORMS/CLP 788

9
ICP SERIAL DILUTIONS

SAMPLE NO.

BC_006L

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Matrix (soil/water): WATER

Level (low/med): LOW

Concentration Units: ug/L

Analyte	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Differ- ence	Q	M
Aluminum		-		-		-	-
Antimony		-		-		-	-
Arsenic		-		-		-	-
Barium		-		-		-	-
Beryllium		-		-		-	-
Cadmium		-		-		-	-
Calcium		-		-		-	-
Chromium		-		-		-	-
Cobalt		-		-		-	-
Copper		-		-		-	-
Iron		-		-		-	-
Lead		-		-		-	-
Magnesium		-		-		-	-
Manganese	30.76	-	25.46	B	17.2	-	P
Mercury		-		-		-	-
Nickel		-		-		-	-
Potassium		-		-		-	-
Selenium		-		-		-	-
Silver		-		-		-	-
Sodium		-		-		-	-
Thallium		-		-		-	-
Vanadium		-		-		-	-
Zinc	3344.10	-	3274.53	-	2.1	-	P

AR309210

1 079

FORM IX - IN

7/88

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ENVIROFORMS/CLP 788

10

INSTRUMENT DETECTION LIMITS (QUARTERLY)

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_0C

ICP ID Number:

PLASMA2

Date: 03/13/91

Flame AA ID Number:

Furnace AA ID Number:

Analyte	Wave-length (nm)	Back-ground	CRDL (ug/L)	IDL (ug/L)	M
Aluminum	308.22				
Antimony	206.85				
Barium	455.41				
Beryllium	313.04				
Cadmium	226.50				
Calcium	317.93				
Chromium	267.73				
Cobalt	228.80				
Copper	324.76				
Iron	259.94				
Lead					
Magnesium	279.56				
Manganese	257.62		15.0	2.0	P
Mercury					
Nickel	231.60				
Potassium	766.49				
Selenium					
Silver	328.08				
Sodium	589.59				
Thallium					
Vanadium	292.40				
Zinc	213.87		20.0	7.0	P
Cyanide					

AR309211

Comments:

1 080

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ENVIROFORMS/CLP 788

10
INSTRUMENT DETECTION LIMITS (QUARTERLY)

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 86B-006

SAS No.:

SDG No.: BC_001

ICP ID Number:

Date: 01/15/91

Flame AA ID Number: 5000

Furnace AA ID Number:

Analyte	Wave-length (nm)	Back-ground	CRDL (ug/L)	IDL (ug/L)	M
Aluminum					
Antimony					
Arsenic					
Barium					
Beryllium					
Cadmium					
Calcium					
Chromium					
Cobalt					
Copper					
Iron					
Lead					
Magnesium					
Manganese					
Mercury	253.70		0.2	0.2	CV
Nickel					
Potassium					
Selenium					
Silver					
Sodium					
Thallium					
Vanadium					
Zinc					
Cyanide					

Comments:

AR309212

1 081

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ENVIROFORMS/CLP 788

11A
ICP INTERELEMENT CORRECTION FACTORS (Annually)

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_00

ICP ID Number: PLASMA2

Date: 01/16/90

Analyte	Wave-length (nm)	Interelement Correction Factors for:			
		Al	Ca	Fe	Mg
Aluminum					
Antimony					
Arsenic					
Barium					
Beryllium					
Cadmium					
Calcium					
Chromium					
Cobalt					
Copper					
Iron					
Lead					
Magnesium					
Manganese					
Mercury					
Nickel					
Potassium					
Selenium					
Silver					
Sodium					
Thallium					
Vanadium					
Zinc					
Cyanide					

Comment:

THE PERKIN-ELMER PLASMA II IS A SEQUENTIAL INSTRUMENT THAT DOES REQUIRE INTERELEMENT CORRECTION

77309213

FORM XI (PART 1) - IN

7/E

1 082

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ENVIROFORMS/CLP 788

12
ICP LINEAR RANGE (QUARTERLY)

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

ICP ID Number: PLASMA2

Date: 10/03/90

Analyte	Integ. Time (Sec.)	Concentration (ug/L)	M
Aluminum	1.50	550000.0	P
Antimony	9.00	50000.0	P
Arsenic			NR
Barium	1.25	100000.0	P
Beryllium	1.25	5000.0	P
Cadmium	4.50	10000.0	P
Calcium	1.25	550000.0	P
Chromium	10.00	10000.0	P
Cobalt	1.75	30000.0	P
Copper	2.00	50000.0	P
Iron	1.25	250000.0	P
Lead			NR
Magnesium	1.25	550000.0	P
Manganese	1.25	5000.0	P
Mercury			NR
Nickel	1.50	30000.0	P
Potassium	6.50	50000.0	P
Selenium			NR
Silver	3.50	1000.0	P
Sodium	2.50	200000.0	P
Thallium			NR
Vanadium	1.25	20000.0	P
Zinc	1.50	25000.0	P
Cyanide			NR

Comment:

AR309214

FORM XII - IN

7/88

1 083

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13
Preparation Log

Contract:

SAS No.:

SDG No.: BC_001

Method: P

[illegible]

AR309215

FORM XIII - IN

7/88

1 084

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ENVIROFORMS/CLP 788

13
Preparation Log

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-000

SAS No.:

SDG No.: BC_001

Method: CV

Sample No.	Preparation Date	Weight (gram)	Volume (ml)
BC 013	03/06/91		100
BC 014	03/06/91		100
BC 015	03/10/91		100
BC 016	03/10/91		100
BC 016	03/15/91		100
BC 016D	03/15/91		100
BC 016S	03/15/91		100
BC 017	03/06/91		100
BC 018	03/06/91		100
BC 019	03/10/91		100
BC 019D	03/10/91		100
BC 019S	03/10/91		100
BC 020	03/10/91		100
BC 021	03/06/91		100
BC 021D	03/06/91		100
BC 021S	03/06/91		100
BC 022	03/06/91		100
BC 022	03/14/91		100
BC 022D	03/14/91		100
BC 022S	03/14/91		100
BC 023	03/10/91		100
BC 023	03/19/91		100
BC 023D	03/19/91		100
BC 023S	03/19/91		100
BC 024	03/10/91		100
BC 025	03/06/91		100
BC 026	03/06/91		100
BC 027	03/06/91		100
PBW	03/06/91		100
PBW1	03/10/91		100
PBW2	03/14/91		100
PBW3	03/15/91		100

AR309216

FORM XIII - IN

7/88

1 085

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13
Preparation Log

Contract:

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Method: CV

[illegible]

AR309217

FORM XIII - IN

7/88 1 086

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ENVIROFORMS/CLP 788

14
Analysis Run Log

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-006

SAS No.:

SDG No.: BC_001

Instrument ID Number: Plasma2

Method: P

Start Date: 03/22/91

End Date: 03/22/91

Sample No.	D/F	Time	% R	Analytes															
				A	S	A	B	B	C	C	C	C	F	P	M	M	H	N	K
				L	B	S	A	E	D	A	R	O	U	E	B	G	N	G	I
S	1.00	1541														X			
S	1.00	1543														X			
S	1.00	1545														X			
SO	1.00	1548														X			
SO	1.00	1550														X			
ICV1	1.00	1552														X			
ICB1	1.00	1554														X			
CRI1	1.00	1557														X			
ICSA1	1.00	1559														X			
ICSA81	1.00	1601														X			
LCSW	1.00	1604														X			
PBW	1.00	1606														X			
LCSW	1.00	1609														X			
PBW	1.00	1611														X			
BC 002	1.00	1613														X			
BC 002	10.00	1616														X			
CCV1	1.00	1618														X			
CCB1	1.00	1620														X			
BC 003	1.00	1623														X			
BC 003	10.00	1625														X			
BC 004	1.00	1627														X			
BC 004D	1.00	1630														X			
BC 004S	1.00	1632														X			
BC 004L	5.00	1635														X			
BC 005	1.00	1638														X			
BC 005D	1.00	1640														X			
BC 005S	1.00	1643														X			
BC 005L	5.00	1646														X			
CCV2	1.00	1648														X			
CCB2	1.00	1651														X			
BC 006	1.00	1653														X			
BC 006D	1.00	1656														X			

AR309218

FORM XIV - IN

7/88

1 087

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14
Analysis Run Log

End Date: 03/22/91

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FORM XIV - IN

1 088

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ENVIROFORMS/CLP 788

14
Analysis Run Log

Lab Name: IEA, INC. Contract:
Lab Code: IEA Case No.: 868-006 SAS No.: SDG No.: BC_001
Instrument ID Number: 5000 Method: CV
Start Date: 03/06/91 End Date: 03/06/91

Sample No.	D/F	Time	% R	Analytes															
				A	S	B	B	C	C	C	F	P	M	M	H	N	K	S	I
				L	B	S	A	E	D	A	R	O	U	E	B	G	N	G	I
				E	G	A	L	N	N										
S0	1.00	1455														X			
S0.5	1.00	1456														X			
S1.0	1.00	1457														X			
S3.0	1.00	1459														X			
S5.0	1.00	1500														X			
S10.0	1.00	1502														X			
ICV1	1.00	1504														X			
ICB1	1.00	1505														X			
CRA1	1.00	1507														X			
CCV1	1.00	1508														X			
CCB1	1.00	1510														X			
PRW	1.00	1511														X			
BC 013	1.00	1512														X			
BC 014	1.00	1513														X			
BC 017	1.00	1515														X			
BC 018	1.00	1516														X			
BC 021	1.00	1517														X			
BC 021D	1.00	1519														X			
BC 021S	1.00	1520														X			
CCV2	1.00	1521														X			
CCB2	1.00	1523														X			
BC 022	1.00	1524														X			
BC 025	1.00	1525														X			
BC 026	1.00	1526														X			
BC 027	1.00	1528														X			
CCV3	1.00	1529														X			
CCB3	1.00	1530														X			
77777	5.00	1532																	
77777	5.00	1533																	
77777	5.00	1535																	
77777	5.00	1536																	
77777	5.00	1537																	

AR309220

FORM XIV - IN

7/BE

1 089

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ENVIROFORMS/CLP 788

14
Analysis Run Log

Lab Name: IEA, INC.

Lab Code: IEA

Instrument ID Number: 5000

Start Date: 03/06/91

Case No.: 868-006

Contract:

SAS No.:

Method: CV

End Date: 03/06/91

SDG No.: BC_001

Sample No.	D/F	Time	% R	Analytes																							
				A	S	A	B	B	C	C	C	C	F	P	M	H	N	K	S	A	N	T	I	V	Z	C	
				L	B	S	A	E	D	A	R	O	U	E	B	G	N	G	I	E	G	A	L	N	I	N	
77777	5.00	1538																									
77777	5.00	1539																									
77777	5.00	1541																									
77777	5.00	1542																									
CCV4	1.00	1543																X									
CCB4	1.00	1545																X									
77777	5.00	1546																									
77777	5.00	1547																									
77777	1.00	1548																									
77777	1.00	1549																									
77777	1.00	1551																									
77777	1.00	1552																									
77777	1.00	1553																									
77777	1.00	1554																									
77777	1.00	1555																									
CCV5	1.00	1557																X									
CCB5	1.00	1558																X									
77777	1.00	1559																									
77777	1.00	1600																									
77777	1.00	1601																									
77777	1.00	1603																									
77777	1.00	1604																									
77777	1.00	1605																									
77777	1.00	1606																									
77777	1.00	1608																									
CCV6	1.00	1609																X									
CCB6	1.00	1610																X									

AR309221

FORM XIV - IN

1 090

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14
Analysis Run Log

End Date: 03/10/91

[illegible]

1 091

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ENVIROFORMS/CLP 788

14
Analysis Run Log

Lab Name: IEA, INC. Contract: .
Lab Code: IEA Case No.: 868-006 SAS No.: SDG No.: BC_001
Instrument ID Number: 5000 Method: CV
Start Date: 03/14/91 End Date: 03/14/91

Sample No.	D/F	Time	% R	Analytes															
				A	S	A	B	B	C	C	C	C	F	P	M	H	N	K	S
				L	B	S	A	E	D	A	R	O	U	E	B	G	N	G	I
90	1.00	2025														X			
90.5	1.00	2026														X			
91.0	1.00	2028														X			
93.0	1.00	2030														X			
95.0	1.00	2032														X			
910.0	1.00	2034														X			
ICV5	1.00	2036														X			
ICB4	1.00	2040														X			
CRA3	1.00	2042														X			
CCV9	1.00	2045														X			
CCB10	1.00	2046														X			
77777	2.00	2047														X			
77777	1.00	2048														X			
77777	1.00	2049														X			
77777	1.00	2051														X			
77777	1.00	2052														X			
77777	1.00	2054														X			
77777	1.00	2055														X			
77777	1.00	2056														X			
CCV10	1.00	2058														X			
CCB11	1.00	2059														X			
77777	1.00	2100														X			
77777	1.00	2102														X			
77777	1.00	2103														X			
77777	1.00	2105														X			
CCV11	1.00	2106														X			
CCB12	1.00	2108														X			
77777	1.00	2109														X			
77777	1.00	2110														X			
77777	1.00	2112														X			
77777	1.00	2114														X			
77777	1.00	2115														X			

AR309223

FORM XIV - IN

1 092

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ENVIROFORMS/CLP 788

14

Analysis Run Log

Lab Name: IEA, INC.

Contract:

Lab Code: IEA

Case No.: 868-CO6

SAS No.:

SDG No.: BC_001

Instrument ID Number: 5000

Method: CV

Start Date: 03/14/91

End Date: 03/14/91

Sample No.	D/F	Time	% R	Analytes															
				A	S	A	B	B	C	C	C	C	F	P	M	M	H	N	K
				L	B	S	A	E	D	A	R	O	U	E	B	G	N	G	I
ZZZZZ	1.00	2116																	
ZZZZZ	1.00	2118																	
ZZZZZ	1.00	2119																	
ZZZZZ	1.00	2121																	
CCV12	1.00	2122																	
CCB13	1.00	2124																	
ZZZZZ	5.00	2125																	
ZZZZZ	5.00	2127																	
ZZZZZ	5.00	2128																	
ZZZZZ	5.00	2130																	
ZZZZZ	5.00	2131																	
ZZZZZ	5.00	2133																	
ZZZZZ	5.00	2135																	
ZZZZZ	5.00	2136																	
CCV13	1.00	2138																	
CCB14	1.00	2139																	
ZZZZZ	5.00	2140																	
ZZZZZ	5.00	2141																	
ZZZZZ	1.00	2143																	
ZZZZZ	1.00	2144																	
ZZZZZ	1.00	2145																	
CCV14	1.00	2148																	
CCB15	1.00	2150																	
ZZZZZ	1.00	2151																	
ZZZZZ	1.00	2152																	
ZZZZZ	1.00	2153																	
CCV15	1.00	2156																	
CCB16	1.00	2157																	
PBW2	1.00	2159																	
BC 022	1.00	2201																	
BC 022D	1.00	2203																	
BC 022S	1.00	2204																	

AR309224

FORM XIV - IN

7/BE

1 093

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14
Analysis Run Log

SDG No.: BC_001

AR309225

1 094

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14
Analysis Run Log

Contract:
SAS No.: SDG No.: BC_001
Method: CV
End Date: 03/15/91

[illegible]

1 095

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14
Analysis Run Log

Start Date: 03/19/91

End Date: 03/19/91

SDG No.: BC_001

AR309227

FORM XIV - IN

1 096

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IEA, INC.
CASE 868-006
SDG BC_001
METALS RAW DATA

1 OF 1

AR309228

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ICP RAW DATA

AR309229

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2 0001

BC-001#1
 DATA SHEET
 DATE 3-22-91
 TIME 8:00-6
 TEST DATE 3-5-91 / 3-13-91

3-22-91
 BC-001#1
 3-22-91

ANALYST	DATE	SAMPLE	TEST	INITIAL	FINAL	REMARKS
1	1	WCLD STD				
2	2	WCLD STD				
3	3	STD				
4	4	blank				
5	5	blank				
6	6	ICV 1				
7	7	ICB 1				
8	8	CRT 1				
9	9	ICSA 1				
10	10	ICSA 1				
11	11	LCSW	64-4			
12	12	PRW	↓			
13	13	LCSW	72-4			
14	14	PRW	↓			
15	15	RC-002	64-4			
16	16	RC-002	↓			1:10
17	17	CLV 1				
18	18	CLV 1				
19	19	RC-003	64-4			
20	20	RC-003				1:10
21	21	RC-004				
22	22	RC-004D				
23	23	RC-004S				
24	24	RC-004L				5x
25	25	RC-005				
26	26	RC-005D				
27	27	RC-005S				
28	28	RC-005L	↓			5x
29	29	CLV 2				
30	30	CLV 2				
31	31	RC-006	72-4			
32	32	RC-006D				R300230
33	33	RC-006S				
34	34	RC-006L	↓			5x
35	35	RC-007	64-4			

ANALYST: M. H. H. DATE: 3-25-91
 TEST: TRP

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2 0002

ICP DATA SHEET
(CONT.)

ANALYSIS	QTY.	PARCEL ID#	WEIGHT	INITIAL WEIGHT/ VOLUME	FINAL VOLUME	DIFFERENCE WEIGHT
36	36	RC-008	64-4			
37	37	RC-009	↓			
38	38	RC-010				
39	39	BLK				
40	40	CCV 3				
41	41	CCR 3				
42	42	RC-011	64-4			
43	43	RC-012	↓			
44	44	RC-025				
45	45	RC-026	↓			
46	46	RC-027				
47	47	TCSA 2				
48	48	TCSAB 2				
49	49	CRI 2				
50	50	CCV-2 nd 4				
51	51	CCR 4				
52						
53						
54						
55						
56						
57						
58						
59						
60						
61						
62						
63						
64						
65						
66						
67						
68						
69						

AR309231

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Method Name: CN_00101

Comment:

Read Delay : 40 sec

Replicates : 2 Format name : display

Analysis Number	AS Posn	Sample ID	Sequence Name	Dilution	Weight / Volume
1	1	wcal std	bikseq		
2	2	wcal std	bikseq		
3	3	std	bikseq		
4	4	blank	bikseq		
5	5	blank	bikseq		
6	6	ICV1	bikseq		
7	7	ICB1	bikseq		
8	8	CRI1	bikseq		
9	9	ICSA1	bikseq		
10	10	ICSA1	bikseq		
11	11	LCSW	bikseq		
12	12	FBW	bikseq		
13	13	LCSW	bikseq		
14	14	FBW	bikseq		
15	15	EC_002	Mn		
16	16	EC_002	Mn	1:10	
17	17	CCV1	bikseq		
18	18	CCB1	bikseq		
19	19	EC_003	Mn		
20	20	EC_003	Mn	1:10	
21	21	EC_004	bikseq		
22	22	EC_004D	bikseq		
23	23	EC_004S	bikseq		
24	24	EC_004L	bikseq		
25	25	EC_005	bikseq		
26	26	EC_005D	bikseq		
27	27	EC_005S	bikseq		
28	28	EC_005L	bikseq		
29	29	CCV2	bikseq		
30	30	CCB2	bikseq		
31	31	EC_006	bikseq		
32	32	EC_006D	bikseq		
33	33	EC_006S	bikseq		
34	34	EC_006L	bikseq		
35	35	EC_007	bikseq		
36	36	EC_008	bikseq		
37	37	EC_009	bikseq		
38	38	EC_010	bikseq		
39	39	BLK	bikseq		
40	40	CCV3	bikseq		
41	41	CCB3	bikseq		
42	42	EC_011	Mn		
43	43	EC_012	Mn		
44	44	EC_023	Mn		
45	45	EC_024	Mn		
46	46	EC_027	Mn		
47	47	ICSA2	bikseq		
48	48	ICSA2	bikseq		
49	49	CRI2	bikseq		
50	50	CCV4	bikseq		
51	51	CCB4	bikseq		

AR309232

2 0003

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ICP STANDARDS LOG

Method# BC-001*1 3-26-11 Instrument# ICP
SDG# BC-001 3-26-11 Analyst Thel. Fells
IEA# ABG-6

Prep date: 35-91/3-13-91

Date: 3-22-91

[illegible]

Reviewed by: Michael Cal Date: 3-25-91

Reviewed by: _____ Date: _____

COMMENTS:

USE: Zn, Mn

AR309233

2 0004

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03/22/91 15:42

wcal std	rep	1	ZNa213	em	17809.9	conc	2000.0
	rep	1	MNa257	em	24672.7	conc	1500.0
	rep	2	ZNa213	em	18314.2	conc	2000.0
	rep	2	MNa257	em	27007.5	conc	1500.0

wcal std

03/22/91 15:42

ZNa213	av	18042.0	sd	356.56	%cv	1.97	conc	2000.0
MNa257	av	24840.1	sd	234.68	%cv	0.88	conc	1500.0

03/22/91 15:44

wcal std	rep	1	ZNa213	em	20237.5	conc	2000.0
	rep	1	MNa257	em	29088.8	conc	1500.0
	rep	2	ZNa213	em	20027.9	conc	2000.0
	rep	2	MNa257	em	29135.8	conc	1500.0

wcal std

03/22/91 15:44

ZNa213	av	20132.7	sd	148.25	%cv	0.74	conc	2000.0
MNa257	av	29112.3	sd	33.21	%cv	0.11	conc	1500.0

03/22/91 15:46

std	rep	1	ZNa213	em	22119.2	conc	2000.0
	rep	1	MNa257	em	31330.9	conc	1500.0
	rep	2	ZNa213	em	22040.4	conc	2000.0
	rep	2	MNa257	em	31282.2	conc	1500.0

std

03/22/91 15:47

ZNa213	av	22079.8	sd	55.77	%cv	0.25	conc	2000.0
MNa257	av	31306.5	sd	34.37	%cv	0.11	conc	1500.0

03/22/91 15:49

blank	rep	1	ZNa213	em	21.3
	rep	1	MNa257	em	23.7
	rep	2	ZNa213	em	15.6
	rep	2	MNa257	em	49.1

blank

03/22/91 15:49

ZNa213	av	18.5	sd	4.02	%cv	21.78
MNa257	av	34.4	sd	17.96	%cv	49.38

03/22/91 15:51

blank	rep	1	ZNa213	em	34.9
	rep	1	MNa257	em	22.7
	rep	2	ZNa213	em	51.7
	rep	2	MNa257	em	39.9

blank

03/22/91 15:51

ZNa213	av	44.3	sd	10.44	%cv	23.56
MNa257	av	31.3	sd	12.18	%cv	38.95

03/22/91 15:53

ICV1	rep	1	ZNa213	conc	770.5	ppb
	rep	1	MNa257	conc	600.5	ppb
	rep	2	ZNa213	conc	747.8	ppb
	rep	2	MNa257	conc	587.1	ppb

ICV1

03/22/91 15:53

ZNa213	av	769.2	ppb	sd	1.88	%cv	0.24
MNa257	av	593.8	ppb	sd	9.48	%cv	1.59

AR309234

2 0005

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03/22/91 15:55						
ICBI	rep	1	ZNa213	conc	1.1 ppb	
	rep	1	MNa257	conc	-1.4 ppb	
	rep	2	ZNa213	conc	-1.5 ppb	
	rep	2	MNa257	conc	0.3 ppb	
ICBI						
03/22/91 15:56						
ZNa213	av		-0.2 ppb	sd	1.84 %av	1010.5
MNa257	av		-0.5 ppb	sd	1.19 %av	216.48
03/22/91 15:58						
CRI1	rep	1	ZNa213	conc	38.7 ppb	
	rep	1	MNa257	conc	28.3 ppb	
	rep	2	ZNa213	conc	40.0 ppb	
	rep	2	MNa257	conc	27.9 ppb	
CRI1						
03/22/91 15:58						
ZNa213	av		39.3 ppb	sd	0.98 %av	2.49
MNa257	av		28.1 ppb	sd	0.24 %av	0.91
03/22/91 16:00						
ICSA1	rep	1	ZNa213	conc	74.3 ppb	
	rep	1	MNa257	conc	18.3 ppb	
	rep	2	ZNa213	conc	70.7 ppb	
	rep	2	MNa257	conc	19.3 ppb	
ICSA1						
03/22/91 16:00						
ZNa213	av		72.5 ppb	sd	2.55 %av	3.51
MNa257	av		18.9 ppb	sd	0.57 %av	3.04
03/22/91 16:02						
ICSA1	rep	1	ZNa213	conc	953.2 ppb	
	rep	1	MNa257	conc	478.3 ppb	
	rep	2	ZNa213	conc	955.2 ppb	
	rep	2	MNa257	conc	478.7 ppb	
ICSA1						
03/22/91 16:03						
ZNa213	av		954.2 ppb	sd	1.38 %av	0.14
MNa257	av		478.6 ppb	sd	0.15 %av	0.03
03/22/91 16:05						
LCSW	rep	1	ZNa213	conc	741.2 ppb	
	rep	1	MNa257	conc	576.1 ppb	
	rep	2	ZNa213	conc	751.2 ppb	
	rep	2	MNa257	conc	571.1 ppb	
LCSW						
03/22/91 16:05						
ZNa213	av		746.2 ppb	sd	7.08 %av	0.93
MNa257	av		573.6 ppb	sd	3.58 %av	0.62
03/22/91 16:07						
FW	rep	1	ZNa213	conc	4.3 ppb	
	rep	1	MNa257	conc	0.4 ppb	
	rep	2	ZNa213	conc	2.7 ppb	
	rep	2	MNa257	conc	-0.6 ppb	
FW						
03/22/91 16:07						
ZNa213	av		3.3 ppb	sd	1.14 %av	32.33
MNa257	av		-0.1 ppb	sd	0.71 %av	456.28

AR309235

2-0005

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03/22/91 16:09

LCSW	rep	1	ZNa213	conc	743.3 ppb
	rep	1	MNa257	conc	580.4 ppb
	rep	2	ZNa213	conc	754.2 ppb
	rep	2	MNa257	conc	574.1 ppb

LCSW

03/22/91 16:10

ZNa213	av	749.8 ppb	sd	9.09 %av	1.21
MNa257	av	578.4 ppb	sd	3.19 %av	0.55

03/22/91 16:12

PBW	rep	1	ZNa213	conc	2.4 ppb
	rep	1	MNa257	conc	0.5 ppb
	rep	2	ZNa213	conc	3.1 ppb
	rep	2	MNa257	conc	-1.2 ppb

PBW

03/22/91 16:12

ZNa213	av	2.9 ppb	sd	0.35 %av	12.13
MNa257	av	-0.4 ppb	sd	1.24 %av	353.17

03/22/91 16:14

BC_002	rep	1	MNa257	conc	25475.4 ppb
	rep	2	MNa257	conc	25147.0 ppb

BC_002

03/22/91 16:14

MNa257	av	25311.3 ppb	sd	232.37 %av	0.92
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03/22/91 16:16

BC_002	rep	1	MNa257	conc	23478.2 ppb
	rep	2	MNa257	conc	23108.9 ppb

BC_002

03/22/91 16:17

MNa257	av	23393.5 ppb	sd	402.40 %av	1.72
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03/22/91 16:19

CCV1	rep	1	ZNa213	conc	957.3 ppb
	rep	1	MNa257	conc	1020.1 ppb
	rep	2	ZNa213	conc	942.0 ppb
	rep	2	MNa257	conc	990.9 ppb

CCV1

03/22/91 16:19

ZNa213	av	959.7 ppb	sd	3.34 %av	0.35
MNa257	av	1005.5 ppb	sd	20.49 %av	2.04

03/22/91 16:21

CCB1	rep	1	ZNa213	conc	1.1 ppb
	rep	1	MNa257	conc	0.5 ppb
	rep	2	ZNa213	conc	0.5 ppb
	rep	2	MNa257	conc	0.9 ppb

CCB1

03/22/91 16:21

ZNa213	av	0.8 ppb	sd	0.47 %av	59.47
MNa257	av	0.7 ppb	sd	0.29 %av	39.47

03/22/91 16:24

BC_003	rep	1	MNa257	conc	21774.5 ppb
	rep	2	MNa257	conc	21414.7 ppb

BC_003

03/22/91 16:24

MNa257	av	21774.6 ppb	sd	254.48 %av	1.17
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AR309236

2.0007

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03/22/91 14:24						
HC_003	rep	1	MNa257	conc	20307.4 ppb	
	rep	2	MNa257	conc	20234.5 ppb	
HC_003						
03/22/91 14:24						
MNa257	av		20271.0 ppb	sd	51.53 %av	0.25
03/22/91 14:28						
HC_004	rep	1	ZNa213	conc	3291.1 ppb	
	rep	1	MNa257	conc	28.8 ppb	
	rep	2	ZNa213	conc	3277.0 ppb	
	rep	2	MNa257	conc	25.9 ppb	
HC_004						
03/22/91 14:28						
ZNa213	av		3284.1 ppb	sd	9.97 %av	0.30
MNa257	av		27.3 ppb	sd	2.00 %av	7.33
03/22/91 14:31						
HC_004D	rep	1	ZNa213	conc	3245.2 ppb	
	rep	1	MNa257	conc	25.2 ppb	
	rep	2	ZNa213	conc	3251.8 ppb	
	rep	2	MNa257	conc	25.4 ppb	
HC_004D						
03/22/91 14:31						
ZNa213	av		3258.5 ppb	sd	9.47 %av	0.29
MNa257	av		25.4 ppb	sd	0.24 %av	1.02
03/22/91 14:33						
HC_004E	rep	1	ZNa213	conc	3583.4 ppb	
	rep	1	MNa257	conc	451.7 ppb	
	rep	2	ZNa213	conc	3534.1 ppb	
	rep	2	MNa257	conc	453.2 ppb	
HC_004E						
03/22/91 14:34						
ZNa213	av		3558.7 ppb	sd	34.83 %av	0.98
MNa257	av		452.4 ppb	sd	1.01 %av	0.22
03/22/91 14:34						
HC_004L	rep	1	ZNa213	conc	428.5 ppb	
	rep	1	MNa257	conc	3.3 ppb	
	rep	2	ZNa213	conc	430.8 ppb	
	rep	2	MNa257	conc	4.5 ppb	
HC_004L						
03/22/91 14:34						
ZNa213	av		429.7 ppb	sd	1.48 %av	0.24
MNa257	av		3.9 ppb	sd	0.81 %av	20.74
03/22/91 14:39						
HC_005	rep	1	ZNa213	conc	2287.3 ppb	
	rep	1	MNa257	conc	30.5 ppb	
	rep	2	ZNa213	conc	2155.4 ppb	
	rep	2	MNa257	conc	29.7 ppb	
HC_005						
03/22/91 14:39						
ZNa213	av		2182.4 ppb	sd	38.15 %av	1.75
MNa257	av		30.1 ppb	sd	0.57 %av	1.88
03/22/91 14:41						
HC_005D	rep	1	ZNa213	conc	2341.9 ppb	

AR309237

2 0008

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	rep	1	MNa257	conc	34.4 ppb		
	rep	2	ZNa213	conc	2348.9 ppb		
	rep	2	MNa257	conc	34.3 ppb		
BC_005D							
03/22/91 16:41							
ZNa213	av		2355.4 ppb	sd	9.17 %av	0.39	
MNa257	av		34.3 ppb	sd	0.06 %av	0.19	
03/22/91 16:44							
BC_005B							
	rep	1	ZNa213	conc	2902.0 ppb		
	rep	1	MNa257	conc	522.3 ppb		
	rep	2	ZNa213	conc	2904.8 ppb		
	rep	2	MNa257	conc	522.4 ppb		
BC_005B							
03/22/91 16:44							
ZNa213	av		2904.4 ppb	sd	3.40 %av	0.12	
MNa257	av		522.5 ppb	sd	0.18 %av	0.03	
03/22/91 16:46							
BC_005L							
	rep	1	ZNa213	conc	383.4 ppb		
	rep	1	MNa257	conc	4.0 ppb		
	rep	2	ZNa213	conc	385.8 ppb		
	rep	2	MNa257	conc	4.5 ppb		
BC_005L							
03/22/91 16:47							
ZNa213	av		384.6 ppb	sd	1.69 %av	0.44	
MNa257	av		4.3 ppb	sd	0.33 %av	7.64	
03/22/91 16:49							
CCV2							
	rep	1	ZNa213	conc	943.8 ppb		
	rep	1	MNa257	conc	1004.2 ppb		
	rep	2	ZNa213	conc	979.5 ppb		
	rep	2	MNa257	conc	995.4 ppb		
CCV2							
03/22/91 16:49							
ZNa213	av		971.6 ppb	sd	11.05 %av	1.14	
MNa257	av		999.9 ppb	sd	6.05 %av	0.61	
03/22/91 16:52							
CCB2							
	rep	1	ZNa213	conc	0.1 ppb		
	rep	1	MNa257	conc	0.6 ppb		
	rep	2	ZNa213	conc	-2.6 ppb		
	rep	2	MNa257	conc	0.1 ppb		
CCB2							
03/22/91 16:52							
ZNa213	av		-1.3 ppb	sd	1.95 %av	155.50	
MNa257	av		0.3 ppb	sd	0.35 %av	105.23	
03/22/91 16:54							
BC_006							
	rep	1	ZNa213	conc	3384.0 ppb		
	rep	1	MNa257	conc	30.9 ppb		
	rep	2	ZNa213	conc	3304.2 ppb		
	rep	2	MNa257	conc	30.4 ppb		
BC_006							
03/22/91 16:55							
ZNa213	av		3344.1 ppb	sd	56.44 %av	1.69	
MNa257	av		30.8 ppb	sd	0.17 %av	0.56	
03/22/91 16:57							
BC_006D							
	rep	1	ZNa213	conc	3335.7 ppb		

AR309238

2 0009

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	rep	1	MNa257	conc	31.3 ppb		
	rep	2	ZNa213	conc	3342.2 ppb		
	rep	2	MNa257	conc	32.9 ppb		
BC_004D							
03/22/91	16:57						
	ZNa213	av	3339.0 ppb	sd	4.63 %av	0.14	
	MNa257	av	32.1 ppb	sd	1.13 %av	3.53	
03/22/91 16:59							
BC_004E	rep	1	ZNa213	conc	3709.2 ppb		
	rep	1	MNa257	conc	524.8 ppb		
	rep	2	ZNa213	conc	3691.2 ppb		
	rep	2	MNa257	conc	533.2 ppb		
BC_004E							
03/22/91	17:00						
	ZNa213	av	3700.2 ppb	sd	12.74 %av	0.34	
	MNa257	av	530.0 ppb	sd	4.53 %av	0.84	
03/22/91 17:02							
BC_004L	rep	1	ZNa213	conc	457.5 ppb		
	rep	1	MNa257	conc	5.0 ppb		
	rep	2	ZNa213	conc	452.3 ppb		
	rep	2	MNa257	conc	5.2 ppb		
BC_004L							
03/22/91	17:02						
	ZNa213	av	454.9 ppb	sd	3.45 %av	0.54	
	MNa257	av	5.1 ppb	sd	0.11 %av	2.23	
03/22/91 17:05							
BC_007	rep	1	ZNa213	conc	2236.8 ppb		
	rep	1	MNa257	conc	32.2 ppb		
	rep	2	ZNa213	conc	2246.8 ppb		
	rep	2	MNa257	conc	31.0 ppb		
BC_007							
03/22/91	17:05						
	ZNa213	av	2241.8 ppb	sd	7.07 %av	0.32	
	MNa257	av	31.4 ppb	sd	0.90 %av	2.83	
03/22/91 17:07							
BC_008	rep	1	ZNa213	conc	12.0 ppb		
	rep	1	MNa257	conc	0.9 ppb		
	rep	2	ZNa213	conc	9.0 ppb		
	rep	2	MNa257	conc	-0.1 ppb		
BC_008							
03/22/91	17:07						
	ZNa213	av	10.5 ppb	sd	2.15 %av	20.43	
	MNa257	av	0.1 ppb	sd	0.75 %av	177.89	
03/22/91 17:10							
BC_009	rep	1	ZNa213	conc	5.4 ppb		
	rep	1	MNa257	conc	1.8 ppb		
	rep	2	ZNa213	conc	7.4 ppb		
	rep	2	MNa257	conc	0.1 ppb		
BC_009							
03/22/91	17:10						
	ZNa213	av	4.4 ppb	sd	1.39 %av	17.75	
	MNa257	av	0.9 ppb	sd	1.20 %av	134.57	
03/22/91 17:12							
BC_010	rep	1	ZNa213	conc	18.9 ppb		

AR309239

2 0010

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	rep	1	MNa257	conc	1.0 ppb	
	rep	2	ZNa213	conc	13.9 ppb	
	rep	2	MNa257	conc	0.4 ppb	
HC_010						
03/22/91 17:12						
	av		16.4 ppb	sd	3.54 %cv	21.53
	av		0.8 ppb	sd	0.29 %cv	35.09
03/22/91 17:14						
HLK	rep	1	ZNa213	conc	0.8 ppb	
	rep	1	MNa257	conc	-1.3 ppb	window edge
	rep	2	ZNa213	conc	2.7 ppb	
	rep	2	MNa257	conc	0.2 ppb	
BLK						
03/22/91 17:15						
	av		1.8 ppb	sd	1.34 %cv	74.51
	av		-0.5 ppb	sd	1.04 %cv	192.36
03/22/91 17:17						
CCV3	rep	1	ZNa213	conc	941.6 ppb	
	rep	1	MNa257	conc	987.1 ppb	
	rep	2	ZNa213	conc	981.8 ppb	
	rep	2	MNa257	conc	973.8 ppb	
CCV3						
03/22/91 17:17						
	av		971.7 ppbv	sd	14.29 %cv	1.47
	av		980.5 ppbv	sd	9.40 %cv	0.96
03/22/91 17:19						
CCB3	rep	1	ZNa213	conc	0.6 ppb	
	rep	1	MNa257	conc	-2.1 ppb	window edge
	rep	2	ZNa213	conc	-0.2 ppb	
	rep	2	MNa257	conc	0.6 ppb	
CCB3						
03/22/91 17:20						
	av		0.2 ppb	sd	0.57 %cv	282.43
	av		-0.8 ppb	sd	1.91 %cv	254.27
03/22/91 17:22						
HC_011	rep	1	MNa257	conc	749.0 ppb	
	rep	2	MNa257	conc	747.2 ppb	
HC_011						
03/22/91 17:22						
	av		748.1 ppb	sd	1.24 %cv	0.16
03/22/91 17:24						
HC_012	rep	1	MNa257	conc	648.8 ppb	
	rep	2	MNa257	conc	645.0 ppb	
HC_012						
03/22/91 17:24						
	av		646.9 ppb	sd	2.71 %cv	0.42
03/22/91 17:24						
HC_025	rep	1	MNa257	conc	-0.1 ppb	AR309260
	rep	2	MNa257	conc	-1.6 ppb	window edge
HC_025						
03/22/91 17:26						
	av		-0.8 ppb	sd	1.07 %cv	125.40
03/22/91 17:28						

2 0011

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HC_024	rep	1	MNa257	conc	0.3 ppb		
	rep	2	MNa257	conc	-0.1 ppb		
HC_024							
03/22/91 17:28							
MNa257	av		0.1 ppb	sd	0.29 %av	347.41	
03/22/91 17:30							
HC_027	rep	1	MNa257	conc	-0.7 ppb		
	rep	2	MNa257	conc	1.7 ppb		
HC_027							
03/22/91 17:30							
MNa257	av		0.5 ppb	sd	1.73 %av	350.06	
03/22/91 17:32							
ICBA2	rep	1	ZNa213	conc	70.5 ppb		
	rep	1	MNa257	conc	21.0 ppb		
	rep	2	ZNa213	conc	73.6 ppb		
	rep	2	MNa257	conc	18.9 ppb		
ICBA2							
03/22/91 17:33							
ZNa213	av		72.0 ppb	sd	2.17 %av	3.01	
MNa257	av		20.0 ppb	sd	1.52 %av	7.62	
03/22/91 17:35							
ICBA2	rep	1	ZNa213	conc	943.7 ppb		
	rep	1	MNa257	conc	471.3 ppb		
	rep	2	ZNa213	conc	1014.3 ppb		
	rep	2	MNa257	conc	467.8 ppb		
ICBA2							
03/22/91 17:35							
ZNa213	av		990.0 ppb	sd	34.41 %av	3.48	
MNa257	av		469.6 ppb	sd	2.52 %av	0.54	
03/22/91 17:37							
CR12	rep	1	ZNa213	conc	34.8 ppb		
	rep	1	MNa257	conc	28.1 ppb		
	rep	2	ZNa213	conc	38.7 ppb		
	rep	2	MNa257	conc	29.8 ppb		
CR12							
03/22/91 17:37							
ZNa213	av		36.8 ppb	sd	2.78 %av	7.55	
MNa257	av		28.9 ppb	sd	1.22 %av	4.22	
03/22/91 17:39							
CCV4	rep	1	ZNa213	conc	953.3 ppb		
	rep	1	MNa257	conc	979.7 ppb		
	rep	2	ZNa213	conc	980.8 ppb		
	rep	2	MNa257	conc	978.2 ppb		
CCV4							
03/22/91 17:40							
ZNa213	av		967.0 ppb	sd	19.43 %av	2.01	
MNa257	av		978.9 ppb	sd	1.11 %av	0.11	
03/22/91 17:41							
CCB4	rep	1	ZNa213	conc	-0.2 ppb		
	rep	1	MNa257	conc	-0.3 ppb		
	rep	2	ZNa213	conc	-2.3 ppb		
	rep	2	MNa257	conc	0.3 ppb		
CCB4							
03/22/91 17:42							

AR309241

2 0012

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ZNa213	av	-1.3 ppb	sd	1.45 %av 114.65
MNa257	av	0.0 ppb	sd	0.47 %av 3483.8

AR309242

2 0013

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MERCURY

AR309243

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MERCURY PREP and ANALYSIS

00162

Prep date 6 MAR 91 Analysis Date 6 MAR 91
 Bath Temp 95°C Lamp Energy 61
 Spike Conc. 1ppb Vendor MALINCROFT Corr Coeff. 0.99964

Sample ID #	Weight	Volume	Dil. factor	ABS.	Time	Value from curve	Reported Value	% Recovery
S 0				0.001	1455			
S 0.5 (A)				0.017	1456			
S 1.0				0.032	1457			
S 3.0				0.094	1459			
S 5.0				0.152	1500			
S 10.0				0.290	1502			
TCV (A) 4.0 ppb				0.119	1504	3.99		99.7%
TCR				0.001	1505			
GRA(E) 0.2 ppb				0.009	1507	0.19		92.7%
CCV 1 (E) 6.0 ppb				0.174	1508	5.89		98.1%
CCB 1				0.001	1510	-0.09		
PRW				0.001	1511	-0.09		
BC013		100mL		0.001	1512	-0.09	<0.0005	
BC014				0.001	1513	-0.09	<0.0005	
BC017				0.067	1515	2.19	0.0022	
BC018				0.065	1516	2.12	0.0021	
BC021				0.001	1517	-0.09	<0.0005	
BC021 D				0.001	1519	-0.09	<0.0005	
BC021 S				0.031	1520	0.95	—	94.6%
CCV 2				0.170	1521	5.75		95.8%
CCB 2				0.001	1523	-0.09		
BC022		100mL		0.001	1524	-0.09	<0.0005	
BC025				0.001	1525	-0.09	<0.0005	
BC026				0.001	1526	-0.09	<0.0005	
BC027				0.001	1528	-0.09	<0.0005	
CCV 3				0.170	1529	5.75		95.8%
CCB 3				0.001	1530	-0.09		

Reviewed by: Dianne Hunder

Date: 3-9-91

Prep Analyst: MSM

Analyst: MB AR309244

* STD # 333

+ STD # 336

2 0014

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MERCURY PREP and ANALYSIS

00101

Prep date 6 MAR 91

Bath Temp 95°C

Spike Conc. 1 ppb

Vendor MILLIKRAT

Analysis Date 6 MAR 91

Lamp Energy 95°C 61

Corr Coeff. 0.99964

Sample ID #	Weight	Volume	Dil. factor	ABS.	Time	Value from curve	Reported Value	% Recovery
T267-58(0)-1 (5X)		100mL	5X	0.001	1532	-0.09	<0.0025	
-1S				0.090	1533	2.99	—	74.6%
✓ T747-27-1				0.001	1535	-0.09	<0.0025	
-1D				0.001	1536	-0.09	<0.0025	
-1S				0.098	1537	3.26	—	81.5%
✓ T1167-1-4				0.002	1538	-0.06	<0.0025	
-4S				0.113	1539	3.78	—	94.5%
T-BLANK-198				0.002	1541	-0.06	<0.0025	
-198S	↓	↓	↓	0.115	1542	3.85	—	96.2%
✓ CCV4				0.172	1543	5.82		96.9%
CCB4				0.001	1545	-0.09		
T-BLANK-199 (5X)		100mL	5X	0.001	1546	-0.09	<0.0025	
-199S	↓	↓	↓	0.110	1547	3.68	—	91.9%
✓ E860-102(0)-1		100mL		0.001	1548	-0.09	<0.0005	
-1D				0.001	1549	-0.09	<0.0005	
-2				0.001	1551	-0.09	<0.0005	
-2S				0.031	1552	0.95	—	94.6%
E-BLANK-82				0.001	1553	-0.09	<0.0005	
✓ 204-428(0)-1E				0.001	1554	-0.09	<0.0005	
-2E		↓		0.001	1555	-0.09	<0.0005	
✓ CCV5				0.171	1557	5.78		96.4%
CCB5				0.001	1558	-0.09		
✓ 204-429(0)-1		100mL		0.001	1559	-0.09	<0.0005	
-2				0.001	1600	-0.09	<0.0005	
-3				0.001	1601	-0.09	<0.0005	
-3D				0.001	1603	-0.09	<0.0005	
-3S		↓		0.032	1604	0.98	—	98.0%

Reviewed by: Diana Hander

Date: 3-9-91

† STD # 336

Prep Analyst: MSS

Analyst: MSS

2 0015

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00164

Prep date 6 MAR 91 Analysis Date 6 MAR 91
 Bath Temp 95°C Lamp Energy 61
 Spike Conc. 1 ppb Vendor MALLINCKRODT Corr Coeff. 0.99964

Prep Analyst: MB

Analyst: MSK

+ STA # 336

2 0016

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00169

Corr Coeff. 0.9997

AR3092

Prep Analyst: Brian [Signature]
Analyst: Brian [Signature]

2 0017

† Std # 356

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MERCURY PREP and ANALYSIS

00172

Prep date 14 MAR 91

Bath Temp 95°C

Spike Conc. 1 ppb

Vendor MALINCKROTT

SDG: GN-155

Analysis Date 3-14-91

Lamp Energy 61

Corr Coeff. 0.9999

Sample ID #	Weight	Volume	Dil. factor	ABS.	Time	Value from curve	Reported Value	% Recovery
S 0				0.001	20:25			
S 0.5 (A)				0.015	20:26			
S 1.0				0.029	20:28			
S 3.0				0.081	20:30			
S 5.0				0.130	20:32			
S 10.0				0.264	20:34			
+ ICB (E) 4.0 ppb				0.108	20:36	4.07		101.6
ICB				0.001	20:40			
+ CRA (E) 0.2 ppb				0.006	20:42	0.166		82.9
+ CCV1 (E) 6.0 ppb				0.153	20:45	5.79		96.4
CCB1				0.001	20:46	-0.025		
LCSS (E) 209	0.25g	50mL	2X	0.137	20:47	5.12		
PBS (F) 6651	0.20g			0.000	20:49	-0.064		
GN-155	0.21g			0.007	20:49	0.209		
GN-156	0.23g			0.002	20:51	0.013		
GN-157	0.22g			0.003	20:52	0.051		
GN-158	0.24g			0.003	20:54	0.051		
GN-159	0.23g			0.005	20:55	0.128		
GN-160	0.25g			0.008	20:56	0.242		
+ CCV2				0.149	20:58	5.633		93.9
CCB2				0.003	20:59	0.051		
GN-161	0.25g			0.011	21:00	0.357		
GN-162	0.21g			0.004	21:02	0.089		
GN-162 D	0.25g			0.004	21:03	0.089		
GN-162 S	0.25g			0.027	21:05	0.969		
+ CCV3				0.150	21:06	5.67		94.5
CCB3				0.001	21:08	-0.025		

Reviewed by: Diana H. H. H.

Date: 3-15-91

Prep Analyst: M. S. H.

Analyst: M. S. H.

* STD # 379

+ STD # 382

BC-001 total Hg

2 0018

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MERCURY PREP and ANALYSIS

00

Prep date 14 MAR 91

Analysis Date 3-14-91

Bath Temp 95°C

Lamp Energy 61

Spike Conc. 1 ppb Vendor MALINCKRODT

Corr Coeff. 0.9991

Sample ID #	Weight	Volume	Dil. factor	ABS.	Time	Value from curve	Reported Value	% Recovery
GN-163-PBW				0.001	21:09	-0.025		
GN-163-163		100 mL		0.005	21:10	0.128		
GN-165-164				0.010	21:12	3.76		
GN-165				0.001	21:14	-0.025		
GN-166				0.002	21:15	0.013		
GN-167				0.005	21:16	0.128		
GN-168				0.005	21:18	0.128		
GN-168 D				0.005	21:19	0.128		
GN-168 S		↓		0.024	21:21	1.836		
CCV4				0.151	21:22	5.709		95.0
CCB4				0.004	21:24	0.089		
✓ T186-271-1 (5X)		100 mL	5X	0.001	21:25	-0.025	<0.0025	
-1 S				0.097	21:27	3.645		91%
✓ T186-272-1				0.000	21:28	-0.064	<0.0025	
-1 S				0.099	21:30	3.722		93%
✓ T381-733-1				0.001	21:31	-0.025	<0.0025	
-1 S				0.106	21:33	3.989		99%
✓ T747-28-1				0.001	21:35	-0.025	<0.0025	
-1 S	↓	↓	↓	0.097	21:36	3.645		91%
✓ CCV5				0.150	21:38	5.671		94.5
CCB5				0.003	21:39	0.051		
T186-271-1 (5X)		100 mL	5X	0.000	21:40	-0.064	<0.0025	
-1 S	↓		↓	0.013	21:41	4.773	8.93	10.8%
✓ 860-110-2				-0.001	21:43	-0.446	<0.0005	
-2 O				0.000	21:44	-0.064	↓	0% = 0.0
-2 S		↓		0.020	21:45	0.701		70%
✓ CCV6				0.145	21:48	5.480		91.3

Reviewed by: Dianna Hunder

Prep Analyst: MBJ

Date: 3-15-91

Analyst: MBJ 249

+ STO # 382

2 0019

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0017.1

Analysis Date 3-14-91

Lamp Energy 41

Vendor MALLINCKRODT

Corr Coeff. 0.99991

+

Prep Analyst: MSG

Analyst: Mr. ARB 9250

2 0020

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06.75

Corr Coeff. 0.99998

2 0021

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00173

Analysis Date 19 MAR 91

Lamp Energy 61

Corr Coeff.	<u>0.9990</u>
-------------	---------------

~~AR 309262~~

Prep Analyst: MBR

Analyst: MBW

igneous

unfiltered sample @ C.

BC-001

2 0022

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Balance serial #05761
 Analyst: Robert H. Mader
 Date: March 05, 1991

Book # 3
 Page # 103

1:1 H₂O₂ lot # 044052, conc. H₂O₂ Reg. # 6
 1:1 HCl lot # D34044
 #H₂O₂ 311 # H₂O₂ 103 # 103 conc. H₂O₂ 100-5 01 H₂O in tagline
 @ 103 311 # 264 addend

Boxed 64-4

IRA Sample #	EPA Sample #	Matrix	pH	Volume/Weight	Final Volume	Method H or N	Filtered	Color		Clarity		Texture / Artifacts
								Before	After	Before	After	
2-2-6-2A	EC003	H ₂ O	2	100.0	100.0	HCl	X	colorless	yellow	clear	clear	X
-3A	EC003		2				yes	colorless	yellow	clear	clear	X
-4B	EC004		2				yes	colorless	colorless	clear	clear	X
Dip -4B	EC0040		2				yes	colorless	colorless	clear	clear	X
Spl -4B	EC0045		2				yes	colorless	colorless	clear	clear	X
-5A	EC005		2				X	colorless	colorless	clear	clear	X
Dip -5A	EC0050		2				X	colorless	colorless	clear	clear	X
Spl -5A	EC0055		2				X	colorless	colorless	clear	clear	X
-6D	EC006		2				yes	colorless	colorless	clear	clear	X
-7A	EC007		2				X	colorless	colorless	clear	clear	X
-8B	EC008		2				X	colorless	colorless	clear	clear	X
-9A	EC009		2				X	colorless	colorless	clear	clear	X
-10B	EC010		2				yes	colorless	colorless	clear	clear	X
-11B	EC011		2				yes	colorless	colorless	clear	clear	X
-12B	EC012		2				yes	colorless	yellow	cloudy	clear	X
-25A	EC025		2				yes	colorless	colorless	cloudy	clear	X
-26D	EC026		2				yes	colorless	colorless	cloudy	clear	X
-27C	EC027		2				yes	colorless	colorless	clear	clear	X
EC028	EC028		X				X	X	X	X	X	X
EC029	EC029		X				X	X	X	X	X	X
EC030	EC030		X				X	X	X	X	X	X
EC031	EC031		X				X	X	X	X	X	X
EC032	EC032		X				X	X	X	X	X	X
EC033	EC033		X				X	X	X	X	X	X
EC034	EC034		X				X	X	X	X	X	X
EC035	EC035		X				X	X	X	X	X	X
EC036	EC036		X				X	X	X	X	X	X
EC037	EC037		X				X	X	X	X	X	X
EC038	EC038		X				X	X	X	X	X	X
EC039	EC039		X				X	X	X	X	X	X
EC040	EC040		X				X	X	X	X	X	X
EC041	EC041		X				X	X	X	X	X	X
EC042	EC042		X				X	X	X	X	X	X
EC043	EC043		X				X	X	X	X	X	X
EC044	EC044		X				X	X	X	X	X	X
EC045	EC045		X				X	X	X	X	X	X
EC046	EC046		X				X	X	X	X	X	X
EC047	EC047		X				X	X	X	X	X	X
EC048	EC048		X				X	X	X	X	X	X
EC049	EC049		X				X	X	X	X	X	X
EC050	EC050		X				X	X	X	X	X	X
EC051	EC051		X				X	X	X	X	X	X
EC052	EC052		X				X	X	X	X	X	X
EC053	EC053		X				X	X	X	X	X	X
EC054	EC054		X				X	X	X	X	X	X
EC055	EC055		X				X	X	X	X	X	X
EC056	EC056		X				X	X	X	X	X	X
EC057	EC057		X				X	X	X	X	X	X
EC058	EC058		X				X	X	X	X	X	X
EC059	EC059		X				X	X	X	X	X	X
EC060	EC060		X				X	X	X	X	X	X
EC061	EC061		X				X	X	X	X	X	X
EC062	EC062		X				X	X	X	X	X	X
EC063	EC063		X				X	X	X	X	X	X
EC064	EC064		X				X	X	X	X	X	X
EC065	EC065		X				X	X	X	X	X	X
EC066	EC066		X				X	X	X	X	X	X
EC067	EC067		X				X	X	X	X	X	X
EC068	EC068		X				X	X	X	X	X	X
EC069	EC069		X				X	X	X	X	X	X
EC070	EC070		X				X	X	X	X	X	X
EC071	EC071		X				X	X	X	X	X	X
EC072	EC072		X				X	X	X	X	X	X
EC073	EC073		X				X	X	X	X	X	X
EC074	EC074		X				X	X	X	X	X	X
EC075	EC075		X				X	X	X	X	X	X
EC076	EC076		X				X	X	X	X	X	X
EC077	EC077		X				X	X	X	X	X	X
EC078	EC078		X				X	X	X	X	X	X
EC079	EC079		X				X	X	X	X	X	X
EC080	EC080		X				X	X	X	X	X	X
EC081	EC081		X				X	X	X	X	X	X
EC082	EC082		X				X	X	X	X	X	X
EC083	EC083		X				X	X	X	X	X	X
EC084	EC084		X				X	X	X	X	X	X
EC085	EC085		X				X	X	X	X	X	X
EC086	EC086		X				X	X	X	X	X	X
EC087	EC087		X				X	X	X	X	X	X
EC088	EC088		X				X	X	X	X	X	X
EC089	EC089		X				X	X	X	X	X	X
EC090	EC090		X				X	X	X	X	X	X
EC091	EC091		X				X	X	X	X	X	X
EC092	EC092		X				X	X	X	X	X	X
EC093	EC093		X				X	X	X	X	X	X
EC094	EC094		X				X	X	X	X	X	X
EC095	EC095		X				X	X	X	X	X	X
EC096	EC096		X				X	X	X	X	X	X
EC097	EC097		X				X	X	X	X	X	X
EC098	EC098		X				X	X	X	X	X	X
EC099	EC099		X				X	X	X	X	X	X
EC100	EC100		X				X	X	X	X	X	X

Reference serial # 05761

Analyst: Bobo Turner

Date: 3-19-91

Batch 78-1

+ 0.5 ml std 204 added

conc. HNO_3 R6
1:1 HNO_3 R6
1:1 HCl L2 #
D11010

Book # 4

Page # 6

EPA Sample #	EPA Sample #	Matrix	pH	Volume/Weight	Final Volume	Method H or N	Filtered	Color		Clarity		Texture/Artifacts
								Before	After	Before	After	
668-6-60	BC-0008	H ₂ O	2	50ml	50ml	HCl	X	colorless	colorless	clear	clear	
668-6-60	BC-0006.D		2				X	colorless	colorless	clear	clear	
668-6-60	BC-0006.5		2				X	colorless	colorless	clear	clear	
RLC5W	78-1		X				X					
PAW	78-1		X				X					

AR309254

2 0024

X 2 ml std # 155, 0.5 ml conc HNO_3 , Teflon Beaker used

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U.S. EPA - CLP
13
PREPARATION LOG

Lab Name: _____ Contract: 868-6

Lab Code: Meguna Case No.: SAS No.: SDG No.: BC-001

Method: CV

[illegible]

BC-019 - unfiltered inorganic QC
BC-023 - unfiltered inorganic QC
BC-016 - filtered inorganic QC

FORM XIII - IN

3/90

BC-021 - unfiltered total OC
BC-022 - filtered total OC

AR309255

2 0025

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868-4

118
PA

BL 013

2

BL 014

2

BL 015

2

BL 016

2

BL 017

2

BL 018

2

BL 019

2

BL 020

2

BL 021

2

BL 022

2

BL 023

2

BL 024

2

BL 027

2

AR309256

2 0026

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an environmental testing company

P.O. Box 12848

Research Triangle Park, North Carolina 27709

(919) 877-0090

FAX (919) 877-0427

April 3, 1991

APR 4 1991

Peg Bonaker
BCM Engineers, Inc.
One Plymouth Meeting Hall
Plymouth Meeting, PA 19462

M. Bonaker
A. Davis

Reference IEA Report No.: 868006(0), 868007, & 868008
Project ID: BCM 00-6012-02

Dear Ms. Bonaker,

Transmitted herewith are the results of analyses on 19 samples submitted to our laboratory.

Please see the enclosed reports for your results.

Very truly yours,

INDUSTRIAL & ENVIRONMENTAL ANALYSTS, INC.

Lillian L. Neme

Linda F. Mitchell
Director, Technical Support Services

State Certification:

Alabama - #40210
Georgia - #816
Kansas - #E-158

New Jersey - #67719
Tennessee - #00296
Virginia - #00179

South Carolina - #99021
North Carolina - #80900

184

Monroe,
Connecticut
203-261-4494

Miramar,
Florida
305-998-0828

Schaumburg,
Illinois
708-708-0740

N. Billerica,
Massachusetts
617-872-8212

Whippany,
New Jersey
201-438-6181

Essex Junction,
Vermont
802-878-6136

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IEA LABORATORY RESULTS

IEA Project #: 868-006(0)
Client Name: BCM Engineers, Inc.

Sample #	Client ID	Parameter	Results	Date Analyzed
1	26A	Ammonia-N	<0.01 mg/L	03/12/91
2	33A	Ammonia-N	0.1 mg/L	03/12/91
3	33AD	Ammonia-N	<0.01 mg/L	03/12/91
4	Equipment Blank	Ammonia-N	<0.01 mg/L	03/12/91
1	26A	Biochemical Oxygen Demand	<2.0 mg/L	02/20/91
2	33A	Biochemical Oxygen Demand	<2.0 mg/L	02/20/91
3	33AD	Biochemical Oxygen Demand	<2.0 mg/L	02/20/91
4	Equipment Blank	Biochemical Oxygen Demand	<2.0 mg/L	02/20/91
1	26A	Chloride	15 mg/L	03/01/91
2	33A	Chloride	14 mg/L	03/01/91
3	33AD	Chloride	14 mg/L	03/01/91
4	Equipment Blank	Chloride	<1.0 mg/L	03/01/91
1	26A	Chemical Oxygen Demand	32 mg/L	03/15/91
2	33A	Chemical Oxygen Demand	<25 mg/L	03/15/91
3	33AD	Chemical Oxygen Demand	<25 mg/L	03/15/91
4	Equipment Blank	Chemical Oxygen Demand	<25 mg/L	03/15/91
1	26A	Nitrite	0.02 mg/L	02/21/91
2	33A	Nitrite	<0.02 mg/L	02/21/91
3	33AD	Nitrite	<0.02 mg/L	02/21/91
4	Equipment Blank	Nitrite	<0.02 mg/L	02/21/91
1	26A	Nitrate	<0.02 mg/L	02/21/91
2	33A	Nitrate	10 mg/L	02/21/91
3	33AD	Nitrate	7.4 mg/L	02/21/91
4	Equipment Blank	Nitrate	<0.02 mg/L	02/21/91
1	26A	Sulfate	46 mg/L	03/11/91
2	33A	Sulfate	46 mg/L	03/11/91
3	33AD	Sulfate	43 mg/L	03/11/91
4	Equipment Blank	Sulfate	<3.0 mg/L	03/11/91
1	26A	Sulfide	<1.0 mg/L	02/26/91
2	33A	Sulfide	<1.0 mg/L	02/26/91
3	33AD	Sulfide	1.2 mg/L	02/26/91
4	Equipment Blank	Sulfide	<1.0 mg/L	02/26/91

AR309258

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METHOD BLANK SUMMARY

Parameter: Ammonia
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.1 mg/L	03/12/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM MELKSUM Rev 022790

AR309259

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

D

IEA Project No.: 866-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
866-006(0)-1	Ammonia	<0.1	<0.1	0	EPA #350.3

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

866-006(0)-1
866-006(0)-2
866-006(0)-3
866-006(0)-4

FORM IQCSUM Rev 062990

AR309260

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-006(0)-1	03/12/91	Ammonia	20	<0.1	19	95	EPA #350.3

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM QCSFK Rev 082890

AR309261

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METHOD BLANK SUMMARY

Parameter: Biochemical Oxygen Demand
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	2.0 mg/L	02/20/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM MBLSUM Rev 022790

AR309262

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
296-236-1	BOD	680	790	15	Std. Method #507

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM IQCSUM Rev 082990

AR309263

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METHOD BLANK SUMMARY

Parameter: Chloride
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	1.0 mg/L	03/01/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

AR309264

FORM MBLKSUM Rev 022790

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-006-4	Chloride	8.0	8.0	0	EPA #325.3

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM IQCSUM Rev 082990

AR309265

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-008-4	03/01/91	Chloride	50	8.0	58	100	EPA #325.3

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM QCSFK Rev 082890

AR309266

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METHOD BLANK SUMMARY

Parameter: Chemical Oxygen Demand
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	25 mg/L	03/15/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM MELSUM Rev 022790

AR309267

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
860-006(0)-1	COD	32	36	12	EPA #410.4

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM IQCSUM Rev 082990

AR309268

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-006(0)-2	03/15/91	COD	250	<25	240	96	EPA #410.4

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM QCSPE Rev 082890

AR309269

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METHOD BLANK SUMMARY

Parameter: Nitrate/Nitrite
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.02 mg/L	02/21/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM MELKSUM Rev 022790

AR309270

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
860-006(0)-3	Nitrite/Nitrate	7.4	7.2	3	EPA #353.2

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM IQCSUM Rev 082990

AR30927

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)			%R	METHOD
			SA	SR	SSR		
196-221-2	02/21/91	Nitrate/Nitrite	0.25	0.03	0.27	96	EPA #353.2

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM QCSPK Rev 082890

AR309272

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METHOD BLANK SUMMARY

Parameter: Sulfate
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	3.0 mg/L	03/11/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM MBLXSUM Rev 022790

AR309273

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
860-006(0)-3	Sulfate	43	46	7	Std. Methods #426C

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM IQCSUM Rev 082990

AR309274

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INORGANIC QC SUMMARY
SPIKE RESULTS

IFA Project No.: 868-006(0)
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-006(0)-1	03/11/91	Sulfate	*	46	*	*	Std. Methods #426C

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

*Percent recovery not calculated due to the sample concentration being greater than four times the spiking concentration.

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM QCSFK Rev 082890

AR309275

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METHOD BLANK SUMMARY

Parameter: Sulfide
IEA Project No.: 868-006(0)

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	1.0 mg/L	02/26/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM MBLXSUM Rev 022790

AR309276

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
860-006(0)-1	Sulfide	<1.0	<1.0	0	EPA #376.1

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-006(0)-1
868-006(0)-2
868-006(0)-3
868-006(0)-4

FORM IQCSUM Rev 082990

AR309277

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868-00614)

Sample Management Chain of Custody

26A

CC/Khachan

21 20/

Bottle Letters Available:

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

In it

RF

3/27/90

Date _____

REV.

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Case No. : 868-006 (P)

Sample Management Chain of Custody

EP A ID: 33A

Log-in By: CBK

Date: 2/20/91

Bottle Letters Available: 2A-H

Notes: *Applicable Codes Are

ST	=	Storage
TR	=	Transfer
EX	=	Extraction
DI	=	Dispose

*Refer to SAM generated Work List for Bottle definition.

Verified by:

Init

Date _____

REV. 10/50

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Case No. :

Sample Management Chain of Custody

ΕΡΑ ΙΟ:

33AD

Log-in By:

CP/Kutub

Date: 2/20/91

Bottle Letters Available: 3A-H

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

Date _____

AR30928

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Case No. :

868-006 (4)

Sample Management Chain of Custody

ΕΡΑ ΙΘ:

Equipment Blank

Log-in By:

CC/K

Date: 2/20/91

Bottle Letters Available: 4A-H

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definition

Verified by:

In it

Date _____

REV. 10/90

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-1	BOD	<2.0	<2.0	0	Std. Methods #507

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM IQCSUM Rev 082990

AR309283

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METHOD BLANK SUMMARY

Parameter: Biochemical Oxygen Demand
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	2.0 mg/L	02/22/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM MBLSUM Rev 022790

AR309284

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IEA LABORATORY RESULTS

IEA Project #: 868-007
Client Name: BCM Engineers, Inc.

Sample #	Client ID	Parameter	Results	Date Analyzed
1	9A	Ammonia-N	<0.01 mg/L	03/12/91
2	39A	Ammonia-N	0.2 mg/L	03/12/91
3	9AD	Ammonia-N	<0.01 mg/L	03/12/91
4	9B	Ammonia-N	<0.01 mg/L	03/12/91
5	22A	Ammonia-N	<0.01 mg/L	03/12/91
6	MWI-1-43	Ammonia-N	<0.01 mg/L	03/12/91
7	5A	Ammonia-N	<0.01 mg/L	03/12/91
8	5B	Ammonia-N	<0.01 mg/L	03/12/91
9	Equipment Blank	Ammonia-N	<0.01 mg/L	03/12/91
10	48A	Ammonia-N	<0.01 mg/L	03/12/91
1	9A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
2	39A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
3	9AD	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
4	9B	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
5	22A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
6	MWI-1-43	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
7	5A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
8	5B	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
9	Equipment Blank	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
10	48A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
1	9A	Chloride	15 mg/L	03/01/91
2	39A	Chloride	150 mg/L	03/01/91
3	9AD	Chloride	15 mg/L	03/01/91
4	9B	Chloride	13 mg/L	03/01/91
5	22A	Chloride	15 mg/L	03/01/91
6	MWI-1-43	Chloride	17 mg/L	03/01/91
7	5A	Chloride	3.0 mg/L	03/01/91
8	5B	Chloride	9.0 mg/L	03/01/91
9	Equipment Blank	Chloride	<1.0 mg/L	03/01/91
10	48A	Chloride	8.0 mg/L	03/12/91
1	9A	Chemical Oxygen Demand	<25 mg/L	03/12/91
2	39A	Chemical Oxygen Demand	<25 mg/L	03/12/91
3	9AD	Chemical Oxygen Demand	<25 mg/L	03/12/91
4	9B	Chemical Oxygen Demand	<25 mg/L	03/12/91
5	22A	Chemical Oxygen Demand	<25 mg/L	03/12/91
6	MWI-1-43	Chemical Oxygen Demand	<25 mg/L	03/12/91
7	5A	Chemical Oxygen Demand	<25 mg/L	03/12/91
8	5B	Chemical Oxygen Demand	<25 mg/L	03/12/91
9	Equipment Blank	Chemical Oxygen Demand	<25 mg/L	03/12/91
10	48A	Chemical Oxygen Demand	<25 mg/L	03/12/91
1	9A	Nitrite	<0.02 mg/L	02/22/91
2	39A	Nitrite	0.05 mg/L	02/22/91
3	9AD	Nitrite	<0.02 mg/L	02/22/91
4	9B	Nitrite	<0.02 mg/L	02/22/91
5	22A	Nitrite	<0.02 mg/L	02/22/91

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IEA LABORATORY RESULTS

IEA Project #: 868-007
Client Name: BCM Engineers, Inc.

Sample #	Client ID	Parameter	Results	Date Analyzed
6	MWI-1-43	Nitrite	<0.02 mg/L	02/22/91
7	5A	Nitrite	<0.02 mg/L	02/22/91
8	5B	Nitrite	<0.02 mg/L	02/22/91
9	Equipment Blank	Nitrite	<0.02 mg/L	02/22/91
1	9A	Nitrate	1.6 mg/L	02/22/91
2	39A	Nitrate	0.35 mg/L	02/22/91
3	9AD	Nitrate	1.6 mg/L	02/22/91
4	9B	Nitrate	11 mg/L	02/22/91
5	22A	Nitrate	9.0 mg/L	02/22/91
6	MWI-1-43	Nitrate	9.4 mg/L	02/22/91
7	5A	Nitrate	0.61 mg/L	02/22/91
8	5B	Nitrate	3.9 mg/L	02/22/91
9	Equipment Blank	Nitrate	<0.02 mg/L	02/22/91
1	9A	Sulfate	39 mg/L	03/13/91
2	39A	Sulfate	<3.0 mg/L	03/13/91
3	9AD	Sulfate	38 mg/L	03/13/91
4	9B	Sulfate	48 mg/L	03/13/91
5	22A	Sulfate	24 mg/L	03/13/91
6	MWI-1-43	Sulfate	3.7 mg/L	03/13/91
7	5A	Sulfate	50 mg/L	03/13/91
8	5B	Sulfate	22 mg/L	03/13/91
9	Equipment Blank	Sulfate	<3.0 mg/L	03/13/91
10	48A	Sulfate	26 mg/L	03/13/91
1	9A	Sulfide	<1.0 mg/L	02/26/91
2	39A	Sulfide	<1.0 mg/L	02/26/91
3	9AD	Sulfide	<1.0 mg/L	02/26/91
4	9B	Sulfide	<1.0 mg/L	02/26/91
5	22A	Sulfide	<1.0 mg/L	02/26/91
6	MWI-1-43	Sulfide	<1.0 mg/L	02/26/91
7	5A	Sulfide	<1.0 mg/L	02/26/91
8	5B	Sulfide	<1.0 mg/L	02/26/91
9	Equipment Blank	Sulfide	<1.0 mg/L	02/26/91
10	48A	Sulfide	<1.0 mg/L	02/26/91

AR309286

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METHOD BLANK SUMMARY

Parameter: Ammonia
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.1 mg/L	03/12/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM MBLKSUM Rev 022790

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-1	Ammonia	<0.1	<0.1	0	EPA #350.3

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM IQCSUM Rev 082990

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-007
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-007-1	03/12/91	Ammonia	20	<0.1	18	90	EPA #350.3

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM QCSFK Rev 082890

AR309289

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METHOD BLANK SUMMARY

Parameter: Chloride
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	1.0 mg/L	03/01/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM MBLKSUM Rev 022790

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-1	Chloride	15	15	0	EPA #325.3

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	
868-007-4	
868-007-5	

FORM IQCSUM Rev 082990

AR309291

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-007
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-007-1	03/01/91	Chloride	50	15	65	100	EPA #325.3

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-007-1 868-007-6
868-007-2 868-007-7
868-007-3
868-007-4
868-007-5

FORM QCSPE Rev 082890

AR309292

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-8	Chloride	9.0	10	10	EPA #325.3

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-8
868-007-9
868-007-10

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AR3092

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-007
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)			%R	METHOD
			SA	SR	SSR		
868-007-8	03/01/91	Chloride	50	9.0	60	102	EPA #325.3

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-007-8
868-007-9
868-007-10

FORM QCSFK Rev 082890

AR309294

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METHOD BLANK SUMMARY

Parameter: Nitrate
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.02 mg/L	02/22/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	

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AR309295

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-8	Nitrate	3.9	3.9	0	EPA #353.2

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	

FORM IQCSUM Rev 082990

AR309296

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-007
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SA	SPIKE RESULTS (mg/L)			METHOD
				SR	SSR	%R	
868-007-4	02/22/91	Nitrate	*	11	*	*	EPA #353.2

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

*Percent recovery not calculated due to the sample concentration being greater than four times the concentration of the spiking solution.

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	

FORM QCSFK Rev 082890

AR309297

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METHOD BLANK SUMMARY

Parameter: Nitrite
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.02 mg/L	02/22/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	

FORM MELKSUM Rev 022790

AR309298

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-8	Nitrite	<0.02	<0.02	0	EPA #353.2

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	

FORM IQCSUM Rev 082990

AR309299

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-007
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SA	SPIKE RESULTS (mg/L)		%R	METHOD
				SR	SSR		
868-007-4	02/22/91	Nitrite	0.25	<0.02	0.24	96	EPA #353.2

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

$$\%R = (SSR - SR) / (SA) * 100$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	

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AR309300

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METHOD BLANK SUMMARY

Parameter: Sulfate
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	3.0 mg/L	03/13/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

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AR309301

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-1	Sulfate	39	37	5	Std. Methods #426C

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM IQCSUM Rev 082990

AR309302

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-007
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-007-4	03/13/91	Sulfate	10	12	22	100	Std. Methods #426C

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM QCSFK Rev 082890

AR309303

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METHOD BLANK SUMMARY

Parameter: Sulfide
IEA Project No.: 868-007

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	1.0 mg/L	02/26/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM MELSUM Rev 022790

AR309304

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-007
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-6	Sulfide	<1.0	<1.0	0	Std. Methods #426C

$$\text{RPD} = \frac{S-D}{(S+D)/2} \times 100$$

Corresponding Samples:

868-007-1	868-007-6
868-007-2	868-007-7
868-007-3	868-007-8
868-007-4	868-007-9
868-007-5	868-007-10

FORM IQCSUM Rev 082990

AR309305

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**ANALYSTS, INC.
3000 WESTON PKWY.
CARY, N.C. 27513**

REGULATORY CLASSIFICATION - PLEASE SPECIFY

☐ NPDES ☐ DRINKING WATER ☐ RCRA ☒ OTHER CERCLA/SARA

NO: 14019

PROJECT #		PROJECT NAME		MATRIX		CONTAINERS		REQUESTED PARAMETERS		IEA RUSH NO.			
SAMPLE ID	DATE	TIME	STATION LOCATION	#	OF	SOIL	WATER	17 PL	18 PL	19 PL	20 PL		
00-602-02	2/20/01	0815	39A	12	X			1	3	4	2		
		0155	9A	28				2	6	8			
		0165	9AD	16				1	3	4			
		1205	9B	16				1	3	4			
		1405	22A	8				1	3	4			
		1410	MWI-1-43	8				1	3	4			
		1550	5A	8				1	3	4			
		1535	5B	8				1	3	4			
		1655	48A	6				1	3				
		1730	Trp Bence	4									
		1110	2quipped Bence	16				1	3	4	2		
RECEIVED BY				DATE		TIME		IEA QUOTE NO.				IEA RUSH NO.	
Russell McBeth				2/21/01		1500		Security Treatment					
RECEIVED FOR LAB BY				DATE		TIME		PROJECT MANAGER (PLEASE PRINT)				P.O. NO.	
Russell McBeth				2/21/01		1830		C. K. K.					
IEA REMARKS				IEA REMARKS		IEA REMARKS		IEA REMARKS		IEA REMARKS		IEA REMARKS	
IEA # 805-006-CLP				IEA # 805-007-04N		IEA # 805-008-04N		IEA # 805-009-04N		IEA # 805-010-04N		IEA # 805-011-04N	

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Case No. : 868-007

Sample Management Chain of Custody

ΕΡΑ ΙΟ:

9A

Log-in By:

CDK 1

Date: 2/21/91

Bottle Letters Available: 1A-P

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

3/27/90

Date _____

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Case No. : 868-007

Sample Management Chain of Custody

EPÀ ID: 394

Log-in By: CDK

Date: 2/21/91

Bottle Letters Available: 2A-14

[illegible]

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
OI = Dispose

AR309308

*Refer to SAM generated Work List for Bottle definitions

Verified by:

In it

527 HC

Date

REV. 10/50

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Case No. : 868-007

Sample Management Chain of Custody

EPA ID:

9AD

Log-in By:

Date: 2/21/91

Bottle Letters Available: 3A - H

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose

AR309300

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Int

3/27/90

Date _____

REV. 10

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868-007

Sample Management Chain of Custody

22A

Log-in By:

Date:

2/21/11

Bottle Letters Available: 5A - H

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

Date:

REV. to,

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Case No. : 868-007

Sample Management Chain of Custody

EPA ID: MWI-1-43

Log-in By: Chandra

Date: 2/21/91

Bottle Letters Available: GA-H

[illegible]

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

AR309312

Date _____

REV. 10/90

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Case No. :

868-007

Sample Management Chain of Custody

ΕΡΑ ID:

5A

Log-in By:

C. B. Knecht

Date:

2 / 21 /

Bottle Letters Available:

7A-H

[illegible]

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DX = Dispose

***Refer to SAM generated Work List for Bottle definitions**

Verified by:

In it

Date _____

AR309313

REV. to

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Case No. :

868-007

Sample Management Chain of Custody

ΕΡΑ ΙΟ:

5B

Log-in By:

CQK.L

Date: 2/21/91

Bottle Letters Available: 8A-H

[illegible]

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

In it

AR309314

3/27/90

Date _____

REV. 10/50

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Case No. : 868-007

Sample Management Chain of Custody

EPA ID: Equipment Blank

Log-in By: CDK

Date: 2/2/14

Bottle Letters Available: 9A -H

[illegible]

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose'

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Info

Date _____

AR 309315

REV. to/

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Case No. :

868-007

Sample Management Chain of Custody

ΕΡΑ ΙΟ:

48A

Log-in By:

CD Krishna

Date:

2 1 2 1 9 1

Bottle Letters Available: 10A-F

10A-F

[illegible]

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definition

Verified by:

Init

Date

1A0309316

3/27/90

REV. 10/90

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IEA LABORATORY RESULTS

IEA Project #: BCM Engineers, Inc.
Client Name: 868-008

Sample #	Client ID	Parameter	Results	Date Analyzed
2	24A	Ammonia-N	<0.01 mg/L	03/12/91
3	Field Blank	Ammonia-N	<0.01 mg/L	03/12/91
4	19A	Ammonia-N	0.3 mg/L	03/12/91
5	16A	Ammonia-N	0.1 mg/L	03/12/91
2	24A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
3	Field Blank	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
4	19A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
5	16A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
2	24A	Chloride	21 mg/L	03/01/91
3	Field Blank	Chloride	<1.0 mg/L	03/01/91
4	19A	Chloride	8.0 mg/L	03/01/91
5	16A	Chloride	5.0 mg/L	03/01/91
2	24A	Chemical Oxygen Demand	<25 mg/L	03/15/91
3	Field Blank	Chemical Oxygen Demand	<25 mg/L	03/15/91
4	19A	Chemical Oxygen Demand	<25 mg/L	03/15/91
5	16A	Chemical Oxygen Demand	<25 mg/L	03/15/91
1	48A	Nitrite	<0.02 mg/L	02/22/91
2	24A	Nitrite	<0.02 mg/L	02/22/91
3	Field Blank	Nitrite	<0.02 mg/L	02/22/91
4	19A	Nitrite	<0.02 mg/L	02/22/91
5	16A	Nitrite	<0.02 mg/L	02/22/91
1	48A	Nitrate	6.6 mg/L	02/22/91
2	24A	Nitrate	1.6 mg/L	02/22/91
3	Field Blank	Nitrate	<0.02 mg/L	02/22/91
4	19A	Nitrate	6.5 mg/L	02/22/91
5	16A	Nitrate	1.2 mg/L	02/22/91
2	24A	Sulfate	68 mg/L	03/11/91
3	Field Blank	Sulfate	<3.0 mg/L	03/11/91
4	19A	Sulfate	39 mg/L	03/11/91
5	16A	Sulfate	23 mg/L	03/11/91
2	24A	Sulfide	<1.0 mg/L	02/26/91
3	Field Blank	Sulfide	<1.0 mg/L	02/26/91
4	19A	Sulfide	<1.0 mg/L	02/26/91
5	16A	Sulfide	<1.0 mg/L	02/26/91

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IEA LABORATORY RESULTS

IEA Project #: 868-008
Client Name: BCM Engineers, Inc.

Sample #	Client ID	Parameter	Results	Date Analyzed
2	24A	Ammonia-N	<0.01 mg/L	03/12/91
3	Field Blank	Ammonia-N	<0.01 mg/L	03/12/91
4	19A	Ammonia-N	0.3 mg/L	03/12/91
5	16A	Ammonia-N	0.1 mg/L	03/12/91
2	24A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
3	Field Blank	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
4	19A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
5	16A	Biochemical Oxygen Demand	<2.0 mg/L	02/22/91
2	24A	Chloride	21 mg/L	03/01/91
3	Field Blank	Chloride	<1.0 mg/L	03/01/91
4	19A	Chloride	8.0 mg/L	03/01/91
5	16A	Chloride	5.0 mg/L	03/01/91
2	24A	Chemical Oxygen Demand	<25 mg/L	03/15/91
3	Field Blank	Chemical Oxygen Demand	<25 mg/L	03/15/91
4	19A	Chemical Oxygen Demand	<25 mg/L	03/15/91
5	16A	Chemical Oxygen Demand	<25 mg/L	03/15/91
1	48A	Nitrite	<0.02 mg/L	02/22/91
2	24A	Nitrite	<0.02 mg/L	02/22/91
3	Field Blank	Nitrite	<0.02 mg/L	02/22/91
4	19A	Nitrite	<0.02 mg/L	02/22/91
5	16A	Nitrite	<0.02 mg/L	02/22/91
1	48A	Nitrate	6.6 mg/L	02/22/91
2	24A	Nitrate	1.6 mg/L	02/22/91
3	Field Blank	Nitrate	<0.02 mg/L	02/22/91
4	19A	Nitrate	6.5 mg/L	02/22/91
5	16A	Nitrate	1.2 mg/L	02/22/91
2	24A	Sulfate	68 mg/L	03/11/91
3	Field Blank	Sulfate	<3.0 mg/L	03/11/91
4	19A	Sulfate	39 mg/L	03/11/91
5	16A	Sulfate	23 mg/L	03/11/91
2	24A	Sulfide	<1.0 mg/L	02/26/91
3	Field Blank	Sulfide	<1.0 mg/L	02/26/91
4	19A	Sulfide	<1.0 mg/L	02/26/91
5	16A	Sulfide	<1.0 mg/L	02/26/91

AR309318

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METHOD BLANK SUMMARY

Parameter: Ammonia
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.1 mg/L	03/12/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-2
868-008-3
868-008-4
868-008-5

AR309319

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-006(0)-1	Ammonia	<0.1	<0.1	0	EPA #350.3

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCSUM Rev 082990

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-008
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)			%R	METHOD
			SA	SR	SSR		
868-006(0)-1	03/12/91	Ammonia	20	<0.1	19	95	EPA 350.3

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

$$\%R = (SSR - SR) / (SA) * 100$$

Corresponding Samples:

868-008-2
868-008-3
869-008-4
868-008-5

FORM QCSFK Rev 082890

AR30932

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METHOD BLANK SUMMARY

Parameter: Biochemical Oxygen Demand
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	2.0 mg/L	02/22/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-2
868-008-3
868-008-4
868-008-5

FORM MELKSUM Rev 022790

AR309322

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-008-2	BOD	<2.0	<2.0	0	Std. Methods #507

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCSUM Rev 1182990

AR309323

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METHOD BLANK SUMMARY

Parameter: Chloride
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	1.0 mg/L	03/01/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-2
868-008-3
868-008-4
868-008-5

FORM MBLKSUM Rev 022790

AR309324

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-008-4	Chloride	8.0	8.0	0	EPA #325.3

$$\text{RPD} = \frac{S-D}{(S+D)/2} \times 100$$

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCSUM Rev 082990

AR309325

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-008
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SA	SPIKE RESULTS (mg/L)			METHOD
				SR	SSR	%R	
868-008-4	03/01/91	Chloride	50	8.0	58	100	EPA #325.3

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM QCSFK Rev 082890

AR309326

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METHOD BLANK SUMMARY

Parameter: Chemical Oxygen Demand
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	25 mg/L	03/15/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-2
868-008-3
868-008-4
868-008-5

AR309327

FORM MELKSUM Rev 022790

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
860-008-4	COD	<25	<25	0	EPA #410.4

$$\text{RPD} = \frac{S-D}{(S+D)/2} \times 100$$

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCSUM Rev 082990

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-008
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SA	SPIKE RESULTS (mg/L)			METHOD
				SR	SSR	%R	
868-008-5	03/15/91	COD	250	<25	250	100	EPA #410.4

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM QCSFK Rev 082890

AR309383

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METHOD BLANK SUMMARY

Parameter: Nitrate
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.02 mg/L	02/22/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-1
868-008-2
868-008-3
868-008-4
868-008-5

FORM MBLSUM Rev 022790

AR309330

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-008-1	Nitrate	6.6	6.6	0	EPA #353.2

$$\text{RPD} = \frac{S-D}{(S+D)/2} \times 100$$

Corresponding Samples:

868-008-1
868-008-2
868-008-3
868-008-4
868-008-5

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-008
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SA	SPIKE RESULTS (mg/L)			METHOD
				SR	SSR	%R	
868-008-5	02/22/91	Nitrate	0.25	1.2	1.5	120	EPA #353.2

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-008-1
868-008-2
868-008-3
868-008-4
868-008-5

FORM QCSFK Rev 082890

AR309332

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METHOD BLANK SUMMARY

Parameter: Nitrite
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	0.02 mg/L	02/22/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-1
868-008-2
868-008-3
868-008-4
868-008-5

FORM MBLKSUM Rev 022790

AR30933

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-008-1	Nitrite	<0.02	<0.02	0	EPA #353.2

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-008-1
868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCSUM Rev 082990

AR309334

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-008
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SA	SPIKE RESULTS (mg/L)		%R	METHOD
				SR	SSR		
868-008-5	02/22/91	Nitrite	0.25	<0.02	0.24	96	EPA #353.2

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-008-1
868-008-2
868-008-3
868-008-4
868-008-5

FORM QCSFK Rev 082890

AR309335

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METHOD BLANK SUMMARY

Parameter: Sulfate
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	3.0 mg/L	03/11/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-2
868-008-3
868-008-4
868-008-5

FORM MBLKSUM Rev 022790

AR309336

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-008-2	Sulfate	68	68	0	Std. Methods #426C

$$\text{RPD} = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCMM Rev 082990

AR309337

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INORGANIC QC SUMMARY
SPIKE RESULTS

IEA Project No.: 868-006(0)
Matrix: Water

IEA REF. NO.	ANALYSIS DATE	TEST PARAMETER	SPIKE RESULTS (mg/L)				METHOD
			SA	SR	SSR	%R	
868-006(0)-1	03/11/91	Sulfate	*	46	*	*	Std. Methods #426C

$$\%R = (SSR - SR) / (SA) * 100$$

SA = Spike Added
SR = Sample Results
SSR = Spike Sample Results

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM QCSFK Rev 082890

AR309338

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METHOD BLANK SUMMARY

Parameter: Sulfide
IEA Project No.: 868-008

BLANK RESULTS	QUANTITATION LIMIT	DATE ANALYZED
BQL	1.0 mg/L	03/26/91

BQL = Below Quantitation Limit

CORRESPONDING SAMPLES:

868-008-2
868-008-3
868-008-4
868-008-5

AR309339

FORM MBLSUM Rev 022790

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INORGANIC QC SUMMARY
DUPLICATE ANALYSIS

IEA Project No.: 868-008
Matrix: Water

IEA REFERENCE NO.	TEST PARAMETER	DUPLICATE RESULTS		RPD (%)	METHOD
		Sample (mg/L)	Duplicate (mg/L)		
868-007-6	Sulfide	<1.0	<1.0	0	Std. Methods #426C

$$RPD = \frac{S-D}{(S+D)/2 \times 100}$$

Corresponding Samples:

868-008-2
868-008-3
868-008-4
868-008-5

FORM IQCSUM Rev 082990

AR309340

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CHAIN OF CUSTODY RECORD

REGULATORY CLASSIFICATION - PLEASE SPECIFY

☐ NPDES ☐ DRINKING WATER ☐ RCRA ☒ OTHER CERCLA/SARA

NO: 12134

[illegible]

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Case No. :

828-008

Sample Management Chain of Custody

ΕΡΑ ID:

48A

Log-in By:

CAK

Date: 2/22/91

Bottle Letters Available: 1A-B

[illegible]

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

In it

3/27/90

Date _____

AR309342

REV. 10/50

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Case No.: 868-008

Sample Management Chain of Custody

EPA ID: 24A

Log-in By: Chakrabarti

Date: 2/22/91

Bottle Letters Available: 2A - H

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

In it

3/27/90
Date

AR309343

REV. 10/50

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Case No. : 868-008

Sample Management Chain of Custody

EPA ID: Field Blank

Log-in By: CEK

Date: 2/22/91

Bottle Letters Available: 3A - H

[illegible]

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

Date _____

REV. 10/50

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Case No. :

868-005

Sample Management Chain of Custody

ΕΡΑ ID:

19A

Log-in By:

CEK-ah

Date: 2/22/71

Bottle Letters Available:

44-44

[illegible]

Notes: *Applicable Codes Are

ST = Storage

TR = Transfer

EX = Extraction

DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

AR309345

3/27/90

Date _____

REV. 10/20

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Case No. : 868-008

Sample Management Chain of Custody

EPA ID:	16A
---------	-----

Log-in By: CA/K

Date: 2/22/91

Bottle Letters Available: SA - H

[illegible]

Notes: *Applicable Codes Are

ST = Storage
TR = Transfer
EX = Extraction
DI = Dispose

*Refer to SAM generated Work List for Bottle definitions

Verified by:

Init

Date _____

AR309346

3/27/90

REV. 10/50

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AR309347

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D

DATA VALIDATION REPORT
CHEM-SOLV, INC.
CHESWOLD, DELAWARE
GROUNDWATER SAMPLING EVENT

BCM PROJECT NO. 00-6012-02
APRIL, 1991

PREPARED BY

Atwood F. Davis
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AR309348

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DATA VALIDATION REPORT
INORGANIC DATA
GROUNDWATER SAMPLING EVENT
CHEM-SOLV INC.
CHESWOLD, DELAWARE

INTRODUCTION

This data validation report addresses inorganic data from the February 1991 groundwater sampling event at the Chem-Solv, Inc. site in Cheswold, Delaware. Case 868-006 contains six samples. Each has a filtered and unfiltered aliquot. Filtered and unfiltered samples 26A, 33A, 33AD, and 39A were analyzed for manganese. Filtered and unfiltered samples 33A and 33AD were analyzed for zinc. Filtered and unfiltered samples 9B and 9A were analyzed for both total and inorganic mercury. The sample set included two equipment blanks, filtered and unfiltered, and two unfiltered trip blanks. All samples were analyzed by a single subcontracted laboratory according to the United States Environmental Protection Agency (EPA) Contract Laboratory Program (CLP) Statement of Work (SOW) for Inorganic Analysis, July 1988, revisions 2/89 and 6/89.

These data were reviewed according to the EPA Laboratory Data Validation Functional Guidelines for Evaluating Inorganic Analyses, June 13, 1988, and EPA Region III Modifications to the Inorganic Functional Guidelines, June 1988. The review of the data included holding times, blank analysis results, initial and continuing calibrations, matrix spike results, laboratory and field duplicate results, detection limits, quantitative calculations, and a general review of analytical data. All data have been validated with regards to usability. Areas of concern with respect to usability are listed in the following section according to the seriousness of the problem. The data summary tables located at the end of this report have been annotated with the appropriate qualifier codes. Appendix A of this document summarizes the specifics of this review.

FINDINGS/QUALIFIERS

All Clean Water Act (40 CFR 136) holding times were met as specified. Laboratory and field blanks exhibited no contamination. Laboratory control samples and calibration verification samples were within control limits. Laboratory and field duplicates exhibited good reproducibility. Matrix spike recoveries were precise.

It is recommended that the data from Inorganic Case 868-006 be used with the following qualifier statement:

1. Filtered samples 33ADF (BC005), 33AF (BC007), and equipment blank 2/19/91 (BC009) are qualified as estimated for zinc due to serial dilution results that do not agree within 10 percent, indicating a physical or chemical interference or laboratory specific problem.

AR309349

BCM

It should be noted that unfiltered sample 98U was analyzed for both total and inorganic mercury. The inorganic mercury result of 2.6 ug/l was slightly higher than the total mercury result of 2.2 ug/l. A modified method 245.1 CLP-M was used to analyze for inorganic mercury. This method deleted certain steps in the digestion procedure that liberate the organic mercury and eliminate interferences. This was done to measure only the inorganic mercury. Since the complete digestion procedure was not performed there may be possible organic interferences causing the inorganic mercury result to be higher than the total mercury. Alternatively, the difference between the two results is so small that it may just represent the analytical variability of the method.

Filtered sample 33ADF (BC005) had a manganese result of 30.1 ug/l while the unfiltered sample 33ADU (BC004) had a manganese result of 27.3 ug/l.

The linear range for manganese and zinc was last determined in October 1990. Although this is non-compliant with SOM 788, which states a linear range analysis must be determined every 3 calendar months for each element, there is no significant impact on the data.

SUMMARY

The laboratory performed the analysis in compliance with CLP including the required quality control samples.

Holding times were met for all samples. Analyses of laboratory and field blanks showed no contamination. Laboratory control and calibration verification samples were within acceptable control limits. Laboratory and field duplicates exhibited good reproducibility. Matrix spike recoveries were within control limits. A serial dilution result was outside the control limit indicating a possible matrix interference.

All data should be used with the limitations imposed by the assigned qualifier. For specific review forms see Appendix A.

AR309350

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DATA SUMMARY

ECM

Page 1 of 3

Site Chem-Solv, Inc.

Case No. 868-006

Matrix Aqueous

Laboratory TEA

Units Mg/L

Date of Sample 2/19/91 + 2/20/91

LAB ID.	FIELD ID.	26AL	26AF	33ADU	33ADF	33AU	33AF	84BKA	84BKF	TOPBKA	TOPBKF
LAB NO.	LAB ID.	8C002	8C003	8C004	8C005	8C006	8C007	8C008	8C009	8C010	8C011
	Aluminum										
	Antimony										
	Arsenic										
	Barium										
	Beryllium										
	Cadmium										
	Calcium										
	Chromium										
	Cobalt										
	Copper										
	Iron										
	Lead										
	Magnesium										
15	Manganese	33402	20320	27.3	30.1	31.8	31.6				7108
0.2	Mercury										
	Nickel										
	Potassium										
	Selenium										
	Silver										
	Sodium										
	Thallium										
	Vanadium										
20	Zinc			3280	2180	3340	2240	770.5			770.4
	Cyanide										

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Matrix aqueousUnits me18

DATA SUMMARY

Chico-Sol Inc.

868-006

Laboratory IEA

Date of Sample 2/19/91 2/20/91

[illegible]

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BCM

DATA SUMMARY

Page 3 of 3
 Matrix Aqueous
 Units mg/L

Site Chem-Solv Inc.
 Case No. 868-006
 Laboratory IEA

Date of Sample 2/20/91

CRDL	LAB IDL	FIELD ID.	9AF	9AILL	9AIF	88CKU	88BKF	TBIRLL
	LAB IDL	LAB ID.	BC033	BC023	BC034	BC035	BC024	BC027
	Aluminum							
	Antimony							
	Arsenic							
	Barium							
	Beryllium							
	Cadmium							
	Calcium							
	Chromium							
	Cobalt							
	Copper							
	Iron							
	Lead							
	Magnesium							
15	Manganese							
0.2	Mercury							
	Nickel							
	Potassium							
	Selenium							
	Silver							
	Sodium							
	Thallium							
	Vanadium							
20	Zinc							
	Cyanide							

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ECM

QUALIFIER CODES

J - Estimated value

UJ - Estimated detection limit

AR309354

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ECM

DATA VALIDATION REPORT
VOLATILE ORGANICS DATA
GROUNDWATER SAMPLING EVENT
CHEM-SOLV INC.
CHESWOLD, DELAWARE

INTRODUCTION

This data validation report addresses volatile organic data from the February 1991 groundwater sampling event at the Chem-Solv, Inc. site in Cheswold, Delaware. Case 868-006 contains four samples, an equipment blank, and a trip blank. All samples were unfiltered and preserved with hydrochloric acid. The findings offered in this review are based upon a general review of sample data, analytical holding times, gas chromatography/mass spectrometry (GC/MS) tunes, initial and continuing calibrations, blank analysis results, matrix spike/matrix spike duplicate results (MS/MSD), field duplicate analysis results, surrogate spike results, internal standards performance, TCL volatile compound identifications, tentatively identified compound (TIC) identification, and quantitative calculations.

The samples in this report were analyzed by a subcontracted laboratory using the United States Environmental Protection Agency (EPA) Contract Laboratory Program (CLP) Statement of Work (SOW) for Organic Analyses, February 1988 with revisions 9/88 and 4/89. The quality assurance review was prepared in accordance with EPA Laboratory Data Validation Functional Guidelines for Evaluating Organic Analysis, February, 1988.

The support documentation appendix of this report summarizes the specifics of this review. The data summary lists only compound which were reported to be present. The constituents detected represent a small portion of the total analytical protocol. Analytical problems were minimal and are outlined in the following Findings/Qualifier Section.

FINDINGS/QUALIFIERS

No results were qualitatively questioned.

Clean Water Act (40 CFR 136) analytical holding times and CLP contract required holding times were met for all samples. GC/MS tune, internal standards performance, surrogate recovery, and MS/MSD results met criteria. Laboratory and field blanks were found to be free of contamination, the lab blank met CLP criteria for all target compounds. Target compound positive results were properly identified and quantitatively correct. Required quantitation limits were met. The field duplicate samples 33A (BC006) and 33AD (BC004) exhibited acceptable results.

AR309355

BCM

Initial calibration response factors, average response factors, and percent relative standard deviations were spot checked (all compounds with positive results), no errors were found. Continuing calibrations response factors and percent differences were spot checked (all compounds with positive results and all compounds with percent differences greater than 25 percent), no errors were found. Systems performance check compounds and continuing calibration compound met criteria for both initial and continuing calibrations.

It is recommended that the data from Organic Case 868-006 be used with the following qualifier statement:

1. Due to continuing calibration percent differences greater than 25 percent, quantitation limits for bromomethane, acetone, and 2-butanone are estimated in all samples.

It should be noted that this effect is probably minor for acetone (28 percent) and 2-butanone (42 percent due to low response factors), especially since these compounds are common laboratory artifacts which may also contaminate standards causing the variability noted in the response factors.

The reviewer noted that sample 26A (BC002) which exhibited a positive result for benzene (29 ug/l) was analyzed immediately after the MS/MSD analysis (to which benzene is spiked at 50 ug/l). There is a possibility that the benzene result in 26A (BC002) is due to carryover; however, the reviewer assessment is that benzene is present. Its presence is based upon the finding that similar volatile TICs were found to be present in 26A (BC002) and that the likelihood of a volatile carryover at greater than 50 percent of the spike component concentration is remote.

Sample 26A (BC002) was the only sample which exhibited TICs. The TIC spectra were reviewed and, in general, the reviewer concurred with the laboratory's interpretation. Reviewer comments on TIC interpretation are included in the attached support documentation appendix.

SUMMARY

The volatile organics analysis was performed in compliance with the referenced CLP SOW. All contractual/SOW criteria were met. No results were qualitatively questioned, no positive results were quantitatively questioned.

This review has identified continuing calibration percent differences as a concern for three compound quantitation limits. For specifics relating to this review, see the attached support documentation in Appendix B.

AR309356

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BCM Number 00-6012-02
SDG Number BC001 CASE 863-006

Site Name Chern Solv
Date of Sample 2-19-91

☒ Organic

Vowels Only

Compounds Detected

[illegible]

NOTE: For a review of this data and non-target, tentatively identified compounds, please see the Analytical Quality Assurance section of this report.

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QUALIFIER CODES

J - Estimated value below contract required quantitation limit.

/22100

AR309358

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DATA VALIDATION REPORT
WET CHEMISTRY DATA
GROUNDWATER SAMPLING EVENT
CHEM-SOLV., INC.
CHESWOLD, DELAWARE

INTRODUCTION

This data validation report addresses wet chemistry data from the February 1991 groundwater sampling event at the Chem-Solv., Inc. site in Cheswold, Delaware. Case 868-006 contained two groundwater samples, an equipment blank, and a field duplicate sample. Case 868-007 contained eight groundwater samples, an equipment blank, and a field duplicate sample. Case 868-008 contained three samples and an equipment blank. All samples were analyzed by a single subcontracted laboratory according to the United States Environmental Protection Agency (EPA) Methods for Chemical Analysis of Water and Wastes, March 1983.

These data have been reviewed according to EPA Laboratory Data Validation Functional Guidelines for Evaluating Inorganic Analyses, June 13, 1988, and EPA Region III Modifications to the Inorganic Functional Guidelines, June 1988. The review of the data included holding times, blank analysis results, laboratory and field duplicate results and matrix spike results. All data have been validated with regards to usability. Areas of concern with respect to usability are listed in the following section. The data summary tables located at the end of this report have been annotated with the appropriate qualifier codes.

FINDINGS/QUALIFIERS

All Clean Water Act (40 CFR 136) holding times were met as specified. Laboratory and equipment blanks exhibited no contamination. Laboratory duplicates showed good reproducibility. Matrix spike recoveries were precise.

It is recommended that the data from Case 868-006 be used with the following qualifier statement:

1. Nitrate results for field duplicate samples 33A and 33AD exceeded the 20 percent RPD acceptance limits for duplicate precision. Reported results are qualified as estimated. The reported results may not reflect the true average concentration.

It should be noted that the laboratory reported a quantitation limit of 0.01 mg/l for ammonia on the laboratory results form, while on the method blank summary sheet a quantitation limit of 0.1 mg/l for ammonia is reported.

AR309355

BCM

The laboratory did not include a method blank, duplicate analysis, or a matrix spike result summary for C.O.D. for Case 868-007.

The reported sample, blank, duplicate, and matrix spike results cannot be verified due to an incomplete data package.

SUMMARY

Holding times were met for all samples. Analyses of laboratory and equipment blanks exhibited no contamination. Laboratory duplicates showed good reproducibility. Matrix spike recoveries were within control limits.

/22100

AR309360

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QUALIFIER CODES

J - Estimated value

UJ - Estimated detection limit

AR30936

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Site Name Onion-Solv. Inc.
Date of Sample 2/14/91
Laboratory IEA

Method

IPA Number 805-0000

[illegible]

AR309362

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TSD Number BLB-007
 EPA Number BLB-007
 Site Name Chem-Solv, Inc.
 Date of Sample 2/22/91
 Laboratory ZBA

SAMPLE DATA SUMMARY

Sample #	Sample Description and Location	Phase	Units	AMMONIA-N					CHLORIDE					C.O.D					NITRITE					NITRATE					SULFATE					SULFIDE					Remarks	
				B.O.D	CHLORIDE	C.O.D	NITRITE	NITRATE	SULFATE	SULFIDE																														
9A			mg/L		15			1.6	39																															
39A			mg/L	0.2	150		0.05	0.50																																
9AD			mg/L		15			1.6	38																														field duplicate	
9B			mg/L		13			11	48																															
22A			mg/L		15			9	24																															
MWH-45			mg/L		17			9.4	3.7																															
5A			mg/L		3			0.21	50																															
5B			mg/L		9			3.9	22																															
42A			mg/L		8			6.6																																
Blk			mg/L						26																															equipment Blk 2/22/91

AR309383

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ANYTIME VJ

INQUEST VJ2

810-3-13

Slit Name Cheon-Silly Dox

Date of Sample 3/21/61

LEA

[illegible]

AR 809364

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BCM

APPENDIX A

AR309365

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BCM ENGINEERS, INC.
INORGANIC DATA VALIDATION SUMMARY
ICP ANALYSES

Project Name Chem - Solid - 200 Case No. 868-006
BCM Project No. 6012-02 Sampling Date(s) 2/19/91 + 2/20/91
Field ID 26AU, 26AF, 33ADU, 33ADF Date Review Completed 3/26/91
33AU, 33AF, Equip. BLK 42/19 + 2/20 Reviewer D. McGuire
Eg. BLK F 2/19 + 20, Trip BLK 42/19 + 20

MATRIX	CONCENTRATION			MATRIX RELATED COMMENTS
	Low	Med.	High	
Soil/solid				
Aqueous	✓			2 Equip BLKs U+F / date Trip BLK U
Other				

QC CHECK	OK	FYI	ACTION	COMMENTS
Holding Time	✓			3/22
Calibration Blanks	✓			
Initial Calibration	✓			
Continuing Calibration	✓			
Preparation Blank	✓			
CRDL Standard (CRI)	✓			
Interference Check Sample	✓			Did not run for any of the
Lab Control Sample	✓			
Lab Duplicate	✓			
Matrix Spike	✓			
Serial Dilution	✓			Zn 005 11.9

REVIEWER'S COMMENTS:

Linear range 10/13/90 over 3 months
ICSA + ICSAB only run for Mn & Zn thereafter form 11 AR309566
completed.
Zn - BL 005 11.9 serial dil J, 009 UJ (associated BLK)

BCM

Page 2 of 3

BCM ENGINEERS, INC.
INORGANIC DATA VALIDATION SUMMARY
MERCURY ANALYSES

Project Name Chem-Solv Inc Case No. 868-006
BCM Project No. 6012-02 Sampling Date(s) 2/19, 20
Field ID 26AU, 26AF, 28AU Date Review Completed 3/26/91
33AU, 33AF, 33BK, 7.12.12 Reviewer D McGuire

MATRIX	CONCENTRATION			MATRIX RELATED COMMENTS
	Low	Med.	High	
Soil/solid				
Aqueous	/			
Other				

QC CHECK	OK	FYI	ACTION	COMMENTS
Holding Time	/			
Calibration Blank	/			
Initial Calibration	/			
Continuing Calibration	/			
CRDL Standard (CRA)	/			
Preparation Blank	/			
Lab Duplicate	/			
Matrix Spike	/			

r = 0.9996

REVIEWER'S COMMENTS:

13, 14, 17, 19, 21, 22 Total
15, 16, 19, 20, 23, 24 Inorg.

019 - Inorg mercury > Total

AR309367

BCM

APPENDIX B

AR309368

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BCM ENGINEERS INC.

VOLATILE/SEMI-VOLATILE DATA VALIDATION SUMMARY

Project Name Chem Solu CASE 862-006
SDG BL001 Sampling Dates 2/19/91
BCM Project No. 00-6012-02 Date Review Completed 4/29/91
Field ID 26A, 41A, 33A, 33AD, Reviewer Arthur F. Davis
TRIP BLANK, EQUIP BLANK

MATRIX	CONCENTRATION			MATRIX-RELATED COMMENTS
	Low	Med	High	
Soil/Solid	—	—	—	—
Aqueous	<u>X</u>	—	—	—
Other	—	—	—	—

<u>VOLATILE/SEMI-VOLATILE</u>	<u>OK</u>	<u>FYI</u>	<u>ACTION</u>	<u>COMMENTS</u>
Holding Time	<u>X</u>	—	—	—
GC/MS Tuning	<u>X</u>	—	—	—
Calibration	—	<u>X</u>	—	<u>ICAL RAF & BASD OK</u> <u>CCAL 3 compounds %D >25% SPEC OK</u>
Blanks	<u>X</u>	—	—	—
Surrogate Recovery	<u>X</u>	—	—	—
MS/MSD	<u>X</u>	—	—	—
Field Duplicates	<u>X</u>	—	—	<u>Agree well, TCE MD = 8.7%</u> <u>1,1,1-TCA = 22.2% (low), PCE MS 4.</u> <u>OK due to low level results.</u>
Internal Standards	<u>X</u>	—	—	—
TCL Compound ID	<u>X</u>	—	—	<u>Benzene was identified in BC002</u> <u>run after MS/MSD @ 60 ug/L each identified</u> <u>See attached review TICS</u>
TIC	—	<u>X</u>	—	—
Quantitation and Detection Limits	<u>X</u>	—	—	—

* Possible compound of benzene in BC002; however, BC002 also has TICs of similar structures (pentachlorobenzene). 50% compound not likely - reviewer assessed positive for benzene in sample.

SOP WORK SHEETS FOR VOLATILES

HOLDING TIMES														
LAB	SAMPLE NO. FIELD	HCP PRESENTED ATTACHED		CONC LEVEL/MATRIX	DATE SAMPLED	DATE RECEIVED	DATE ANALYZED	TIME ANALYZED	INSTRUMENT ID.	10 days UTA CONTRACT HOLDING TIME NET **		14 days UTA 40 CFR 136 HOLDING TIME NET		ACTION
		YES	NO							YES	NO			
BC001	41A	✓		L/A	2/17/91	2/20/91	3/2	1:05	MSDS	✓	✓			
BC002	26A	✓					3/2	4:07		✓	✓			
BC004	33AD	✓					3/1	23:10		✓	✓			
BC006	33A	✓					3/1	22:36		✓	✓			
BC008	EBLK	✓					3/1	23:50		✓	✓			
BC010	TEBLK	✓					3/2	0:26		✓	✓			
BC001MS	41A	✓					3/2	3:32		✓	✓			
BC001MSD	41A	✓		✓	✓	✓	3/2	2:29	✓	✓	✓	✓		
VBK				L/A	NA	NA	3/5/91	22:01	MSDS					

** SAMPLES REC'D 2/20 @ 9:10 A, 10 days UTA = 3/2/91

AR309370

* INCLUDE MATRIX SPIKES, BLANKS AND RE-RUNS HERE

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Chem-Solv, Inc. SFC
 Chewood, Delaware
 Project No. 00-6012-02
 Property of: BCM Engineers Inc.
 1 Plymouth Meeting, PA 19462
 (210) 925-3800

Volume I - Field activities
 October 1987 - March 1990

Volume II - Field activities

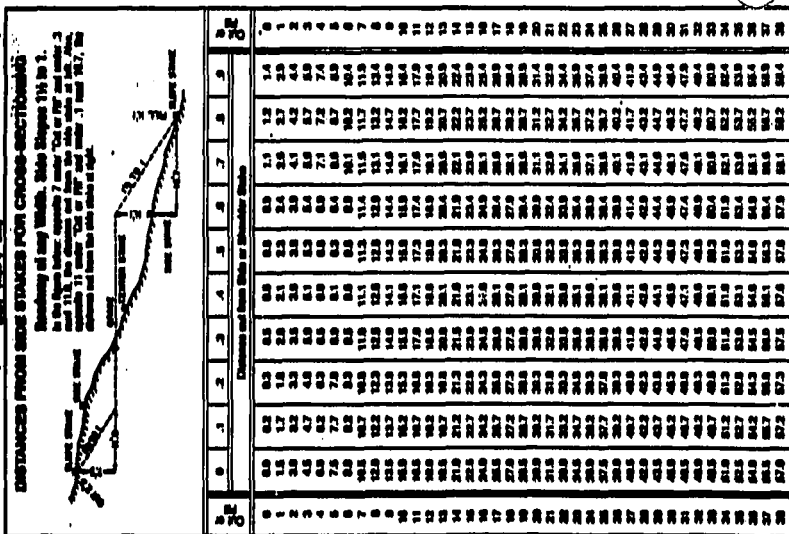
Volume III - Field activities
 March 1990 -

Volume IV - Groundwater Sampling

ATTACHMENT A



The paper in this book is
 made of 50% high grade rag stock with
 a WATER RESISTING surface lining.



AR309371

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7A
VOLATILE CONTINUING CALIBRATION CHECK

Lab Name: IEA Contract: 68-W8-0045
 Lab Code: IEA Case No.: 868-6 SAS No.: SDG No.: BC001
 Instrument ID: msd5 Calibration Date: 3/ 1/91 Time: 21:24
 Lab File ID: 0301E01 Init. Calib. Date(s): 2/20/91 2/20/91
 Matrix:(soil/water) WATER Level:(Low/med): LOW Column:(pack/cap) CAP
 Min RRF50 for SPCC(%) = .300 (0.250 for Bromoform) Max %D for CCC(*) = 25.0%

COMPOUND	RRF	RRF50	%D
CHLOROMETHANE	.842	.672	20.2
BROMOMETHANE	.991	.540	45.5
VINYL CHLORIDE	1.032	.889	13.9
CHLOROETHANE	.664	.692	4.2
METHYLENE CHLORIDE	1.202	1.329	10.5
ACETONE	.138	.177	28.1
CARBON DISULFIDE	2.677	2.668	.4
1,1-DICHLOROETHENE	1.002	1.031	2.9
1,1-DICHLOROETHANE	2.488	2.730	9.7
1,2-DICHLOROETHENE (TOTAL)	1.288	1.437	11.6
CHLOROFORM	2.598	2.982	14.8
1,2-DICHLOROETHANE	1.401	1.716	22.5
2-BUTANONE	.065	.092	42.0
1,1,1-TRICHLOROETHANE	.426	.477	12.0
CARBON TETRACHLORIDE	.424	.476	12.3
VINYL ACETATE	.372	.433	16.4
BROMODICHLOROMETHANE	.617	.715	15.9
1,2-DICHLOROPROPANE	.385	.421	9.3
CIS-1,3-DICHLOROPROPENE	.553	.608	10.0
TRICHLOROETHENE	.395	.435	10.0
DIBROMOCHLOROMETHANE	.517	.612	18.5
1,1,2-TRICHLOROETHANE	.298	.343	14.9
BENZENE	.743	.822	10.6
TRANS-1,3-DICHLOROPROPENE	.433	.466	7.7
BROMOFORM	.387	.465	20.1
4-METHYL-2-PENTANONE	.204	.250	22.5
2-HEXANONE	.138	.168	21.9
TETRACHLOROETHENE	.429	.473	10.1
1,1,2,2-TETRACHLOROETHANE	.572	.642	12.3
TOLUENE	.607	.618	1.8
CHLOROBENZENE	.885	.901	1.7
ETHYLBENZENE	.381	.384	.8
STYRENE	.742	.789	6.4
XYLENE (TOTAL)	.448	.466	3.9
TOLUENE-d8	1.080	1.057	2.1
BROMOFLUOROBENZENE	.944	1.002	6.2
1,2-DICHLOROETHANE-D4	1.335	1.514	13.4

AR309373

FORM VII VOA

1 0072

1/8

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Initial Cal $\overline{RRF}$ check

ChemSolo

6012-02

4/29/91 *ADD*

Trichloroethane

$$RRF\ 20 = \frac{165944}{905772} \times \frac{50}{20} = .458 \checkmark OK$$

$$RRF\ 50 = \frac{370729}{906608} \times \frac{50}{50} = .4089 (.409) \checkmark OK$$

$$RRF\ 100 = \frac{742320}{947948} \times \frac{50}{100} = .3915 (.392) \checkmark OK$$

$$RRF\ 150 = \frac{1026782}{932511} \times \frac{50}{150} = .367 \checkmark OK$$

$$RRF\ 200 = \frac{1272464}{906061} \times \frac{50}{200} = .3511 (.351) \checkmark OK$$

Check

$$\%RSD = \frac{\frac{.0415}{.3954} \times 100}{.3954} = 10.4957 (10.5) \checkmark OK$$

$$\overline{x} = .395 \checkmark OK$$

1,1,1-TCA

$$RRF\ 20 = \frac{162167}{405172} \times \frac{50}{20} = .4474 (.447) \checkmark OK$$

$$RRF\ 50 = \frac{339745}{906608} \times \frac{50}{50} = .4299 (.430) \checkmark OK$$

$$RRF\ 100 = \frac{814948}{947948} \times \frac{50}{100} = .4298 (.430) \checkmark OK$$

$$RRF\ 150 = \frac{1169114}{932511} \times \frac{50}{150} = .4181 (.418) \checkmark OK$$

$$RRF\ 200 = \frac{1477457}{906061} \times \frac{50}{200} = .4066 (.407) \checkmark OK$$

$$\%RSD = \frac{\frac{.0150}{.4264} \times 100}{.4264} = 3.5177 \checkmark OK \text{ rounding diff.}$$

$$\overline{x} = .4264 = .426 \checkmark OK$$

AR309374

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ICAL Check

Benzene

$$RRF_{20} = \frac{813399}{905772} \times \frac{50}{20} = .8152 (.915) \checkmark OK$$

$$RRF_{50} = \frac{679931}{906608} \times \frac{50}{50} = .750 \checkmark OK$$

$$RRF_{100} = \frac{1410180}{947948} \times \frac{50}{100} = .7438 (.744) \checkmark OK$$

$$RRF_{150} = \frac{142269}{932611} \times \frac{50}{150} = .705 \checkmark OK$$

$$RRF_{200} = \frac{2532970}{906061} \times \frac{50}{200} = .6989 (.699) \checkmark OK$$

$$\%RSD = \frac{.0464}{.7436} \times 100 = 6.11 \quad (6.3 \checkmark OK \text{ round diff})$$

$$\bar{x} = .7426 = .743 \checkmark OK$$

Tetrachloroethane

$$RRF_{20} = \frac{159119}{758347} \times \frac{50}{20} = .5244 (.524) \checkmark OK$$

$$RRF_{50} = \frac{339338}{769034} \times \frac{50}{50} = .4409 (.441) \checkmark OK$$

$$RRF_{100} = \frac{684515}{783711} \times \frac{50}{100} = .4176 (.418) \checkmark OK$$

$$RRF_{150} = \frac{897585}{746804} \times \frac{50}{150} = .4007 (.401) \checkmark OK$$

$$RRF_{200} = \frac{1055461}{726087} \times \frac{50}{200} = .3634 (.363) \checkmark OK$$

$$\%RSD = \frac{.0601}{.4294} \times 100 = 13.996 (14) \checkmark OK$$

$$\bar{x} = .4294 \checkmark OK$$

Chem

6012-02

4/29/91

AR309375

CCAL Check

Chem Solo

6012-02 ASD

4/25/91

TCE

$$RRF_{SD} = \frac{376067}{864705} \times \frac{SD}{SD} = .4349 \checkmark OK$$

$$\%D = \frac{.395 - .435}{.395} \times 100 = -10.1 \checkmark OK \text{ Round diff.}$$

1,1,1-FTCA

$$RRF_{SD} = \frac{412762}{864705} \times \frac{SD}{SD} = .4773 \checkmark OK$$

$$\%D = \frac{.426 - .477}{.426} \times 100 = -11.97 (12.0) \checkmark OK$$

Benzene

$$RRF_{SD} = \frac{710410}{864705} \times \frac{SD}{SD} = .8216 \checkmark OK$$

$$\%D = \frac{.743 - .822}{.743} \times 100 = -10.63 (10.6) \checkmark OK$$

Tetrachloroethane

$$RRF_{SD} = \frac{.429 - .473}{.429} \times 100 = -10.256 (10.1) \text{ Round off } \checkmark OK$$

$$RRF_{SD} = \frac{362867}{767497} \times \frac{SD}{SD} = .4729 \checkmark OK$$

$\%D > 25\%$

Bromomethane

$$RRF_{SD} = \frac{104211}{192911} \times \frac{SD}{SD} = .5405 (.540) \checkmark OK$$

$$\%D = \frac{.991 - .540}{.991} \times 100 = 45.51 (45.5) \checkmark OK$$

$> 25\%$

Acetone

$$RRF_{SD} = \frac{34054}{192911} \times \frac{SD}{SD} = .1766 (.177) \checkmark OK$$

$$\%D = \frac{.138 - .177}{.138} \times 100 = -28.3 \checkmark OK$$

$> 25\%$ AR309376 effect

2-Butanone

$$RRF_{SD} = \frac{17815}{192911} \times \frac{SD}{SD} = .0924 (.092) \checkmark OK$$

$$\%D = \frac{.065 - .092}{.065} \times 100 = -41.54 (42) \checkmark OK > 25\%$$

SOP WORK SHEETS FOR VOLATILES

[illegible]

AR309377

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SOP WORK SHEETS FOR VOLATILES

[illegible]

AR309378

* INCLUDE MATRIX SPIKES, BLANKS AND RE-RUNS HERE

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SOP WORK SHEETS FOR VOLATILES

[illegible]

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1E
VOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDS

EPA SAMPLE NO.

BC002

Lab Name: IEA

Contract: 68-W8-0045

Lab Code: IEA

Case No.: 868-6

SAS No.:

SDG No.: BC001

Matrix: (soil/water) WATER

Lab Sample ID:

Sample wt/vol: 5.000 (g/mL) ML

Lab File ID: 0301E11

Level: (low/med) LOW

Date Received: 2/20/91

% Moisture: not dec. 100.

Date Analyzed: 3/ 2/91

Column: (pack/cap) CAP

Dilution Factor: 1.00

CONCENTRATION UNITS:
(ug/L or ug/Kg) UG/L

Number TICs found: 10

CAS NUMBER	COMPOUND NAME	RT	EST. CONC.	Q	Rev COMI
1. 78-78-4	Butane, 2-methyl-	5.49	100.	J	OK
2. -	UNKNOWN	5.66	100.	J	OK
3. 96-14-0	Pentane, 3-methyl-	8.67	100.	J	OK
4. 96-37-7	Cyclopentane, methyl-	10.43	100.	J	OK
5. -	UNKNOWN HYDROCARBON	11.72	100.	J	OK
6. -	UNKNOWN	12.44	100.	J	OK
7. 108-87-2	Cyclohexane, methyl-	13.78	40.	J	OK
8. -	UNKNOWN HYDROCARBON	14.53	30.	J	(4)
9. 560-21-4	Pentane, 2,3,3-trimethyl-	14.78	40.	J	(5)
10. 767-58-8	1H-Indene, 2,3-dihydro-1-met	25.58	50.	J	(5)
11.					
12.					
13.					
14.					
15.					
16.					
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28.					
29.					
30.					

- (1) No match in library
- (2) Appears to be hydrocarbons
- (3) Match qualities poor
- (4) Possible trimethylpentane isomer
- (5) Could also be an ethenyl, ethyl benzene isomer.

FORM I VOA-TIC

1/87 Rev.

AR309380

1 0013

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AR309381

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APPENDIX N

AR309382

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D

APPENDIX N

GROUNDWATER ANALYTICAL RESULTS AND QUALITY ASSURANCE REVIEW
EPA SPLIT SAMPLES
FEBRUARY 1991

AR309383

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

REGION III
CENTRAL REGIONAL LABORATORY
839 WESTGATE ROAD
ANNAPOLIS, MARYLAND 21401
(301) 286-8180

DATE: March 18, 1991

SUBJECT: NH_3N Determination of Chem Solv Samples
Superfund-Enforcement, (2/26/91 - 3/7/91), 910221-01,03,05

FROM: Wanda Boyd *WB*
Environmental Specialist

TO: Joseph L. Slayton
Acting Chief, Laboratory Branch

THRU: Khin C. Thuang *KCT*
Chief, Inorganic Section

Chem Solv samples, 910221-01,03,05 were analyzed for the determination of NH_3N (EPA Method 350.1). All samples were preserved and then their pH was adjusted to 6-7 using a 6N solution of sodium hydroxide. The results appear below.

Additional quality control data is available for review upon request.

Sample Description:

<u>Lab No.</u>	<u>Description</u>
910221-01	Chem Solv, Monitoring Well 9B, Sta. MW9B
910221-03	Chem Solv, Monitoring Well 9B-D, Sta. MW-9B-D
910221-05	Chem Solv, Field Blank, Sta. FB

Results:

<u>Sample No.</u>	<u>NH_3N (mg/L)</u>
910221-01	<0.040* (114%)
910221-03	<0.040
910221-05	<0.040

* Analyzed in duplicate, both values below specified detection limit.
Number in parenthesis is a spike recovery.

WB:nc

cc: Norman Fritsche *NF*

AR309384

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

REGION III
CENTRAL REGIONAL LABORATORY
838 BERTGATE ROAD
ANNAPOLIS, MARYLAND 21401
(301) 266-9180

DATE : March 25, 1991

SUBJECT: Mercury Determination of Chem Solv, Superfund Enforcement (TG803NPN5),
(2/28/91 - 3/14/91), 910221-01-06

FROM : Thomas H. Reppert ^R
Environmental Scientist

TO : Frederick Dreisch
Chief, Laboratory Branch

THRU : Khin C. Thaug ^{KCT}
Chief, Inorganic Section

The results of the mercury analyses are by automated cold vapor atomic absorption spectroscopy, EPA Method 245.2.

Additional quality control results are available upon request.

The mercury analyses provided by the CRL are virtually all for Total Hg. Our method, therefore, to determine separate organic and inorganic values was somewhat modified to accommodate this data request. Organic Hg values were to represent the difference between Total Hg values, which are obtained through analysis of digested sample, and inorganic values which are obtained by analysis of undigested sample.

The values for Total Hg and Inorganic Hg are listed below. The generally lower values for Total Hg can be attributed to the natural variability inherent in the two methods used, the digestion process, and the auto analyzer-detection equipment. While the actual Organic Hg percentage cannot be consistently quantified by our analysis methods, it's apparent that nearly all of the Total Hg is inorganic. The detection of the small amount of Organic Hg remaining is lost in the method precision variability. Analysis scans were repeated several times with the same results.

If you would like to discuss the analysis data further, please contact Tom Reppert at (301) 266-9180.

AR309385

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Sample Description and Results:

Lab No.	Description
910221-01	Chem Solv, Monitoring Well 9B, Sta. MW9B
-02	Chem Solv, Monitoring Well 9B, Sta. MW9B
-03	Chem Solv, Monitoring Well 9B-D, Sta. MW-9B-D
-04	Chem Solv, Monitoring Well 9B-D, Sta. MW-9B-D
-05	Chem Solv, Field Blank, Sta. FB
-06	Chem Solv, Field Blank, Sta. FB

Results:

Lab No.	Total Hg(ug/L)	Filtered Total Hg(ug/L)	Inorganic Hg(ug/L)	Filtered Inorganic Hg(ug/L)
910221-01	1.7	---	2.6[75%] RPD=4.9	---
-02	---	1.9	---	2.3[106%] RPD=3.5
-03	1.6[93%] RPD=2.9	---	2.5	---
-04	---	1.8[95%] RPD=1.9	---	2.2
-05	<0.2	---	<0.2	---
-06	---	<0.2	---	<0.2

*Analyzed in duplicate, both values below specified detection limit.

**Analyzed in duplicate, one value below and one above detection limit.

Numbers in brackets are pre-digestion [matrix] spike recoveries.

RPD = Method duplicates are reported as the Mean and the Relative Percent Difference. $(| \text{Replicate 1} - \text{Replicate 2} | / \text{Mean}) * 100$

THR:nc

cc: Ronald H. Altman *TH*

AR309386

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

Chem Solv

¹ Methods for Chemical Analysis of Water and Waste, EPA-600/4-79-020

RPD - values reported are the average of method duplicates and relative percent difference.

*Samples are analyzed in duplicate; both values below the detection limit.

RHA/sjc

cc: Norman Fritsche *mf*

AR309388

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY
REGION III
CENTRAL REGIONAL LABORATORY
839 BESTGATE ROAD
ANNAPOLIS, MARYLAND 21401

DATE : March 5, 1991

SUBJECT : BOD, Determination of Chem Solv, 910221-01,03,05, Superfund Enforcement TGB03NI
(2/21/91-2/26/91)

FROM : Norman Fritsche *NF*
Environmental Scientist

TO : Joseph L. Slayton
Acting Chief, Laboratory Branch

THRU : Khin C. Thaug *KCT*
Chief, Inorganic Section

BOD, Determinations (910221-01,03,05) EPA Method 405.1* of Chem Solv are listed below.

Additional quality control results are available upon request.

Sample Description and Results:

<u>Lab No.</u>	<u>Description</u>
910221-01	Chem Solv, Monitoring Well 9B, Sta. MW9B
-03	Chem Solv, Monitoring Well 9B-D, Sta. MW-9B-D
-05	Chem Solv, Field Blank, Sta. FB

<u>Lab No.</u>	<u>BOD, mg/L</u>
910221-01	<1.0**
-03	<1.0
-05	1.5

<1.0 - Detection limit of BOD, test using 300 mL of sample.

*Methods for Chemical Analysis of Water and Wastes, EPA 600/4-79-020.

**Sample analyzed in duplicate, both values below detection limit.

NF:sjc

cc: Ronald Altman *RA*

AR309389

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2588A RIVA ROAD
SUITE 300
ANNAPOLIS, MD 21401
PHONE: 301-266-9887

DATE: March 13, 1991

SUBJECT: Chemical Oxygen Demand (COD) of Chem Solv samples for
Superfund-Enforcement. (TGB03NPN5). (02/25/91 thru
03/11/91). 910221-01,03,05.
ESAT TID #03910207. ESAT Task #2298.

FROM: Behrooz Khoshkhoo
ESAT Chemist

Tim Dawson *T. Dawson*
ESAT Chemist

TO: Joseph Slayton
Acting Chief, Laboratory Branch

THRU: Richard Dresser *RD for RD*
ESAT Team Manager

The ESAT Laboratory Staff was assigned three (3) aqueous samples from Chem Solv for Chemical Oxygen Demand. All samples were successfully analyzed within the twenty eight (28) day holding time. Sample descriptions and results are presented in the following tables.

SAMPLE DESCRIPTIONS:

910221-01	Chem Solv, Monitoring Well 9B, Sta. MW9B
910221-03	Chem Solv, Monitoring Well 9B-D, Sta. MW-9B-D
910221-05	Chem Solv, Field Blank, Sta. FB

AR309390

WESTEN

Site Name: Chem Solv
Sample Nos.: 910221-01,03,05.

Page 2 of 3

SAMPLE RESULTS:

<u>Lab No.</u>	<u>Chemical Oxygen Demand (mg/L)</u>
910221-01	15.2*
910221-03	15.4
910221-05	36.8

O.C. RESULTS:

Duplicates:

<u>Lab.No.</u>	<u>Sample Result (mg/L)</u>	<u>Duplicate Result (mg/L)</u>	<u>SD</u>
910221-01	15.4	15.0	0.3

Spikes:

<u>Lab.No.</u>	<u>Sample Result (mg/L)</u>	<u>Spike Result (mg/L)</u>	<u>Spike Added (mg/L)</u>	<u>% Recovery</u>
910221-01	15.2	50.4	37.5	94

Audit:

<u>Lab No.</u>	<u>Found Value (mg/L)</u>	<u>True Value (mg/L)</u>	<u>% Recovery</u>
WP1284 (conc.1) (Range 7.27-22.3)	15.4	14.9	103

NOTE:

* The result for sample 910221-01 is the average of duplicate result.

AR309391

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WESTERN

Site Name: Chem Solv
Sample Nos.: 910221-01,03,05.

Page 3 of 3

NOTE: The Q.C. samples provide an estimate of combined sample preparation and measurement variability and/or bias.

cc: Ron Altman, EPA Task Monitor - *RL*
Terry Simpson, ESAT DPO

BK103A03.CHE

AR309392

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2566A RIVA ROAD
SUITE 300
ANNAPOLIS, MD 21401
PHONE: 301-266-9887

DATE: March 22, 1990

SUBJECT: Volatile Gas Chromatography/Mass Spectrometry Analyses
of Samples from the Chem Solv Site for Superfund-
Enforcement (TGB03NPN5). (02/25/91 to 03/08/91);
Samples: 910221-07 thru 910221-11; ESAT TID No.:
03910214; ESAT Task No.: 2299.

FROM: Sue Raupuk *S.R.*
Analytical Chemist

Fredrick Foreman *FF*
Senior Analytical Chemist

TO: Frederick Dreisch
Chief, Laboratory Branch

THRU: Richard D. Dresser *RDE*
ESAT Team Manager

OVERVIEW

Five (5) aqueous samples were assigned to ESAT on February 25, 1991 for analyses of both hazardous substance list (HSL) and priority pollutant list (PPL) volatile organic compounds. They were analyzed on February 25 and 26, 1991 following methods from the "USEPA Contract Laboratory Program Statement of Work for Organics Analysis", February, 1988 and EPA Method 624, "Purgeables", Federal Register, October, 1984. Instrumentation utilized consisted of a purge and trap apparatus interfaced to a gas chromatograph/mass spectrometer equipped with a fused silica capillary column. The samples that were received for analyses are described in the following table.

SAMPLE IDENTIFICATION:

<u>Sample No.</u>	<u>Description</u>
910221-07	Chem Solv, Monitoring Well 26A, Sta. MW-26A
910221-08	Chem Solv, Monitoring Well 33A, Sta. MW-33A
910221-09	Chem Solv, Monitoring Well 33D, Sta. MW-33D
910221-10	Chem Solv, Trip Blank, Sta. TB
910221-11	Chem Solv, CRL Lab Blank

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WESTEN

Site Name: Chem Solv
Sample Nos.: 910221-07 thru 910221-11

page 2 of 3

NOTES

- o The target compounds' concentrations were calculated using relative response factors based on the closest eluting internal standard. The internal standard and surrogate mixes were added immediately prior to analyses.
- o All HSL and PPL compounds which were positively identified are reported on the volatile analysis data sheet. The contaminants found below their nominal quantitation limit (NQL) are qualified "J" for estimated.
- o All other eluting peaks with an area above ten percent of the nearest internal standard were compared to the NBS mass spectral library for tentative identification.
- o All compounds which were found in both a method blank and/or a field blank and a sample were qualified "B" if the concentration of the compound in the sample was less than ten times (10x) the compound's concentration in the blank.
- o All samples were analyzed only after correct system performance was confirmed to meet EPA approved tuning criteria for Bromofluorobenzene (BFB) (See Form Vs in Appendix C).
- o All analyses exhibited surrogate recoveries within Region III CRL Q.C. limits. (See the Volatile Surrogate Recovery Data Sheet in Appendix C). A matrix spike and matrix spike duplicate was analyzed with this sample set and all recoveries and %RSDs were within Region III CRL Q.C. limits (See Matrix Spike %Recovery Data Sheets).
- o Several compounds exhibited a %D greater than twenty-five (25) percent when comparing the daily calibration standard and the initial calibration curve. Also, the %RSDs for several compounds in the initial calibration standards were above thirty (30) percent. These compounds are either common laboratory contaminants or were not found to be present in these samples. They are qualified "UJ" in the affected samples (See support documentation in Appendix C).

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Site Name: Chem Solv
Sample Nos.: 910221-07 thru 910221-11

page 3 of 3

Attachments

1. Appendix A - Glossary of Data Qualifier Codes
2. Appendix B - Data Summary; Results as Reported by the Laboratory and Qualified
3. Appendix C - Support Documentation

cc: Sue Warner, EPA Task Monitor
Terry Simpson, ESAT DPO

SW

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APPENDIX A

Glossary of Data Qualifier Codes

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GLOSSARY OF DATA QUALIFIER CODES (ORGANIC)

CODES RELATING TO IDENTIFICATION

(Confidence concerning presence or absence of compounds)

NO CODE = Confirmed identification

U = Not Detected. The associated number indicates approximate sample concentration necessary to be detected.

B = Not detected substantially above the level reported in the laboratory or field blanks.

R = Unreliable results. Analyte may or may not be present in the sample. Supporting data necessary to confirm results.

N = Tentative identification. Consider present. Special methods may be needed to confirm its presence or absence in future sampling efforts.

CODES RELATED TO QUANTITATION

(Can be used for both positive results and sample quantitation limits)

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

K = Analyte present. Reported value may be biased high. Actual value is expected to be lower.

L = Analyte present. Reported value may be biased low. Actual value is expected to be higher.

UJ = Not detected. Quantitation limit may be inaccurate or imprecise.

UL = Not detected. Quantitation limit is probably higher.

OTHER CODES

Q = No analytical results.

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APPENDIX B

Data and TIC Summary

AR309398

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E.P.A. Region III Central Regional Laboratory

Volatile Organics Analysis Data Sheet

Site: Chem Solv
Program: Superfund-Enforcement
Units: ug/L

			Date Sampled:	02/19/91	02/19/91	02/19/91	02/19/91	02/25/91	
			Date Analyzed:	02/25/91	02/25/91	02/25/91	02/25/91	02/25/91	
			Dilution Factor =	(1)	(1)	(1)	(1)	(1)	(1)
				Sta. MW-26A 910221-07	Sta. MW-33A 910221-08	Sta. MW-33D 910221-09	Trp. Blk., Sta. TB 910221-10	CRL Lab Blank 910221-11	
NGL	CAS #	Compound							
5	75-71-8	Dichlorodifluoromethane							
5	76-87-3	Chloromethane	UJ	UJ	UJ	6.0J	UJ		
5	75-01-6	Vinyl Chloride							
5	74-83-9	Bromomethane							
5	75-00-3	Chloroethane							
5	75-69-4	Trichlorofluoromethane	UJ	UJ	UJ	UJ	UJ		
5	75-35-4	1,1-Dichloroethene	UJ	UJ	UJ	UJ	UJ		
5	75-15-0	Carbon Disulfide							
10	67-66-1	Acetone				8.3J			
5	75-09-2	Methylene Chloride	UJ	UJ	UJ	UJ	5.6J		
5	156-60-5	trans-1,2-Dichloroethene							
5	75-34-3	1,1-Dichloroethane							
10	108-05-4	Vinyl Acetate							
5	594-20-7	2,2-Dichloropropane							
10	78-93-3	2-Butanone	3.48	UJ	UJ	2.4J	UJ		
5	74-97-5	Bromochloromethane							
5	156-59-2	cis-1,2-Dichloroethene		1.4J	1.5J				
5	65-66-3	Chloroform		1.48	1.38	5.2			
5	71-35-6	1,1,1-Trichloroethane		11.6	11.6				
5	56-23-5	Carbon Tetrachloride							
5	563-58-6	1,1-Dichloro-1-propene							
5	71-43-2	Benzene	32.0						
5	107-06-2	1,2-Dichloroethane							
5	79-01-6	Trichloroethane		142	142				
5	78-87-5	1,2-Dichloropropane							
5	74-95-3	Dibromomethane							
5	75-27-4	Bromodichloromethane				0.4J			
10	100-73-8	2-Chloroethylvinyl ether							
5	10061-02-6	trans-1,3-Dichloropropene							
10	108-10-1	4-Methyl-2-pentanone							
5	106-68-3	Toluene							
5	10061-01-5	cis-1,3-Dichloropropene							
5	79-00-5	1,1,2-Trichloroethane	1.3J	3.5J	3.5J				
5	127-18-4	Tetrachloroethane							
5	142-28-9	1,3-Dichloropropane							
10	591-78-6	2-Hexanone							
5	124-48-1	Dibromochloromethane							
5	106-93-4	1,2-Dibromoethane (EDB)							
5	108-90-7	Chlorobenzene							
5	630-20-6	1,1,1,2-Tetrachloroethane							
5	100-41-4	Ethylbenzene							

NGL = Nominal Quantitation Limit

Actual Quantitation Limit = Dilution Factor x NGL

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E.P.A. Region III Central Regional Laboratory

Volatile Organics Analysis Data Sheet

Site: Chem Solv
Program: Superfund-Enforcement
Units: ug/L

			Date Sampled:	02/19/91	02/19/91	02/19/91	02/19/91	02/25/91
			Date Analyzed:	02/25/91	02/25/91	02/25/91	02/25/91	02/25/91
			Dilution Factor =	(1)	(1)	(1)	(1)	(1)
				Sta. MW-26A	Sta. MW-33A	Sta. MW-33D	Trp. Btk. Sta. TS	CAL Lab Blank
NGL	CAS #	Compound		910221-07	910221-08	910221-09	910221-10	910221-11
-5		(m & p)-Xylene isomers	0.4J					
5	95-47-6	o-Xylene						
5	100-42-5	Styrene						
5	75-23-2	Bromoform						
-5	98-82-8	Isopropylbenzene	15.8					
5	108-86-1	Bromobenzene						
5	79-34-5	1,1,2,2-Tetrachloroethane						
5	96-18-6	1,2,3-Trichloropropane						
5	103-65-1	n-Propylbenzene						
5	95-49-8	2-Chlorotoluene						
5	106-43-4	4-Chlorotoluene						
5	108-67-8	1,3,5-Trimethylbenzene						
-5	98-06-6	tert-Butylbenzene	0.4J					
5	93-63-6	1,2,4-Trimethylbenzene	0.4J					
5	133-98-8	sec-Butylbenzene	8.8					
5	541-73-1	1,3-Dichlorobenzene						
5	106-46-7	1,4-Dichlorobenzene						
5	99-87-6	p-Isopropyltoluene						
5	95-50-1	1,2-Dichlorobenzene						
-5	104-51-8	n-Butylbenzene	6.2					
5	96-12-8	1,2-Dibromo-3-chloropropane						
5	120-82-1	1,2,4-Trichlorobenzene						
5	91-20-3	Naphthalene						
5	87-68-3	Hexachlorobutadiene						
5	87-61-6	1,2,3-Trichlorobenzene						

IL = Nominal Quantitation Limit

Actual Quantitation Limit = Dilution Factor x NGL

AR309400

B-2

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WESTERNVOLATILE ORGANICS ANALYSIS DATA SHEET
TENTATIVELY IDENTIFIED COMPOUNDSCase Name: Chem Solv Contractor: Region III CRL-ESATSample ID: 910221-07_Stn. MW-26A Lab File ID: EA799Date Rec: 02/23/91 Date/Time Analyzed: 02/23/91 13:49Matrix: water Level: lowNumber TIC's found: 18 Concentration units: ug/L

CAS NUMBER	COMPOUND IDENTIFICATION	R.T.(min)	EST. CONC.
*****	Unknown, m/z = 43	9.09	12.0
78784	Butane, 2 methyl-(8C19C1)	10.41	150
75832	Butane, 2,2-dimethyl-(8C19I)	11.67	20.0
*****	Unknown, m/z = 43	12.40	45.0
*****	Unknown, m/z = 42	12.48	61.0
*****	Unknown, m/z = 57	12.86	69.0
96377	Cyclopentane, methyl-(8C19C1)	14.23	64.0
*****	Unknown, m/z = 84	15.30	26.0
*****	Unknown, m/z = 56	15.41	26.0
*****	Unknown, m/z = 57	16.06	22.0
*****	Unknown, m/z = 55	17.31	32.0
563733	Pentane, 2,3,4-trimethyl-(8C19C1)	18.14	24.0
560214	Pentane, 2,3,3-trimethyl-(8C19C1)	18.38	31.0
533773	Benzene, 1-methyl-3-(1-methylethyl)- (9C1)	30.03	20.0
767388	1H-Indene, 2,3-dihydro-1-methyl-(9C1)	30.23	33.0
767388	1H-Indene, 2,3-dihydro-1-methyl-(9C1)	32.24	17.0
*****	Unknown aromatic, m/z = 119	32.59	6.0
17059482	1H-Indene, 2,3-dihydro-1,4-dimethyl- (9C1)	33.13	10.0

FORM 1 VOA-TIC

AR309401

B-3

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APPENDIX C

Support Documentation

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E.P.A. Region III Central Regional Laboratory

Volatile Surrogate Recovery Data Sheet

Site : Chem Solv
Program : Superfund-Enforcement
Date Sampled: 02/19/91, 02/25/91 & 02/26/91
Date Analyzed: 02/25/91 & 02/26/91
Units : % Recovery

SAMPLE I.D.	1,2-Dichloro- ethane-d4	Fluoro- benzene	Toluene-d8	Bromofluoro- benzene
1. 910221-07	112	100	106	106
2. 910221-08	113	100	102	104
3. 910221-09	114	101	103	101
4. 910221-10	112	101	103	108
5. 910221-11 CRL Lab Blank	106	99	99	102
6. 910221-08MS	105	97	102	105
7. 910221-08MSD	99	97	108	94
8. VOA Method Blank (02/26/90)	98	99	98	97
CRL LIMITS:	76-114	80-120	88-110	86-115

% REC: 0 out of 32 outside limits

AR309404

c-1

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E.P.A. Region III Central Regional Laboratory

Matrix Spike % Recovery Data Sheet

Site: Chem Solv
Program: Superfund-Enforcement

Sample I.D.: 910221-08
Date Sampled: 02/19/91
Date Analyzed: 02/26/91

CAS #	COMPOUND	SPIKE		MATRIX SPIKE		MATRIX SPIKE DUP	
		CONC.	REC.	CONC.	REC.	CONC.	REC.
75-35-6	1,1-Dichloroethene	50	51.7	103	51.0	102	
71-43-2	Benzene	50	48.3	97	50.5	101	
79-01-6	Trichloroethene	50	50.3	101	50.3	101	
108-88-3	Toluene	50	50.6	101	57.4	115	
108-90-7	Chlorobenzene	50	51.6	103	54.0	108	

CAS #	COMPOUND	RPO	RPO		RECOVERY	
			QC LIMITS		QC LIMITS	
75-35-6	1,1-Dichloroethene	1	14		61-145	
71-43-2	Benzene	4	11		76-127	
79-01-6	Trichloroethene	0	14		71-120	
108-88-3	Toluene	13	13		76-125	
108-90-7	Chlorobenzene	5	13		75-130	

RPO = Relative percent difference

REC.: 0 out of 10 outside limits

RPO: 0 out of 5 outside limits

AR309405

C-2

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WESTERN

GC/MS TUNING AND MASS CALIBRATION

4-Bromofluorobenzene (BFB)

Case No.: Chem Solv

Contractor: Region III CRL (ESAT)

Instrument ID: HP-5970MSD

Date/Time: 02/20/90 09:52

Lab File ID: >EA761

Date Release Authorized By: _____

m/z	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of the base peak	15.68 OK
75	30.0 - 60.0% of the base peak	38.15 OK
95	Base peak, 100% relative abundance	100.00 OK
96	5.0 - 9.0% of the base peak	6.80 OK
173	Less than 2.0% of mass 174	0.00 OK
174	Greater than 50.0% of the base peak	56.71 OK
175	5.0 - 9.0 % of mass 174	6.54 OK (7.55)#1
176	Between 95.0 and 101.0% of mass 174	56.86 OK (100.18)#1
177	5.0 - 9.0% of mass 176	5.53 OK (6.37)#2

THIS PERFORMANCE TUNE APPLIES TO THE FOLLOWING
SAMPLES, BLANKS AND STANDARDS.

#1 - Value in parentheses is % mass 176

#2 • Value in parentheses is % says 176

[illegible]

FORM V

7/85

AR309406

C-3

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GC/MS TUNING AND MASS CALIBRATION

4-Bromofluorobenzene (BFB)

Case No.: Chem Solv

Contractor: Region III CRL (ESAT)

INSTRUMENT ID: HP-5970MSD

Date/Time: 02/26/91 11:40

Lab File ID: PEAS09

Date Release Authorized By: _____

m/z	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of the base peak	22.26 OK
75	30.0 - 60.0% of the base peak	48.88 OK
95	Base peak, 100% relative abundance	100.00 OK
96	5.0 - 9.0% of the base peak	6.48 OK
173	Less than 2.0% of mass 174	0.00 OK
174	Greater than 50.0% of the base peak	86.40 OK
175	5.0 - 9.0 % of mass 174	6.36 OK (7.59)#1
176	Between 95.0 and 101.0% of mass 174	82.67 OK (95.68)#1
177	5.0 - 9.0% of mass 176	5.19 OK (6.28)#2

THIS PERFORMANCE TIME APPLIES TO THE FOLLOWING
SAMPLES, BLANKS AND STANDARDS.

#1 - Value in parentheses is % mass 174

#2 - Value in parentheses is % mass 176

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FORM V

7/85

AR309408

4-5

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ISr SS Mix prepared :
STO Mix prepared 2/1

Initial Calibration Data
HSL Compounds

Case No: _____ Instrument ID: HP 5970 MSD
Contractor: REGION III - CRL Calibration Date: ~~02-22-91~~ 02/20/91 SR
Contract No: ESAT

Minimum RF for SPCC is 0.300 Maximum % RSD for CCC is 30.0%

Compound	RF	RF	RF	RF	RF	RRT	RF	% RSD	CCC	SPCC
	10.00	20.00	50.00	100.00	200.00					
Dichlorodifluoromethane	.12809	.22429	.28658	.30216	.22189	.113	.23260	29.920		
Chloromethane	.09252	.14976	.18487	.16572	.11692	.541	.14196	26.212	**	
Vinyl Chloride	.10537	.17290	.23164	.23021	.16149	.565	.18032	29.279		
Bromomethane	.22254	.34103	.44682	.42000	.26803	.610	.34368	26.921		
Chloroethane	.08501	.12702	.17823	.17051	.11663	.625	.13548	28.652		
Trichlorofluoromethane	.41900	.69209	.92291	.93082	.62321	.654	.71760	20.077		
1,1-Dichloroethane	.06188	.09141	.13367	.12950	.08716	.704	.10072	18.167	*	
Carbon Disulfide	.46848	.73594	1.02650	.97905	.66581	.718	.77516	23.720		
Acetone	.08229	.08956	.11180	.10148	.07208	.716	.09144	17.096		
ethylene Chloride	.38997	.40351	.40751	.35244	.22578	.753	.35584	21.325		
Trans-1,2-Dichloroethane	.15048	.23623	.30404	.29915	.21047	.777	.24007	26.753		
1,1-Dichloroethane	.41222	.58984	.76785	.76577	.55682	.816	.61850	24.409	**	
Vinyl Acetate	.04653	.05494	.05657	.06105	.06447	.826	.05672	12.016		
2,2-Dichloropropane	.25806	.35205	.40263	.40822	.27161	.870	.33851	20.942		
Cis-1,2-Dichloroethane	.23724	.32707	.40027	.41960	.30673	.872	.33818	21.623		
2-Butanone	.14707	.15500	.23037	.16874	.11419	.879	.15507	10.921		
Bromochloromethane	.27635	.38304	.46661	.44890	.34561	.895	.38410	20.208		
Chloroform	.56657	.74920	.93339	.98563	.76582	.905	.80012	20.772	*	
1,1,1-Trichloroethane	.47839	.66440	.80672	.85005	.65994	.919	.69190	21.138		
Carbon Tetrachloride	.52501	.71739	.87988	.93642	.72026	.935	.79499	21.258		
1,1-Dichloro-1-propene	.34772	.45561	.53303	.58626	.47214	.936	.47895	18.748		
1,2-Dichloroethane-d4(SURR)	.19316	.21950	.23368	.23659	.21241	.992	.21907	8.625		(Conc=50.0,50.0,5)
Benzene	.49027	.60793	.68268	.72930	.64120	.957	.63028	14.361		
1,2-Dichloroethane	.36793	.45288	.53208	.52799	.48714	.960	.47392	14.261		
Fluorobenzene(SURR)	.33700	.56088	.97397	1.01117	.97517	.987	.97165	2.767		(Conc=50.0,50.0,5)
Trichloroethane	.57446	.56177	.59367	.58511	.55636	1.025	.59347	8.844		
1,2-Dichloropropane	.28847	.36431	.43594	.46744	.33270	1.050	.37777	19.457	*	
Dibromomethane	.40061	.54226	.76461	.72057	.47935	1.064	.58148	26.859		
Bromodichloromethane	.62324	.86140	1.28119	1.24989	.85609	1.083	.97436	29.033		
2-Chloroethylvinylether	.15859	.26740	.34273	.29529	.21461	1.121	.24772	27.443		

RF - Response Factor (Subscript is amount in ug/L)

RRT - Average Relative Retention Time (RT Std/RT Istd)

RF - Average Response Factor

RPSD - Percent Relative Standard Deviation

CCC - Calibration Check Compounds (*) SPCC - System Performance Check Compounds (**)

Exam III Page 1 of 1

AR3094C

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Initial Calibration Data
HSL Compounds

Case No: _____ Instrument ID: HP 5970 MSD
Contractor: REGION III - CRL Calibration Date: 02/20/91
Contract No: ESAT

Minimum RF for SPCC is 0.300 Maximum % RSD for CCC is 30.00%

Compound	EA764 RF 10.00	EA765 RF 20.00	EA766 RF 50.00	EA767 RF 100.00	EA768 RF 200.00	RRT	RF	% RSD	CCC	SPCC
Trans-1,3-Dichloropropene	.55957	.88787	1.12848	1.05274	.70739	1.135	.86721	27.245		
4-Methyl-2-pentanone	.12786	.14395	.14888	.16715	.19284	.856	.15612	15.922		
Toluene-d8(SURR)	1.09082	1.06096	1.11816	1.18774	1.23931	.862	1.13940	6.402		(Conc=50.0,50.0,50.0,50.0)
Toluene	.50377	.55910	.57011	.64622	.72638	.868	.60111	14.390		
Cis-1,3-Dichloropropene	.36064	.40475	.43202	.46630	.53996	.891	.44065	15.310		
1,1,2-Trichloroethane	.38686	.44038	.44190	.48760	.56407	.908	.46416	14.278		
Tetrachloroethane	.78600	.89287	.86407	.95062	1.10584	.918	.91988	13.010		
1,3-Dichloropropene	.54350	.61196	.63976	.70433	.81705	.923	.66332	15.606		
2-Hexanone	.12385	.14152	.14928	.16254	.19158	.934	.15295	16.734		
Dibromochloromethane	1.00952	1.18930	1.24966	1.36901	1.50634	.944	1.26477	14.808		
1,2-Dibromoethane (ED8)	.74674	.82195	.84515	.92408	1.06777	.954	.88114	13.846		
Bromobenzene	.89744	.98504	1.02179	1.10049	1.26169	1.003	1.05329	13.052		**
1,1,1,2-Tetrachloroethane	.61410	.66571	.71444	.76680	.85363	1.012	.72294	12.790		
Ethylbenzene	1.15468	1.30403	1.32059	1.44001	1.43385	1.015	1.33063	8.763		*
m,p-Xylenes	.87228	.99926	.97177	1.08286	1.29605	1.028	1.04364	15.310		
o-Xylene	.85596	.98215	.97462	1.05144	1.15206	1.069	1.00324	10.853		
Styrene	1.48696	1.67994	1.70272	1.82123	2.09688	1.071	1.75753	12.767		
Bromoform	1.09299	1.22846	1.27830	1.38673	1.52845	1.089	1.30299	12.617		**
Isopropylbenzene	1.28645	1.45197	1.50237	1.67850	1.92281	1.109	1.56842	15.459		
Bromofluorobenzene(SURR)	.97492	1.01005	.97626	1.03002	1.15168	1.124	1.02859	7.064		(Conc=50.0,50.0,50.0,50.0)
Bromobenzene	.88089	.95170	1.05068	1.03667	1.11292	.881	1.00657	9.019		
1,1,2,2-Tetrachloroethane	.89910	.98582	1.05676	1.12997	1.26567	.885	1.06747	13.108		**
1,2,3-Trichloropropane	.58213	.71233	.80714	.77011	.89201	.887	.75725	15.349		
n-Propylbenzene	.49570	.51111	.55679	.56464	.64797	.892	.54724	12.994		
2-Chlorotoluene	.52691	.57473	.63171	.62536	.67337	.898	.60641	9.331		
4-Chlorotoluene	.54297	.60585	.61119	.63395	.70592	.907	.61990	9.468		
1,3,5-Trimethylbenzene	1.50781	1.65777	1.78122	1.76492	1.89876	.908	1.72250	8.551		
tert-Butylbenzene	1.45592	1.79010	1.88611	1.90797	2.07945	.935	1.82315	12.594		
1,2,4-Trimethylbenzene	1.47842	1.64607	1.78269	1.78109	1.95060	.940	1.72777	10.205		
sec-Butylbenzene	2.00252	2.26833	2.38224	2.53806	2.72399	.954	2.38295	11.493		

RF - Response Factor (Subscript is amount in ug/L)

RRT - Average Relative Retention Time (RT Std/RT Istd)

RF - Average Response Factor

RSD - Percent Relative Standard Deviation

CCC - Calibration Check Compounds (*) SPCC - System Performance Check Compounds (**)

AR309410

If the page filmed in this frame is not as readable or legible as this label, it is due to substandard color or condition of the original page.

Initial Calibration Data
HSL Compounds

Case No: _____ Instrument ID: HP 5970 MSD
Contractor: REGION III - CRL Calibration Date: ~~02-23-77~~ 02/20/91 SR
Contract No: ESAT

Minimum RF for SPCC is 0.300 Maximum % RSD for CCC is 30.00%

Compound	Laboratory ID: >EA764 >EA765 >EA766 >EA767 >EA768					RRT	RF	% RSD	CCC	SPCC
	RF	RF	RF	RF	RF					
	10.00	20.00	50.00	100.00	200.00					
1,3-Dichlorobenzene	1.37901	1.52028	1.58100	1.63094	1.80803	.962	1.58385	9.907		
1,4-Dichlorobenzene	1.49374	1.59017	1.70873	1.69968	1.81448	.970	1.69256	7.828		
p-Isopropyltoluene	1.70331	1.88936	2.01737	2.04193	2.15215	.968	1.96082	8.752		
1,2-Dichlorobenzene	1.33637	1.49998	1.50999	1.49694	1.98871	1.002	1.47840	6.238		
n-Butylbenzene	1.54097	1.72701	1.78624	1.80043	2.02146	1.004	1.77122	9.716		
1,2-Dibromo-3-chloropropane	.28529	.32039	.39984	.34792	.40020	1.071	.34273	12.567		
1,2,4-Trichlorobenzene	1.29610	1.41285	1.39396	1.35316	1.43925	1.145	1.37826	3.986		
Naphthalene	1.73240	1.89722	1.98662	1.92292	2.10027	1.166	1.92781	6.976		
Hexachlorobutadiene	1.17996	1.31583	1.24935	1.16952	1.17049	1.162	1.21943	5.310		
1,2,3-Trichlorobenzene	1.23074	1.31192	1.39004	1.23165	1.24169	1.189	1.27921	4.289		

RF - Response Factor (Subscript is amount in ug/L)

RRT - Average Relative Retention Time (RT Std/RT Istd)

RF - Average Response Factor

%RSD - Percent Relative Standard Deviation

CCC - Calibration Check Compounds (*) SPCC - System Performance Check Compounds (**)

AR309411

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IS + SS Mix prepared
2/13/91 SR

STD Mix prepared
2/14/91

Continuing Calibration Check
HSL Compounds

Case No: _____ Calibration Date: 02/25/91
Contractor: REGION III - CRL Time: 10:17
Contract No: ESAT Laboratory ID: 1EA795
Instrument ID: HP 5970 MSD Initial Calibration Date: 02/27/91 02/30/91 SR

Minimum RF for SPC is 0.300 Maximum % Diff for CCC is 25.00%

Compound	RF	RF	% Diff	CCC SPC
Dichlorodifluoromethane	.23260	.18269	21.46	
Chloromethane	.14196	.10414	26.64	**
Vinyl Chloride	.18032	.13529	24.97	*
Bromomethane	.34368	.26307	23.46	
Chloroethane	.13548	.10854	19.89	
Trichlorofluoromethane	.71760	.59660	22.44	
1,1-Dichloroethane	.10072	.08201	18.58	*
Carbon Disulfide	.77516	.61377	20.82	
Acetone	.09144	.10699	17.00	
Methylene Chloride	.35984	.23692	35.53	
Trans-1,2-Dichloroethane	.24007	.19203	20.01	
1,1-Dichloroethane	.61850	.50428	18.47	**
Vinyl Acetate	.05672	.05770	1.74	
2,2-Dichloropropane	.33891	.33610	.71	
Cis-1,2-Dichloroethane	.33818	.28393	16.04	
2-Butanone	.15507	.17609	13.56	
Bromochloromethane	.38410	.31150	18.90	
Chloroform	.80012	.74749	6.58	*
1,1,1-Trichloroethane	.69190	.65086	5.93	
Carbon Tetrachloride	.75499	.71843	4.84	
1,1-Dichloro-1-propene	.47895	.45677	4.63	
1,2-Dichloroethane-d4(SURR)	.21907	.21198	3.24	
Benzene	.63028	.58416	7.32	
1,2-Dichloroethane	.47352	.49176	3.85	
Fluorobenzene(SURR)	.97165	.94113	3.14	
Trichloroethene	.55347	.52259	5.98	
1,2-Dichloropropane	.37777	.36925	18.14	*
Dibromomethane	.58148	.49848	15.63	
Bromodichloromethane	.57436	.85438	12.31	
2-Chloroethylvinylether	.24772	.19444	21.91	
Trans-1,3-Dichloropropene	.84721	.68817	21.97	
4-Methyl-2-pentanone	.15612	.15742	1.73	

RF - Response Factor from daily standard file at 50.00 ug/L

AR309412

RF - Average Response Factor from Initial Calibration Form VI

% Diff - % Difference from original average or curve

CCC - Calibration Check Compounds (*) SPC - System Performance Check Compounds (**)

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Continuing Calibration Check
HSL Compounds

Case No: _____ Calibration Date: 02/25/91
Contractor: REGION III - CRL Time: 10:17
Contract No: ESAT Laboratory ID: 1EA295
Instrument ID: HP 5970 MSD Initial Calibration Date: ~~02/21/91~~ 02/20/91 SR

Minimum RF for SPCC is 0.300 Maximum % Diff for CCC is 25.00%

Compound	RF	RF	%Diff	CCC SPCC
Toluene-d8 (SURR)	1.13940	1.14561	.55	
Toluene	.60111	.62204	3.48	*
Cis-1,3-Dichloropropene	.44065	.47109	6.91	
1,1,2-Trichloroethane	.44416	.49276	2.46	
Tetrachloroethane	.91988	.89042	3.20	
1,3-Dichloropropane	.66332	.66101	.35	
2-Hexanone	.19295	.16076	5.11	
Dibromochloromethane	1.26477	1.30229	2.97	
1,2-Dibromoethane (EDB)	.88114	.89088	3.43	
Chlorobenzene	1.05329	1.01753	3.40	**
1,1,1,2-Tetrachloroethane	.72294	.74688	3.31	
Ethylbenzene	1.33063	1.40423	5.53	*
m,p-Xylenes	1.04364	1.05362	.96	
o-Xylene	1.00324	1.01645	1.32	
Styrene	1.79793	1.80869	2.91	
Bromoform	1.30299	1.25331	3.81	**
Isopropylbenzene	1.56842	1.57436	.38	
Bromofluorobenzene (SURR)	1.02859	.98482	4.25	
Bromobenzene	1.00657	1.02649	1.98	
1,1,2,2-Tetrachloroethane	1.04747	1.10443	3.46	**
1,2,3-Trichloropropane	.79275	.71615	4.86	
n-Propylbenzene	.54724	.56328	2.93	
2-Chlorotoluene	.60641	.61331	1.14	
4-Chlorotoluene	.61998	.62254	.43	
1,3,5-Trimethylbenzene	1.72298	1.79833	4.40	
tert-Butylbenzene	1.82315	1.94851	6.44	
1,2,4-Trimethylbenzene	1.72777	1.89576	7.41	
sec-Butylbenzene	2.38295	2.55822	7.02	
1,3-Dichlorobenzene	1.98385	1.61997	2.28	
1,4-Dichlorobenzene	1.69256	1.78384	3.10	
p-Isopropyltoluene	1.94082	2.12697	8.47	
1,2-Dichlorobenzene	1.47848	1.53848	3.92	

RF - Response Factor from daily standard file at 50.00 ug/L

RF - Average Response Factor from Initial Calibration Form VI

%Diff - % Difference from original average or curve

CCC - Calibration Check Compounds (*) SPCC - System Performance Check Compounds (**)

AR309413

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Continuing Calibration Check
HSL Compounds

Case No: _____ Calibration Date: 02/25/91
Contractor: REGION III - CRL Time: 10:17
Contract No: ESAT Laboratory ID: EA795
Instrument ID: HP 5970 MSD Initial Calibration Date: ~~02/21/91~~ 02/20/91SR

Minimum RF for SPCC is 0.300

Maximum % Diff for CCC is 25.00%

Compound	RF	RF	%Diff	CCC	SPCC
n-Butylbenzene	1.77122	1.91458	8.09		
1,2-Dibromo-3-chloropropane	.34273	.35895	4.73		
1,2,4-Trichlorobenzene	1.37826	1.40376	1.85		
Naphthalene	1.92781	1.90823	1.02		
Hexachlorobutadiene	1.21543	1.28740	5.92		
1,2,3-Trichlorobenzene	1.27321	1.25507	1.42		

RF - Response Factor from daily standard file at 50.00 ug/L

- Average Response Factor from Initial Calibration Form VI

Diff - % Difference from original average or curve

CC - Calibration Check Compounds (*) SPCC - System Performance Check Compounds (**)

AR309414

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IS+SS Mix prep
2/13

STD Mix prep
2/14

Continuing Calibration Check
HSL Compounds

Case No: _____ Calibration Date: 02/26/91
Contractor: REGION III - CRL Time: 12:28
Contract No: ESAT Laboratory ID: EAB10
Instrument ID: HP 5970 MSD Initial Calibration Date: ~~12/11/90~~ 02/26/91 SR

Minimum RF for SPC is 0.300 Maximum % Diff for CCC is 25.00%

Compound	RF	RF	%Diff	CCC	SPC
Dichlorodifluoromethane	.23260	.20149	13.37		
Chloromethane	.14196	.09757	31.27	**	
Vinyl Chloride	.18032	.13997	24.60	*	
Bromomethane	.34368	.25803	24.92		
Chloroethane	.13548	.10639	21.47		
Trichlorofluoromethane	.71760	.59931	16.49		
1,1-Dichloroethane	.10072	.08059	19.99	*	
Carbon Disulfide	.77516	.62683	19.14		
Acetone	.09144	.07124	22.09		
Methylene Chloride	.39984	.27827	23.04		
Trans-1,2-Dichloroethane	.24007	.19827	17.41		
1,1-Dichloroethane	.61890	.52651	14.87	**	
Vinyl Acetate	.09672	.09766	1.66		
2,2-Dichloropropane	.33851	.35999	9.16		
Cis-1,2-Dichloroethane	.33818	.29393	13.08		
2-Butanone	.19507	.13946	10.67		
Bromochloromethane	.38410	.29403	23.45		
Chloroform	.80012	.79890	9.15	*	
1,1,1-Trichloroethane	.69190	.69411	.32		
Carbon Tetrachloride	.79499	.76446	1.92		
1,1-Dichloro-1-propene	.47899	.47788	.22		
1,2-Dichloroethane-d4(SURR)	.21987	.23393	6.40		
Benzene	.63028	.61096	3.06		
1,2-Dichloroethane	.47392	.52783	11.47		
Fluorobenzene(SURR)	.97145	.96345	.84		
Trichloroethane	.59347	.48358	12.43		
1,2-Dichloropropane	.37777	.32763	13.43	*	
Dibromomethane	.58148	.49663	14.99		
Bromodichloromethane	.97436	.83239	14.57		
2-Chloroethylvinylether	.24772	.20913	17.19		
Trans-1,3-Dichloropropane	.86721	.78498	10.72		
4-Methyl-2-pentanone	.19612	.15377	1.50		

RF - Response Factor from daily standard file at 50.00 ug/L

RF - Average Response Factor from Initial Calibration Form VI

%Diff - % Difference from original average or curve

CCC - Calibration Check Compounds (*) SPC - System Performance Check Compounds (**)

AR309415

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Continuing Calibration Check
HSL Compounds

Case No: _____ Calibration Date: 02/26/91
Contractor: REGION III - CRL Time: 12:28
Contract No: ESAT Laboratory ID: 1E9810
Instrument ID: HP 5970 MSD Initial Calibration Date: ~~02/21/91~~ 02/26/91 SR

Minimum RF for SPC is 0.300

Maximum % Diff for CCC is 25.00%

Compound	RF	RF	%Diff	CCC SPC
Toluene-d8(SURR)	1.13940	1.18045	3.60	
Toluene	.60111	.61651	2.56	*
Cis-1,3-Dichloropropene	.44065	.46324	5.13	
1,1,2-Trichloroethane	.46416	.44948	3.16	
Tetrachloroethane	.91988	.80500	12.49	
1,3-Dichloropropene	.66332	.67273	1.42	
2-Hexanone	.15295	.14190	7.22	
Dibromochloromethane	1.26477	1.22492	3.15	
1,2-Dibromoethane (EDB)	.88114	.83839	4.85	
Chlorobenzene	1.05329	.99610	5.43	**
1,1,1,2-Tetrachloroethane	.72294	.72166	.18	
Ethylbenzene	1.33063	1.42948	7.43	*
m,p-Xylenes	1.04364	1.04228	.13	
o-Xylene	1.00324	1.02106	1.78	
Styrene	1.75753	1.83791	4.57	
Bromoform	1.30299	1.19514	8.28	**
Isopropylbenzene	1.56842	1.66033	5.86	
Bromofluorobenzene(SURR)	1.02859	1.00971	1.83	
Bromobenzene	1.00457	1.01932	.87	
1,1,2,2-Tetrachloroethane	1.06747	1.20341	12.74	**
1,2,3-Trichloropropane	.79275	.89489	10.88	
n-Propylbenzene	.54724	.60231	10.06	
2-Chlorotoluene	.60441	.64851	6.94	
4-Chlorotoluene	.61990	.63244	2.02	
1,3,5-Trimethylbenzene	1.72250	2.00800	16.57	
tert-Butylbenzene	1.82315	2.04453	12.14	
1,2,4-Trimethylbenzene	1.72777	2.02677	17.31	
sec-Butylbenzene	2.38295	2.78740	17.06	
1,3-Dichlorobenzene	1.50305	1.61596	2.03	
1,4-Dichlorobenzene	1.65256	1.73925	5.25	
p-Isopropyltoluene	1.96082	2.24654	14.57	
1,2-Dichlorobenzene	1.47840	1.53286	3.68	

RF - Response Factor from daily standard file at 50.00 ug/L

- Average Response Factor from Initial Calibration Form VI

%Diff - % Difference from original average or curve

CCC - Calibration Check Compounds (*) SPC - System Performance Check Compounds (**)

AR309416

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Continuing Calibration Check
HSL Compounds

Case No: _____ Calibration Date: 02/26/91
Contractor: REGION III - CRL Time: 12:28
Contract No: ESAT Laboratory ID: EAB10
Instrument ID: HP 5970 MSD Initial Calibration Date: ~~02/21/91~~ 02/26/91 *SR*

Minimum $\overline{RF}$ for SPCC is 0.300 Maximum % Diff for CCC is 25.00%

Compound	$\overline{RF}$	RF	%Diff	CCC	SPCC
n-Butylbenzene	1.77122	2.17089	22.56		
1,2-Dibromo-3-chloropropane	.34273	.39095	14.07		
1,2,4-Trichlorobenzene	1.37826	1.33477	3.16		
Naphthalene	1.92781	1.92632	.08		
Hexachlorobutadiene	1.21943	1.27906	4.91		
1,2,3-Trichlorobenzene	1.27321	1.27692	.29		

RF - Response Factor from daily standard file at 50.00 ug/L

$\overline{RF}$ - Average Response Factor from Initial Calibration Form VI

%Diff - % Difference from original average or curve

CCC - Calibration Check Compounds (*) SPCC - System Performance Check Compounds (**)

Form VII Page 3 of 3

C-144

AR309417

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QUANT REPORT

Operator ID: ESAT Quant Rev: 6 Quant Time: 910225 14:30
 Output File: ^EA799:1:QT Injected at: 910225 13:49
 Data File: >EA799:1:03 Dilution Factor: 1.00000
 Name: 910221-07 CHEM SOLU
 Misc: MON W 26 A, 5ML SAMPLE + 5UL IS+SS

ID File: VQAID1:ME
 Title: DB-624 DIRECT 6PSI
 Last Calibration: 910225 11:30

	Compound	R.T.	Q ion	Area	Conc	Units	q
1)	*1,4-Difluorobenzene	16.58	114.0	75909	50.00	ug/L	68
3)	Chloromethane	10.43	50.0	3096	24.39	ug/L	97
8)	Acetone	11.67	43.0	15496	95.41	ug/L	61
7)	2-Butanone	14.55	43.0	910	3.40	ug/L	90
9)	Chloroform	15.00	83.0	1174	1.03	ug/L	89
13)	1,2-Dichloroethane-d4(SURR)	15.79	67.0	18060	112.24	ug/L	99
14)	Benzene	15.86	78.0	28374	31.99	ug/L	99
5)	1,2-Dichloroethane	15.92	62.0	10570	1.42	ug/L	96
16)	Fluorobenzene(SURR)	16.36	96.0	71645	100.29	ug/L	86
3)	*Chlorobenzene-d5	22.43	117.0	59646	50.00	ug/L	97
1)	Toluene-d8(SURR)	19.35	98.0	72151	105.59	ug/L	98
9)	Tetrachloroethene	20.58	166.0	1393	1.31	ug/L	89
6)	Ethylbenzene	22.85	91.0	862	.51	ug/L	74
1)	Isopropylbenzene	24.87	105.0	29736	15.83	ug/L	92
2)	Bromofluorobenzene(SURR)	25.22	95.0	62030	105.60	ug/L	92
3)	*1,2-Dichlorobenzene-D4	29.00	152.0	44863	50.00	ug/L	91
8)	1,3,5-Trimethylbenzene	26.14	105.0	753	.47	ug/L	100
1)	tert-Butylbenzene	27.12	119.0	608	.35	ug/L	93
2)	1,2,4-Trimethylbenzene	27.24	105.0	730	.44	ug/L	92
3)	sec-Butylbenzene	27.67	105.0	20073	8.77	ug/L	89
8)	n-Butylbenzene	29.12	91.0	10684	6.22	ug/L	100
1)	Naphthalene	33.83	128.0	236	.55	ug/L	100

* Compound is ISTD

AR309418

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QUANT REPORT - False positives deleted, #47 names
integrated and 2nd TIC set

Operator ID: ESAT
Output File: ^EA799::QT
Data Filter: >EA799::03
Name: 910221-07 CHEM SOLU
Misc: MON W 26 A, 5ML SAMPLE + 5UL IS+SS

Quant Rev: 6 Quant Time: 910225 14:50
Injected at: 910225 13:49
Dilution Factor: 1.00000

ID File: UDAID1::ME
Title: DB-624 DIRECT 6PSI
Last Calibration: 910225 11:30

	Compound	R.T.	Scan#	Area	Conc	Units
1)	*1,4-Difluorobenzene	16.58	1208	79909	50.00	ug/L
17)	2-Butanone	14.55	1034	910	3.40	ug/L
23)	1,2-Dichloroethane-d4(SURR)	15.79	1141	18060	112.24	ug/L
24)	Benzene	15.86	1147	28374	31.99	ug/L
26)	Fluorobenzene(SURR)	16.36	1189	71645	100.29	ug/L
33)	*Chlorobenzene-d5	22.43	1709	59646	50.00	ug/L
35)	Toluene-d8(SURR)	19.35	1445	72151	105.59	ug/L
39)	Tetrachloroethane	20.58	1951	1393	1.31	ug/L
47)	m&p-Xylenes	23.06	1763	249M	.40	ug/L
51)	Isopropylbenzene	24.87	1918	29736	15.83	ug/L
52)	Bromofluorobenzene(SURR)	25.22	1948	62030	105.60	ug/L
53)	*1,2-Dichlorobenzene-D4	29.00	2272	44863	50.00	ug/L
61)	tert-Butylbenzene	27.12	2111	608	.35	ug/L
62)	1,2,4-Trimethylbenzene	27.24	2121	730	.44	ug/L
63)	sec-Butylbenzene	27.67	2158	20073	8.77	ug/L
68)	n-Butylbenzene	29.12	2282	10684	6.22	ug/L

* Compound is 1STD

AR309419

C-16

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QUANT REPORT

Operator: ID: ESAT

Quant Rev: 6

Quant Time: 910225 15:17

Output File: ^EAB000::QT

Injected at: 910225 14:36

Data File: >EAB000::D3

Dilution Factor: 1.00000

Name: 910221-08 CHEM SOLV

Misc: MON W 33 A, 5ML SAMPLE + 5UL IS+SS

ID File: UDAID1::ME

Title: DB-624 DIRECT 6PS1

Last Calibration: 910225 11:30

	Compound	R.T.	Q ion	Area	Conc	Units	q
1)	*1,4-Difluorobenzene	16.57	114.0	75713	50.00	ug/L	70
16)	Cis-1,2-Dichloroethene	14.46	96.0	595	1.38	ug/L	93
19)	Chloroform	14.99	83.0	1579	1.40	ug/L	75
20)	1,1,1-Trichloroethane	15.23	97.0	11456	11.62	ug/L	94
21)	Carbon Tetrachloride	15.22	117.0	1494	1.37	ug/L	89
23)	1,2-Dichloroethane-d4(SURR)	15.79	67.0	18186	113.31	ug/L	98
26)	Fluorobenzene(SURR)	16.35	96.0	71255	100.00	ug/L	89
27)	Trichloroethene	17.00	130.0	112137	141.71	ug/L	97
33)	*Chlorobenzene-d5	22.41	117.0	58113	50.00	ug/L	96
35)	Toluene-d8(SURR)	19.32	98.0	68093	102.22	ug/L	89
9)	Tetrachloroethene	20.57	166.0	3568	3.45	ug/L	83
2)	Bromofluorobenzene(SURR)	25.21	95.0	59479	103.93	ug/L	86
33)	*1,2-Dichlorobenzene-D4	28.98	182.0	43320	50.00	ug/L	92

* Compound is ISTD

AR309420

C-17

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QUANT REPORT

Operator ID: ESAT
 Output File: ^EAB01::QT
 Data File: >EAB01::D3
 Name: 910221-09 CHEM SOLU
 Misc: MON W 330, 5ML SAMP * 5UL IS+SS

Quant Rev: 6 Quant Time: 910225 16:05
 Injected at: 910225 15:24
 Dilution Factor: 1.00000

ID File: UDAID1::ME
 Title: DB-624 DIRECT 6PSI
 Last Calibration: 910225 11:30

	Compound	R.T.	Scan#	Area	Conc	Units
1)	*1,4-Difluorobenzene	16.54	1208	73845	50.00	ug/L
16)	Cis-1,2-Dichloroethene	14.42	1026	607	1.45	ug/L ✓
19)	Chloroform	14.97	1073	1460	1.32	ug/L ✓
20)	1,1,1-Trichloroethane	15.19	1092	11163	11.61	ug/L ✓
21)	Carbon Tetrachloride	15.20	1093	1482	1.40	ug/L ²⁵ 0.0001
23)	1,2-Dichloroethane-d4(SURR)	15.76	1141	17867M	114.14	ug/L
26)	Fluorobenzene(SURR)	16.32	1189	70114	100.89	ug/L
27)	Trichloroethene	16.98	1245	109906	142.40	ug/L ✓
33)	*Chlorobenzene-d5	22.38	1708	58293	50.00	ug/L
35)	Toluene-d8(SURR)	19.29	1443	69111	103.49	ug/L
39)	Tetrachloroethene	20.55	1551	3669	3.53	ug/L
52)	Bromofluorobenzene(SURR)	25.18	1948	58075	101.16	ug/L
53)	*1,2-Dichlorobenzene-D4	28.96	2272	44457	50.00	ug/L

* Compound is ISTD

AR30942

C-18

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QUANT REPORT

Operator: TD: ESAT Quant Rev: 6 Quant Time: 910225 13:42
 Output File: \EA798::QT Injected at: 910225 13:01
 Data File: >EA798::D3 Dilution Factor: 1.00000
 Name: 910221-10 CHEM SOLV
 Misc: TRIP BLANK, 5ML SAMPLE & 5UL IS + SS

ID File: UOAI01::ME
 Title: DB-624 DIRECT 6PSI
 Last Calibration: 910225 11:30

	Compound	R.T.	Q ion	Area	Conc	Units	q
1)	*1,4-Difluorobenzene	16.44	114.0	74221	50.00	ug/L	70
3)	Chloromethane	8.79	50.0	929	6.01	ug/L ✓	80
.0)	Acetone	11.76	43.0	1322	8.32	ug/L ✓	97
.7)	2-Butanone	14.42	43.0	618	2.36	ug/L ✓	87
.9)	Chloroform	14.86	83.0	5714	5.15	ug/L ✓	93
13)	1,2-Dichloroethane-d4(SURR)	15.66	67.0	17650	112.78	ug/L	96
16)	Fluorobenzene(SURR)	16.23	96.0	70462	100.87	ug/L	89
10)	Bromodichloromethane	17.84	83.0	526	.41	ug/L ✓	93
13)	*Chlorobenzene-d5	22.30	117.0	58938	50.00	ug/L	93
15)	Toluene-d8(SURR)	19.20	98.0	69346	102.70	ug/L	97
2)	Bromofluorobenzene(SURR)	25.12	95.0	62489	107.66	ug/L	89
13)	*1,2-Dichlorobenzene-D4	28.94	152.0	43864	50.00	ug/L	89
18)	n-Butylbenzene	29.07	91.0	873	.52	ug/L	98.100 1/441

* Compound is ISTD

AR309422

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QUANT REPORT

Operator ID: ESAT Quant Rev: 6 Quant Time: 910225 12:46
 Output File: ^EA797:1QT Injected at: 910225 12:05
 Data File: ^EA797:1D3 Dilution Factor: 1.00000
 Name: UOA METHOD BLANK-7/022/-11-CAL Lab Blank, R.
 Misc: 5UL IS+SS IN 5ML OFW

ID File: UOAID1:ME
 Title: DB-624 DIRECT 6PSI
 Last Calibration: 910225 11:30

Compound	R.T.	Q ion	Area	Conc	Units
1) *1,4-Difluorobenzene	16.58	114.0	78872	50.00	ug/L
11) Methylene Chloride	12.44	84.0	2091	5.60	ug/L ✓
23) 1,2-Dichloroethane-d4(SURR)	15.78	67.0	17771	106.29	ug/L
26) Fluorobenzene(SURR)	16.36	96.0	73173	98.58	ug/L
33) *Chlorobenzene-d5	22.41	117.0	62346	50.00	ug/L
35) Toluene-d8(SURR)	19.32	98.0	70940	99.32	ug/L
52) Bromofluorobenzene(SURR)	25.21	95.0	62493	101.78	ug/L
53) *1,2-Dichlorobenzene-D4	28.99	152.0	46159	50.00	ug/L

* Compound is ISTD

AR309423

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QUANT REPORT

Operator: 40: ESAT Quant Rev: 6 Quant Time: 910226 17:09
 Output File: ^EAB13:1QT Injected at: 910226 16:28
 Data File: >EAB13:1O3 Dilution Factor: 1.00000
 Name: 910221-08MS CHEM SOL
 Misc: MON W 33 A, 5ML SAMPLE + 5UL IS+SS + 5UL STD MIX

ID File: VQA101:ME
 Title: DB-624 DIRECT 6PS1
 Last Calibration: 910226 13:53

	Compound	R.T.	Q ion	Area	Conc	Units	q
1)	*1,4-Difluorobenzene	16.63	114.0	70546	50.00	ug/L	67
2)	Dichlorodifluoromethane	7.36	85.0	5443^	19.15	ug/L	93
3)	Chloromethane	8.55	50.0	6578	47.78	ug/L	96
4)	Vinyl Chloride	9.11	62.0	9194	47.92	ug/L	93
5)	Bromomethane	9.97	94.0	17108	46.99	ug/L	84
6)	Chloroethane	10.26	64.0	7273	48.45	ug/L	98
7)	Trichlorofluoromethane	10.79	101.0	47549	56.23	ug/L	97
8)	1,1-Dichloroethane	11.66	96.0	5880	51.71	ug/L	94
9)	Carbon Disulfide	11.88	76.0	42541	48.10	ug/L	100
10)	Acetone	11.88	43.0	4932	49.07	ug/L	97
11)	Methylene Chloride	12.50	84.0	14646	37.90	ug/L	74
12)	Trans-1,2-Dichloroethene	12.91	96.0	13335	47.67	ug/L	85
13)	1,1-Dichloroethane	13.56	63.0	38338	51.61	ug/L	95
14)	Vinyl Acetate	13.74	86.0	4012	49.32	ug/L	77
15)	2,2-Dichloropropane	14.47	77.0	26406	52.57	ug/L	90
16)	Cis-1,2-Dichloroethene	14.49	96.0	21183	51.08	ug/L	91
17)	2-Butanone	14.60	43.0	9610	48.84	ug/L	96
18)	Bromochloromethane	14.88	128.0	20995	50.61	ug/L	81
19)	Chloroform	15.04	83.0	57056	53.29	ug/L	96
20)	1,1,1-Trichloroethane	15.28	97.0	61965	63.27	ug/L	78
21)	Carbon Tetrachloride	15.55	117.0	55587	51.40	ug/L	97
22)	1,1-Dichloro-1-propene	15.57	75.0	34507	51.18	ug/L	96
23)	1,2-Dichloroethane-d4(SURR)	15.84	67.0	17311	105.08	ug/L	95
24)	Benzene	15.91	78.0	41629	48.29	ug/L	99
25)	1,2-Dichloroethane	15.97	62.0	39382	52.88	ug/L	97
26)	Fluorobenzene(SURR)	16.41	96.0	66020	97.13	ug/L	88
27)	Trichloroethane	17.05	130.0	131036	192.05	ug/L	91
28)	1,2-Dichloropropane	17.46	63.0	22761	49.33	ug/L	94
29)	Bromodichloromethane	18.00	83.0	61697	52.53	ug/L	91
30)	Trans-1,3-Dichloropropene	18.87	75.0	48394	29.20	ug/L	98
31)	*Chlorobenzene-d5	22.46	117.0	52105	50.00	ug/L	97
32)	4-Methyl-2-pentanone	19.25	58.0	8339	52.04	ug/L	85
33)	Toluene-d8(SURR)	19.38	98.0	62823	102.14	ug/L	92
34)	Toluene	19.50	92.0	32495	50.58	ug/L	95
35)	Cis-1,3-Dichloropropene	20.02	75.0	26614	77.18	ug/L	97
36)	1,1,2-Trichloroethane	20.40	97.0	25095	50.58	ug/L	99
37)	Tetrachloroethane	20.64	166.0	45797	54.59	ug/L	93
38)	1,3-Dichloropropane	20.73	76.0	37453	53.42	ug/L	87
39)	2-Hexanone	20.99	58.0	8069	54.57	ug/L	78
40)	Dibromochloromethane	21.21	129.0	68100	53.35	ug/L	97
41)	1,2-Dibromoethane (EDB)	21.42	107.0	46348	53.05	ug/L	95
42)	Chloroethane	22.53	112.0	53546	51.58	ug/L	72
43)	1,1,1,2-Tetrachloroethane C-11	22.75	131.0	78007	50.57	ug/L	97

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	Compound	R.T.	Q ion	Area	Conc	Unit
46)	Ethylbenzene	22.81	91.0	76118	51.10	ug/L
47)	m&p-Xylenes	23.10	106.0	54695	100.71	ug/L
48)	o-Xylene	24.00	106.0	54910	103.21	ug/L
49)	Styrene	24.06	104.0	95482	99.70	ug/L
50)	Bromoform	24.46	173.0	64333	51.65	ug/L
51)	Isopropylbenzene	24.91	105.0	86812	50.17	ug/L
52)	Bromofluorobenzene (SURR)	25.25	95.0	55463	105.42	ug/L
53)	*1,2-Dichlorobenzene-D4	25.83	152.0	37620	50.00	ug/L
54)	Bromobenzene	25.57	156.0	37743	49.41	ug/L
55)	1,1,2,2-Tetrachloroethane	25.70	83.0	47498	52.46	ug/L
56)	1,2,3-Trichloropropane	25.75	75.0	31520	46.81	ug/L
57)	n-Propylbenzene	25.91	120.0	22419	49.47	ug/L
58)	2-Chlorotoluene	26.08	126.0	24267	49.73	ug/L
59)	4-Chlorotoluene	26.36	126.0	24996	52.53	ug/L
60)	1,3,5-Trimethylbenzene	26.37	105.0	75548	50.00	ug/L
61)	tert-Butylbenzene	27.16	119.0	73103	47.52	ug/L
62)	1,2,4-Trimethylbenzene	27.27	105.0	77722	50.97	ug/L
63)	sec-Butylbenzene	27.71	105.0	104907	49.98	ug/L
64)	1,3-Dichlorobenzene	27.93	146.0	62391	51.31	ug/L
65)	1,4-Dichlorobenzene	28.16	146.0	64240	49.09	ug/L
66)	p-Isopropyltoluene	28.09	119.0	88380	52.29	ug/L
67)	1,2-Dichlorobenzene	29.07	146.0	56754	49.21	ug/L
68)	n-Butylbenzene	29.13	91.0	81995	50.20	ug/L
69)	1,2-Dibromo-3-chloropropane	31.12	75.0	15306	52.03	ug/L
70)	1,2,4-Trichlorobenzene	33.25	180.0	49427	49.22	ug/L
71)	Naphthalene	33.87	128.0	74921	51.69	ug/L
72)	Hexachlorobutadiene	33.76	225.0	45031	46.94	ug/L
73)	1,2,3-Trichlorobenzene	34.53	180.0	47492	49.43	ug/L

* Compound is ISTD

AR309425

C-22

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QUANT REPORT

Operator ID: ESAT Quant Rev: 6 Quant Time: 910226 18:45
 Output File: ^EAB15::QT Injected at: 910226 18:03
 Data File: >EAB15::D3 Dilution Factor: 1.00000
 Name: 910221-08MSD CHEM SO
 Misc: MON W 33A, 5ML SAMP + 5UL IS+SS + 5UL STD MIX

ID File: UDAID1:IME
 Title: DB-624 DIRECT 6PSI
 Last Calibration: 910226 13:53

	Compound	R.T.	Scan#	Area	Conc	Units	q
1)	*1,4-Difluorobenzene	16.60	1209	67842	50.00	ug/L	67
2)	Dichlorodifluoromethane	7.38	419	10278	37.59	ug/L	94
3)	Chloromethane	9.03	561	6862	51.83	ug/L	97
4)	Vinyl Chloride	9.50	601	9359	50.73	ug/L	99
5)	Bromomethane	10.26	666	17200	49.13	ug/L	89
6)	Chloroethane	10.50	687	7482	51.83	ug/L	95
7)	Trichlorofluoromethane	10.98	728	48725	59.92	ug/L	99
8)	1,1-Dichloroethane	11.79	797	5579	51.02	ug/L	93
9)	Carbon Disulfide	12.00	815	40730	47.89	ug/L	100
10)	Acetone	11.96	812	5974	61.80	ug/L	90
11)	Methylene Chloride	12.57	864	14285	38.44	ug/L	83
12)	Trans-1,2-Dichloroethene	12.94	896	13552	50.38	ug/L	91
13)	1,1-Dichloroethane	13.58	951	38283	53.59	ug/L	99
14)	Vinyl Acetate	13.74	964	3867	49.43	ug/L	76
15)	2,2-Dichloropropane	14.47	1027	22561	46.71	ug/L	92
16)	Cis-1,2-Dichloroethene	14.49	1029	19878	49.84	ug/L	89
17)	2-Butanone	14.59	1037	10031	53.01	ug/L	91
19)	Chloroform	15.02	1074	53727	52.18	ug/L	95
20)	1,1,1-Trichloroethane	15.25	1094	54152	57.50	ug/L	75
21)	Carbon Tetrachloride	15.52	1117	49182	47.29	ug/L	92
22)	1,1-Dichloro-1-propene	15.53	1118	32988	50.88	ug/L	97
23)	1,2-Dichloroethane-d4(SURR)	15.83	1143	15682	98.98	ug/L	97
24)	Benzene	15.88	1148	41857	50.49	ug/L	99
25)	1,2-Dichloroethane	15.95	1154	36098	50.40	ug/L	96
26)	Fluorobenzene(SURR)	16.39	1191	63680	97.43	ug/L	91
27)	Trichloroethene	17.03	1246	126104	192.19	ug/L	92
28)	1,2-Dichloropropane	17.44	1281	23344	52.61	ug/L	93
30)	Bromodichloromethane	17.97	1327	62074	54.96	ug/L	93
32)	Trans-1,3-Dichloropropene	18.84	1401	47169	29.59	ug/L	98
33)	*Chlorobenzene-d5	22.40	1706	47659	50.00	ug/L	98
34)	4-Methyl-2-pentanone	19.21	1433	8296	56.60	ug/L	84
35)	Toluene-d8(SURR)	19.33	1443	60902M	108.25	ug/L	96
36)	Toluene	19.47	1455	33733	52.40	ug/L	97
37)	Cis-1,3-Dichloropropene	19.97	1498	26000	82.40	ug/L	96
38)	1,1,2-Trichloroethane	20.34	1530	23854	55.68	ug/L	99
39)	Tetrachloroethene	20.58	1550	46401	60.47	ug/L	86
40)	1,3-Dichloropropene	20.68	1559	37483	58.45	ug/L	87
41)	2-Hexanone	20.94	1581	7397	54.69	ug/L	74
42)	Dibromochloromethane	21.15	1599	62843	53.82	ug/L	95
43)	1,2-Dibromoethane (EOB)	21.36	1617	45887	57.42	ug/L	97
44)	Chlorobenzene	22.47	1712	51308	54.04	ug/L	77
45)	1,1,1,2-Tetrachloroethane	22.49	1710	36000	50.00	ug/L	90

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48)	o-Xylenes	22.70	1837	47227	101.10 ug/L
49)	Styrene	23.99	1842	87588	99.99 ug/L
50)	Bromoform	24.39	1876	58804	51.62 ug/L
51)	Isopropylbenzene	24.84	1915	78147	49.38 ug/L
52)	Bromofluorobenzene (SURR)	25.19	1945	45334	94.21 ug/L
53)	*1,2-Dichlorobenzene-O4	28.99	2269	36048	50.00 ug/L
54)	Bromobenzene	25.50	1971	32068	43.81 ug/L
55)	1,1,2,2-Tetrachloroethane	25.64	1983	43542	50.19 ug/L
56)	1,2,3-Trichloropropane	25.70	1988	30427	47.16 ug/L
57)	n-Propylbenzene	25.85	2001	18845	43.40 ug/L
58)	2-Chlorotoluene	26.01	2015	20087	42.96 ug/L
59)	4-Chlorotoluene	26.30	2039	20513	44.99 ug/L
60)	1,3,5-Trimethylbenzene	26.32	2041	64410	44.49 ug/L
61)	tert-Butylbenzene	27.10	2108	71990	48.84 ug/L
62)	1,2,4-Trimethylbenzene	27.23	2119	73528	50.32 ug/L
63)	sec-Butylbenzene	27.65	2155	99972	49.71 ug/L
64)	1,3-Dichlorobenzene	27.87	2174	54405	46.70 ug/L
65)	1,4-Dichlorobenzene	28.11	2194	64474	51.42 ug/L
66)	p-Isopropyltoluene	28.05	2189	85531	52.81 ug/L
67)	1,2-Dichlorobenzene	29.02	2272	55503	50.22 ug/L
68)	n-Butylbenzene	29.09	2278	82339	52.61 ug/L
69)	1,2-Dibromo-3-chloropropane	31.04	2445	16197	57.47 ug/L
70)	1,2,4-Trichlorobenzene	33.18	2628	49042	50.96 ug/L
71)	Naphthalene	33.80	2681	77695	55.94 ug/L
72)	Hexachlorobutadiene	33.67	2670	48415	52.67 ug/L
73)	1,2,3-Trichlorobenzene	34.44	2736	45590	49.52 ug/L

* Compound is ISTD

AR309427

QUANT REPORT

Operator ID: ESAT
 Output File: ^EAB12::QT
 Data File: >EAB12::D3
 Name: VOA METHOD BLANK
 Misc: 5UL IS + SS MIX IN 5ML OFW

Quant Rev: 6 Quant Time: 910226 16:21
 Injected at: 910226 15:39
 Dilution Factor: 1.00000

ID File: VOAID1::ME
 Title: DB-624 DIRECT 6PSI
 Last Calibration: 910226 13:53

	Compound	R.T.	Q ion	Area	Conc	Units	q
1)	*1,4-Difluorobenzene	16.46	114.0	69835	50.00	ug/L	69
11)	Methylene Chloride	12.32	84.0	2677	7.00	ug/L	75
23)	1,2-Dichloroethane-d4(SURR)	15.65	67.0	16048	98.40	ug/L	96
26)	Fluorobenzene(SURR)	16.23	96.0	66759	99.22	ug/L	91
33)	*Chlorobenzene-d5	22.34	117.0	55874	50.00	ug/L	94
35)	Toluene-d8(SURR)	19.22	98.0	64798	98.24	ug/L	93
52)	Bromofluorobenzene(SURR)	25.16	95.0	54467	96.54	ug/L	89
53)	*1,2-Dichlorobenzene-D4	28.97	152.0	39719	50.00	ug/L	91

* Compound is ISTD

AR309428

QUANT REPORT

Operator ID: ESAT
Output File: ^EA795::QT
Data File: >EA795::D3
Name: UQA 50 STD
Misc: 5UL IS+SS & 5UL STD MIX IN 5ML OFW

Quant Rev: 6 Quant Time: 910225 11:11
Injected at: 910225 10:17
Dilution Factor: 1.00000

ID File: UQAID1::ME
Title: DB-624 DIRECT 6PSI
Last Calibration: 910225 11:30

	Compound	R.T.	Scan#	Area	Conc	Units
1)	*1,4-Difluorobenzene	16.56	1207	83722	50.00	ug/L
2)	Dichlorodifluoromethane	7.35	417	15295M	50.00	ug/L
3)	Chloromethane	8.48	514	8719M	50.00	ug/L
4)	Vinyl Chloride	9.04	562	11327	50.00	ug/L
5)	Bromomethane	9.90	636	22025	50.00	ug/L
6)	Chloroethane	10.19	661	9087	50.00	ug/L
7)	Trichlorofluoromethane	10.72	706	46600	50.00	ug/L
8)	1,1-Dichloroethene	11.60	782	6866	50.00	ug/L
9)	Carbon Disulfide	11.80	799	51386	50.00	ug/L
10)	Acetone	11.80	799	8957	50.00	ug/L
11)	Methylene Chloride	12.43	853	19802M	50.00	ug/L
12)	Trans-1,2-Dichloroethene	12.83	887	16077	50.00	ug/L
13)	1,1-Dichloroethane	13.49	944	42219	50.00	ug/L
14)	Vinyl Acetate	13.66	958	4831	50.00	ug/L
15)	2,2-Dichloropropane	14.40	1022	28139	50.00	ug/L
16)	Cis-1,2-Dichloroethene	14.43	1024	23771	50.00	ug/L
17)	2-Butanone	14.54	1034	14743	50.00	ug/L
18)	Bromochloromethane	14.80	1056	26079	50.00	ug/L
19)	Chloroform	14.98	1071	62581	50.00	ug/L
20)	1,1,1-Trichloroethane	15.20	1090	54491	50.00	ug/L
21)	Carbon Tetrachloride	15.47	1113	60148	50.00	ug/L
22)	1,1-Dichloro-1-propene	15.49	1115	38242	50.00	ug/L
23)	1,2-Dichloroethane-d4 (SURR)	15.77	1139	17747	100.00	ug/L
24)	Benzene	15.83	1144	48907	50.00	ug/L
25)	1,2-Dichloroethane	15.90	1150	41171	50.00	ug/L
26)	Fluorobenzene (SURR)	16.33	1187	78793	100.00	ug/L
27)	Trichloroethene	16.98	1243	43752	50.00	ug/L
28)	1,2-Dichloropropane	17.39	1278	25891	50.00	ug/L
29)	Dibromomethane	17.62	1297	41064	50.00	ug/L
30)	Bromodichloromethane	17.94	1325	71530	50.00	ug/L
31)	2-Chloroethylvinylether	18.59	1380	16279	50.00	ug/L
32)	Trans-1,3-Dichloropropene	18.81	1399	56945	30.00	ug/L
33)	*Chlorobenzene-d5	22.39	1706	64973	50.00	ug/L
34)	4-Methyl-2-pentanone	19.19	1432	9968	50.00	ug/L
35)	Toluene-d8 (SURR)	19.32	1443	74434	100.00	ug/L
36)	Toluene	19.45	1454	40415R30600	50.00	ug/L
37)	Cis-1,3-Dichloropropene	19.96	1498	30608	50.00	ug/L
38)	1,1,2-Trichloroethane	20.34	1530	29417	50.00	ug/L
39)	Tetrachloroethene	20.57	1550	57853	50.00	ug/L
40)	1,3-Dichloropropene	20.68	1559	42948	50.00	ug/L
41)	2-Hexanone	20.93	1581	10445	50.00	ug/L
42)	Dibromochloromethane	21.14	1599	84614	50.00	ug/L
43)	1,2-Dibromoethane (EDB)	21.35	1617	55284	50.00	ug/L
44)	Chlorobenzene	22.48	1713	64112	50.00	ug/L
45)	1,1,1,2-Tetrachloroethane	22.48	1713	40577	50.00	ug/L

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48)	o-Xylene	23.95	1839	63042	100.00 ug/L	93
49)	Styrene	24.01	1844	117516	100.00 ug/L	100
50)	Bromoform	24.39	1877	81431	50.00 ug/L	96
51)	Isopropylbenzene	24.86	1917	102291	50.00 ug/L	87
52)	Bromofluorobenzene (SURR)	25.21	1947	63987	100.00 ug/L	84
53)	*1,2-Dichlorobenzene-D4	29.01	2272	47870	50.00 ug/L	91
54)	Bromobenzene	25.53	1974	49138	50.00 ug/L	86
55)	1,1,2,2-Tetrachloroethane	25.64	1984	52869	50.00 ug/L	97
56)	1,2,3-Trichloropropane	25.70	1989	34282	50.00 ug/L	95
57)	n-Propylbenzene	25.87	2003	26964	50.00 ug/L	92
58)	2-Chlorotoluene	26.03	2017	29399	50.00 ug/L	94
59)	4-Chlorotoluene	26.31	2041	29801	50.00 ug/L	97
60)	1,3,5-Trimethylbenzene	26.32	2042	86086	50.00 ug/L	100
61)	tert-Butylbenzene	27.11	2109	92892	50.00 ug/L	99
62)	1,2,4-Trimethylbenzene	27.25	2121	88835	50.00 ug/L	98
63)	sec-Butylbenzene	27.68	2158	122079	50.00 ug/L	98
64)	1,3-Dichlorobenzene	27.89	2176	77548	50.00 ug/L	96
65)	1,4-Dichlorobenzene	28.12	2196	81563	50.00 ug/L	93
66)	p-Isopropyltoluene	28.06	2191	101818	50.00 ug/L	100
67)	1,2-Dichlorobenzene	29.05	2275	73264	50.00 ug/L	97
68)	n-Butylbenzene	29.11	2280	91651	50.00 ug/L	100
69)	1,2-Dibromo-3-chloropropane	31.08	2449	17183	50.00 ug/L	88
70)	1,2,4-Trichlorobenzene	33.22	2632	67198	50.00 ug/L	100
71)	Naphthalene	33.84	2685	91347	50.00 ug/L	100
72)	Hexachlorobutadiene	33.72	2675	61628	50.00 ug/L	100
73)	1,2,3-Trichlorobenzene	34.49	2741	60080	50.00 ug/L	100

* Compound is ISTD

AR309430

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QUANT REPORT

Operator ID: ESAT
Output File: ^EAB10::QT'
Data File: ^EAB10::O3
Name: VOA 50 STD
Misc: 5UL IS+SS + 5UL STD MIX IN 5ML OFW

Quant Rev: 6 Quant Time: 910226 13:54
Injected at: 910226 12:28
Dilution Factor: 1.00000

ID File: VOAID1::ME
Title: OB-624 DIRECT 6PSI
Last Calibration: 910226 13:53

	Compound	R.T.	Scan#	Area	Conc	Units
1)	*1,4-Difluorobenzene	16.55	1206	76260	50.00	ug/L
2)	Dichlorodifluoromethane	7.53	433	15366M	50.00	ug/L
3)	Chloromethane	8.62	526	7441	50.00	ug/L
4)	Vinyl Chloride	9.14	571	10369	50.00	ug/L
5)	Bromomethane	9.98	643	19677M	50.00	ug/L
6)	Chloroethane	10.26	667	8113	50.00	ug/L
7)	Trichlorofluoromethane	10.76	710	45703	50.00	ug/L
8)	1,1-Dichloroethene	11.64	785	6146	50.00	ug/L
9)	Carbon Disulfide	11.83	802	47802	50.00	ug/L
10)	Acetone	11.85	803	5433	50.00	ug/L
11)	Methylene Chloride	12.45	855	20885	50.00	ug/L
12)	Trans-1,2-Dichloroethene	12.85	889	15120	50.00	ug/L
13)	1,1-Dichloroethane	13.50	945	40152	50.00	ug/L
14)	Vinyl Acetate	13.68	960	4397	50.00	ug/L
15)	2,2-Dichloropropane	14.39	1021	27148	50.00	ug/L
16)	Cis-1,2-Dichloroethene	14.42	1024	22415	50.00	ug/L
17)	2-Butanone	14.54	1034	10635	50.00	ug/L
18)	Bromochloromethane	14.80	1056	22423	50.00	ug/L
19)	Chloroform	14.97	1071	57874	50.00	ug/L
20)	1,1,1-Trichloroethane	15.19	1090	52933	50.00	ug/L
21)	Carbon Tetrachloride	15.48	1114	58450	50.00	ug/L
22)	1,1-Dichloro-1-propene	15.49	1115	36443	50.00	ug/L
23)	1,2-Dichloroethane-d4(SURR)	15.78	1140	17809	100.00	ug/L
24)	Benzene	15.84	1145	46592	50.00	ug/L
25)	1,2-Dichloroethane	15.90	1150	40252	50.00	ug/L
26)	Fluorobenzene(SURR)	16.34	1188	73473	100.00	ug/L
27)	Trichloroethene	16.98	1243	36878	50.00	ug/L
28)	1,2-Dichloropropane	17.40	1279	24939	50.00	ug/L
29)	Dibromomethane	17.61	1297	37873	50.00	ug/L
30)	Bromodichloromethane	17.95	1326	63478	50.00	ug/L
31)	2-Chloroethylvinylether	18.59	1381	15643	50.00	ug/L
32)	Trans-1,3-Dichloropropene	18.83	1401	53756	50.00	ug/L
33)	*Chlorobenzene-d5	22.44	1710	59302	50.00	ug/L
34)	4-Methyl-2-pentanone	19.20	1433	9119	50.00	ug/L
35)	Toluene-d8(SURR)	19.32	1443	70003	50.00	ug/L
36)	Toluene	19.46	1455	36560	50.00	ug/L
37)	Cis-1,3-Dichloropropene	19.97	1499	27471	70.00	ug/L
38)	1,1,2-Trichloroethane	20.36	1532	26655	50.00	ug/L
39)	Tetrachloroethene	20.58	1551	47738	50.00	ug/L
40)	1,3-Dichloropropene	20.68	1560	39894	50.00	ug/L
41)	2-Hexanone	20.97	1584	8415	50.00	ug/L
42)	Dibromochloromethane	21.18	1602	72540	50.00	ug/L
43)	1,2-Dibromoethane (EDB)	21.37	1619	49718	50.00	ug/L
44)	Chlorobenzene	22.48	1714	59071	50.00	ug/L

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47)	m&p-Xylenes	23.26	1763	61809	100.00 ug/L	83
48)	o-Xylene	23.98	1842	60551	100.00 ug/L	97
49)	Styrene	24.03	1846	108992	100.00 ug/L	100
50)	Bromoform	24.41	1879	70874	50.00 ug/L	97
51)	Isopropylbenzene	24.88	1919	98461	50.00 ug/L	90
52)	Bromofluorobenzene (SURR)	25.23	1949	59878	100.00 ug/L	91
53)	*1,2-Dichlorobenzene-D4	29.03	2274	41384	50.00 ug/L	91
54)	Bromobenzene	25.54	1976	42018	50.00 ug/L	91
55)	1,1,2,2-Tetrachloroethane	25.67	1987	49802	50.00 ug/L	94
56)	1,2,3-Trichloropropane	25.72	1991	37034	50.00 ug/L	95
57)	n-Propylbenzene	25.90	2006	24926	50.00 ug/L	97
58)	2-Chlorotoluene	26.05	2019	26838	50.00 ug/L	90
59)	4-Chlorotoluene	26.33	2043	26173	50.00 ug/L	94
60)	1,3,5-Trimethylbenzene	26.35	2045	83099	50.00 ug/L	100
61)	tert-Butylbenzene	27.15	2113	84611	50.00 ug/L	94
62)	1,2,4-Trimethylbenzene	27.26	2123	83876	50.00 ug/L	97
63)	sec-Butylbenzene	27.70	2160	115440	50.00 ug/L	94
64)	1,3-Dichlorobenzene	27.91	2178	66875	50.00 ug/L	93
65)	1,4-Dichlorobenzene	28.14	2198	71977	50.00 ug/L	99
66)	p-Isopropyltoluene	28.09	2194	92971	50.00 ug/L	100
67)	1,2-Dichlorobenzene	29.08	2278	63436	50.00 ug/L	95
68)	n-Butylbenzene	29.14	2283	89840	50.00 ug/L	100
69)	1,2-Dibromo-3-chloropropane	31.11	2452	16179	50.00 ug/L	78
70)	1,2,4-Trichlorobenzene	33.26	2636	55238	50.00 ug/L	100
71)	Naphthalene	33.88	2689	79719	50.00 ug/L	100
72)	Hexachlorobutadiene	33.76	2679	52767	50.00 ug/L	100
73)	1,2,3-Trichlorobenzene	34.51	2743	52844	50.00 ug/L	100

* Compound is ISTD

AR309432

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ENVIRONMENTAL PROTECTION AGENCY
Office of Enforcement

CHAIN OF CUSTODY RECORD

PROJECT NAME CS				STATION LOCATION		NO. OF CONTAINERS		REMARKS	
STA. NO.	DATE	TIME	INITIALS	STATION LOCATION	NO. OF CONTAINERS	NO.	OF CONTAINERS	REMARKS	
HW-33A	01/11/84	0845	X	Montgomery, WA 19101A	3	3	3	91022107	
HW-33A	01/11/84	1700	X	Montgomery, WA 19101A (Gravel)	9	9	9	91022108, 09	
HW-33A	01/11/84	1900	X	Montgomery, WA 19101A (Gravel)	3	3	3	91022109	
HW-33A	01/11/84	0730	X	Leip, Grange	3	3	3	91022110	
HW-33A	01/11/84	1500	X	Montgomery, WA 19101A	1	1	1	91022101	
HW-33A	01/11/84	1600	X	Montgomery, WA 19101A (Gravel)	1	1	1	91022103	
HW-33A	01/11/84	0730	X	Field, Grange, 600	1	1	1	91022105	
HW-33A	01/11/84	1215	X	Montgomery, WA 19101A	1	1	1	91022101	
HW-33A	01/11/84	1215	X	Montgomery, WA 19101A (Gravel)	1	1	1	91022103	
HW-33A	01/11/84	0730	X	Field, Grange, 600	1	1	1	91022105	
HW-33A	01/11/84	1215	X	Montgomery, WA 19101A	1	1	1	91022101	
HW-33A	01/11/84	1215	X	Montgomery, WA 19101A (Gravel)	1	1	1	91022103	
HW-33A	01/11/84	0730	X	Field, Grange - Sulfate	1	1	1	91022105	

Relinquished by: (Signature)	Date / Time	Received by: (Signature)	Date / Time
<i>Randy Allen</i>	01/11/84 1430		
Relinquished by: (Signature)	Date / Time	Received by: (Signature)	Date / Time
Relinquished by: (Signature)	Date / Time	Received by: (Signature)	Date / Time

Received for Laboratory by: (Signature) *Randy Allen* Date / Time 01/11/84 1430

Remarks: Unusual via Federal Express

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Office of Enforcement

PROJ. NO. PROJECT NAME

CHAIN OF CUSTODY RECORD

REGION 3
Curtis Bldg., 6th & Walnut Sts.
Philadelphia, Pennsylvania 19108

PROJ. NO. PROJECT NAME		SAMPLERS: (Signature) <i>Richard E. Schaeffer</i> <i>Rocky Area</i>		STATION LOCATION		NO. OF CONTAINERS	CHAIN OF CUSTODY			REMARKS
STA. NO.	DATE	TIME	GRAB	DATE	TIME		CHLORIDE	NITRATE/NITRITE	TOTAL NITROGEN	
14-98	12/27/15	1215	X	Monitoring well 98		1			91022101	
14-98	12/27/15	1215	X	Monitoring well 98 (duplicate)		1			91022101	
14-98	12/27/15	0730	X	Field Blank Chloride		1			91022105	
14-98	12/27/15	1215	X	Monitoring well 98		1			91022101	
14-98	12/27/15	1215	X	Monitoring well 98 (duplicate)		1			91022103	
14-98	12/27/15	0730	X	Field Blank Sulfate		1			91022105	
14-98	12/27/15	1215	X	Monitoring well 98		1			91022101	
14-98	12/27/15	1215	X	Monitoring well 98 (duplicate)		1			91022103	
14-98	12/27/15	0730	X	Field Blank Nitrate/Nitrite		1			91022105	
14-98	12/27/15	1215	X	Monitoring well 98		1			91022101	
14-98	12/27/15	1215	X	Monitoring well 98 (duplicate)		1			91022103	
14-98	12/27/15	0730	X	Field Blank Nitrogen		1			91022105	
14-98	12/27/15	1200	X	Monitoring well 98		1			91022101	
14-98	12/27/15	1200	X	Monitoring well 98 (duplicate)		1			91022103	
14-98	12/27/15	1115	X	Field Blank Total N		1			91022105	
<i>Rocky Area</i> Relinquished by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i> Received by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i>						Relinquished by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i> Received by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i>		Relinquished by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i> Received by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i>		
Relinquished by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i> Received by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i>						Relinquished by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i> Received by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i>		Relinquished by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i> Received by: (Signature) <i>[Signature]</i> Date / Time <i>2/26/16 1430</i>		

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ENVIRONMENTAL PROTECTION AGENCY
Office of Enforcement

CHAIN OF CUSTODY RECORD

PROJ. NO.		PROJECT NAME		STATION LOCATION		NO. OF CONTAINERS	REMARKS
STA. NO.	DATE	TIME	DATE	TIME			
SAMPLERS: (Signature) <i>William E. Strawn</i> <i>Redy, Steve</i>							
14-19B	3/20/84	1300	X	Monit. bery. (del 9B)	1	Received Laboratory Integrity Integrity	3-21-1051 91022102
14-19B	3/20/84	1300	X	Monit. bery. (del 9B) (Signature)	1		3-21-1051 91022104
14-19B	3/20/84	1115	X	Field blank Dissolved Pb	1		3-21-1051 91022106
14-19B	3/20/84	1300	X	Monit. bery. (del 9B)	1		3-21-1051 91022101
14-19B	3/20/84	1300	X	Monit. bery. (del 9B) (Signature)	1		3-21-1051 91022103
14-19B	3/20/84	1115	X	Field blank Integrity Pb	1		3-21-1051 91022105
14-19B	3/20/84	1300	X	Monit. bery. (del 9B)	1		3-21-1051 91022102
14-19B	3/20/84	1300	X	Monit. bery. (del 9B) (Signature)	1		3-21-1051 91022104
14-19B	3/20/84	1115	X	Field blank Dissolved Integrity Pb	1		3-21-1051 91022105
14-19B	3/20/84	1300	X	Monit. bery. (del 9B)	1		
Relinquished by: (Signature) <i>Redy, Steve</i> Date / Time 3/21/84 1430 Received by: (Signature) Relinquished by: (Signature) Date / Time Received by: (Signature) Relinquished by: (Signature) Date / Time Received by: (Signature)							

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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

REGION III
CENTRAL REGIONAL LABORATORY
838 BERTGATE ROAD
ANNAPOLIS, MARYLAND 21401
(301) 266-9100

DATE : April 15, 1991

SUBJECT : Region III Data QA Review

FROM : Theresa A. Simpson *TAS*
Region III ESAT DFO (3ES23)

TO : Stephanie Dehnhard
Regional Project Manager (3HW25)

Attached is the inorganic data review for the Chem/Solv Site (Case15915) completed by the Region III Environmental Services Assistance Team (ESAT) contractor under the direction of Region III ESD.

If you have any questions regarding this review, please call me.

Attachment

cc: Cathy Garria, CDM
Edward Kantor, EMSL-LV
Regional CLP TPO: Stevie Wilding Region: III Lab Code: ITPA

TID File: 03910219 Task 2338

AR309436

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2568A RIVA ROAD
SUITE 300
ANNAPOLIS, MD 21401
PHONE 301-266-9887

DATE: 11 APRIL 1991

SUBJECT: INORGANIC DATA VALIDATION, Case 15915
SITE: CHEM/SOLV

FROM: PETE CHAPMAN *PC* MARSHA BURRELL *mb*
INORGANIC DATA REVIEWER SENIOR DATA REVIEWER

TO: TERRY SIMPSON
ESAT DEPUTY PROJECT OFFICER

THRU: RICHARD D. DRESSER *RD*
ESAT TEAM MANAGER

OVERVIEW

The set of samples for Case 15915 contained five (5) unfiltered aqueous and five (5) filtered aqueous samples which were analyzed through the Contract Laboratory Program (CLP) Routine Analytical Services. Included in the sample set was an unfiltered aqueous equipment blank, a filtered field blank, an unfiltered field duplicate pair, and a filtered field duplicate pair. Samples MCES39 and MCES41 exceeded the 10-day Chemical Health Advisory Level for the Pb analyte. The advisory level for Pb is 20.0 ug/L and the results for these samples were 173 ug/L and 88.7 ug/L, respectively.

SUMMARY

All analytes were successfully analyzed in all samples. Areas of concern with respect to data usability are listed according to the seriousness of the problem. These include:

MINOR ISSUES

The preparation blank had results >IDL for the Ca, Fe, K, Na and Zn analytes in the aqueous samples and the unfiltered field blank had results >IDL for the Tl and Zn analytes in the unfiltered samples. The reported results for these analytes in the affected samples which are <5x the blank concentration may be biased and therefore, have been qualified "B".

The ICP serial dilution results for the K, Na and Zn analytes in the samples were outside of the 10% control limit. The reported results for these analytes in the samples have been qualified estimated, "J". However, the reported result for sample MCES45 in the Zn analyte has been superceded by the qualifier "B" as previously mentioned.

The matrix spike recovery was high (>125%) for the Hg analyte in the samples. The reported result for this analyte in sample MCES45 may be biased high, and therefore, has been qualified "K".

The analytical spike recovery was high (>115%) for the Tl analyte in sample MCES43. The reported result for this sample may be biased high, and therefore, it has been qualified "K".

NOTE:

The data was reviewed in accordance with the National Functional Guidelines for Evaluating Inorganic Analyses.

INFORMATION REGARDING REPORT CONTENT

Table 1A is a summary of qualifiers added to the laboratory's results during evaluation.

ATTACHMENTS

TABLE 1A	SUMMARY OF QUALIFIERS ON DATA SUMMARY AFTER DATA VALIDATION
TABLE 1B	CODES USED IN COMMENTS COLUMN
TABLE 2	GLOSSARY OF DATA QUALIFIER CODES
TABLE 3	DATA SUMMARY FORMS
APPENDIX A	RESULTS REPORTED BY LABORATORY FORM 1a
APPENDIX B	TPO REPORT

PC104A02.CSO

AR309438

WESTERN

TABLE 1A
SUMMARY OF QUALIFIERS ON DATA SUMMARY
AFTER DATA VALIDATION

ANALYTE	SAMPLES AFFECTED	NON-		BIAS	COMMENTS*
		POSITIVE	DETECTED		
		VALUES	VALUES		
Ca	MCES44	B		High	A (18.5 ppb)
Fe	MCES43, MCES45, MCES46	B		High	A (42.4 ppb)
Hg	MCES45	K		High	B (126%)
K	All aqueous except MCES43, MCES44	J			C (10.1%)
	MCES43, MCES44	B		High	A (79.2 ppb)
Na	All aqueous except MCES43, MCES44	J			C (11.4%)
	MCES43	B		High	A (27.4 ppb)
Tl	MCES41	B		High	D (1.0 ppb)
	MCES43	K		High	E (140%)
Zn	All aqueous except MCES43, MCES44, MCES45	J			C (43.3%)
	MCES43, MCES44	B		High	A (1.9 ppb)
	MCES45	B		High	D (4.9 ppb)

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TABLE 1B

CODES USED IN COMMENTS COLUMN

- A - The preparation blank had a result that was >IDL (the result is in parentheses) and the reported results were <5x the blank. The reported results may be biased high.
- B - Due to a high matrix spike recovery (% recovery is in parentheses), the reported result may be biased high.
- C - The ICP serial dilution result was outside of the control limit (the result is in parentheses), the reported results are estimated.
- D - The field blank had a result that was >IDL (the result is in parentheses) and the reported results were <5x the blank. The reported results may be biased high.
- E - Due to a high analytical spike recovery (% recovery is in parentheses), the reported result may be biased high.

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TABLE 2

GLOSSARY OF DATA QUALIFIER CODES (INORGANIC)

CODES RELATED TO IDENTIFICATION

(confidence concerning presence or absence of analytes):

U = Not detected. The associated number indicates approximate sample concentration necessary to be detected.

(NO CODE) = Confirmed identification.

B = Not detected substantially above the level reported in laboratory or field blanks.

R = Unreliable result. Analyte may or may not be present in the sample. Supporting data necessary to confirm result.

CODES RELATED TO QUANTITATION

(can be used for both positive results and sample quantitation limits):

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

K = Analyte present. Reported value may be biased high. Actual value is expected to be lower.

L = Analyte present. Reported value may be biased low. Actual value is expected to be higher.

[] = Analyte present. As values approach the IDL the quantitation may not be accurate.

UJ = Not detected, quantitation limit may be inaccurate or imprecise.

UL = Not detected, quantitation limit is probably higher.

OTHER CODES

Q = No analytical result.

AR309441

DATA SUMMARY FORM: INORGANICS

Site Name: Chem-Solv

WATER SAMPLES

(P5/L)

Case #: 15915 Sampling Date(s): 2/19-20/91

Due to dilution, sample quantitation limit is affected. See dilution table for specific.

Sample No. Dilution Factor Location	MCES37 I.D. MW-2LA	MCES38 I.D. MW-2LA Filtered	MCES39 I.D. MW-3LA Unfiltered	MCES40 I.D. MW-3LA Filtered	MCES41 I.D. MW-33D Unfiltered	MCES42 I.D. MW-33D Filtered	MCES43 I.D. FB Field Blank	MCES44 I.D. FB Filtered Field Blank	MCES45 I.D. MW-9B	MCES46 I.D. MW-9B Filtered
ANALYTE										
Aluminum	153.01	154.51	380 190		1421				180.01	
Antimony			150.51	151.61	149.71	149.31		13.91	138.81	131.71
Barium	11501	11611								
Beryllium										
Cadmium	32800	32000	16700	16900	16700	16700		34.01	14500	14900
Calcium										
Chromium	13.51	112.71	1541	543	3331	532				
Cobalt	251	7530	3170	173	28.7	4.9			184	1341
Copper	2110	20300	1710	690	6820	1670			76	3.2
Iron	1131	20300	1710	690	6820	1670			14200	4280
Lead	23400	22400	448	312	415	33.7			39.8	38.3
Magnesium									2.5	K
Manganese										
Mercury										
Nickel	15100	15100	5880	5900	5810	5950		11721	32100	31900
Potassium										
Selenium										
Silver										
Sodium	26800	27000	18100	17900	17700	18000		12101	21500	21100
Thallium										
Tin	1231									
Vanadium	58.5	30.2	6620	2510	1080	2470		1371	22.2	193
Zinc										
Cyanide										

CDL = Contract Required Detection Limit

Action Level Exits

SEE NARRATIVE FOR CODE DEFINITION

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APPENDIX A
RESULTS REPORTED BY LABORATORY
FORM I'S

AR309443

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INORGANIC ANALYSES DATA SHEET

000003
EPA SAMPLE NO.

Name: ITAS_PITTSBURGH

Contract: 13-03-0087

MC2327

Order: ITPA

Case No.: 15313

CPS No.: _____

SDS No.: MC2327

Sample (soil/water): WATER

Lab Sample ID: MC2327

Flow/med): LOW

Date Received: 02/21/91

Order: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration (C)	Q	M
7429-90-5	Aluminum	53.318		P
7440-39-2	Antimony	3.01U		P
7440-33-2	Arsenic	3.01U		P
7440-29-3	Barium	15013		P
7440-41-7	Beryllium	1.01U		P
7440-43-9	Cadmium	3.01U		P
7440-70-2	Calcium	32300		P
7440-47-3	Chromium	3.01U		P
7440-48-4	Cobalt	13.318		P
7440-50-8	Copper	3.318		P
7439-89-6	Iron	8140		P
7439-92-1	Lead	1.318		F
7439-95-4	Magnesium	30600		P
7439-96-5	Manganese	23400		P
7439-97-5	Mercury	0.231U	N	CV
7440-02-0	Nickel	4.01U		P
7440-09-7	Potassium	15100	E	P
7782-49-2	Selenium	2.01U		F
7440-22-4	Silver	3.01U		P
7440-23-5	Sodium	26800	E	P
7440-28-0	Thallium	1.01U	W	F
7440-62-2	Vanadium	2.318		P
7440-66-6	Zinc	53.3	E	P
	Cyanide			NR

Before: COLORLESS

Clarity Before: CLEAR

Texture: _____

After: COLORLESS

Clarity After: CLEAR

Artifacts: _____

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INORGANIC ANALYSES DATA SHEET

000004
EPA SAMPLE NO.

MC2328

Lab Name: ITAS_PITTSBURGH Contract: 12-09-0087

Lab Code: ITPA Case No.: 12315 EAS No.: SDG No.: MC2327

Matrix (Solid/Liquid): WATER

Lab Sample ID: MC2328

Level (Low/Med): LOW

Date Received: 02/21/91

Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
7439-90-5	Aluminum	54.318			P
7440-26-0	Antimony	3.01U			P
7440-28-2	Arsenic	3.01U			P
7440-29-3	Barium	15112			P
7440-41-7	Beryllium	1.01U			P
7440-43-9	Calcium	2.01U			P
7440-70-2	Calcium	330001			P
7440-47-2	Chromium	3.01U			P
7440-48-4	Cobalt	12.718			P
7440-50-8	Copper	2.01U			P
7439-99-6	Iron	75301			P
7439-92-1	Lead	1.01U			P
7439-95-4	Magnesium	302001			P
7439-96-3	Manganese	224001			P
7439-97-6	Mercury	0.301U	N		ICV
7440-02-0	Nickel	4.01U			P
7440-09-7	Potassium	134001	E		P
7792-49-2	Selenium	2.01U			P
7440-22-4	Silver	3.01U			P
7440-23-5	Sodium	270001	E		P
7440-28-0	Thallium	1.01U			P
7440-62-2	Vanadium	2.01U			P
7440-66-6	Zinc	30.21	E		P
	Cyanide				NR

Color Before: COLORLESS Clarity Before: CLEAR Texture: _____

Color After: COLORLESS Clarity After: CLEAR Artifacts: _____

Comments:

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1
INORGANIC ANALYSES DATA SHEET

EPA SAMPLE NO.

Name: ITAS_PITTSBURGH

Contract: 68-09-0097

MCE339

Code: ITRA

Case No.: 15512

SAS No.: 1

SDG No.: MCE337

In (soil/water): WATER

Lab Sample ID: MCE339

I (low/med): LOW

Date Received: 02/21/91

I (d): 0.0

Concentration Units (ug/L or mg/kg dry weight): ug/L

CAS No.	Analyte	Concentration	C	Q	M
7439-90-5	Aluminum	380			P
7440-36-0	Antimony	9.019			P
7440-38-2	Arsenic	2.01U			P
7440-39-2	Barium	20.518			P
7440-41-7	Beryllium	1.01U			P
7440-43-9	Cadmium	2.01U			P
7440-70-2	Calcium	16700			P
7440-47-3	Chromium	2.01U			P
7440-48-4	Cobalt	2.01U			P
7440-50-8	Copper	5.418			P
7439-99-6	Iron	3970			P
7439-92-1	Lead	173			P
7439-95-4	Magnesium	6910			P
7439-96-5	Manganese	46.8			P
7439-97-3	Mercury	0.201U	N		CV
7440-02-0	Nickel	4.01U			P
7440-09-7	Potassium	3880	E		P
7782-49-2	Selenium	2.01U			P
7440-22-4	Silver	3.01U			P
7440-23-5	Sodium	18100	E		P
7440-29-0	Thallium	1.01U			P
7440-62-2	Vanadium	2.01U			P
7440-66-6	Zinc	5620	E		P
	Cyanide				NR

Before: COLORLESS

Clarity Before: CLEAR

Texture: _____

After: COLORLESS

Clarity After: CLEAR

Artifacts: _____

nts:

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EPA SAMPLE NO.

INORGANIC ANALYSES DATA SHEET

MC2340

Lab Name: ITAS PITTSBURGH

Contract: 62-09-0087

Lab Code: ITPA

Case No.: 12315

SAS No.:

SOS No.: MC2337

Matrix (soil/water): WATER

Lab Sample ID: MC2340

Level (low/med): LOW

Date Received: 02/21/91

% Solids: 0.0

Concentration Units (ug/L or ug/kg dry weight): UG/L

CAS No.	Analyte	Concentration	Q	M
7439-90-5	Aluminum	9.01U		P
7440-36-0	Antimony	9.01U		P
7440-38-3	Arsenic	3.01U		P
7440-39-3	Barium	54.61B		P
7440-41-7	Beryllium	1.01U		P
7440-43-4	Cadmium	3.01U		P
7440-70-2	Calcium	159001		P
7440-47-3	Chromium	5.01U		P
7440-48-4	Cobalt	3.01U		P
7440-50-8	Copper	2.01U		P
7439-89-6	Iron	5431		P
7439-92-1	Lead	7.71		P
7439-95-4	Magnesium	69801		P
7439-96-5	Manganese	34.21		P
7439-97-6	Mercury	0.201U	N	CV
7440-02-0	Nickel	4.01U		P
7440-09-7	Potassium	59001	E	P
7782-49-2	Selenium	2.01U		P
7440-22-4	Silver	3.01U		P
7440-23-5	Sodium	179001	E	P
7440-29-0	Thallium	1.01B		P
7440-62-2	Vanadium	2.01U		P
7440-66-6	Zinc	23101	E	P
	Cyanide			NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

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INORGANIC ANALYSES DATA SHEET

EPA SAMPLE NO.

MC2341

Name: ITAS PITTSBURGH Contract: 68-02-0037

Code: ITAA Case No.: 15315 SAS No.: SDG No.: MC2337

ix (still/water): WATER

Lab Sample ID: MC2341

I (low/med): LOW

Date Received: 02/21/91

lids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	Q	M
7439-90-5	Aluminum	14318		P
7440-36-0	Antimony	8.01U		P
7440-33-2	Arsenic	3.01U		P
7440-39-2	Barium	49.718		P
7440-41-7	Beryllium	1.01U		P
7440-43-9	Cadmium	3.01U		P
7440-70-2	Calcium	16700		P
7440-47-3	Chromium	3.01U		P
7440-48-4	Cobalt	3.01U		P
7440-50-8	Copper	3.318		P
7439-99-6	Iron	2330		P
7439-92-1	Lead	88.71		P
7439-95-4	Magnesium	6820		P
7439-96-5	Manganese	41.51		P
7439-97-6	Mercury	0.201U	N	CY
7440-02-0	Nickel	4.01U		P
7440-09-7	Potassium	5840	E	P
7782-49-2	Selenium	2.01U		P
7440-22-4	Silver	3.01U		P
7440-23-5	Sodium	17700	E	P
7440-29-0	Thallium	1.018		P
7440-62-2	Vanadium	2.01U		P
7440-66-6	Zinc	6080	E	P
	Cyanide			NR

Before: COLORLESS Clarity Before: CLEAR Texture: _____

After: COLORLESS Clarity After: CLEAR Artifacts: _____

its:

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INORGANIC ANALYSES DATA SHEET

EPA 000000000000

MC2342

Lab Name: ITAS_PITTSBURGH Contract: 68-09-0087

Lab Code: ITRA Case No.: 12913 SAS No.: SDG No.: MC2327

Matrix (soil/water): WATER

Lab Sample ID: MC2342

Level (low/med): LOW

Date Received: 02/21/91

Solids: 0.0

Concentration Units (ug/L or ug/kg dry weight): UG/L

CAS No.	Analyte	Concentration	C	Q	M
17429-90-2	Aluminum	9.01U			P
17440-36-0	Antimony	9.01U			P
17440-38-2	Arsenic	2.01U			F
17440-39-3	Barium	49.21B			P
17440-41-7	Beryllium	1.01U			P
17440-43-9	Cadmium	2.01U			P
17440-70-2	Calcium	167001			P
17440-47-3	Chromium	2.01U			P
17440-48-4	Cobalt	3.01U			P
17440-50-8	Copper	2.01U			P
17439-99-6	Iron	3321			P
17439-92-1	Lead	9.91			F
17439-95-4	Magnesium	68701			P
17439-96-5	Manganese	33.71			P
17439-97-5	Mercury	0.301U	N		CV
17440-02-0	Nickel	4.01U			P
17440-09-7	Potassium	39301	E		P
17782-49-2	Selenium	2.01U			F
17440-22-4	Silver	3.01U			P
17440-23-5	Sodium	180001	E		P
17440-28-0	Thallium	1.01U			F
17440-62-2	Vanadium	2.01U			P
17440-66-6	Zinc	24701	E		P
	Cyanide				NR

Color Before: COLORLESS Clarity Before: CLEAR Texture: _____

Color After: COLORLESS Clarity After: CLEAR Artifacts: _____

Comments:

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INORGANIC ANALYSES DATA SHEET

EPA SAMPLE NO.

Name: ITAS PITTSBURGH

Contract: 68-09-Q087

MCES42

Code: ITPA

Case No.: 15915

SAS No.: 1

SDG No.: MCES37

Mk (soil/water): WATER

Lab Sample ID: MCES42

sl (low/med): LOW

Date Received: 02/21/91

slids: 10.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

ICPS No.	Analyte	Concentration	C	Q	IM
7439-90-3	Aluminum	9.0IU			P
7440-16-0	Antimony	3.0IU			P
7440-18-2	Arsenic	3.0IU	W		P
7440-39-2	Barium	1.0IU			P
7440-41-7	Beryllium	1.0IU			P
7440-43-9	Cadmium	3.0IU			P
7440-70-2	Calcium	3.0IU			P
7440-47-3	Chromium	5.0IU			P
7440-48-4	Cobalt	3.0IU			P
7440-50-8	Copper	3.0IU			P
7439-89-6	Iron	11.413			P
7439-92-1	Lead	1.0IU			F
7439-95-4	Magnesium	12.0IU			P
7439-96-5	Manganese	1.0IU			P
7439-97-6	Mercury	0.20IU	N		CV
7440-02-0	Nickel	4.0IU			P
7440-09-7	Potassium	16818	E		P
7732-49-2	Selenium	2.0IU			F
7440-22-4	Silver	3.0IU			P
7440-23-5	Sodium	10318	E		P
7440-28-0	Thallium	1.018	W		F
7440-62-2	Vanadium	2.0IU			P
7440-66-6	Zinc	4.918	E		P
	Cyanide				NR

r Before: COLORLESS

Clarity Before: CLEAR

Texture:

r After: COLORLESS

Clarity After: CLEAR

Artifacts:

ents:

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INORGANIC ANALYSES DATA SHEET

EPA SAMPLE NO.

MCE344

Lab Name: ITAS_PITTSBURGH Contract: 33-29-0087

Lab Code: ITA Case No.: 13915 SAS No.: SDG No.: MCE337

Matrix (solid/liquid): WATER

Lab Sample ID: MCE344

Level (low/med): LOW

Date Received: 02/21/91

Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

ICAS No.	Analyte	Concentration	C	Q	M
7440-90-2	Aluminum	9.01U			P
7440-35-0	Antimony	3.01U			P
7440-33-2	Arsenic	3.01U		W	F
7440-39-3	Barium	3.91B			P
7440-41-7	Beryllium	1.01U			P
7440-43-9	Cadmium	2.01U			P
7440-70-2	Calcium	34.61B			P
7440-47-3	Chromium	3.01U			P
7440-48-4	Cobalt	3.01U			P
7440-50-9	Copper	2.01U			P
7439-89-7	Iron	4.01U			P
7439-92-1	Lead	1.01U			F
7439-95-4	Magnesium	12.01U			P
7439-96-3	Manganese	1.01U			P
7439-97-6	Mercury	0.301U		N	CV
7440-02-0	Nickel	4.01U			P
7440-09-7	Potassium	1721B		E	P
7782-49-2	Selenium	2.01U			F
7440-22-4	Silver	3.01U			P
7440-23-5	Sodium	2161B		E	P
7440-28-0	Thallium	1.01U			P
7440-62-2	Vanadium	2.01U			P
7440-66-6	Zinc	3.71B		E	P
	Cyanide				INR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture: _____

Color After: COLORLESS

Clarity After: CLEAR

Artifacts: _____

Comments:

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(INORGANIC ANALYSES DATA SHEET)

EPA SAMPLE NO.

Name: STAS_PITTSBURGH

Contract: 33-C9-2007

MCE345

Code: ST94

Case No.: 15915

CAS No.:

SDS No.: MCE337

IX (soil/water): WATER

Lab Sample ID: MCE345

I (low/mid): LOW

Date Received: 02/21/91

Iids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	Q	M
7439-90-2	Aluminum	20.018		P
7440-26-0	Antimony	3.01U		P
7440-33-2	Arsenic	3.01U		P
7440-39-3	Barium	29.318		P
7440-41-7	Beryllium	1.01U		P
7440-43-9	Cadmium	3.01U		P
7440-70-2	Calcium	439013		P
7440-47-3	Chromium	3.01U		P
7440-48-4	Cobalt	3.01U		P
7440-50-8	Copper	3.01U		P
7439-89-6	Iron	1841		P
7439-92-1	Lead	7.61		P
7439-95-4	Magnesium	422013		P
7439-96-5	Manganese	39.31		P
7439-97-6	Mercury	3.31	N	CV
7440-02-0	Nickel	4.01U		P
7440-09-7	Potassium	321001	E	P
7782-49-2	Selenium	3.01U		P
7440-22-4	Silver	3.01U		P
7440-23-5	Sodium	315001	E	P
7440-28-0	Thallium	1.01U		P
7440-62-2	Vanadium	3.01U		P
7440-66-6	Zinc	22.21	E	P
	Cyanide			NR

Before: COLORLESS

Clarity Before: CLEAR

Texture: _____

After: COLORLESS

Clarity After: CLEAR

Artifacts: _____

its:

AR309452

INORGANIC ANALYSES DATA SHEET

EPA SAMPLE NO.

Lab Name: ITAS_PITTSBURGH Contract: 61-09-0087

MCE346

Lab Code: ITRA Case No.: 15913 SAS No.: SDG No.: MCE337

Matrix (soil/water): WATER

Lab Sample ID: MCE346

Level (low/med): LOW

Date Received: 02/21/91

Solids: 0.0

Concentration Units (ug/L or mg/kg dry weight): UG/L

CAS No.	Analyte	Concentration	Q	M
7429-90-5	Aluminum	9.01U		P
7440-36-0	Antimony	9.01U		P
7440-33-2	Arsenic	2.01U		P
7440-39-3	Barium	31.718		P
7440-41-7	Beryllium	1.01U		P
7440-43-9	Cadmium	2.01U		P
7440-70-2	Calcium	490018		P
7440-47-3	Chromium	5.01U		P
7440-48-4	Cobalt	3.01U		P
7440-50-8	Copper	2.01U		P
7439-89-6	Iron	8.418		P
7439-92-1	Lead	3.21	S	P
7439-92-4	Magnesium	429018		P
7439-96-5	Manganese	38.31		P
7439-97-6	Mercury	0.201U	N	CV
7440-02-0	Nickel	4.01U		P
7440-09-7	Potassium	319001	E	P
7782-49-2	Selenium	2.01U		P
7440-22-4	Silver	3.01U		P
7440-23-5	Sodium	214001	E	P
7440-29-0	Thallium	1.01U	W	P
7440-62-2	Vanadium	2.01U		P
7440-66-6	Zinc	19.318	E	P
	Cyanide			NR

Color Before: COLORLESS

Clarity Before: CLEAR

Texture:

Color After: COLORLESS

Clarity After: CLEAR

Artifacts:

Comments:

AR309453

WESTERN

APPENDIX B

TPO REPORT

AR309454

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WESTERN

Page 1 of 1

TPO: [] ACTION [X] FYI

Region III

INORGANIC REGIONAL DATA ASSESSMENT SUMMARY

CASE NO: 15915
SDG NO: MCES37
SOW: 7/88
NO. OF SAMPLES: five (5)

LABORATORY: ITPA
DATA USER: Stavia Wilding
REVIEW COMPLETION DATE: 4/5/91
MATRIX: Unfiltered aqueous

REVIEWER: ESAT

	ICP	AA	Hg	CYANIDE
1. HOLDING TIMES	_O_	_O_	_O_	--
2. INITIAL CALIBRATIONS	_O_	_O_	_O_	--
3. CONTINUING CALIBRATIONS	_O_	_O_	_O_	--
4. FIELD BLANKS (F=NOT APPLICABLE)	_X_	_X_	_O_	--
5. LABORATORY BLANKS	_X_	_O_	_O_	--
6. ICS	_O_			
7. LCS	_O_	_O_		
8. DUPLICATE ANALYSIS	_O_	_O_	_O_	--
9. MATRIX SPIKE	_O_	_O_	_X_	--
10. MSA		_O_		
11. SERIAL DILUTION	_M_			
12. SAMPLE VERIFICATION	_O_	_O_	_O_	--
13. REGIONAL QC (F=NOT APPLICABLE)	_F_	_F_	_F_	--
14. OVERALL ASSESSMENT	_M_	_X_	_X_	--

0 = No problem or minor problems that do not affect data usability
X = No more than about 5% of the data points are qualified as either estimated or unusable.
M = More than about 5% of the data points are qualified as estimated.
F = More than about 5% of the data points are qualified as unusable.
TPO action requested; use in conjunction with one of the above codes.

TPO ACTION ITEMS:

AREAS OF CONCERN:

AR309-55

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WESTERN

Page 2 of 2

TPO: [] ACTION [X] FYI

Region III

INORGANIC REGIONAL DATA ASSESSMENT SUMMARY

CASE NO: 15915
 DG NO: MCES37
 OW: 7/88
 O. OF SAMPLES: five (5)

LABORATORY: ITPA
 DATA USER: Stavia Wilding
 REVIEW COMPLETION DATE: 4/5/91
 MATRIX: Filtered aqueous

VIEWER: ESAT

	ICP	AA	Hg	CYANIDE
. HOLDING TIMES	_O_	_O_	_O_	--
. INITIAL CALIBRATIONS	_O_	_O_	_O_	--
. CONTINUING CALIBRATIONS	_O_	_O_	_O_	--
. FIELD BLANKS (F=NOT APPLICABLE)	_O_	_O_	_O_	--
. LABORATORY BLANKS	_X_	_O_	_O_	--
. ICS	_O_			
. LCS	_O_	_O_		
. DUPLICATE ANALYSIS	_O_	_O_	_O_	--
. MATRIX SPIKE	_O_	_O_	_O_	--
. MSA		_O_		
. SERIAL DILUTION	_M_			
. SAMPLE VERIFICATION	_O_	_O_	_O_	--
. REGIONAL QC (F=NOT APPLICABLE)	_F_	_F_	_F_	--
. OVERALL ASSESSMENT	_M_	_O_	_O_	--

O = No problems or minor problems that do not affect data usability
 X = No more than about 5% of the data points are qualified as either estimated or unusable.
 M = More than about 5% of the data points are qualified as estimated.
 Z = More than about 5% of the data points are qualified as unusable.
 A = TPO action requested; use in conjunction with one of the above codes.

O ACTION ITEMS:

EAS OF CONCERN:

AR309456

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BCM**INTEROFFICE**

TO: Peg Bonaker

DATE: May 13, 1991

cc: M.M. Mang

FROM: Atwood Davis/Denise McGuire

SUBJECT: Chem-Solv
Split Sample Comparison
BCM Project No. 00-6012-02INTRODUCTION

This memo compares the split sample results obtained by EPA Region III's ESAT subcontractor to those obtained by BCM Laboratory. In addition, a summary review of the EPA subcontractor data was performed and is offered in this memo. Three monitoring well samples were collected on February 19, 1991 at the following locations, MW26A, MW33A, and MW33A (field duplicate). These samples were split between BCM and the EPA subcontractor onsite. In addition, a trip blank was collected. However, from the information available, it was not possible to ascertain if the trip blanks were from the same source or if BCM and the EPA contractor each supplied their own trip blank.

BCM's data package was validated in accordance with EPA Laboratory Data Validation Functional Guidelines for Evaluating Organics Analyses, February, 1988. The results of the validation have been prepared and are available as a validation report 1 April, 1991 for the Chem-Solv site under BCM Project Number 00-6012-02.

REVIEW OF EPA SUBCONTRACTOR SAMPLE DATA PACKAGES

The inorganic split sample data package was reviewed under a separate memo of May 7, 1991, by Denise McGuire of BCM's QA Department (Attached).

A review the volatile organic data packages supplied by ESAT indicated the following:

1. Samples were analyzed within acceptable holding times.
2. Surrogate spike recoveries were reported to be within acceptable ranges.
3. Matrix spike and matrix spike duplicate recoveries and relative percent differences (RPDs) were reported to be within acceptable ranges.
4. GC/MS Tune criteria was found to be acceptable for initial calibration and analyses.
5. Initial calibration and continuing calibration response factors (RF) were all above 0.05. All system performance check compounds (SPCC) met average RF criteria, except chloromethane, in both initial and continuing calibrations.

AR3094

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6. All continuing calibration check (CCC) compounds met percent relative standard deviation (RSD) criteria except 1,1-dichloroethene (30.167) which was a minor exceedance in the initial calibration. All percent differences for CCCs in the continuing calibrations met acceptance criteria.
7. Under CLP protocol another initial calibration would be required due to the criteria exceeded in points 5 and 6. However, these criteria were not exceeded to the extent of adversely impacting the reported results.
8. Several percent differences exceeded 30 percent in continuing calibrations. Methylene chloride on 2/25 (probably due to lab contaminants) and chloromethane on 2/26 (a minor exceedance affecting only matrix spike and matrix spike duplicates which had no effect on reported sample data).
9. Several percent RSDs exceeded 30 percent in the initial calibration. All were minor. 2-butanone results and detection limits could be considered estimated. Detection limits for trichlorofluoromethane and 1,1-dichloroethene are also estimated.
10. Some low level trip and lab blank contaminants were reported in the split samples analyzed by the EPA subcontractor.
11. Internal standard areas and retention times were reported to be within criteria.

Several points were also noted in the review of volatiles which have some impact on the comparison of these results. BCM analyzed the samples using CLP-SOM protocol 2/88 revised 4/90, the EPA subcontractor analyzed using CLP-SOM 2/88 and EPA Method 624. This was not fully explained by the EPA subcontractor in the laboratory case narrative; however, it was noted that the GC/MS instrument was calibrated for many more compounds than found on the TCL volatiles and the Method 624 analyte list. This analyte difference caused the EPA splits to contain some reported compounds which were not analyzed by BCM's protocol. These compounds are tabulated on page 2 of the data summary attached to this memo. It should also be noted that the data package provided by the EPA subcontractor for this review was a summary package, not a full CLP deliverable. Therefore, this review is limited to the summary information only and was not verified against raw data (except internal standard areas from the quantitation lists supplied).

The overall assessment of the data supplied by EPA's subcontractor is that, with the exception of minor calibration problems and low level blank contaminants which question some sample results, the reported results appear to be consistent with BCM results for the major contaminants which were analyzed by both parties.

COMPARISON OF SPLIT SAMPLE RESULTS

Volatile Organics

A comparison of EPA subcontractor to BCM subcontractor (BCM) results is shown on page 1 of the attached data summary. The positive results for benzene in MW26 agree well as do the positive results for trichloroethene and 1,1,1-trichloroethane in MW33A and MW33 duplicates. Both laboratories also detected tetrachloroethene in MW33 duplicate below the required quantitation limit.

Blank contaminants which were reported but were not detected in the actual sample are not of consequence to the comparison. However, 2-butanone and chloroform were detected in the EPA split blank at levels similar to or greater than those detected in the samples. These sample results were flagged by the EPA subcontractor laboratory with a "B" indicating the possibility of false positives.

All other positive results in the well samples reported by the EPA subcontractor were not detected in the BCM analyses. In most cases these detects were low level below CLP quantitation limits. As previously noted, due to difference in analyte calibrated, the results shown on page 2 of the attached data summary were compounds not tested for by BCM. As a result, no comparison can be made for these compounds.

Inorganics - Mercury

A comparison of BCM subcontractor (BCM), EPA Central Regional Laboratory (CRL) and EPA Contract Laboratory Program Routine Analytical Services (CLP RAS) subcontractor results is shown on page 3 of the attached data summary. Total and inorganic mercury was analyzed for the unfiltered and filtered sample MW9B.

The unfiltered total mercury results vary (EPA CLP RAS results are highest). The EPA CLP RAS reported a high matrix spike recovery for mercury indicating this result may be biased high. BCM and the EPA CRL filtered total mercury results agree well although EPA CLP RAS reported filtered total mercury below the detection limit. Without backup raw data, this value cannot be verified.

The unfiltered inorganic mercury results agree very well, although the results are higher than the total mercury results. The filtered inorganic mercury results greatly vary. The filtered inorganic mercury duplicates analyzed by EPA CRL exhibited good reproducibility, although the inorganic mercury results are greater than the total mercury results. The higher values for inorganic mercury are probably attributed to the variability in the two methods used, specifically the different digestion procedures. Since the complete digestion procedure is not performed for the determination of inorganic mercury, there may be possible organic interferences causing the inorganic results to be evaluated.

AR309459

Met Chemistry

A comparison of BCM and EPA CRL results is shown on page 5 of the attached data summary. The positive results for chloride, nitrate, and sulfate in sample MH98 and its duplicate agree well. Positive COD results reported by EPA were not detected by BCM. These detects were below BCM subcontractor's detection limits. As a result, no comparison can be made.

SUMMARY

Volatile organic split sample results appear to agree very well except at levels below required quantitation limits. Several compounds detected in MH 26A by the EPA subcontractor were not analyzed for by BCM.

Sample MH98 unfiltered total and inorganic mercury results appear to agree. Although the EPA CLP RAS subcontractor results for total mercury is slightly higher, a high matrix spike recovery indicates the result is biased high (actual value is lower) and, therefore, closer in value to the other results. The BCM subcontractor and EPA CRL total mercury results appear to agree, although the EPA CLP RAS subcontractor reported results below the detection limit. The filtered inorganic mercury results vary considerably. Due to the method used, organic interferences may be present causing variability between the sample results.

Met chemistry split sample results appear to agree very well. Chloride was detected by the EPA CRL at levels below the BCM subcontractor's detection limits.

For specifics concerning this comparison, please see the attached memorandum. Data validation reports and analytical data are available from file 00-6012-02.

AR309460

DATA SUMMARY

UNIS ~~4412~~

CROL	LAB IDL	FIELD I.D.	QBU	QBL	QA	QB	QC	QD
	LABELD.		EPA 9102-2	EPA 9102-2	EPA 9102-2	BCL BCL	BCL BCL	EPA 9102-2
NR	Aluminum							
NR	Antimony							
NR	Arsenic							
NR	Barium							
NR	Beryllium							
NR	Cadmium							
NR	Calcium							
NR	Chromium							
NR	Cobalt							
NR	Copper							
NR	Iron							
NR	Lead							
NR	Magnesium							
NR	Manganese							
0.2	Mercury	2.3	1.7	1.6	2.5 K	2.6	2.6	2.5
NR	Nickel							
NR	Potassium							
NR	Selenium							
NR	Silver							
NR	Sodium							
NR	Titanium							
NR	Zinc							
NR	Cyanide							

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Case No. 8128-006

Lab. No. _____ Date of Sample 2/20/91
 Laboratory BCU, EPA, EPACI PRAS

Matrix Agave
Units 1318

CODL	LAB ID.	FIELD I.D.	QBF ROM B09W	QBF EPA 816B-13	QBF EPA 816B-13	QBF EPA 816B-13	QBT EPA 816B-13	QSTF dup EPA 816B-13
	LAB ID.	FIELD L.D.	QBF	QBF	QBF	QBF	QBT	QSTF dup
NR	Aluminum							
NR	Arsimony							
NR	Arsenic							
NR	Barium							
NR	Beryllum							
NR	Cadmium							
NR	Calcium							
NR	Chromium							
NR	Cobalt							
NR	Copper							
NR	Iron							
NR	Lead							
NR	Magnesium							
NR	Manganese							
OZ	Mercury	2.1	1.9	1.8	0.27	2.3	2.2	
NR	Nickel							
NR	Potassium							
NR	Selenium							
NR	Silver							
NR	Sodium							
NR	Thallium							
NR	Vanadium							
NR	Zinc							
NK	Cyanide							
E	% Solids							

$\text{EPA} = \text{TEH} \cdot \text{EPA} = \text{EPA} \cdot \text{CLPRAS} = \text{ITAS}$

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Chem-Sol Inc.
819-007

Chem--Solv Inc
2120131

Page 5

SAMPLE DATA SUMMARY

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MR 309465

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BCM**INTEROFFICE**

TO: M. Bonaker

DATE: May 7, 1991

cc: M. Mang
A. Davis

FROM: Denise McGuire DM

SUBJECT: Chem-Solv., Inc.

This data review addresses the wet chemistry data and inorganic data from the split groundwater sampling event at the Chem-Solv., Inc. site. Two groundwater samples and one field blank were analyzed for several wet chemistry analyses by the EPA Central Regional Laboratory and Weston. Three groundwater samples and a field blank were analyzed for manganese, mercury, and zinc by the Delaware Department of Natural Resources and Environmental Control. Three unfiltered groundwater samples, three filtered groundwater samples, an unfiltered field blank, a filtered field blank, an unfiltered field duplicate sample and a filtered field duplicate sample were analyzed for TAL metals by Weston.

Wet Chemistry Analyses:

Field blanks exhibited contamination for some analyses (B.O.D and C.O.D) laboratory and field blanks showed good reproducibility. Matrix spike recoveries were within control limits.

It should be noted that the laboratory reported a C.O.D. result by averaging the sample and its duplicate. This is non-compliant with SOW 788, the original sample result should have been reported. The laboratory also calculated the C.O.D. matrix spike recovery using the average value. Although this is non-compliant with SOW 788, there is no significant impact on the data; the actual matrix spike recovery is within control limits.

Inorganic Analyses: Manganese, mercury, and zinc.

holding times were met for all samples. Laboratory blanks showed contamination for zinc analysis. All calibration verification and laboratory control sample results were within control limits. Laboratory duplicates exhibited good reproducibility. Matrix spike recoveries were within control limits. Several serial dilution results were outside control limits indicating a possible matrix interference.

It should be noted that the laboratory used the field blank for the spiked sample analysis. This is non-compliant with SOW 788.

The quality control parameters could not be verified due to the lack of a complete package.

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Memorandum
May 7, 1991
Page 2

Inorganic Analyses: TAL Metals

Holding times were met for all samples. Laboratory and field blanks showed some contamination. Several serial dilution results and matrix spike recoveries were outside of control limits indicating a possible matrix interference.

Not all quality control parameters could be reviewed due to the lack of complete packages. Also laboratory duplicates and matrix spike results can not be verified.

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APPENDIX O

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BCM

APPENDIX O

DNR/C GROUNDWATER ANALYTICAL RESULTS
MARCH 1991

AR309469

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